Physical Science Techniques for Non-Invasive, Quantitative Measurements within the Living Human Body

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## Copies of Publications
The research in this thesis focuses on the use of a range of physical science techniques to perform quantitative measurements within the living human body in a non-invasive manner, for the purpose of understanding disease processes, diagnosing abnormality or monitoring therapeutic responses. The research publications cover work which has been undertaken over the last thirty years in the Universities of Edinburgh, London and Leeds. The research has not concentrated on one aspect of the physical sciences but has taken advantage of scientific and technological developments, encompassing neutron, X and γ ionising radiation, laser and THz non-ionising radiation, magnetism, radio-waves and a range of mathematical and computing techniques. Clinical applications have encompassed a wide range of diseases and conditions and include osteoporosis and other metabolic bone diseases, cardiac disease, vascular disease, rheumatoid arthritis and cancer.
Declaration and Acknowledgements

The nature of the research that I undertake often requires multidisciplinary collaboration between scientists and clinicians. The publications I have submitted are either purely my own work, the result of collaborative research where I have made a major contribution as joint researcher, or output from a research group. In the case of the latter I have either led or co-led the research group and have been responsible for both the strategic direction of the research and the day to day oversight of the research.

As a consequence, I have had the pleasure and privilege of working with many talented and inspirational colleagues whose support and contribution to my work I would like to acknowledge:

In Edinburgh:
Dr Peter Tothill, who started me on my research career, Prof Richard Eastell, Prof David Reid and Prof George Nuki in the area of Bone Measurement.
Dr John Ridgway, Prof Tony Bell, Dr Martin Been, Dr David Kean, Dr Lindsay Turnbull and Prof Jonathan Best in the area of Magnetic Resonance Imaging.

In London:
Dr Stephen Keevil, Dr Soundrie Padayachee, Dr Ted Baker, Prof Ray Gosling, Prof Michael Maisey and Prof Mike Richards in the area of Magnetic Resonance Imaging and Spectroscopy.

In Leeds:
Dr John Sutcliffe in the area of Body Composition Measurement
Dr John Truscott in the area of Bone Measurement
Dr Elizabeth Berry in the area of Image Analysis, Evidence Based Imaging and Terahertz Imaging and Sensing
Dr John Ridgway, Dr Steve Tanner and Dr Jim Meaney in the area of Magnetic Resonance Imaging
Prof Martyn Chamberlain (who also co-led the research group) in the area of Imaging and Sensing

I would also like to acknowledge my PhD students during this period: John Ridgway, Alison McKie, Maria Tarnowski, David Porter, Martin Graves, Stephen Keevil, Sylvester Yankuba, John Trustcott, Nasser Dennaoui, Sasha Radjenovic, Niall MacDougall, David Brettle, David Higgins and Gillian Walker. I would also like to acknowledge research fellows who have worked with me: Steve Kelly, Dr Tony Fitzgerald, Dr Graham Woodrow, Brian Oldroyd and Dr Chris Twelves.
Background

The publications which I have submitted for consideration for the award of DSc cover work which has been undertaken over the last thirty years during which I have worked in the Universities of Edinburgh, London and Leeds in association with NHS Teaching Hospitals. My research has focused on the use of physical sciences to perform quantitative measurements within the living human body in a non-invasive manner, for the purpose of understanding disease processes, diagnosing abnormality or monitoring therapeutic responses.

This underlying scientific theme has remained constant throughout my research career but it has been mediated by two other factors which I believe to be important in the area of applied medical science research. Both are associated with the transition of the development of an original idea to its eventual widespread availability in the health sector. I had always envisaged that any new medical technique should go through a natural, and logical, progression from basic scientific research to its eventual provision and availability in the health sector. Given that part of my personal motivation was to develop new techniques which would have an impact on healthcare, it was natural that I should take an increasing interest in the subsequent stages of this process after my initial contribution. The two factors to which I refer were therefore the clinical validation of new techniques and their commercial development.

At the start of my research career I had thought that my expertise and responsibility would, and indeed should, be concentrated in the early stages of this process, ie in the development of new techniques, and that appropriate processes and funding mechanisms were in place to investigate clinical effectiveness and efficacy. I subsequently recognised that this was naïve, a fact which I attribute to youthful inexperience, and I became aware that if I did not extend my research involvement further then appropriate validation would not take place. In addition this recognition of the importance of the need to evaluate new techniques stemmed from personal experience, as I recognized that the acceptance of new scientific techniques by health sectors was not necessarily evidence based. I became aware that the route from the assessment of the feasibility of the use of a new scientific technique in clinical practice, to finally a rigorous analysis of the efficacy and effectiveness of an investigative technique in routine clinical use, generally did not occur. This compromised the rapid implementation of new techniques into health provision.

In a number of cases the eventual outcome of my research has been associated with a medical device which has had commercial potential. From the start of my career I considered this to be part of the ‘research pipeline’, a concept which I have always had, long before ‘knowledge transfer’ became a significant part of University life. This was driven by the recognition that if any of the developments with which I was associated were to become widely available in the medical and health sector, they would need to be commercially produced and supported. This extension of my research into the commercial sphere exposed me to an additional critical environment which encompassed economic and financial issues as well as perceived user acceptance.
In my chosen field there has always been a close and synergistic link between the science and the technology. Often, particularly early in my career, there was a need to identify innovative scientific solutions to overcome problems which, we can now see with hindsight, could have been easily solved with later technology or as the technology matured. Obvious examples include computer power, firmware and software packages, non-magnetic materials, image storage devices, data communication protocols and micro-fabrication techniques. As I consider and review my early research I am conscious that I am doing so from the perspective of current technology and that initially I had a tendency to be somewhat dismissive of some of my early innovative work which had to overcome technological deficiencies. I have made every effort not to fall into this trap but include such work and the associated publications.

Because of the ongoing nature of developments in science and technology my research has not concentrated on one aspect of the physical sciences but has encompassed many, utilising neutron, X and γ ionising radiation, laser and THz non-ionising radiation, magnetism, radio-waves and a range of mathematical techniques. Likewise the clinical applications have also varied and include osteoporosis and other metabolic bone diseases, cardiac disease, vascular disease, rheumatoid arthritis and cancer. This variety is neither dilettantism nor simple opportunism on my part. Throughout my career I have recognised when it is time for a physical scientist such as myself to reduce my involvement in a subject area and pass the baton on to clinicians or the commercial sector. In such cases I have often retained a significant involvement but have transferred the main focus of my research attention to another area. Also my desire to pursue research at a high level has resulted in my willingness to transfer my attention to more promising scientific methods, when the opportunity arises, and to choose the clinical applications which are most important or which provide a challenging exemplar for the new techniques.

In addition to some of the scientific reasons for the evolution of my research, I would also like to highlight the two other factors which have had an impact. The first was the impact of career changes which have imposed both constraints and opportunities. These are generally associated with the equipment and facilities available and, probably more importantly, scientific and clinical colleagues with whom it is essential to undertake collaborative research. This collaborative approach to research is essential for high quality research into the non-invasive use of physical science in medicine. Throughout my career I have formed research teams and sometimes the presence or absence of a key team member has influenced the direction of the research. In my acknowledgements I highlight my key collaborators. The second factor was the external influence of funding priorities and opportunities. My research area requires significant equipment and recurrent expenditure on facilities and, as a consequence, is highly dependent on external funding. Generally the evolution of my research into a new area anticipated a change in funding priorities of which I was then able to take advantage (magnetic resonance imaging in the 1980s, evidence based medicine in the 1990s). Other research areas were affected by a relative lack of funding opportunities (neutron activation analysis after the 1970s, physical science aspects of bone measurement...
in the 1980s, and magnetic resonance imaging in the 1990s). Far from criticising these changes in funding priorities, I would support the necessary change of emphasis into areas which offer, or potentially offer, better outcomes. In addition the funding priorities also reflect the correct emphasis away from 'near market' research.

In this review of my research it will be apparent that though the non-invasive use of physical science in medicine has remained constant, the primary focus of my research has changed with time. In addition, as with other things in life, though I became involved in new areas, association with pre-existing areas continued; as a consequence there is considerable overlap in time between different research areas. For ease of understanding I have tried to indicate the time-line of my research in the table below.

*Number of papers in each research area during approximate five year periods*

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The table above includes the total number of my refereed publications in the different research areas. In my DSc submission I have not included all my research publications but have been selective and included those which contain the major research outputs. A small number of publications are referred to in the text in {these} brackets. These include patents, which relate to the research described but would not be appropriate as part of my DSc submission, and large systematic reviews, which are in the form of substantial monographs. For reasons of space hard copies have not been included, but electronic copies have been included on a accompanying CD.

The publications I am submitting for my DSc are listed chronologically under broad category headings, rather than in a continuous chronological list. I have only included a small number of published conference papers; these have all been refereed and the specific conferences are a recognised route for publication, often in a rapidly evolving field of research. A small number of publications arose following my PhD studies at the University of Edinburgh; these are indicated in the publication list with the symbol (*PhD) and I have included them for completeness.
Research Areas

Bone Measurement

At the start of my research career in Edinburgh in the early 1970s, one of the small number of physics techniques which had the potential to enable quantitative measurements to be made in-vivo was neutron activation analysis (NAA), and a number of research groups were investigating its development and application. My initial research focused on the development of techniques which could be used in practice on patients, in a clinical environment. Many other researchers were using neutron sources such as research reactors and cyclotrons, which may have been appropriate for the development of a research tool, but were clearly impractical for clinical use. I developed precise techniques for measuring bone calcium and other elements in the forearm and spine and applied them to the investigation of osteoporosis and bone disease associated with renal dialysis [1, 2, 3]. My scientific research had to overcome problems of relatively low neutron yield from the small $^{252}\text{Cf}$ sources, minimise the effect of the variation in sensitivity with depth in tissue, design and construct detection systems for the relatively high $\gamma$ radiation emitted and develop a computer technique for spectral analysis using matrix inversion. I also undertook the preliminary clinical evaluation, designing and managing the small trial in renal dialysis patients.

Whilst developing this technique I became aware of the limitations of studying a part of the body where there was insufficient accurate knowledge of the location of the area of measurement. Hitherto the research I had undertaken relied on accurate repositioning and sequential measurements or assumptions of internal localisation in the human body. Also the variation between people of different sizes was clearly a limitation in a fixed geometry and was particularly important in performing absolute measurements and determining the percentage difference from normal. This led me to become involved with two different approaches to resolve this underlying problem, both of which I pursued by attracting external funding to support additional research support staff.

The first was to extend NAA measurements to the whole human body [4, 11, 22] using a clinical cyclotron which had just been installed in the hospital. The spectral analysis programme I had developed was extended to measure a wider range of body elements including Ca, Na, Cl, P and N. Absolute measurements on patients were obtained by normalising the measurement using an equation based on the patient's stature; this enabled the results to be expressed as a percentage of normal. The seasonal variation in total body calcium was also measured [15]. The whole body technique was applied to a range of clinical conditions, particularly osteoporosis [12] and osteoarthritis [13, 20], rheumatoid arthritis [7, 17, 18], ankylosing spondylitis [16] and other osteopenic disorders [14, 19].

The second, and more far reaching, approach was to recognize the opportunities that medical imaging offered to localise accurately the site in the body where quantitative measurements were being performed. It was this which led me into the area of medical imaging, I was less interested in developing new imaging
techniques *per se* but rather I recognised the powerful combination of performing quantitative measurements and identifying where the information was coming from.

Whilst performing NAA measurements to measure bone calcium I had previously implemented a simple method to measure bone mineral of the wrist by measuring the absorption of γ radiation from a radionuclide source as it traversed the wrist. I had used it to measure demineralization in patients with Turner's syndrome [6] and had compared the changes measured using the two techniques [36]. It was natural to extend this work into two dimensions to obtain quantitative measurement of bone mineral in the spine. Two radionuclide energies were used, to correct for the body thickness; the technique was known as dual photon absorptiometry (DPA) and it was able to obtain both quantitative measurements and relatively low resolution images to obtain accurate and reproducible localization [8, 10]. This method was also developed further to measure bone mineral in the whole hand [21].

Whilst developing DPA I became aware of a potentially significant problem associated with the interference of one energy with the other which I termed the 'crossover correction' [5]. A consequence of an error in the crossover correction could, I hypothesised, be responsible for the variation in the decrease in bone mineral with age which was apparent at different international locations [9].

The focus of my research on bone and the increasing national interest on osteoporosis led me to consider methods of monitoring or measuring fracture healing using radionuclides. As a consequence I developed and evaluated a nuclear medicine technique to predict fracture healing in a relatively large clinical study [23].

After moving from Edinburgh I maintained my research interest in bone measurement, both in London where I offered some scientific advice and support to clinicians, but more so later in Leeds where I provided leadership to the Bone and Body Composition Research Group. The technology had advanced and commercially available dual energy X-ray based systems (DPX) were available to measure bone mineral. My research therefore followed the natural path of investigating the variation of bone mineral with age in large populations, measured using DPX [24, 28] and applying the technique to the study of a variety of bone diseases [27, 30, 32, 33].

In Leeds I also become interested in newer techniques to measure bone mineral. These included the use of ultrasound for measuring bone parameters and its relationship with measurements using photon absorptiometry [25]. To support this work it was necessary to resolve the difficult problem of simulating bone for ultrasound and so a phantom material for ultrasound was developed [29] and patented [34]. Availability of new technology enabled me to extend the technique of single photon absorptiometry, using a radiation sensitive CCD camera, to the challenging application of the measurement of neonatal bone mineral [26, 31].
Statistics

Whilst in the early phase of my research career my mathematical background and computing experience, coupled with my involvement in the clinical evaluation of the physical science techniques as well as their scientific development, caused me to become increasing involved in the statistical and data analysis aspect of research. Initially this led me to consider statistical issues associated with the comparison of techniques and the level of correlation which was possible due to inherent precision values [36].

Consideration of a possible clinical study of bone loss after surgery highlighted a potentially more important problem for research. This was the question of survival after surgery and whether there was a link to smoking status. An almost unique patient data set, recorded over several decades, had been maintained by a surgeon. This, coupled with the recently available computer power (even though the data were still entered on punch cards), enabled me to undertake a rigorous statistical analysis and demonstrate the relationship between survival after gastric surgery and smoking status [37, 38, 39].

Collaborative research with colleagues also led to a number of publications of a statistical nature [40, 41, 42] whilst I was in Edinburgh and London.

Magnetic Resonance

At the beginning of the 1980s, whilst still in Edinburgh, I recognized the limitations of neutron activation analysis in the clinical environment and also I felt that there was limited further physical science research to be done in the area of photon absorptiometry. I had become aware of exciting developments in in-vivo nuclear magnetic resonance and I therefore made a conscious decision to change the physical science techniques I would use for quantitative measurements in-vivo. Clearly a major difficulty was access to the very expensive equipment which was still in the early stages of development. I became involved with a proposal to seek funding to install the first pre-production prototype clinical MRI system in Scotland. As an MRC senior research fellow I was responsible for the scientific research of the programme. I therefore became actively involved in research to develop new techniques of magnetic resonance imaging (MRI) using the low field magnetic resonance system, and also to explore potential areas of clinical applications.

I published papers on some of the practical aspects of MRI [43, 46, 51, 59] and collaborated with radiologists on exploring its use for qualitative imaging of the brain, having extended the system to obtain sagittal images. However my main research interests were firstly the quantitative measurement of the magnetic resonance relaxation rate $T_1$, particularly in the brain and heart and secondly the quantitative measurement of velocity and flow.

In MRI the $T_1$ value is related to the ratio of free to bound water and, as a consequence, MRI offered a method to measure water content of tissue. With colleagues I studied changes in brain water in chronic alcoholic patients [44, 48,
62, 68] and, more importantly, related $T_1$ measurements to direct measurements of brain water content in patients with brain tumours [53, 56].

Though elsewhere the use of MRI was generally being investigated in the brain and CNS, we started investigating its use in the heart [45]. During this time my research group produced the first quantitative $T_1$ measurements in the human heart [49] which required the synchronisation of the MRI acquisition with the movement of the subject's heart. Qualitative cardiac images had recently been acquired elsewhere but there was much greater complexity involved in obtaining gated quantitative $T_1$. The technique was used to investigate acute myocardial infarction [47, 55, 60, 61] and monitor changes with time; there was a clear increase in $T_1$ in recent infarcts.

At the time there was some debate about the ability of $T_1$ and $T_2$ to be tissue specific and I contributed to this debate, based on our experience of quantitative measurements [50]. There was also a clinical interest in measuring the breast using MRI but the MRI system was insufficiently sensitive because of its vertical magnetic field and large body coil. Other MRI systems had horizontal magnetic fields which easily enabled specialist orthogonal vertical field surface coils to be developed. I developed an new surface coil for breast imaging for vertical field magnets [54] which was the subject of two patents [86, 87].

The influence of movement on the phase of the NMR signal was known and, utilising this relationship, my research group was one of the first to develop a quantitative technique to measure fluid velocity in-vivo. We used two separate gradient pulse sequences which were subtracted to eliminate spatial phase variations and so produced an image which was directly related to velocity. We were already using gradient echoes for imaging so were able to develop MRI to produce velocity maps [52]. In addition to measuring blood velocity we explored its uses at low velocity levels, demonstrating the ability to measure the pulsatility of CSF [57] and its value for identifying communicating syringomyelia [58].

Having contributed significantly to the development of MRI science and technology at a low field strength I was approached to move to Guy's Hospital in London in 1986 to take on a senior academic position and continue research using one of the first high field MRI systems which had the field strength to offer the potential of in-vivo spectroscopic measurements.

Whilst in London I continued and developed my earlier interest in cardiac imaging and flow measurement, using the increased sensitivity and resolution of a much higher field strength magnet. I extended my cardiac research to the investigation of cardio-vascular abnormalities in infants [65, 66, 67] which concentrated on the accurate special localisation of the defect. I extended my research into MR flow measurements to the development of MR angiography [64, 70] and extending the quantification of it to the measurement of time-average flow [72, 73] and its evaluation against Doppler ultrasound.

The high field strength magnet offered the opportunity to measure, and potentially quantify, phosphorus metabolism. Previous research in the field had used surface coils to localise measurements, predominantly in muscle. My research, along with
a small number of other centres internationally, focused on image localised spectroscopic techniques [63] focusing on the effectiveness and characterisation of the localisation in-vivo [71, 75]. The technique was applied to breast tumours [69, 74, 77, 78] using customised breast coils.

Though this was the early days of MR spectroscopy, I made the decision not to continue in that area as I felt there were inherent scientific problems in utilising phosphorous MR spectroscopy for clinical usage. This also corresponded to a time, in the late 80s, where it was becoming increasingly difficult, if not impossible, for researchers in an academic environment to make the necessary scientific and technological changes to commercially produced MR systems to pursue research in certain areas. In addition this coincided with my view that image analysis would have an increasing role in the future and that a number of developments in imaging techniques, including MR, would require an image analysis approach.

This awareness coincided with my decision to move to Leeds in 1989 to take up the chair in Medical Physics at the University of Leeds where, once I had raised funding to purchase MRI systems, I continued with applied MR research. I continued research in the area of angiography [76, 79] but also extended this to the development and implementation of perfusion measurements, particularly in infants, including neonates [80, 82]. The difficult task of attempting to measure perfusion in cardiac tissue was pursued by my research group, the first stage being the development of quantitative $T_1$ in different phases of the cardiac cycle in a single breath-hold [84]. Another potential method for estimating perfusion is the quantification of pharmacokinetic parameters and my research group developed a practical clinical technique [85] which is being used for cardiac and rheumatologic applications.

The use of MRI for quantitative radiation dose measurements had been around for many years. I felt that the technique could have practical use, with the increasing availability and usage of conformal radiotherapy, but I was concerned about the relatively poor quality of the evidence which I saw in published papers. On that basis my research group undertook to review the evidence and evaluate the performance of the latest generation of polymer gels used for dosimetry [81, 83].

During the period of my research in MRI, the technology had evolved from MRI systems in the laboratory which I could modify as required, through more restricted commercial systems which I could configure differently, to finally relatively closed commercial systems which could not be modified without extensive negotiation and delay. The opportunity for significant physics research in MRI was therefore severely restricted for someone like myself, working in the university and health sectors. I had recognised this growing restriction in the late 1980s at the same time as I recognised that many research problems may best be resolved by image analysis methods rather than by developments in MRI techniques. For this reason my research in image analysis, which only started to flourish in the early 1990s after my move to Leeds, has often focused on MRI; this work is described in the later section about image analysis.
Body Composition Measurement

In the 1980s there was interest in the measurement of body composition in vivo, particularly body fat and muscle mass, initially to monitor parenteral nutrition and the after effects of surgery; of particular interest was the measurement of nitrogen as a better indicator of muscle mass. After moving from London to Leeds in 1989 I was able to utilise my experience in both neutron activation analysis and dual photon absorptiometry to provide leadership to a newly formed Bone and Body Composition Research Group.

I had previously been aware of the potential advantages of prompt neutron activation analysis to measure body nitrogen and initiated a research project whilst in Edinburgh [88] but recognised problems with that method associated with sensitivity. My move to Leeds in 1989 offered the unique opportunity to explore the use of the high and controllable output of neutron generators for body composition measurement. We developed a much improved pulsed neutron technique to measure body carbon, hydrogen and oxygen [89, 90] and considered whether it could be used to measure glycogen [91]. This work was extended to consider the measurement of total body water, fat and protein from the measurement of the ratio of the major elements [93].

We also undertook research into a recently published whole body electromagnetic method for body composition measurement which we thought may overcome some of the problems of bio-impedance methods [95]. However we found major scientific flaws in the technique [96], which to my knowledge is now no longer in use.

The commercially available techniques such as dual energy X-ray absorptiometry and bio-impedance analysis and laboratory techniques such as total body potassium offered an array of measurements of total body fat, fat-free mass, water, and lean-tissue mass. Research was undertaken to understand the relationship between the different methodologies [92, 94] with a particular focus on patients on renal dialysis [97, 98, 99, 100, 101, 102, 103, 105].

Image Analysis

My move to Leeds in 1989 coincided with my emerging view that the development of medical image analysis was of growing importance in the next stage in extracting quantitative information from MRI and other imaging techniques; indeed I had set up a medical image analysis group at Guy's Hospital just prior to my departure.

As outlined earlier I had become concerned that there should be a natural progression from basic scientific research, through applied research to a rigorous analysis of the efficacy and effectiveness of an investigative technique in routine clinical use. These three fundamental stages in research underpinned the creation of a multidisciplinary Centre of Medical Imaging Research (CoMIR), which I set up in Leeds in 1992 and of which I was Director until 1998, to undertake research in all these three areas. My own personal research which I pursued from the early 1990s in medical imaging was in magnetic resonance imaging (described above),
image analysis (described in this section) and evidence based imaging which developed beyond expectations and is described in the next section.

The possibility of using the spatial distribution of mineralisation in low resolution hip images obtained using DXA to improve the prediction of fracture risk was studied [106] and the image analysis methods developed demonstrated the benefit of such an approach. Unfortunately, however, it was beyond the scope of my research group to test the technique further by implementing it in a sufficiently large prospective clinical study.

New developments in MR offered the opportunity of superimposing darker grids onto tissue, so-called ‘tagging’; these offered greater opportunity to quantify the movement of cardiac muscle during the cardiac cycle. Research was undertaken to perform the necessary simulation before the technique could be used for accurate quantification of heart movement [107, 109]. The increased usage of 3D data sets from MR and the need to combine them led to research to develop a mathematically robust method for 3D registration [110].

Familiarity with techniques for the extraction of quantitative information, such as texture from medical images, led to research into how, in effect, a reversal of the process could be used to create better psychophysical tests to access an individual’s ability to view medical images [111]. The outcome of the evaluation of this method has just been accepted for publication.

Evidence Based Imaging

Over a decade before my first publication in evidence based imaging I had become concerned about issues associated with the assessment of health technologies [112], and I had become familiar with the debate around evidence based medicine. With the creation of CoMIR soon after my move to Leeds I ensured that there was a focus in the area of evidence based imaging, both in terms of primary research and also secondary research, ie the scientific evaluation of existing published material.

My emphasis on evidence based imaging was planned to coincide with changes in funding policy which offered the opportunity to focus staff and resources appropriately. Initial research highlighted, somewhat unexpectedly, the relatively poor quality of evidence in medical imaging research and the presence of bias in many studies [113]. It was clear that a simple focus on randomised controlled trials, as used in the Cochrane Collaboration, could not be applied to the evaluation of medical imaging research, and that we would need to undertake research into methodological issues.

For example it was necessary to develop a decision analytic model for the role of ultrasound in primary coronary stenting before a full review could be undertaken. In addition research had to be undertaken into the identification of studies for systematic reviews in medical imaging [114] and also how to undertake a review in a rapidly evolving field [116].

Extensive systematic reviews were undertaken in four areas of medical imaging: endoscopic ultrasound in gastro-oesophageal cancer [115] {118}; spiral and
electron beam computed tomography \(\{119\}\); intravascular ultrasound-guided interventions in coronary artery disease \(\{120\}\); and magnetic resonance angiography of carotid artery stenosis and peripheral vascular disease \(\{117\}\ \{121\}\). These systematic reviews \(\{118, 119, 120, 121\}\) are large published monographs, each running to over 100 pages long, and their length precludes the inclusion of hard copies; instead copies have been included on a CD to be found on the back cover.

In 1998 I was President of the British Institute of Radiology and chose the subject of ‘Evidence Based Imaging’ for my Presidential Lecture giving me the opportunity to stimulate a wider debate in this area. However although I felt I had made significant progress in evidence based imaging I came to the conclusion that it was strategically appropriate to begin to withdraw from the field in the late 1990s for a number of reasons. Firstly it was difficult to continue our secondary research in medical imaging because the funding policy was to concentrate support in centres which specialised in systematic reviews, rather than specialist research centres such as CoMIR which also undertook systematic reviews; thus it was very difficult to recruit and retain the necessary expertise. Secondly we were unable to obtain and maintain the appropriate clinical support for good quality primary research.

**Terahertz Imaging and Sensing**

Throughout my career I had recognised the value of imaging for the localisation of quantitative measurements but I also recognised there were limitations about what could be measured utilising certain physical science techniques. In the late 1990s this led me to consider other parts of the electromagnetic spectrum where, perhaps, any signal variation could potentially be influenced more by cellular or molecular changes. As a consequence I developed an interest in the use of terahertz radiation \(\{122\}\) which could potentially act both as an imaging technique \(\{123\}\) and also a sensor in a localised region. I recognised that instead of using large volume images of a human subject to localise an abnormality and perform a relatively crude quantitative measurement, it may be possible to develop techniques that could be delivered using minimal invasive surgery techniques. Though these may only provide imaging information over a relatively small volume, they could produce much more valuable information about tissue at a highly localised point.

My research has been involved in understanding some of the science behind terahertz radiation and the subject is still in its early days of development. Research was undertaken to model the propagation of terahertz radiation through tissue \(\{127, 129\}\) and also catalogue the optical properties of tissue at terahertz frequencies \(\{125, 132\}\). In developing an imaging technique we were surprised to identify gaps in data used to define safety criteria which could influence issues of safety in the terahertz region \(\{126, 130\}\). As terahertz systems were developed to image tissue, we undertook research to determine how to evaluate image quality using test objects \(\{124\}\) and also the effect of frequency on image quality \(\{131\}\). Multispectral clustering techniques were used for the extraction of quantitative information from the imaging data to classify histopathology \(\{128\}\).
Summary
The value and importance of the physical sciences for measurement of health and disease in-vivo has been demonstrated by my research. I recognise that I have been fortunate that my research has coincided with significant technological developments, opening up avenues of research that would have been unimaginable to my predecessors.

Working over a span of three decades I have contributed to fundamental and applied knowledge, changing my approach and methodology as opportunities present themselves, whilst retaining my focus on quantitative, non-invasive measurements in the human body.
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Development of Apparatus to Measure Calcium Changes in the Forearm and Spine by Neutron Activation Analysis using Californium-252

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Abstract. Techniques were developed to measure small changes of calcium in the forearm and spine in vivo by neutron activation analysis using two sources of 252Cf in a hospital environment. Using purpose-built part-body counters and bilateral irradiation with 7-5 cm premoderation between the sources and the bone, peripheral bone was measured with a total source strength eventually as low as 50 mCi. Two methods of spectral analysis were used and compared. Patient studies of the forearm were successfully undertaken, with a precision of 2-6% which included patient movement, and an annual bone dose of less than 10 rem and skin dose of 25 rem from six measurements.

Two 100 mCi sources were used for measurements of the lumbar spine. Care was taken to minimise the problems of non-uniformity of activation which are present using unilateral irradiation. Emphasis was placed on measuring the bodies of the vertebrae with adequate sensitivity and uniformity, and the spinous processes and arches with low sensitivity. A whole body counter was used for the bilateral detection of the induced activity. The precision of the method was 3-0% with an annual peak bone dose of 2-1 rem and skin dose of 18 rem from three measurements.

I. Introduction

Since the first measurements in vivo on human volunteers by neutron activation analysis (Anderson, Osborn, Tomlinson, Newton, Rundo, Salmon and Smith 1964) the field has expanded to include both whole-body and part-body measurements of a range of body elements. The element that is of most clinical interest is calcium, and apparatus for measuring part-body calcium changes can be constructed relatively cheaply for use in a hospital environment.

Most part-body calcium measurements on patients have been performed on peripheral bone, and although large neutron sources have been used (Comar, Riviere, Raynaud and Kellershohn 1968), most studies have been performed with compact radionuclide neutron sources using the 48Ca(n,γ)49Ca reaction. Measurements of the hand have been carried out using a single 25 Ci Am–Be source (Catto, McIntosh and MacLeod 1973), a single 75 mCi 241Cf source (Guey, Leitienne, Zech and Traegar 1978) and a combination of two 100 mCi 241Cf and four 10 Ci Pu–Be sources (Maziere and Comar 1976). No activation measurements have been performed on the forearm, the site which is used for bone mineral determination by photon absorptiometry (Cameron and Sorenson 1967). In order to make a direct comparison with absorptiometry and because
the long bones of the forearm could be easily immobilised during any measurements, the forearm was chosen as the site for our investigations of peripheral bone loss.

Although a method of utilising twelve $\alpha$ Ci Pu–Be sources has been used for calcium measurements of the trunk (McNeill, Thomas, Sturtridge and Harrison 1973), part-body measurements of the spine alone have only been performed using a cyclotron (Al-Hiti, Thomas, Al-Tikrity, Ettinger, Fremlin and Dabek 1976). We have developed apparatus which could be used in a hospital to measure the calcium changes in the lumbar spine using $^{252}$Cf sources.

2. Neutron sources

Two $^{252}$Cf sources were obtained for activation analysis. They had previously been housed for three years at the Scottish Universities Research and Reactor Centre where some physical evaluation had taken place (Boddy, Robertson and Glaros 1974) and a pneumatic delivery system developed (Glaros 1975), before being moved to Edinburgh. Although the ideal source strength for patient measurements has been stated as 200 mCi (Boddy et al. 1974) or between 500 mCi and 1500 mCi (Evans, Le Blanc and Johnson 1976), the total combined source strength that was available for patient measurements over a 2½ year period was eventually as low as 50 mCi. At this point the sources were replaced by two 104 mCi sources, mainly for the purpose of spine measurements.

An activation chamber was constructed inside a room by building an L-shaped baffle of concrete blocks and wax. An access 0.75 m wide allowed entry into the 2.5 m × 1.4 m irradiation chamber where the sources were stored 3 m below ground.

3. Forearm measurements

3.1. Activation

Two designs of apparatus were used for activation measurements of the forearm, both using a bilateral irradiation geometry. The sources and limb were surrounded by a hydrogenous material, which acted as a neutron 'reflector', to increase the thermal flux through the bone.

The premoderator thickness was chosen not to give the maximum thermal flux, which occurs at a depth of approximately 1.5 cm, but to give the minimum absorbed radiation dose for a given induced activity. Combining data from the variations in dose and thermal flux with thickness of premoderator and activation measurements of a cadaver limb, it was possible to calculate the dose per fixed count for a range of premoderator thicknesses (fig. 1). The shape of the curve is a function of the activity of the source when patient measurements are made of radiomucides with short half-lives.

The first forearm irradiation geometry (A), 19-75 cm source separation, 6 cm Perspex premoderator and wax blocks above and below the limb has been previously described (Smith and MacPherson 1977, Smith, Tothill, Simpson, MacPherson, Merrick, Strong and Boddy 1978). This was improved upon and
The second irradiation geometry (B) consisted of the two sources inside cylindrical wax reflectors each side of a water bath into which the patient's arm was fixed at an angle of 45° (fig. 2). The dimensions of the tank were

Fig. 1. Curve 1 shows the relative dose per unit activation for radionuclides with a long half-life. For induced radionuclides with a half-life comparable with the irradiation time, long irradiation times are not rewarding. Curve 2 shows the dose in rem required to obtain a statistical coefficient of variation of 2% when activating $^{48}$Ca with a total $^{252}$Cf activity of about 55 mCi.

Fig. 2. Forearm irradiation geometry (B) incorporating a water bath to act as combined premoderator and reflector. The dotted lines indicate the position of the second wax reflector.
50 cm × 45 cm × 17 cm so that the water acted as a premoderator, giving a total of about 7.5 cm premoderation between source and bone, as well as a reflector. With the water bath the sensitivity of activation over a 25 cm length was found to be improved by a factor of 1.8 compared with the first irradiation geometry. Part of this improvement, about 30%, was due to the smaller source separation, but the major increase in sensitivity was due to the elimination of air gaps around the forearm. The latter also reduced the variation in calcium counts from patient to patient due to overlying soft tissue, increasing the possibility of absolute measurements.

During the 10 min activation period the patient gripped a Perspex rod and the forearm was restrained in a mould constructed from butyrate sheet. The variation in thermal fluence across the central 5 cm was ±4.2% and the full width at half maximum (FWHM) of the variation along the arm was 16.2 cm, measured using 1 cm diameter 0.025 cm thick gold foils.

Dose measurements were performed in a phantom using gas flow ionisation chambers and a GM counter (Law, Lawson and Porter 1974). During a 10 min irradiation with a total source strength of 65 mCi the subject received a neutron dose of 0.68 rad and a gamma ray dose of 0.59 rad to the skin surface and a neutron dose of 0.20 rad and a gamma ray dose of 0.26 rad to the bone. Combining the variation of RBE with energy and the calculated neutron spectrum, a mean quality factor (QF) of 7.8 was obtained for the neutrons after 6 cm premoderation. The dose equivalent to the skin surface was therefore 5.8 rem and 1.8 rem to the bone. The gonad dose was less than 14 mrem taking the value QF = 10. For patient studies the irradiation time was kept constant so the maximum annual dose to the patient from six measurements was less than 10 rem to the bone or 4.5 rem averaged over the forearm.

3.2. Detection

A part-body counter (fig. 3) was built for the detection of induced activity in peripheral bones and the thyroid. Two 15 cm × 10 cm NaI detectors were encased in 5.0 cm thick lead and could be used in a bilateral geometry with or without a large central shield which could surround a limb such as the forearm. When used with the central shield there was a thickness of at least 7.5 cm of lead in all directions around the crystal except in the direction of the limb entrances of the central shield. All three sections could be moved around a large steel table 0.84 m above the floor. The background in the calcium region, 2.9 MeV to 3.8 MeV, with a 9 cm separation between the detectors in the central shield was 0.09 cpr and with a 12 cm separation without the central shield was 0.13 cpr. The variation in sensitivity at 2.75 MeV in the central 5 cm between the detectors for these separations was ±6% and ±5.8% respectively. The FWHM for the variation in detection sensitivity along the arm was 21.8 cm.

At the end of the irradiation period the patient was transferred to the part-body counter with a delay of 1 min. The forearm was counted for 1000 s
in the central shield of the counter with a detector separation of 9 cm, the forearm being constrained in the same way as for activation. The combination of activation and detection had a FWHM of 12.7 cm centred 13.5 cm from the centre of the fist (fig. 4).

Fig. 3. Part-body counter. One detector had been removed from the central shield to show the position of the forearm support.

Fig. 4. Combined activation and detection profiles to show region of measurement of the forearm. The curves illustrate the improved sensitivity of calcium measurement using the water bath.

4. Data analysis and precision

The spectrum obtained from a forearm after a 10 min irradiation, 1 min delay and 1000 s count is shown in fig. 5. It was not possible to wait for the calcium to decay in order to calculate the contributions from $^{36}$Cl and $^{24}$Na to the calcium peak, because during this time there would be a significant decrease in the $^{24}$Na peak due to redistribution in blood and soft tissue. Two methods were used to calculate the $^{49}$Ca contributions to the spectrum. In the first the patient spectrum and standard spectra of $^{24}$Na and $^{49}$Ca were integrated over the regions of the two photopeaks. Solutions of the two
simultaneous equations gave the sodium and calcium contributions in the two regions.

The second method used a computer to fit the standard spectra of $^{34}$Cl, $^{24}$Na and $^{49}$Ca to the patient spectrum. Matrix algebra was used to fit simultaneously the three standard spectra using a least-squares technique. The intrinsic precision of the two methods was compared by taking a single patient spectrum and randomly shifting each point in a Gaussian distribution within the limits of counting statistics. The coefficient of variation (cv) of the two methods based on manipulated spectra from a single patient spectrum was 2.4% and 2.1% respectively.

Precision measurements on a cadaver limb showed that with the low source activity, small changes in a patient could best be detected by performing sets of three repeated measurements at the beginning and end of a time period (e.g. six months) rather than six individual measurements performed throughout that time period. In addition, this irradiation programme enabled accurate repeatability figures to be obtained, which could include the effect of patient movement. From the many sets of repeated patient measurements the error due to movement (cv$_{\text{m}}$), which was independent of the number of counts measured, was calculated. This could then be combined with the statistical cv to give the overall precision (total cv) for each patient. Table 1 shows values for both irradiation geometries and both data processing techniques. The precision figures quoted were based on the mean calcium counts for both methods and the least significant change was calculated from the standard error of the mean of the sets of measurements, taking the level of significance at $P < 0.05$.
Neutron Activation Analysis of Part-body Calcium

Table 1. Reproducibility of the two forearm irradiation apparatuses and methods of spectral analysis based on sets of repeated patient measurements

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<th>CVm (%)</th>
<th>Precision (%)</th>
<th>Least significant change (%)</th>
<th>No. of sets of patient measurements</th>
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<td>3.45</td>
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5. Spine measurement

It is recognised that $^{252}$Cf is not the ideal source for measurements of the spine because of its relatively low mean neutron energy. However, extensive experiments were performed to determine the optimum irradiation geometry to measure changes in calcium in the lumbar spine using $^{252}$Cf.

Changes in the bodies of the vertebrae are of most clinical interest, so good uniformity of activation and detection was sought in the region 5 cm to 9 cm below the skin surface, which is the approximate position of the region of interest (Brinkley and Masters 1967). In addition, effort was made to measure the region 0 cm to 5 cm deep with as low a sensitivity as possible so that calcium counts from the spinous processes and arches of the vertebrae contributed as little as possible to the total calcium counts detected. Thus changes in the bodies of the vertebrae, where the calcium losses are most important, would be most likely to be detected.

5.1. Activation

The variation in thermal neutron flux with depth was investigated in fourteen different irradiation geometries. Reflector position, thickness of premoderator and source to skin distance were varied and the effect of a thermal neutron absorber was investigated. The efficiency of activation, using both phantoms and a cadaver, and the dose were then measured in the geometries with reasonable thermal neutron flux profiles.

The flux profiles of the two irradiation geometries chosen as most suitable are shown in fig. 6. Geometry 1, to be used for the majority of patients, gave the best thermal fluence uniformity. The sources were situated 6 cm from the skin surface and no premoderator or reflector was used. Geometry 2 would only be used for patients of very small stature, where the bodies of the vertebrae were likely to be in the region 3 cm to 6 cm below the skin surface, and so would be measured fairly uniformly and with higher sensitivity. For this geometry the sources were situated 6 cm from the skin surface and surrounded
Neutron Activation Analysis of Part-body Calcium

5.2. Detection

Counting was performed bilaterally with the patient supine in a shadow shield whole body counter incorporating four 15 cm \times 10 \text{ cm} \ NaI detectors, two above and two below, separated by 41 cm. In this position, a background of 0.25 \text{ cps} was obtained in the 40Ca region. The variation in sensitivity in the approximate region of the vertebrae, 6 cm to 9 cm above the bed, was \( \pm 8.0\% \). The variation along the spine 7 cm above the bed was 43-2 cm (fwhm) which was compatible with the activation profile 7 cm below the skin surface of 36-0 cm (fwhm).

A pilot study was carried out using the spine irradiation apparatus (geometry 1), sets of three measurements being performed on ten volunteers. The results showed that using two 100 mCi sources, the lumbar spine could be measured with a precision of around 3.0\%. The dose, measured using nuclear emulsion films from a 200 \text{ s} irradiation, would be 6 rem to the skin, 0.7 rem to the bone (0.4 rem averaged over the whole spine) and 0.2 rem to the female gonads. The mean calcium count from this dose would be approximately 2000 counts.

The 1000 \text{ s} counting time was split into three periods of 200 \text{ s}, 200 \text{ s} and 600 \text{ s}, the first two of these being used to determine the feasibility of phosphorus measurements, via the \( ^3\text{P} \,(\text{n}, \alpha) ^{28}\text{Al} \) reaction (threshold 2.0 MeV, \( T_1 = 2.3 \text{ min} \)). The number of counts from \( ^{28}\text{Al} \) varied enormously, the statistical error being greater than 10\% in five out of nine volunteers. Thus, as would be expected, the neutron energy of \( ^{252}\text{Cf} \) is too low for phosphorus activation.

6. Discussion

Apparatus has been developed to use the technique of neutron activation analysis to monitor calcium changes for use in a hospital. Sequential measurements of the forearm have been performed now for two years on over 90 patients and the technique has been proved to be a useful research tool. Significant changes both in individual patients and in groups of patients with respect to treatment have been detected and new forms of treatment evaluated (Winney, Tothill, Robson, Abbot, Lidgard, Cameron, Smith, MacPherson and Strong 1977).

The use of a water bath acting as premoderator and reflector, and the matrix method of spectral analysis have been shown to give the best precision. The advantage of this latter technique is, however, marginal when used with the water bath, and so the simultaneous equations method could be used with almost equal success for clinical measurements.

The characteristics of the forearm irradiation and detection apparatus compare favourably with other methods of monitoring sequential changes of calcium at peripheral sites. The precision value of 2.6\% for a single measurement is better than those achieved by Catto et al. (1973) and Maziere and Comar (1976) who quote values of 5\% and 3\% respectively. The bone dose is also less, due both to the lower mean neutron energy and the relatively large amount of premoderator used. The precision of activation analysis of the forearm also
Haus verwendet wurden. Es wurden Methoden zur Messung der Änderungen der Kalziumwerte im Vorderarm und im Rückgrat entwickelt. Sie wurden am labenden Organismus, d.h. in vivo, durchgeführt und beruhen auf Neutronenaktivierung als analytische Methodik, bei der zwei \( ^{252} \text{Cf} \)-Quellen im Krankenhaus verwendet wurden. Unter Einsatz speziell gebauter Zähler zur Messung der Radioaktivität...
in bestimmten Körperräumen und einer zweiseitigen Strahleninwirkung bei einem, 7,5 cm langen Strahlenbremsschirm zwischen den Quellen und der zu messenden Haut wurden die Peripheriezonen von Knochen bei einer Gesamtstärke der Quelle, die zum Schluss nur noch 50 mCi betrug, gemessen. Angewandt wurden zwei verschiedene Methoden der Spektralanalyse und die so erhaltenen Werte wurden mit einander verglichen. Die Messung des Kalziumstoffwechsels im Vorderarm von Patienten wurde somit erfolgreich durchgeführt, indem eine Genauigkeit von 2,4% erreicht wurde, sofern man die Bewegung des Patienten noch mitberücksichtigt. Die jährliche Dosis für die Knochen lag unter 10 rem und diejenige für die Haut bei 35 rem. Diese Werte verstehen sich für sechs Messungen.

Zwei Strahlenquellen mit je 100 mCi wurden zur Messung des Kalziumwechsels im Rückgrat in der Lendenwirbelsäule eingesetzt. Hierbei wurde vorsichtig darauf geachtet, diejenigen Schwierigkeiten so gering wie möglich zu halten, welche sich aus der Ungleichförmigkeit der Aktivierung bei Verwendung einer einseitig wirksamen Bestrahlung ergeben. Das Schwingungswirke wurde darauf gelegt, die Wirbelsäulentente mit ausreichender Empfindlichkeit und Gleichförmigkeit sowie die Veränderungen der Wirbelsäule und der Wölbungen mit niedriger Empfindlichkeit zu messen. Zur Messung der induzierten Radioaktivität bei zweiseitiger Strahleninwirkung wurde ein Zählrohr für die Knochen eingesetzt, das den ganzen Körper erfassend. Die Genauigkeit der Methode betrug 3,0% bei einer jährlichen Pizotendosis für die Haut von 2,1 rem und einer Dosis für die Haut von 18 rem. Diese Werte gelten für drei Messungen.

References

Short Communication

Measurement of spinal calcium by in vivo neutron activation analysis in osteoporosis

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1. Introduction

Osteoporosis is the most common disease of bone, affecting in particular the elderly female population. The lumbar spine is of particular interest in the study of this disease for two reasons. Firstly, lumbar vertebral crush fracture is a common manifestation of osteoporosis, suggesting that the spine appears to be particularly vulnerable. Secondly, the normal turnover rate of bone at such trabecular sites is 8% per annum compared with 3% per annum in the cortical bone of the metacarpals or the radius (ICRP 23 1975). Any change or response to treatment may thus be made more apparent by measuring the lumbar spine than by using conventional techniques of metacarpal index or photon absorptiometry of the radius.

Methods for measuring changes in calcium content of the lumbar spine by in vivo neutron activation analysis have been described using $^{252}$Cf (Smith and Tothill 1979), and elsewhere using a cyclotron (Al Hiti et al 1976). No report of changes in spinal calcium content in response to treatment for osteoporosis has yet been published.

This study set out to evaluate the technique of part-body neutron activation analysis of the spine as a method for monitoring calcium changes in elderly osteoporosis patients.

2. Methods and patients

The apparatus used consisted of a modified dental chair that could tilt back to minimise patient movement. Two $^{252}$Cf sources, each initially 2.96 GBq (80 mCi), were positioned behind the patient 6 cm from the skin surface and 20 cm apart, centred on the iliac crest. No premoderator or hydrogenous reflector was used. The patient was irradiated for 3.5 min. The induced activity was measured in a whole-body counter containing four 15 cm x 10 cm NaI detectors for 1000 s (Smith and Tothill 1979). As there was no collimation of the neutron flux, parts of the thoracic spine, sacrum and pelvis were also activated, in addition to the lumbar spine.

Twenty-three women presenting with fractured neck of femur were studied. Their mean age was 69.5 years (range 43-80 years) and all were post-menopausal. Twenty of these patients had normal blood biochemistry and were randomly allocated to one of four treatment groups: (a) placebo, (b) ethinyloestradiol (oestrogen) and calcium
supplement (Ca), (c) 1α-hydroxycholecalciferol (1αOHD₃) and Ca, (d) 1αOHD₃, oestrogen and Ca (Eastell et al 1980). Three patients with possible osteomalacia, as indicated by blood biochemistry, were given 1αOHD₃ and Ca.

The patients were measured at intervals of four months for one year. In addition to the measurement of spinal calcium, the mineral content of the non-dominant forearm was measured using photon absorptiometry at a site 5 cm proximal to the tip of the styloid process of the radius. Patients were assessed radiographically at their first and last visit to detect the occurrence of vertebral crush fractures during the period of the study. It was recognised that an increase in the thickness of soft tissue over the spine would decrease the efficiency of activation, so patients who showed a consistent weight increase of more than 5 kg during the year were omitted from the study.

3. Results

An average of 997 counts was measured in the ⁴⁸Ca photopeak in these patients. After correcting for decay of the ²⁵⁵Cf, the annual rate of change in spinal calcium content was calculated by fitting a regression line to the four measurements. The combined residual sum of squares from the regression lines gave an overall estimate of reproducibility (coefficient of variation) of 5% for all the patient data.

Three patients showed a consistent weight increase (probably as a result of the oestrogen therapy), one suffered a crush fracture during the study and one was found to suffer from rheumatoid arthritis. The results of the remaining fifteen osteoporotic patients are shown in figure 1. A one-way analysis of variance (Kruskal and Wallis 1952) suggested that the results were not of the same population (p = 0.6). The treatment that was expected to be of most benefit, 1αOHD₃, Ca and oestrogen (Marshall and Nordin 1977, Nordin et al 1980) was significantly better than placebo (P < 0.05). There were no other significant differences between the groups. The predictable response to vitamin D therapy in the three osteomalacic patients was demonstrated by an increase in spinal calcium in all three patients, the increases being significant (P < 0.05) in two of the three.
Similar analysis of the photon absorptiometry measurements of the forearm gave a figure for the reproducibility of 4.4%. Analysis of the data suggested that all the results from the osteoporotic patients came from the same population \((p = 0.45)\) and no significant differences were seen between groups. No significant correlation was found between changes in the forearm and spine \((r = 0.21)\).

4. Discussion

The results of this study demonstrate that NAA of the spine using \(^{252}\text{Cf}\), perhaps the most convenient and inexpensive neutron source, can be successfully used to monitor calcium changes due to osteoporosis and its treatment in the elderly female population. The reproducibility figure of 5% is worse than the figure of 3% which had been achieved in a previous small pilot study (Smith and Tothill 1979) on a group of patients not suffering from osteoporosis. This can be accounted for by the lower calcium levels in the osteoporotic patients and hence a larger error due to counting statistics, plus their difficulty in sitting still during the activation period, due to pain in their recently fractured hip. The assessment of reproducibility in the osteoporotic group assumes a linear change with time; such an assumption may lead to an over-pessimistic figure for reproducibility.

The expected 'best' treatment regime, Ca, \(1\alpha\text{OHDA}\), and oestrogen was shown to be significantly better \((P < 0.05)\) than placebo by spine NAA. This result was not demonstrated by forearm absorptiometry, even though the latter had better reproducibility. The lack of correlation between changes in bone mineral in different parts of the skeleton confirms other findings (Aloia et al 1975, Cohn et al 1975, 1976, Dabek et al 1977, Harrison et al 1974).

The patients in the study were selected from a group of 180 patients presenting with fractured neck of femur over a seven month period. It had been hoped that about half of these would be suitable for the study, hence the four treatment groups, but the high morbidity and mortality rates associated with this fracture reduced the available number considerably. The low admittance rate into the study was not due to difficulties associated with the NAA measurement technique.

Nonetheless there is one drawback of the technique. This is that an increase in body tissue in the region of the spine will affect the sensitivity of the activation. As some patients steadily increase their weight as a result of receiving oestrogen, these patients must be eliminated from any analysis of the results or, preferably, experiments must be performed to correct for variations in activation and detection efficiency.

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Long-Term Effect of Dialysate Calcium and 1α-Hydroxycholecalciferol on Bone Calcium Content in Haemodialysis Patients as Measured by Neutron Activation Analysis of the Forearm

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Key Words. Bone calcium · Haemodialysis · 1α-Hydroxycholecalciferol · Dialysate calcium concentration · Neutron activation analysis

Abstract. The effect of different concentrations of dialysate calcium and treatment with 1α-OHD₃ on bone calcium content in haemodialysis patients was investigated over a 2½-year period. Part-body neutron activation analysis of the forearm with ²⁵²Cf was used to measure both an initial absolute calcium content and changes in the bone calcium. A highly significant inverse correlation (r = -0.84, p < 0.001) was found between the initial calcium content and the duration of previous dialysis utilising a dialysate calcium concentration of 1.75 mmol/l. No significant difference was found between the changes in bone calcium in patients who continued using a dialysate calcium of 1.75 mmol/l and those whose dialysate calcium was reduced to 1.375 mmol/l. There was a significant increase of 16.7% (p < 0.01) in bone calcium over the 2½-year period in the patients receiving 1α-OHD₃. The results of the study suggest that bone calcium loss is not necessarily influenced by dialysate calcium concentration but that 1α-OHD₃ has a beneficial long-term effect on bone calcium content.

Introduction

Renal osteodystrophy in haemodialysis patients is a complex mixture of osteomalacia, secondary hyperparathyroidism and osteoporosis. While it is now recognised that disturbance of vitamin D metabolism has a central role in the pathogenesis of renal osteodystrophy, the concentration of calcium in the dialysis fluid has also been thought to be of importance in haemodialysis patients [1]. However, while using a dialysate calcium concentration of 1.5 mmol/l or higher may prevent hypocalcaemia and improve calcium balance [2-5] there is still controversy regarding the effect of dialysate calcium concentrations on bone disease [6,7]. By contrast, short-term treatment of renal osteodystrophy with synthetic active metabolites of vitamin D has produced promising improvement in renal bone disease [8] except when this is aluminium-induced [9].

The end result of all forms of renal osteodystrophy is loss of bone mineral and calcium which may result in bone pain and fractures. Since bone mineral content may change slowly as renal bone disease progresses, assessment of its response to treatment requires a means of detecting small changes in bone mineral or bone calcium content. Skeletal radiography is a qualitative technique which detects only severe changes while bone histology requires expertise for quantification of changes and since it is traumatic for the patient, is not ideal for frequent analyses. Neutron activation analysis (NAA) is a non-invasive and precise technique which measures calcium content and would be of value in quantitating small changes in the calcium content of bone. Its usefulness has been confirmed by a number of studies where use of this technique has detected changes in calcium content paralleling biochemical, radiological and histological changes [10].

This study assesses the effect of dialysate calcium and treatment with 1α-hydroxycholecalciferol (1α-OHD₃) on the calcium content of bone in haemodialysis patients over a 2½-year period using NAA to measure changes in calcium content of the forearm.

Patients and Methods

21 patients, aged 22-56 years (mean 40 years) had been established on haemodialysis from 2 to 31 years (mean 4.5 years). These were the
entire population of patients on home dialysis who had not been treated previously with vitamin D. Patients were dialysed for 5–8 h twice or thrice weekly using either Gambro 1M², 1M³ coil or Cordis Dow 1.3M² hollow fibre. Prior to this study, dialysis on all patients was conducted with a dialysate calcium concentration of 1.75 mmol/l and magnesium concentration of 0.75 mmol/l; their diet contained 70 g of protein, 800 mg (20 mmol) of calcium and 1,100 mg (35.5 mmol) of phosphorus daily. In addition, patients received aluminium hydroxide if necessary to maintain the pre-dialysis plasma phosphate in the range 1.6–2.0 mmol/l.

The patients were randomly allocated into one of three treatment groups. Group 1 continued their treatment unchanged with a dialysate calcium concentration of 1.75 mmol/l; 2 patients failed to attend for the study leaving only 3 in the group. Group 2 contained 7 patients who were treated with a low dialysate calcium concentration of 1.375 mmol/l. Group 3 also contained 7 patients who were treated with a low dialysate calcium concentration of 1.375 mmol/l but in addition were given 1α-OHd3 in an initial dose of 2 μg daily, adjusted later if necessary to avoid hypercalcaemia. The initial characteristics of each group are shown in table I. There were no significant differences between the groups at the start of the study.

The calcium content of the non-fistula forearm was measured by NAA at time zero before the treatment alteration, then at 8, 16 and 30 months after the start of the trial. 2 patients in Group 1 did not complete the study: 1 died of cardiac failure after 17 months and 1 had a fistula inserted in the arm being measured after 18 months. In Group 2, 2 patients also did not complete the study, 1 after 9 months due to hypocalcaemia and 1 after 17 months when 1α-OHd3 treatment commenced. In Group 3, 1 patient died just before the final measurement whilst receiving treatment of oesophageal carcinoma. Throughout the period of this study, skeletal radiographs of the ankle, hand, pelvis, chest, lumbar spine and skull were taken to monitor excessive calcium loss.

The measurement of calcium content of bone using NAA involves irradiating the region of interest with neutrons which react with the isotope ⁴⁰Ca in bone to form unstable ⁴⁰Ca. This induced radioactivity, which has a half-life of 8.8 min, can then be measured with a gamma-ray detector to provide a relative estimate of the calcium content of the bone, expressed as measured ⁴⁰Ca counts. In our apparatus [11,12] the forearm is irradiated with two neutron sources of ⁶⁰Co and the activity measured with two 15 × 10 cm sodium iodide detectors placed one on either side of the forearm. A triple irradiation procedure was adopted at each visit giving a precision of 1.8%, the same apparatus and procedure being used on each occasion. The NAA results over the 2½ years of the study were expressed as a percentage of the initial value.

<table>
<thead>
<tr>
<th>Group</th>
<th>Number in group</th>
<th>Age, years</th>
<th>Duration of dialysis months</th>
<th>Plasma calcium mmol/l</th>
<th>Plasma alkaline phosphatase U/l</th>
<th>Plasma parathyroid hormone μg/l</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>4</td>
<td>42 (27-56)</td>
<td>2.65 ± 0.11</td>
<td>130 ± 60</td>
<td>1.03 ± 0.90</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>5</td>
<td>42 (22-52)</td>
<td>2.69 ± 0.31</td>
<td>134 ± 64</td>
<td>1.29 ± 0.72</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>4</td>
<td>36 (25-56)</td>
<td>2.53 ± 0.19</td>
<td>156 ± 113</td>
<td>1.19 ± 1.30</td>
</tr>
</tbody>
</table>

At the time of the last attendance at 30 months, irradiation apparatus using a water bath had been developed so an additional measurement was performed which enabled a more absolute calcium determination to be made using a normalisation procedure to correct for variations due to the patient's size [13]. This value, combined with the changes over the 2½-year period, enabled an initial absolute forearm calcium content to be established in the 14 patients who completed the study. A group of 18 patients awaiting treatment with lithium carbonate were used to define a normal range, so the initial forearm calcium in the renal patients could be expressed as a percentage of normal.

Analysis of data was made using non-parametric statistics unless otherwise stated and p > 0.05 was taken to be not significant.

**Results**

For the dialysis patients as a group the initial calcium content in the forearm prior to the start of the study was significantly lower than normal (mean 87%, p < 0.05, t test). A highly significant inverse correlation was found (r = -0.84, p < 0.001) between the initial bone calcium level and the duration of previous haemodialysis treatment (fig. 1).

![Fig. 1. Relationship between initial calcium content of the forearm, expressed as a percentage of normal, and duration of previous haemodialysis.](image-url)
Bone Calcium in Haemodialysis Patients

Fig. 2. Percentage changes in forearm calcium in individual patients in the three groups over a period of 2½ years expressed as a percentage of the initial value.

Table II. Mean percentage change (±1 SD) in the non-fistula forearm calcium as measured by NAA in the three groups

<table>
<thead>
<tr>
<th></th>
<th>0-8 months</th>
<th>8-16 months</th>
<th>16-30 months</th>
<th>0-30 months</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Group 1</strong></td>
<td>-1.54 (±2.80)</td>
<td>-2.84 (±2.37)</td>
<td>+1.36 (±8.75)</td>
<td>-5.29 (±9.20)</td>
</tr>
<tr>
<td>(n = 5)</td>
<td></td>
<td></td>
<td>(n = 3)</td>
<td>(n = 5)</td>
</tr>
<tr>
<td><strong>Group 2</strong></td>
<td>+2.10 (±5.85)</td>
<td>-0.94 (±3.48)</td>
<td>-6.22 (±6.73)</td>
<td>-3.97 (±7.42)</td>
</tr>
<tr>
<td>(n = 7)</td>
<td></td>
<td></td>
<td>(n = 5)</td>
<td>(n = 5)</td>
</tr>
<tr>
<td><strong>Group 3</strong></td>
<td>+9.26 (±4.82)</td>
<td>+0.63 (±3.31)</td>
<td>+6.49 (±5.37)</td>
<td>+16.65 (±9.42)</td>
</tr>
<tr>
<td>(n = 7)</td>
<td></td>
<td></td>
<td>(n = 6)</td>
<td>(n = 6)</td>
</tr>
</tbody>
</table>

n = Number of patients.

The percentage changes in calcium content of the non-fistula forearm in individual patients in the 3 groups over the period of study are shown in figure 2. The mean changes in NAA calcium in each group expressed as a percentage change over the different time periods are given in table II.

In the patients in Group 3 treated with 1α-OHDA, there was a consistent increase in calcium content which was significantly greater than that in Group 1 (p < 0.01) and Group 2 (p < 0.02) over the initial period of 8 months [14], Group 2 (p < 0.01) over the 16- to 30-month period and in both groups (p < 0.01) over the complete time period of 2½ years. The change in calcium content in Groups 1 and 2 was variable but any differences between these two groups were not significant.

There was no increase in soft tissue or vascular calcification over the period of the study on skeletal radiographs.

Discussion

The reduced forearm calcium content in these patients at the onset of the study, as well as the clear relationship between the initial forearm calcium and the duration of haemodialysis treatment, indicates that prolonged use of a 'high' dialysate calcium concentration of 1.75 mmol/l did not prevent bone calcium loss in these patients. In addition we have also failed to detect any significant difference between changes in calcium content in patients treated with either a high or a low dialysate calcium concentration. These findings contrast with those in earlier studies which indicated that use of a dialysate calcium of 1.5 mmol/l or higher might prevent the loss of bone mineral [1, 3, 4], but confirm similar more recent observations [6, 7]. Thus, while the use of a dialysate calcium of 1.5 mmol/l or higher may
improve calcium balance and prevent hypocalcaemia [2, 5], the role of dialysate calcium in the pathogenesis of renal bone disease requires further study.

The significant increase in forearm calcium content in patients treated with 1α-OHD₃ compared to the untreated groups demonstrates the beneficial long-term effect of 1α-OHD₃ on bone calcium in haemodialysis patients. The rate of increase in bone calcium in patients treated with 1α-OHD₃ in this study was equivalent to 7.1% per annum over the whole period of observation. This compares with estimated increases of 3% per annum using whole body NAA over periods of 10–18 months [5] and 8.7% in 3 months using NAA measurements of the hand [15]. The increases in hand calcium by NAA in the short-term demonstrated by Catto et al. [15], however, did not continue in the long term [16; Catto pers. commun.]. While the reason for these differences in results of NAA using these two part-body sites is not clear, they may reflect regional differences in the skeletal response to 1α-OHD₃.

The increase in bone calcium in this study in response to 1α-OHD₃ occurred despite the use of a dialysate calcium concentration of 1.375 mmol/l at which calcium loss into the dialysate fluid might be expected. The 'low' dialysate calcium was chosen in this study in the light of previous experience when the use of a dialysate calcium of 1.75 mmol/l restricted the dosage of 1α-OHD₃ due to hypercalcaemia [17]. Since calcium absorption improves with 1α-OHD₃, it may be unnecessary, indeed undesirable, to use a 'high' dialysate calcium during treatment with 1α-OHD₃, particularly in view of the uncertainty about the influence of dialysate calcium on bone disease.

This study indicates that part-body in vivo NAA is a useful means of monitoring renal osteodystrophy. The hand is commonly chosen for NAA measurements on renal patients [18–20] but since NAA measurements of the hand have failed to demonstrate a long-term beneficial effect of 1α-OHD₃ on bone calcium, it is suggested that the forearm, which in any event contains more calcium, is more reliable and should be regarded as a more suitable site for monitoring renal bone disease using part-body NAA.

Acknowledgements

The authors would like to thank the following: Prof. J. S. Robson for very valuable initial discussion, Mr. W. R. McBeath and Mr. W. R. Harvey for technical assistance, Mrs. P. Williams for performing some of the NAA measurements, Mrs. M. Catling for computer analysis of much of the data, Mrs. P. Holis, Mrs. M. Proven and Mrs. H. D. McInnes for secretarial services, and the Scottish Home and Health Department for funding the project.

References


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Total body neutron activation analysis of calcium: calibration and normalisation

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Abstract. An irradiation system has been designed, using a neutron beam from a cyclotron, which optimises the uniformity of activation of calcium. Induced activity is measured in a scanning, shadow-shield whole-body counter. Calibration has been effected and reproducibility assessed with three different types of phantom. Corrections were derived for variations in body height, depth and fat thickness. The coefficient of variation for repeated measurements of an anthropomorphic phantom was 1.8% for an absorbed dose equivalent of 13 mSv (1.3 rem). Measurements of total body calcium in 40 normal adults were used to derive normalisation factors which predict the normal calcium in a subject of given size and age. The coefficient of variation of normalised calcium was 6.2% in men and 6.6% in women, with the demonstration of an annual loss of 1.5% after the menopause. The narrow range should make single measurements useful for diagnostic purposes.

1. Introduction

Total body neutron activation analysis is now established as one of the most important methods of determining the elemental composition of the human body. It can be used to determine both the absolute amount and changes with time of certain body elements, in particular calcium, sodium, chlorine, phosphorus and nitrogen (Cohn 1981).

In the technique, the subjects are exposed to a beam of partially moderated fast neutrons which induces amongst others the reactions $^{48}\text{Ca}(n, \gamma)^{49}\text{Ca}$, $^{23}\text{Na}(n, \gamma)^{24}\text{Na}$, $^{37}\text{Cl}(n, \gamma)^{38}\text{Cl}$, $^{31}\text{P}(n, \alpha)^{28}\text{Al}$ and $^{14}\text{N}(n, 2n)^{13}\text{N}$. The radioactive isotopes produced then decay, emitting $\gamma$-rays which can be readily detected in a whole-body counter.

If the reproducibility of the method were good then small changes in the body elements with time might be detected relatively easily. Absolute quantification of the elements is more difficult, because of the natural variation in body dimensions and the difficulty of obtaining a suitable variable phantom of known composition for calibration. To overcome these problems several different approaches have been made. Nelp et al (1972) made repeated measurements on five cadavers of different sizes and then ashed the skeletons to determine the calcium content. Cohn and Dombrowski (1971) and Cohn et al (1972) used only one size of phantom for activation and depended on the invariant response of a 54-detector whole-body counter (Cohn et al 1969). Three sizes of phantom were used by Williams et al (1978); their patients
were then graded into three groups according to body habitus. Two different sizes of phantom were used by Spinks et al. (1977) and the effects of varying thickness or adding wax around these phantoms were studied by Spinks (1979). The effect of varying the thickness of the trunk and thighs was also studied by McNeil et al. (1974). Oxby and Brooks (1979) have tackled the problem of varying body dimensions by designing a complex phantom of variable shape and content.

This paper describes the technique used in Edinburgh to measure total body calcium (TBCa) with a high degree of precision and also our methods of overcoming the problems mentioned above. The technique of evaluating whether a subject contains a normal amount of calcium is in two parts. Firstly, corrections to eliminate the effects of variations in activation and detection efficiency due to varying body dimensions are made. This enables the absolute amount of calcium in the subject in grams to be evaluated and requires a variety of phantom measurements. Secondly, the normal predicted total body calcium is calculated using a formula derived from the results of a group of healthy normal volunteers aged 40–70 years. These results were used to obtain a range of normal values of the body elements of interest.

Total body calcium measurements of normal volunteers have also been made by Nelp et al. (1972), Cohn et al. (1976), Aloia et al. (1978), and Chesnut et al. (1981). Where appropriate, the results of these studies and the data presented in this paper will be compared and discussed.

2. Method

2.1. Neutron activation

The neutron source used for patient activation was a cyclotron producing neutrons of mean energy 6.5 MeV by the reaction of 15 MeV deuterons on a beryllium target (Williams et al. 1979). During irradiation the subject was positioned in a moderator kiosk made from 3 cm thick polyethylene sheets mounted on a turntable. The thickness of the premoderator was chosen, after preliminary measurements, in order to minimise the variation in the thermal neutron flux through the patient. This was measured using a $^{235}$U fission chamber with and without a cadmium shield; in a phantom 24 cm deep...
Total body neutron activation analysis of calcium

Figure 2. The irradiation enclosure. The rear wall is brought forward until the subject's trunk is in contact with both the front and rear walls. The depth of the perspex window was chosen so that the effect on the thermal neutron profile was equal to that of the 3 cm thick polyethylene walls.

deep this variation was ±7% (figure 1) and in a limb phantom 12.5 cm in diameter the variation was ±13%. The back wall of the kiosk was movable and was adjusted so that the subject's trunk was in contact with both the front and back walls (figure 2). This contact kept the subject in the correct position and gave extra support. The fixed neutron dose was given to the patient in two successive irradiations each lasting about 22 s, with the kiosk rotating through 180° in 18 s after the first irradiation and before the second. These times were checked with a stopwatch and a decay correction applied if necessary. The target to subject distance was 5.4 m.

Using a Geiger–Müller tube sheathed in 3 mm of 6LiF, the photon absorbed dose was measured as 0.4 mGy. The total dose, measured at the position of the pelvis using a tissue-equivalent ionisation chamber, was 1.68 mGy, and was nearly constant with depth through the patient. Using a neutron quality factor of 10 therefore, the total dose equivalent received by the patient was 13.2 mSv. This dose was controlled by the neutron monitor chamber of the beam and the irradiation calibrated by activating simultaneously a container of Na2CO3 in a polyethylene jacket mounted on the side of the moderator kiosk. Possible long term variations in the neutron beam distribution were monitored by weekly measurements of the Bush phantom.

2.2. Measurement of induced activity

After irradiation, the patient was transferred to a shadow-shield whole-body counter containing four 15 cm diameter x 10 cm thick sodium iodide detectors, two above and two below the bed, arranged in line across the patient. The separation of the upper and lower counters was 40 cm. Four 200 cm scans at constant speed, each lasting five minutes, were then performed. The scan direction was simply reversed at the end of each scan. The time from the end of the irradiation to the start of the first scan was measured and was usually about 6 min.
The patient spectrum, covering the energy range 0–4 MeV, was recorded on a multichannel analyser. A twenty minute background count, measured using an inactive phantom, was then subtracted from the patient spectrum and the difference was punched out on paper tape for computer analysis.

2.3. Spectral analysis

The data were analysed by fitting five standard spectra, those of calcium, sodium, chlorine, phosphorus and potassium to the smoothed subject spectrum in the energy range 1.0–3.4 MeV, using a least squares technique (Smith et al 1976, Smith and Tothill 1979). The standard spectra were obtained by separately filling a phantom of human dimensions with appropriate solutions, activating the phantom and then recording the spectra on the whole-body counter. When the best possible fit to the subject's spectrum had been obtained, the amount of each standard required to make up the spectrum was listed. These amounts were then divided by the count from a twenty minute scan of the Na₂CO₃ standard. The values for each element were then corrected for radioactive decay.

In addition to the major elements of interest, some nuclides are activated which interfere with the measurement of the elements under analysis. The main interfering reaction in the measurement of calcium is the \(^{37}\text{Cl}(n,p)^{37}\text{S}\) reaction, since the \(^{37}\text{S}\) produced has a 5.06 min half-life and a gamma-ray energy of 3.1 MeV. The correction applied to the calcium content, obtained by activating chlorine in a water-filled phantom was 3.1% of the chlorine count corrected back to the reference time. By similar methods, the reaction \(^{39}\text{K}(n, 2n)^{38}\text{K}^\ast\) was found to contribute 0.9% of the total counts in the chlorine region (2.00 to 2.32 MeV) and the \(^{24}\text{Mg}(n, p)^{24}\text{Na}\) reaction contributed 0.8% of the total counts measured in the sodium region (2.57 to 2.92 MeV) when the composition of reference man (IRCP 1975), was assumed. The interferences from the last two reactions were regarded as negligible.

3. Results

3.1. Reproducibility

The reproducibilities of the measurements of calcium, sodium, chlorine and phosphorus were investigated by repeatedly measuring a phantom which consisted of hollow polyethylene sections containing known amounts of the elements of interest (Bush 1946). The water-filled weight of the phantom was 68.5 kg and its height 172 cm. Three separate filling mixtures were used (see table 1) and these mixtures were irradiated 3, 3 and 7 times respectively. In the final mixture, the total amounts

<table>
<thead>
<tr>
<th>Element</th>
<th>Phantom composition</th>
<th>Reference man (IRCP 23)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium</td>
<td>600 802 1009</td>
<td>1000</td>
</tr>
<tr>
<td>Sodium</td>
<td>60 70 84 100</td>
<td>100</td>
</tr>
<tr>
<td>Chlorine</td>
<td>50 61 70</td>
<td>95</td>
</tr>
<tr>
<td>Phosphorus</td>
<td>455 455 455</td>
<td>780</td>
</tr>
<tr>
<td>Potassium</td>
<td>120 120 130</td>
<td>140</td>
</tr>
</tbody>
</table>

Table 1. Contents of phantoms (in g).
of the various elements were similar to those in reference man (IRCP 1975) with the exception of phosphorus where more recent results imply that the quoted content is too high. There was no significant dependence of the counts obtained per gram of element on the ratio of the various elements in the phantom. The reproducibilities (coefficients of variation (CV)) of the measurements of the various elements, obtained from the 13 irradiations of the phantom, using the same dose as given to the patient, were ±1.7% for calcium, ±2.2% for sodium, ±4.3% for chlorine and ±4.3% for phosphorus. The values of the CV for calcium, sodium and chlorine reflect almost entirely the counting statistics for these elements. For a small female with only 500 g of calcium the CV for calcium would increase to about 3.0%. The reproducibility of the measurement of calcium was also investigated by repeatedly measuring a phantom consisting of a human skeleton with approximately 700 g calcium encapsulated in tissue-equivalent rubber. The reproducibility in this case, determined from 10 measurements, was 1.8% for calcium and 2.1% for sodium.

3.2. Measurement corrections

In this section the corrections required to allow for the influence of body habitus on the combined activation and counting efficiency are determined. These corrections are essential for accurately calculating the absolute amount of calcium in the subject.

3.2.1. Effect of height. The thermal neutron fluence at a depth of 5 cm, over an area 185 cm high and 60 cm wide at the subject irradiation position, was uniform to within ±8% with the maximum incident neutron flux around 85 cm above the chamber floor. The effect of varying height was tested by altering the height of the water filled phantom over the range 150 cm to 193 cm by inserting wax spacers between the various sections. The phantom was measured twice at each of five different positions. The range 150 cm to 172 cm was investigated by removing the pelvis section. The results obtained were fitted by the power law:

\[
\% \text{ loss of counts} = 0.5 \times (\text{subject height} - 1.5)^2 \times 100 \quad \text{height} > 1.5 \text{ m.}
\]

3.2.2. Effect of body build and body fat. The effect of variations in thickness of the subject, measured along the beam direction, was investigated using sections of the solution-filled phantoms of different sizes, but similar heights. Two repeated measurements were made at each of five different thicknesses i.e. 11, 14, 16, 18 and 24 cm. The decrease in measurement efficiency was 1.8% per cm increase in thickness of the chest section and 1.4% per cm increase in diameter of the limb section.

3.2.3. Effect of wall separation. Ideally the front and back premoderator sections of the activation chamber are positioned just to touch the subject's trunk at the front and back. The separation of the walls is dependent, therefore, on the subject's depth measured along the beam direction (figure 2). Due to patient apprehension in some cases this is not possible and the walls are positioned further back than normal. The effect of this extra separation was investigated by repeatedly measuring both the solution-filled and the skeletal phantoms, increasing the wall separation each time up to an air gap of 10 cm. Two repeated measurements were made at each of three different positions. The decrease in activation efficiency was 1.0% per cm increase in the air gap.
3.2.4. Absolute calibration of calcium. Using the results from sections 3.2.1 to 3.2.3, the mass of calcium in the subject in grams was calculated from the equation:

\[ TBCa = \text{measured calcium} \times \text{height correction} \times \text{depth correction} \times \text{fat correction} \times \text{wall correction} \]

where:

- height correction = 0.5 \((\text{subject height} - 1.50)^2 + 1\) \text{ height} > 1.5 m
- depth correction = 0.6 \((\text{Subject depth} - \text{TST} - 0.2) + 1\)
- fat correction = \(3.6 \left( \frac{\text{TST} + \text{LST}}{2} \right) + 1\)
- wall correction = 1.0 \((\text{wall separation} - \text{subject depth}) + 1\).

All measurements are in metres and \(\text{TST} = \text{mean torso skinfold thickness of subject, i.e. the mean of the subscapula and iliac crest measurements}; \text{LST} = \text{mean limb skinfold thickness of subject, i.e. the mean of the biceps and triceps measurement} \); 1.6 = mean fractional loss per m increase in depth; 3.6 = mean fractional loss per m increase in fat; 0.2 = reference depth, i.e. the depth of the trunk section of the solution filled phantom. The depth correction makes the assumption that the depths of the various limbs scale in proportion to the depth of the trunk.

The number of counts obtained per gram of element was checked by repeatedly activating three phantoms of different sizes similar to the Bush phantom but with the calcium and phosphorus contained in solid rods which simulated the human skeleton (Batty et al 1971). The results obtained for sodium and chlorine agreed within experimental error. However the calcium was on average 3.1% low and the phosphorus result 11.2% low compared with elements dispersed in solution. This effect might well be explained by the difference in the combined efficiency of activating and detecting the calcium and phosphorus when they are uniformly distributed in the phantom compared with when they are contained in a simulated skeleton. This effect has also been noted by Spinks et al (1977). The necessary change was therefore made to the conversion factor for calcium to give counts per gram of the element. The difference between the results of calcium and phosphorus was due to the fact that the latter undergoes a fast neutron reaction.

4. Normalisation procedures

To establish the range of normal values for total body calcium, 40 normal volunteers were measured: twenty males (six aged 40-49, eight aged 50-59 and six aged 60-69) and twenty females (two aged 40-49, ten aged 50-59 and eight aged 60-69). All the subjects were active and in good health. None had any history of disease that might be expected to influence calcium metabolism, and all had normal blood and urine biochemistry. There was a large variation in the total body calcium results for both groups (table 2). The females ranged from 596 to 1036 g with a mean of 821 ± 15.2% (CV), while the males ranged from 886 to 1428 g with a mean of 1143 ± 11.7%. Since the total body composition of the various elements is clearly related to the subject's build, a mean value based on sex alone is inadequate. Therefore the dependence of the measured \(TBCa\) on height, span, weight, lean body mass (calculated from the
skinfold thickness measurement), age and number of years post-menopause was investigated by multiple linear regression analysis.

For males, this gave a formula for predicted total body calcium in grams (TbCa$_p$),

$$\text{TbCa}_p = 347 \text{ (height)}^{2.07} \quad (r = 0.84, P < 0.001).$$

Total body calcium was also found to be strongly dependent on span

$$\text{TbCa}_p = 384 \text{ (span)}^{1.80} \quad (r = 0.74, P < 0.001).$$

Height and span are measured in metres. Perhaps because the spread in age was small no significant dependence of total body calcium on age was observed. Weight and lean body mass also proved to be unimportant. The biological variation in normal TbCa due to size could be reduced by expressing the subjects result in terms of the ratio TbCa/TbCa$_p$. When height was used in the formula the cv in the male controls was reduced to 6.2%.

For the female controls, multiple regression gave

$$\text{TbCa}_p = 399 \text{ (span)}^{1.60} \exp(-0.015 Y) \quad (r = 0.90, P < 0.001)$$

where span was measured in metres and $Y$ is the subject's years post-menopause. This gave a highly significant correlation with an annual loss of calcium after the

---

**Figure 3.** Total body calcium of normal women, normalised using the span (s), plotted against years post-menopause. The line representing a loss of 1.5% per annum is plotted with confidence and tolerance limits. $\alpha$ is a constant = 399; $\bullet$, normal women; ---, confidence limits, $P = 0.05$; ---, tolerance limits, $P = 0.10$. 

---

**Table 2.** Data for normal controls.

<table>
<thead>
<tr>
<th></th>
<th>Male (cv)</th>
<th>Female (cv)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>Height (m)</td>
<td>1.770 (4.7)</td>
<td>1.622 (4.2)</td>
</tr>
<tr>
<td>Span (m)</td>
<td>1.843 (4.7)</td>
<td>1.655 (4.4)</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>81.8 (13.8)</td>
<td>63.7 (17.2)</td>
</tr>
<tr>
<td>Calcium (g)</td>
<td>1143 (11.7)</td>
<td>821 (15.1)</td>
</tr>
</tbody>
</table>
menopause of 1.5% (figure 3). An equally good correlation was obtained if a linear relationship between calcium and years post-menopause was assumed. Total body calcium, for the female controls, was also found to be strongly dependent on height

\[ \text{TB}Ca_p = 443 \times \text{height}^{1.57} \times \exp(0.016 \times Y) \quad (r = 0.87, P < 0.001). \]

Since total body calcium was more strongly correlated with span than with height and as height may be significantly reduced in patients with osteoporosis, it was decided to use span rather than height as a variable for normalisation for women. The coefficient of variation of total body calcium in normal women \( \frac{\text{TB}Ca}{\text{TB}Ca_p} \), after normalisation improved to 6.6%.

The possibility was investigated that the correction factors, applied to the raw data, described in section 3.2.4 could be replaced with an extra normalisation factor, namely weight. Regression analysis performed on the uncorrected data for the male controls gave a low correlation with height, \( r = 0.40 \), compared to the corrected data, \( r = 0.84 \). The introduction of weight as a normalisation factor improved the correlation considerably

\[ \text{TB}Ca_p = C_M \times \text{height}^{2.15} \times \text{weight}^{-0.84} \quad (r = 0.80) \]

where the superscript indicates that this equation refers to the raw data, uncorrected for the height, build and body fat corrections derived in section 3.2.4.

A similar effect was obtained, using the uncorrected female data, after the introduction of a normalisation factor for weight

\[ \text{TB}Ca_p = C_F \times \text{span}^{1.66} \times \text{weight}^{-0.18} \times \exp(-0.016 \times Y) \quad (r = 0.89) \]

where \( C_M \) and \( C_F \) are constants.

This contrasts with the case of the corrected data, both male and female, where weight proved to be unimportant. Therefore to a large extent the correction factors described in section 3.2.4 and the introduction of weight as a normalisation factor fulfil a similar function. The CV of the \( \frac{\text{TB}Ca}{\text{TB}Ca_p} \) ratio was 7.0% for both men and women. When the ratio is used, the units of \( \text{TB}Ca_p \) can be left as \(^{40}\text{Ca} \) counts. However, using the results from the calculations of \( \text{TB}Ca_p \), the values \( C_M = 1447 \) and \( C_F = 870 \) will give \( \text{TB}Ca_p \) in grams of calcium.

5. Discussion

The technique of total body neutron activation analysis presented in this paper with its precision of 1.8% on phantoms is suitable for detecting small changes in total body calcium. The process of correcting for varying body dimensions is most important for the accurate determination of the absolute amount of calcium in the body. When the corrections described in section 3.2.4 were applied to the raw data for our normal male volunteers they increased the estimated body calcium by 16% on average. The importance of such corrections has also been emphasised by Spinks (1979). Using a similar irradiation system, he found correction factors for total thickness and wax layer thickness which were close to ours. McNeill et al (1974) also studied the effect of body thickness on the efficiency of activation and measurements of calcium in the trunk. They found a reduction of calcium counts of 5% cm\(^{-1}\). Their figure was somewhat higher than ours, probably because their Pu-Be sources were a fixed distance apart and fairly close to the subject's skin. Correction for body stature is important not only for absolute calcium determinations, but also for studying changes over a
period if the subject is likely to alter in weight. Some treatments which lead to osteoarthritis, or which are designed to avoid the condition, are liable in themselves to change the amount of fat in the body.

The establishment of a normal range of total body calcium for men and women is vital if full use is to be made of the absolute values. Total body calcium measurements of normal volunteers have been made by Nelp et al. (1972), Cohn et al. (1976), Aloia et al. (1978) and Chesnut et al. (1981). Partial body (trunk) measurements have been made by Harrison et al. (1975) and by McNeill and Harrison (1977). The most comprehensive of these studies is that of Cohn et al. (1976). They derived equations for predicting normal values based on measurements on 79 normal individuals, 48 in the age range covered by this study. Their whole body counter corrects for variations in response due to different body dimensions, but no attempt was made to separate variations in activation efficiency from the biological variations of actual calcium content in the normal population studied.

The equations developed by Cohn et al. (1976), which incorporate height, age and total body potassium levels, therefore included a consideration of measurement errors, as well as statistical variations. The coefficient of variation of normalised total body calcium was 7.8% for men and 7.1% for women. As a result of the different approach to normalisation and because Cohn et al. (1976) assumed a uniform age of menopause at 55, our values of total body calcium are not directly comparable with theirs. As in this study, Cohn et al. (1976) found total body calcium in males to be dependent approximately on height squared. These results are in slight disagreement with the total body calcium study of Nelp et al. (1972) and the partial body studies (trunk) of Harrison et al. (1975) who found a near cubic relationship between calcium and height.

The percentage of calcium lost by normal women after the menopause of 1.5% per year from this study is slightly higher than that of 1.1% determined by Cohn et al. (1976). They also calculated a rate of calcium loss of 0.4% per annum between the ages of 35 and 55. This is probably the combination of a lower (if existent) pre-menopausal bone loss and a higher post-menopausal loss from those normal women who were post-menopausal before the age of 55.

Although multiple regression analysis on our male data gave a mean loss of calcium with age of 0.1% per year, this result was not significantly non-zero. Cohn et al. (1976) reported a mean annual loss of calcium for the average man of 0.70% after 50 years of age. However it is interesting to note that their result, which was obtained by omitting their data for men over 80, leads to such a low predicted value of calcium for older men that the majority of their own male subjects aged 67 and over have TBCa/TBCap values substantially greater than one.

In section 4 it was shown that the introduction of weight as a normalisation factor applied to the raw data, rather than using the correction factors described in section 3.2.4, implied that total body calcium was dependent partly on (weight)$^{-0.34}$ for males and (weight)$^{-0.18}$ for females. These values are in broad agreement with the exponent determined by Cohn et al. (1976) of $-0.25$ (see appendix Cohn et al. 1976, substituting $K_a$ for $K$). We obtained no significant dependence on weight after multiple regression analysis was performed on the final data because of the correction factors used in section 3.2.4. For the determination of a narrow normal range of calcium values the use of weight as an extra normalisation factor would appear to be almost as good, and would be simpler than determining and applying the various correction factors for height, body fat and thickness. The success of weight as a normalisation factor presumably depends on its close correlation with body thickness. However the body
correction factors described in section 3.2.4 are important for the accurate determination of the absolute amount of calcium in grams in individuals whose size or shape differs, even by only a few centimetres, from the phantom used for calibration. The values of the correction factors are only applicable to the experimental arrangement used in this study. However if other research groups were able to calculate accurately the absolute amount of calcium in grams in their patients then the equations for \( \text{TBCa}_n \) derived from our normal controls could be used to calculate the \( \text{TBCa}/\text{TBCa}_p \) ratio.

We believe that the best approach is the one we have adopted, of first trying to establish an accurate measure of whole body calcium, making correction for all the variables likely to affect activation and detection efficiency, and then deriving normalisation factors which take into account variations of calcium with size, age and sex. The narrow ranges for normal subjects that we have observed, with coefficients of variation of 6.2% for men and 6.6% for women, may allow the technique of total body neutron activation analysis of calcium to be a useful diagnostic procedure.

Acknowledgments

We wish to express our gratitude to the staff of the MRC Cyclotron Unit at Edinburgh, in particular Mr T Saxton and Mr J Williams, for their considerable cooperation and assistance, Mr L Mackie and Mr H Easton for skillful workshop services and Mrs Elisabeth Law for technical assistance. We should also like to thank Mr T J Spinks and Professor K Boddy for allowing us to use their phantoms.

This work was supported by grants from the Scottish Home and Health Department and from the Medical Research Council.

Résumé

Analyse par activation neutronique du calcium de corps entier: calibration et normalisation.

Nous avons réalisé un système d'irradiation, utilisant un faisceau de neutrons émis par un cyclotron, qui optimise l'uniformité de l'activation du calcium. L'activité induite est mesurée par un compteur corps entier à balayage 'shadow shield'. Nous avons calibré les mesures et testé leur reproductibilité à partir de trois types différents de fantômes. Nous en avons déduit des corrections pour les variations de la taille du corps, de la profondeur et de l'épaisseur de la graisse. Le coefficient de variation pour les mesures répétées sur un fantôme anthropomorphe est de 1.8% pour une dose absorbée équivalente à 13 mSv (1.3 rem). Nous avons utilisé les mesures du calcium global du corps chez 40 adultes normaux pour déduire les facteurs de normalisation qui prédissent le taux normal du calcium chez un sujet d'âge et de taille donnés. Le coefficient de variation du calcium normalisé est de 6.2% chez l'homme et de 6.6% chez la femme avec une perte annuelle de 1.5% après la ménopause. Cette faible dispersion des valeurs rendrait utile une seule mesure à des fins diagnostiques.

Zusammenfassung

Ganzkörper-Neutronenaktivierungsanalyse von Kalzium: Kalibrierung und Normierung.

Total body neutron activation analysis of calcium

abzuleiten, die den normalen Kalziumgehalt eines Menschen bei vorgegebener Größe und Alter vorhersagen können. Der Variationskoeffizient von normiertem Kalzium variiert 6.2% bei Männern und 6.6% bei Frauen, mit einem jährlichen Verlust von 1.5% nach der Menopause. Der eingeschränkte Bereich könnte einzelne Messungen für diagnostische Zwecke brauchbar machen.

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Scientific Note

The crossover correction in dual photon absorptiometry with $^{153}\text{Gd}$

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1. Introduction

The measurement of the bone mineral content (BMC) by single photon absorptiometry has long been an established technique for the forearm. However, in the study of bone disease, the part-body site of most clinical interest is the lumbar spine. Recently, efforts have been made to measure the BMC using dual photon absorptiometry (DPA), the second photon energy being used essentially to correct for varying amounts of soft tissue within the trunk. The energy combinations that have been used are $^{241}\text{Am}$ and $^{137}\text{Cs}$ with energies at 60 keV and 660 keV and $^{153}\text{Gd}$ with energies at 44 keV and 100 keV, the latter being generally considered the most suitable energy combination for measurements of the spine.

Although reasonable figures of 2–3% for the precision (or reproducibility) have been obtained by the various methods, there is a degree of disagreement about the absolute measurements of BMC. This is particularly the case with cross-sectional data on normal volunteers, i.e. single BMC measurements on subjects of different ages. Two groups (Madsen 1977, Riggs et al 1981) have reported no increase in the rate of loss of BMC from the spine at the time of the menopause. The expected increase in the rate of loss after the menopause, as shown using total body neutron activation analysis (Cohn et al 1976, Smith et al 1980) and forearm absorptiometry (Mazess and Cameron 1974) has, however, been shown in the spine by two other groups (Krohner and Pors Nielsen 1980, Smith et al 1981) in cross-sectional studies and also in a longitudinal study (Krohner and Pors Nielsen 1981), where the bone loss was determined from repeated measurements on individual subjects over a period of time. It is not known whether this discrepancy between the various methods is due to differences in methodology or in the populations studied. Before accepting the latter the possibility of the former must be thoroughly investigated.

All but two of the methods of DPA (Roos 1974, Condon et al 1979) which have been used for patient or normal volunteer studies have used $^{153}\text{Gd}$ as the radionuclide source. Obviously there are several factors that influence the accuracy of the technique. This note considers one aspect of the methodology of DPA which is specifically important when using $^{153}\text{Gd}$ and which will affect the absolute determination of BMC. This is the contribution to the 44 keV energy window from the 100 keV energy caused by Compton events within the crystal and scatter within the patients. This will be referred to as the ‘crossover correction’ ($x_{0}$) and is expressed as a percentage of the 100 keV photopeak.
2. Gadolinium-153

The radionuclide source $^{153}\text{Gd}$ has a half-life of 242 days with gamma-ray energies (plus approximate abundance) of 103 keV (23%), 97 keV (31%) and 70 keV (2.6%) and Eu K x-rays with energies 41 keV (86%) and 47 keV (23%). The peaks at 103 keV and 97 keV and at 41 keV and 47 keV cannot be resolved by NaI detectors and so are considered as 100 keV and 44 keV peaks respectively. The ratio of the 44 keV peak to the 100 keV peak is approximately 2:1 though this falls to about 1.4:1 from an encapsulated sealed source measured by a Ge(Li) detector, due mainly to the stainless steel window specified as 0.2–0.25 mm for our source, and to 1.1:1 from the same source measured by a 10 cm diameter, 5 cm thick NaI detector, additional attenuation arising in the crystal housing. The measurements described in this paper were performed using a 12 mm diameter sealed $^{153}\text{Gd}$ source (ref no X.92/1, Amersham 1978) of approximate activity 4 GBq, spectra of which are shown in figure 1.

![Figure 1. $^{153}\text{Gd}$ spectra measured by a NaI detector and by a GeLi detector.](image)

3. Bone mineral calculation

The basic equations for calculating the BMC from two photon energies are quoted elsewhere (Roos and Sköldborn 1974) but can be simplified to

$$\text{BMC} \propto \ln(I_{100}) - \frac{\mu_{ST,100}}{\mu_{ST,44}} \ln(I_{44})$$

where $I$ is the count rate of 100 keV or 44 keV radiation, $\mu_{ST}$ is the attenuation coefficients through soft tissue at 100 keV or 44 keV, and $I_{44}$ is (measured count rate in 44 keV region) – (xo/100) × $I_{100}$ with xo the crossover correction (per cent).

The two criteria for accurate measurements of BMC by the technique of DPA using the above equation are that (i) the BMC must be independent of the thickness of soft tissue and (ii) the measured BMC must be proportional to the actual BMC. The xo will affect both the estimate of $I_{44}$ and $\mu_{ST,44}$ in the above equation. The mean supine
anterior to posterior thickness (with range) at the lumbar region in 180 normal subjects and patients was found to be 22 cm (15-29 cm) in men and 19.8 cm (15-29 cm) in women. Criterion (i) must be verified over an appropriate range of soft tissue thicknesses. For criterion (ii) to be fulfilled the basic assumption that both 44 keV and 100 keV radiation have exponential attenuation in bone and soft tissue must be true. In particular the attenuation of the corrected 44 keV radiation through bone mineral or a similar material such as aluminium must be exponential.

4. Crossover correction

Experiments to investigate the xo were performed using a 13 mm diameter source collimator and a 20 mm square detector collimator with a separation between the two of 0.4 m. This collimation, which we have used for spine BMC measurements (Smith et al 1981), is slightly larger than that used by Wilson and Madsen (1977) and smaller than that used by Peppler and Mazess (1981). Some experiments were also performed using 6 mm diameter source and detector collimators, the size used by Riggs et al (1981) and smaller than that used by Krölner and Pors Nielsen (1980). The NaI detector used was 10 cm diameter, 5 cm thick, with an energy resolution of 20-25% over the energy range used.

The standard method adopted in the past to calculate the xo has been to use copper to filter out the 44 keV peak at the source collimator. This method was used with a 3.3 mm Cu filter and energy windows from 27-61 keV and 83-121 keV. Using the larger collimation the xo was found to vary with the thickness of soft tissue equivalent material (STEM) (figure 2) but the two criteria mentioned previously were not fulfilled (figure 3) due to an apparent underestimation of the xo. The possible reasons were investigated.
The theoretical attenuation was calculated assuming narrow beam attenuation (McMaster et al. 1969). The error due to background may be negligible when measuring the BMC, but becomes significant when calculating the xo values. More important than the room background were the Compton scatter contributions from $^{152}$Eu contamination in the $^{153}$Gd source. Lead 4 mm thick was used to filter out the $^{153}$Gd to assess the contribution from the higher energy peaks of $^{152}$Eu to the 44 keV and 100 keV channels. The variation in background, including $^{152}$Eu, with thickness of STEM was
measured and used to correct subsequent data. No corrections were ever required for counting losses.

The $^{153}$Gd spectrum with STEM and aluminium in place to simulate the transmission through the lumbar spine was compared with the spectrum of the same arrangement with the Cu filter, both spectra having the same number of counts in the 100 keV channel (figure 4). It can be seen that the filtered $^{153}$Gd spectrum appears to under estimate the contribution to the 44 keV channel. It seemed likely that the low abundance 70 keV peak in the $^{153}$Gd spectrum contributes appreciably to the lower energy channel, but is severely attenuated by the copper used to eliminate the 44 keV radiation.

The possibility was investigated by using monoenergetic X-rays to simulate the upper portion of the $^{153}$Gd spectrum. Between 4 and 70 GBq of $^{99}$Tc$^{m}$, absorbed onto ion exchange resin, was used to produce the characteristic 70 keV and 100 keV X-rays from mercury and uranium respectively (figure 5). The Hg and U were placed separately with the $^{99}$Tc$^{m}$ in the position of the $^{153}$Gd source so that the geometry remained the same. The counts in the 44 keV region were expressed as a percentage of the counts in the respective photopeaks. The proportion of the 70 keV peak compared with the 100 keV peak from the $^{153}$Gd source in air was 4.6% as measured by a Ge(Li) detector. This value combined with the relative attenuations of the 70 keV and 100 keV Hg and U peaks through STEM enabled the $\rho_0$ contribution from the 70 keV $^{153}$Gd peak to be expressed as a percentage of the 100 keV peak. The results are shown in table 1 and the total $\rho_0$ value plotted in figure 2.

It was found empirically that if the $\rho_0$ derived from the use of the copper filter was multiplied by the ratio of the portions of the two spectra in figure 4 between the peaks (60–85 keV) the two criteria required in the calculation of bone mineral were fulfilled. The values of the $\rho_0$ calculated in this way for the 13 mm source and 20 mm detector collimation are shown in figure 2 for various thicknesses of STEM.

It can be seen that both the new methods of calculating the $\rho_0$ give the same result (figure 2). When these $\rho_0$ values were used for measurements on 10 mm Al in depths of water ranging from 8 cm to 27 cm no variation in the apparent mineral
content was found. In addition the attenuation of the corrected counts in the 44 keV channel through Al in 20 cm scattering medium was exponential (figure 3).

5. Discussion

The importance of the correct calculation of the \( x_0 \) value for absolute measurements of spine BMC should not be underestimated. Although the actual percentage value may seem low, a \( x_0 \) value of only 3% would reduce the counts in the 44 keV region by 19% in a subject of average thickness (22 cm) and average bone density, and by 25% in a subject of thickness 27 cm (approximate normal mean \( \pm 2\sigma \)) and high bone density (approximate normal mean \( \pm 2\sigma \)). In a subject 22 cm thick the use of a \( x_0 \) value of 2.8%, obtained using the standard Cu filter technique, rather than the true value of 4.4%, will produce an error of 15% in the BMC measurement of a normal subject.

The \( x_0 \) values quoted vary considerably. Mazess et al (1974) quote 1.9%, Wilson and Madsen (1977) 3.0%, Wahner and Dunn (1980) 6.0% and Riggs et al (1981) 4–6% depending on the source. The commercial apparatus used by Krølner and Pors Nielsen (1980) has a \( x_0 \) of ‘about 1%’ (personal communication). Only Peppler and Mazess (1980) mention that \( x_0 \) varies with depth, 1.8% at 14 cm and 3.4% at 25 cm; Wilson and Madsen (1976) specifically state that it does not. Although, as we have shown, the \( x_0 \) will depend on the size of the beam, the variation in \( x_0 \) values quoted by different authors cannot be attributed to variations in collimator size and should not depend on the source. It has been suggested that ‘beam hardening’ could account for any deviation from an exponential by the 44 keV radiation (Krølner and Pors Nielsen 1980). We would suggest that an underestimation of the \( x_0 \) value could be the true cause. Similarly the non-linear relationship between measured BMC and actual BMC which sometimes has to be corrected for (Krølner and Pors Nielsen 1980) is due to the same effect.

The underestimation of the \( x_0 \) is due to the significant contribution from the small 70 keV \(^{153}\)Gd peak which may be completely filtered out by Cu. This photopeak will produce an iodine escape peak at 40 keV. Thus, for example, in a 20 cm thick patient 70% of the total \( x_0 \) is due to the 100 keV peak and 30% due to the 70 keV peak.

The \( x_0 \) has been shown to vary with the thickness of the patient though when small collimation is used this variation is small. If, as is often the case, larger collimators need to be used to achieve acceptable count rates then a \( x_0 \) value appropriate for the patient’s thickness must be used. The previous standard method of using Cu to filter the lower energy channel has been shown to be inadequate. It is suggested that

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**Table 1.** \( x_0 \) values for 44 keV region using Hg and U to simulate the upper part of the \(^{153}\)Gd spectrum with 13 mm source and 20 mm detector collimators.

<table>
<thead>
<tr>
<th>Thickness of STEM (cm)</th>
<th>0</th>
<th>5</th>
<th>10</th>
<th>15</th>
<th>20</th>
<th>25</th>
</tr>
</thead>
<tbody>
<tr>
<td>Using U, Percentage of 100 keV photopeak</td>
<td>2.0</td>
<td>1.9</td>
<td>2.3</td>
<td>2.5</td>
<td>3.1</td>
<td>3.4</td>
</tr>
<tr>
<td>Using Hg, Percentage of 70 keV photopeak</td>
<td>13.2</td>
<td>13.5</td>
<td>16.6</td>
<td>20.6</td>
<td>25.0</td>
<td>27.2</td>
</tr>
<tr>
<td>Combined U and Hg to simulate (^{153})Gd</td>
<td>2.5</td>
<td>2.5</td>
<td>3.1</td>
<td>3.5</td>
<td>4.3</td>
<td>4.7</td>
</tr>
<tr>
<td>Percentage of 100 keV photopeak</td>
<td>70%</td>
<td>70%</td>
<td>70%</td>
<td>70%</td>
<td>70%</td>
<td>70%</td>
</tr>
</tbody>
</table>
the true xo should be calculated by simulating the 100 keV and 70 keV peaks of the \textsuperscript{153}Gd spectrum over a range of stem from 12 cm to 28 cm.

Acknowledgments

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Bone demineralisation in patients with Turner's syndrome

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SUMMARY The hypothesis that the demineralisation associated with gonadal dysgenesis is analogous to post-menopausal osteoporosis was investigated. Bone mineral content of the distal forearm was measured in 11 adult patients with Turner's syndrome aged 18 to 57 years. As a group these patients were significantly demineralised (p<0.001) when compared with normal subjects. A bimodal distribution of bone mineral was demonstrated, the eight patients below the normal range having a bone mineral content 75% of normal. This may be the usual bone mineral content for a large proportion of Turner's patients. No steady reduction in mineralisation with age was demonstrated. The number of osteoporotic type fractures was obtained from the records of 36 adult patients with Turner's syndrome. From the cumulative total years at risk (770 patient years) from the age of 15 years, it was found that the number of fractures of the distal radius corresponded to the normal pre-menopausal rather than post-menopausal fracture incidence. The absence of any reduction in bone mineral content with age and no clear evidence of an increase in frequency of fractures both suggest that the demineralisation associated with Turner's syndrome is not analogous to post-menopausal osteoporosis. The regular use of long term oestrogen therapy as a treatment for 'osteoporosis' in these patients is therefore not justified.

Qualitative assessment of skeletal radiographs from patients with Turner's syndrome has demonstrated that in this condition osteoporosis occurs with a frequency of between 60 and 80%.1-4 In affected children bone histology has shown increased bone resorption with normal bone formation,5 while radiographs of the second metacarpals show reduced total diameters and increased medullary diameters.2 6 The degree of demineralisation owing to osteoporosis can be assessed quantitatively using the technique of photon absorptiometry. The linear density of bone is obtained from the degree of attenuation of radiation from a monoenergetic radionuclide source.7 This technique has been used to demonstrate demineralisation of the radius and os calcis in children and young adults.5 8 Oestrogen inhibits bone resorption in osteoporosis and has been advocated for the treatment of post-menopausal osteoporosis.9 The rationale behind this treatment is the prevention of the normal loss of bone mineral of about 1% per annum,10 thus reducing the likelihood of debilitating fractures in later life. There are, however, disadvantages since there is evidence that when it is terminated the mineral content falls to the level at which it would have been had no treatment been given.12 Thus, to achieve a long term benefit, oestrogen therapy may have to be continuous. Other disadvantages include associated side effects such as increased incidences of abnormal uterine bleeding and endometrial carcinoma.13-16 The treatment of osteoporosis in patients with Turner's syndrome might depend on whether osteoporosis associated with gonadal dysgenesis is analogous to post-menopausal osteoporosis in normal women. If it is, there should be a steady decrease in bone mineral after the age of about 15 years combined with a raised incidence of fractures. In these circumstances long term oestrogen therapy would be indicated. The present study was undertaken to investigate the bone mineral content (BMC) in adult patients with Turner's syndrome, and to assess the severity and age relationship of any deficit in bone mineralisation. The incidence of osteoporotic fractures has also been examined and compared with the incidence in the general population.
Bone demineralisation in patients with Turner's syndrome

Patients and methods

The bone mineral content was studied in 11 adult patients with Turner's syndrome over a wide age range (18 to 57 years). These were available patients with Turner's syndrome in the south-east of Scotland who were over the age of 18 years and who had been registered with the MRC Cytogenetics Unit. They had all given informed consent to the investigation. Their mean height was 140.8 cm. Other details of the patients are given in Table 1. For comparison we studied a group of 16 healthy premenopausal women of presumed normal karyotype aged 18 to 44 years. Within the group of 16 there was a subgroup of 11 short women with a mean height of 154.0 cm whose BMC measurements were used to establish the normal range. The remaining five were significantly taller with a mean height of 169.1 cm, and their BMC measurements were used to determine the effect of a height difference of 15 cm.

The BMC of the non-dominant forearm was measured by photon absorptiometry using a commercial 'Novo Osteodensitometer' which had a reproducibility of 1.4%. The source and detector scan the forearm in a rectilinear motion from the distal end. The BMC is measured for each traverse, but is only stored once the gap between the radius and ulna exceeds 6 mm. A further five scans, 4 mm apart, are performed and the total BMC of the radius and ulna are recorded. The width of the radius and the ulna on the third and sixth traverse were measured from a chart recorder output and their sum total (W) used as a normalisation factor.

The medical records of the 11 patients with Turner's syndrome and of 25 others with the same diagnosis were examined. Their mean age was 36.4±8.4 years (±SD). The number of fractures since the age of 15 years of the type associated with osteoporosis, that is, of the distal radius and of the neck of the femur, was noted and compared with figures for a normal Scottish population. The cumulative total years at risk in the group of 36 patients with Turner's syndrome was 770 patient years.

Results

The results of BMC measurements are displayed in Fig 1. For three patients with Turner's syndrome they were within the normal range and for eight they were significantly low (five patients p<0.05, three patients p<0.01). The mean for the group of patients with Turner's syndrome was 2.31±0.44 (arbitrary units±SD) which is significantly lower (p<0.01) than for the group of normal subjects, whose mean was 2.85±0.27. The results of the group of taller normals (mean BMC = 2.94) demonstrated that a height difference of 15 cm did not significantly affect the normalised BMC values. It can be seen in Fig 1 that the eight patients with values outside the normal range are perhaps a subgroup in themselves with a mean value of 2.07±0.15 which is 73% of normal. Indeed, the three patients (CM, SG, MS) in the normal range were found to be biochemically hypothyroid, a condition which may be expected to increase bone mineral. One of these three patients (SG aged 36) was still menstruating spontaneously.

The values of BMC are plotted against age in

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### Table 1 Details of patients with Turner's syndrome on whom the BMC of the distal forearm was measured

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age</th>
<th>Karyotype</th>
<th>Oestrogen therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>ES</td>
<td>18</td>
<td>45.X/46,X(i(Xq))</td>
<td>1978</td>
</tr>
<tr>
<td>JC</td>
<td>26</td>
<td>45.X/46.XY</td>
<td>Never</td>
</tr>
<tr>
<td>CM</td>
<td>26</td>
<td>45.X</td>
<td>Never</td>
</tr>
<tr>
<td>MV</td>
<td>27</td>
<td>45.X</td>
<td>1972–1973</td>
</tr>
<tr>
<td>IC</td>
<td>35</td>
<td>45.X</td>
<td>1965–1977</td>
</tr>
<tr>
<td>SG</td>
<td>36</td>
<td>45.X/46.XY</td>
<td>1964–1965</td>
</tr>
<tr>
<td>HB</td>
<td>39</td>
<td>46.X(i(Xq))</td>
<td>1974</td>
</tr>
<tr>
<td>MS</td>
<td>42</td>
<td>45.X</td>
<td>1956–1967</td>
</tr>
<tr>
<td>FS</td>
<td>47</td>
<td>45.X</td>
<td>1956–1964</td>
</tr>
<tr>
<td>ML</td>
<td>45</td>
<td>45.X</td>
<td>Irregularly until 1977</td>
</tr>
<tr>
<td>HS</td>
<td>51</td>
<td>45.X</td>
<td>Never</td>
</tr>
</tbody>
</table>

---

**Fig 1** BMC values in patients with Turner's syndrome compared with the normal range. Note the narrow spread in BMC values of patients below the normal range.
FIG 2. BMC values in patients with Turner's syndrome plotted against age. The normal rate of loss in post-menopausal women is shown.

There were only two osteoporotic type fractures in the group of 36 Turner patients. One patient had fractured the neck of the femur aged 32 years, and the other the distal radius aged 35 years. The incidence of fractures of the distal radius in the normal population is 0.55% per annum and 0.15% per annum for post-menopausal and premenopausal women, respectively. Applying these values to the accumulated Turner patient years at risk, the expected number of fractures of the distal radius would be four at the post-menopausal fracture rate and one at the premenopausal rate. Similar analysis of neck of femur fractures could not be performed because the number of fractures based on the normal incidences was less than one for both pre- and post-menopausal periods. The patient with a fracture of the lower end of the radius (MS) had a normal BMC.

Discussion

The results are in good agreement with those obtained in another two studies of bone mineral content in Turner's syndrome. In both these studies the mean BMC was significantly reduced, and in both the values fell into two clear groups, those in the normal range and those which were distinctly low. All three studies therefore suggest a bimodal distribution, with the mean for the lower values being in close agreement in the three studies (table 2). This suggests that this may be the usual bone mineral content for the larger proportion of Turner patients. The proportion of patients with low values (73%) also agrees with the prevalence of 'osteoporosis' found radiologically. The BMC values of the patients bore little or no relationship to whether oestrogen therapy had been administered previously (table 1). This was not surprising as it is known that any gain in bone mineral because of oestrogen therapy disappears once treatment is stopped, and only two of the patients in this present study are currently receiving oestrogen (ES and HB). The values for both these patients are at the upper end of the range for the sub-group with low BMC values.

The failure to demonstrate a reduction in bone mineral content with age (fig 2) in this study is substantiated by measurements of metacarpal cortical thicknesses in 22 patients aged 18 to 56 years. No reduction in cortical thickness in the older patients was observed although it was not commented on in the paper. This is contrary to the findings in post-menopausal women in whom there is a progressive reduction in BMC with age. If BMC had been reduced by 1% per annum in the patients with Turner's syndrome we would have expected to see it even with the relatively small numbers studied. It had not been our clinical impression that the fracture rate in Turner patients was noticeably different from normal subjects of similar age. Unfortunately, it is not possible to draw firm conclusions from the analysis of fractures in this study because of the small numbers. However, the incidence of fractures of the distal radius does appear to be lower than the incidence in post-menopausal women and is comparable, in fact, to the incidence in women of similar age. A lower than expected fracture rate was also noted by Pregel et al in a group of over 50 Turner patients.

In conclusion, this study quantifies the bone mineral content in patients with Turner's syndrome. The absence of any reduction in BMC with age and no clear evidence of an increase in frequency of fractures both suggest that if osteoporosis is a

TABLE 2. Comparison of all quantitative studies of BMC in Turner's patients. Note the close agreement in the results of the three studies.

<table>
<thead>
<tr>
<th>BMC site</th>
<th>No of patients</th>
<th>Age range (yr)</th>
<th>No. significantly demineralised</th>
<th>Mean BMC of demineralised patients (% of normal)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radius</td>
<td>8</td>
<td>9-19</td>
<td>6</td>
<td>71% Brown et al</td>
</tr>
<tr>
<td>Os calcis</td>
<td>10</td>
<td>17-28</td>
<td>3</td>
<td>75% Risch et al</td>
</tr>
<tr>
<td>Distal femur</td>
<td>11</td>
<td>18-57</td>
<td>8</td>
<td>73% Present study</td>
</tr>
</tbody>
</table>
complication of Turner's syndrome it should not be thought analogous to the osteoporosis that occurs in post-menopausal women. It would not, therefore, be possible to justify the regular use of long term oestrogen therapy in patients with Turner's syndrome on the grounds of 'osteoporosis'.

We wish to thank Novo Diagnostic Systems for the loan of the Osteodensitometer, Mrs D McKinna for secretarial services, and Miss D Lytton of the Department of Medical Illustration.

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Total body calcium in rheumatoid arthritis: effects of disease activity and corticosteroid treatment

D M REID, N S J KENNEDY, M A SMITH, P TOTHILL, G NUKI

Abstract

Rheumatoid arthritis may be associated with generalised as well as periarticular osteoporosis. To assess the extent of bone loss and the influence of corticosteroid treatment total body calcium was measured by in-vivo neutron activation analysis in 63 patients with rheumatoid arthritis treated with non-steroidal anti-inflammatory drugs alone and 31 treated with additional low-dose corticosteroids. The results were compared with those in 40 normal controls matched for age, sex, and menopausal state. There were significant reductions in mean total body calcium in the group treated with non-steroidal anti-inflammatory drugs (5.3% in men; 6.8% in women) and greater reductions in the corticosteroid-treated patients (11.5% in men, 15.5% in women). The reduction was correlated with disease duration and activity in the patients treated with non-steroidal anti-inflammatory drugs alone. Measured total body calcium was signifi-
cantly less than the values predicted when this relation was used in the corticosteroid-treated patients. The data suggest that increased bone loss in patients with rheumatoid arthritis treated with corticosteroids is attributable to drug treatment rather than disease activity.

Many patients with rheumatoid arthritis treated with low-dosage corticosteroids and some postmenopausal women with the disease are likely to be at risk from the complications of osteoporosis.

Introduction

Rheumatoid arthritis is thought to be associated with generalised as well as periarticular osteoporosis, but considerable uncertainty exists regarding the extent of bone loss and its relation to the duration of disease, disease activity, and additional corticosteroid treatment. Most previous studies of osteoporosis in rheumatoid arthritis have used radiological or photon absorptiometric methods. These techniques have the disadvantage of using single bone sites and may be particularly misleading in a disease associated with severe and variable localised periarticular osteoporosis. The only accurate method of measuring total bone mass in vivo is to measure total body calcium using neutron activation analysis. To assess bone mass in rheumatoid arthritis and its relation to duration of disease, disease activity, and drug treatment we measured total body calcium in sizable groups of patients treated with non-steroidal anti-inflammatory drugs alone and with additional corticosteroids and compared the results with those obtained in controls matched for age, sex, and menopausal state.

Subjects and methods

We measured total body calcium in 63 patients (29 men, 34 women) with definite or classical rheumatoid arthritis treated with non-steroidal anti-inflammatory agents alone and in 31 patients (nine men, 22 women) who were receiving in addition low-dosage corticosteroids (<10 mg prednisolone/day) (see table). The results were compared with those from 40 normal control subjects (20 men, 20 women). Serum calcium, phosphate, and protein concentrations and alkaline phosphatase activity were measured in all subjects, and those with abnormalities of calcium metabolism or medical conditions associated with secondary osteoporosis were excluded.

Total body calcium was measured by in-vivo neutron activation analysis. Patients were irradiated for 20 seconds from front and back, while standing in a rigid polyethylene activation enclosure, using
neutrons from the Edinburgh MRC cyclotron. They were then transferred to a shadow shield whole-body counter, where the gamma radiation from calcium-49 induced from stable calcium-48 by neutron capture was measured for 20 minutes. The patient's total body calcium (in grams) was calculated by comparison with the energy spectrum from an activated anthropomorphic phantom containing a known quantity of calcium. Repeated measurements of the phantom gave a long-term precision of 1.8% for a radiation dose of 13 mSv (133 rem). Mean total body calcium (±1 SD) in the control population was 1142.7 ± 133.8 g in men and 820.5 ± 124.5 g in women. Individual results were expressed as a percentage of the expected normal value for the patient's skeletal size (arm span) and menopausal state. The spread in the normal range was 7.8% in male controls and 6.5% in female controls.

Clinical assessments undertaken at the time of measurement of total body calcium included measurements of articular index, functional state, Westergren erythrocyte sedimentation rate, duration of early morning stiffness, and rheumatoid factor titre. Statistical analysis of results was performed using Student's t test for unpaired variables and by multiple linear regression analysis. The Mann-Whitney test was used for data not normally distributed and for small samples. All patients gave informed consent, and ethical approval was obtained from local and national committees.

Results

The table shows details of age, duration of disease, variables of disease activity, and corticosteroid dosage in the patients studied. Comparison of the results obtained in the control and patient groups (fig 1) showed a significant reduction in mean total body calcium in patients receiving non-steroidal anti-inflammatory drugs alone (men 5.3%, p<0.05; women 6.8%, p<0.01). Mean total body calcium was further reduced in patients receiving low-dosage corticosteroids compared with controls (men 11.5%, p<0.01; women 15.5%, p<0.001) and in women receiving low-dosage corticosteroids compared with those receiving only non-steroidal anti-inflammatory drugs (8.7%, p<0.02).

In patients receiving non-steroidal anti-inflammatory drugs total body calcium was not significantly correlated with any single index of disease activity or duration. Multiple linear regression analysis, however, showed a weak but significant correlation between total body calcium and a composite index of articular index, functional state, erythrocyte sedimentation rate, rheumatoid factor titre, and duration of disease (r=0.34, p<0.01) in 56 patients who had not received gold. A similar correlation was shown in 26 of the patients receiving corticosteroids (r=0.64, p<0.01), in whom there was also an inverse correlation between total body calcium and articular index (r=−0.60, p<0.01).

When the equation obtained by multiple linear regression analysis in patients receiving only non-steroidal anti-inflammatory drugs was used the mean total body calcium in patients receiving additional
Mean age and variables of disease in patients studied (ranges given in parentheses)

<table>
<thead>
<tr>
<th></th>
<th>Men NSAID* (n = 29)</th>
<th>Steroids (n = 9)</th>
<th>Women NSAID* (n = 34)</th>
<th>Steroids (n = 22)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>51.7 (29-70)</td>
<td>59.8 (35-70)</td>
<td>52.4 (25-70)</td>
<td>56.1 (34-66)</td>
</tr>
<tr>
<td>Duration of disease (years)</td>
<td>8-5 (0-8-10)</td>
<td>7-5</td>
<td>9-2 (0-6-30)</td>
<td>12-3 (3-33)</td>
</tr>
<tr>
<td>Erythrocyte sedimentation rate (mm in first h)</td>
<td>35-9 (1-115)</td>
<td>31-9 (2-104)</td>
<td>39-5 (2-165)</td>
<td>35-0 (4-177)</td>
</tr>
<tr>
<td>Rheumatoid factor (1/titre)</td>
<td>392 (0-1280)</td>
<td>366 (0-1280)</td>
<td>345 (0-2480)</td>
<td>332 (0-1280)</td>
</tr>
<tr>
<td>Articular index</td>
<td>10.6 (0-21)</td>
<td>10.6 (0-18)</td>
<td>14.8 (4-31)</td>
<td>10.6 (1-23)</td>
</tr>
<tr>
<td>Functional state</td>
<td>2.14 (1-8)</td>
<td>1.96</td>
<td>1.99 (1-2-5)</td>
<td>2.00 (1-3)</td>
</tr>
<tr>
<td>Prednisolone (mg)</td>
<td>6.1 (4-6)</td>
<td>6.1</td>
<td>6.3 (4-10)</td>
<td>6.5 (4-10)</td>
</tr>
<tr>
<td>Duration of steroid treatment (years)</td>
<td>2.7 (1-6)</td>
<td>6.6 (2-22)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*NSAID = Non-steroidal anti-inflammatory drugs.

FIG 1—Total body calcium expressed as percentage of expected normal value (mean ± 1 SD) in male and female controls and patients with rheumatoid arthritis (RA) treated with non-steroidal anti-inflammatory drugs (NSAID) and corticosteroids.
corticosteroids predicted from the duration of disease and variables of disease activity was found to be 93.6±3.7% (range 87.5-102.3%). This was significantly higher (p<0.005) than the measured value in this group and similar to the measured value (93.2±10.9%) for men and women in the group taking non-steroidal anti-inflammatory agents alone. No significant differences were found in erythrocyte sedimentation rate, functional state, and rheumatoid factor titre between the group receiving non-steroidal anti-inflammatory drugs alone and additional corticosteroids, although the duration of disease was 25% higher (p<0.02) and the articular index 20% lower (p<0.05) in the group receiving corticosteroids.

In patients receiving corticosteroids total body calcium was closely correlated with the mean daily dose of prednisolone (r=−0.66, p<0.001) (fig 2) but not with duration of treatment or the total steroid dose (daily dose x duration of treatment).

Discussion

These studies show a significant reduction in mean total body calcium in patients with rheumatoid arthritis who had not received corticosteroid drugs and strongly suggest that the reduction in bone mass is an integral feature of the disease. The only previous studies of total body calcium in patients with rheumatoid arthritis showed conflicting results. In a small preliminary study Zanzi et al found no reduction in seven women who had not received corticosteroids, while Kennedy et al found an 18% reduction in 32 women compared with hypertensive controls. No details of drug treatment preceding the study were given in the latter paper.
By using localised methods of skeletal measurement several previous studies have suggested a relation between bone loss and duration of disease in rheumatoid arthritis, while others have been unable to confirm this. Most studies in which radiological or absorptiometric methods have been used, however, have suggested a relation between bone loss and variables of disease activity including erythrocyte sedimentation rate and rheumatoid factor titre.

Unfortunately, there is no good method by which overall disease activity over a period of time can be assessed. The composite index used in this study included the erythrocyte sedimentation rate and rheumatoid factor titre as indicators of disease activity at the time of measurement of total body calcium as well as functional state and duration of disease, which also reflect the clinical state over a prolonged period. Although the correlation between total body calcium and the composite index of disease duration and activity in patients who had not received corticosteroid treatment was relatively weak, it did permit total bone mass in the steroid-treated patients to be predicted. The striking discrepancy between measured and predicted total body calcium in the steroid-treated group suggests that loss of bone mass in these patients is directly attributable to the drug treatment and not a function of more prolonged or severe underlying disease. Some previous studies using less sophisticated methods have failed to show a relation between osteoporosis and corticosteroid treatment in patients with rheumatoid arthritis, and it has been suggested that low-dosage treatment may be free from this risk and that improved mobility attributable to the use of corticosteroids may have the effect of reducing bone loss. Most studies, however, have shown evidence of increased bone loss in patients with rheumatoid arthritis treated with corticosteroids, although this has not been correlated with steroid dosage. The finding that daily corticosteroid dosage is correlated with reduction in total body calcium while duration of corticosteroid treatment is not is consistent with the hypothesis that the histochemical changes and catabolic effects of corticosteroids on bone are most pronounced early in the course of treatment.

Previous measurements of total body calcium in women with symptomatic postmenopausal osteoporosis have shown mean reductions of 16 and 18% greater than expected values. This would seem to suggest that despite low mean daily doses of prednisolone (men 6-1 mg; women 6-3 mg) some male patients with rheumatoid arthritis treated with corticosteroids (mean reduction in total body calcium of 11-5%) and many female patients (mean reduction 15-5%) are at risk of osteoporotic complications. As a group patients treated with non-steroidal anti-inflammatory drugs alone, with a mean reduction in total
body calcium of only 5-3% in men and 6-8% in women, are probably not at high clinical risk of complications of osteoporosis. Since, however, the range of the reduction was wide (−20% to +12% in men; −33% to +19% in women) and many patients with rheumatoid arthritis are postmenopausal women, probably many individual patients who have not received corticosteroids will sustain clinically important reductions in skeletal mass. Although there has been an isolated report of spontaneous fractures in patients with rheumatoid arthritis who had not received corticosteroids, there is clearly a need for epidemiological studies of the incidence of fracture in patients with this disease.

These studies also suggest that it will be important to try and establish the pathological basis of reduced bone mass in patients with rheumatoid arthritis as well as to determine to what extent this is the result of local and generalised osteoporosis.

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References


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Comparison between $^{153}$Gd and $^{241}$Am, $^{137}$Cs for dual-photon absorptiometry of the spine

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Abstract. The bone mineral content (BMC) of the spine can be measured from the attenuation of two photon beams. An equation was derived for the theoretical precision of BMC measurement and validated experimentally. Though $^{153}$Gd was found to possess the better energy combination, the precision due to the photon energy levels using $^{241}$Am, $^{137}$Cs was only 1.5 times the value obtained from $^{153}$Gd. The available photon outputs from $^{241}$Am and $^{153}$Gd were investigated and with a 12 mm diameter disc source a theoretical precision of 1% could be obtained with source strengths 20 GBq $^{241}$Am or 5 GBq $^{153}$Gd in a 40 min and 20 min patient scan respectively. There was no great advantage in either energy combination when the problems of fat or patient dose were considered. Given the disadvantages of $^{153}$Gd, its cost and availability, $^{241}$Am, $^{137}$Cs can be used as a practical alternative to measure spine BMC from a reconstructed bone mineral image.

1. Introduction

Dual-photon absorptiometry (DPA) to measure bone mineral content (BMC) was first proposed by Reed (1966). Subsequently interest concentrated on the forearm and single-photon absorptiometry became the method of choice (Cameron and Sorensen 1963, Christiansen and Rødbro 1977). More recently the technique of DPA has been used to measure the BMC of the spine, the site of most clinical interest, (Mazess et al 1974, Roos and Sköldborn 1974, Wilson and Madsen 1976, Price et al 1977, Condon et al 1979, Krølner and Pors Nielsen 1980, Wahner and Dunn 1980, Smith et al 1983) and whole body (Price et al 1977, Peppier and Mazess 1981), the second energy being used to compensate for variations in the thickness of overlying soft tissue.

The photon energies at 60 keV and 660 keV from $^{241}$Am and $^{137}$Cs respectively have been used by Roos and Sköldborn (1974) and Condon et al (1979). More commonly the radionuclide $^{153}$Gd, with effective energies at 44 keV and 100 keV has been used (Mazess et al 1974, Wilson and Madsen 1976, Krølner and Pors Nielsen 1980, Wahner and Dunn 1980, Smith et al 1983). The reason for this is that it has become generally accepted that $^{153}$Gd possesses the optimum energy combination for such work. Mazess et al (1974) state that $^{153}$Gd is optimal for bones larger than the radius and ulna, citing the theoretical work of Judy (1971). However Judy (1971) concludes that the optimum lower energy for DPA is between 30 keV and 40 keV for a thickness of soft tissue of only 5 g cm$^{-2}$. Peppier and Mazess (1981) state that $^{153}$Gd is optimal, citing the theoretical work of Hanson (1974) and Watt (1975). Hanson (1974), however, compared $^{153}$Gd with $^{125}$I, $^{241}$Am for thicknesses of soft tissue ranging

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up to 10 g cm\(^{-2}\). Watt (1975) does show the optimum lower energy to be 40 keV, but for a total sample thickness of only 15 g cm\(^{-2}\) and an upper energy of 400 keV. The suggestion that \(^{153}\)Gd possesses the optimum lower energy is therefore interpreted from calculations using thicknesses that are of the order of half that found in human subjects in the region of the lumbar spine. In addition, none of the theoretical analyses included all the variables associated with the bone mineral measurement.

Although \(^{153}\)Gd may have nearly the optimum energy for BMC measurements of the spine, it has a relatively short half-life of 240 d compared with 458 yr for \(^{241}\)Am. In addition, \(^{153}\)Gd is more expensive and less readily available than \(^{241}\)Am. The cost of \(^{153}\)Gd, which has to be replaced annually, is therefore considerably greater than \(^{241}\)Am.

Two factors, however, cause \(^{241}\)Am to have a much lower photon output per unit volume than \(^{153}\)Gd. The first is its much longer half-life. The second is the photons emitted per disintegration, 36% for \(^{241}\)Am and 110% for the lower energy of \(^{153}\)Gd. For a given output at the lower energy a much larger volume of \(^{241}\)Am is therefore required and self absorption becomes significant. It is accepted that to reduce the error due to repositioning without excessively increasing the patient dose, the best method to measure spine BMC is to scan the patient and construct a bone mineral image which includes the region of the spine to be measured. The exact region of interest, for example the lumbar vertebrae L2, L3 and L4, can then be selected from the computer-processed bone mineral image. Such a technique requires sufficient resolution and sensitivity to produce an adequate bone mineral image. If narrow collimation of the source is used to produce the image then the maximum possible photon output from \(^{241}\)Am is much lower than from a \(^{153}\)Gd source, the activity of which is typically 37 GBq (1 Ci). However previous work using \(^{153}\)Gd with lower activity of 7.4 GBq (200 mCi) (Smith et al 1983), showed that sufficient resolution to distinguish the separate lumbar vertebrae could be obtained with collimator sizes larger than had been used previously allowing the possibilities of higher activities of \(^{241}\)Am to be considered.

This study was undertaken firstly to establish the optimum energy combination for DPA of the spine at realistic soft tissue and bone thicknesses, secondly to compare the relative merits of the energy combinations from \(^{153}\)Gd and \(^{241}\)Am, \(^{137}\)Cs and thirdly, given that adequate images can be produced with larger collimation than has been generally used, to determine whether \(^{241}\)Am can produce a high enough photon output for DPA. The results would then determine whether \(^{241}\)Am, \(^{137}\)Cs could be used in practice as an acceptable alternative to \(^{153}\)Gd.

2. Methods

The content of either component of a two-phase system (e.g., bone and soft tissue) can be obtained from the attenuation of two suitable monoenergetic photon beams through the system, combined with the unattenuated count rates. For in vivo measurements of the spine a third phase, fat, is present. The assumption is made that the fat content is evenly distributed across the body. The BMC of the spine is therefore calculated using a region adjacent to the spine as the ‘background’ region, rather than the unattenuated through-air count rate (Roos et al 1980).

The relative merits of different energy combinations, irrespective of the actual count rate, were obtained using the following equations for the theoretical variance of the BMC measurement:
The measurement of BMC is obtained from four values

\[ I_1 = \text{total counts from energy 1 (lower) over the spine} \]
\[ I_2 = \text{total counts from energy 2 (upper) over the spine} \]
\[ I_{1s} = \text{total counts from energy 1 over the soft tissue background} \]
\[ I_{2s} = \text{total counts from energy 2 over the soft tissue background} \]

using the equation

\[ M_b = K_a \ln(I_{1s} / I_1) - K_b \ln(I_{2s} / I_2) \]  (1)

where

\[ M_b \text{ is the BMC in g cm}^{-2} \]
\[ K_a = \mu_{s2} / (\mu_{b1} \mu_{s2} - \mu_{b2} \mu_{s1}) \]
\[ K_b = \mu_{s1} / \mu_{s2} \]
\[ \mu \text{ is the mass attenuation coefficient (cm}^2 \text{ g}^{-1}). \]

Subscripts s, b, 1 and 2 refer to soft tissue, bone and the lower and upper energies respectively. Also

\[ I_1 = *I_1 - \chi_2 \]
\[ I_{1s} = *I_{1s} - \chi_1 \]

where \(*I_1\) and \(*I_{1s}\) are the actual measured counts and \(\chi_2\) is the contribution from the upper energy in the region of the lower energy referred to as the crossover correction (Smith and Tothill 1982).

The variance of the BMC measurement is given by

\[ \sigma^2(M_b) = \left( \frac{\partial M_b}{\partial I_1} \right)^2 \sigma^2(*I_1) + \left( \frac{\partial M_b}{\partial I_2} \right)^2 \sigma^2(*I_2) + \left( \frac{\partial M_b}{\partial I_{1s}} \right)^2 \sigma^2(*I_{1s}) + \left( \frac{\partial M_b}{\partial I_{2s}} \right)^2 \sigma^2(*I_{2s}) \]

\[ = \left( \frac{\partial M_b}{\partial I_1} \right)^2 *I_1 - \left( \frac{\partial M_b}{\partial I_2} \right)^2 *I_2 + \left( \frac{\partial M_b}{\partial I_{1s}} \right)^2 *I_{1s} - \left( \frac{\partial M_b}{\partial I_{2s}} \right)^2 *I_{2s} \]

\[ = \frac{K_a^2}{I_1^2} \left[ (C + A) (1 + 2 \chi_2 K_b) + \frac{C^2}{D} \frac{A^2}{B} \right] (\chi_2 + \chi_1) \]  (2)

where

\[ I_{1s}^\text{air} = \text{total counts at energy 1 through air—similarly } I_{2s}^\text{air} \]
\[ A = \exp(\mu_{s1} T_s + \mu_{b1} t_b) = I_{1s}^\text{air} / I_1 \]
\[ B = \exp(\mu_{s2} T_s + \mu_{b2} t_b) = I_{2s}^\text{air} / I_2 \]
\[ C = \exp(\mu_{s1} T_s) = I_{1s}^\text{air} / I_{1s} \]
\[ D = \exp(\mu_{s2} T_s) = I_{2s}^\text{air} / I_{2s} \]
\[ t_b = \text{thickness of bone (g cm}^{-2}); \text{ this is equivalent to } M_b \text{ in equation (1)} \]
\[ t_s = \text{thickness of soft tissue overlying bone} \]
\[ T_s = \text{thickness of soft tissue adjacent to bone} \]
\[ z = I_{2s}^\text{air} / I_{1s}^\text{air}. \]
To obtain the optimum value of $z$

\[
\frac{\partial (\sigma^2(M_b))}{\partial z} = 0
\]

\[
z = \left( \frac{K_b(D+B)}{(C^2/D+A^2/B)(xo-xo^2)} \right)^{1/2}
\]  \hspace{1cm} (3)

As would be expected the variance of the BMC is a function of sensitivity; in the case of the equation above, of the unattenuated lower energy count rate. However the equation for the variance of the BMC (equation 2) can be written as

\[
\sigma^2(M_b) = \frac{\phi^2}{I_{\text{av}}}
\]

where $\phi$ will be referred to as the comparative error function (CEF) and which is independent of the count rate. The CEF will be proportional to the coefficient of variation (CV) of the BMC measurement, which is usually defined as the precision or reproducibility.

The theoretical error analysis of spine BMC measurement using the equations above was performed for a range of energies, bone mineral values and soft tissue thicknesses. The range of the lower energy was 30 keV to 80 keV, the range of the upper energy 100 keV to 800 keV. Soft tissue values ranged from 5 g cm$^{-2}$ to 30 g cm$^{-2}$ and the bone mineral from 0.6 g cm$^{-2}$ to 1.4 g cm$^{-2}$. The values for the bone mineral covered the range detected in patients and normal volunteers (Hansson and Roos 1980, Riggs et al 1981). The mass attenuation coefficients, derived from the data of Sorenson and Cameron (1974), are listed in table 1, the soft-tissue values being 85% muscle and 15% fat. When simulating $^{153}$Gd, energies of 44 keV and 100 keV were used with $z = 1$. The value of $z = 1$ is higher than from the theoretical emissions but is the approximate ratio that has been observed from encapsulated sources (Peppier and Mazess 1981, Smith and Tothill 1982).

| Table 1. Mass attenuation coefficients (cm$^2$ g$^{-1}$) |
|----------------|----------------|
| $^{153}$Gd       | $^{241}$Am,$^{137}$Cs |
| 44 keV           | 100 keV          | 60 keV  | 660 keV |
| $\mu_p$         | 0.790            | 0.203   | 0.402   | 0.076   |
| $\mu_s$ (15% fat)| 0.241            | 0.170   | 0.204   | 0.086   |
| $\mu_m$         | 0.245            | 0.170   | 0.205   | 0.086   |
| $\mu_t$         | 0.216            | 0.170   | 0.195   | 0.086   |

To examine whether $^{153}$Gd possesses the ideal energies for spine BMC measurement, the CEF was determined over the energy ranges mentioned previously. The bone mineral values and soft tissue thicknesses used were more realistic than those used in previous analyses. The relative advantage, if any, of $^{153}$Gd over $^{241}$Am,$^{137}$Cs due to the photon energy levels was investigated by studying the ratio of the CEFs, $\phi_{Am,137Cs}/\phi_{Gd}$. The effects of the bone mass, soft tissue mass, xo and method of background calculation on the CEF ratio and the optimum value of $z$ for $^{241}$Am,$^{137}$Cs were investigated.
The theoretical precision of spine BMC measurement using the $^{153}$Gd and $^{241}$Am, $^{137}$Cs was compared with experimental reproducibility measurements on phantoms and normal volunteers. A 12 mm diameter sealed $^{153}$Gd disc source (ref. no X.92/1, Amersham 1981) of activity 2.5–5.5 GBq(68–150 mCi) was used. The stainless steel ‘window’ for the source was specified as 0.2–0.25 mm. The $^{241}$Am,$^{137}$Cs combination used a 5 mm diameter spherical $^{241}$Am source of activity 7.4 GBq(200 mCi) (ref. no X.108, Amersham 1981). Again, the window for the source was stainless steel, 0.2–0.3 mm thick. The $^{137}$Cs was a 2 mm diameter, 0.93 GBq(25 mCi) spherical bead placed behind the $^{241}$Am. Bone mineral content measurements of the spine were obtained using a dual headed rectilinear scanner, the J&P Multipoise II, with NaI crystals 10 cm diameter and 5 cm thick. The top crystal was retained in use whilst the source was placed in the position of the lower one. The minimum scan speed was 5 mm s$^{-1}$ with a stepping distance of 4 mm, so an average spine scan of 12 cm by 20 cm took 20 min. Two matrices of the high and low-energy transmission scans were stored on computer. At the end of the scan these could be combined to produce a two-dimensional bone mineral image of the lumbar spine on a colour TV display. Regions of interest around the vertebrae and a suitable background region could then be chosen. A variety of collimator sizes were used to vary the count rate and resolution though the separation remained constant at 0.4 m. In all cases there was sufficient resolution to define a region of interest around L2, L3 and L4.

Two types of phantom were used for experimental reproducibility measurements. The first consisted of the bones of a spine and pelvis encased in paraffin wax 15 cm thick. The bones were obtained from a cadaver and had a normal BMC (1.0 g cm$^{-2}$). The second was an anthropomorphic phantom consisting of a human skeleton encapsulated in tissue-equivalent rubber with a front to back thickness in the lumbar spine region of 22.5 cm. The BMC was lower than normal (0.8 g cm$^{-2}$) as was also confirmed by total body neutron activation analysis (Kennedy et al 1982). The precision of BMC measurements on human subjects was obtained from sets of two repeated measurements, separated by at least a week, on a number of normal volunteers. A single experienced operator analysed the scans used for the precision estimation so that the results would not contain any inter-observer error. Using equation (2) and measured values of A, B, C and D, the theoretical precision was calculated in all cases for comparison.

When comparing the two energy combinations the error due to fat must be considered. In equation (1) the assumption is made that the percentage of fat in the soft tissue is the same in both the background region and over the spine. The value for $\mu_\ell$ used assumes 15% fat and 85% muscle. If, however, there is a varying proportion of fat in the two regions then an error is introduced into the BMC calculation. To estimate the error in the BMC due to fat, associated with the two different energy combinations, the following equations were used in which the components of soft tissue, fat and muscle were considered separately:

$$I_1 = I_{1air} \exp(-\mu_1 f_m - \mu_1 b_0 - \mu_1 f(t))$$
$$I_2 = I_{2air} \exp(-\mu_2 f_m - \mu_2 b_0 - \mu_2 f(t))$$
$$I_1^{soft} = I_{1air} \exp(-\mu_{1,0} T_m - \mu_{1,0} T)$$
$$I_2^{soft} = I_{2air} \exp(-\mu_{2,0} T_m - \mu_{2,0} T).$$

Substituting in equation (1)

$$M_0 = K_0 \{b_0 (\mu_{b1} - K_{b1b2}) + (T_m - T) (\mu_{m1} - K_{b1b2}) \}$$
where

\[ T_m, T_l = \text{thickness of muscle and fat (g cm}^{-2}\text{) adjacent to the bone} \]
\[ t_m, t_l = \text{thickness of muscle and fat overlying the bone} \]
\[ \mu_m, \mu_l = \text{mass attenuation coefficients of muscle and fat at energies 1 and 2.} \]

Other factors in the equation have been defined previously.

The mass attenuation coefficients listed in table 1 were used and the values \( T_m, T_l, t_m \) and \( t_l \) were varied. The calculated BMC was compared with the true value \( t_0 \) for both energy combinations.

The two energies can also be used to estimate the percentage fat in the background region adjacent to the spine. The error in the fat estimation due to counting statistics was calculated using equation (2), substituting the appropriate bone mineral parameters with those for fat and the soft tissue parameters with those for muscle. This enabled the precision of the percentage fat estimation to be obtained. The major source of error in the accuracy of the fat estimation is due to inaccuracies in the mass attenuation coefficients which are known typically to three decimal places. The constant \( K_o \), which in the estimation of fat content becomes \( \mu_{m2}/(\mu_1\mu_{m2} - \mu_{(3\mu_1m1)}) \), is affected most by errors in \( \mu \). The maximum variation in the estimation of the fat content was therefore calculated using the values of \( \mu \) in table 1 plus and minus 0.0005.

The patient skin dose from the sources described previously was measured. An ionisation chamber was placed on the bed and surrounded by tissue-equivalent material. The dose was measured whilst the scanner operated in the same mode as was used for spine measurements.

Having obtained a comparison between \(^{153}\text{Gd}\) and \(^{241}\text{Am},^{137}\text{Cs}\) intrinsic to the photon energy levels, experimentally validated the theoretical precision estimation and considered the effect of fat and dose, the question of available count rate was investigated. The error due to counting statistics in DPA is determined principally by the lower energy count rate. For spine BMC measurements the source would have a single circular hole collimator, so a disc shaped source would give the highest output per unit volume. The degree of self absorption is negligible in a \(^{153}\text{Gd}\) source but significant in a \(^{241}\text{Am}\) source. The photon output for various sizes of \(^{241}\text{Am}\) and \(^{153}\text{Gd}\) sources of different activities was investigated. The photon output was measured using the two sources and detector described previously plus in addition a 12 mm diameter 11.1 GBq (300 mCi) \(^{241}\text{Am}\) source (ref. no X.92/0, Amersham 1981) with 0.2–0.25 mm stainless steel window.

Previous research into DPA of the spine had shown that adequate resolution, sufficient to define the space between the lumbar vertebral on a bone mineral image, could be obtained using a 12 mm diameter source collimator, a 20 mm square detector collimator and a separation between the two of 0.4 m (Smith et al 1983). The theoretical precision of spine BMC measurement for different activities of \(^{153}\text{Gd}\) and \(^{241}\text{Am}\) was calculated using values of \( I_{70}^{77} \) that could be obtained from commercially available sealed sources. The variation of precision with source activity was calculated assuming a 12 mm diameter disc source and source collimator, 20 mm square detector collimator, 0.4 m separation and a scan area of 12 cm by 20 cm. The time taken to scan this area was assumed to be 20 min or 40 min. The average area over L2, L3 and L4 was calculated from six volunteer scans, and theoretical precision was calculated for both a subject with normal spine bone mineral and soft tissue thickness and the 'worst' type of patient, one with dimineralised bone and a large thickness of soft tissue.
3. Results

The optimum energy combination to minimise the CEF for spine BMC measurements, assuming the optimum value of $z$, was found to be approximately 43 keV and 180 keV for a bone mineral of 1.0 g cm$^{-2}$ and a soft tissue thickness of 20 g cm$^{-2}$. The variation in CEF is shown in figure 1 for a range of lower and upper photon energies. It can be seen that though there is a well defined minimum at approximately the same lower energy level, the upper energy level has little effect on the CEF.

The results of the CEF ratio for $^{241}$Am, $^{137}$Cs and $^{153}$Gd are shown in figure 2. A $\chi^2$ value of 4% was used and the assumption was made that the area to define the background level on the bone mineral image was the same as that used for the bone mineral region. Also shown in figure 2 is a histogram showing the distribution of

Figure 1. Variation in CEF for a range of lower and upper energy values. Curve A, 100 keV; B, 200 keV; C, 400 keV; D, 600 keV.

Figure 2. Ratio of CEF for a range of soft tissue thicknesses. BMC values of 0.6, 1.0 and 1.4 g cm$^{-2}$ are used. Dashed line: $\chi^2 = 1$ for $^{241}$Am, $^{137}$Cs and $^{153}$Gd. Dotted line: $\chi^2 = 1$ for $^{153}$Gd, optimum $\chi^2$ for $^{241}$Am, $^{137}$Cs. A histogram showing the variation in soft tissue thickness is also shown.
anterior to posterior thicknesses in the lumbar region of 180 patients and normal subjects. It can be seen that for measurements of the spine, though $^{153}$Gd may possess nearly the optimum lower photon energy, the intrinsic error due to the photon energies using the energy combinations from $^{241}$Am, $^{137}$Cs is only about 1.5 times greater than the error obtained using $^{153}$Gd. The value of the $\chi_0$ was found to have little effect on the CEF ratio; for example if the bone mineral is 1.0 g cm$^{-2}$ and the soft tissue thickness is 20 g cm$^{-2}$ then a $\chi_0$ of 6% will increase the CEF ratio by 3.8% and a $\chi_0$ of 2% will reduce it by 6.4%.

Figure 3 shows the variation in $z$, the optimum ratio of the two values.

![Figure 3](image.png)

**Figure 3.** Variation in optimal value of $z$ with soft tissue thickness. BMC value of 1.0 g cm$^{-2}$ and $\chi_0$ values of 2%, 4% and 6% are used.

The results in table 2 show how well the theoretical error agrees with the experimental reproducibility. The large 4 cm diameter detector collimator used with $^{241}$Am and the wax phantom showed the effect of increasing the $^{241}$Am photon output. With this size collimator it was possible to distinguish the lumbar vertebrae on the bone mineral image of the phantom but not of the human subjects.

The error due to fat is shown in figures 4(a) and 4(b). It can be seen that with both a constant proportion of fat and different amounts of soft tissue and with different proportions of fat and the same amount of soft tissue the error in BMC is worse using $^{153}$Gd by a factor of 1.3 compared with $^{241}$Am, $^{137}$Cs. The precision of the fat estimation (CV) using the two energies in the region adjacent to the spine improves as the total soft tissue thickness decreases and the percentage fat increases. For a total soft tissue thickness of 20 g cm$^{-2}$, 15% fat and $\rho_{w} = 10^8$, the precision of the fat estimation using $^{153}$Gd is 1.7% and using $^{241}$Am, $^{137}$Cs is 3.8%. The maximum error in the fat estimation due to the assumed inaccuracies of the mass attenuation coefficients varies with the percentage fat content. With 15% fat the maximum error is 8% and 20% and with 30% fat the maximum error is 2.5% and 6.5% for $^{153}$Gd and $^{241}$Am, $^{137}$Cs respectively.

The skin dose was found to be 123 $\mu$Sv when the $^{153}$Gd activity was 10.6 GBq (286 mCi). The dose from the 7.4 GBq (200 mCi) $^{241}$Am plus 0.93 GBq (25 mCi) $^{137}$Cs combination was 14 $\mu$Sv.
Table 2. Theoretical and experimental precision using phantoms and normal volunteers.

(i) Anthropomorphic phantom using $^{153}$Gd.

<table>
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<th>Collimation</th>
<th>CV (%)</th>
<th>Number of measurements</th>
</tr>
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<tr>
<td>Detector (cm x cm)</td>
<td>Source (mm)</td>
<td>Measured</td>
</tr>
<tr>
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<td>13</td>
<td>1.1</td>
</tr>
<tr>
<td>2 x 2</td>
<td>10</td>
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</tr>
<tr>
<td>2 x 2</td>
<td>6</td>
<td>2.5</td>
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</table>

(ii) Wax phantom using $^{153}$Gd and $^{241}$Am, $^{137}$Cs.

<table>
<thead>
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<th>Source</th>
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<th>CV (%)</th>
<th>Number of measurements</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{241}$Am, $^{137}$Cs</td>
<td>4 cm</td>
<td>0.6</td>
<td>0.90</td>
</tr>
<tr>
<td>$^{153}$Gd</td>
<td>2 cm x 2 cm</td>
<td>0.4</td>
<td>0.37</td>
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</tbody>
</table>

(iii) Normal subjects using $^{153}$Gd and $^{241}$Am, $^{137}$Cs

<table>
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<th>Source</th>
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<th>Number of measurements</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{241}$Am, $^{137}$Cs (Condon et al 1979)</td>
<td>2-3</td>
<td>2.64</td>
</tr>
<tr>
<td>$^{153}$Gd (Tothill et al 1981)</td>
<td>0.95</td>
<td>0.96</td>
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Figure 4. Variation in calculated BMC with non-uniform distribution of fat in background and over spine.

(a) True BMC 1.0 g cm$^{-2}$, attenuation coefficients assume 15% fat. Percentage fat is constant over bone and in background but amount of soft tissue in background is 25 g cm$^{-2}$ and over bone is 15 g cm$^{-2}$. (b) Amount of soft tissue is constant, 20 g cm$^{-2}$, and true BMC 1.0 g cm$^{-2}$. Percentage fat over bone and in background is different. In each case, curve A, $^{153}$Gd; curve B, $^{241}$Am, $^{137}$Cs.
Various factors were found to affect the detected count rate from $^{241}\text{Am}$ and $^{153}\text{Gd}$. It can be seen from figure 5 that the theoretical self-absorption within a disc source is a major problem with $^{241}\text{Am}$. The problem of self-absorption is not significant however for $^{153}\text{Gd}$, there being less than 1% with a source activity of 56 GBq(1.5 Ci). The stainless steel window of the source encapsulation was assumed to be 0.225 mm thick giving an attenuation of 19% and 39% at 60 keV and 44 keV respectively. An additional source of photon attenuation in the source is caused by the fact that $^{241}\text{Am}$ is mixed with a ceramic which does not form a perfect disc when set but has one concave face. This results in a reduction of approximately 15% in the photon output. An improvement in the output from a sealed source can be achieved if a 1 mm Be window is used. This will cause an attenuation of only 3% for both radionuclides. The photon output from the sealed source is also significantly attenuated in the housing around the NaI crystal, approximately 15% and 35% at 60 keV and 44 keV respectively.

The theoretical variation of precision with source activity is shown in figure 6. The value of $I_1^\nu$ used is obtained from the theoretical output assuming attenuation by the factors described above and with a 1 mm Be window rather than stainless steel. The time spent over L2, L3 and L4 was taken as 357 s and 750 s for 20 min and 40 min patient scans respectively. It can be seen that with the collimation used, $^{153}\text{Gd}$ has an acceptable theoretical precision of less than 1.0% at activities down to 5 GBq(135 mCi). This is also the case when an activity of more than 20 GBq(540 mCi) of $^{241}\text{Am}$ is used with a 40 min scan time.

4. Discussion

The theoretical analysis of the variance of the bone mineral estimation is presented in a more comprehensive form than in previous studies. Wooten (1971) ignored the effect of $x_0$ and assumed four different optimum count times through bone and the adjacent background at the two energies, and also different optimum values of $z$. Hanson (1974) in his analysis ignored the $x_0$, the error due to the background estimation and the ratio of the two energies. Roos (1974) dealt more thoroughly with
the various factors but his analysis was very specific to his particular technique and he did not present a single equation of the total error that was generally applicable.

The relative error function (REF) proposed by Judy (1971) and Watt (1975) also ignored the $x_0$, assumed that the background is estimated from the unattenuated beams and that the ratio of the two energies was unity. In fact the REF is a special case of the CEF presented in this paper with $x_0 = 0$, $z = 1$ and $C = D = 1$. When these values were inserted into the equation for the CEF, identical results to those given in the paper by Watt (1975) were found.

The theoretical analysis confirms that $^{153}$Gd possesses the ideal lower energy for spine BMC measurement, though the upper energy is rather lower than the optimum. Given that the upper energy is 100 keV, however, the actual ratio of the photon outputs is reasonably close to the optimum value. Though intrinsically the energy combination of $^{153}$Gd is superior to $^{241}$Am, $^{137}$Cs, the precision in BMC measurement is only increased by a factor of 1.5 if the latter is used. This suggests that $^{241}$Am, $^{137}$Cs is a practical proposition if a high enough output can be obtained.

Neither $^{153}$Gd nor $^{241}$Am, $^{137}$Cs showed any clear advantage when the problem of fat was considered. Though $^{153}$Gd estimates the fat content in the background region with greater precision and accuracy than $^{241}$Am, $^{137}$Cs, a greater error is introduced in the BMC calculation using $^{153}$Gd if the fat is not uniformly distributed over the spine.

There is also no particular advantage between the two energy combinations when the patient dose, which is very low in both cases, is considered. The skin dose to the patient from DPA of less than 150 µSv is far lower than other quantitative techniques, such as neutron activation analysis, where the dose can be of the order of 30–60 mSv (Al-Hiti et al 1976, Smith and Tothill 1979).

For clinical studies into metabolic disorders of the bone, such as osteoporosis, the precision of the technique needs to be around 2%, so the theoretical precision based on counting statistics should be ideally around 1%. Using the 2 cm square detector collimator and a 12 mm diameter disc source, which has been shown to give adequate resolution of the BMC image, theoretical and experimental results (figure 6 and

![Figure 6. Variation in theoretical precision (CV) with activity of $^{153}$Gd or $^{241}$Am. (a) 20 g cm$^{-2}$ soft tissue, 1.0 g cm$^{-2}$ bone, 20 min scan. (b) 25 g cm$^{-2}$ soft tissue, 0.8 g cm$^{-2}$ bone, 20 min scan. (c) 25 g cm$^{-2}$ soft tissue, 0.8 g cm$^{-2}$ bone, 40 min scan.](image)
Table 2 showed that this degree of precision could be obtained with a 20 min scan if at least 5 GBq(135 mCi) \(^{153}\text{Gd}\) were used. Clearly the sensitivity of \(^{241}\text{Am}\) could be increased by reducing the scan speed, but it is our experience that 40 min is about the maximum time that a patient will endure without becoming restless. Using the same collimation and a 40 min scan time, it is possible to achieve this precision if at least 20 GBq(540 mCi) \(^{241}\text{Am}\) are used.

The cost of \(^{241}\text{Am}\) is considerably less than \(^{153}\text{Gd}\), though the cost of the latter does vary in different countries. In Great Britain the cost of a 37 GBq(1 Ci) \(^{241}\text{Am}\) 12 mm disc source with a Be window would be of the order of £2000 compared with £7000 for the same activity of \(^{153}\text{Gd}\). Given that \(^{153}\text{Gd}\) needs to be replaced about every 12 months, there is a considerable financial advantage in developing apparatus which uses \(^{241}\text{Am}\). In addition the availability of \(^{153}\text{Gd}\) is irregular.

The results in this paper show that activities down to 5 GBq(135 mCi) \(^{153}\text{Gd}\) can be used suggesting that a 37 GBq(1 Ci) \(^{153}\text{Gd}\) source could be used for about two years. To use activities over this range, the size of the collimation would have to be increased during the two year period to avoid dead-time errors at high activity and to give a high enough count rate at low activity. This would alter the geometry and also necessitate the use of different \(X_0\) values (Smith and Tothill 1982) which could introduce errors into the BMC measurement. If longitudinal patient studies are to be undertaken, such as the study of osteoporosis covering a period of many years, then it would be desirable to use a technique whose parameters remain constant.

Acknowledgments

The authors would like to thank Mr J Wilson and Dr J Hannan for technical assistance and advice, the Radiochemical Centre, Amersham for technical information and the loan of one source and Miss D Lytton for drawing the figures. The research was partially funded by the Scottish Home and Health Department.

Résumé

Etude comparée du \(^{153}\text{Gd}\) et du couple \(^{241}\text{Am}/^{137}\text{Cs}\) pour les mesures d’absorption dans la colonne vertébrale en double énergie.

Le contenu minéral de l’os (BMC) peut être mesuré dans le rachis à partir de l’atténuation de deux faisceaux de photons. Une formule a été établie pour l’estimation théorique de l’incertitude qui affecte la mesure du BMC, et elle a été validée expérimentalement. Bien que le \(^{153}\text{Gd}\) présente la meilleure combinaison d’énergies de photons, l’incertitude due aux niveaux d’énergies des photons du couple \(^{241}\text{Am}/^{137}\text{Cs}\) est seulement 1,5 fois plus importante que celle correspondant au \(^{153}\text{Gd}\). Les flux de photons disponibles avec le \(^{241}\text{Am}\) et le \(^{153}\text{Gd}\) ont été mesurés et, pour une source ayant la forme d’un disque de 12 mm de diamètre, une incertitude théorique de 1% peut être obtenue lors de l’examen d’un patient avec 20 GBq de \(^{241}\text{Am}\) (durée 40 min) et 5 GBq de \(^{153}\text{Gd}\) (durée 20 min). Aucun avantage important n’apparaît avec l’une ou l’autre des combinaisons d’énergies pour la dose reçue par le patient ou les problèmes posés par la graisse. Aussi, compte-tenu des inconvénients du \(^{153}\text{Gd}\), en particulier son prix et sa disponibilité, il peut être remplacé en pratique par le couple \(^{241}\text{Am}/^{137}\text{Cs}\) pour la mesure du BMC dans la colonne vertébrale à partir d’une image reconstruite du contenu minéral de l’os.

Zusammenfassung

Vergleich zwischen \(^{153}\text{Gd}\) und \(^{241}\text{Am}/^{137}\text{Cs}\) für 2-Photonenabsorptiometrie der Wirbelsäule.

Der Knochenmineralgehalt (KMG) der Wirbelsäule kann durch die Schwächung zweier Photonenstrahlen gemessen werden. Eine Gleichung für die theoretische Genauigkeit von KMG-Messungen wurde abgeleitet.
und experimentell bestätigt. Obwohl $^{153}$Gd die günstigere Energiekombination besitzt, ist die Genauigkeit, die durch die Photonenenergieniveaus bei Verwendung von $^{241}$Am/$^{137}$Cs erreicht wird 1.5 mal größer als die für $^{153}$Gd erhaltenen Werte. Die verfügbaren Photonemissionen von $^{241}$Am und $^{153}$Gd wurden untersucht und mit einer Flächenquelle von 12 mm Durchmesser konnte eine theoretische Genauigkeit von 1% erreicht werden mit Quellstärken von 20 GBq $^{241}$Am oder 5 GBq $^{153}$Gd in einem 40- bzw. 20-minütigen Patienten-Scan. Es gab für beide Energiekombinationen keine großen Vorteile bei der Berücksichtigung der Fett- oder Patientendosis. Wegen der Nachteile von $^{153}$Gd, Kosten und Verfügbarkeit, kann $^{241}$Am/$^{137}$Cs als praktische Alternative zur Messung des KMB der Wirbelsäule aus einem rekonstruierten Bild des Knochenminerals verwendet werden.

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but they have misrepresented the procedures of others and have alluded to intra-laboratory differences where none have been shown to exist.

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24 February 1983

Reference


Intra-laboratory variations using dual-photon absorptiometry

The Editor,

Sir,

We welcome the letter by Mazess et al (previous letter), which provides further justification for our belief that technical problems in dual-photon absorptiometry (DPA) need to be explored. In our recent scientific note (Smith and Tohill 1982) we stated that there were intra-laboratory differences between cross-sectional studies of vertebral bone mineral content (BMC) in normal women as measured by DPA. Two groups have reported an increase in the rate of loss of BMC from the spine at the time of the menopause (Krølner and Pors Nielsen 1982, Smith et al 1983) whereas two other groups have not (Madsen 1977, Riggs et al 1981). Examples from two groups are shown in figure 1 illustrating this fairly fundamental disparity. Our own normal data (Tohill et al 1983) show that the units used to express the BMC, whether in grams (Krølner and Pors Nielsen 1980) or g cm⁻² (Riggs et al 1981), do not alter any age relationship. Nowhere in our scientific note did we state that the differences illustrated in figure 1 were due to the crossover corrections; we simply stated that 'It is not known whether this discrepancy between the various methods is due to differences in methodology or in the populations studied. Before accepting the latter the possibility of the former must be thoroughly investigated.'

We considered just one technical aspect of BMC measurement by DPA, the crossover correction, and though it was recognised that this would be influenced by both the NaI crystal and scatter within the patient it was considered unnecessary to separate the two components. There is a dearth of information in the literature about methods
used to determine the crossover correction. Mazess et al (1974) state that copper and aluminium were used to filter out the lower energy radiation. Other authors however (Wilson and Madsen 1977, Krohner and Pors Nielsen 1980, Wahner and Dunn 1980, Peppier and Mazess 1981, Riggs et al 1981) do not mention the absorber used to determine the crossover correction. It is not true that aluminium does not preferentially attenuate the 70 keV peak (Mazess et al 1983). Aluminium may be superior to copper but 5 cm of the former will attenuate the 70 keV peak 2.6 times more than the 100 keV peak, hence our rather complicated simulation of the upper end of the $^{153}$Gd spectrum. Our empirical technique provided a satisfactory alternative, but would need to be validated for other apparatus.

Although previous literature of DPA techniques omitted many technical details, contradictions do appear to exist in the crossover correction values. Peppier and Mazess (1981) used a detector collimation of 8 mm x 25 mm and stated that the crossover correction varied with patient thickness from 1.8% to 3.4% whereas Wilson and Madsen (1977), using an even larger detector collimator of 12.5 mm x 37.5 mm, stated that the crossover correction was constant at 3%. Wahner and Dunn (1980) and Riggs et al (1981) used 6 mm detector collimation, the smallest used for DPA, but obtained crossover corrections of 6% and 4–6% respectively. The variation in crossover correction of 4–6% (Riggs et al 1981) was stated to be dependent on the source although no reason is apparent. The high value of 6% (Wahner and Dunn 1980) was due to the thin 1 mm NaI crystal used. If a thicker crystal were used which detected the 100 keV peak efficiently then this value would be reduced to about 2.5%. In the apparatus used by Krohner and Pors Nielsen (1980) however, which has a large detector collimator of 13 mm, a crossover correction of only 1% was used (personal communication).
In our original scientific note (Smith and Tothill 1982) and in this letter we do not intend criticism of any particular method of DPA for vertebral BMC measurements. Explanations may exist for many of the intra-laboratory differences but they need to be investigated with a degree of scientific thoroughness which hitherto has been absent.

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17 March 1983

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Dual photon absorptiometry of the spine with a low activity source of gadolinium 153


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Abstract

Apparatus and data-processing techniques were developed to measure the bone mineral content (BMC) of the lumbar spine. $^{153}$Gd was used as the dual photon source with an activity down to 7 GBq, lower than that adopted by other workers. The compromise between resolution and sensitivity was optimised. A reproducibility in normal subjects of 1.5% (coefficient of variation) was obtained. Normalisation procedures using parameters of the scan image and other indicators of body size such as span were developed, and reduced the coefficient of variation between normal subjects to 10% for men and 8% for pre-menopausal women. The absolute values of BMC are similar to those reported by others. Cross-sectional studies in women demonstrated an accelerated loss of BMC after the menopause.

Absorptiometry is well established as a method for measuring bone mineral. If a uniform overall thickness can be achieved, for example by using a water bath, and a two-phase system assumed, a single photon technique is adequate. Dual photon techniques have been used for some time to allow for varying thicknesses or an extra phase, and more recently have been applied to the study of the spine, the area of trabecular bone of most interest in diseases such as osteoporosis.

The first measurements of bone mineral content (BMC) of the spine Roos and Sköldborn (1974) used the 60 keV radiation from $^{241}$Am and 660 keV gamma rays from $^{137}$Cs. They did not rely on retrospective choice of the site of analysis from a bone mineral image, but identified the scan position from a prior radiograph. Subsequent spine BMC measurements have generally used the radiations from $^{153}$Gd, complex mixtures of X and γ rays effectively giving energies of 44 and 100 keV (Mazess et al., 1974; Wilson & Madsen, 1977; Price et al., 1977; Kroehne & Pros Nielsen, 1980; Wahner & Dunn, 1980; Smith et al., 1983a). These authors have mostly selected the appropriate vertebrae from the processed image.

We set out to develop a scanning and data processing system for dual photon absorptiometry (DPA) of the spine. We were able to obtain only a lower-activity, larger-diameter $^{153}$Gd source than had been used by previous workers and so had to investigate different compromises between sensitivity and resolution. An appropriate collimation having been chosen, measurements were made of spinal BMC in a group of normal volunteers to establish the degree of variation and to provide a baseline against which to judge abnormality. Methods of normalisation of the data for body size were investigated. The results are presented here and compared with others from the literature.

Methods

A standard dual-headed radionuclide scanner (J. & P. Engineering Ltd.) was adapted for transmission scanning. The $^{153}$Gd source was mounted in the lower head and was initially of 18.5 GBq (500 μCi); its activity had decayed to about 7 GBq when the measurements on normal volunteers were made. The source was a disc, diameter 12 mm, in a stainless steel capsule (Amersham International X 92/1). The initial photon output was quoted by the supplier as $7 \times 10^8$ photons/s/steradian at 100 keV. The output at 44 keV was found to be about the same, there being appreciable attenuation in the 0.25 mm thick stainless steel exit face.

The detector was the standard 10 × 5 cm overhead NaI crystal of the scanner, which moved in synchronism with the source. The collimator face separation was 40 cm. The minimum reliable scanner speed of 5 mm s$^{-1}$ was used, with a stepping distance between scans of 4 mm. A variety of single parallel-hole collimators, each of length 5 cm, was fitted to the source for measurements on a phantom, but once the optimum conditions had been determined a collimator of 13 mm diameter was used for all measurements on subjects. The detector collimator was 7.5 cm long and fixed at 20 mm square.

Two pulse-height analysers with window settings 32–56 keV and 75–125 keV were used to select the gamma-ray energies. Some of the results were obtained by recording data on a Digidek tape recorder for off-line transfer to a PDP 12 computer. Later a Cromexcom microcomputer and TV monitor were interfaced to the scanner. Programs were written to correct for dead-time when necessary and the contribution from the higher energy radiation to the lower energy detection channel (the cross-over). The latter arises partly from scatter in the subject and partly from interactions in the detector.
crystal; the correction was found to vary with patient thickness. To derive the cross-over with $^{153}$Gd it is necessary to filter out the lower energy peak; the presence of a low abundance 70 keV peak creates complications (Smith & Tothill, 1982). The corrections were determined for each collimator combination used.

The bone mineral was calculated for each 4 mm cell using the equation derived by Roos and Sköldborn (1974)

$$ M_b = \frac{\mu_{s2} \ln (I_{01}/I_1) - \mu_{s1} \ln (I_{02}/I_2)}{\mu_{s2}\mu_{b2} - \mu_{s1}\mu_{b2}} $$

(1)

where

- $M_b$ = bone mineral mass per unit area
- $\mu$ = mass attenuation coefficient of soft tissue (s) and bone (b) at the lower energy (1) and higher energy (2)
- $I_0$ = intensity of unattenuated beam
- $I$ = intensity of transmitted beam

Re-arranging equation (1),

$$ (\mu_{s1} \ln I_2 - \mu_{s2} \ln I_1) - (\mu_{s1} \ln I_{02} - \mu_{s2} \ln I_{01}) $$

$$ M_b = \frac{\mu_{s1}\mu_{b2} - \mu_{s2}\mu_{b1}}{\mu_{s2}\mu_{b2} - \mu_{s1}\mu_{b1}} $$

(2)

In the soft-tissue area lateral to the spine, $M_b = 0$, and the second bracketed term containing the unattenuated beam intensities is equal to the first bracketed term. That area can be taken to constitute a background level above which the bone mineral values are calculated, and the expression then simplifies to

$$ M_b = \frac{\mu_{s1} \ln I_2 - \mu_{s2} \ln I_1}{\mu_{s2}\mu_{b1} - \mu_{s1}\mu_{b2}} $$

(3)

The equations as presented here assume that soft tissue is homogeneous, whereas in practice it consists of mixtures of lean and adipose tissue which vary between patients. If the thickness of fat were uniform, the only error introduced would be in the values of $\mu_s$ chosen. Such uniformity is most unlikely, and non-uniformity leads to an incorrect choice of baseline (Roos et al, 1980). However, in common with other workers, we ignore the errors introduced by this factor.

The attenuation coefficients used in our calculations were derived from attenuation measurements through water and hydroxyapatite, and were as follows:

$$ \mu_{s1} = 0.218, \mu_{s2} = 0.152, \mu_{b1} = 0.71, \mu_{b2} = 0.21 $$

(all in cm$^{-1}$)

The ratio $\mu_{s2}/\mu_{s1}$ is particularly important, as it was found from phantom measurements that the adoption of an incorrect ratio in the calculations led to an apparent change of bone mineral with water depth. Our ratio of 0.697 is close to the values used by other workers, even though individual values of attenuation coefficients may differ, perhaps due to different assumptions about the composition of soft tissue and bone mineral.

The bone mineral image is displayed on a colour monitor (Fig. 1). The original scan, over an area of about 12 cm wide and 20 cm long (taking about 20 min to perform), is positioned to include part of the pelvis, to facilitate identification of vertebrae on the image. Three lumbar vertebrae are included, cursors being positioned manually at the top of L2 and the bottom of L4. All cells in a vertical column between these limits are summed and a horizontal profile plotted (Fig. 2). Horizontal baselines are defined either side of the vertebrae, their selection adjacent to the bone edge being facilitated by vertical cursors. The bone mineral values are then integrated above the line joining the baselines. The results are expressed as a total in the three vertebrae, as bone mineral per unit length or unit area.

After reproducibility studies had been performed with anthropomorphic phantoms the optimum collimations were then used for repeated measurements on nine normal volunteers aged 21–58 years. Two measurements were made on separate days on each
subject. Each bone mineral image was analysed by three operators.

The $^{153}$Gd source with optimum collimation has been used to measure spine BMC in 24 normal women, aged 21–61 years, and 18 normal men aged 21–58 years. As there is a measure of subjectivity in the selection of the area of interest in the bone mineral image, all scans were analysed by the same operator.

The radiation dose to the skin was measured with a 1 litre ionisation chamber and found to be 140 $\mu$Gy when the $^{153}$Gd activity was 12 GBq.

**RESULTS**

From the experiments on models it was found that the precision improved as the sensitivity was increased by using larger diameter collimators, in spite of the worsening resolution (Table I). The experimental coefficients of variation were derived from 10 repeated measurements on an anthropomorphic phantom. The theoretical precision was based on counting statistics, taking into account the photon outputs at the two energies, the collimator geometries, the thickness of soft tissue, the bone mineral content, the detector efficiencies and the crossover correction. Details of the calculation are given elsewhere (Smith et al, 1983b).

**TABLE I**

**EXPERIMENTAL AND THEORETICAL PRECISION USING AN ANTHROPOMORPHIC MODEL OF A HUMAN SKELETON IN TISSUE-EQUIVALENT RUBBER, 10 MEASUREMENTS. DETECTOR COLLIMATOR FIXED AT 20 x 20 mm**

<table>
<thead>
<tr>
<th>Source collimator diameter, mm</th>
<th>Coefficient of variation, %</th>
<th>Measured</th>
<th>Predicted</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>2.5</td>
<td>2.4</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>2.1</td>
<td>1.8</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>2.0</td>
<td>1.5</td>
<td></td>
</tr>
<tr>
<td>13</td>
<td>1.1</td>
<td>1.2</td>
<td></td>
</tr>
</tbody>
</table>

The results of the repeated measurements on normal subjects, using the collimators that gave the highest precision in phantoms, were calculated as the average coefficients of variation (CV) obtained by dividing the differences between two measurements on each subject, expressed as a percentage of the mean, by 1.128 (Crow et al, 1960). The mean CV for operator 1, who had more experience than the others, was 1.0%, for operator 2, 1.5% and for operator 3, 3.3%.

The BMC values for the 42 normal subjects are summarised in Table II. In column 4 is the total mass of bone mineral in grams (B) in lumbar vertebrae 2, 3 and 4 as defined by the rectangle selected from the bone mineral image. For column 5 the total BMC values were divided by the length (L) of the rectangle in cm (along the spine) to give $B/L$. A further division by the width of the rectangle yielded the figures for BMC per unit area ($B/A$) in column 6. The best reproducibility was found by selecting the boundaries of the rectangle at the edge of the profile peak as illustrated in Fig. 2. The width of the rectangle was then such as to exclude the extremities of the transverse processes. A small amount of bone mineral was therefore included in the soft-tissue background. This leads to an inaccuracy analogous to that arising from non-uniform fat distribution.

The use of $B/L$ or $B/A$ gives a measure of normalisation for body-size, but further relationships were sought, in the hope of minimising the variations in the values for the normal groups. The best correlation for the men and for the women aged 50 or less was between $B/L$ and span. This linear relationship was used to derive formulae to give predicted BMC ($B/L$) for normal subjects.

For women, $B/L = 5.42 \times S - 3.50$, $r = 0.81, p < 0.001$.

For men, $B/L = 5.73 \times S - 4.61$, $r = 0.48, p < 0.05$.

where $S$ = span in metres.
We can define a bone mineral index (BMI) as the ratio of B/L to (B/L)p for individual subjects. The mean values of the BMI are included in the last column of Table II. Normalisation using span was slightly better for B/L than for B.

When BMC was plotted against age for men, no correlation was found. For women there was a progressive loss of BMC with age above about 50 years. For example, B/L is plotted against age in Fig. 3, without normalisation for span. As there are relatively few points and a fair amount of scatter, a degree of smoothing has been obtained by calculating the mean BMC cm⁻¹ for each decade. The same method of presentation, without data points or error bars, is used in Fig. 4A, to illustrate the different normalisation procedures and for comparison with other published results, the means for the first decade being set to 100%.

A further indication of age-related bone loss in women was obtained by correlating BMC with years post-menopause (Y). Only for the ratio BMI was the correlation significant (r = -0.63, p < 0.05, n = 11), the linear regression equation being BMI = 0.995 - 0.017X + Y, corresponding to a rate of loss of 1.7% per annum soon after the menopause.

**DISCUSSION**

The main difference between our measurements of spine BMC and those of others using a similar technique was that the activity of our 153Gd source was much lower, only about 7 MBq when most of the measurements on normal subjects were performed, compared with about 40 GBq in common use. Krohner and Pors Nielsen (1982b) quote the lowest activity employed, considering their source to be usable down to 15 GBq. We had already shown by theoretical analysis and measurements on models that an acceptable precision (around 1%) could be obtained with this lower activity, using a coarser collimation to maintain sensitivity (Smith et al, 1983b). The results presented here confirm this finding in repeated measurements on normal subjects, the coefficients of variation of 1.0% and 1.5% for operators 1 and 2 being close to the theoretical CV based on counting statistics. The CV of 3.3% obtained by operator 3 is less acceptable, but he was inexperienced in the analysis, and we have found that practice brings a rapid improvement in the consistency of area selection.

The acceptability of precision obtained with our activity means that a 20 GBq source could be used for about a year and 50 GBq for twice as long.

Another way in which our source differed was that it had an active diameter of 12 mm. The specific activity of 153Gd is such that self-absorption of the X rays is not a limitation with smaller sources, and other workers have used lower diameters, giving a better resolution. Some have also improved resolution by using a smaller detector collimation. Our demonstration that satisfactory results can be obtained with a 12 mm source is not so important for 153Gd, but is vital if 241Am is considered for the lower energy photon beam. The long half-life of 241Am, which makes it so attractive and economical a source, means that its specific activity is low. It is only possible to obtain sufficient photon output from 241Am to give the required statistical precision if a diameter of at least 12 mm is used (Smith et al, 1983b).

Our measured radiation doses are similar to those
published by others. For example, Krolner and Pors Nielsen (1980) quote a skin dose of 0.1 mGy and Wilson and Madsen (1977) approximately 0.02 mGy. In any case the doses are low enough to permit repeated observations in a wide category of subjects.

The BMC results for normal subjects are presented in Table II in different forms to allow comparison with other published results. Each form has its merits. As Krolner and Pors Nielsen (1980) have pointed out, the total BMC in a number of vertebrae (usually the second, third and fourth) is influenced least by vertebral crush fractures, which compress the vertebrae into a shorter length. On the other hand, no normalisation for body size is provided by this presentation and the consistency of the measurement is most critically dependent on the choice of boundaries between vertebrae. The total height of the three vertebrae is about 100 mm, corresponding to 25 scans at our 4 mm spacing. An inconsistency of boundary selection by one line, which is difficult to avoid, leads to an error of about 4%. This error can be much reduced, and some measure of normalisation for body size obtained, by expressing results as BMC per unit length. The penalty of this method is that crushed vertebrae may thereby appear to have a raised bone mineral content.

An advantage in expressing the result in terms of BMC per unit length is that the length is easily defined and so intercomparison of results using different techniques can be made. This may not be so if the result is expressed in terms of BMC per unit area, because the width of the vertebrae can be expressed in several ways. Our method was chosen to give the best reproducibility of selection of region of interest. Increasing the width of the chosen rectangle will not, beyond a certain level, affect the BMC value but will progressively decrease the BMC per unit area. Some centres do not use a rectangle but an irregular region of interest, calculated using an edge detection programme. Thus though it may be possible for the different techniques to obtain comparable BMC or BMC per unit length values, there is likely to be considerable variation between values of BMC per unit area. This is illustrated in the results shown in Tables II and III.

Using our apparatus, the mean width, which was affected by the relatively coarse collimation, was about 5.5 cm; this is somewhat greater than the average width of the vertebral bodies, but less than the overall width of the transverse processes. By this exclusion we slightly underestimate the total bone mineral, but include all the trabecular bone of the bodies. The selection of a width large enough to encompass the whole bones would reduce the precision and the value of normalisation by width. The use of a finite beam width and scanning speed leads to an additional underestimate of BMC (West & Reed, 1970; Watt, 1973) but in common with others making measurements of the spine, we do not attempt to apply corrections.

Table II shows that the CV for the normal population studied was highest when total BMC was used (column 4). A reduction was obtained, as might be expected, when normalisation by length or area was applied (columns 5 and 6). The least variation was found when the additional factor of span was incorporated to give the predicted BMC per unit length, with which the measured value could be compared (column 7).

Our results can be compared with others from the literature. As all investigators have demonstrated a loss of BMC with age in women, it was thought best to present the mean values for the populations studied, together with the mean ages. These results are summarised in Table III, which can be compared with our Table II.

Roos et al (1980) relied on prior X-radiography to select a single vertebra and performed a single profile scan with a combined $^{241}$Am/$^{137}$Cs beam.

The remaining authors used $^{153}$Gd, but Madsen (1977) did not select the vertebrae from a bone mineral image, relying instead on a fixed scanning length of 10 cm starting at the level of the iliac crests. Riggs et al (1981) used lumbar vertebrae 1–4 and determined the area used for normalisation from an edge-detection programme applied to the images. Krolner and Pors Nielsen (1982a) integrated profiles to obtain total BMC in vertebrae L2–4; although they express their results in arbitrary units with the dimensions of mass, it is clear from an earlier paper (Krolner & Pors Nielsen, 1980) that these units are equivalent to grams of hydroxyapatite.

It is seen from Tables I and II that our results are very similar to those already published.

### TABLE III

<table>
<thead>
<tr>
<th>Authors</th>
<th>Sex</th>
<th>No.</th>
<th>Mean age</th>
<th>Vertebrae studied</th>
<th>B/cm²</th>
<th>B/L/cm²</th>
<th>B/A/cm²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Madsen (1977)</td>
<td>F</td>
<td>41</td>
<td>61</td>
<td>10 cm length</td>
<td>4.5</td>
<td>0.85</td>
<td></td>
</tr>
<tr>
<td>Roos et al (1980)</td>
<td>M</td>
<td>18</td>
<td>63.4</td>
<td>L3</td>
<td>4.5</td>
<td>0.85</td>
<td></td>
</tr>
<tr>
<td>Krolner &amp; Pors Nielsen (1982a,b)</td>
<td>F</td>
<td>70</td>
<td>50.5</td>
<td>L2–4</td>
<td>44.5</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

833
An examination of the loss of bone mineral from the spine with increasing age in women is important, as somewhat different patterns have been reported from the various cross-sectional studies considered here. Madsen (1977) fitted a linear regression to $B/A$ over the whole age range, with a slope of 0.40% per annum at age 60. However, by re-analysing the data from his graph, it can be seen that there is no evidence of a loss below the age of about 50. For women over 50 a linear regression has a correlation coefficient of 0.565, $n = 31$, $p < 0.001$, with a slope of 0.75% per annum at age 60.

Hansson and Roos (1980) measured the bone mineral per unit length in single vertebrae removed at autopsy. They correlated $B/L$ with age using a linear regression, showing a loss of 1.0% per annum at age 60. However, only 4 of the 18 women were below 50 years of age and the data do not permit deductions about possible changes of slope.

Riggs et al. (1981) also used a single linear regression, this time of $B/A$ against age, but justified it statistically, there being much larger numbers and no significant difference in the slope pre- and post-menopause. The rate of loss was 1.0% per annum at age 60.

Krauner and Pors Nielsen (1982a) considered that the distribution of lumbar BMC of normal pre-menopausal women showed skewness, as logarithmic transformation reduced the moment coefficient. However, analysis of their data shows that the skewness was not statistically significant. They found that BMC of pre-menopausal women was not related to age, whereas a significant reduction was found in the post-menopausal group, with an annual loss of 1.4%. They investigated several linear and polynomial regression equations, but chose to fit data to a gamma variate function, although a better fit was not obtained thereby. The function serves well enough at higher ages, but has the disadvantage that it introduces a maximum in the curve at about the age of 35. This is not in accord with the data and it seems a pity to distort the possible conclusions.

The data plotted in the four publications reviewed above have been analysed in the same way as our own data, mean values of BMC for each decade of age being plotted against the mean age in Fig. 4b. Our own results, such as those presented in Figs. 3 and 4a, do not contain sufficient numbers or cover a wide enough age range for detailed analysis, but support the view that there is an increased loss of lumbar bone mineral after the menopause. Fig. 4a demonstrates that the normalisation procedure adopted does not affect the pattern of loss, suggesting that differences in Fig. 4b are due to some other cause.

When sufficient normal women have been measured, it should be possible to define the predicted normal lumbar BMC for a given age and size, as has been done for total body calcium (Cohn et al., 1976; Kennedy et al., 1982). The annual losses of lumbar bone mineral in post-menopausal women summarised here are not greatly different from those reported for total body calcium, 1.1% (Cohn et al., 1976) and 1.5% (Kennedy et al., 1982).

The age-dependence of lumbar bone mineral in men has been reported by fewer authors. Hansson and Roos (1980), again using autopsy specimens from 15 subjects, found the slope of a linear regression of $B/L$ against age to be the same as for women in terms of g cm$^{-2}$. Taking into account the higher mean BMC of the men, the proportional loss was lower, 0.76% per annum at age 60.

Riggs et al. (1981) analysed the variation of lumbar BMC with age in 82 men. They found a small loss, of less than 0.2% per annum, with a significance of $p < 0.05$. With smaller numbers studied and a more restricted age range, it is not surprising that we demonstrated no change with age. Once again these results are similar to those from total body calcium determinations (Kennedy et al., 1982).

ACKNOWLEDGMENTS

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**Book review**


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For many years tumour tissues have been known to localise porphyrins, although the biochemical reasons for this are not understood. Nevertheless this finding is being exploited clinically both by using haematoporphyrins as photosensitisers as a means of eradication of tumours (PRT) and to aid tumour localisation. In 1981 a workshop was held in which workers involved in fundamental studies on porphyrin chemistry, physics and photobiology, along with those using porphyrins clinically, were brought together. This book contains the contributions of the participants and is an extremely useful source of information on the topic for all scientists and clinicians interested in this field of knowledge.

A series of articles on clinical aspects of haematoporphyrin photoradiation therapy is followed by a paper concerning the dosimetry of light in tissue by Doiron et al and a general review of photobiological considerations in photoradiation therapy by J. A. Parrish, both of which should be of widespread interest. The latter half of the book includes papers on the photochemical and photobiological mechanism of action of haematoporphyrin derivatives, including a report of a study by Michael Berns and co-workers using fluorescent analysis of cells with a laser light source. Each chapter ends with a useful list of references, so that people with no prior knowledge of the subject can use this as an original source of information.

D. J. G. Davies.
Normal levels of total body sodium and chlorine by neutron activation analysis

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Abstract. In vivo neutron activation analysis was used to measure total body sodium and chlorine in 18 male and 18 female normal adults. Corrections for body size were developed. Normalisation factors were derived which enable the prediction of the normal levels of sodium and chlorine in a subject. The coefficient of variation of normalised sodium was 5.9% in men and 6.9% in women, and of normalised chlorine 9.3% in men and 5.5% in women. In the range examined (40–70 years) no significant age dependence was observed for either element. Total body sodium was correlated with total body chlorine and total body calcium. Sodium excess, defined as the amount of body sodium in excess of that associated with chlorine, also correlated well with total body calcium. In females there was a mean annual loss of sodium excess of 1.2% after the menopause, similar to the loss of calcium.

1. Introduction

Total body neutron activation analysis can be used to determine the absolute amounts of certain body elements and also changes in these levels with time. At present, elements which may be measured by this technique include calcium, sodium, chlorine, phosphorus and nitrogen.

Absolute levels of total body sodium (TBNa) with or without total body chlorine (tBCl) in normal adults have been determined using neutron activation analysis by Chamberlain et al (1968), Rudd et al (1972), Ellis et al (1976) and Aloia et al (1980). Where appropriate the results of these studies and the data presented in this paper will be compared and discussed. TBNa has been shown to change in several disease states, e.g., renal failure (Cohn et al 1972), hypertension (Boddy et al 1978) and osteomalacia (Hosking et al 1972), and so techniques which allow the accurate evaluation of TBNa and tBCl are of considerable interest. For these to be of diagnostic significance it is necessary to establish normal ranges and develop normalisation procedures.

Most of the sodium in the body is extracellular, fairly rapidly exchangeable and associated with an equivalent amount of chlorine. The sodium in bone exchanges less rapidly and the chlorine concentration is much lower. The presence of a slowly-exchangeable pool of sodium has been demonstrated by the differences between the

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values for exchangeable sodium as measured by isotope dilution techniques and total body sodium as measured by chemical analysis (Forbes and Lewis 1956) or by neutron activation analysis (Chamberlain et al. 1968, Boddy et al. 1978).

2. Methods

In the technique, the subjects are exposed to a beam of partially moderated fast neutrons which induces amongst others the reactions, $^{23}$Na$(n, \gamma)^{24}$Na, $^{37}$Cl$(n, \gamma)^{38}$Cl, $^{48}$Ca$(n, \gamma)^{49}$Ca, $^{31}$P$(n, \alpha)^{28}$Al and $^{14}$N$(n, 2n)^{13}$N. The neutron source used for patient activation was a cyclotron producing neutrons of mean energy 6.5 MeV (Williams et al. 1979). During irradiation the subject was positioned in a moderator kiosk which optimises the uniformity of activation of sodium and chlorine. The subject was then transferred to a shadow-shield whole-body counter where the $\gamma$-ray spectrum, covering the energy range 0 to 4 MeV, was recorded on a multichannel analyser. The data were analysed by fitting five standard spectra obtained from the activation of sodium, chlorine, calcium, phosphorus and potassium to the smoothed subject spectrum in the energy range 1.0 to 3.4 MeV, using a least squares technique (Smith and Tothill 1979).

Corrections to eliminate the effects of variations in activation and detection efficiency with body size and position were then made, allowing the $TBNa$ and $TBCl$ content in grams to be determined accurately.

The technique of total body neutron activation analysis used in this study has been described in detail, with reference to measurements of calcium, by Kennedy et al. (1982). The corrections were determined for each element using phantoms of several sizes, but there were no significant differences between those for calcium, sodium and chlorine. The coefficient of variation (cv) for repeated measurements of an anthropomorphic phantom was 2.1% for sodium and 4.3% for chlorine for an absorbed dose equivalent of 13 mSv. The phantom contained known amounts of sodium, chlorine, calcium, phosphorus, potassium and nitrogen, and there was no significant dependence of the counts obtained per gram of element on the ratio of the various elements in the phantom. Although no attempt was made in this study to determine total body nitrogen levels, a physiological amount of nitrogen (160 g) was contained in the phantom. The importance of including nitrogen has recently been emphasised by Spinks and Bewley (1982).

Although this technique requires little patient preparation, it is important to check that the subject is not wearing thermal underwear which may contain significant levels of chlorine.

Since the chlorine space approximates extracellular fluid volume and the ratio of Na to Cl in the extracellular fluid has been estimated, the amount of sodium in excess of that associated with the chlorine space may be defined as the sodium excess value, $Na_{ES}$ (after Ellis et al. 1976) and can be expressed in grams as

$$Na_{ES} = TBNa - 0.78 TBCl$$

where $TBNa$ and $TBCl$ are in grams. It is assumed that $Na_{ES}$ represents most of the sodium resident in bone.

3. Results

3.1. Normal values

$TBNa$ and $TBCl$ measurements were made on a group of 36 normal volunteers, 18 male and 18 female, between 40 and 70 years of age. All the subjects were active
Normal levels of total body sodium and chlorine

and in good health. None had any history of disease that might be expected
influence their TBNa and TBCl levels, and all had normal blood and urine biochemistry.
There was a considerable spread in the TBNa and TBCl results for both groups (see
Table 1). For the males the TBNa ranged from 74.8 g to 99.4 g with a mean of
83.7 g±8.2% (cv) while the TBCl ranged from 61.8 g to 93.4 g with a mean of
71.7 g±11.3%. The TBNa for the females ranged from 55.1 g to 75.9 g with a mean
of 62.0 g±9.2% while the TBCl ranged from 46.5 g to 65.8 g with a mean of 53.4 g±
10.2%.

<table>
<thead>
<tr>
<th>Mean (cv)</th>
<th>Male</th>
<th>Female</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td>18</td>
<td>18</td>
</tr>
<tr>
<td>Height (m)</td>
<td>178.3 (4.8)</td>
<td>160.9 (4.0)</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>81.3 (12.0)</td>
<td>62.5 (17.7)</td>
</tr>
<tr>
<td>Lean body mass (kg)</td>
<td>59.3 (11.7)</td>
<td>39.9 (12.6)</td>
</tr>
<tr>
<td>Age (y)</td>
<td>54.3 (16.5)</td>
<td>56.7 (10.9)</td>
</tr>
<tr>
<td>Sodium (g)</td>
<td>83.7 (8.2)</td>
<td>62.0 (9.2)</td>
</tr>
<tr>
<td>Chlorine (g)</td>
<td>71.7 (11.3)</td>
<td>53.4 (10.2)</td>
</tr>
<tr>
<td>Sodium excess NaES (g)</td>
<td>27.7 (12.2)</td>
<td>20.5 (19.9)</td>
</tr>
<tr>
<td>NaES/kg calcium (g)</td>
<td>24.4 (9.1)</td>
<td>24.9 (7.5)</td>
</tr>
</tbody>
</table>

3.2. Normalisation procedures

As the total composition of the various elements in the body is related to the subject’s
build, the dependence of the measured TBNa and TBCl on height, weight, lean body
mass (LBM) (calculated from the weight and skinfold thickness measurements), age
and number of years post-menopause was investigated by multiple linear regression
analysis.

For males, this gave a formula for predicted total body sodium (TBNap) in grams
\[ TBNap = 19.2 \times (LBM)^{0.26} \times (\text{height})^{0.71} \]  
\( r = 0.72, P < 0.001 \)

and predicted total body chlorine (TBClp) in grams
\[ TBClp = 11.3 \times (LBM)^{0.37} \times (\text{height})^{0.57} \]  
\( r = 0.58, P < 0.05 \).

Height is measured in metres and LBM in kilograms. Similar equations were obtained
when weight was used instead of LBM.

\[ TBNap = 19.1 \times (\text{weight})^{0.23} \times (\text{height})^{0.81} \]  
\( r = 0.69, P < 0.001 \)

\[ TBClp = 8.9 \times (\text{weight})^{0.40} \times (\text{height})^{0.56} \]  
\( r = 0.60, P < 0.05 \).

No significant dependence of TBNa and TBCl on age was observed. The biological
variation in normal TBNa and TBCl due to body build could be reduced by expressing
the results in terms of the ratios TBNa/TBNap and TBCl/TBClp. This reduced the CV
in the male controls to 5.8% for sodium and 9.2% for chlorine.

For the female controls, multiple regression analysis gave,

\[ TBNap = 13.1 \times (LBM)^{0.34} \times (\text{height})^{0.63} \]  
\( r = 0.72, P < 0.001 \)

\[ TBClp = 5.2 \times (LBM)^{0.59} \times (\text{height})^{0.33} \]  
\( r = 0.84, P < 0.001 \).
As with the male controls, similar equations were obtained when weight was substituted for LBM

\[ \text{TBN}_p = 19.1(\text{weight})^{0.16}(\text{height})^{1.09} \quad (r = 0.72, P < 0.001) \]
\[ \text{TBCl}_p = 9.8(\text{weight})^{0.31}(\text{height})^{0.88} \quad (r = 0.81, P < 0.001). \]

No significant dependence on age was observed for either element. Using the ratios \( \text{TBN}_a/\text{TBN}_p \) and \( \text{TBCl}/\text{TBCl}_p \) the CV for sodium was 6.9% and for chlorine was 5.5%.

The \( \text{TBN}_a \) and \( \text{TBCl} \) results for both male and female subjects are presented in figure 1. There was a good correlation between the \( \text{TBN}_a \) and the \( \text{TBCl} \) results for males \( (r = 0.89, P < 0.001) \) and females \( (r = 0.70, P < 0.005) \) and when the sexes were combined \( (r = 0.94, P < 0.001) \).

![Figure 1. Relationship between total body sodium and total body chlorine. The regression line is calculated from male and female data combined.](image)

Total body sodium was also found to be dependent on the subject's total body calcium (TBCa) \( (r = 0.67, P < 0.01) \). The TBCa results for these subjects were reported by Kennedy et al (1982). Significant dependence for TBCl on TBCa was found only for men \( (r = 0.49, P < 0.05) \).

There was a correlation between NaES and TBCa (figure 2). The relationship derived was

\[ \text{NaES} = 19.9 \text{ TBCa} + 4.5 \quad (r = 0.79, P < 0.001) \]

where NaES is measured in g and TBCa in kg, and the mean NaES per kg of TBCa was found to be 24.4 g for males and 24.9 g for females.

Although no correlation was found between TBNa and age, NaES was found to be correlated with years post-menopause in females \( (r = 0.50, P < 0.05) \) with an annual loss of NaES after the menopause of 1.2%. No significant dependence of NaES on age was observed for the male volunteers.

4. Discussion

The mean values of TBNa g/kg body weight for males, 1.03, and females, 0.99, are in good agreement with those values determined by Ellis et al (1976) of 1.08 for
Normal levels of total body sodium and chlorine

Figure 2. Relationship between sodium excess and total body calcium. The regression line is calculated from male and female data combined.

males and 0.98 for females. However the mean values of tbCl g/kg body weight for males, 0.88, and females, 0.85, are lower than those of Ellis et al (1976) of 0.99 for both sexes.

No dependence of either tbNa or tbCl on age was observed for either sex. Over the age range covered by this study, 40 to 70 years, this is largely in agreement with Ellis et al (1976), which is the most comprehensive of the other studies, although they did observe a slight decrease in tbNa in elderly females. Even when the varying body build of the subjects in this study was taken into account by plotting $\frac{tbNa}{tbNa_p}$ and $\frac{tbCl}{tbCl_p}$ against age, no significant correlation was obtained for either sex. The annual loss of Na$_{ES}$ after the menopause of 1.2% is not significantly different from the loss of 1.5% per annum in TbCa in the same subjects (Kennedy et al 1982) and implies that an approximately equal percentage of calcium and sodium is lost from bone per year after the menopause. The lack of significant dependence of Na$_{ES}$ on age for the male volunteers agrees with a similar result for TbCa.

The mean level of 24.3 g of Na$_{ES}$ per kg of TbCa for males agrees well with that determined by Ellis et al (1976), for a similar age range, of 24.0 g/kg TbCa. However, their mean value of Na$_{ES}$ for females, aged 40–70 y of 15.3 g/kg TbCa, and that of a later study by the same group, of 16.0 g/kg TbCa for females aged 50 to 59 y (Aloia et al 1980) are considerably lower than the 24.9 g/kg TbCa of our study. Ellis et al do not comment on the surprising difference they found between the sexes of the relationship of sodium excess to total body calcium.

The results from this study imply that the sodium excess represents around 33% of the tbNa for both male and female normal volunteers. This compares with values of 27% for males by Rudd et al (1972), 18% for males by Chamberlain et al (1972) and 27% for males and 23% for females by Ellis et al (1976).

The total body levels of sodium and chlorine determined in this study are in broad agreement with other studies, e.g., Anderson et al (1968), Rudd et al (1972) and Ellis et al (1976), and again suggest that the levels quoted by the International Commission
on Radiological Protection (ICRP 23) of 100 g for a 70 kg man for sodium and 95 g for chlorine are overestimates.

The narrow ranges for normal subjects that we have observed and the equations for predicting $\text{tnNa}$ and $\text{tnCl}$ should allow the technique of total body neutron activation analysis of sodium and chlorine to be a useful diagnostic procedure.

Acknowledgments

We should like to thank Professor J A Strong, Dr D Reid, Dr J D Simpson, Mr C M Ferrington, Mrs E M Law and the staff of the MRC Cyclotron Unit, Edinburgh. The research was partly funded by the Scottish Home and Health Department and the Medical Research Council.

Résumé

Niveaux normaux du sodium et du chlore du corps entier par analyse par activation neutronique.

Nous avons utilisé l'analyse par activation neutronique in vivo pour mesurer le sodium et le chlore du corps entier chez des adultes normaux (18 hommes et 18 femmes). Nous avons introduit des corrections tenant compte des dimensions du corps. Nous en avons déduit des facteurs de normalisation permettant de prédire les niveaux normaux du sodium et du chlore chez un sujet. Le coefficient de variation du sodium normalisé est de 5.9% chez l'homme et 6.9% chez la femme, et celui du chlore normalisé est de 9.3% chez l'homme et de 5.5% chez la femme. Pour les deux éléments, nous n'avons observé aucune dépendance significative vis à vis de l'âge dans la tranche d'âge examinée (40-70 ans). Le sodium du corps entier est corrélé avec le chlore et le calcium du corps entier. L'excès de sodium, défini comme la quantité de sodium du corps en excédant par rapport à celui associé au chlore, est aussi bien corrélé avec le calcium du corps entier. Chez les femmes, on observe une perte moyenne annuelle du sodium en excès de 1.2% après la ménopause, similaire à la perte en sodium.

Zusammenfassung

Normaler Ganzkörper-Natrium- und -Chlorspiegel mit Hilfe der Neutronenaktivierungsanalyse.


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Normal levels of total body sodium and chlorine

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The Assessment of Postmenopausal Osteoporosis by Total Body Neutron Activation Analysis

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Abstract

Total body calcium (TBCa) was measured using a cyclotron for in vivo neutron activation analysis (IVNAA) in 20 healthy women, 15 women with vertebral compression fractures, and 8 women with wrist fractures. The precision of the technique, using phantoms, was 1.8% for a dose of 13 mSv. A formula for predicted TBCa (TBCap) was derived from the 20 normal women based on span and years postmenopause. The coefficient of variation of TBCa after normalization in the normal women was 6.8%. The mean TBCa values for the vertebral and wrist fracture groups were 69% and 84% of TBCap for women at the time of the menopause. The low TBCa in the wrist fracture group was attributable to postmenopausal bone loss. Of the low TBCa in the vertebral fracture group, about half the loss could be attributed to postmenopausal age and half to other factors.

Key Words: Activation—Analysis—Osteoporosis.

Introduction

In vivo neutron activation analysis (IVNAA) is an established technique for measuring total body calcium (TBCa) and has been described by five other groups (Chamberlain et al., 1968; Cohn et al., 1976a; Palmer et al., 1968; Spinks et al., 1977; Kennedy et al., 1979) as well as by ourselves (Kennedy et al., 1982). Since 99% of TBCa is present in bone (Heany, 1963), IVNAA can be used to measure bone mass and in this way avoid the errors in sampling from different parts of the skeleton. Two other groups have reported TBCa in women with postmenopausal osteoporosis (Chesnut et al., 1977, Cohn et al., 1974).

In this report we describe a formula for predicting TBCa from the span and postmenopausal age based on a study of 20 normal women. This formula is then used to compare the bone mass of women with vertebral or wrist fractures with their expected bone mass. The degree of osteopenia associated with these two types of fracture is also calculated.

Patients and Methods

Twenty women aged 46–66 years were selected from volunteers who answered requests circulated in the hospital, in a factory, in the local press, and on BBC Radio Scotland. The criteria for selection were based on age (40–70 years) and the absence of a past history of back pain, fragility fractures, artificial menopause, rheumatic, hepatic, renal, or endocrine diseases, exposure to steroid or anticonvulsant therapy, or excessive alcohol intake. Screening tests to exclude occult malabsorption and disorders of calcium metabolism or of renal or hepatic function were undertaken as follows: serum or plasma calcium, phosphate, magnesium, alkaline phosphatase, albumin, parathyroid hormone, 25-hydroxyvitamin D, iron, iron-binding capacity, ferritin, vitamin B12 and folate, urinary calcium, phosphate, magnesium, hydroxyproline and creatinine clearance, and a full blood count.

Eight women with wrist fracture, aged 56–69 years, were selected from a consecutive series of patients attending the orthopedic clinic. Selection was made on the same basis as for the volunteers. Measurements of TBCa and of plasma and urine as for the normal women were made 1 year after the fracture.

Fifteen women aged 48–71 years were referred to the endocrine clinic for medical therapy of their vertebral osteoporosis. Back pain had been present for less than 2 years in 11 of the 15 women. At least two collapsed vertebral were seen on a lateral radiograph of the thoracolumbar spine in all patients. No selection was made of these patients, and the same investigations as already described were performed together with iliac bone scans and bone biopsy when indicated.

TBCa was measured by IVNAA using the MRC cyclotron in Edinburgh (Williams et al., 1979) as the source of neutrons. The induced radiation was measured using four sodium iodide detectors mounted in a whole-body counter. Computer analysis of the spectrum and the value of a standard irradiated simultaneously with the subjects gave the TBCa in Ca counts. The long-term precision, based on anthropomorphic phantom measurements, was 1.8% (coefficient of variation, CV) for a dose of 13 mSv (1.3 rem) (Kennedy et al., 1982). The effect of measurement efficiency due to variation in body size was determined experimentally. Calcium counts were converted to grams of calcium using a correction formula from phantom measurements. Biologic variation due to size and age was then reduced using multiple regression analysis of height, arm span, weight, age and years postmenopause, and lean body mass (% body fat was obtained using skin calipers) (Durnin and Womersley, 1974). The factors relating most closely to TBCa were found to be arm span and years postmenopause.

Informed consent was obtained from all patients. The study protocol was approved by the Hospital Ethics Committee and by ARSAC (Administration of Radiocactive Substances Advisory Committee).
Results

The TBCa in normal women was 820 ± 124 g (mean ± standard deviation, SD). This gave a CV of 15.1% before any allowance was made for span or postmenopausal age. The greatest reduction in variance was achieved using the following formula to obtain the predicted TBCa (TBCaρ):

\[ TBCaρ = \alpha s^{1.68} e^{-0.10y} (r = 0.90, P < 0.001) \]

where:

\[ TBCaρ \] is in grams
\[ \alpha = 399 \]
\[ s = \text{arm span in meters} \]
\[ y = \text{years postmenopause} \]

The arm span not only related more closely to TBCa than height but also eliminated the error due to loss of height associated with vertebral fractures. The formula applied to perimenopausal women and to women up to 22 years postmenopause. The regression is based on an exponential loss of bone mass starting at the time of the menopause. The estimated rate of bone loss was 1.59% per year. The ratio TBCa/TBCaρ has been defined by Cohn et al. as the calcium ratio (CaR) (Cohn et al., 1974) and for our normal women was 0.87 with a CV of 6.8%.

The TBCa in women with vertebral and wrist fractures was 149 ± 87 g and 787 ± 72 g, respectively (mean ± SD). The values of span and postmenopausal age are shown in Table 1. In the women with vertebral fractures the CaR was 1.00 ± 0.10, which was normal (Fig. 1). Six of the 15 women with vertebral fracture had postmenopausal ages greater than 22 years. This was the greatest postmenopausal age among the normal women; the formula for TBCaρ cannot be extrapolated to cover the women in this group who were more than 22 years past the menopause. In the remaining 9 women the CaR was 0.87 ± 0.06, significantly lower than in normal women (P < 0.01). Factors that may have contributed to this excessive bone loss were identified in 5 of the 9 patients, namely, prednisolone therapy for polymyalgia rheumatica, phenytoin therapy for epilepsy, previous history of thyrotoxicosis, atrophic gastritis, and moderate impairment of renal function (creatinine clearance 20 ml/min).

When TBCa was normalized for span alone (TBCa/s1.68), i.e., y = 0 in the equation for TBCaρ, an estimate of osteopenia in the patients with fractures could be made by comparing the values with the CaR of the normal women (Fig. 2). The CaR/s1.68 for women with vertebral and wrist fractures and for normal women were 0.69 ± 0.06, 0.84 ± 0.08, and 0.8 ± 0.11, respectively (mean ± SD). All the women with vertebral fractures had values less than 0.78, but 3 of the 8 women with wrist fractures and 5 of the 20 normal women also had values below 0.78.

Table 1. Scan and postmenopausal age in normal women and women with vertebral and wrist fractures. The years postmenopause in the vertebral fracture group were significantly higher than in the normal women (P < 0.002).

<table>
<thead>
<tr>
<th>Group</th>
<th>Number</th>
<th>Span (m) Mean ± SD</th>
<th>Postmenopausal Age (yr) Mean ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal women</td>
<td>20</td>
<td>1.65 ± 0.07</td>
<td>8.9 ± 7.0</td>
</tr>
<tr>
<td>Vertebral</td>
<td>15</td>
<td>1.66 ± 0.07</td>
<td>21.9 ± 8.4</td>
</tr>
<tr>
<td>Wrist fracture</td>
<td>8</td>
<td>1.66 ± 0.06</td>
<td>12.0 ± 5.9</td>
</tr>
</tbody>
</table>

Fig. 1. Calcium ratio (CaR) in normal women (○), women with wrist fracture (△), and women with vertebral fracture (□). Horizontal lines are mean CaR = 2 SD for normal women.
Discussion

TBCa for the normal women (820 ± 124 g) was similar to that reported by Cohn et al. (1976b), namely, 804 ± 106 g for normal women aged 50–59 years. The TBCa for women with vertebral fractures (649 ± 87 g) resembled that described by Chesnut et al. (1977) (mean 688 g ± 94 g) and Cohn et al. (1974) (mean 590 g ± 108 g). However, some of the patients in the latter two groups did not have vertebral compression fractures but only demineralized vertebral.

Different formulae for TBCa, have been reported. Cohn et al. (1976b) use age from birth, height, and total body potassium: the latter two also may be age dependent. Nelp et al. (1972) predict TBCa from the cube of the patient's height. In the present report it was noted that span was more effective than height in reducing variance, and we would prefer to use span, as it is unaffected by the height loss due to vertebral compression fractures.

The CV of TBCa, in normal women was 6.6% as compared with 7.1% reported by Cohn et al. (1976b). The latter group found that the rate of bone loss after the age of 55 years was 1.1% per year compared with the rate of 1.5% per year after the menopause in our group.

The Ca, was 0.57 in the women with vertebral fractures, similar to the value of 0.82 found by Cohn et al. (1974). An early menopause in these two groups of women may explain in part their osteopenia. Additional factors, including a low TBCa at the time of the menopause or an accelerated loss of bone after the menopause, must also have been present. The normal Ca, in the women with wrist fractures suggests that they were not specifically prone to fracture compared with other postmenopausal women.

Expressions such as TBCa/81 as1.26 to describe the degree of osteopenia have not been reported previously. Although the sample size of 15 was small, in this group of women vertebral crush fractures occurred when total body calcium was below 80% of that predicted from span alone, i.e., TBCa/81 less than 0.79. However there was some overlap of these values with the other groups of women. It may be possible, therefore, to predict those women likely to sustain a vertebral fracture and to recommend treatment, estrogens, for example, aimed at preventing further loss of bone mass.

Acknowledgement. We would like to thank our orthopedic colleagues at the Western General Hospital for allowing us to study their patients.

We also thank the volunteers, in particular the WRVS, Mrs. E. Law and the MRC cyclotron staff assisted with patient measurements.

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RÉSUMÉ

Le calcium corporel total (Ca CT) a été mesuré par activation neutronique in vivo avec l'aide d'un cyclotron chez 20 femmes en bonne santé, 15 femmes ayant un tassement vertébral et 8 femmes atteintes de fracture de l'extrémité inférieure du radius. La précision de la technique, évaluée sur des fantômes, est de 1,8 pour cent pour une dose de 13 mSv. Une formule permettant de prédire le CaCT à partir du délai écoulé depuis la ménopause a été calculée pour les 26 femmes normales. Le coefficient de variation du CaCT après normalisation est de 5,6 pour cent dans la population témoign. Les valeurs moyennes du CaCT dans le groupe avec fracture vertébrale et dans celui avec fracture du radius étaient respectivement égales à 69 et 84 pour cent du CaCT des femmes au moment de leur ménopause. Le CaCT réduit dans le groupe avec fracture du radius peut être attribué à la perte osseuse post-ménopausique. La réduction du CaCT dans le groupe avec tassements vertébraux peut être imputée pour moitié environ à l'état post-ménopausique et pour moitié à d'autres facteurs.

Résumé

Le calcium corporel total (Ca CT) a été mesuré par activation neutronique in vivo avec l'aide d'un cyclotron chez 20 femmes en bonne santé, 15 femmes ayant un tassement vertébral et 8 femmes atteintes de fracture de l'extrémité inférieure du radius. La précision de la technique, évaluée sur des fantômes, est de 1,8 pour cent pour une dose de 13 mSv. Une formule permettant de prédire le CaCT à partir du délai écoulé depuis la ménopause a été calculée pour les 26 femmes normales. Le coefficient de variation du CaCT après normalisation est de 5,6 pour cent dans la population témoign. Les valeurs moyennes du CaCT dans le groupe avec fracture vertébrale et dans celui avec fracture du radius étaient respectivement égales à 69 et 84 pour cent du CaCT des femmes au moment de leur ménopause. Le CaCT réduit dans le groupe avec fracture du radius peut être attribué à la perte osseuse post-ménopausique. La réduction du CaCT dans le groupe avec tassements vertébraux peut être imputée pour moitié environ à l'état post-ménopausique et pour moitié à d'autres facteurs.
Bone mass in nodal primary generalised osteoarthrosis

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SUMMARY Previous studies of patients with primary osteoarthrosis of the hip have suggested an increase in bone mass compared with control populations. Nodal primary generalised osteoarthrosis is known to have a strong familial tendency. To test the hypothesis that this tendency might also lead to increased bone mass, total body calcium has been measured by in-vivo neutron activation analysis and cortical area calculated from measurements of metacarpal indices in 15 female patients with primary generalised osteoarthrosis. The results have been compared with those from 12 healthy controls matched for age, menopausal status, and skeletal size. No significant differences were noted in the total body calcium or cortical area measurements between the 2 groups either before or after correction for skeletal size and menopausal status. No relationship was found between the grade of radiological osteoarthrosis in the hand and either bone mass parameter. Bone mass would not appear to be an important factor in the aetiopathogenesis of nodal primary generalised osteoarthrosis.

Osteoarthrosis (OA) is not a single disease but rather a pattern of biomechanical failure of joints which may be secondary to a variety of disorders of bone or articular cartilage. Although most theories of the pathogenesis of OA are based on primary alterations in the articular cartilage,7 Radin et al.4 have suggested that the progressive wear of fibrillated articular cartilage seen in 'primary' OA results from stiffening of the subchondral bone. Clinical support for such a hypothesis comes from the observation that pathological changes of OA are unusual in femoral heads removed from patients with fractured necks of femur2 and that bone mass appears to be increased in patients with primary OA of the hip when assessed by measurements of metacarpal indices6 or photon absorptiometry.7

To examine this hypothesis further we have assessed bone mass in female patients with nodal primary generalised osteoarthrosis by measuring total body calcium and metacarpal indices and comparing the results with controls matched for age, skeletal size, and menopausal status.

Patients and methods

Fifteen female patients with nodal primary generalised osteoarthrosis (PGOA) fulfilling the criteria defined by Kellgren and Moore9 and 12 asymptomatic healthy women matched for age, skeletal size, and menopausal status have been studied. Serum calcium, phosphate, alkaline phosphatase, and albumin were measured in all subjects by standard methods, and persons with abnormalities of calcium metabolism or medical conditions known to be associated with secondary osteoporosis were excluded.

Total body calcium (TBCa) was measured by in-vivo neutron activation analysis. Patients were irradiated for 20 seconds from front and rear, while standing in a rigid polyethylene activation enclosure, by means of neutrons from the Edinburgh Medical Research Council Cyclotron. The patients were then transferred to a shadow-shield whole-body counter, where the gamma radiation from calcium 43Ca induced from stable 47Ca by neutron capture, was measured for 20 minutes. The patient's TBCa in grams was calculated by comparison with the energy spectrum from an activated anthropomorphic phantom of human dimensions containing a known quantity of calcium. Repeated measurements of the phantom gave a long term precision of 1.8% for a radiation dose of 13 mSv (1.3 rem).9 The mean TBCa ± SD in the control population was 842 ± 142 g. Individual results were expressed both in grams and
Table 1 Mean age, menopausal status, span, height, and osteoarthrosis score in patients with PGOA and controls; range given in parentheses

<table>
<thead>
<tr>
<th>Age (years)</th>
<th>PGOA n = 15</th>
<th>Controls n = 12</th>
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<tbody>
<tr>
<td>38-5</td>
<td>55-5</td>
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<tr>
<td>(47-68)</td>
<td>(46-65)</td>
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<th>Years after menopause</th>
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<tbody>
<tr>
<td>8-5</td>
<td>6-9</td>
<td></td>
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<tr>
<td>(0-20)</td>
<td>(0-22)</td>
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<th>Arm span (cm)</th>
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<th>Controls n = 12</th>
</tr>
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<tr>
<td>166-2</td>
<td>165-1</td>
<td></td>
</tr>
<tr>
<td>(155-181)</td>
<td>(150-175)</td>
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</tbody>
</table>

<table>
<thead>
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<th>Height (cm)</th>
<th>PGOA n = 15</th>
<th>Controls n = 12</th>
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<tr>
<td>162-2</td>
<td>159-8</td>
<td></td>
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<td>(156-5-170)</td>
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<th>Osteoarthrosis score</th>
<th>PGOA n = 15</th>
<th>Controls n = 12</th>
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<tr>
<td>36</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>(11-64)</td>
<td>(0-9)</td>
<td></td>
</tr>
</tbody>
</table>

Discussion

These studies suggest that there are no significant differences in total bone mass (measured by TBCa) or local bone mass (measured by cortical area) in patients with PGOA compared with matched controls. Statistical analysis of the data shows that mean differences in TBCa normalised for skeletal size and menopausal status ≥7.2% and cortical area ≥9.9% would have been significant at the 95% level in groups of this size.

Previous studies have only measured bone mass in patients with primary osteoarthrosis of the hip. Foss and Byers found an increase in periosteal diameter (external cortical diameter) and cortical area in primary OA of the hip compared with an age matched normal range, but the differences in females may well have been attributable to increased skeletal size. More recent studies have failed to confirm such increases in bone density using measurements of cortical area and cortical thickness or cortical area related to cross-sectional area.

Table 2 TBCa expressed in grams and as a percentage of normal and cortical area in PGOA patients and controls. Mean values ± SD with the range given in parentheses

<table>
<thead>
<tr>
<th>TBCa in grams</th>
<th>PGOA n = 15</th>
<th>Controls n = 12</th>
</tr>
</thead>
<tbody>
<tr>
<td>836-6±129-9</td>
<td>842±142-7</td>
<td></td>
</tr>
<tr>
<td>(590-2-1053-7)</td>
<td>(596-0-1036-1)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>TBCa % of normal</th>
<th>PGOA n = 15</th>
<th>Controls n = 12</th>
</tr>
</thead>
<tbody>
<tr>
<td>69±5-11-1</td>
<td>100±6-31</td>
<td></td>
</tr>
<tr>
<td>(83-0-120-1)</td>
<td>(88-7-110-4)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Cortical area in mm²</th>
<th>PGOA n = 15</th>
<th>Controls n = 12</th>
</tr>
</thead>
<tbody>
<tr>
<td>47-0±9-4</td>
<td>46-3±5-2</td>
<td></td>
</tr>
<tr>
<td>(36-1±5-6)</td>
<td>(37-5±5-9)</td>
<td></td>
</tr>
</tbody>
</table>
Photon absorptiometric methods have been used to estimate metacarpal and radial bone mineral content in patients with primary OA of the hip. The results have been conflicting: one study appeared to show an increase in bone mineral content of 13% at cortical and 23% at trabecular sites, while another showed an increase at a trabecular site alone, and one showed no increase at either site.

There are several possible explanations for the differences shown in these studies. The use of a stick as support might cause an increase in local bone mass, and this was eliminated in only 2 of the studies. More important, patients and study groups have not been closely matched for skeletal size in any of the studies where increased bone mineral has been shown. In 2 of them the osteoarthrotic groups were indeed taller than the controls.

No previous studies have corrected for menopausal status. As it is recognised that bone loss in females occurs at a rate of 1-1% to 1-5% per annum after the menopause and at a much slower rate before (0-37%), the small changes in the bone mineral content of the skeleton shown in some of the above studies may simply be related to different menopausal status.

Our failure to show increased total skeletal bone mass in patients with PGOA makes it very unlikely that bone mineral content is important in the aetiology of this genetically determined condition. Nevertheless, these findings do not rule out the possibility of local increases in bone density in areas adjacent to affected joints, or very small increases in total bone mass.

This work was supported by grants from the Scottish Hospital Endowment Research Trust and the Arthritis and Rheumatism Council. We are indebted to the Medical Research Council for use of the cyclotron at the Western General Hospital, Mrs Joan Gilsob for radiographic assistance, and Mrs Elizabeth Law for help with the total body calcium measurements.

References
Changes in Total Body Calcium after Renal Transplantation
Effect of Low-Dose Steroid Regime

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Key Words. Activation analysis - Homologous transplantation

Abstract. Total body calcium was measured in 8 men and 4 women, aged 20–51 years, undergoing kidney transplantation. The initial measurement was made within 8 weeks of operation and subsequent measurements up to 33 months postoperatively. Transplant rejection was prevented by low-dose prednisolone therapy (20 mg/day). 2 patients underwent parathyroidectomy for hypercalcaemia, and their total body calcium increased by 29 g (3%) and 66 g (8%). In the remainder the mean annual change was -0.9% (3.7, SD) over an average follow-up period of 17 months. This fall in total body calcium was statistically insignificant and was smaller than that previously described in patients treated with higher doses of steroids.

Introduction

Total body calcium (TBCa) can be used to measure long-term changes in calcium balance. We have applied this technique to 12 patients undergoing kidney transplantation. Previous prospective studies have shown that bone mass falls by between 5 and 7% in the 1st year after transplantation [Lindsay et al., 1976; Aird and Pierides, 1977]. However, these studies used higher doses of steroids than those used in the present study.

Patients and Methods

8 men and 4 women aged 20–51 years (mean 35.9) were studied. They had been on maintenance haemodialysis for 4–54 months (mean 23). The causes of renal failure were chronic glomerulonephritis (6), polycystic kidney disease (2), chronic pyelonephritis (1), obstructive nephropathy (1), malignant hypertension (1), and Goodpasture's syndrome (1). 2 patients had hypercalcaemia due to secondary hyperparathyroidism at the time of operation, 1 had aluminium-induced osteomalacia, and 1 had osteomalacia treated with alfacalcidol. 7 patients had arterial wall calcification.

9 patients received kidneys from cadavers and 3 from related donors. Antirejection treatment was given as described by McGeown et al [1977]. Azathioprine was given at a dose of 3 mg/kg/day and prednisolone at a dose of 20 mg/day. Subacute rejection episodes (average of 1 per patient, range 0–3) were treated with prednisolone 200 mg/day and reduced to 20 mg/day over 10 days. This dose was reduced to 10 mg/day over 10–33 months, unless chronic rejection developed (2 cases), and then the dose was increased to 20 mg/day.

TBCa was measured by in vivo neutron activation analysis using a cyclotron as the source of neutrons [Kennedy et al., 1982]. This method has a precision in anthropomorphic models of 1.8% for a dose of 13 mSv. Formulae have been described for predicting TBCa (TBCap) from height in men and from arm span and years postmenopausal in women [Kennedy et al., 1982]. The ratio of TBCa to TBCap is the calcium ration (CaR) and the mean value ± standard deviation (SD) for 20 healthy men was 1.000±0.062 and for 20 women 1.000±0.066. Initial measurements were made in the week prior to surgery in the 3 patients receiving kidneys from related donors and within 8 weeks of surgery in the remainder. Subsequent measurements were made at about 6-month intervals for up to 33 months (mean 17 months). Patients gave informed written consent, and the study was approved by the hospital ethics committee and the Administration of Radioactive Substances Advisory Committee.

Plasma calcium values were available immediately prior to operation and were subsequently measured monthly along with magnesium, protein, and creatinine. Plasma phosphate values were also available from before operation, and plasma parathyroid hormone (PTH), and plasma 25-hydroxyvitamin D were measured when the initial TBCa measurement was made; measurements were repeated at the end of the study. Radiographs were taken when considered appropriate by the physician in charge.

A paired t test was used to analyze the changes following kidney transplantation. Comparison with controls were made using the unpaired t test. Plasma alkaline phosphatase and 25-hydroxyvitamin measurements were log2-transformed prior to statistical analysis.
Table I. Initial TBCa results in patients undergoing renal transplantation.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Men</th>
<th>Women</th>
</tr>
</thead>
<tbody>
<tr>
<td>TBCa, g</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Renal failure</td>
<td>1,006</td>
<td>685</td>
</tr>
<tr>
<td>Controls</td>
<td>1,143</td>
<td>821</td>
</tr>
<tr>
<td>CaR</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Renal failure</td>
<td>0.93*</td>
<td>0.82***</td>
</tr>
<tr>
<td>Controls</td>
<td>1.00</td>
<td>1.00</td>
</tr>
</tbody>
</table>

*p < 0.05; ***p < 0.001 (unpaired t test).

Table II. Initial plasma biochemistry and changes following renal allotransplantation in 10 patients.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Initial value</th>
<th>Mean change</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alkaline phosphatase, U/l</td>
<td>72.2</td>
<td>-54</td>
<td>NS</td>
</tr>
<tr>
<td>Phosphate, mmol/l</td>
<td>1.69</td>
<td>-51</td>
<td>&lt;0.02</td>
</tr>
<tr>
<td>Calcium, mmol/l</td>
<td>2.34</td>
<td>8</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>Magnesium, mmol/l</td>
<td>1.26</td>
<td>-29</td>
<td>&lt;0.002</td>
</tr>
<tr>
<td>Parathyroid hormone, g/l</td>
<td>0.82</td>
<td>-71</td>
<td>NS</td>
</tr>
</tbody>
</table>

NS = Not significant.

Results

Total Body Calcium

Initial values for TBCa and CaR are shown in table I. The mean initial CaR values in the 8 men showed a 7% reduction below the mean for male controls and in the 4 women an 18% reduction below the mean for female controls. 3 of the 4 women and 2 out of the 8 men had CaR values less than 2 SD below the mean for the controls.

The 2 men who underwent parathyroidectomy 4 and 20 months after renal transplantation had increases in TBCa of 66 g (8%) over 10 months and 29 g (3%) over 18 months. In the remainder the mean change was calculated from the initial and final measurements and was -0.9%/year, with 95% confidence intervals of -3.2–1.4%/year (fig. 1).

Biochemistry.

The initial biochemistry results are shown in table II. The elevated plasma creatinine fell gradually, but never into the normal range (fig. 2). 2 patients underwent parathyroidectomy, and so their data are excluded from the following results.

Plasma alkaline phosphatase and phosphate (table II) and plasma magnesium (fig. 3) all fell into the normal range. Plasma calcium gradually increased over the first 6
months such that the mean value was at the upper limit of the reference range for the 30-month follow-up period (fig. 3).

**Discussion**

The annual reduction in TBCa of 0.9% was not statistically significant and was much smaller than the 7% fall in metacarpal bone density described by Lindsay et al. [1976] in the 1st year after renal transplantation. Aird and Pierides [1977] showed a fall of 5.1–5.4% in lower femur density over a similar period. Both groups used a higher dose of steroid, and this could account for the more rapid bone loss compared with the changes in the present study.

The insignificant change in TBCa contrasts with the dramatic biochemical changes. The time course of the return of plasma alkaline phosphatase, phosphate, and magnesium to normal, along with a reduction in plasma PTH almost to normal, corresponds with the bone biopsy findings of Bortolotti et al. [1977]. They showed that the marrow fibrosis, excess osteoid, and increased bone resorption had resolved by 16–20 months after renal allotransplantation.

The persistent mild hypercalcaemia has been described in 21% of the 386 patients reported by Chatterjee et al. [1976], David et al. [1973], Ibel et al. [1978], and Lee et al. [1973]. It is associated with normal or raised plasma PTH levels [Kleerekoper et al., 1975]. This elevation may be due to the persistently impaired renal function. Le-gnore et al. [1979] have shown that plasma PTH rises when creatinine clearance falls below 50–80 ml/min (0.83–1.33 ml/s).

The initial low TBCa differs from the findings of others. Cohn et al. [1975] and Denney et al. [1973] reported normal mean TBCa values in patients on haemodialysis for chronic renal failure. The type of bone disease was reported to affect TBCa, the lowest values being found in those with osteomalacia and normal or high values in those with hyperparathyroidism. In the present series the low TBCa may be due to a higher frequency of osteomalacia, but unfortunately bone biopsies were not performed.

The healing of renal osteodystrophy after renal allotransplantation might be expected to be associated with an increase in TBCa, particularly in the patients with a predominantly osteomalacia type of bone disease. In the present series this effect was countered by the osteopaenic effect of the steroids. Ectopic calcification was present in more than half the subjects, but was always confined to blood vessel walls and was unlikely to have contributed much to TBCa. Since once the renal osteodystrophy has healed, the action of the steroids is unopposed and may be potentiated by the commonly associated mild hyperparathyroidism. This might account for the findings of Andrensen and Nielsen [1982] of a low bone density in patients 67 months after renal transplantation; the mean value of bone mineral concentration of the radius and ulna was 25% below age- and sex-matched controls.

**Acknowledgements**

We would like to thank the surgeons and physicians of the Nuffield Transplantation Surgery Unit for their co-operation in this study. The biochemical estimations were performed in the laboratories of Dr. D. B. Horn, Dr. P. J. Ashby, and Dr. J. Seth. R. Eastell was funded by the Medical Research Council and N. S. J. Kennedy by the Scottish Home and Health Department.

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Lemmann, XXXXX.


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The seasonal variation of total body calcium

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Abstract. Total body calcium was measured using in vivo neutron activation analysis in 156 patients with rheumatic diseases at six-month intervals. Evidence of seasonal variation was sought by relating deviation from a linear change to the month in which the measurement was made. A cyclic regression fitted to the data had an amplitude of 0.25% but a significance of only $P = 0.08$. The calcium balance studies of Malm were re-analysed to quantify the seasonal variation he had demonstrated in one group of men. The deduced variations of body calcium were highly significant, with a phase and amplitude very similar to the regression of our data. Both analyses differed from two previous reports of seasonal variations of part-body bone mineral of much greater amplitude and different phase.

1. Introduction

In 1958 Malm demonstrated a seasonal variation of calcium balance in a group of subjects maintained on a fixed intake over a long period (Malm 1958). This finding is compatible with subsequent studies of the seasonal variation of associated measurements of calcium biochemistry (Smith et al 1964, Stamp and Round 1974, Robertson et al 1974).

There have been two reports of direct measurements of bone mineral in limited parts of the skeleton which also showed seasonal variation (Aitken et al 1973, Krämer 1983). There do not appear to have been any studies of such an effect in the whole skeleton. As we have been carrying out longitudinal studies of total body calcium over an appropriate period it seemed worthwhile to examine the data for evidence of seasonal variation. Such a variation would be of direct physiological interest and also relevant to the interpretation of longitudinal studies of total body calcium.

2. Method

Total body calcium was measured by in vivo neutron activation analysis using a cyclotron source (Kennedy et al 1982). Corrections were applied for body size. The long term precision with anthropomorphic models was 1.8% and with patients 2.9% (Tothill et al 1984) for a dose-equivalent of 13 mSv. Three or four measurements of total body calcium were made at intervals of six months in 156 patients with various rheumatic diseases treated with a variety of agents (Reid et al 1984). 96 patients had rheumatoid arthritis, 9 psoriatic arthropathy, 19 primary generalised osteoarthrosis, 14 polymyalgia rheumatica and 18 ankylosing spondilitis. Permission was obtained from the local ethical committee and the Administration of Radioactive Substances Advisory Committee and the patients gave informed consent.
The rate of change of total body calcium was assessed by fitting a linear regression to the points for each patient. The possibility of a seasonal variation of total body calcium was investigated by relating the deviation of each point from the regression line to the month in which the measurement was made. Comparisons were made between mean deviations for monthly and three-monthly periods. In addition, a cyclic regression was fitted to the data.

Malm (1958) had demonstrated the existence of a seasonal variation of calcium balance by relating the proportion of 'better, poorer and intermediate' retentions to the month in which the observations were made. Such a method of analysis does not lend itself to the assessment of the significance or magnitude of the differences, so his data were subjected to re-analysis by the same techniques as were applied to our own.

3. Results

Our study commenced in mid 1980 and continued through 1983, although only 27 measurements were made in the last year. When mean deviations for each month of the study were plotted, no consistent pattern of variation was seen. As any seasonal changes were likely to be small compared with the spread of the data, results for each calendar month throughout the whole period were combined. The mean results for all subjects are shown in figure 1. Average deviations for each month are given, with standard errors. In addition, mean values are given for three-month periods. There is no significant difference between the mean values for any two months or quarters; the standard deviations and numbers of subjects were such that a difference of 0.7% in mean total body calcium between two three-month periods would have been significant at $P = 0.05$.

An expression of the form $x = a + b \cos(t + c)$ was fitted to all 590 observations using a least-squares method. $x$ is the percentage deviation, $t$ the month of the year and $a$, $b$ and $c$ are the parameters of the equation. $a$ represents the mean value of $x$, $b$ the amplitude of the oscillation and $c$ the phase angle. The best-fit curve was

$$x = -0.002 + 0.253 \cos(30t - 19.37).$$

The correlation coefficient was $r = 0.070$, with a significance of $P = 0.08$.

The correlation coefficient provides a measure of the amount by which the residual variance of the data is reduced by the regression; in this case the reduction is barely significant.
A curve was also fitted to the mean monthly deviations
\[ x = 0.032 + 0.187 \cos(30t - 8.93). \] (2)
The regression \((r = 0.44)\) had a significance of \(P > 0.1\), reflecting the undue weight placed on the results from July, a month in which fewer than average measurements were made.

The data were also considered separately for women before and after the menopause and for men. No significant differences (at \(P < 0.05\)) between monthly or three-monthly mean values were seen and cyclic regressions were not significant. As mean rates of change were different in some of the disease and treatment groups (Reid et al 1984) the possibility of seasonal variation was also examined in each of the groups separately. No evidence of such variation was found.

Malm (1958) considered seasonal variations of calcium balance in two groups of men studied for at least eight months. The first consisted of 16 men on a daily calcium intake of 950 mg. Over the whole period of study all but one of the subjects was in positive calcium balance. To quantify the variation, the mean of all 14-day calcium balances determined in a particular month was evaluated, together with its standard error. The results are plotted in figure 2(a). The three-month totals are also included. The numbers of observations varied slightly, averaging 31 per month. The difference between the highest positive balance in August and the lowest in April was significant \((P < 0.05)\). When quarterly totals are considered the mean balances in the period July–September were greater than those in January–June, also with \(P < 0.05\). The significance of the differences was calculated using tables of the Studentised range (Snedecor and Cochran 1980).

A cyclic regression was fitted to the data, the equation being
\[ y = 73.9 + 45.0 \cos(30t + 86.6) \] (3)
where the units of \(y\) are mg d\(^{-1}\) of calcium and \(t\) is in months. The correlation coefficient

![Figure 2. Malm's calcium balance data. Points are mean balances for each calendar month, with standard errors. Blocks relate to means and standard errors for three-month periods. (a) 16 men maintained for at least eight months with an intake of 950 mg d\(^{-1}\); the full curve is the cyclic regression fitted to the data. (b) 10 men maintained for at least eight months with an intake of 450 mg d\(^{-1}\).](image-url)
is \( r = 0.90 \) and the regression is highly significant \( (P < 0.001) \); the amplitude is 45 mg d\(^{-1}\). The curve is included in figure 2(a).

Variations in total body content of calcium can be obtained by integration of equation (3), giving

\[
z = 2.25t + 2.58 \sin(30t + 86.6)
\]

where \( z \) is the change in calcium content in g since the beginning of the year. The linear term reflects the overall positive balance of the subjects. The cyclic term has an amplitude of 2.6 g, with the phase shifted by three months compared with equation (3), so that the maximum positive deviation from a linear change occurs in December and the maximum negative deviation in June. The cyclic part of the regression is plotted in figure 3.

\[
\text{Figure 3. Cyclic regressions fitted to deviations from a linear change: A, total body calcium from Malm's balance data; B, total body calcium measured by us; C, metacarpal bone mineral reported by Aitken et al.; D, spine bone mineral reported by Kröliner.}
\]

The other group in which Malm claimed to have observed seasonal variation of calcium balance contained ten men maintained on a diet of only 450 mg d\(^{-1}\). The mean monthly and three-monthly balances are plotted in figure 2(b). None of the differences between means was significant. The best-fit cyclic regression, \( y = -44.5 + 7.73 \cos(30t + 91.7) \), was also not significant \( (r = 0.29, P > 0.1) \). Although this regression was not statistically significant, it may be noted that the phase was very similar to that observed for the high-level-intake regression.

4. Discussion

The first suggestion of possible seasonal variation of calcium balance came from McCance and Widdowson (1943). However, they made the finding in only three subjects. Malm (1958) demonstrated such an effect with more certainty by long-term balance studies on groups of subjects in a Norwegian state penitentiary, using closely controlled calcium intakes. Our study was the first to examine seasonal changes of total body calcium by direct measurement. The results suggest such a variation, but do not demonstrate it with a high degree of statistical significance.
Seasonal variation of total body calcium

Our re-analysis of Malm's data shows the magnitude and significance of seasonal variation based on long-term balance measurements. For the group of 16 men with a high calcium intake, the seasonal variation was highly significant. The maximum deviations of total body calcium from a linear change were 2.6 g. The calcium content of Reference Man is 1000 g (ICRP 1975) and our own measured mean value is 1143 g (Kennedy et al 1982) so that such an amplitude represents only about 0.25% of total body calcium. The dispersion of our results of \( \text{TBCa} \) measurements explains the difficulty that we have in detecting a change of this magnitude with any degree of significance. Nevertheless, although of marginal statistical significance, the curve fitted to our data is very similar in amplitude and phase to that derived from Malm's data. Both curves are included in figure 3.

Malm's observation of an apparent seasonal variation in the group of ten men with a low calcium intake was not, in fact, statistically significant.

The previous reports of the direct measurement of seasonal changes of bone mineral in humans have examined only a part of the skeleton. Aitken et al (1973) showed that the metacarpal mineral content of 48 post-menopausal women was 3.8% higher in the period May-August than in November-February. A cyclic regression fits very well to their data, with amplitude 2.3% and a maximum in July (figure 3).

Krölnér (1983) similarly reported an increase of 1.7% in the lumbar spine bone mineral content of 26 normal women (nine of them post-menopausal) during July-September compared with January-March. There was a mean annual loss of 1% in this group and Krölnér assessed the seasonal variation from deviations from a linear regression in the same way as we have. He fitted a cyclic regression equation, the amplitude of which corresponded to 1% of the spine bone mineral, with a maximum in August. This curve is also plotted in figure 3, which demonstrates that the two observations of bone mineral variation in parts of the body exhibit a greater amplitude than our deductions about total body calcium from Malm's balance data. They also show a different phase, with the maximum occurring about five months earlier.

Our results are not incompatible with Malm's observations. They are, however, at variance with the part-skeleton results; we could easily have detected changes of the magnitude reported by Aitken et al (1973) and Krölnér (1983) if they had applied to the whole skeleton. It may be that seasonal variations are only apparent at certain bone sites. The greater variation in the spine than in the whole body might be thought to be due to the more rapid turnover of trabecular bone. However, the amplitude was even greater in the cortical bone of the metacarpals. It would perhaps be unwise to draw too many conclusions from different populations examined by different techniques.

We have to consider whether the likelihood of seasonal variation of \( \text{TBCa} \) in our patients might be different from that for normal subjects by virtue of their disease. Our population consisted of groups of patients, the majority of whom suffered from rheumatoid arthritis or primary generalised osteoarthritis. Previous studies have shown that patients with these diseases do not have any disturbance of vitamin D metabolism (Bird et al 1980, 1982) and none of our patients had any abnormality of elementary calcium biochemistry. All were ambulant out-patients and it seems unlikely that the rheumatic diseases could affect any seasonal variation in calcium balance by either of the mechanisms that have been suggested, i.e. variable exposure to sunlight (Aitken et al 1973) or reduced exercise in winter (Krölnér 1983).

One of our motives for undertaking this study was concern that seasonal variations might influence the interpretation of our longitudinal studies of \( \text{TBCa} \). With the numbers
Acknowledgments

We should like to thank Mr Martin Connell for assistance in statistical computation and Mrs Norma Brown for much help with the research which was supported by the Scottish Home and Health Department and the Arthritis and Rheumatism Council.

Résumé

La variation saisonnière du contenu en calcium des tissus. Les auteurs ont mesuré, à six mois d'intervalle, le contenu en calcium des tissus par analyse de l'activation neutronique in vivo sur 156 patients présentant des troubles rhumatismaux. Une variation saisonnière évidente a été recherchée en reliant l'écart à un changement linéaire au mois durant lequel la mesure était faite. Une régression cyclique ajustée aux données a présenté une amplitude de 0,25% mais une significativité de seulement \( P = 0,08 \). Les études par Malm de l'équilibre en calcium ont été analysées de nouveau pour quantifier la variation saisonnière qu'il avait démontré chez un groupe d'hommes. Les variations déduites du contenu en calcium des tissus étaient hautement significatives, avec une phase et une amplitude tout à fait comparables à celles obtenues en ajustant la régression à nos données. Les deux analyses différaient de deux précédents rapports sur les variations saisonnières du contenu minéral des os dans certaines parties du corps, présentant une amplitude beaucoup plus grande et une phase différente.

Zusammenfassung


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Stamp T C B and Round J M 1974 Seasonal changes in human plasma levels of 25 hydroxyvitamin D Nature 247 563-5
Bone Mass in Ankylosing Spondylitis

DAVID M. REID, JEREMY J. NICOLL, NORMAN S. J. KENNEDY, MICHAEL A. SMITH,
PETER TOTHILL, and GEORGE NUKI

Abstract. To assess bone mass in ankylosing spondylitis (AS) we have measured total body calcium, bone mineral content of the lumbar spine and metacarpal indices in groups of patients with AS. Mean total body calcium was reduced by 5.3% (p < 0.05) in 20 patients compared with controls. The mean annual loss of bone, assessed over an 18 month period in 17 patients, was 2.9% (p < 0.001). Compared to controls, bone mineral content was increased by 28% (p < 0.05) in 8 male patients while metacarpal indices were normal in 18 male and female patients. The results of total body calcium measurements give support to the hypothesis of a minor increase in bone turnover in AS. The increased bone mineral content in the male patients may relate to syndesmophyte formation. (J Rheumatol 1986;13:932-935)

Key Indexing Terms:
ANKYLOSING SPONDYLITIS
TOTAL BODY CALCIUM

Although vertebral osteoporosis was originally considered to be an early radiological sign of ankylosing spondylitis (AS)1-2, it is now recognized to occur as a late feature of the disease process3-4. Spontaneous fractures, a common consequence of vertebral osteoporosis, appear to be uncommon in AS, occurring only once in 212 patients in one report5. However, another group of investigators found 2 spontaneous fractures in a series of 50 patients, and the majority of subjects were said to have mild axial and appendicular osteoporosis6.

To shed light on the presence and site of bone loss in AS we have measured total bone mass by total body calcium in patients with AS and related the results to those of vertebral bone mass measured by dual photon absorptiometry, peripheral bone mass measured by metacarpal indices and radiographic disease status measured by syndesmophyte count.

MATERIALS AND METHODS

Patients. We measured total body calcium in 20 randomly selected patients (14 men, 6 women) with AS as defined by the New York criteria7 and compared the results with 52 age matched normal control subjects (27 men, 25 women). From within the total group 10 patients with AS (8 men, 2 women) had a single estimation of lumbar spine bone mineral content, 18 (13 men, 5 women) had a single measurement of metacarpal indices and 8 (7 men, 1 woman) had a syndesmophyte count. These patients were selected solely on willingness to attend for bone mass assessment. Bone mineral content results were compared with 36 normal control subjects (18 men, 18 women) and metacarpal indices with 45 normal controls consisting of 24 asymptomatic volunteers (12 men, 12 women) and 21 patients with primary generalized osteoarthritis (5 men, 18 women) who had been shown to have normal total bone mass8.

Total body calcium measurement was repeated on 3 or 4 occasions over an 18 month period in 17 patients with AS (13 men, 4 women).

Serum calcium, phosphate, proteins and alkaline phosphatase were measured in all subjects and those with abnormalities of calcium metabolism or medical conditions associated with secondary osteoporosis were excluded.

Methods. Total body calcium was measured by in vivo neutron activation analysis as described9. Neutrons produced from the Medical Research Cyclotron were directed at a patient standing in a rigid activation enclosure. The gamma rays produced by the reaction46 Ca(n,γ)48 Ca were measured in a whole body counter and compared with the results from a phantom to give a figure for total body calcium in grams. For longitudinal studies the measurement was repeated 3 or 4 times over an 18-month period and a regression line drawn through the values of total body calcium. From the slope of the regression line, the rate of change of total body calcium was calculated and expressed as a percentage of the initial value. In the initial studies the figure for total body calcium in grams was normalized for skeletal size (arm span) for both sexes and menopausal status in women using the regression formulae previously published9. The in vivo precision of the technique was 2.9%10 for a radiation dose of 15 mSv.

Bone mineral content of the lumbar spine was measured by dual photon absorptiometry with an adapted dual head rectilinear scanner using11 Gd as a dual energy source12. The bone mineral image was displayed on a colour monitor and cursors placed at the top of lumbar vertebrae 2 (L2) and at the bottom of L4. The bone mineral content of the area within the cursors was calculated and divided by the vertical length to give bone mineral content in g/cm. The in vivo precision was reduced to 1.0% by an experienced operator. Predicted bone mineral content in patients with AS was calculated using the formula derived from the linear relationship found in normals between span and bone mineral content/unit length13.

Metacarpal indices were measured by a variation of the technique described by Dequeker14. From nomenclature radiographs of the hands taken at a fixed 1 mm source to film distance, the length (L) of the 2nd, 3rd and 4th metacarpals of the right hand were measured. At the mid-point of each metacarpal the external diameter (D) and the internal diameter (d) of the cortex were measured using a Vernier caliper. The mean cortical area as a percentage

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of surface area \( [(D^2-d^2)/DL] \) was then calculated. The precision of the technique, evaluated from 2 radiographs taken at daily intervals in 10 young healthy controls was 2.0%.

The formation of bony bridging between the lateral and anterior borders of adjacent lumbar vertebrae from L1 to L5 was measured from posterior and lateral films of the lumbar spine. Syndesmophytes were graded on a 0-3 scale with 3 equal to complete bony bridging. A final syndesmophyte count was recorded as a mean of the values at the 12 sites measured. Hence, no syndesmophytes would score 0 and a bony spine 3.

Statistical analysis was performed by Student’s t test for unpaired variables. For small numbers and data not normally distributed, Wilcoxon’s rank sum test and Kendall’s rank correlation test were used.

**RESULTS**

Details of sex and age in controls and patients undergoing each assessment of bone mass are shown in Table 1. The results of normalized total body calcium, bone mineral content and metacarpal indices expressed as a percentage of normal are shown in Figure 1. Total bone mass was reduced by 5.3% (p<0.05) in AS compared with controls, when both sexes were considered together. The reduction was greater in the 6 female patients (9.7%, p<0.05) than in the males (2.4%), although the difference between the sexes did not reach statistical significance.

Bone mineral content of the lumbar spine was increased by 28% (p<0.05) in the 8 male patients measured but reduced in the 2 females. Metacarpal indices were normal in male and female patients with AS (Figure 1).

Details of disease duration and activity are shown in Table 2. None of the bone mass indices were related to disease duration or activity as measured by duration of early morning stiffness and erythrocyte sedimentation rate (ESR). The rate of change of total body calcium did not correlate with the change in the disease activity indices and was not related to the duration of disease. In addition, none of the bone mass indices was significantly related to the syndesmophyte count (mean ± 1SD = 1.15 ± 0.85), when male and female patients were considered together. As syndesmophytes might be expected to increase spinal bone mineral, the difference in percentage of expected bone mass at lumbar and total bone sites (i.e., % bone mineral content – % total body calcium) was compared to the syndesmophyte count and a significant correlation was found (tau= –0.571, p<0.01).

The annual rate of loss of total body calcium in the 17 patients with AS completing the longitudinal study (Table 3) was 2.9% (p<0.01), a figure not entirely explained by the large loss in the 4 female patients (5.2%), the male reduction also being significant.

---

**Table 2. Results of disease activity indices expressed as the mean initial and mean change (± 1 SD) in male and female patients with AS undergoing measurements of total body calcium.**

<table>
<thead>
<tr>
<th></th>
<th>Male</th>
<th>Female</th>
</tr>
</thead>
<tbody>
<tr>
<td>Duration of disease (years)</td>
<td>12.1 ± 6.8</td>
<td>14.3 ± 10.4</td>
</tr>
<tr>
<td>Initial EMS (min)</td>
<td>20.0 ± 25.9</td>
<td>114.2 ± 107.7</td>
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<tr>
<td>Change in EMS (min)</td>
<td>+3.5 ± 14.3</td>
<td>−46.2 ± 41.9</td>
</tr>
<tr>
<td>Initial ESR (mm/h)</td>
<td>24.8 ± 23.0</td>
<td>48.7 ± 40.4</td>
</tr>
<tr>
<td>Change in ESR (mm/h)</td>
<td>−5.5 ± 19.5</td>
<td>−31.7 ± 35.0</td>
</tr>
</tbody>
</table>

(EMS = duration of early morning stiffness; ESR = erythrocyte sedimentation rate)

---

**Table 1. Sex, mean age and years post menopause (YPM) ± 1 SD in groups of patients with AS and controls undergoing total body calcium (TBCa), bone mineral content of the lumbar spine (BMC), metacarpal indices (MI) and syndesmophyte count (SC).**

<table>
<thead>
<tr>
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<th>Controls</th>
</tr>
</thead>
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<tr>
<td></td>
<td>Sex</td>
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</tr>
<tr>
<td>TBCa</td>
<td></td>
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<tr>
<td>Males</td>
<td>14</td>
<td>41.7 ± 9.7</td>
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<tr>
<td>Females</td>
<td>6</td>
<td>39.7 ± 8.1</td>
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<tr>
<td>BMC</td>
<td></td>
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<tr>
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<td>8</td>
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</tr>
<tr>
<td>Females</td>
<td>2</td>
<td>40.0 ± 11.3</td>
</tr>
<tr>
<td>MI</td>
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<td>Females</td>
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<td>SC</td>
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<td></td>
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<td>Males</td>
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<td>44.0 ± 10.4</td>
</tr>
<tr>
<td>Females</td>
<td>1</td>
<td>32</td>
</tr>
</tbody>
</table>

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DISCUSSION

In vivo methods of bone mass assessment have not previously been reported in patients with AS. However, radiographic studies have suggested that osteoporosis occurs both in the axial and appendicular skeletons, and using the technique of 24-h whole body retention of diphosphonate, bone turnover has been shown to be increased in our study. Total body calcium has been shown to be slightly, but significantly, reduced in patients with AS although much of the reduction occurred in the small group of female patients. In general the female patients had slightly more active disease than the males as measured by ESR and duration of early morning stiffness. However, as the differences did not reach statistical significance and the indices were not related to the results of total body calcium, it is not clear whether the slightly more active disease in females accounts for the lower bone mass. Although this small group of female patients was younger than controls, the age difference did not reach statistical significance using nonparametric tests. Total bone mass falls with age, but we have failed to show a relationship between age and total body calcium in normal premenopausal females either in this study or in previous work. The lack of such a relationship has been suggested by other workers who only include an age correction factor for those women over the age of 55. Accordingly it seems very unlikely that the difference in mean age between patients and controls accounts for the substantial differences in total body calcium.

Longitudinal total body calcium assessment gave support to the theory of a minor absolute reduction in bone mass as there was a significant annual loss of bone in males (2.2%) despite the young mean age of the group (42.6 years). Ethical considerations precluded the collection of control data for sequential measurements of bone mass, although previous work with the technique has shown an estimated annual reduction of total body calcium of only 0.7% in men over the age of 50 with no recorded loss before that age. The rate of bone loss was not related to the activity of disease which tended to diminish during the study period.

If there is a reduction in total bone mass and an increased rate of bone loss, it is of interest to speculate where this might occur. Bone mineral content of the lumbar spine in 6 male patients was increased by 28% compared to controls but this increase is almost certainly explained by excess bone formation. The syndesmophyte count was correlated with the difference between the expected bone mass in the vertebrae and the total body while it was not related to lumbar spine bone mass alone. Hence, there is a strong suggestion that the increase in vertebral bone mineral content is caused by the presence of syndesmophytes. Using our current methods it is impossible to eliminate the effect on vertebral bone mass of syndesmophytes which become ossified when fully developed. It remains possible that the vertebral bodies themselves are relatively osteopenic as suggested in pathological and radiological studies. The bone mineral content of the lumbar spine of the 2 female patients was markedly reduced and this is in keeping with the reduced total bone mass of the female patients.
The reduction in total body calcium in AS does not appear to be explained by reduced vertebral bone mass or by appendicular bone loss as metacarpal indices were normal in our patients. In order to be more certain of the site of bone loss it would be desirable to assess total, axial and appendicular bone mass simultaneously in the same groups of patients with AS and controls. This was not possible in our study.

The causes of increased bone loss in AS are unknown. Vertebral immobilization and changes in blood flow could be implicated, but the humoral factors responsible for increased turnover of vertebral bone have yet to be identified. Both immunoreactive parathyroid hormone (PTH) and biologically active PTH have been found to be elevated in some patients with AS but, in the absence of overt signs of hypoparathyroidism, both studies suggested that end organ failure might be responsible for the biochemical abnormality. We did not measure PTH values in our patients but we did not find any evidence of abnormal calcium biochemistry in our patients, and the only patient found to have a transient elevation in serum calcium was omitted from the analysis reported above. Many other biological mediators of bone resorption have recently been identified. From among these bone resorbing lympokinies, interleukin I, prostaglandins and vitamin D metabolites are all mediators that deserve further investigation in chronic inflammatory disorder such as AS.

ACKNOWLEDGMENT

We are indebted to the Medical Research Council for use of the cyclotron at the Western General Hospital, Edinburgh, Mrs. N. Brown for data collection and to Mrs. E. Law and Mr. C. Ferrington for assistance with some of the measurements of total body calcium.

REFERENCES

Corticosteroids and bone mass in asthma: comparisons with rheumatoid arthritis and polymyalgia rheumatica

D M Reid, J J Nicoll, M A Smith, B Higgins, P Tottill, G Nuki

Abstract
Bone mass has previously been shown to be reduced at peripheral bone sites in patients with bronchial asthma receiving corticosteroids. To assess whether total bone mass is reduced in asthma total body calcium was measured by in vivo neutron activation analysis in patients receiving various treatments for asthma and compared with results from normal controls and patients with rheumatoid arthritis and polymyalgia rheumatica. Compared with controls total body calcium was reduced by 13-6% (p<0-001) in patients with asthma receiving daily oral corticosteroids but by only 9-0% (p<0-005) in a similar group of patients who had received oral calcium supplements at the start of their corticosteroid treatment. Total body calcium was also reduced in a group of patients receiving only inhaled corticosteroids (8-8%; p<0-001) but not significantly reduced in a small group of patients with asthma who had never received these drugs. When compared with controls a group of patients matched for age and for dose of corticosteroids given for rheumatoid arthritis had a similar reduction in total body calcium to the patients with asthma receiving daily oral treatment (17-7%; p<0-001), but no such reduction was shown in patients with polymyalgia rheumatica.

These findings suggest that the risk of bone loss with low dose oral corticosteroids is similar in asthma and rheumatoid arthritis. Further work is required to assess the clinical relevance of small losses of bone associated with the use of inhaled corticosteroids.

Introduction
Since Cushing's original description of the syndrome in 1932 it has been recognised that supraphysiological concentrations of endogenous corticosteroids are associated with excess bone loss. The introduction of cortisone as a therapeutic agent was followed very shortly by case reports of spontaneous fractures during treatment, and subsequently radiological osteoporosis was reported at various skeletal sites. The cause of corticosteroid induced osteoporosis is now better understood but there remains controversy about the dose and duration of treatment with synthetic corticosteroids necessary to induce osteoporosis. Whether or not bone loss can be prevented by altering the route or timing of drug administration remains uncertain. There is also continuing controversy whether patients with some diseases may be less susceptible.

As there is a relative paucity of data on bone mass in asthmatics we have carried out measurements of total body calcium in groups of patients with bronchial asthma treated with oral and inhaled corticosteroids and compared the results with those in people who have not received steroids. The results were also compared from those patients with rheumatoid arthritis and polymyalgia rheumatica receiving regular oral corticosteroid treatment and from controls matched for age and sex.

Patients and methods
We measured total body calcium in 70 patients with asthma (28 men, 42 women). The total group was subdivided according to current and previous treatment. Group 1 consisted of 23 patients (eight men, 15 women) who were receiving regular oral corticosteroids and had been doing so for at least two years. Group 2 consisted of a further 13 patients (two men) who were also receiving regular oral corticosteroids but who had received oral calcium supplements as calcium lactate gluconate three tablets a day (elemental calcium dose 1 2 g daily) at the start of their corticosteroid treatment and for two to 20 years thereafter. Group 3 consisted of 22 patients (11 men) whose only regular corticosteroid treatment was given by the inhaled route either as beclometasone 100 µg four times daily or as betamethasone 200 µg four times daily. Sixteen patients (six men) in group 3 had also received a mean of 2-2 booster courses of oral corticosteroids, usually beginning with 40 mg prednisolone daily and reducing to nothing over seven to 10 days. Group 4
consisted of 12 patients (seven men) who had never received corticosteroids either by inhalation or by the oral route. The results from the four groups of patients with asthma were compared with those from 40 controls (20 men). The results from patients in group 1 were further compared with those from 27 patients (eight men) with definite or classical rheumatoid arthritis, matched for daily dose of prednisolone, and 12 patients (3 men) with polymyalgia rheumatica who were receiving similar daily doses of oral corticosteroids. All patients and controls who had abnormalities of calcium biochemistry or medical conditions known to be associated with secondary osteoporosis were excluded.

The corticosteroid dose for each patient was calculated retrospectively from the case records, supplemented when necessary by information from the patient. The total dose prescribed was expressed as the daily dose of prednisolone or equivalent averaged over the duration of treatment. Data on booster courses of the drugs were taken both from the case records and by direct questioning of the patient. Total body calcium was measured by in vivo neutron activation analysis using the Edinburgh Medical Research Council cyclotron. Patients were irradiated with neutrons for 40 seconds while standing in a rigid polyethylene activation enclosure. Patients were then transferred to a shadow shield whole body counter, where γ radiation induced by the reaction calcium-48(n,γ) calcium-49 was measured for 20 minutes. The patient's total body calcium was calculated in grams by comparison with the energy spectrum from an activated anthropomorphic phantom of human dimensions containing a known quantity of calcium. Repeated measurements of the spectrum gave a long term precision of 1.8% for a radiation dose of 13 mSv (1.3 rem). Normalisation was carried out by expressing individual results as a percentage of the expected normal value for the patient’s skeletal size (arm span) in both sexes and menopausal state in women. The in vivo-precision of the technique was 2.9%. Statistical methods—Results were analysed by Student's t-test for unpaired variables. The Mann-Whitney test was used for data not normally distributed and for small samples. By using Chauvenet’s criterion—a statistical technique designed to eliminate extreme values considered to occur by chance—two grossly aberrant total body calcium values from a woman in each of groups 1 and 2 were removed, leaving a total of 65 patients with asthma, whose results are reported.

Results

Table 1 gives details of age, duration of disease, corticosteroid and oral calcium treatment, and total body calcium values in grams in the four groups of patients with asthma. Patients in group 4 were significantly younger than those in the other three groups. Duration of disease was longer in group 2 than in group 3 (by 14.8 years; p<0.05) and group 4 (by 6.8 years; p<0.01). Patients in group 1 had received a lower dose of edisolone (by 2 mg/day; p<0.01) and for a shorter period (by 5-5 years; p<0.05) than those in group 2. Prenormalisation values for total body calcium in controls were 1142-7 (SD 133-8) g for men and 820-5 (124-7) g r women, as previously reported.18

Comparison of normalised results obtained in the control and asthma groups (fig 1) showed a reduction in mean total body calcium in group 1 (13.9%; p<0.001), group 2 (9.0%; p<0.005), and group 3 (8.8%; p<0.001). There was also a significant difference in mean total body calcium between groups 1 and 4 (8.1%; p<0.05). No significant differences were found between men and women.

Table 2 describes the sex ratio, age, corticosteroid treatment, and total body calcium values in group 1 patients with asthma and patients with rheumatoid arthritis and polymyalgia rheumatica. No significant differences were found between the patient groups with regard to age or corticosteroid treatment except that the patients with asthma had received oral steroids for longer than the patients with rheumatoid arthritis (7-4 years; p<0.001) and polymyalgia rheumatica (8-8 years; p<0.001).

Comparison of normalised results in these three groups showed a reduction in mean total body calcium in patients with rheumatoid arthritis (17.7%; p<0.001) but none in those with polymyalgia rheumatica (fig 2). Compared with the group with polymyalgia bone mass was reduced in the patients with asthma (13.2%; p<0.02) and rheumatoid arthritis (17.3%; p<0.001). The apparent difference in mean total body calcium between the patients with asthma and those with rheumatoid arthritis did not reach statistical significance (0.1%; p<0.05).

The reduction in total body calcium was not related to the duration of treatment or the total steroid dose (daily dose x duration) in any of the groups but was correlated with the mean daily dose of prednisolone in those patients with rheumatoid arthritis (r=0.605; p<0.001).

Discussion

Oral corticosteroid treatment has previously been associated with reduced bone mass in patients with bronchial asthma.19 These
studies compared treated patients with normal controls using different measurements of forearm bone mass. Though unlikely, the possibility that bone mass might be reduced in asthma as a consequence of the disease itself was not considered. As bones contain 99% of the body's calcium content it is a reasonable assumption that total body calcium is a measurement of total bone mass, particularly as it is extremely well correlated with the total bone mineral content of the skeleton measured by dual photon absorptiometry.2 Hence this study confirms that reduced total bone mass does occur in patients with bronchial asthma treated with daily oral corticosteroids (group 1) when compared both with age matched normal controls and with young asthmatics who had never received steroids (group 4). Though this latter group was significantly younger than the controls, other studies have failed to show a relation between total body calcium and age in men or premenopausal women, suggesting that the loss of bone in group 1 was attributable to steroid treatment rather than to age or disease.

The reduction in mean total body calcium of 13-6% in group 1 was substantial. Postmenopausal patients with crush fractures have a mean reduction in total body calcium of 16-20%,2,24 suggesting that many patients with asthma treated with steroids are at risk of osteoporotic fractures. Indeed two patients in group 1 had sustained at least one atraumatic crush fracture before the measurement of total body calcium and a further patient had recurrent problems with multiple rib fractures.

Prevention of corticosteroid induced osteoporosis in patients with asthma, rheumatoid arthritis, and other diseases remains a severe therapeutic problem.3 The use of alternate day corticosteroid regimens has not been shown to be advantageous.3,37 In this study five patients in groups 1 and 2 who had received an alternate day regimen of corticosteroids for the first two to five years of treatment had a total bone mass which did not differ from that of patients who had received continuous daily treatment.

Despite higher daily doses and a longer duration of corticosteroid treatment a small group of patients (group 2) who had received oral calcium supplements at the start of treatment had a slightly greater mean total body calcium than those in group 1 (p<0.01). This suggests that the use of oral calcium supplements alone given at the start of corticosteroid treatment should be further investigated as a possible means of preventing osteoporosis. Other studies aimed at overcoming the calcium malabsorption known to be associated with corticosteroid treatment3 have shown some benefit with 25-hydroxy vitamin D but not with 1,25-dihydroxy vitamin D.27

The use of inhaled corticosteroids in asthma is almost certainly associated with a reduction in systemic side effects compared with oral treatment2,26 but suppression of the hypothalamic-adrenal axis may occur with large doses.2,26 There is little doubt that the main portion of the aerosol dose is deposited in the mouth and swallowed, allowing gastrointestinal as well as pulmonary absorption. The finding that total body calcium was reduced in patients in group 3 by a mean of 8.8% compared with controls is therefore of concern. Sixteen of the 22 patients had received booster courses of oral corticosteroids, and such intermittent treatment has been associated with minor reductions of bone mass in patients with asthma.3 The six patients who had never received oral treatment, however, had some of the lowest individual values of total body calcium (fig 1). This suggests that inhaled corticosteroids might be at least in part responsible for the reduction in group 3. The widespread and increasing use of high dose corticosteroid inhalers suggests that further studies of bone mass in patients so treated should be undertaken. The pending development of inhaled corticosteroids which have an improved topical to systemic ratio should be encouraged.

The effects of corticosteroids on bone mass in patients with asthma, rheumatoid arthritis, and polymyalgia rheumatica were compared (fig 2). Both patients with asthma and those with rheumatoid arthritis had reduced bone mass compared with patients treated for polymyalgia rheumatica but we were unable to confirm previous studies,12-14 which suggested that corticosteroid treated patients with rheumatoid arthritis were more susceptible to bone loss than patients with asthma treated with similar daily doses. This study shows a small reduction in mean total body calcium (4.3%) in steroid treated patients with rheumatoid arthritis compared with asthmatics. As we have previously shown that total body calcium is reduced by 5-0-6.8% in patients with rheumatoid arthritis who have never received corticosteroids,14 this reduction in bone mass may be attributable to the effects of the disease process, which causes appreciable bone loss at peripheral sites.14

Total body calcium is inversely related to the daily dose of prednisolone in patients with rheumatoid arthritis (r=-0.605; p<0.001). This finding raised the hypothesis that corticosteroid induced bone loss might occur early in the course of treatment,15 and a recent longitudinal study has lent support to this theory.27 Lack of a similar relation between total body calcium and daily steroid dose in asthma may reflect the wider fluctuations in individual steroid doses in these patients.

The explanation for the normal bone mass in patients with polymyalgia rheumatica is not clear. Women with polymyalgia rheumatica were older than those in the two other groups and the magnitude of the corticosteroid induced bone loss may not be apparent because of the substantial yearly 1-5% correction after the mean age.

At present there is little convincing evidence for increased or decreased susceptibility to corticosteroid induced bone loss in patients with asthma or rheumatoid arthritis, though the position in patients with polymyalgia rheumatica may require further studies including larger numbers of subjects.

This work was supported by grants from the Scottish Hospital Endowment Research Trust and the Arthritis and Rheumatism Council with further funding from Sanofi UK Ltd. We are indebted to the Medical Research Council for use of the cyclotron and to Mrs E Law and Mr C Ferrington for helping with some of the measurements of total body calcium. This study would not have been completed without the invaluable work of Mrs N Brown, research assistant. We are grateful to Drs G Crompton and I Grant for permission to study patients under their care.

References


Total and peripheral bone mass in patients with psoriatic arthritis and rheumatoid arthritis


Rheumatic Diseases Unit, Northern General Hospital*, Department of Medicine, Western General Hospital* and Department of Medical Physics and Medical Engineering**, University of Edinburgh, Edinburgh, UK.

SUMMARY Psoriatic arthritis is thought to be associated with periarticular osteoporosis while rheumatoid arthritis may be associated with generalised as well as periarticular bone loss. To assess the extent of total and peripheral bone loss in these two diseases, total body calcium was measured by in vivo neutron activation analysis and peripheral bone mass was assessed by metacarpal indices in age-matched patients with psoriatic arthritis and rheumatoid arthritis treated with nonsteroidal anti-inflammatory drugs alone. In comparison with age and sex-matched normal controls, total and peripheral bone mass was normal in psoriatic arthritis. There were significant reductions in total (6.2% in men; 7.9% in women) and peripheral (10.9% in men; 12.8% in women) bone mass in patients with rheumatoid arthritis compared with controls. Peripheral bone mass was significantly correlated with the degree of radiographic damage in male and female patients with rheumatoid arthritis. The mean annual loss of total body calcium was insignificant in psoriatic arthritis (0.6% in men; 1.9% in women) but markedly greater in rheumatoid arthritis (4.4% in men; 2.7% in women). The data suggested that total and peripheral bone loss is greater in rheumatoid arthritis than psoriatic arthritis. Substantial reductions in peripheral bone mass in patients with rheumatoid arthritis not receiving corticosteroids may account in part for the small reductions in total bone mass.

Key words: Psoriatic Arthritis, Rheumatoid Arthritis, Bone Mass, Total Body Calcium, Metacarpal Indices.

INTRODUCTION

There is little published information on the bone mass of patients with psoriatic arthritis (PA), although local osteoporosis is a recognised radiological feature (1). This is not the experience of all investigators (2) and it is certainly limited in extent compared to that occurring in rheumatoid arthritis (RA) (3), where periarticular osteoporosis is an early radiological sign (4). Generalised osteoporosis is not thought to be a feature of PA (5) although it may occur in RA even in the absence of corticosteroids (6,7). To our knowledge no investigators have applied the in vivo methods of bone mass assessment to
patients with PA. As part of a large survey of bone mass in rheumatic diseases, we have measured total bone mass, by total body calcium, and peripheral bone mass, by metacarpal indices, in a small group of patients with psoriatic arthritis and compared their results with those from a group of age and sex-matched patients with rheumatoid arthritis treated with nonsteroidal anti-inflammatory drugs alone.

PATIENTS AND METHODS

Twelve patients (6 males, 6 females) with PA, meeting the suggested diagnostic criteria of Wright and Moll (3), were enrolled in the study. All patients had a peripheral arthritis conforming to the PA1, PA3 and PA4 sub-groups recognised by Moll and Wright (8). No patient had clinical evidence of spondylitis or sacro-iliitis (sub-group PA5) nor were there any patients with arthritis mutilans (sub-group PA2). All patients were sero-negative for rheumatoid factor and were receiving therapy with nonsteroidal anti-inflammatory drugs (NSAID) alone. No patient had received oral corticosteroids, suppressive anti-rheumatic drugs or immunosuppressive therapy.

Initial total body calcium (TBCa) has been assessed in the 12 patients with PA and compared with that of 45 patients (23 males, 22 females) with RA who were receiving NSAID alone and 36 controls (19 males, 17 females) (Table I).

The annual rate of change of TBCa has been assessed in 9 of the PA patients (5 males, 4 females) and compared with that of 13 age-matched patients (6 males, 7 females), with RA receiving NSAID alone (Table II).

Metacarpal indices have been measured in 11 of the PA patients and have been compared with results from 37 age and sex-matched patients (17 males, 20 females) with RA receiving NSAID alone and 27 controls (13 males, 14 females) (Table II).

TBCa was measured by in vivo neutron

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Male/Female ratio, mean (± 1 standard deviation)</th>
<th>age, years post menopause</th>
<th>and disease activity in controls and patients with psoriatic arthritis (PA) and rheumatoid arthritis (RA) included in initial total body calcium measurement. Ranges given in parentheses. Statistical differences between PA and RA patients shown when significant.</th>
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</thead>
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<tr>
<td>Controls (n=36)</td>
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<tr>
<td>Articular Index</td>
<td></td>
<td>5.4±6.9* (0-22)</td>
<td>12.4±5.7* (0-21)</td>
</tr>
</tbody>
</table>

[ESR = erythrocyte sedimentation rate; EMS = duration of early morning stiffness; * = P<0.01].
Table II  Mean age and years post menopause in controls and patients with psoriatic arthritis (PA) and rheumatoid arthritis (RA) undergoing measurements of metacarpal indices (MI) and then rate of change in total body calcium (cTBCa).

<table>
<thead>
<tr>
<th></th>
<th>Controls</th>
<th>Psoriatic Arthritis</th>
<th>Rheumatoid Arthritis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Men</td>
<td>Women</td>
<td>Men</td>
</tr>
<tr>
<td>MI</td>
<td>(n = 13)</td>
<td>(n = 14)</td>
<td>(n = 6)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>44.1</td>
<td>45.4</td>
<td>42.7</td>
</tr>
<tr>
<td>Pre/post-menopause</td>
<td>—</td>
<td>8/6</td>
<td>—</td>
</tr>
<tr>
<td>Years post menopause</td>
<td>—</td>
<td>5.8</td>
<td>—</td>
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</table>

<table>
<thead>
<tr>
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<th>Psoriatic Arthritis</th>
<th>Rheumatoid Arthritis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Men</td>
<td>Women</td>
<td>Men</td>
</tr>
<tr>
<td>cTBCa</td>
<td>—</td>
<td>(n = 5)</td>
<td>(n = 6)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>—</td>
<td>46.8</td>
<td>46.8</td>
</tr>
<tr>
<td>Pre/post-menopause</td>
<td>—</td>
<td>4/0</td>
<td>—</td>
</tr>
<tr>
<td>Years post menopause</td>
<td>—</td>
<td>0</td>
<td>—</td>
</tr>
</tbody>
</table>

Range shown in parentheses

Activation analysis. Patients were irradiated for 40 seconds while standing in a rigid polyethylene activation enclosure, with neutrons from the Edinburgh Medical Research Council Cyclotron. The patients were then transferred to a shadow-shield whole-body counter, where gamma radiation induced by the reaction $^{48}$Ca(n, $\alpha$)$^{49}$Ca was measured for 20 minutes. The patients' TBCa in grams was calculated by comparison with the energy spectrum from an activated anthropomorphic phantom of human dimensions containing a known quantity of calcium. Repeated measurements of the phantom gave a long term precision of 1.8% for a radiation dose of 13mSv (1.3 rem) (9). Individual results were expressed as a percentage of the expected normal value for the patient's skeletal size (arm span) in both sexes and menopausal status in women (9).

The rate of change of TBCa was calculated from 3 or 4 measurements of TBCa undertaken at 6-monthly intervals over an 18-month period. The TBCa in grams was plotted against time and from the slope of the regression line, the rate of change of TBCa was calculated and expressed as a percentage of the initial value. The in vivo precision of the technique, calculated from the deviation of individual points from the lines of regression representing the change in TBCa with time, was 2.9% in a large group of patients with rheumatic diseases (10).

Metacarpal indices were measured by a variation of the technique described by Dequeker (11). A single postero-anterior radiograph of the hands was taken at a constant 1 metre tube-to-film distance using non-screen film. Morphometric measurements were performed at the right 2nd, 3rd, and 4th metacarpals. The length of each metacarpal was determined with a millimetre rule and the
The results of initial TBCa measurement in males, females and both sexes together are shown in Table III. Patients with PA had a mean TBCa value which did not differ significantly from controls. Although the RA patients had reduced TBCa compared with PA patients, the results did not reach statistical significance. The RA patients did have reduced TBCa compared with controls and this reached significance in females and both sexes when considered together. No relationship was found between age and TBCa in controls, patients with PA or patients with RA after the effect of the menopause was excluded.

The annual rate of change of TBCa in PA and RA patients is shown in Table IV. There was a significant mean annual loss of TBCa in the RA patients (-3.7%, p<0.005). The numbers in the PA group were small but the mean rate of change of TBCa was not significant in the male (-0.6%), the female (-1.9%) or the total group (-1.1%). The apparent difference between the rate of change of TBCa in RA and PA patients just failed to reach statistical significance (0.1 > p > 0.05). Annual change in TBCa was not related to disease duration or any one index of disease activity in the PA or RA groups.

The mean metacarpal indices for controls, PA and RA patients are shown in the figure. Cortical area as a percentage of surface area \((D^2-d^2)/DL\) was normal in the small groups of male and female PA patients but significantly reduced in the groups of male and

### Table III

<table>
<thead>
<tr>
<th></th>
<th>Controls</th>
<th>Psoriatic Arthritis</th>
<th>Rheumatoid Arthritis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Males</strong></td>
<td>100.4 ± 11.4</td>
<td>98.5 ± 7.9</td>
<td>94.2 ± 10.2</td>
</tr>
<tr>
<td><strong>Females</strong></td>
<td>99.0 ± 8.0*</td>
<td>99.8 ± 11.1</td>
<td>91.1 ± 10.8*</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>99.7 ± 9.9**</td>
<td>99.1 ± 9.7</td>
<td>93.3 ± 10.5**</td>
</tr>
</tbody>
</table>

\[* = P<0.02; ** = P<0.005\]

Statistical difference between controls and patients with rheumatoid arthritis shown were significant.
Table IV  Mean initial total body calcium (TBCa) expressed in grams and as a percentage of the expected normal value in patients with psoriatic arthritis and rheumatoid arthritis. Mean annual change in TBCa expressed as a percentage of the initial value in male, pre-menopausal and post-menopausal female patients. Range for each measurement shown in parentheses.

<table>
<thead>
<tr>
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<th>Psoriatic arthritis (n = 9)</th>
<th>Rheumatoid arthritis (n = 13)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial TBCa</td>
<td>1022.4 (821-1275)</td>
<td>913.5 (638-1196)</td>
</tr>
<tr>
<td>Initial % TBCa</td>
<td>100.4 (90.1 - 121.4)</td>
<td>92.7 (74.3 - 106.2)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Psoriatic arthritis (n = 9)</th>
<th>Rheumatoid arthritis (n = 13)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annual change in TBCa</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Males</td>
<td>-0.6% (+3.6 to -4.1)</td>
<td>-2.2% (+0.9 to -6.0)</td>
</tr>
<tr>
<td>Pre-menopausal females</td>
<td>-1.9% (+0.5 to -4.3)</td>
<td>-2.4% (-0.1 to -4.4)</td>
</tr>
<tr>
<td>M-Post-menopausal females</td>
<td>-</td>
<td>-4.8% (-2.6 to -7.1)</td>
</tr>
<tr>
<td>Males + females</td>
<td>-1.1%</td>
<td>-3.7%</td>
</tr>
</tbody>
</table>

Female RA patients both in comparison with controls (males: 10.9%, p<0.01; females: 12.8%, p<0.01) and patients with PA (males: 10.6%, p<0.02; females: 14%, p<0.02). $D^2-d^2/DL$ was not related to disease duration or any one index of disease activity in PA or RA with the exception of the Steinbrocker X-ray status (13) which was inversely related to the index in RA patients ($r = -0.35$, $p < 0.05$). Metacarpal indices were not related to age or years post-menopause in controls or patients with the exception of the female RA patients where there were significant relationships with age ($r = -0.50$, $p < 0.05$) and years post-menopause ($r = -0.65$, $p < 0.01$).

**DISCUSSION**

The results of initial total body calcium measurements strongly suggest that total bone mass is normal in patients with psoriatic arthritis. Such a suggestion is in keeping with a previously published view (5) and is given further support by the longitudinal study where the mean change in TBCa in the 9 patients with PA was only 1.1%. This was not statistically significant and as the figure also includes any loss of bone with age, the lack of significant change gives support to the hypothesis that psoriatic arthritis is not usually associated with generalised osteoporosis.

Local osteoporosis is a radiological feature of psoriatic arthritis (1) but we were unable to detect any local bone as measured by metacarpal indices (Fig. 1). However, in the patients with rheumatoid arthritis local bone mass was markedly reduced and was statistically less than that in psoriatic arthri-
Total and peripheral bone mass in patients with PA and RA

FEMALES

<table>
<thead>
<tr>
<th></th>
<th>Control (n=14)</th>
<th>PA (n=5)</th>
<th>RA (n=20)</th>
</tr>
</thead>
</table>

MALES

<table>
<thead>
<tr>
<th></th>
<th>Control (n=13)</th>
<th>PA (n=6)</th>
<th>RA (n=17)</th>
</tr>
</thead>
</table>

Fig. 1: Cortical area as a percentage of surface area [(D^2-d^2)/DL] of the 2nd, 3rd and 4th metacarpal bones in male (▲), pre-menopausal (●) and post-menopausal (○) female controls and patients with psoriatic and rheumatoid arthritis. Results expressed as a percentage of the mean of the controls.

Similar trends were seen in the cross-sectional measurement of TBCa and also in the rate of change of TBCa, with RA patients showing reduced bone mass and a greater rate of bone loss than patients with PA. These results give support to the hypothesis that PA is a more benign condition as far as bone is concerned (3). Our PA patients had significantly less active disease than the RA patients and this may account for the differences in bone mass measurements, raising the possibility that biological mediators of bone resorption may be present in greater concentration in RA than in PA.

The reduction in local bone mass in RA is greater, in percentage terms, than the reduction in TBCa. Although the patients and controls having both bone mass measurements carried out were substantially the same, there were some who did not undergo both measurements especially amongst the controls. Nevertheless, the substantial reduction in metacarpal indices (males: 10.9%, females: 12.8%) suggests a marked degree of local bone loss in RA which, if present in all peripheral bones, might account in part for the reduction in total bone mass shown in this study (males: 6.2%; females: 7.9%). A previous study where peripheral and total bone mass has been measured in the same group of patients showed an 8% reduction in total body bone mineral but a 19% reduction in the bone mineral content of the forearm (14). We have previously reported the lack of relationships between TBCa and any one index of disease duration and activity in patients with RA (15). In addition no
relationship was found in this study between TBCa and Steinbrocker X-ray status. However, metacarpal indices have been shown to relate to X-ray status in RA (16) and this has been confirmed in our present study.

Peripheral bone loss in RA does seem to be related to the long-term disease activity and may account for some, if not all, of the reduction of TBCa. However, further work using improved methods of local bone mass assessment and relating any changes noted to total bone mass in the same groups of patients and controls will be required before this suggestion can be substantiated.

Acknowledgements: This work was supported by grants from the Scottish Hospital Endowment Research Trust and the Arthritis and Rheumatism Council. We are indebted to the Medical Research Council for use of the cyclotron at the Western General Hospital, Edinburgh, to Mrs. N. Brown for data collection and to Mrs. E. Law and Mr. C. Ferrington for assistance with some measurements of total body calcium.

REFERENCES

Changes in Total Body Calcium Following Surgery for Primary Hyperparathyroidism

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Abstract

The aims of this study were to measure the deficit in total body calcium in patients with primary hyperparathyroidism and to observe whether this deficit was reversed by parathyroidectomy. Total body calcium was measured in five women and three men preoperatively, and the mean was found to be 11% below that of age- and sex-matched controls after normalization for postmenopausal age and body size (P < 0.01). Following parathyroidectomy, repeat total body calcium measurements showed an increase of 7.1% over an average period of 14 months (P < 0.03). In patients with primary hyperparathyroidism, total body calcium returns toward normal following surgical removal of the adenoma.

Key Words: Primary Hyperparathyroidism—Parathyroidectomy—Total Body Calcium—In Vivo Neutron Activation Analysis

Introduction

Bone mass may be reduced in primary hyperparathyroidism. This has been found using techniques, such as in vivo total body neutron activation analysis (Cohn et al., 1973), photon absorptiometry of the radius (Pak et al., 1975, Pozzi-Mucelli et al., 1983; Posen et al., 1985), the lumbar spine (Seeman et al., 1982), and the third metacarpal (Geriant et al., 1973). X-ray spectrophotometry of the radius and ulna (Dalen and Ijern, 1974) and quantitative computed tomography of the first and second lumbar vertebrae (Pozzi-Mucelli et al., 1983). These invasive techniques have shown mean reductions in bone of about 10% compared with age-matched controls. Low mean bone mass has been described in bones that are predominantly trabecular (Seenan et al., 1982; Pozzi-Mucelli et al., 1983) as well as in bones that are predominantly cortical in type. However, two studies have shown normal trabecular bone volume in bone biopsy specimens taken from postmenopausal women with primary hyperparathyroidism (Marcus et al., 1984; Charhon et al., 1982).

The skeleton is made up of approximately 80% cortical bone and 20% trabecular bone (Johnson, 1964). Therefore, total body calcium measurements are affected by changes in both trabecular and cortical bone, and the application of this measurement (=total body calcium) to patients with primary hyperparathyroidism should be appropriate. However, two studies of the effect of surgery for primary hyperparathyroidism on total body calcium have shown no overall beneficial effect (Cohn et al., 1973; Hosking et al., 1972). This is in contrast to prospective studies of regional bone mass, which have shown a beneficial effect of parathyroidectomy on bone density (Dalen and Hjern, 1974; Geriant et al., 1973; Leppila et al., 1982). We report here the effect of parathyroidectomy on total body calcium of patients with primary hyperparathyroidism.

Patients and Methods

Patients

Eight patients with surgically proven primary hyperparathyroidism were studied. Age, sex, and clinical information are shown in Table I. Four of the five women were postmenopausal (1-11 years). The weight of the adenoma ranged from 0.17 to 1.99 g. Subperiosteal erosions were present on the hand radiograph of only one patient.

Twenty healthy women (aged 57.5 ± 6.5, SD) and 20 healthy men (aged 54.4 ± 8.6, SD) acted as controls.

Methods

Height was measured by wall-mounted ruler, arm span by wall-mounted graph paper, weight by Avery beam balance, and skinfold thickness by Harpenden calipers. These measurements were used in the normalization of total body calcium (Kennedy et al., 1982).

Biochemistry. Plasma and urinary calcium were measured by atomic absorption spectrophotometry and plasma alkaline phosphatase by the aminoantipyrene method. Immunoreactive parathyroid hormone (PTH) was measured by a modification of the method of Woo and Singer (1974) (hormone antiserum complex precipitated by donkey antiguaena pig antiserum), which used polyvalent antihormone PTH antibody directed mainly against N-terminal PTH.

Total body calcium (TBCa). TBCa was measured by in vivo neutron activation as previously described (Kennedy et al., 1982). The method has a reproducibility of 1.8% (coefficient of variation of repeated measurements on phantoms) and a radiation dose equivalent of 1.3 rem (13 mSv). The reproducibility in vivo is 2.9% (coefficient of variation of repeated measurements on humans.
Tateill et al., 1984). TBCa can be predicted from height in men and arm span in women (Kennedy et al., 1982). The TBCa is expressed as a ratio of the observed to the predicted value (calcium ratio, CaR). The reference range of CaR was calculated from the 40 control subjects (mean ± 2 SD) and was 1.00 ± 0.13.

Statistical methods. The patient and control groups were compared by the Wilcoxon Rank Sum test. Changes following parathyroidectomy were compared using the paired t-test. The relationships between TBCa and biochemical measurements were assessed by linear correlation. Data for alkaline phosphatase and iPTH were analyzed after in-transformation.

Results

Biochemistry. The results are shown in Table I, in which patients are compared with age-matched controls. In addition to the expected elevation in plasma calcium and iPTH, there were also increases in plasma alkaline phosphatase activity and urinary calcium excretion. Following parathyroidectomy plasma calcium fell into the normal range or below (1 patient), and plasma alkaline phosphatase activity fell by a mean of 22 IU/l (P < 0.05).

Total body calcium. The mean TBCa was 828 g (± 206 g, SD). The mean CaR was 0.890 (± 0.015, SD), that is, after normalization for body size and postmenopausal age there was an 11% reduction in bone mass (P < 0.01). Two of the eight patients had results below the 95% confidence intervals of CaR in controls (Fig. 1).

TBCa was remeasured 6–24 months (mean 14 months) after parathyroidectomy (Fig. 1). The mean increase was 59 g (range: 29–142 g), equivalent to a mean rise of 7.1% in TBCa (P < 0.03). Only two of these eight patients had elevated plasma alkaline phosphatase activity preoperatively.

The weak relationships between the change in TBCa and initial urinary calcium excretion (r = 0.40), plasma alkaline phosphatase activity (r = 0.50), plasma iPTH (r = 0.22), and CaR (r = −0.41) did not reach statistical significance (P > 0.05).

Discussion

Cohn et al. (1973) found a mean CaR of 0.89 in nine patients with primary hyperparathyroidism, a result identical to that reported here. However, in the five patients they remeasured following parathyroidectomy there was no overall rise in TBCa. Similarly, Hosking et al. (1972) measured TBCa in 10 patients with primary hyperparathyroidism, and following parathyroidectomy half the patients showed a fall in TBCa, and in one this reduction was 30%. The patients studied by these groups had more severe hypercalcemia [mean plasma calcium 11.9 (Cohn et al., 1973) and 12.3 (Hosking et al., 1972), compared with 10.6 mg/dl in the present study] and higher plasma alkaline phosphatase activity. In patients with more severe hypercalcemia the osteopenia may not be reversible.

Three groups have reported increases in regional bone density following parathyroidectomy. In each study the size of the increase was similar to that reported in the present study. Pak et al. (1975) measured bone density of the distal radius by single photon absorptiometry and found normal values in men and premenopausal women and a 22% reduction in the bone density of postmenopausal women. The same group (Leppila et al., 1982) reported that after parathyroidectomy in postmenopausal women, as in the group of patients with primary hyperparathyroidism as a whole, there was a rise in bone density of 6.4% at 1 year. Over the following 2 years there was no further rise.

Genant et al. (1973) measured bone density of the third digit by photon absorptiometry and found a mean reduction of 13%. One to three years following parathyroidectomy there was a 7% rise in bone density. Dalen and Hjern (1974) measured bone density of the distal radius and ulna by x-ray spectrophotometry and found a mean reduc-
were unable to measure changes. However, measurements have shown that domestic animals in recruitment of bone mass is associated with an increase in bone mass, which is restored almost completely after the osteoblastic formation phase has finished. This ordinarily occurs in 2–3 months. In states of increased resorption there is a predictable increase in this transient pool of missing bone. Parathyroidectomy would cause an abrupt decrease in recruitment of resorption units, but those units already engaged in resorption would complete their remodeling cycles. This would explain why Leppla et al. (1982) found that the rise in bone density was almost complete at 6 months after parathyroidectomy. We were not able to delineate the time course of the change in total body calcium, since the dose of radiation used precludes multiple measurements.

Can the change in bone mass following surgery be predicted? No attempt was made by the three groups who have shown changes in regional bone mass to relate the changes to biochemical variables or initial bone mass. We were unable to predict changes in TBCa from baseline biochemical measurements or CaR.

The reduction in bone mass in patients with primary hyperparathyroidism may be associated with fragility fractures, such as vertebral crush fractures. For example, we found that the mean CaR in 14 women with postmenopausal osteoporosis and vertebral crush fractures was 0.87 (Eastell et al., 1983), a value similar to that found in the present study. Dauphine et al. (1975) found an increased prevalence of vertebral crush fractures in a case-control study of patients with primary hyperparathyroidism. Most of the patients with vertebral crush fractures in the latter study were postmenopausal women. The increase in total body calcium following parathyroidectomy may lower the risk of fracture.

Although all the patients in the present study had plasma calcium values of less than 12 mg/dl, most of them had complications of primary hyperparathyroidism. The results of this study may not apply to patients with mild, asymptomatic primary hyperparathyroidism. However, they do emphasize the need for a randomized controlled trial of the effect of parathyroidectomy on bone mass in such patients.

Acknowledgement: We acknowledge the help given by Dr. J.P. Ashby, Mr. N.S. Brown, and Dr. D.B. Horn in performing the biochemical estimations and thank Professor J.A. Strong, Dr. P.L. Padfield, Dr. J.D. Baird, and Mr. D.W. Hamer-Hodges for allowing us to study patients under their care. R. Eastell was supported by a grant from the Medical Research Council and N.S.J. Kennedy by a grant from the Scottish Home and Health Department.

References


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BONE LOSS IN RHEUMATOID ARTHRITIS AND PRIMARY GENERALIZED OSTEOARTHRITIS: EFFECTS OF CORTICOSTEROIDS, SUPPRESSIVE ANTIRHEUMATIC DRUGS AND CALCIUM SUPPLEMENTS

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Rheumatic Diseases Unit, Northern General Hospital, Department of Medicine, Western General Hospital, and Department of Medical Physics and Medical Engineering, University of Edinburgh, UK

SUMMARY

The annual rate of bone loss in rheumatoid arthritis (RA) and primary generalized osteoarthrosis (PGOA) was determined by measurement of total body calcium (TBCa). The mean annual rate of bone loss in 24 patients with RA treated with nonsteroidal anti-inflammatory drugs (NSAIDs) alone was 3.4%. This rate of bone loss was not reduced in ten RA patients responding to suppressive antirheumatic drugs (4.3%) or seven patients receiving oral calcium supplements (4.5%). The mean annual rate of loss of TBCa in 19 patients with PGOA was 1.6%, a figure which probably represents age-associated bone loss. The rate of bone loss in PGOA was significantly less than that in RA patients not receiving corticosteroids. The mean annual rate of change of TBCa in 30 RA patients receiving corticosteroids (+0.7%) was significantly less than that in any of the other RA groups despite an initial normalized bone mass which was significantly less than in those RA patients receiving NSAIDs alone. The data supported the hypothesis that bone loss occurred early in the course of corticosteroid therapy and thereafter the drugs might have a protective effect on the loss of bone in RA.

KEY WORDS: Bone loss, Total body calcium, Rheumatoid arthritis, Osteoarthrosis, Corticosteroids, Suppressive antirheumatic drugs.

TOTAL bone mass has been shown to be reduced in rheumatoid arthritis (RA) [1-3] and further reduced by low-dose corticosteroid therapy [2, 3]. However, there is no literature on the rate of total bone loss in this disease although loss of bone from metacarpal sites has been shown to occur at rates of 2-4% per annum in patients receiving nonsteroidal anti-inflammatory drugs (NSAIDs), corticosteroids or penicillamine [4, 5].

Initial cross-sectional studies reported by us gave some support to the hypothesis that excess bone loss induced by corticosteroids occurred early in the course of therapy [2]. To test this hypothesis and to examine the effects on the rate of bone loss of oral calcium supplements and suppression of the disease process with antirheumatic drugs, we now report the results of an 18-month longitudinal study of total body calcium in RA. The results have been compared with those from patients with primary generalized osteoarthrosis (PGOA) in whom total bone mass has been found to be normal [6].

METHODS

Patients

Seventy-one patients (30 males, 41 females) with definite or classical RA [7] underwent three or four measurements of total body calcium (TBCa) at 6-monthly intervals over 18 months. Twenty-four patients continued NSAIDs alone, 30 patients continued treatment with low-dose corticosteroids, ten patients commenced therapy with suppressive antirheumatic drugs (SARDs) in the form of gold or penicillamine and seven patients commenced oral calcium supplements (OCS) in the form of calcium lactate gluconate (Sandocal) one tablet t.d.s., giving an elemental calcium dose of 1.2 g/day. Those treated with SARDs responded clinically to the medication. All patients in the RA groups were receiving NSAIDs. A group of 19 patients (2 males, 17 females) with PGOA also had three or four measurements of TBCa and have been included as a 'disease control group'. Eight of these were receiving NSAIDs during the study.
TABLE I

<table>
<thead>
<tr>
<th></th>
<th>RA + NSAIDs</th>
<th>RA + SARDs</th>
<th>RA + steroids</th>
<th>RA + OCS</th>
<th>PGOA</th>
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<tbody>
<tr>
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<td>10</td>
<td>30</td>
<td>7</td>
<td>19</td>
</tr>
<tr>
<td>Male/female</td>
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<td>5/5</td>
<td>10/20</td>
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<td>2/17</td>
</tr>
<tr>
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<td>4/1</td>
<td>5/15</td>
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<td>2/15</td>
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<tr>
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<td>(25-62)</td>
<td>(25-69)</td>
<td>(48-55)</td>
<td>(47-68)</td>
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<td>(0-22)</td>
<td>(0-12)</td>
<td>(0-30)</td>
<td></td>
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</tbody>
</table>

Male/female ratio, mean age and menopausal status of patients with primary generalized osteoarthritis (PGOA) and rheumatoid arthritis (RA) treated with nonsteroidal anti-inflammatory drugs (NSAIDs) alone, additional suppressive antirheumatic drugs (SARDs), additional corticosteroids or additional oral calcium supplements (OCS) completing the 18-month study period (range shown in parentheses).

while the other 11 were treated with simple analgesics. All patients gave informed consent, and ethical approval was obtained from local and national committees.

Clinical and serological measurements

These were undertaken at each 6-monthly visit for TBCa estimation and included measurements of duration of early-morning stiffness, functional status [8], articular index [9], ESR and rheumatoid-factor titre [10]. A Steinbrocker radiographic grading [8] was determined from a hand radiograph at the beginning of the study. All patients had renal and liver function tests and calcium and phosphate biochemistry at the beginning of the study.

Total body calcium measurements

Our method for a single measurement of total body calcium (TBCa) has been fully reported [2, 11]. TBCa was carried out by in-vivo neutron activation analysis and calibration obtained by comparison with an anthropomorphic phantom. A figure for TBCa in grams was produced with an in-vitro precision of 1.8% [11] and an in-vivo precision over a period of 18 months of 2.9% [12]. Individual results in the cross-sectional studies were reported as a percentage of the expected normal value for the patient's skeletal size (arm span) and menopausal state derived from measurements in 40 normal subjects [11].

Longitudinal analysis was undertaken by calculating a best-fit linear regression for the three or four measurements of TBCa in grams. After deviant points were omitted using Chauvenet's criterion [13], the rate of change of TBCa was calculated from the slope of the regression line and expressed as a percentage of the initial value. Analysis demonstrated that the assumption of a linear fit was reasonable. In the longitudinal analysis no correction was made for the patient's skeletal size or menopausal status as the subjects acted as their own controls.

Statistical analysis

This was performed using Student's t test for paired and unpaired variables. The Mann-Whitney test was used for small samples.

RESULTS

The male/female ratio, age and menopausal status of the 71 RA and 19 PGOA patients who completed the longitudinal study are shown in Table I. There was a preponderance of females in all groups with the exception of the RA + OCS group. Patients in the RA + SARDs group were significantly younger than the NSAIDs group (13.4 years, p < 0.02), the steroid (9.8 years, p < 0.05) and the PGOA (15.9 years, p < 0.005) groups.

Indices of disease duration and activity as well as details of corticosteroid therapy are displayed in Table II. The mean disease duration in the SARDs group was shorter than the other treatment groups but not significantly. However, the initial mean ESR in the SARDs group was significantly higher than that in the NSAIDs (34 mm/h, p < 0.025), steroid (32.7 mm/h, p < 0.05) and OCS (41.4 mm/h, p < 0.01) groups. The ESR fell significantly in the SARDs group during the study period (−32.8 mm/h, p < 0.001) but none of the indices changed significantly in the other groups.

The results of initial TBCa, expressed as a percentage of the expected normal values, are shown in Fig. 1. All the patients who continued with NSAIDs and who were to commence SARDs or OCS are contained in the RA plus no steroid group. The steroid-treated group had a significantly lower initial TBCa than the group...
of the differences between OCS: 0.01; significant.

Four RA groups had previously received NSAIDs: 4.5% and OCS (4.5%); SARDs: 4.3% (p < 0.01); OCS: 4.5%; p < 0.01) although the latter group did show a significant annual loss of TBCa (−1.6%, p < 0.01).

The corticosteroid-treated patients have been sub-divided into those taking a low dose of prednisolone (≤5 mg/day) and a moderate dose of the drug (5.1–10 mg/day). Details of disease activity indices and corticosteroid therapy for the two subgroups are shown in Table III. All disease activity indices were greater in the moderate-dose group. The initial bone mass of the two groups was not significantly different but there was a significant difference between the rates of change of TBCa (Fig. 3). The mean rate who had not received corticosteroids (4.6%, p < 0.05) and the patients with PGOA (12.5%, p < 0.01). Both RA groups had a significantly reduced mean TBCa compared with the 40 normal controls previously reported [2, 11].

Comparison of the rate of change of TBCa in the four RA groups (Fig. 2) showed that they all had significant mean annual losses of TBCa (NSAIDs: 3.4%; p < 0.001; SARDs: 4.3%; p < 0.01; OCS: 4.5%; p < 0.01) with the exception of the steroid group. There were no significant differences between the annual rates of change of the NSAIDs, SARDs and OCS groups but all of these lost more calcium than the steroid group (NSAIDs: 4.1%, p < 0.05; SARDs: 5.0%; p < 0.01; OCS: 5.2%, p < 0.02). There was no tendency for the rate of bone loss to diminish in the NSAIDs or OCS groups. In the steroid group there was no significant relationship between the rate of change of TBCa and the daily dose or duration of steroid therapy. The mean rate of change of TBCa was similar in males and females in all of the RA groups.

The 41 RA patients not treated with corticosteroids (i.e. the NSAIDs, SARDs and OCS groups combined) had a significantly increased rate of bone loss compared to the steroid group (−4.3%, p < 0.001) and the PGOA group (−2.0%, p < 0.025) although the latter group did show a significant annual loss of TBCa (−1.6%, p < 0.01).

Results of disease activity indices expressed as the mean initial and mean change in the index with the standard deviation. No change in functional status or radiographic grade occurred during the study.

NSAIDs = nonsteroidal anti-inflammatory drugs; SARDs = suppressive antirheumatic drugs; OCS = oral calcium supplements; ESR = erythrocyte sedimentation rate (mm/h); A.I. = articular index; E.M.S. = duration of early morning stiffness in minutes; F.S. = functional status.

TABLE II

<table>
<thead>
<tr>
<th>Changes of Disease Activity</th>
<th>RA+ NSAIDs</th>
<th>RA+ SARDs</th>
<th>RA+ steroids</th>
<th>RA+ OCS</th>
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</thead>
<tbody>
<tr>
<td>Number</td>
<td>24</td>
<td>10</td>
<td>30</td>
<td>7</td>
</tr>
<tr>
<td>Duration (months)</td>
<td>10</td>
<td>6.2</td>
<td>10.3</td>
<td>10.2</td>
</tr>
<tr>
<td>Disease (range)</td>
<td>(1-35)</td>
<td>(1.5-15)</td>
<td>(1-38)</td>
<td>(2-27)</td>
</tr>
<tr>
<td>Initial ESR</td>
<td>27.7</td>
<td>61.7</td>
<td>29</td>
<td>20.3</td>
</tr>
<tr>
<td>Change in ESR</td>
<td>±20.8</td>
<td>±38.0</td>
<td>±27.1</td>
<td>±13.1</td>
</tr>
<tr>
<td>Initial A.I.</td>
<td>12.7</td>
<td>14</td>
<td>8.7</td>
<td>8.9</td>
</tr>
<tr>
<td>Change in A.I.</td>
<td>±2.7</td>
<td>±7.9</td>
<td>±6.4</td>
<td>±2.7</td>
</tr>
<tr>
<td>Initial E.M.S.</td>
<td>45.4</td>
<td>113</td>
<td>45.3</td>
<td>91.7</td>
</tr>
<tr>
<td>Change in E.M.S.</td>
<td>±21.2</td>
<td>±107</td>
<td>±76.6</td>
<td>±124</td>
</tr>
<tr>
<td>Change in E.M.S. (mg/day)</td>
<td>±4.8</td>
<td>±36</td>
<td>±3.6</td>
<td>±57</td>
</tr>
<tr>
<td>Initial F.S.</td>
<td>±2.0</td>
<td>±2.3</td>
<td>±1.8</td>
<td>±1.6</td>
</tr>
<tr>
<td>Radiographic grade</td>
<td>±0.3</td>
<td>±0.5</td>
<td>±0.5</td>
<td>±0.5</td>
</tr>
<tr>
<td>Prednisolone (mg/day)</td>
<td>±2.6</td>
<td>±2.4</td>
<td>±2.6</td>
<td>±2.4</td>
</tr>
<tr>
<td>Duration of therapy (years)</td>
<td>±1.0</td>
<td>±0.8</td>
<td>±1.1</td>
<td>±0.8</td>
</tr>
</tbody>
</table>

Fig. 1. Initial normalized total body calcium (TBCa) expressed as a percentage of normal in male (▲) and female (●) patients with rheumatoid arthritis (RA) who were receiving nonsteroidal anti-inflammatory drugs (NSAIDs) alone or additional oral corticosteroids at the onset of the study. TBCa of patients with primary generalized osteoarthritis (PGOA) are shown for comparison. Normal range displayed in the hatched area as the mean ±1 standard deviation.
There was bone loss in these studies with NSAIDs alone and further reduced (by 5-8%) in patients treated with nonsteroidal anti-inflammatory drugs (NSAIDs), suppressive anti-inflammatory drugs (SARDs), oral corticosteroids or oral calcium supplements (OCS) as well as in patients with primary generalized osteoarthrosis (PGOA).

The annual rate of change was not related to disease duration or any one index of disease activity in any of the four groups. There was no difference between the rate of change of bone in those patients with seropositive or seronegative disease.

Annual change in TBCa was not significantly correlated with age or years postmenopause in females or age in males. However, in the females, when considering 5-year postmenopause cohorts, the highest mean annual loss of TBCa was in the 5 years immediately after the menopause (0.1-5 years) both in the RA patients who did not receive corticosteroids (n = 6; 4.0%) and in the PGOA patients (n = 5; 3.2%).

**DISCUSSION**

Total bone mass has been shown to be reduced (by 5-8%) in patients with RA treated with NSAIDs alone and further reduced (by 11-15%) with low-dose corticosteroid therapy [2, 3]. Both these studies were cross-sectional and the rate of bone loss was difficult to estimate, particularly as there was no relationship between duration of disease and total bone mass. Our longitudinal study reported here is the first account of the rate of loss of total bone mass in RA. The mean annual rate of loss of calcium in patients treated with NSAIDs alone was a substantial 3.4% per annum. Although the rate is similar to the 2-4% previously shown in longitudinal studies of photon absorptiometry of the forearm [4] and metacarpal indices [4, 5], these bone sites are close to areas of potentially inflamed synovium and therefore might lose bone at a faster rate than the total skeleton.

The annual rate of loss of TBCa in the RA patients receiving NSAIDs alone was greater than one would have expected from our cross-sectional study where TBCa was reduced by 5.3% in males and 6.8% in females [2]. However, in this study, the annual rate of loss of TBCa was relatively increased by expressing it as a percentage of the individual’s initial value and also by omitting the age and menopausal corrections of our original studies [2, 6]. Furthermore, RA patients with more advanced, destructive disease may have an increased rate of bone loss [4] and hence loss of TBCa would tend to accelerate with time.

Loss of bone with age is greater in women than
The number of men measured was too small to comment on the rate of bone loss. It is reassuring to note that the mean loss of 1.5% per annum in women was very similar to the annual reduction in total bone mass in women found in cross-sectional studies of TBCa [11, 16].

The rate of bone loss in RA patients not treated with corticosteroids was significantly greater than in patients with PGOA, suggesting that bone loss, in excess of that due to ageing, is a feature of the former disease.

There were no significant differences between the annual loss of TBCa in the three nonsteroid-treated RA groups. The failure to demonstrate a reduction in the rate of bone loss in those responding to gold or penicillamine, or those taking calcium supplements was supported by finding no diminution in the rate of bone loss. The failure to prevent bone loss with SARDs is disappointing. Preliminary data had suggested that both gold [18-20] and penicillamine [21] might limit the degree of radiological destruction after a year of treatment. Indeed penicillamine may reverse the local bone loss associated with RA as measured by metacarpal indices [5, 22]. Penicillamine is a lathyritic agent which inhibits collagen cross-linking and synthesis, and theoretically could hasten the osteoporotic process by inhibiting collagen maturation in bone [23]. However, there was no significant difference in our study between the men [14], particularly immediately after the menopause [11, 15]. Rather surprisingly the mean rate of loss in our study was similar in males and females in all groups (Fig. 2). There was a tendency for those nearest to the menopause to have the greatest rate of loss of TBCa.

In the light of these observations and because the groups were reasonably matched for years postmenopause, normalization for postmenopause time was not justified. The explanation for the substantial loss of bone in male RA patients is not clear. While there is a slow fall in bone mass with age in men it has been estimated at only 0.7% per annum after the age of 50 [16].

As the rate of bone loss is likely to be linear in males [17] and therefore we were not surprised to find no relationship between bone loss rates and age in men. Nevertheless, the rate of bone loss in males with RA must include an effect of ageing. We have shown total bone mass to be normal in PGOA [6], therefore the rate of bone loss in this condition was attributed to ageing.
rate of loss of TBCa in those treated with penicillamine and gold.

The failure to prevent bone loss with oral calcium supplements is less surprising although oral calcium is thought to reduce the rate of bone loss in postmenopausal osteoporosis [24] and in corticosteroid-treated patients with rheumatoid arthritis [25]. The difference may be due to calcium malabsorption in postmenopausal osteoporosis [26] and in corticosteroid-induced bone loss [27] but not in RA [28]. However, a recent study has shown a mild degree of calcium malabsorption in early disease [29]. All our patients had RA for at least one year. Alternatively, most of the bone loss in RA patients treated with NSAIDs alone may take place at peripheral skeletal sites which may not be amenable to reversal by oral calcium supplements.

The data from the corticosteroid-treated patients give strong support to our hypothesis that bone loss due to prednisolone or its derivatives occurs early in the course of therapy [2]. This hypothesis is consistent with studies of the catabolic and histochemical effects of corticosteroids on bone in both animals [30] and humans [31, 32] and also with longitudinal studies of metacarpal indices [4] and bone histomorphometry [33] in RA and other diseases.

The failure of the steroid-treated patients to lose bone (Fig. 2) despite their initial reduced bone mass (Fig. 1) is perhaps surprising. It might reflect a tendency for those individuals with the lowest initial TBCa to lose the least bone [4]. However, we were unable to detect a relationship between the initial bone mass and the rate of loss in the steroid- or nonsteroid-treated patients. A more exciting possibility is that corticosteroid therapy might limit the degree of local bone damage at periarticular sites by reducing disease activity. Such a suggestion was made initially in the Medical Research Council trials of corticosteroids [34, 35] and has been given support in more recent studies [36, 37]. The subgroup of steroid-treated patients who had received moderate doses of prednisolone (5.1-10 mg/day) had significantly greater mean loss of bone than the low-dose treated group (Fig. 3). The moderate-dose group was significantly older, had more severe disease and were significantly less mobile than the low-dose subgroup. These facts may explain the greater mean loss of TBCa in the former group of patients rather than their steroid therapy. Despite long-standing severe disease, the small group of patients receiving more than 5 mg of prednisolone daily still had a lower rate of bone loss than the other RA patients. It is conceivable that corticosteroids, used in the treatment of RA, could induce bone loss early in the course of therapy, perhaps at an axial site, and thereafter prevent destructive bone loss at appendicular bone sites. Longitudinal studies assessing bone mass at both these bone sites and relating any loss to alteration of erosion formation will be necessary to explore such a hypothesis.

Acknowledgements

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References


Measurement of hand bone mineral content using single-photon absorptiometry

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Abstract. A single photon absorption imaging technique has been developed to assess the bone mass of the hand, especially in patients with rheumatoid arthritis or bronchial asthma. A modified rectilinear scanner images the hand by transmission scanning in a water bath with a 7.4 GBq $^{125}$I source. A microcomputer is used to calculate the bone mineral distribution, and the total bone mineral content (BMC) of the hand is determined from that distribution. The precision (coefficient of variation) of the measurement is 1.9%. A control population of 20 men and 58 women has been studied to determine normal variations in hand bone mineral content with age, sex, body size, hand volume and years since menopause. The normal men are found to have an average hand BMC of 25.1 g with a coefficient of variation (CV) of 22%, which is reduced to 12% by normalising for body size using span. The normal women had an average hand BMC of 18.0 g ± 15%. The CV is reduced to 13% by normalising for span and years post-menopause.

1. Introduction

Single-photon absorptiometry (SPA) is an established technique for the measurement of forearm bone mineral content (BMC) using $^{125}$I as the isotope source (Cameron et al 1968), but until now it had not been applied to the measurement of hand bone mineral. Neutron activation analysis is also used to measure bone in vivo and has been applied to the hand (Catto et al 1973, Maziere et al 1979). Maziere et al (1979) compared activation analysis of hand calcium using $^{252}$Cf (132 controls, 45 osteoporotics) with SPA measurements of BMC in the radial epiphyses (37 controls, 13 osteoporotics) and found a strong correlation between the two. They preferred the NAA technique, which gave a coefficient of variation of 2.4% for repeated measurements on ten controls compared with 3.6% for the SPA. Some SPA measurements on patients were abandoned because results could not be reproduced. Maziere's preference for NAA of the hand was based upon the site of measurement rather than upon the technique. The radiation absorbed dose equivalent to the hand was 7.5 mSv. Smith et al (1981) compared part-body neutron activation analysis of the forearm with SPA of the radius 5 cm proximal to the styloid process, and found the two techniques to be equally sensitive in vivo methods for monitoring changes in calcified bone.

Total bone mass has been shown to be reduced by 5-8% in patients with rheumatoid arthritis who have been treated with non-steroidal anti-inflammatory drugs alone and further reduced in those patients receiving additional corticosteroid therapy.

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(Reid et al 1982, AIS et al 1983). The site of bone loss in rheumatoid arthritis remains in doubt, although substantial reductions in peripheral bone mass may in part account for small reductions in the total bone mass in patients who have not received corticosteroids (Reid et al 1986). Substantial reductions in peripheral bone mass may relate both to periaricular osteoporosis and marginal bone erosion, which are radiological signs of rheumatoid arthritis (Bywaters 1960), especially in regions of the skeleton where small joints are involved, such as the hand. A method of evaluating hand bone mass could be useful not only to clarify the site of bone loss in RA but also as a potential measurement of long-term disease activity and to monitor treatment.

Apparatus was therefore designed to apply the technique of SPA with its established potential for precision and low radiation doses and its simple nucleonic requirements to the measurement of hand bone mineral.

2. Method

A J & P Multipoise dual-headed rectilinear scanner (J & P Engineering Ltd) is rotated until the scanning frames are vertical and modified by the addition of a rigid frame attached to one NaI detector (10 cm diameter, 5 cm thick) carrying a collimated $^{125}I$ source. The 1 mm diameter sealed source has an activity of 7.4 GBq on delivery and is used until it has decayed to about 1.5 GBq. The positions and orientation of the collimated source are adjusted to maximise the detected count rate. The detector collimator is circular with a diameter of 4 or 5 mm; the larger diameter being used to maintain the count rate as the source decays. The change of collimator does not degrade the image. No dead time correction is necessary. The hand is placed in a water bath at 35 °C between the source and detector (figure 1). Various widths of rigid air-filled perspex boxes are placed in the tank alongside the hand to reduce the water thickness as much as possible for an individual. Consideration was given to problems associated with rheumatoid arthritis when designing the apparatus, the most important feature being that some patients would be unable to straighten their fingers. For this reason the (flexed) hand is held immobile in the tank by water-filled balloons, rather than a rigid frame, for the duration of the scan. The apparatus is designed to scan the left hand at a speed of approximately 0.5 cm s$^{-1}$ and a cell size of 4 mm, giving a total scan time of around 15 min.

The detected transmitted counts are collected into a 64 × 64 matrix by a Cromemco microcomputer. This transmission image (shown in figure 2(a)) is used to calculate a bone mineral image (figure 2(b)) by applying the following equation pixel by pixel:

$$M_b = \frac{\rho_b}{(\mu_b \rho_b - \mu_s \rho_s)} \ln I_0/I$$

where $M_b$ is the bone mass per unit area, $\rho_b$ and $\rho_s$ are densities (g cm$^{-3}$) (the subscript ‘B’ indicating bone mineral and ‘S’ soft tissue or water) and $\mu_b$ and $\mu_s$ are mass absorption coefficients (cm$^2$ g$^{-1}$) of the bone mineral and soft tissue or water, respectively. The values used are those given by Cameron et al (1968). $I_0$ is the background intensity; this is determined from three regions of interest as shown in figure 2(a), and $I$ is the transmitted intensity.

A rubber tourniquet is loosely applied around the wrist, 3 cm proximal to the radial styloid. The position is found using a specially made set of dividers. This band is clearly seen in the images (figure 2) and is used to define the limit of the hand. Total hand BMC is the sum of all pixels inside a region of interest, as shown in figure 2(b).

Hand volume, distal to the marker band, is measured by the displacement of water.
3. Experimental results

Two to six repeat measurements were performed on ten volunteers on separate occasions over the life of a source (approximately five months), giving a precision (coefficient of variation) of 1.9%. The absorbed dose, measured using thermoluminescent dosimetry, was 6 \( \mu \)Gy per scan.

The limit of measurement, 3 cm proximal to the radial styloid, was chosen arbitrarily. Figure 3 shows the variation in total BMC by movement of the upper limit for a typical male hand. Hand bone mineral is not unduly influenced by the position of the limit, and the relationship is linear. Errors introduced in the result by repositioning the marker band are incorporated in the precision measurement.

The measurement of hand BMC is quite a sensitive function of the measured background level; a 1% change in background giving a 2% change in hand BMC for an average female hand with a BMC of 19 g. The background count rate is well estimated; the counts in the three background regions give a coefficient of variation of 0.1% and the integral non-uniformity is less than 0.5%. This slight non-uniformity results from bowing of the tank walls when it is filled with water. More importantly, fat attenuates less than water or soft tissue and fat around the hand will give a locally raised background level. Increased transmission around the wrist may be clearly seen on some transmission images. The worst case seen, in a 109 kg female patient, shown in figure 4, gave an increased transmission of 15% at the side of the wrist. Using a simple cylindrical anatomy this gives a maximum localised error over the bone of 1.5% in transmitted counts or 3% in the BMC. However, this underestimate of the BMC is non-uniform, being directly dependent on an individual's fat distribution, and cannot be accurately measured. The effect is not included in the precision measurement as it predominantly affects the accuracy rather than the reproducibility.
Figure 2. (a) Regions used to determine the background level on the transmission image. (b) Region used to determine hand BMC using the bone mineral image. The apparently very dense band around the wrist is the marker.
Hand bone mineral content using absorptiometry

Figure 3. Variation of hand BMC with position of wrist limit marker.

Figure 4. Transmission image of the hand of a 109 kg female showing increased transmission due to fat around the wrist.

4. Normalisation procedures

To establish a range of normal values for hand BMC, 78 normal volunteers were measured: 20 males (four aged 20–29, seven aged 30–39, two aged 40–49, four aged 50–59 and three aged 60–69) and 58 females (thirteen aged 20–29, seven aged 30–39, fourteen aged 40–49, twelve aged 50–59, eight aged 60–69 and four over 70). All the subjects were active, in good health and without any history of disease that might be expected to influence bone metabolism. The variation in hand BMC and in size is shown in table 1. The females ranged from 12.9 to 25.6 g with a mean of $18.0 \pm 15\%$ (CV), while the males ranged from 14.6 to 34.9 g with a mean of $25.1 \pm 22\%$ (CV).
The dependence of measured hand BMC upon size parameters (hand volume, arm span, height and weight) and age was investigated using multiple regression analysis. The more significant results for men are shown in table 2(a) and for women in table 2(b). For women, years post-menopause is considered as an alternative age parameter. The biological variation in BMC due to size and age could be reduced by expressing the subjects' results as a ratio BMC/BMCP. BMCp is a predicted value obtained by using equations derived from the multiple regression analyses and is given in tables 2(a) and (b). These tables also contain the coefficients of variation (CV) of BMC/BMCP for the control population when these predictive expressions are applied.

For men (table 2(a)), hand BMC is found to correlate equally well with hand volume and arm span. However, the correlation with age alone is poor (r = 0.25, P > 0.3), and the addition of age does not improve the correlation with either span or volume.

Amongst the female controls the biological variation was initially smaller (CV = 15%) (table 1), but correlations with size and age parameters were much weaker and hence a much smaller reduction in the CV was achieved by normalisation. Hand BMC correlates with span (r = 0.36, P < 0.01) or height (r = 0.32, P < 0.02) alone but not with volume alone. For women, both age (r = 0.33, P < 0.02) and years post-menopause (r = 0.31, P < 0.02) correlate with hand BMC and may be combined with a size parameter to improve the result. When this is done, age results in the lowest CV, giving 12.5% with volume and 12.2% with arm span. However, the most significant correlation (r = 0.48, P = 0.001) is for span and years post-menopause, a combination which gives a CV of 12.8%.

Figure 5 is the result of plotting hand BMC normalised using the expression given for span in table 2(b) (BMCp = 9.125 S1.25) against age. By chance, all women over 50 are post-menopausal. The linear regressions are shown for the whole population, the pre-menopausal women and the post-menopausal women. The population as a whole shows a loss of 0.25% hand BMC per year with a significance P < 0.05. The pre-menopausal group show an increase which is not significant and the post-menopausal group show a loss of 0.4% hand BMC per year which is also insignificant.

5. Discussion

Single-photon absorptiometry can be usefully employed to make precise measurements of hand BMC and hence to study local bone loss. It is relatively insensitive to variations in placing and replacing the marker band, the effect of which is incorporated in the precision estimate. Fat deposits around the hand or wrist will result in a small inaccuracy in some patients.
Table 2. (a) Normalisation of male hand BMC for age and size parameters, showing the equations for the predicted hand BMC (BMC_p) with coefficients of variation (CV) of the normalised controls, correlation coefficients (r) and significances (P). Correlation with age alone was poor (P>0.3) and has been omitted.

<table>
<thead>
<tr>
<th>Body size parameters</th>
<th>Age parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Age (A) in years</td>
</tr>
<tr>
<td>Hand Volume in litres</td>
<td>BMC_p = 2.12 \times 10^{-2} V^{1.7}</td>
</tr>
<tr>
<td></td>
<td>CV = 11.8%</td>
</tr>
<tr>
<td></td>
<td>r = 0.81</td>
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<td>P &lt; 0.0001</td>
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<tr>
<td>Span (S) in metres</td>
<td>BMC_p = 3.07 S^{3.42}</td>
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<tr>
<td></td>
<td>CV = 11.9%</td>
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<tr>
<td></td>
<td>r = 0.80</td>
</tr>
<tr>
<td></td>
<td>P &lt; 0.0001</td>
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<tr>
<td>Height (H) in metres</td>
<td>BMC_p = 3.28 H^{3.6}</td>
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<tr>
<td></td>
<td>CV = 15.0%</td>
</tr>
<tr>
<td></td>
<td>r = 0.67</td>
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<tr>
<td></td>
<td>P &lt; 0.0001</td>
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</table>

(b) Normalisation of female hand BMC for age and size parameters, showing the equations for the predicted hand BMC (BMC_p) with coefficients of variation (CV), correlation coefficients (r) and significances (P).

<table>
<thead>
<tr>
<th>Body size parameters</th>
<th>Age (A) in years</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Years post-menopause (Y) in years</td>
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<tr>
<td>Height in metres</td>
<td>BMC_p = 11.03 V^{1.12}</td>
</tr>
<tr>
<td></td>
<td>CV = 25.4%</td>
</tr>
<tr>
<td></td>
<td>r = 0.07</td>
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<tr>
<td></td>
<td>P &gt; 0.2</td>
</tr>
<tr>
<td>Span (S) in metres</td>
<td>BMC_p = 9.12 S^{1.29}</td>
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<tr>
<td></td>
<td>CV = 13.8%</td>
</tr>
<tr>
<td></td>
<td>r = 0.36</td>
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<tr>
<td></td>
<td>P &lt; 0.01</td>
</tr>
<tr>
<td>Hand volume in litres</td>
<td>BMC_p = 12.45 S^{0.91}</td>
</tr>
<tr>
<td></td>
<td>CV = 12.2%</td>
</tr>
<tr>
<td></td>
<td>r = 0.43</td>
</tr>
<tr>
<td></td>
<td>P &lt; 0.01</td>
</tr>
</tbody>
</table>

The variation in the control populations may be reduced by normalising for size and age. In the male controls there is no significant loss of hand BMC with age, and normalising for either span or hand volume gives an appreciable reduction in the coefficient of variation. We had thought that, since any variation due to mispositioning...
Figure 5. Variation of hand BMC normalised using span alone with age in female controls. •, pre-menopausal—the full line represents the best fit to the data and has a gradient = 0.16% yr⁻¹, significance \( P > 0.2 \); O, post-menopausal—the full line representing the best fit with a gradient = -0.41% yr⁻¹, significance \( P > 0.2 \). The broken line, representing the best fit to all the data, has a gradient = -0.24% yr⁻¹, significance \( P < 0.05 \).

of the marker band would be present in both the hand BMC and volume measurements, normalising for volume would help eliminate this source of error. However, if such an effect is present it is masked by the greater biological variation (15%) in this parameter, probably due to differences in soft tissue. The benefits of normalising the female control results are much less, with only a 2-3% reduction in cv although, since there is less biological variation initially, the normalised male and female controls have very similar cv.

The range of hand BMC obtained agrees well with the results found by Maziere et al (1979) who found 4.5-16 g of calcium distal to the radial and ulna styloid. Catto et al (1973) gave results for four patients in the range 5.9-13.6 g.

The normalisation procedures used by Maziere and co-workers are more direct, being based on an estimate of hand bone volume from a plane radiograph, and in the 20-60 age range they find a cv of 9% in men and 11% in women. However, this normalisation procedure is not open to us since we cannot expect to be able to flatten our patients' hands to obtain a suitable radiograph.

Age related loss of hand BMC is more apparent amongst the females although, as shown in figure 5, the rate of loss is small and the statistical significance is poor. These loss rates are much less than those reported previously for total body calcium where a post-menopausal loss of 1.5% per year was observed (Kennedy et al 1982). The age decade data given by Maziere et al (1979) for hand calcium suggest a loss rate of 1.3% per year amongst post-menopausal women if the menopause is assumed to occur at an age of 50. Dequeker (1976) used various hand bone indices from radiogrammetry
which suggest a loss rate of just under 1% per year following the menopause. However, none of these measurements are directly comparable, nor is it possible to calculate the significances from the published data.

The apparent change in the hand BMC loss rate at menopause, together with the improved correlation, leads us to use span and years post-menopause for normalising our female hand BMC. Span alone will be used for the male data.

Acknowledgments

We wish to express our gratitude to Miss Elizabeth A Jones and Dr David W Pye for their assistance, Mr George Campbell for his workshop skills and most importantly to our 78 'normal' volunteers.

This work was supported by a grant from the Arthritis and Rheumatism Council.

Résumé

Mesure du contenu minéral des os de la main par absorptiométrie à photon unique.

Les auteurs ont développé une technique d'imagerie d'absorption à photon unique pour évaluer la masse osseuse de la main, en particulier pour des patients atteints de polyarthrite rhumatoïde ou d'asthme bronchique. Les images de la main ont été obtenues sur un scintigraphe à balayage modifié, par des mesures de transmission dans un bac d'eau utilisant une source de 125I de 7,4 GBq. Un micro-ordinateur permet de calculer la distribution minérale dans l'os, et le contenu minéral total des os (BMC) de la main est déterminé à partir de cette distribution. La dispersion des mesures est de 1,9% (coefficient de variation). Une population de contrôle de 20 hommes et de 58 femmes a été étudiée afin de déterminer les variations normales du contenu minéral osseux de la main en fonction de l'âge, du sexe, de la taille du corps, du volume de la main, et du temps écoulé après la ménopause. Les hommes normaux (témoins) ont un BMC moyen de la main de 25,1 g avec un coefficient de variation (cv) de 22% qui peut être amené à 12% en normalisant par rapport à la taille du corps, estimée avec l'empan. Les femmes normales (témoins) ont un BMC moyen de la main de 18,0 g ± 15%. Le cv est réduit à 13% en normalisant par rapport à l'empan et le temps écoulé après la ménopause.

Zusammenfassung

Messung des Knochenmineralgehaltes der Hand mit Hilfe der Einzel-Photon-Absorptiometrie.


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**Scientific note**

**In vivo** precision of total body calcium and sodium measurements by neutron activation analysis

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1. Introduction

**In vivo** neutron activation analysis is an important established technique for determining body elemental composition. It may be used to determine both the absolute amount of some body elements or changes with time (Cohn 1981).

The MRC cyclotron in Edinburgh has been used as a neutron source for total body neutron activation analysis (TBNA) of calcium (Kennedy et al 1982), sodium and chlorine (Kennedy et al 1983) using the reactions $^{48}$Ca($n, \gamma$)$^{49}$Ca, $^{23}$Na($n, \gamma$)$^{24}$Na and $^{37}$Cl($n, \gamma$)$^{38}$Cl respectively.

We have derived absolute quantities of calcium and sodium by identifying and correcting for the body dimensions that affect the efficiency of activation and gamma-ray measurement (Kennedy et al 1982). However, it is often of more importance, particularly when measuring calcium, to determine changes within a patient over a long period of time.

The ability to detect such changes in body elements depends upon the reproducibility or precision of the method. This is defined as the coefficient of variation of repeated measurements over varying periods of time. The precision quoted by various centres making TBNA estimations of total body calcium (TBCa), based on measurements of phantoms or cadavers, is shown in table 1. No measurements have yet been published of the long term **in vivo** precision of TBNA based on repeated patient measurements. Since the typical dose is about 10 mSv it would not be ethical to activate human subjects repeatedly solely to obtain this information. However if patients are being measured over a long period of time to monitor either their bone loss as a result of disease or a change in bone calcium as a result of treatment, it is possible to use these results to obtain a measurement of the long term precision of TBNA **in vivo**.

2. Method

One hundred and fifty nine male and female subjects participated in a clinical study to investigate bone loss in various rheumatic diseases. These patients attended for TBNA on three or four visits, six months apart, over a period of up to 18 months. On
Table 1. Precision and dose for TBCa and TBNa measurements.

<table>
<thead>
<tr>
<th>Neutron source</th>
<th>TBCa precision (%)</th>
<th>Method</th>
<th>TBNa precision (%)</th>
<th>Dose equivalent (mSv)</th>
<th>Authors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cyclotron</td>
<td>2-3</td>
<td>8 measurements</td>
<td>—</td>
<td>15*</td>
<td>Chamberlain et al</td>
</tr>
<tr>
<td></td>
<td></td>
<td>of 1 cadaver</td>
<td></td>
<td></td>
<td>(1968)</td>
</tr>
<tr>
<td>Neutron generator</td>
<td>1.7</td>
<td>5 measurements</td>
<td>—</td>
<td>6.4*</td>
<td>Cohn et al (1970)</td>
</tr>
<tr>
<td>14 Pu-Be</td>
<td>1.1</td>
<td>Phantom</td>
<td>—</td>
<td>2.8*</td>
<td>Cohn et al (1972, 1976)</td>
</tr>
<tr>
<td>Two neutron</td>
<td>2.9</td>
<td>35+ measurements</td>
<td>—</td>
<td>10</td>
<td>Boddy et al (1973)</td>
</tr>
<tr>
<td>generators</td>
<td></td>
<td>of various phantoms</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cyclotron</td>
<td>2</td>
<td>Cadaver</td>
<td>—</td>
<td>10</td>
<td>Spinks et al (1977)</td>
</tr>
<tr>
<td>Cyclotron</td>
<td>1.8</td>
<td>13 measurements</td>
<td>2.1</td>
<td>13*</td>
<td>Kennedy et al (1982, 1983)</td>
</tr>
<tr>
<td>Neutron generator</td>
<td>3.6</td>
<td>43 measurements</td>
<td>—</td>
<td>5</td>
<td>Burkinshaw (1985)</td>
</tr>
<tr>
<td>generator</td>
<td></td>
<td>of 1 phantom</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>over 14 months</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neutron generator</td>
<td>3.0</td>
<td>5 measurements</td>
<td>1.9</td>
<td>10</td>
<td>Shafai et al (1983)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>of 1 phantom</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* A quality factor or relative biological effectiveness of 10 is given in the references cited.

Each occasion they attended, simultaneous measurement was made of TBCa, total body sodium (TBNa), and total body chlorine (TBCl). The patients were activated in their normal clothing. This means that no reliable interpretation can be placed on the TBCl estimates as these are influenced by the chlorine content of their clothes, especially due to the use of thermal underwear in winter.

 Corrections were applied for patient height, body thickness and fat layer and for air gap in the irradiation chamber (Kennedy et al 1982, 1983). Observed counts were related to those from a sodium standard irradiated at the same time and overall reproducibility was monitored by periodic activation of an anthropomorphic model.

As a consequence of their disease or treatment, the patients' TBCa and TBNa might be expected to change with time and this was observed in most patient groups (Reid et al 1984). In calculating the in vivo precision, it was assumed that the changes would be linear over the period concerned. The results of patients who attended three or four times were analysed by fitting a linear regression to the TBCa and TBNa results plotted against time. The coefficient of variation of the deviation of the points from the regression line was calculated assuming a normal distribution and (n-2) degrees of freedom, where n is the number of visits for that patient. Individual points which deviated by more than 3.4 standard deviations from the regression line were eliminated by applying Chauvenet's criterion (Documenta Geigy 1956). The mean of these coefficients of variation is taken to be the in vivo precision.

3. Results

After elimination for non-attendance and the application of Chauvenet's criterion, which removed 15 points, 590 points remained representing 156 patients with three or four TBCa results. The distribution of coefficients of variation is shown in figure 1.
In vivo precision of TBCa and TBNa

The in vivo precision is 2.9% compared with a precision of 1.8% obtained by repeated activation of an anthropomorphic model over a period of three months. 158 subjects with three or four TBNa results remained. The distribution of coefficients of variation is shown in figure 2. This gives an in vivo precision of 3.6% compared with a precision of 2.1% obtained from measurements of an anthropomorphic model.

4. Discussion

TBNA is in regular use for determining body composition. In longitudinal studies it has successfully demonstrated changes in total body minerals as a result of disease or its treatment (Eastell et al 1985, Gruber et al 1984). Longitudinal studies have been over a variety of periods mainly in the range of six months to two years. The application of TBNA, however, requires a knowledge of the precision of the estimate. In the past this has been obtained from phantoms activated within a relatively short period of time. These models must represent the patient both in chemical composition and in the distribution of the elements. This has led to the development of complicated models, such as those of Oxby and Brooks (1979). Our technique of estimating the precision overcomes the limitations of anthropomorphic models. All the uncertainties in a single estimate are incorporated together with any long term variations in patient parameters (e.g. weight) and in the equipment. The assumption of linear changes in calcium and sodium seems to be reasonable (Tothill et al 1984) but if the changes are non-linear then the method is likely to overestimate the coefficient of variation. The true precision would then be better than that estimated by this.
method. The distribution of the deviations from the regression lines are shown in figures 1 and 2. The use of the mean to describe the coefficient of variation is a conservative choice.

Inspection of table 1 shows that our in vitro precision compares favourably with that obtained by other authors. The difference observed between long term measurements in vivo and measurements based on the phantom (2.9 and 1.8% respectively for TBCa, and 3.6 and 2.1% for TBNa) is not surprising. Corrections for patient size and movement, neutron beam shape, and irradiation and measurement timing may not be completely adequate. The precision is still very satisfactory and allows the study of the effects of disease or treatment to be made with relatively small groups of subjects. We cannot tell how much deterioration of precision to be expected with other systems as no previous estimates have been made in vivo. Cohn (1982) has reported a precision of 2.6% over a 4–5 year patient study compared with a phantom precision of 1% but without details of how this was measured. Some centres are able to use anthropomorphic models to monitor precision over long periods (Burkinshaw 1985) but this may not fully allow for effects due to changes in the patients' weight or body build between activations.

Acknowledgments

We should like to thank Mrs Norma Brown, Mrs E M Law, Mr J R Williams and the staff of the MRC Cyclotron Unit in Edinburgh. The research was funded by the Scottish Hospitals Endowment Research Trust and by the Arthritis and Rheumatism Council.

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The uptake of $^{99m}$Tc-MDP was studied in 73 patients after a tibial fracture. The image obtained five minutes after injection during a period between one and four weeks after fracture was found to be related to the incidence of non-union after six months. A ratio of 1.3 between the uptake at the fracture site and at normal bone adjacent to it predicted non-union in an individual patient with a sensitivity of about 70%, and a specificity of 90%.

It has been recognised for many years that there is an increased uptake of bone scanning agents at the site of a fracture and there have been several reports on bone scans as indicators of fracture healing. Various radionuclides such as $^{32}$P (Tucker 1950), $^{82}$Sr (Bauer and Wendeborg 1959; Muheim 1973), $^{87m}$Sr (Illingworth and Schiess 1971; Johansen 1973) and $^{18}$F (Riggens et al. 1974) have been used to investigate fracture healing without a great deal of success. Technetium $^{99m}$-phosphate compounds such as $^{99m}$Tc-Sn-pyrophosphate (Lund et al. 1978) have also been investigated but it is the compound $^{99m}$Tc-methylene diphosphonate (MDP) which seems the most promising in the investigation of fracture healing (Hughes 1980).

Most of the research using MDP has concentrated on the qualitative assessment of scans recorded at two to four hours after injection. Mutin (1979) and Desai et al. (1980), however, showed that there was increased uptake at fractures sites. Auchincloss and Watt (1982), in a study of fractures of the lower leg, demonstrated a significant relationship between uptake at the fracture site and time to union; they suggested that the two-hour scan may be of value in the prediction of delayed healing, though there was a considerable degree of overlap between the groups in their study. However, Jacobs et al. (1981) and Gregg, Barsoum and Clayton (1983) came to the opposite conclusion. The only previous report on MDP uptake very soon after injection was a small study (Jacobs et al. 1981) which demonstrated a relationship between the rate of uptake and time to union.

It has been shown that the MDP uptake at two to four hours after injection is a function of both regional blood flow and bone formation (Lavender et al. 1979). Given the relatively slow transfer of MDP from blood through the bone fluids to bone itself with a peak at about one hour (Hughes et al. 1978; Maklerand Charkes 1980), the uptake of MDP soon after injection should be largely dependent on blood flow alone. If the interruption of blood flow in the bone is the reason for the number of tibial fractures which fail to unite, then MDP uptake, particularly soon after injection, could be a valuable predictor of non-union. If bone blood flow is the predominant factor, then it is important that studies be made when the reactive increase in blood flow is maximal.

Previous studies using MDP to investigate fracture healing (Jacobs et al. 1981; Auchincloss and Watt 1982) have demonstrated that various indices are related to the time to union. Our aim was to conduct a prospective study to investigate the differences in early uptake of MDP between patients obtaining union in normal time and those developing non-union. We hoped to develop a technique which could predict non-union soon after fracture in order to allow consideration of an early prophylactic operation.

**PATIENTS AND METHODS**

All patients aged from 16 to 55 years with a fractured tibia and a normal contralateral limb admitted to the Royal Infirmary, Edinburgh from January 1982 to July
considered able were plateau. time respect flow is followed immediate chosen for six and 12 weeks after 2, four and 12 weeks for the clinical clinical methods accepted all of management. There had been declined to of clinical because of interest series, but 13 were subsequently withdrawn because of clinical problems or failure to attend. We accepted all types of tibial fracture (Table 1) and all methods of management. There were 27 patients aged 15 to 19 years, 36 aged 20 to 29 years, 12 aged 30 to 39 years and 11 aged from 40 to 55 years.

In addition to the routine of management, each patient was examined by the same orthopaedic surgeon at monthly intervals in order to assess the state of healing of the fracture. He had no access to the results of the bone scan but, using a combination of radiographic and clinical criteria, recorded each patient as showing union, delayed union or non-union. Normal union was recorded for the fractures which had united within four months, and delayed union where fractures had united between four and six months. Non-union was recorded for those fractures which had not united by six months.

Our aim was to perform quantitative bone scans at two, six and 12 weeks after the injury. These times were chosen for a number of reasons. After a fracture there is an immediate decrease in blood supply to that bone; this is followed by an increase which reaches a maximum after 10 to 14 days in the canine tibia (Paradis and Kelly 1975; McCarthy and Hughes 1984). This increased blood flow is associated with new bone formation as the fracture heals. Two weeks after injury is a suitable time with respect to the mobility of the patient, and is also the time when the increased blood flow should reach a plateau. Operation for failure to unite is commonly considered after about 12 weeks; Jacobs et al. (1981) were able to show a difference in uptake between cases of delayed and of non-union at this time. We therefore performed a scan at 12 weeks and also at an intermediate time to assess the healing process. The work of Lund et al. (1978) suggested that six weeks was an appropriate time.

At each visit, each patient was given an intravenous dose of 200 MBq of 99mTc-MDP; this is one-third of the dose normally given for a routine bone scan. Before the injection the radioactivity of the syringe was measured by scintillation counter against a 99mTc-MDP standard which was approximately 1% of the injected activity. The patient was positioned supine on a scanning bed below which a gamma camera had both lower legs in its field of view. A Technicare 110 gamma camera with a high sensitivity collimator and an MCS-560 computer processor were used. The 99mTc-MDP was administered as a bolus injection in the right antecubital fossa and was flushed through with 20 ml of saline. A dynamic series of 30 × 10 second images were obtained from the time of injection for 300 seconds. After this a static image was collected from 300 to 800 seconds post-injection. This static image was termed the "early uptake". About two hours the patient returned to the scanning room and was positioned as before. A 300 second static scan was then collected which was termed the "late uptake". This late uptake scan was obtained with the same gamma camera but with a medium resolution, lower sensitivity collimator. After each early and late uptake scan the 99mTc-MDP standard was placed in the centre of the collimator field and measured for 100 seconds so that the absolute uptake at the fracture site could be determined.

The results from the bone scans were analysed using the computer to measure the radioactivity within regions of interest (Fig. 1). These were defined as: A, round the
Table II. The ratios between activity levels on "early uptake" and "late uptake" scans at the three visits for each of the groups (mean ± s.d.)

<table>
<thead>
<tr>
<th>Ratios</th>
<th>A/B</th>
<th>A/C</th>
<th>C/D</th>
<th>A/B ± C/D</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>U</td>
<td>D</td>
<td>NU</td>
<td>U</td>
</tr>
<tr>
<td><strong>First visit</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early</td>
<td>2.78</td>
<td>1.95</td>
<td>2.38</td>
<td>1.09</td>
</tr>
<tr>
<td></td>
<td>±1.18</td>
<td>±0.67</td>
<td>±0.74</td>
<td>±0.39</td>
</tr>
<tr>
<td>Late</td>
<td>4.97</td>
<td>4.31</td>
<td>4.56</td>
<td>4.25</td>
</tr>
<tr>
<td></td>
<td>±1.76</td>
<td>±1.60</td>
<td>±1.33</td>
<td>±0.99</td>
</tr>
<tr>
<td><strong>Second visit</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early</td>
<td>2.32</td>
<td>2.16</td>
<td>2.03</td>
<td>1.24</td>
</tr>
<tr>
<td></td>
<td>±0.73</td>
<td>±0.61</td>
<td>±0.57</td>
<td>±0.55</td>
</tr>
<tr>
<td>Late</td>
<td>6.78</td>
<td>6.34</td>
<td>5.45</td>
<td>5.23</td>
</tr>
<tr>
<td></td>
<td>±2.32</td>
<td>±2.55</td>
<td>±1.31</td>
<td>±1.32</td>
</tr>
<tr>
<td><strong>Third visit</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early</td>
<td>2.62</td>
<td>2.79</td>
<td>2.10</td>
<td>2.94</td>
</tr>
<tr>
<td></td>
<td>±1.29</td>
<td>±0.99</td>
<td>±0.63</td>
<td>±0.57</td>
</tr>
<tr>
<td>Late</td>
<td>7.54</td>
<td>9.13</td>
<td>6.63</td>
<td>4.24</td>
</tr>
<tr>
<td></td>
<td>±2.61</td>
<td>±2.89</td>
<td>±2.67</td>
<td>±1.79</td>
</tr>
</tbody>
</table>

U, union; D, delayed union; NU, non-union.

RESULTS

The 73 patients admitted into the study, not all attended on all three occasions. Although it had been intended to obtain scans at two weeks, precise timing was not possible, so we analysed scans taken at the "first visit" from between one and four weeks after fracture. Two patients had evidence of infection at the time of scan and these results were excluded since infection is known to increase MDP uptake (Hughes et al. 1986). There were fewer exclusions from the results of scans taken between five and eight weeks after fracture, referred to as the "second visit", or between 10 and 15 weeks after injury, the "third visit".

The mean ratios obtained at each visit for the three groups of patients are listed in Table II. As would be expected, the ratios of most interest (A/B and A/C) tend to decrease in the patients who eventually showed non-union. For these ratios the "non-union" group has a lower mean than the "union" group for both the early...
Table III. The t value and significance of the difference between the groups with union and with non-union (Table II) for the ratios A/C and A/B on early and late uptake

<table>
<thead>
<tr>
<th></th>
<th>Ratio</th>
<th>t value of difference</th>
<th>Significance</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>First visit</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early uptake</td>
<td>A/C</td>
<td>3.2</td>
<td>p &lt; 0.005</td>
</tr>
<tr>
<td></td>
<td>A/B</td>
<td>1.3</td>
<td>NS</td>
</tr>
<tr>
<td>Late uptake</td>
<td>A/C</td>
<td>3.3</td>
<td>p &lt; 0.005</td>
</tr>
<tr>
<td></td>
<td>A/B</td>
<td>0.8</td>
<td>NS</td>
</tr>
<tr>
<td><strong>Second visit</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early uptake</td>
<td>A/C</td>
<td>0.2</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>A/B</td>
<td>1.7</td>
<td>NS</td>
</tr>
<tr>
<td>Late uptake</td>
<td>A/C</td>
<td>2.6</td>
<td>p &lt; 0.05</td>
</tr>
<tr>
<td></td>
<td>A/B</td>
<td>2.2</td>
<td>p &lt; 0.05</td>
</tr>
<tr>
<td><strong>Third visit</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early uptake</td>
<td>A/C</td>
<td>2.1</td>
<td>p &lt; 0.05</td>
</tr>
<tr>
<td></td>
<td>A/B</td>
<td>1.6</td>
<td>NS</td>
</tr>
<tr>
<td>Late uptake</td>
<td>A/C</td>
<td>2.3</td>
<td>p &lt; 0.05</td>
</tr>
<tr>
<td></td>
<td>A/B</td>
<td>0.9</td>
<td>NS</td>
</tr>
</tbody>
</table>

NS, not significant

The results of the early uptake A/C ratio at the first visit related to the time to union. The dashed line represents the ratio 1.3.

Fig. 2

The results for the late uptake A/B ratio at the second visit related to the time to union.

Fig. 3
the late uptake A/C ratio was higher than the corresponding early uptake ratio, this was due to very high A/C values in the union group at late uptake even though separation was not as good. The possibility of improving the separation by using both early and late A/C ratios at the first visit was investigated. The product of these ratios gave no better separation between the groups than the early uptake A/C ratio at the first visit on its own. The late uptake A/B ratio at the second visit is shown in Figure 3.

The absolute uptake in region A, as a percentage of the injected activity, was calculated but no significant differences were found between the union and non-union groups at either the early or late uptakes at any of the three visits. No significant differences between the groups were found when the rate of uptake immediately after injection was calculated from the initial series of dynamic images.

The predictive value of the two-week early uptake A/C ratio was investigated. The intra-observer variation in the analysis of these results was found to be a mean of 6% between the measured uptake at the fracture site on two occasions six months apart. More important, however, is the inter-observer variation, as the investigator's choice of the region C will have important implications. The sensitivity and specificity of the early uptake A/C ratio at two weeks was calculated independently by two of the authors using different A/C cut-off values and the results are given in Table IV. As expected, there are differences in the ratios and these are reflected in the variation in the sensitivity and specificity. It appears that a cut-off value of 1.3 for the ratios provides an acceptable compromise between sensitivity and specificity. Sensitivity varied from 63% to 84% and the specificity was from 91% to 88% for the two operators. Neither the use of longitudinal profiles or of computer programs to define the region of interest provided any improvement over a skilled operator, and, in fact, the results were significantly worse.

The timing of the early scan at the first visit is important in relation to the time of injection but less so with regard to the time since fracture. In eight patients, uptake was measured from a 15-minute dynamic series of images; this showed a steady increase in the activity at the fracture site, and there was little variation from patient to patient (Fig. 4). This finding suggests that the A/C ratios illustrated in Figure 2 are likely to differ significantly if the measurements are performed at a different time after injection of the isotope.

The variation of early uptake A/C ratio with time since fracture in patients having normal union is illustrated in Figure 5. During the period from one week to four weeks after fracture there was no correlation between the A/C ratio and the time since fracture. Four patients had scans even earlier, between one and seven days after fracture, though these results were not included in the main analysis. Two of these patients had
commonly chosen, of the overall leg, the likely time of flow in bones which will be greatest, hypothesis related to be a blood ration in the normal patient individual provide can help bone, normal Our to cold indicative of bone scan concluded that the similar a very low ratio whereas those scanned after six days had results similar to those of the other patients. We concluded that the timing of the first scan after fracture is not particularly crucial to the ratio. Inspection of the bone scan images showed no appearance which was indicative of union or non-union, and in particular, no evident cold spots were seen in the scans of patients who proceeded to non-union.

**DISCUSSION**

Our results suggest that a quantitative MDP bone scan can help to predict non-union, but that the contralateral normal bone, as used by previous workers, does not provide a useful control site. Enough separation between the groups to enable a prediction to be made in an individual patient can only be achieved by using a normal region in the same limb as control.

One hypothesis is that, since disruption of the bone blood flow may be related to non-union, the early uptake result, which is predominantly related to blood flow, will be a better predictor than the late uptake, which is related to both blood flow and bone formation. This hypothesis can be extended, since it also explains why the greatest separation between the increased bone blood flow in bones which will unite, and its reduction in bones which will fail to unite, will be obtained at around the time of the maximum increase of blood flow. This period is likely to be nearer two weeks after injury than six or 12 weeks, and this explains our findings.

After a fracture blood flow increases to the whole of the bone, not just to the fracture site. This is why the use of a control region in the fractured limb rather than, as commonly chosen, a region in the contralateral normal leg, can compensate for variations between patients of the overall increase in blood flow. Thus the A/C ratio will reflect variations in uptake at the fracture site alone. This may explain the smaller spread of the A/C ratios in the group obtaining union (coefficient of variation 23%) as compared with the A/B ratios of 42%.

Our results are in broad agreement with those of other workers. The mean values of the A/B + C/D and C/D ratios agree well with those reported by Aucinholos and Watt (1982), and the inverse relationship they found between the late uptake C/D ratio at six weeks and the time to union is also confirmed. Auchinholos and Watt (1982) state that this finding is "difficult to rationalise and no data are available to explain this observation". A possible explanation is that the fracture produces a general increase in blood flow and activity at C is increased relative to D. As the fracture heals, blood flow decreases and uptake at C decreases; failure to unite may result in the continued higher activity at C, resulting in the increased C/D ratio for those with non-union. This theory is substantiated by our results; the C/D ratio is similar in the union group at two weeks to that in the non-union group at six weeks.

Jacobs et al. (1981) also used a control region in the fractured limb and investigated early uptake, but our uptake curves over the first 15 minutes (Fig. 4) are significantly different from the example given in their paper. We do not agree that the uptake between 7 min and 15 min is a "linear phase" which is "more an index of the affinity of the bone for phosphate and thus correlated with bone formation". Our images of the late uptake from one day to three weeks after fracture confirm the findings of Gregg et al. (1983) that "neither the presence of a cold spot nor any other scintigraphic feature could be correlated with the progress or time to fracture union". It must be noted, however, that we were using a high sensitivity collimator with consequent low resolution.
The early uptake \( \text{A} \div \text{C} \) ratio at the first visit demonstrates a separation between patients proceeding to union and those who develop non-union, and there are acceptable levels of sensitivity and specificity at a cut-off level of 1.3, even when different operators are analysing the results. However, the same data has been used both to develop and to assess the accuracy of the various cut-off values, so the values for sensitivity and specificity may be over-optimistic. There is therefore a need to test the technique in a further prospective study.

There are important practical implications for the timing of measurements. The fact that the best results were achieved as early as two weeks after fracture would allow changes to be made in the management of the patient far earlier than by other methods and these changes could reduce the mean time to union. The routine application of the technique is relatively easy because the "early uptake" can be investigated in only 10 minutes of the patient's time. The proportion of our patients recorded as suffering non-union was higher, at 20\%, than the 13.5\% quoted in a larger study by Ramadier et al. (1981). The reason is probably that we defined non-union as failure at six months; this is an important interval for clinical decision on further management. Aucinclos and Watt (1982) used the terms delayed union or unsatisfactory union for these patients and also had a lower incidence (11.5\%) than in our study. This difference may be due to the higher proportion of severely injured limbs in our study: we had 16\% and 66\% respectively in Groups III and II (Ellis 1958) compared with 9\% and 33\% in Aucinclos and Watts' series.

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REFERENCES


Premenopausal Bone Loss in the Lumbar Spine and Neck of Femur: A Study of 225 Caucasian Women

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Introduction

Age is well recognized as an important determinant of bone density, but the natural history of age-related bone loss remains controversial. In women, cortical bone density shows little or no diminution until the menopause which is followed by a period of rapid bone loss (Riggs et al. 1981). Advances in technology have led to the introduction of dual photon absorptiometry and quantitative CT scanning and these techniques make it possible to measure bone density at the spine and femur, the clinically important sites where osteoporotic fractures tend to occur. These areas have a significant trabecular bone component and evidence is accumulating that cortical and trabecular bone exhibit different patterns of loss.

Three patterns of age-related changes in trabecular bone have been described. Riggs et al. (1981) demonstrated loss commencing in "young adulthood" and continuing in a linear manner throughout life. A second model suggests that trabecular bone behaves in a similar way to cortical bone; it is maintained until the menopause when loss commences (Aloia et al. 1985; Sambrook et al. 1987). Other groups have shown that trabecular bone mineral reaches a peak in the mid-30s, and this is followed by a progressive

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Abstract

Two hundred and twenty-five premenopausal women were studied to evaluate age-related changes in trabecular bone mass. Measurements were made at the lumbar spine and femoral neck by dual photon absorptiometry. It was found that spinal bone density increased significantly from the 20s to reach a peak in the mid-30s. Identical trends were observed in total bone mass and bone mass normalized by length. Bone loss then proceeded at a rate of 1% per year, and by the early 50s, 10% of peak spinal density was lost. There was no peak in femoral neck density; loss commenced in the late 20s and continued at a rate of 0.4% per year. The cumulative premenopausal loss from the femur at 9% was comparable to that in the spine. It is concluded that significant amounts of trabecular bone are lost from both the spine and femoral neck before the menopause. The implications of these findings for the prevention of osteoporosis are discussed.

Key Words: Bone density—Dual photon absorptiometry—Menopause.

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Table I. Characteristics of the study population

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>n</th>
<th>Height (m) mean ± SD</th>
<th>Weight (kg) mean ± SD</th>
<th>Pill use (year) mean</th>
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</thead>
<tbody>
<tr>
<td>18-22</td>
<td>55</td>
<td>1.65 ± 0.06</td>
<td>59.30 ± 8.07</td>
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<tr>
<td>21-25</td>
<td>69</td>
<td>1.65 ± 0.14</td>
<td>58.90 ± 7.11</td>
<td>2.4</td>
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<tr>
<td>24-28</td>
<td>41</td>
<td>1.65 ± 0.06</td>
<td>57.58 ± 12.0</td>
<td>2.9</td>
</tr>
<tr>
<td>27-31</td>
<td>33</td>
<td>1.63 ± 0.21</td>
<td>61.79 ± 10.0</td>
<td>2.9</td>
</tr>
<tr>
<td>30-34</td>
<td>27</td>
<td>1.66 ± 0.06</td>
<td>63.44 ± 9.59</td>
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<td>19</td>
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<tr>
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<td>28</td>
<td>1.62 ± 0.05</td>
<td>57.00 ± 17.0</td>
<td>2.0</td>
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<tr>
<td>39-43</td>
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<td>1.62 ± 0.05</td>
<td>62.20 ± 9.99</td>
<td>1.8</td>
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<tr>
<td>42-46</td>
<td>23</td>
<td>1.63 ± 0.05</td>
<td>64.83 ± 8.36</td>
<td>2.0</td>
</tr>
<tr>
<td>45-49</td>
<td>17</td>
<td>1.60 ± 0.04</td>
<td>65.90 ± 11.2</td>
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<tr>
<td>48-52</td>
<td>15</td>
<td>1.56 ± 0.02</td>
<td>75.00 ± 14.3</td>
<td>1.9</td>
</tr>
</tbody>
</table>
loss (Krolner and Pors-Nielsen 1982; Cann et al. 1985). The aim of the present study is to establish the pattern of age-related alterations in trabecular bone in normal Caucasian women. An understanding of these changes may suggest alternative approaches to the prevention of osteoporosis.

Patients and Methods

Patients

The study population consisted of 225 Caucasian women whose ages ranged from 18 to 52 years. Women were recruited from the hospital staff and from outpatient clinics; approximately 50% of volunteers in each age group were staff members. They were all premenopausal with regular menstrual cycles. Subjects had no relevant medical history, and in particular, there was no history of bone or joint disease. None of the women were taking medication known to influence bone density. Women with a daily calcium intake of less than 500 mg, as assessed by questionnaire, were excluded as were those with a history of anorexia nervosa. In all cases, body mass index was <25 (weight in kilograms/height in meters$^2$). The characteristics of the study population are shown in Table I. Height and weight were similar in all age groups with the exception of the oldest group which was significantly shorter and heavier than other age groups. Current and past pill users were included in this study; we have previously shown that combined oral contraception does not influence bone density (Rodin et al. 1987).

All subjects gave informed consent for the study and approval was granted by the Ethical Committee, Guy’s Hospital, London.

Methods

Individuals attended for densitometry on entry to the study and there was no grouping of measurements with time according to age. All scans were performed and analyzed by the same technician. Bone mineral was measured at the lumbar spine (L2–L4) and at the neck of femur by dual photon absorptiometry using a Novo 22a dual photon absorptiometer. Total bone mineral (BM), bone mineral per unit length (BM/L) and bone mineral per unit area or bone mineral density (BMD) were measured at both sites. The units used were respectively, grams hydroxyapatite (gHA), grams hydroxyapatite per unit distance between the top of L2 and the bottom of L4 (gHA/cm) and grams hydroxyapatite per unit area of L2 to L4 (gHA/cm$^2$) as measured by the edge detection algorithm. Reproducibility data was obtained by repeated measurements in 10 subjects at both sites at 12-week intervals, giving a precision of 2.2% for BM, 2.6% for BM/L and 2.0% for BMD in the lumbar spine, and 2.2% for BMD in the femoral neck.

In the analysis of the data subjects were grouped according to age. Overlapping 5 year groups were formed (18–22 years; 21–25 years; 24–28; 27–31; 30–34; 33–37; 36–40; 39–43; 42–46; 45–49 and 48–52). The mean BMD and standard error of mean were calculated for each group and plotted against the mean age for that group. Non-linear regression analysis was also performed on the whole data set. In order to confirm that the observed trends were inde-
correlation with a linear fit. There was a 5% increase in spinal bone density between the age groups 18–22 and 33–37 which was significant (p = 0.03). This was followed by a progressive decline at a rate of 1.0% per year. By the early 50s, 10% of peak spinal bone mass was lost and this represented a significant fall (p = 0.016) (Fig. 2).

A different pattern was exhibited by the femoral neck, the raw data of which showed a linear correlation with age (r = 0.31, p < 0.001). No improvement in fit was obtained using a second or higher order polynomial. Our study did not demonstrate a peak in femoral bone mass and loss of mineral commenced in the late 20s. The rate of loss was slower than that seen in the lumbar spine at 0.4% per year, however, by the early 50s, 9% of femoral bone mass was lost which represented a significant premenopausal fall (p = 0.012) (Fig. 3).

Discussion

In this study, we have demonstrated that significant premenopausal bone loss occurs from both the lumbar spine and neck of femur. Spinal bone density increases in the 20s to reach a peak in the mid-30s and then falls in an almost linear fashion. The femoral neck shows a different pattern of bone loss: peak density is presumably attained at an earlier age which cannot be defined from the present data. The rate of loss is slower than that seen in the spine, but as bone loss commences at an earlier age, by the late 40s the cumulative loss from peak density at both sites is comparable.

Previous studies offer conflicting evidence about the occurrence of premenopausal bone loss and the patterns of age-related changes in bone mineral. In a cross-sectional study of 105 normal women, Riggs et al. (1981), found that bone loss from the vertebrae begins in young adulthood and is linear. However, the bulk of evidence now conflicts with this early model. Krolner and Pors-Nielsen (1982) examined lumbar spine bone mineral content in 70 women in relation to age. By fitting the data to a gamma variate function they showed a maximum spinal density at 34 years, but this method of statistical handling has been criticized (Tothill et al. 1983). Their own results support the view that there is increased loss of spinal bone mineral after the menopause, but insufficient numbers prevent detailed analysis in premenopausal women. This limitation also applies to other studies. Nilas and Christiansen (1987) carried out a cross-sectional study of 178 healthy women and found no evidence that substantial premenopausal bone loss occurs from any site. Hansson and Roos (1986) in agreement with our findings, noted a continuous age-related decrease of spinal bone density after the age of 35. Mazess et al. (1987) with a heterogeneous study population of 892 women derived from seven centers found a 10% decrease in spinal density in the decade preceding the menopause. A similar pattern was shown for femoral bone density, Schaadt and Bohr (1988), in a recent study, found that the bone mineral content of the femoral neck decreased linearly from the 30s. Premenopausal bone loss is also suggested by histomorphometric studies (Meunier et al. 1973; Marcus et al. 1983) and by postmortem investigations (Weaver and Chalmers 1966; Arnold 1973). The differing descriptions of age-related changes in trabecular bone are probably explained by technological differences, differences in data handling and differences in the sample sizes between the groups. Sambrook et al. (1987) pointed out the pitfalls of using cross-sectional data to draw conclusions about longitudinal changes in a population, and they emphasized the importance of adequate sample size.

Our data and results from previous studies provide a compelling body of evidence that significant loss of trabec-

![Fig. 2. Spinal BMD plotted against age in 225 premenopausal women. Mean BMD was calculated for 5 year overlapping groups and plotted against the mean age for that group.](image)
The finding of a peak in spinal bone density is supported by similar trends in BM and BM/L, suggesting that this phenomenon is independent of area normalization. The significant increase in spinal bone density between the 20s and mid-30s may have important implications for the prevention of osteoporosis in later life. In addition, our study provides further evidence that trabecular bone at different anatomical sites exhibits different patterns of age-related density changes, but the reason for this variation is not fully understood.

The possibility that these patterns of changes in trabecular bone are biased by secular trends needs consideration. It seems unlikely, however, that the rise in spinal density seen between 18 years to the mid-30s could be due to such influences. The relatively short time scale argues against a secular effect. However, we cannot rule out the possibility that the differences between those at the age extremes of the study population are influenced by secular effects.

The mechanism of early bone loss remains unclear. The importance of oestrogen in maintaining skeletal bone mass is well established and there is evidence for accelerated bone loss after oophorectomy (Cann et al. 1980) or following a natural menopause (Gennant et al. 1984). Oestrogen replacement therapy is effective in preventing postmenopausal bone loss (Lindsay et al. 1976). However, the menopause is preceded by a transition phase which is characterized by rising gonadotrophin levels and falling total oestrogen output from the ovaries (Sherman et al. 1976). It therefore seems likely that the patterns of endocrine changes will vary between individuals, and we suggest that genetic and environmental factors interact to influence the timing and profile of the hormonal fluxes associated with the menopause. A proportion of women will therefore become relatively oestrogen deficient before the cessation of their menses, and it is well known that some develop vasomotor instability and genital atrophy during the climacteric. It has been suggested that this relative deficiency of oestrogen may contribute to the loss of trabecular bone in premenopausal women (Johnston et al. 1985). It is possible that this effect contributes to spinal bone loss which commences in the mid-30s, but it does not adequately explain the pattern of loss seen in the femoral neck.

Prevention of osteoporosis is an important goal, and while hormone replacement therapy in the early premenopausal years is of proven efficacy (Lindsay et al. 1976), our findings raise the possibility that prophylaxis in premenopausal women should be considered. Two potential approaches are suggested: (a) spinal bone density increases in the young woman until the mid-30s, and if this can be enhanced the peak spinal bone density can be maximized; (b) if bone loss in the climacteric can be prevented or reduced, women would again reach the menopause with a better reserve. Exercise and calcium supplementation may be suitable strategies, but before any measures can be recommended they must be fully evaluated and ultimately they must be shown to reduce fracture risk.

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References


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Bone ultrasonic attenuation in women: reproducibility, normal variation and comparison with photon absorptiometry

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Abstract. The reproducibility of two methods of measuring broadband ultrasonic attenuation (BUA) in the calcaneus have been studied. An improvement in reproducibility in vivo from 3.6% to 2.8% between old and new techniques has been observed. Measurements of the calcaneus using BUA were correlated with measurements of bone mineral density measured by dual energy x-ray absorptiometry in the lumbar spine, femur and total body and bone mineral content in the distal and proximal forearm measured by single photon absorptiometry. For the older BUA technique the correlation coefficients ranged between \( r = 0.27 \) and \( r = 0.34 \). For the newer BUA technique the correlation coefficients ranged between \( r = 0.49 \) and \( r = 0.62 \) and were all significant (\( P < 0.001 \)).

1. Introduction

Osteoporosis affects an increasing fraction of the elderly population and can result in fractures of the wrist, spine and hip. Bone loss is more rapid from these sites of high trabecular content (Genant and Cann 1981). Non-invasive bone measurement for the management and diagnosis of osteoporosis is currently carried out using single photon absorptiometry (SPA) of the forearm (Cameron and Sorenson 1963), dual photon absorptiometry (DPA) of the spine (Roos and Skoldborn 1974) and more recently by dual energy x-ray absorptiometry (DEXA) of the proximal femur, lumbar spine and total body (Mazess et al 1989). Whilst all these methods are reasonably precise and accurate they do expose the subject to ionising radiation and they only measure the area density (g cm\(^{-2}\)) of the bone.

Langton et al (1984) suggested that measurement of broadband ultrasonic attenuation (BUA, dB MHz\(^{-1}\)) of the calcaneus can give a direct interpretation of the structure of the bone as well as bone density. The calcaneum is chosen as it is a site containing a high percentage of trabecular bone and may have a composition similar to other bones, such as the vertebrae and femur, which are more commonly measured for BMD. Poll et al (1986) later suggested that a reproducibility figure for this method of 3.9% could be obtained by measuring both heels and taking the average BUA value. Since then, other workers have obtained improved reproducibility values \textit{in vitro} for the method of 2.2% (McKelvie et al 1989) and 2.4% (McCloskey et al 1990a). The majority of the BUA measurements reported by other workers were made using the original commercial machine which measured BUA, the Walker Sonix UBA 1001, which has since been superseded by the Walker Sonix UBA 575.
In this study we examine the reproducibility of both the old and new techniques for measuring BUA in the calcaneus and correlate them with SPA of the forearm and DEXA of the lumbar spine, femur and total body. The normal variation of the newer BUA technique and comparison with DEXA of the spine and femur is also assessed in a group of women in whom we measured BUA of the calcaneus and BMD of the femur and spine on the same day.

2. Methods

2.1. Ultrasound equipment

We measured BUA in the right heel using two machines, the Walker Sonix UBA 1001 and the Walker Sonix UBA 575. With the UBA 1001, the patient sits with his/her foot in a room temperature water bath between two fixed 25 mm diameter transducers, both resonant at 1 MHz, one acting as a transmitter, the other as a receiver. The aim was to position the foot accurately so that the calcaneus lay between the two transducers. A series of short bursts of varying ultrasound frequency from 200 - 600 kHz in 16 kHz steps is transmitted through the water without the foot in the bath and the transmitted amplitude at each frequency recorded. A similar series of measurements is made through the water with the heel inserted in the bath. As the difference between the ultrasound signal amplitude for water and for water plus heel at each frequency sample point is due to the attenuation caused by the heel, an attenuation spectrum can be constructed. The slope of the best straight line fitted to this spectrum (dB MHz⁻¹) is the reported measurement of BUA. The measurement is repeated until three consecutive values are obtained that are within 0.5 dB MHz⁻¹ of each other. This usually takes about 3 min, the delay probably being caused by air trapped in the pores which is released in this period.

The fundamental basis of the UBA 575 is the same as the UBA 1001 but several changes in technique have been made. The transducers have been changed to 19 mm diameter with 500 kHz resonant frequency and signal amplitudes are recorded at 12 equally spaced intervals between 200 and 600 kHz. The heel is still placed in a water bath but now the transducers are made to carry out a rectilinear scan of nine sample points on a 3 × 3 grid in a region of 22 mm × 22 mm located around the region of minimum density in the calcaneus. This region was located by the manufacturers from a study of x-ray films of the calcaneus. The calculation of BUA at each of these sample points is done as previously but the nine values obtained are subjected to a weighted averaging algorithm to produce a single value for BUA. The total measurement time is about 5 min.

2.2. Wetting agents

One problem with BUA measurement in water is that air nucleates on the skin and surface hair forming bubbles at the interface between skin and water. These bubbles cause anomalous attenuation and can be a source of unreliability in the results. To overcome this a wetting agent is added to the water and this releases the absorbed air within the water and on the skin. The detergent recommended by the manufacturer is not widely available and we had to select another capable of giving reliable results. The detergents (and concentrations) that we tested were Ark (4%), Ecover (2%,4%), Quadralene (6%,8%), Stardrops (1%,2%) and Triton X100 (2%). The last of these is a commercial wetting agent and the remainder household detergents. Ark and Ecover were abandoned as they produced a precipitate in the water bath which was opaque to ultrasound. Triton X100 was not used after an initial trial as it tended to gel at room temperature. Using each of the other solutions we made five measurements of BUA using the UBA 575 on each of three phantoms of
different BUA (described in section 2.3). In each case a pooled coefficient of variation (cv) was calculated and used as an indicator of reproducibility.

Prior to the reproducibility studies we found two other sources of variability in BUA measurement which could be easily removed. The first of these was due to talcum powder on the feet of patients. This causes a precipitate in the water bath and can be removed by prebathing the foot before immersion in the scanner. The second source of variability was due to body oils or creams applied to the heel which can be removed with an alcohol swab prior to scanning.

2.3. Reproducibility

The in vitro reproducibility studies were performed using phantoms, representing high, medium and low values of BUA which are made by Walker Sonix and are constructed from room temperature vulcanised silicone in which glass beads are suspended. The phantoms were stabilised in the scanner so that there was a minimal change in position and orientation between scans. On the UBA 1001 37 measurements were made on two phantoms (high and medium BUA) over 13 months. On the UBA 575 64 scans were made on all three phantoms over a period of three months.

The in vivo reproducibility studies were performed using cooperative volunteers who were measured repeatedly with a reasonable time interval between measurements. On the UBA 1001 three volunteers were studied on 13 occasions with an interval of about a month between measurements. On the UBA 575, five volunteers were studied on five occasions with an interval of one day between measurements. In addition, the short term reproducibility in vivo was measured from two scans made within five minutes of each other on 30 female volunteers, the foot being removed from the bath and dried between scans. The effects of the various detergent solutions as described in section 2.2. was determined prior to the evaluation of reproducibility of the UBA 575.

2.4. Bone mineral measurement using absorptiometry

We measured BMC in the forearm at two sites using a Nuclear Data 1100A scanner. This SPA technique uses an $^{125}$I source of 5.5 - 7.4 GBq activity. The patients' arm is held in a water bath and the scanner finds the site at which the radius and ulna are separated by an 8 mm gap. From this point five scans are carried out at 4 mm intervals proximally and four scans at 2 mm intervals distally. The proximal and distal groups of scans are each averaged and the measurement at each of these two sites is given as bone mineral content normalised for both fat cover and bone width and expressed in arbitrary units which can be converted to g cm$^{-2}$. Nilas et al (1985) report a reproducibility of 1 - 1.5% (coefficient of variation) for the two forearm scan sites. The skin entrance dose associated with this technique is between 30 and 50 μSv.

To measure BMD in the spine, femur and total body we used a Lunar Radiation DPX scanner. The DEXA technique uses a filtered x-ray tube to obtain the dual energy beam required. The scanning and analysis procedures on this scanner are controlled by the manufacturers software version 3.2. The measurements we use are the bone mineral densities (BMD g cm$^{-2}$) of the lumbar spine (L2-L4), femoral neck, trochanter, Ward's triangle, and total body. The reproducibility obtained by us for the spine, femur and total body in vivo are 1.2%, 1.7% and 0.7% respectively. The skin entrance dose per scan associated with these techniques is 14 μSv each for spine and femur and 1 μSv for a total body scan.
2.5. Statistical methods

For reproducibility measurements the coefficient of variation has been calculated. With sets of paired data this is obtained from an analysis of the squares of the differences. For multiple measurements on subjects the overall cv is obtained from a pooled variance. This analysis and the calculation of the correlation coefficients was obtained using Statgraphics v3.0 as were the tests for normality and the pooled Students t-test.

3. Subjects

On the UBA 1001 we made bua measurements on 264 women aged 19 to 81 years (group 1), comprising 131 asymptomatic normals, 83 patients with endocrine disorders, 12 with asthma and 38 volunteers who were unclassified. Of these women, 261 had spine and femur bmd and 60 had total body bmd measurements. Spa of forearm was also carried out on 249 of these women.

We made bua measurements using the UBA 575 on 191 women aged 22 to 79 years (group 2), comprising 112 asymptomatic normals, 60 patients with endocrine disorders, 6 with osteoarthritis and 13 volunteers who were unclassified. Of these 190 had spine and femur bmd and 114 had total body bmd measurements. Forearm bmc measurements were made on 103 of these women. On the occasions when measurements were made on the same patient using different techniques they were carried out at the same visit.

The above measurements were made for use in a study of the correlation between the different techniques of bone measurement and so no exclusions were considered merited. To study the normal variation of each of the methodologies when applied to the same group, a subset of 70 women aged 50 to 59 years (group 3) was selected from group 2. These women were taking part in an hormone replacement therapy study and had presented for prophylactic screening with no symptoms of bone disease and comprise the whole of this age group. On groups 1 (UBA 1001) and 2 (UBA 575) we performed correlation analysis between bua and each of the bmd and bmc measurement techniques described above. For group 3 we calculated mean and standard deviation (sd) for the measurements obtained using each technique and for each technique we also calculated the coefficient of variation within the group.

4. Results

In the assessment of the wetting agents the pooled values of cv over the three phantoms measured on the UBA 575 for Quadralene (6%,8%) were 0.52% and 1.99% respectively and for Stardrops (1%,2%) were 2.00% and 0.39% respectively. On the basis of these results we determined to use Stardrops (2%) in our future patient work.

The in vitro reproducibility measured in phantoms over a period of 13 months on the UBA 1001 was 3.1%, on the UBA 575 the equivalent reproducibility value measured over a period of 3 months was 2.2%. We measured reproducibility in vivo over a 13 month period on the UBA 1001 to be 9.6%. On the UBA 575 the reproducibility measured over a 5 day period was 2.8%. In addition to the above reproducibility measurements we measured the reproducibility in vivo on the UBA 575 in the short term to be 3.5%.

Summary statistics for subject groups 1 and 2 for each of the measurement techniques used are given in table 1. The smaller samples used for total body bmd in groups 1 and 2 and the forearm spa samples drawn from group 2 were checked to ensure that the mean value of bua in these samples was representative of the bua of the group from which they
Table 1. Summary statistics for group 1 (UBA 1001) and group 2 (UBA 575) for heel BUA (dB MHz⁻¹), BMD (g cm⁻²) at various sites, BMC (g cm⁻²) in the forearm, height (cm) and weight (kg) comprising sample size (n), mean, standard deviation (sd), minimum and maximum values.

<table>
<thead>
<tr>
<th></th>
<th>Group 1</th>
<th>Group 2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Mean</td>
</tr>
<tr>
<td>Heel BUA</td>
<td>264</td>
<td>96.7</td>
</tr>
<tr>
<td>Spine BMD</td>
<td>261</td>
<td>1.054</td>
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<tr>
<td>Femoral neck BMD</td>
<td>261</td>
<td>0.879</td>
</tr>
<tr>
<td>Trochanter BMD</td>
<td>261</td>
<td>0.719</td>
</tr>
<tr>
<td>Ward's triangle BMD</td>
<td>261</td>
<td>0.743</td>
</tr>
<tr>
<td>Total body BMD</td>
<td>60</td>
<td>1.088</td>
</tr>
<tr>
<td>Proximal forearm BMC</td>
<td>249</td>
<td>1.315</td>
</tr>
<tr>
<td>Distal forearm BMC</td>
<td>249</td>
<td>1.369</td>
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<tr>
<td>Height</td>
<td>264</td>
<td>160.5</td>
</tr>
<tr>
<td>Weight</td>
<td>264</td>
<td>63.5</td>
</tr>
</tbody>
</table>

were drawn. Pooled t-tests revealed no significant differences in BUA values between any of these samples and the group mean values. The correlation coefficients of the different techniques for groups 1 and 2 and their significances are given in table 2. None of the variates used in the correlation analysis departed significantly from being normally distributed. Scatter plots of the correlated variates did not depart from linearity confirming the validity of the use of correlation analysis. It can be seen from table 2 that although the correlations between BUA and other measurements in group 1 are significant, they are all relatively weak. Group 2 shows an improvement in the correlation coefficients all of which are highly significant.

The spread of the normal data in group 3 using the different techniques is given in table 3. Examination of the CVs of the various measurements reveals little difference between the spread of the normal range using the different techniques. The CV of heel BUA is, however, the largest.

Table 2. Correlation coefficients (r), for group 1 (UBA 1001) and group 2 (UBA 575), for various bone mineral measurement techniques compared to BUA in the right calcaneus, with significance values (P) and sample sizes (n). BMD is measured in g cm⁻².

<table>
<thead>
<tr>
<th>Measurement technique</th>
<th>Group 1</th>
<th></th>
<th></th>
<th></th>
<th>Group 2</th>
<th></th>
<th></th>
<th></th>
<th></th>
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</thead>
<tbody>
<tr>
<td></td>
<td>r</td>
<td>P</td>
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<td></td>
<td>r</td>
<td>P</td>
<td>n</td>
<td></td>
<td></td>
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<tr>
<td>Spine BMD</td>
<td>0.34</td>
<td>&lt;0.001</td>
<td>261</td>
<td></td>
<td>0.61</td>
<td>&lt;0.001</td>
<td>190</td>
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<tr>
<td>Femoral neck BMD</td>
<td>0.27</td>
<td>&lt;0.001</td>
<td>261</td>
<td></td>
<td>0.60</td>
<td>&lt;0.001</td>
<td>190</td>
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<td></td>
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<tr>
<td>Trochanter BMD</td>
<td>0.33</td>
<td>&lt;0.001</td>
<td>261</td>
<td></td>
<td>0.58</td>
<td>&lt;0.001</td>
<td>189</td>
<td></td>
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<tr>
<td>Ward's triangle BMD</td>
<td>0.29</td>
<td>&lt;0.001</td>
<td>261</td>
<td></td>
<td>0.62</td>
<td>&lt;0.001</td>
<td>190</td>
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<tr>
<td>Total body BMD</td>
<td>0.31</td>
<td>&lt;0.001</td>
<td>60</td>
<td></td>
<td>0.54</td>
<td>&lt;0.001</td>
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<td>Proximal forearm BMC</td>
<td>0.30</td>
<td>&lt;0.001</td>
<td>249</td>
<td></td>
<td>0.49</td>
<td>&lt;0.001</td>
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<tr>
<td>Distal forearm BMC</td>
<td>0.29</td>
<td>&lt;0.001</td>
<td>249</td>
<td></td>
<td>0.55</td>
<td>&lt;0.001</td>
<td>103</td>
<td></td>
<td></td>
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</tbody>
</table>
5. Discussion

The reproducibility figures obtained suggested an improvement with the UBA 575 over the earlier UBA 1001. This was particularly demonstrated by the *in vivo* data. However, the *in vivo* reproducibility data from the two machines are not entirely comparable due to the differences in time intervals used. The possible presence of a gradual variation in BUA using the UBA 1001 over the 13 month period was investigated by a further analysis of the data. The reproducibility was assessed from the differences between successive measurements; this gave a value of 9.4% suggesting that the relatively poor reproducibility was not due to a gradual long term variation. The improvement in precision in the UBA 575 equipment is probably due to several factors including a change in resonant frequency of the transducers, a change to rectilinear scanning, the addition of a foot restraint, the selection of wetting agent and the thorough cleaning of the foot before placement in the water bath.

Many groups have tried to correlate BUA of the calcaneus with different bone measurement techniques at other sites. With SPA of the distal forearm correlation coefficients of $r = 0.8$ (Poll et al 1986), $r = 0.77$ (McCloskey et al 1990b) and $r = 0.64$ (Rossman et al 1989) have been demonstrated. A study has also been reported which showed no correlation between BUA of the calcaneus and SPA of the forearm (Resch et al 1990). For DPA of the femoral neck Rossman et al (1989) report a correlation coefficient of 0.41 whilst for the lumbar spine they give $r = 0.66$. For the spine McCloskey et al (1990b) give $r = 0.72$. For all these previously reported studies reasonable numbers ($n$ between 44 and 72) were used for the correlations.

Quantitative computed tomography (QCT), a technique developed in different forms by Cann and Genant (1980) and Ruegsegger et al (1976) has revealed a quite strong correlation ($r = 0.85$) between BUA of the calcaneus and forearm cortical bone mineral content, but a far weaker relationship ($r = 0.66$) with trabecular bone mineral in the same limb (Hosie et al 1987). However when BUA and QCT are carried out at the same site correlations of $r = 0.92$ in excised os calcis (McKelvie et al 1989) and $r = 0.80$ in calcaneal core samples (McCloskey et al 1990a) have been observed. Both of these groups report correlations of BUA with physical density of approximately 0.8. However McKelvie et al (1989) rejected a large proportion of their calcaneal core samples (15%) before correlating physical density with BUA. In the studies using QCT the group sizes for correlation analysis ranged between 24 and 28.

The 25 mm core samples measured by McCloskey et al (1990a) were the same diameter as the ultrasound transducers which may result in significant diffraction artefacts. The core samples used above were all taken from similar sites where the structure and orientation of trabeculae could be expected to be similar. Evans and Tavakoli (1990) used samples taken from different regions of bovine femora excised at random angles, thus a difference in

<table>
<thead>
<tr>
<th>Measurement technique</th>
<th>Mean</th>
<th>SD</th>
<th>CV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heel BUA</td>
<td>65.51</td>
<td>12.45</td>
<td>18.98</td>
</tr>
<tr>
<td>Lumbar spine BMD</td>
<td>1.06</td>
<td>0.15</td>
<td>14.58</td>
</tr>
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<td>Femoral neck BMD</td>
<td>0.87</td>
<td>0.12</td>
<td>13.96</td>
</tr>
<tr>
<td>Trochanter BMD</td>
<td>0.72</td>
<td>0.11</td>
<td>15.53</td>
</tr>
<tr>
<td>Ward's triangle BMD</td>
<td>0.75</td>
<td>0.13</td>
<td>17.39</td>
</tr>
</tbody>
</table>
structure and orientation between samples could be expected. This is probably why they observed such a weak correlation \( r = 0.33, n = 27 \) between BUA and physical density. Unless substantial scattering of the ultrasound beam is occurring the trabecular orientation should make little difference to the measurement of attenuation. Evans and Tavakoli (1990) also considered samples with a range of density (1.15 - 1.4 g cm\(^{-3}\)) which had a higher upper limit than other workers. At higher densities errors in the measurement increase, which would diminish the correlation further.

Comparison of our correlation coefficients, of BUA with other techniques, for group 1 (UBA 1001) and group 2 (UBA 575) shows a significant improvement in correlation with the newer of the two methods of measuring BUA. This improvement is probably attributable to the improvement in reproducibility leading to less variability in BUA measurement with the newer method. When we compare the correlations of group 2 BUA with other techniques to those correlations reported by other workers, with the exception of the femoral neck (Rossman et al 1989), all our correlations are weaker, although we have substantially larger groups.

The weak correlations that have been obtained in our study could be due to either or both of the following factors: (i) BUA values may reflect, in some way, both density and structure. We may, therefore, not expect the correlation with only bone density to be particularly good. (ii) The correlation between bone density in the calcaneus and other sites in the body may not be particularly strong.

Other methods of assessing the utility of BUA measurements have been suggested in terms of diagnostic predictive power. Several workers observe that those patients at high risk of fracture can be sorted from the normal population (Langton et al 1984, Miller and Porter 1987, Johnson and Porter 1987). Murray et al (1987) showed that patients with osteoporosis could be identified, Fordham et al (1987) that patients with rheumatoid arthritis differed from controls and Jones et al (1987) that active and sedentary groups could, be separated. The problem still remains that, although groups can be separated, given a single measurement of BUA it is often difficult to assign an individual to a group with any degree of confidence.

Acknowledgements

We wish to thank the MRC for supporting this work, and Mr P Townsend for supplying the UBA 575.

References

NOTE

The use of a radiation sensitive CCD camera system to measure bone mineral content in the neonatal forearm: a feasibility study

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1. Introduction

Single photon absorptiometry (SPA) was first suggested by Cameron and Sorenson (1963) for the measurement of bone mineral content (BMC) in the distal forearm in vivo and quickly gained currency as a quantitative method of bone mineral measurement giving a precision of approximately 2% with a radiation dose of approximately 0.15 mGy similar to the dose for a forearm x-ray film exposure. Adaptations of the method to measure the neonatal forearm met with some success (Steichen et al 1980, Greer et al 1982) but until 1986 (James et al 1986b) an instrument designed specifically to measure the forearm bones of premature neonates had not been constructed. With a dose to the subject of only 0.03 mGy and accuracy of 5% this machine enabled growth and mineralization data to be obtained for fetal mineralization (Ryan et al 1988), osteopenia of prematurity (James et al 1986a) and the repletion of mineral in osteopenia of prematurity (Congdon et al 1990).

Unfortunately a subset of preterm babies were too sick to be amenable to this measurement technique as they could not be moved from incubator to measuring instrument. In many situations it is important to measure BMC in these babies at this very early stage and to continue to monitor it in order to evaluate the efficacy of both treatment and feeding regimes. To study such babies in the incubator it would be necessary to construct a hand-held instrument which was portable, compact, light and accurate. Such a device should only expose the baby to a low radiation dose. It was felt that an isotope transmission device with a radiation sensitive charge coupled device (CCD) camera offered the likeliest solution to the problem.

2. Methods

The development of CCD video cameras which are sensitive to a range of x and γ ray energies (Arndt and In'T Veld 1988) suggested a method of measurement of BMC in the forearm using video techniques. The schematic diagram of an idealized version of
such a system is shown in figure 1. The ideal system would consist of an isotope source of monochromatic $\gamma$-ray photons of sufficient intensity located close to the arm to make the system as compact as possible. Correction for geometric distortion could be carried out relatively easily. The forearm would be enclosed in a parallel-sided soft tissue equivalent bolus material to provide a fixed thickness path in which the only changes in transmitted intensity would be due to changes in equivalent bone thickness. The radiation sensitive CCD video camera would convert each $\gamma$-ray photon arriving at the detector into a visible light photon which is converted to a specific charge on the CCD. The video circuitry produces a TV frame every 40 ms in which the grey level of each pixel is representative of the charge per element on the CCD. This information is digitized using a frame grabber under computer control. With the frame grabber operating in real time, the area to be measured could be viewed on the monitor before the data was captured. Each captured frame would then be added into a frame store until there were sufficient counts per pixel to reduce the measurement variance to an acceptable level.

The feasibility of using a CCD camera was investigated using a model of the neonatal forearm and the following apparatus. The CCD video camera was a Phototonic Science X-ray Imager on loan for a day from the manufacturers. This camera has a detector quantum efficiency (DQE) of $10-12\%$ for energies of 5 to 200 keV covering the photopeak of $^{125}\text{I}$ at 27.5 keV. The effective detector pixels are 80 $\mu$m square working into the CCD by light pipe coupling to give $753 \times 581$ pixels. Digitization and sequential frame addition was carried out using a PC Vision Plus 8 bit frame grabber having two $512 \times 512$ pixel frame stores, controlled by an IBM PC using ITEX PC plus software. The limit of an 8 bit frame store meant that in practice in order to avoid overflow we could only collect 32 frames which is equivalent to an integration period of 1.28 s. The source used was an $^{125}\text{I}$ 1 mm point emitter of activity 1.13 GBq.

The forearm was modelled by a water-filled syringe of 21.7 mm diameter located 74.3 mm from the source and 14.0 mm from the detector surface with two aluminium tubes of diameters 3 mm and 2 mm having a wall thickness of 0.15 mm suspended centrally in the water. These sizes were typical of neonatal forearm bone sizes measured on x-rays. The forearm model was measured in air without a soft tissue equivalent bolus to provide a flat-sided soft tissue path. At this stage we make no attempt to
correct for geometric distortion and differences in intensity caused by the cone beam geometry in the system.

3. Theory

With reference to the transmission image, the bone mass per unit area \( (M_B) \) can be calculated pixel by pixel by the application of the following equation:

\[
M_B = \frac{\rho_B}{(\mu_B \rho_B - \mu_S \rho_S)} \ln(I_0/I) \quad M_S = k_S \ln(I_0/I) \tag{1}
\]

where \( k_S = \rho_B/(\mu_B \rho_B - \mu_S \rho_S) \). \( \mu_B \) and \( \mu_S \) are the mass absorption coefficients for bone mineral \((2.40 \text{ cm}^2 \text{ g}^{-1})\) and water or soft tissue \((0.35 \text{ cm}^2 \text{ g}^{-1})\) respectively. \( \rho_B \) and \( \rho_S \) are the values of density for bone mineral \((1.85 \text{ g cm}^{-3})\) and water or soft tissue \((1.00 \text{ g cm}^{-3})\). \( I \) is the intensity through 'bone' and \( I_0 \) is the background intensity. In the neonatal forearm phantom, bone has been replaced by aluminium having a density of \(2.6989 \text{ g cm}^{-3}\) and a mass absorption coefficient of \(4.00 \text{ cm}^2 \text{ g}^{-1}\), thus we can calculate \( M_A \) the mass of aluminium per unit area.

We wished to study the relationship between the precision of the technique and the exposure time so that the minimum investigation time is achieved. This is particularly important in the measurement of preterm neonates. The variance of the mass per unit area is given by

\[
\sigma^2(M_B) = \sigma^2(I) + \frac{\partial M_B}{\partial I_0}^2 \sigma^2(I_0) = \frac{\partial M_B}{\partial I_0}^2 I + \frac{\partial M_B}{\partial I_0}^2 I_0
\]

and so

\[
\sigma(M_B) = k_B \left( \frac{I + I_0}{I_0} \right)^{1/2} \tag{2}
\]

4. Results

The transmission image in figure 2 results from the capture of 32 frames of the neonatal phantom with the experimental system described above. Cursor regions A and C are \(60 \times 200\) pixels \((4.8 \times 16 \text{ mm})\), cursor B is \(80 \times 200\) pixels \((6.4 \times 16 \text{ mm})\) just covering the width of the larger aluminium tube in the phantom. These represent reasonable useful sizes for cursors \textit{in vivo}. The equivalent mass of aluminium was calculated taking the mean of regions A and C for the background intensity \( I_0 \). Substituting into equation (1) produces a value of \(0.0909 \text{ g cm}^{-2}\) for \( M_A \). This compares well with the known value of \( M_A \) of \(0.0864 \pm 0.0093 \text{ g cm}^{-2}\), a difference of 5.3%.

The variation of the intensity across the phantom bone was studied using a region 250 pixels long but only 2 pixels wide. The plot of intensity across the aluminium phantom is shown in figure 3. This may be a more appropriate method of analysis \textit{in vivo} as interpolation using the through tissue counts may give a more accurate estimation of the background count intensity.
Substitution into equation (2) gives a value for \( \sigma(M_A) \) of 0.0009 g cm\(^{-2}\). This is equivalent to a coefficient of variation (cv) of 0.99%. The computed skin entrance dose for the experimental system with a 1.13 GBq source and an integration period of 1 second is 0.046 \( \mu \)Gy, computed from measurements made on a 3.5 GBq source in a similar configuration, several orders of magnitude less than the other techniques mentioned in the introduction. The variation of the theoretical precision versus time and dose for a 3.7 GBq activity source is shown in figure 4. This shows that at 3.7 GBq
it could be possible to reduce the integration period to as little as 3 seconds and still have a theoretical measurement error of less than 0.5%.

5. Discussion

While the physical size of the neonatal forearm means that an increase in cursor size is unlikely to be achieved, reduction in the measurement error could be brought about in a number of ways. The source we used in this evaluation was depleted and use of a fresh source of 7.4 GBq would increase the photon flux by at least a factor of 6. The position of the source in the experimental system is by no means optimal as the distance away from the object tends to make the system too cumbersome for use in an incubator. Moving the source closer to the object would also have the added advantage of increasing the flux and reducing the exposure time required. The use of a CCD camera working at video rate into an 8 bit frame grabber and frame store produces a grey level limit of 256 levels per pixel effectively limiting exposure time to 1.28 s with the photon flux available to us before overflow started to occur. The exposure time will be reduced with a higher flux source but will still only give a grey level resolution of 256 levels. An increase in grey level resolution could only be achieved by the use of a higher resolution frame grabber of 12 or 16 bits. Further, the system that we used in the feasibility experiment is by no means ideal as the addition of each captured frame into the frame store took about 30 seconds. This limit could be overcome in two ways. Firstly, by staying with the same technology and using higher resolution frame grabbers and stores with virtually instantaneous summation facility, which although increasing the cost considerably would allow the retention of real time viewing of the measurement site. The second possibility is to switch from real time video mode to still camera mode where the x-ray sensitive CCD is operated in integrating mode, exposed for a fixed period and then the charge is read off and converted to a digital signal directly representing photon events per pixel. This system loses the ability to view the measurement site at the time of measurement and suffers the further disadvan-
Figure 5. Image of the sum of 32 frames of raw data obtained from measurement of the distal and middle inter-phalangeal joint of the middle finger in an adult.

The advantage of requiring to be cooled to liquid nitrogen temperatures to suppress electronic noise in the device. Both these techniques would allow us to increase exposure time to 10 s which we feel is the limit imposed by subject movement.

It may be possible to avoid using a bolus for the baby's arm altogether if we can adequately fit a line representing the background intensity. If this cannot be done satisfactorily then the tissue equivalent bolus need not be parallel sided. It may be possible to produce a short source to detector distance and a non-linear bolus geometry which could compensate for any variation in path length.

Although the technique was originally envisaged for the measurement of neonatal BMC there is no reason why it should not be extended to the adult appendicular skeleton. Figure 5 shows an image of the distal and middle inter-phalangeal joint of the middle finger of an adult captured using the evaluation system. The manufacturers have indicated that they could reduce the size of the video camera to 164 mm, with the drive electronics housed up to 1 metre away, making it feasible to place such a system in an incubator. In either neonates or adults this technique could provide a quick and accurate method of measuring BMC in the appendicular skeleton which may be adapted to suit many measurement environments and different measurement sites.

Acknowledgments

The authors would like to thank Patricia Tomkins and Photonic Science for the loan of the x-ray CCD camera and their help with its operation. We would also like to thank
Sharan Packer who computed the radiation dose associated with the new technique. This work was supported by the Medical Research Council.

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PRIMARY GENERALIZED OSTEOARTHRITIS AND BONE MASS

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*Rheumatology and Rehabilitation Research Unit and †Centre for Bone and Body Composition Research, Department of Clinical Medicine, University of Leeds, ‡Department of Applied Biology, University of Hull

SUMMARY

The association of OA with increased bone mass is controversial. This study measured BMD at the hip and spine and total body bone mineral (TBBM) by dual energy X-ray absorptiometry, and BMD at the distal forearm by single photon absorptiometry in 20 post-menopausal women with primary generalized OA. The data were compared with those from 89 normal controls. Osteoarthritic women had significantly increased BMD at the spine (P<0.001), distal forearm (P<0.05) and increased TBBM (P<0.01), but no difference was seen at the femoral neck. These differences were not explained by obesity. The influence of mobility is discussed.

KEY WORDS: Dual energy X-ray absorptiometry, Single photon absorptiometry, Spine, Distal forearm, Femoral neck.

Although the negative association between OA of the hip and femoral neck fracture is well recognized [1-4], it is not clear whether patients with OA have greater bone mass than normal people. A number of studies have been performed comparing bone mass in patients with OA of the hip with controls, but results are conflicting, perhaps due to the selection of subjects, the different techniques and sites used for bone measurement, and the heterogeneous nature of OA of the hip [3-6]. Fewer studies have been performed in patients with primary generalized osteoarthritis (PGOA) but results are also conflicting. Studies using metacarpal morphometry and neutron activation analysis to measure total body calcium [7], quantitative computed tomography of the forearm [8] and single photon absorptiometry of the forearm [9] have shown no difference between osteoarthritic subjects and controls. However a relationship between trabecular bone volume in iliac crest biopsies and radiological grade of OA at the hand was found in one post-mortem study [10].

The aim of our study was to assess BMD at potential osteoporotic fracture sites (lumbar spine, distal forearm and hip) and total body bone mineral (TBBM) in women with PGOA and compare this with data from normal women.

PATIENTS AND METHODS

Twenty postmenopausal women aged 52-79 yr (median 68 yr), attending a rheumatology clinic for treatment of PGOA were recruited. All women had clinical and/or radiological OA affecting three or more groups of joints as described by Kellgren and Moore [11]. Details of joint involvement are given in Table I. Women with hysterectomy were excluded as age of menopause was unknown although one woman with simultaneous hysterectomy and bilateral oophorectomy was included. Women with a history of disease or drug therapy potentially affecting bone mass were also excluded e.g. diabetes, thyroid disease, oral steroid therapy, anticonvulsant treatment. Although women with fractures were not specifically excluded, none were known to have vertebral or hip fractures and only one had a previous Colles' fracture. A second patient sustained a Colles' fracture after the study was completed. Six patients used one or more sticks for walking and two of these six also used wheelchairs for outings. Data from 89 normal postmenopausal women aged 50-79 yr (median 62 yr) recruited locally from hospital and university staff and randomly from a list of postmenopausal women attending an inner city general practice were used as controls [12]. Women with hysterectomy, thyroid disease and other endocrine disorders, liver, kidney or chronic gastrointestinal disease, known osteoporotic fracture, impaired mobility or who had received steroid, hormone replacement therapy or other treatment potentially affecting bone, were excluded from the control population. All study and control subjects were of Caucasian origin.

BMD (bone mineral content/bone width, arbitrary units) was measured at the distal forearm by single photon absorptiometry. Bone mineral content (BMC, g) and BMD (g/cm²) were measured at the lumbar spine (L2-L4) and femoral neck by dual energy X-ray absorptiometry using a Lunar DPX Bone Densitometer.

<table>
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<th>Joint Involvement in Osteoarthritic Patients</th>
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<tr>
<td>Joints</td>
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</tr>
<tr>
<td>Proximal interphalangeal</td>
</tr>
<tr>
<td>First carpo-metacarpal</td>
</tr>
<tr>
<td>First metacarpal-phalangeal</td>
</tr>
<tr>
<td>Knee</td>
</tr>
<tr>
<td>Cervical spine</td>
</tr>
<tr>
<td>Lumbar spine</td>
</tr>
<tr>
<td>Hip</td>
</tr>
<tr>
<td>Shoulder</td>
</tr>
<tr>
<td>Ankle</td>
</tr>
</tbody>
</table>

Submitted 8 September 1992; revised version accepted 18 March 1993.

Correspondence to L. D. Hordon, Rheumatology and Rehabilitation Research Unit, 36 Clarendon Road, Leeds LS2 9NZ.
showing this was not simply a local effect due to artefact. Mean lumbar BMD was 7.3 g higher in osteoarthritic patients than controls, whilst mean TBBM was 1.52 g higher in osteoarthritic patients than controls (Table III). The possible contribution of degenerative disease in the lumbar spine to the difference in TBBM is therefore small. In addition there was a strong correlation between lumbar BMD and distal forearm, femoral neck and TBBM measurements.

We conclude that women with PGOA have greater bone mass than controls at the distal forearm and spine and greater TBBM, although bone mass at the femoral neck is not increased. These differences are not due to obesity. It is possible that impaired mobility in the osteoarthritic women could explain the lack of difference in bone mass at the femoral neck. The cause of the increase in bone mass in women with PGOA is unknown, but the results of this study would support the hypothesis that oestrogen excess may be involved in the pathogenesis of OA [20].

**Acknowledgements**

We would like to thank Mr P Constable, Department of Mathematics, University of Leeds, for statistical advice and Mrs D. K. Smith for typing the manuscript.

**References**


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**Table IV: Relationships Between Age, HAQ score and Bone Mineral Measurements in Osteoarthritic Women Given as the Coefficient of Correlation**

<table>
<thead>
<tr>
<th>Age</th>
<th>Spine BMD</th>
<th>Femoral BMD</th>
<th>Forearm BMD</th>
<th>BMC/BW</th>
<th>TBBM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spine</td>
<td>-0.33</td>
<td>N.S.</td>
<td>-0.5</td>
<td>0.71</td>
<td>***</td>
</tr>
<tr>
<td>BMD</td>
<td>0.75</td>
<td>0.52</td>
<td>0.66</td>
<td>0.61</td>
<td>**</td>
</tr>
<tr>
<td>Femoral</td>
<td>-0.08</td>
<td>0.60</td>
<td>-0.60</td>
<td>-0.70</td>
<td>***</td>
</tr>
<tr>
<td>BMC/BW</td>
<td>**</td>
<td>*</td>
<td>**</td>
<td>**</td>
<td>***</td>
</tr>
<tr>
<td>TBBM</td>
<td>**</td>
<td>***</td>
<td>**</td>
<td>**</td>
<td>***</td>
</tr>
</tbody>
</table>

BMC, Bone mineral content; BW, bonewidht; TBBM, total body bone mineral content; HAQ, Stanford Health Assessment Questionnaire.

* P < 0.05; ** P < 0.01; *** P < 0.001.
**TABLE II**

<table>
<thead>
<tr>
<th>Age at Menopause and BMI in Osteoarthritic Women and Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td>Age at menopause (yr)</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
</tr>
</tbody>
</table>

Results are expressed as means ± s.d.

(Lunar Radiation Corporation, Madison, WI, USA). TBBM (g) was also measured by this technique. Osteoarthritic patients completed a Stanford Health Assessment Questionnaire (HAQ) [13] in order to assess their functional ability.

**Statistical analysis**

Within group data were correlated using Spearman rank correlation (rho). Data on age of menopause and BMI from osteoarthritic women and controls were compared by separate variance t-tests. Differences in spine, femoral neck, distal forearm and TBBM measurements between the two groups were compared using an unbalanced analysis of variance with age as covariant and significant difference of the F statistic for the elevation was tested after the regression lines were shown to have similar slopes.

All data were analysed on Unistat IV statistical software.

**RESULTS**

There was no significant difference in age at menopause or BMI between osteoarthritic women and controls (Table II). Women with OA scored a mean of 0.81 ± 0.64 (s.d.) on the HAQ.

BMC and BMD were significantly higher at the spine (P<0.001) and BMD at the distal forearm was also increased (P<0.05) in osteoarthritic women compared with controls. TBBM was higher in the osteoarthritic group (P<0.01). BMC and BMD at the femoral neck were not significantly increased in osteoarthritic women (Table III).

Relationships between bone mineral measurements, age and HAQ score are shown in Table IV. There was a significant negative correlation between HAQ score and spinal, femoral neck, forearm and TBBM measurements. There was a significant positive correlation of HAQ score with age.

**DISCUSSION**

Women with PGOA have more bone at the distal forearm and spine and greater TBBM than controls, although no significant increase in bone mass was seen at the femoral neck in osteoarthritic women. These differences were not due to obesity as BMI was similar in control and study groups.

Some of the osteoarthritic women had impaired mobility and it is possible this may have influenced bone mass at the femoral neck, reducing differences from controls. Certainly the HAQ score correlated negatively with bone mineral measurements, but as age was also related to HAQ score this should not be over interpreted. It is of interest that muscle strength has been shown to predict bone mass in the femur and forearm, but not the spine in healthy postmenopausal women [14]. A regional increase in BMD adjacent to an osteoarthritic joint has been proposed as a mechanism for increased bone mass in OA [15]. If this theory is correct, the low prevalence of OA of the hip in our study population (Table I) could provide an alternative explanation for the lack of difference at the femoral neck from controls. However, this theory would not explain the increased BMD at the distal forearm in osteoarthritic women.

Neither osteoarthritic women nor controls had a routine lateral lumbar spine X-ray. The presence of osteophytes and degenerative disease at the facet joints can produce artefacts affecting lumbar BMD measurements [16], although the extent to which these measurements are affected is controversial [15]. Spinal osteophytes are present in the majority of 'normal' postmenopausal women [17,18], but account for only 4% of the variance in lumbar BMD [17]. However, only half this effect is artefactual and lumbar osteophyte score is related to femoral BMD [17]. Exclusion of subjects from a study population because of degenerative changes may thus potentially bias the study group. In men, osteophytes make a greater contribution to lumbar BMD [19], perhaps because of lower age-related bone loss, or sex differences in osteophyte formation [18].

Although we cannot exclude an influence of degenerative disease on lumbar BMD, not only was bone mass increased at the spine, but also at the distal forearm, and TBBM was also elevated in the osteoarthritic group.

**TABLE III**

<table>
<thead>
<tr>
<th>Comparison between Bone Mineral Measurements at the Spine, Femoral Neck, Distal Forearm and Total Body Bone Mineral in Osteoarthritic Patients and Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td>Spine</td>
</tr>
<tr>
<td>BMC (g)</td>
</tr>
<tr>
<td>BMD (g/cm²)</td>
</tr>
<tr>
<td>Femoral neck</td>
</tr>
<tr>
<td>BMC (g)</td>
</tr>
<tr>
<td>BMD (g/cm²)</td>
</tr>
<tr>
<td>Distal forearm</td>
</tr>
<tr>
<td>BMC/BW (arbitrary units)</td>
</tr>
<tr>
<td>(n = 18)</td>
</tr>
<tr>
<td>Total body bone mineral</td>
</tr>
<tr>
<td>(g)</td>
</tr>
</tbody>
</table>

BMC: Bone mineral content; BW, bone width. Results expressed as means ± s.d.

* P < 0.05; ** P < 0.01; *** P < 0.001.
Variation in lumbar spine and femoral neck bone mineral measured by dual energy X-ray absorption: a study of 329 normal women

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Centre for Bone and Body Composition Research, Institute of Physical Sciences, Department of Clinical Medicine, University of Leeds, Wellcome Wing, The General Infirmary, Leeds LS1 3EX and Department of Applied Biology, University of Hull, Hull, UK

Abstract
Reference ranges used in dual energy X-ray absorptiometry (DXA) have previously used piecewise linear fits to the whole data set for spine or femur bone mineral density (BMD) as a function of age. In a study of 329 Caucasian normal women we present a refinement to the normal range by fitting straight lines between quinquennial mean values of BMD for each site measured (lumbar spine, femoral neck and Ward’s triangle). From the age of 40 years onwards the premenopausal women demonstrated minimal loss of BMD whereas postmenopausal women showed a rapid loss amounting to 27% in the lumbar spine, 27% in the femoral neck and 38% in the Ward’s triangle region in the age range under examination. Comparison of quinquennial means for pre and postmenopausal women in age bands 45–49 years and 50–54 years shows that at these ages postmenopausal BMD is significantly lower than premenopausal BMD ($P < 0.05$). This finding suggests that separate normal ranges should be used for pre and postmenopausal women. As reduction in the production of oestrogen is a major factor in postmenopausal bone loss and oestrogen replacement is related to years since menopause (YSM), a more logical way of displaying postmenopausal normal BMD ranges would be in terms of YSM rather than chronological age. Such data are given in this paper.

The measurement of bone mineral has long been recognized as important for the study of osteoporosis. Originally instruments were developed to measure bone mineral in the forearm (Cameron & Sorenson, 1963) but in the early 1980s techniques were developed to measure the sites of most clinical interest directly, i.e. the spine and femoral neck. The techniques for measuring the spine and femoral neck use the same basic principle; the differential attenuation that occurs between bone and soft tissue when gamma or X-ray photons of two different energies pass through the body. A measurement of the relative attenuation, coupled with the mass attenuation coefficient gives a measurement of the bone mineral. The region is studied by moving the source and detector in a rectilinear motion across the patient and producing an image in which each pixel is representative of the mineral content. The integration of the data over the appropriate region of interest (ROI) gives the bone mineral content (BMC) in grams.

Though the BMC for a given anatomical ROI may be an important parameter in its own right it is clear that its value will be dependent upon the size of the individual; large subjects will tend to have a higher BMC than small subjects. To compensate for this the BMC is divided by the area of the bone obtained from the image to produce bone mineral density (BMD) in units of g cm$^{-2}$. It should be noted that the BMC is not the measurement of volume density but is the area normalized BMC. BMD appears to have become the measurement of choice but care must be taken with its use.

Early instruments used two gamma ray energies obtained from isotope sources, initially a combination of $^{241}$Am and $^{125}$Cs and subsequently $^{153}$Gd which became the basis of a number of commercial systems. The isotope techniques for measuring BMD were referred to as dual photon absorptiometry (DPA) (Reed, 1966). This gave a reasonable precision at about 8% (Tothill et al., 1983) but often the quality of the images was poor and in many patients it was difficult to identify the areas to measure. A significant improvement in signal-to-noise ratio in the image was obtained with the use of X-ray tubes as the photon source (Sartoris et al., 1985). The more recent development is referred to as quantitative digital radiography (QDR) or DXA (Magee et al., 1988). The two abbreviations refer to the same basic principle and we shall adopt the latter irrespective of the commercial system used. The improved photon output of DXA over DPA results in benefits in terms of improved resolution, improved precision or shorter scan times.

Variation in lumbar spine and femoral neck bone mineral measured by DXA

The initial spine BMD results showed a significant variation in age relationship with different systems (Smith et al., 1983). At the time it was thought that there could be some national differences in BMD although technical explanations were also felt to be a possibility. It must be remembered that the BMC is the ratio of the BMD to the area of the bone. It is quite conceivable that different techniques can calculate the BMC reliably but that different algorithms used for edge detection, to calculate the area of the bone, are likely to differ significantly from system to system. As a consequence it may be the denominator in the ratio, i.e. the area, that is the principal cause of variation between different systems.

The variation in BMC of the spine and femoral neck using either DPA or DXA in Caucasian females in several countries is illustrated in Figures 1a and 1b. The data was extracted from the references listed in Table I which also gives the age range of each of the studies and the number of subjects involved. There are discrepancies between the data from each country, which may reflect differences in either the control population or the technique used. However, much depends upon the size of the data set and consequently the statistical model that can justifiably be used. This study was initiated to enable us to obtain a comprehensive normal range of BMC values for Caucasian females in the United Kingdom (UK). It was also recognized that an individual may suffer a crush fracture which is caused by a vertebral body having a lower BMC and may result in a lower than normal vertebral area. In such a situation the BMC, which is the ratio of the two, can appear in the normal range. As a consequence the normal data were analysed to obtain the vertebral area in the normal population which could be used to give an indication as to whether BMD or BMC values should be used in the assessment of that individual.

Methods
Reference subjects
The reference sample for the spine BMD measurements consisted of 329 women aged between 20 and 81 years, of whom 174 were premenopausal and 155 were postmenopausal. The mean age at menopause was 49.1 (4.5) years, the figure in brackets is the standard deviation. For the femur measurements the sample consisted of 327 women from the above set aged between 21 and 81 years of whom 173 were premenopausal and 154 were postmenopausal with a mean age at menopause of 49.1 (4.4) years. The women were recruited mainly from hospital and university staff by advertising locally and from the list of a local general practitioner (GP). The GP contacted every third woman on his list of Caucasian females in the age range 55–80 years and invited those with no known disorders to participate in the study of whom approximately 30% attended. On the basis of responses to a questionnaire women with endocrine disorders, adrenal disorders, malignancy, chronic gastrointestinal or liver disease, Paget's disease, diabetes, rheumatoid arthritis or renal stones were excluded. Women on steroid therapy, hormone replacement therapy, anticonvulsant drugs, sodium fluoride, heparin, thyroxin or with a history of hip, spine or wrist fracture or extended immobilization were all excluded. The exclusion also extended to women who had undergone hysterectomy or oophorectomy and subjects with histories of drug or alcohol abuse. Spine X-rays were not taken in this study for ethical reasons so that some women who may possibly have asymptomatic vertebral wedge or crush fractures were not identified and thus not excluded. It is recognized that the reference data presented in this study is not a statistically representative sample of the local population. In particular there is no representation of non-Caucasian races and there will

Table I. Major studies using DPA or DXA listed chronologically including first author, number of subjects, measurement site and age range

<table>
<thead>
<tr>
<th>First author</th>
<th>Year</th>
<th>Number</th>
<th>Site</th>
<th>Age range (years)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Riggs</td>
<td>1981</td>
<td>105</td>
<td>Spine</td>
<td>20–89</td>
</tr>
<tr>
<td>Kroner</td>
<td>1982</td>
<td>70</td>
<td>Spine</td>
<td>19–88</td>
</tr>
<tr>
<td>Tothill</td>
<td>1983</td>
<td>24</td>
<td>Spine</td>
<td>21–61</td>
</tr>
<tr>
<td>Guesens</td>
<td>1986</td>
<td>156</td>
<td>Spine</td>
<td>20–85</td>
</tr>
<tr>
<td>Hansson</td>
<td>1986</td>
<td>214</td>
<td>Spine</td>
<td>35–90</td>
</tr>
<tr>
<td>Mazess</td>
<td>1987</td>
<td>892</td>
<td>Spine/Femur</td>
<td>20–79</td>
</tr>
<tr>
<td>Nias</td>
<td>1987</td>
<td>178</td>
<td>Spine</td>
<td>29–78</td>
</tr>
<tr>
<td>Elders</td>
<td>1988</td>
<td>286</td>
<td>Spine</td>
<td>46–55</td>
</tr>
<tr>
<td>Pocock</td>
<td>1988</td>
<td>179</td>
<td>Spine/Femur</td>
<td>20–83</td>
</tr>
<tr>
<td>Ribot</td>
<td>1988</td>
<td>510</td>
<td>Spine</td>
<td>25–70</td>
</tr>
<tr>
<td>Schaadt</td>
<td>1988</td>
<td>113</td>
<td>Spine/Femur</td>
<td>20–89</td>
</tr>
<tr>
<td>Stevenson</td>
<td>1989</td>
<td>284</td>
<td>Spine/Femur</td>
<td>21–68</td>
</tr>
<tr>
<td>Alford</td>
<td>1990</td>
<td>376/257</td>
<td>Spine/Femur</td>
<td>20–80</td>
</tr>
<tr>
<td>Elliott</td>
<td>1990</td>
<td>462</td>
<td>Spine/Femur</td>
<td>20–84</td>
</tr>
<tr>
<td>*Hall</td>
<td>1990</td>
<td>165</td>
<td>Spine</td>
<td>4–80</td>
</tr>
<tr>
<td>Rodin</td>
<td>1990</td>
<td>225</td>
<td>Spine/Femur</td>
<td>18–52</td>
</tr>
</tbody>
</table>

All DPA scans except *which is DEXA.

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The patients who obtained normal data that obtained analysis Data triangle regions alcohol, 75% water) tank filled with times per aluminium spine, femoral neck and Ward's triangle. The coefficient of variation with 1.69% and had lumbar subjects (seven male, lesion is determined to instrumentation errors, while in vitro precision is determined by instrumentation error alone. 15 subjects (seven male, eight female; mean age 43 years) had lumbar spine and femur scans on two occasions with an interval of 28–90 days (mean 55 days) between scans. The coefficient of variation (CV) for the lumbar spine, femoral neck and Ward’s triangle was 1.18%, 1.69% and 3.13%, respectively. The manufacturer’s aluminium spine and femur phantoms were scanned five times per day for 7 days in a low attenuation plastic tank filled with a nominal fat solution (25% isopropyl alcohol, 75% water) to a depth of 15 cm to measure the in vitro precision. The CV for spine, femoral neck and Ward’s triangle regions of the phantoms was 0.45%, 1.00% and 0.84%, respectively.

Scanning techniques and scan analysis

Scanning was carried out using a Lunar DPX machine on sites in the femoral neck and lumbar spine (Mazess et al, 1989). Analysis of the scans was carried out using the manufacturer’s software version 3.2. The effective dose equivalent for each examination is 1 μSv and was measured by the local radiation protection service.

Precision

Precision is defined as the ability of a method to obtain the same value with repeated measurements. In vitro precision represents variations in patient positioning and instrumentation errors, while in vitro precision is determined by instrumentation error alone. 15 subjects (seven male, eight female; mean age 43 years) had lumbar spine and femur scans on two occasions with an interval of 28–90 days (mean 55 days) between scans. The coefficient of variation (CV) for the lumbar spine, femoral neck and Ward’s triangle was 1.18%, 1.69% and 3.13%, respectively. The manufacturer’s aluminium spine and femur phantoms were scanned five times per day for 7 days in a low attenuation plastic tank filled with a nominal fat solution (25% isopropyl alcohol, 75% water) to a depth of 15 cm to measure the in vitro precision. The CV for spine, femoral neck and Ward’s triangle regions of the phantoms was 0.45%, 1.00% and 0.84%, respectively.

Data analysis

A number of different analyses were undertaken to obtain normal data that could be used for comparison with patients who may have a degree of crushing of one or more of the vertebrae. If this is not known then the total BMD of L2–L4 may give an erroneous result. The total area of L2 + L3 + L4 was noted. In this paper we present the normal results as means and standard deviations. For comparison with the normal ranges of the commercial company (Lunar Corporation) straight lines were fitted to the normal data. The assumption has been made by the company that there is constant BMD early in life and then two different linear rates of loss later in life. Least squares techniques are used to identify the slopes of these losses and the break points, i.e. the age at which the rate of loss changes.

Results

The results for the normal vertebral BMD and associated areas are given in Table II, grouped into 5 year age bands for all women and separately for pre and postmenopausal women. The data for BMD of the femoral neck and Ward’s triangle regions are given in Table III, grouped in the same way. The BMD data for spine, femoral neck and Ward’s triangle are given, grouped into 5 year bands of years since menopause (YSM), in Table IV. The results of the variation of BMD with age are shown graphically in Figure 2a for the spine, Figure 3a for the femoral neck and Figure 4a for Ward’s triangle. In each case the figure contains the raw data, the quinquennial mean values and straight lines joining those mean values. The results of the variation of BMD with YSM are shown in Figure 2b for the spine, Figure 3b for the femoral neck and Figure 4b for Ward’s triangle. In each case the figure in the lumbar spine, femoral neck and Ward’s triangle. Pooled Student’s t-tests show that in the 40–44 year band there is no significant difference.

Figure 1. BMD (g cm⁻²) in the lumbar spine (a) and the femoral neck (b) of Caucasian women plotted against age (years) gathered by different measurement centres throughout the world. (Leeds 1991 refers to this paper.)

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Lumbar Spine BMD

Figure 2. BMD (g cm\(^{-2}\)) in the lumbar spine plotted against (a) age (years), unbroken line joining means for all women, upper broken line joining means for premenopausal women, lower broken line joining means for postmenopausal women; and (b) years since menopause (YSM) for postmenopausal women.

between the pre and postmenopausal groups, in the 45-49.9 year quinquennia the difference between the two groups at all three sites is highly significant (P < 0.001) and in the 50-54.9 year band is still significant (P < 0.05). The importance of separating reference data into premenopausal and postmenopausal groups is highlighted by this finding.

The results of comparing the linear fits to the data

Table II. Mean and standard deviation values for measurements made on lumbar vertebrae L2 to L4 of area, BMC and BMD listed for all women and for pre and postmenopausal women by age band including the number in each group

<table>
<thead>
<tr>
<th>Ageband (years)</th>
<th>Number</th>
<th>Spine area (cm(^2))</th>
<th>Spine BMD (g cm(^{-2}))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All</td>
<td>Pre</td>
<td>Post</td>
</tr>
<tr>
<td>20-24</td>
<td>15</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td>25-29</td>
<td>32</td>
<td>32</td>
<td></td>
</tr>
<tr>
<td>30-34</td>
<td>38</td>
<td>38</td>
<td></td>
</tr>
<tr>
<td>35-39</td>
<td>24</td>
<td>24</td>
<td></td>
</tr>
<tr>
<td>40-44</td>
<td>22</td>
<td>18</td>
<td>4</td>
</tr>
<tr>
<td>45-49</td>
<td>35</td>
<td>28</td>
<td>7</td>
</tr>
<tr>
<td>50-54</td>
<td>45</td>
<td>19</td>
<td>26</td>
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<td>55-59</td>
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<td>60-64</td>
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<td>65-69</td>
<td>9</td>
<td>9</td>
<td></td>
</tr>
<tr>
<td>70-74</td>
<td>19</td>
<td>19</td>
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</tr>
<tr>
<td>75-81</td>
<td>14</td>
<td>14</td>
<td></td>
</tr>
</tbody>
</table>

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with the commercial company's normal range showed negligible differences between the two data sets (Mean Z-score of differences 0.062). However, it should be noted that the results from Leeds were included in the data base of the commercial company and account for some 36% of the data used to construct their spine normal range and about 43% of the data used in the construction of their femur normal range.

Table III. Mean and standard deviation values for BMD measurements made in the femoral neck and Ward's triangle regions listed for all women and for pre and postmenopausal women by ageband including the number in each group.

<table>
<thead>
<tr>
<th>Ageband (years)</th>
<th>Number</th>
<th>Femoral neck BMD (g cm⁻²)</th>
<th>Ward's triangle BMD (g cm⁻²)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All</td>
<td>Pre</td>
<td>Post</td>
</tr>
<tr>
<td>20–24</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.114</td>
<td>0.114</td>
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<tr>
<td>25–29</td>
<td>31</td>
<td>31</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.134</td>
<td>0.134</td>
</tr>
<tr>
<td>30–34</td>
<td>38</td>
<td>38</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>0.119</td>
<td>0.119</td>
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<tr>
<td>35–39</td>
<td>24</td>
<td>24</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>0.108</td>
<td>0.108</td>
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<td>40–44</td>
<td>22</td>
<td>18</td>
<td>4</td>
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<td></td>
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<td>0.091</td>
<td>0.094</td>
</tr>
<tr>
<td>45–49</td>
<td>35</td>
<td>28</td>
<td>7</td>
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<td>19</td>
<td>26</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.152</td>
<td>0.151</td>
</tr>
<tr>
<td>55–59</td>
<td>38</td>
<td>38</td>
<td></td>
</tr>
<tr>
<td></td>
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<td>60–64</td>
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<td></td>
<td></td>
<td>0.097</td>
<td>0.097</td>
</tr>
<tr>
<td>65–69</td>
<td>9</td>
<td>9</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.110</td>
<td>0.110</td>
</tr>
<tr>
<td>70–74</td>
<td>18</td>
<td>18</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.124</td>
<td>0.124</td>
</tr>
<tr>
<td>75–81</td>
<td>15</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.153</td>
<td>0.153</td>
</tr>
</tbody>
</table>

Figure 3. BMD (g cm⁻²) in the femoral neck plotted against (a) age (years), unbroken line joining means for all women, upper broken line joining means for premenopausal women, lower broken line joining means for postmenopausal women, and (b) years since menopause (YSM) for postmenopausal women.
Discussion

Both the spine and femur data display an effect caused by the menopause with the end point BMD value of the premenopausal data being higher than the starting point data of the postmenopausal data in each case (Figures 2a, 3a, 4a). A small study by Totten et al. (1983) demonstrated an increased loss of bone from the spine after the menopause but this study was not supported by a larger study from Hansson & Roos (1986). By comparing the BMD values of postmenopausal women who are over 45 years of age with the BMD values of postmenopausal women within 5 years of menopause we can establish whether or not this difference is significant. The outcomes of these tests are given in Table V. It can be seen that in every site measured the difference between the two groups is highly significant, ranging from 7.9% to 11.3%.

In this set of data (Table II) peak BMD in the spine occurs in the 35–39.9 year band with very little loss taking place in the premenopausal women from the time at which peak BMD is attained onwards. In the UK this is earlier than the ages reported by Hull et al. (1990) but consistent with the findings of Rodin et al. (1990). In other countries we find agreement only with Elliott et al. (1990), Krohn & Nielsen (1982), Schaadt & Bohr (1988), Guesens et al. (1986) and Ribot et al. (1988) all find that the peak occurs earlier. Postmenopausally the pattern is more confusing (Figure 2b) with a 10 year

Table IV. Mean and standard deviation values for BMD measurements of the lumbar vertebral (L2–L4), femoral neck (FBMD) and Ward’s triangle (WBMD) listed for postmenopausal women by YSM including the number (n) in each group.

<table>
<thead>
<tr>
<th>YSM (years)</th>
<th>n</th>
<th>SBMD (g cm⁻²)</th>
<th>SBMD (SD)</th>
<th>FBMD (g cm⁻²)</th>
<th>FBMD (SD)</th>
<th>WBMD (g cm⁻²)</th>
<th>WBMD (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–4.9</td>
<td>42</td>
<td>1.091</td>
<td>0.155</td>
<td>0.886</td>
<td>0.137</td>
<td>0.771</td>
<td>0.138</td>
</tr>
<tr>
<td>5–9.9</td>
<td>41</td>
<td>0.987</td>
<td>0.157</td>
<td>0.845</td>
<td>0.110</td>
<td>0.716</td>
<td>0.125</td>
</tr>
<tr>
<td>10–14.9</td>
<td>18</td>
<td>0.965</td>
<td>0.218</td>
<td>0.827</td>
<td>0.127</td>
<td>0.673</td>
<td>0.133</td>
</tr>
<tr>
<td>15–19.9</td>
<td>20</td>
<td>1.019</td>
<td>0.128</td>
<td>0.815</td>
<td>0.081</td>
<td>0.665</td>
<td>0.099</td>
</tr>
<tr>
<td>20–24.9</td>
<td>17</td>
<td>1.042</td>
<td>0.117</td>
<td>0.823</td>
<td>0.092</td>
<td>0.658</td>
<td>0.112</td>
</tr>
<tr>
<td>25–29.9</td>
<td>11</td>
<td>0.941</td>
<td>0.193</td>
<td>0.759</td>
<td>0.208</td>
<td>0.602</td>
<td>0.220</td>
</tr>
<tr>
<td>&gt;30</td>
<td>6</td>
<td>0.816</td>
<td>0.212</td>
<td>0.756</td>
<td>0.134</td>
<td>0.623</td>
<td>0.184</td>
</tr>
</tbody>
</table>

Table V. Results of Student’s t-tests comparing spine BMD (SBMD), femoral neck BMD (FBMD) and Ward’s triangle BMD (WBMD) in two groups of women; premenopausal older than 45 years and postmenopausal within 5 years of menopause, for whom percentage difference is given, including mean, standard deviation (SD) and number of subjects (n) for each group.

<table>
<thead>
<tr>
<th>SBMD</th>
<th>Mean (g cm⁻²)</th>
<th>SD</th>
<th>n</th>
<th>Postmenopausal Age</th>
<th>Premenopausal Age</th>
<th>P</th>
<th>% Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1.218</td>
<td>0.138</td>
<td>47</td>
<td>&gt;45</td>
<td>&lt;5</td>
<td>0.00009</td>
<td>10.4</td>
</tr>
<tr>
<td>FBMD</td>
<td>Mean (g cm⁻²)</td>
<td>SD</td>
<td>n</td>
<td>Postmenopausal Age</td>
<td>Premenopausal Age</td>
<td>P</td>
<td>% Difference</td>
</tr>
<tr>
<td></td>
<td>0.962</td>
<td>0.121</td>
<td>47</td>
<td>&gt;45</td>
<td>&lt;5</td>
<td>0.007</td>
<td>7.9</td>
</tr>
<tr>
<td>WBMD</td>
<td>Mean (g cm⁻²)</td>
<td>SD</td>
<td>n</td>
<td>Postmenopausal Age</td>
<td>Premenopausal Age</td>
<td>P</td>
<td>% Difference</td>
</tr>
<tr>
<td></td>
<td>0.869</td>
<td>0.135</td>
<td>47</td>
<td>&gt;45</td>
<td>&lt;5</td>
<td>0.002</td>
<td>11.3</td>
</tr>
</tbody>
</table>
band of loss followed by a 10 year period of apparently little change, then loss taking place from this time onwards. When we look at this data set as a whole (Figure 2a) the pattern is simpler, showing a rise of BMD during the 20–40 year age range with peak density achieved in the 35–39.9 year band. This is followed by a steady loss amounting to about 27% in the following 42.5 years. This is consistent with the total bone loss of 25% reported by Hall et al (1990) in UK populations covering the same age range as our data. In other countries where Caucasian female populations covering the same age range have been studied, percentage bone losses have been closely comparable with ours with one striking exception. Rigg et al (1981) reported a loss of 47% in the BMD of lumbar vertebrae L1 to L4 but the age range in their study is from 20 to 90 years.

In the femoral neck (Table III) peak BMD is attained in the 25–29.9 year band with bone loss proceeding at a negligible rate before the menopause. Postmenopausally we observe a steady loss with years since menopause, the total loss in the period considered being some 27%. In the UK the total bone loss of 26% reported by Stevenson et al (1989) agrees with our data although they do report the loss phase starting at less than 30 years of age. Elsewhere in the world the data of Mazess et al (1987) (25%) compares closely with this data whereas Elliott et al (1990) report a lower age for both attainment of peak bone mass (20–25 years) and for onset of loss (25–30 years). Pocock et al (1987) also report loss starting earlier (mean age 25 years) than our estimate but proceeding at a comparable rate to achieve a slightly higher total loss (30%) by age 83 years.

In Ward’s triangle, peak BMD is attained in the 25–29.9 year age band with loss commencing in the 30–34.9 year age band. Once again the premenopausal bone loss from this time is negligible. Postmenopausally, however, the bone loss is steady and rapid by comparison with the femoral neck, with total loss in the period amounting to some 38%. This very high rate of loss would make Ward’s triangle an attractive site for measurement if the in vivo precision could be improved.

The data display two stages of bone loss consistent with the classification of Type I and Type II osteoporosis (Riggs & Melton, 1983). Type I is most active in the 15–20 years immediately after the menopause when the rate of loss can be as much as three times higher than normal. The loss is predominantly from trabecular bone and manifests itself in the form of vertebral and Colles’ fractures. Type I osteoporosis is apparently linked to oestrogen deficiency in some, as yet unclear, way. Type II osteoporosis is apparent in women over 75 years of age and presents in the form of vertebral wedge fractures and hip fractures. The mechanism of loss is related to senescence with loss taking place in both cortical and trabecular bone.

The value of considering the vertebral geometry can only be assessed following a study of patients with a variety of established symptoms of osteoporosis; this analysis is currently being pursued. Our preliminary findings suggest that when a patient is referred for a bone mineral measurement the vertebral geometry should be considered in addition to the BMD.

Normal ranges should be developed which split data for pre and postmenopausal women. As reduced oestrogen levels are associated with postmenopausal bone loss and oestrogen levels decline with years since menopause it may be more logical to express postmenopausal normal ranges in terms of YSM rather than chronological age.

Acknowledgements

This work was supported by the Medical Research Council. We are indebted to our friends and colleagues in the hospital and university who formed part of the normal control group and we are grateful to Dr Tom O’Shea, a general practitioner in Leeds, who kindly allowed us to approach individuals registered with him to take part in this study. We are also grateful to the General Infirmary and the Special Trustees of the General Infirmary at Leeds for additional financial support.

References


J G Truscott, R Oldroyd, M Simpson et al


Book reviews


Endosonography—the use of specially designed ultrasound probes for insertion into various body orifices—is a significant advance in the practice of ultrasonography. High frequency transducers can be placed close to the organ of interest in order to generate high resolution images which can dramatically increase the yield of clinically useful information. These probes are now offered by nearly all ultrasound equipment manufacturers and most ultrasound departments are being asked to provide an endosonographic service. There is clearly a need for books which can provide instruction on techniques and advice on image interpretation, and the appearance of these three volumes in the bookshops is timely.

The first book listed above has three authors—Bernaschek, Deutinger and Kratochwil—and is written as a conventional textbook providing a comprehensive coverage of all applications of endosonography in obstetrics and gynecology. It is well written and is systematic in its approach. It is well illustrated and all the scans are presented in a uniform format using a standardized presentation of scan planes, and there is a good selection of references to the literature.

The same publishers, Springer Verlag, have produced the second volume, edited by Feifel, Hildebrandt and Mortensen. This is a collection of six entirely separate review articles, each with its own author or team of authors. As well as covering the history and physics/instrumentation of endosonography, there are chapters on the upper GI tract, rectum, prostate and gynecology. The coverage is selective and both the style and presentation of images are non-uniform.

The third book is edited by Bruno Fornage who has enlisted the help of 17 co-authors. It has a broad scope, covering upper and lower GI tract, urological, gynecological and obstetric applications, but is rather patchy and variable in the depth of its coverage, e.g. transvaginal Doppler techniques receive only cursory mention. The orientation of the scans varies both within and between chapters and the quality of some of the illustrations is disappointing.

In conclusion, most ultrasound departments will be looking for a teaching and reference text on endosonography in obstetrics and gynecology, and the book by Bernaschek, Deutinger and Kratochwil is to be highly recommended. For departments and libraries looking for a book giving an overview of all applications of endosonography, Fornage is recommended, while this reviewer has difficulty in identifying a prospective market for Feifel et al.

H C IRVING
A phantom for quantitative ultrasound of trabecular bone

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Abstract. The propagation mechanisms of ultrasound in trabecular bone are poorly understood and have been the subject of extended debate; also, the reproducibility of ultrasonic measurements on bone in vivo using commercial ultrasound heel-scanning devices is such that the interpretation of the obtained data is difficult. In this paper we describe recent developments in the production of a bone-mimicking material which is well suited to the task of routine monitoring of commercial ultrasound bone scanners. The material, based on a standard epoxy resin is fabricated to a predetermined porosity value by the inclusion of a marrow-mimicking material thereby introducing a known and controlled mean pore size.

Measurements of the velocity and attenuation of the material have been performed over a range of porosity values from 10% to 80% in the frequency range 500–900 kHz; also, broadband ultrasonic attenuation (BUA) values have been obtained from commercial equipment. The material displays velocities in the range 1844–3118 m s\(^{-1}\) and attenuation ranging from 7.0 to 17.7 dB cm\(^{-1}\) at 500 kHz.

1. Introduction

The rapidly increasing use of ultrasound to detect and monitor bone pathology, in particular osteoporosis, has led to the development of several commercially available devices. Such devices typically measure the rate of change of attenuation of ultrasound with frequency (broadband ultrasound attenuation or BUA) between 0.2 and 0.6 MHz, and also the velocity of sound (Langton et al 1984, 1990).

Currently, at least two of the commercial manufacturers offer phantoms for use with their own scanners. However, at best these devices are suitable for monitoring temporal changes in scanner performance. There is no independently validated data on their acoustic parameters and in one case the machine is set in a different mode before the phantom can be used because the acoustic properties differ significantly from those of the os calcis. In neither case is there any attempt to simulate the architecture of the heel nor is it certain that the phantoms themselves have good long-term stability. There is a clear requirement for a phantom, constructed from a material which will mimic acoustically the properties of ultrasound in trabecular bone in this frequency range. Such a phantom would enable intercomparisons to be made between results from different commercial ultrasound bone systems and provide the necessary calibration for long term longitudinal patient studies. Furthermore, the propagation of ultrasound, particularly in trabecular bone, is still poorly understood and so there is a need for a material which could be used to simulate bone in a controlled, reproducible and realistic manner.

This paper describes a new material, made from a mixture of an epoxy resin and gelatine which mimics the speed and attenuation of sound through trabecular bone and which can be formed into a phantom of a specific size and shape.
2. Trabecular-bone-mimicking material

2.1. Requirements of the material

The bones of the human skeleton can be divided into two types: cortical bone and cancellous, or trabecular, bone. The latter has a porous structure made up of cortical trabecules, the pores being filled with bone marrow.

Cortical bone is relatively straightforward to mimic, as it has a homogenous, compact structure. Trabecular bone, having a composite structure is more problematic. The size, shape and concentration of pores varies between skeletal sites; the proportion by volume which is marrow is known as the porosity. Hence there is a requirement for the bone phantom material to be able to mimic porosity in a controllable way. We therefore aimed to have a two component material, one component being ultrasonically equivalent to cortical bone, and the other equivalent to bone marrow. The two components were to be geometrically arranged in a manner comparable to trabecular bone with variable, but controlled, porosity.

The specification for the two materials was that they should match the velocity and attenuation of cortical bone and marrow over as wide a range of conditions as possible. Unfortunately there is significant variation in the reported values for both parameters for cortical bone and almost no data for either quantity in bone marrow. Table 1 summarizes some of the published values, although in all but one case, these refer to animal samples.

<table>
<thead>
<tr>
<th>Material</th>
<th>Velocity (m s⁻¹)</th>
<th>Attenuation at 1 MHz (db cm⁻¹)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cortical bone</td>
<td>3461 ± 104</td>
<td>8.36 ± 1.37</td>
<td>Tavakoli (1991)</td>
</tr>
<tr>
<td>Cortical bone</td>
<td>3250 ± 196*</td>
<td>—</td>
<td>André et al (1980)</td>
</tr>
<tr>
<td>Cortical bone</td>
<td>—</td>
<td>6.9 ± 0.6</td>
<td>McKelvie and Palmer (1987)</td>
</tr>
<tr>
<td>Bone marrow</td>
<td>1435 ± 10</td>
<td>0.50 ± 0.05</td>
<td>Tavakoli (1991)</td>
</tr>
</tbody>
</table>

2.2. Materials to simulate cortical bone and marrow

An extensive search for synthetic materials possessing similar ultrasonic characteristics to those of cortical bone and marrow was made using the techniques described below. This concluded that the optimal values were not achievable. However, materials were found which gave a reasonable match.

The cortical bone mimic is a two-part epoxy compound normally used for the encapsulation of electrical components. It is available commercially as Araldite CW 1302 liquid epoxy resin and is used in conjunction with Araldite HY 1300 hardener in the proportions recommended by the manufacturer. Its ultrasonic velocity is within the published values for bone, although the attenuation coefficient is about 50% less (see table 2 below).

A number of materials were identified as potential marrow mimics but most were rejected since no practical means could be found of creating the required structure when dispersing in the epoxy compound. It was therefore decided to evaluate the use a gelatine/water mixture in the proportions of 11 g of dry gelatine to 150 ml of water. This
concentration was chosen as a compromise between the increased mechanical stiffness at higher concentrations and the lower sound velocities at lower concentrations. This concentration has useful mechanical properties as it can be formed into relatively stable pore-shaped granules, but has the disadvantage that its attenuation coefficient is an order of magnitude too low and its velocity some 6% too high. Nevertheless it was hoped that this would prove to be of little significance as the high acoustic losses in trabecular bone are more dependent on the scatter arising from the geometrical arrangement of its constituents than absorption by the constituents themselves; also, the discrepancy in velocity should be reduced in the composite material. The acoustic properties of the materials are displayed in table 2.

Table 2. The ultrasonic characteristics of the constituents of the bone phantom (Truscott et al 1993). The ranges quoted refer to standard deviation values.

<table>
<thead>
<tr>
<th>Material</th>
<th>Velocity (m s(^{-1}))</th>
<th>Attenuation at 1 MHz (dB cm(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bone mimic</td>
<td>3168 ± 8</td>
<td>3.7 ± 0.3</td>
</tr>
<tr>
<td>Marrow mimic</td>
<td>1519 ± 3</td>
<td>0.051 ± 0.004</td>
</tr>
</tbody>
</table>

2.3. Manufacture of the trabecular bone material

The two components were combined so as to resemble the structure of cancellous bone in the following manner. Firstly the gelatine was extruded twice through a 1.12 mm mesh to produce roughly cubic granules of side approximately 1 mm. This was then added to the liquid epoxy in volumes, calculated by means of weight and density, to give the required porosity and then mixed until a uniform distribution of gelatine particles had been achieved. The material was then de-gassed in a vacuum chamber at 3 Torr to remove any air pockets, then poured into moulds and left to harden.

This process, which is the subject of a patent application, produces a material consisting of gelatine pores enclosed in a matrix of epoxy. This differs from trabecular bone in as much as the gelatine cavities are essentially separated from each other (giving a so-called closed cell structure). In trabecular bone the cells are interconnected giving an open cell structure through which the marrow can flow. Figure 1 compares the cross sections of a sample of 60% porosity, the head of a femur and a lumbar vertebra.

Eight samples having porosities in the range 10% to 80% were made. The volumes of epoxy and gelatine required to give the various porosities were obtained by weighing, these weights being calculated after careful measurement of their densities. Values of the porosity of each sample were then obtained by relating their density to those of their constituents. All density measurements were made using Archimedes' principle.

3. Measurements on the trabecular phantom material

3.1. Instrumentation

Velocity and attenuation measurements at different frequencies were carried out using the experimental set-up shown in figure 2. This consists of a cubic ultrasonic water tank of side 15 cm with two identical 0.75 MHz broadband transducers centrally mounted in cut-outs on opposite faces so as to be in direct contact with the water. A 2% solution of a
proprietary detergent (Stardrops) was added to improve wetting. Transducer T1 is excited by a 1 kV pulse of duration 0.1 $\mu$s. This produces a broadband ultrasound pulse of length approximately 7 $\mu$s. The output from transducer T2 is captured by a digital oscilloscope and then transferred to floppy disc via an IEEE-488 interface and computer.

3.2. Velocity

The velocity of ultrasound in the phantoms was determined using the 'time of flight' method where the difference in the time taken for an ultrasound pulse to travel between the two transducers both with and without a sample of known thickness in place is noted. The velocity can then be calculated using the equation

$$V_s = \frac{d_s V_w}{d_s - V_w \Delta t}$$  (1)

where $V_s$ is the velocity of sound in the sample, $V_w$ is the velocity of sound in water, $d_s$ is the sample thickness and $\Delta t$ is the (time of flight in water only) -- (time of flight with sample present). The measured ultrasonic velocities of the phantom material were compared with two theoretical velocities. The first uses the relationship described by Wyllie et al (1956)
who suggested that the ultrasonic velocity in a porous material could be predicted by the equation

\[
\frac{1}{V_p} = \frac{\phi}{V_b} + \frac{1 - \phi}{V_a}
\]  

where \(V_p\) is the predicted velocity in the material, \(V_a\) is the velocity in the matrix material, \(V_b\) is the velocity in the pore material and \(\phi\) is the porosity of the material.

This equation was derived by analysing the times taken for a pulse to traverse a composite material made up of two slabs possessing different acoustic velocities. The porosity is defined in terms of the slab thicknesses and thus the equation is essentially one-dimensional.

The alternative hypothesis which we propose is that the velocity may change linearly with volume fraction and would therefore be more accurately predicted by the simpler expression

\[
V_p = V_a(1 - \phi) + V_b\phi.
\]  

Here the predicted velocity is directly dependent on the porosity (i.e. the relative volumes of pore and matrix materials) and as such should correspond more closely to the structure of the phantom material. There are potentially a number of other formulations but, in this case, a reasonable agreement with experiment was found with this simple version.

3.3. Attenuation

When assessing the ultrasonic attenuation in a material, allowance has to be made for reflection losses at the water-sample interfaces. This was achieved by measuring the transmitted amplitude for a number of thicknesses of each sample. It was assumed that the reflection losses would be constant for each thickness and hence changes in received amplitude would be due solely to losses within the sample.

The reduction in sample thickness was achieved by milling. This ensured that the sample faces were parallel and hence kept the experimental errors to a minimum; also, care was taken not to reduce the sample thickness to a degree where reverberations of ultrasonic pulses inside the sample would affect the amplitude measurements. A thickness range of 8–20 mm was found to be acceptable.

The received pulses were digitized and subjected to a fast Fourier transform (FFT). This produced a spectrum from which amplitudes at different frequencies could be obtained.

The attenuation coefficient of the phantom was determined graphically. This involved plotting \(20\log_{10}(\text{amplitude})\) at a given frequency against sample thickness in cm. The slope of the resultant straight line should therefore be equal to the attenuation coefficient, \(\alpha\), given in dB cm\(^{-1}\), thus;

\[
\alpha = \frac{20}{d_{s1} - d_{s2}} \log_{10} \left( \frac{A_1}{A_2} \right)
\]  

where \(d_{s1}\) and \(d_{s2}\) are sample thicknesses and \(A_1\) and \(A_2\) are the signal amplitudes at those thicknesses.
3.4. Broadband ultrasonic attenuation (BUA)

Commercial BUA devices generally measure the rate of change of attenuation with frequency between roughly 200 kHz and 600 kHz although the technique has only been applied to specific measurements on the os calcis (Langton et al. 1984). However, although an attempt is made to correct for water and diffraction losses, the sample thickness is not taken into account. Furthermore, there is an implicit assumption that the attenuation is linear with frequency over that range. The equipment used for our attenuation measurements had a poor signal-to-noise ratio below 400 kHz and therefore results are given within the frequency range 500-900 kHz. Although this is not ideal, we were able to confirm the validity of the data by measuring BUA values in a commercial BUA scanner as described below.

In this study the BUA value was measured in a number of our samples using a commercial device (Walker Sonix UBA 575). These samples were all rectangular blocks of dimensions 50 mm x 50 mm x 20 mm and were positioned with the 20 mm axis parallel to the ultrasonic beam.

![Figure 3. The relationship between porosity and velocity for the phantom, and predicted velocities (——) according to Wyllie et al. (1956), and (—-—) proposed. Error bars (indicating ± one standard error of the mean) lie within data symbols.](image)

4. Results and discussion

The variation of velocity in the trabecular phantom material for different values of porosity is shown in figure 3. It can be seen that the measured velocities seem to correspond better to the predicted velocity using equation (3) rather than equation (2). Ideally, the
lower limit of these values would be reduced further to provide a more comprehensive coverage of the range of velocity values quoted for trabecular bone (see table 3) although the range of reported values is large (McKelvie and Palmer 1987, Miller et al 1993). This could be achieved by replacing the gelatine with another material of lower velocity, and we are currently exploring a number of options. Since pulsed ultrasound was used for the measurement of velocities, the possibility of some dispersion cannot be entirely ruled out. Similarly, there was some initial concern about the possible errors due to non-linear propagation. This was investigated by sending pulses with amplitudes differing by 10 dB through the same material under the same conditions. The received signals were Fourier transformed and compared. Above 1 MHz, differences of up to 1 dB were found, the data points below 1 MHz had identical values within the limits of experimental precision and it was concluded that non-linear errors were negligible.

![Figure 4: Example of the determination of attenuation coefficient of the 50% porous sample at (♦) 0.5 MHz and (■) 0.75 MHz. The respective gradients and \( r^2 \) values are (♦) 11.45, 0.99 and (■) 17.37, 0.99.](image)

As stated above, the attenuation coefficient, \( \alpha \), at each frequency, was determined using a range of thicknesses as shown in figure 4. A linear least-squares regression was applied to the data points, the gradient of which gave the attenuation coefficient, \( \alpha \), in dB cm\(^{-1}\). The correlation coefficient \( r^2 \) had values for all samples ranging from 0.83 to 0.99 with a mean value of 0.97. The attenuation coefficients obtained for the various samples over a range of porosities and frequencies, including those of 0% porosity (pure epoxy) and 100% porosity (pure gelatine), are displayed in figure 5. It can be seen that whilst the attenuation
in very low and very high porosity samples is relatively small it increases as the porosity is changed and passes through a peak at roughly 50%.

The results for BUA obtained from the Walker Sonix UBA 575 are displayed in figure 6. The general shape of the data points shows a similarity to those in figure 5, both peaking at around 50% porosity.

It is too early to report on the long-term stability of these phantoms, although we are in the process of carrying out the relevant measurements. However, it is clear that some form of encapsulation is needed to prevent the gelatine from drying out. Present signs indicate that if a thin shell of epoxy is added, it not only provides a suitable seal but it also mimics the outer cortical shell. The temperature dependence of the values is also relevant since some equipment operates at 37 °C while others use room temperature. We have performed tests on the temperature dependence of the 80% porosity phantom which indicate that it has a temperature coefficient of roughly $-1.5 \text{ m s}^{-1} \text{ per } ^\circ\text{C}$ for velocity and $-0.06 \text{ dB MHz}^{-1} \text{ per } ^\circ\text{C}$ for BUA.

As mentioned previously, much work has been done on the ultrasonic characteristics of bone taken from various sources, both human and animal. The most pertinent results, however, are those relating to large volumes of cancellous bone, as it has been suggested that these would be more sensitive to changes in bone metabolism (Langton 1984). A comparison of the results from previous work (McKelvie and Palmer 1987) and those from the present study is shown in table 3. The data were obtained from bone mimics with porosities in the range 47%–76% and cancellous bone with densities quoted as being between 940 kg m$^{-3}$ and 1170 kg m$^{-3}$.
Table 3. A comparison of the ultrasonic characteristics of the bone mimic and real bone.

<table>
<thead>
<tr>
<th></th>
<th>Bone mimic</th>
<th>Trabecular bone</th>
</tr>
</thead>
<tbody>
<tr>
<td>Attenuation at 0.5 MHz (dB cm(^{-1}))</td>
<td>7.0-17.7</td>
<td>6.0-16.0(^a)</td>
</tr>
<tr>
<td>Velocity (m s(^{-1}))</td>
<td>1844-3118</td>
<td>1465(^b)-2084(^c)</td>
</tr>
<tr>
<td>BUA (dB MHz(^{-1}))</td>
<td>24-76</td>
<td>26-99(^d)</td>
</tr>
</tbody>
</table>

\(^a\) Tavakoli (1991).  
\(^b\) Miller et al (1993).  
\(^c\) McKelvie and Palmer (1987).  

Figure 6. The relationship between the porosity and the BUA of the material. Error bars indicate one standard error of the mean.

The peak in the attenuation at 50% porosity implies that the highly attenuating nature of the material is predominantly due to the scatter arising from the geometrical arrangements of its constituents; this is often referred to as the 'architecture' of trabecular bone. This seems not to have been reported in ultrasonic studies of real bone, probably because the full range of porosities is not readily available in biological samples.

The location of the peak in attenuation at 50% porosity could be explained as follows. A standard pore size was used for all samples, the porosity being altered by changing the pore density. It seems likely, therefore, that at high densities there is a degree of connectivity between adjacent pores. Now if, as mentioned above, the attenuation is due to inhomogeneities in the material, we can consider these inhomogeneities to be gelatine pores in an epoxy matrix on one side of the peak (0%-50%) whereas on the other side
(50%-100%) it is the gelatine which forms the continuous phase with particles of epoxy forming the scatterers. It is then not unreasonable to assume that the attenuation in both cases should be a maximum when the number of scatterers per unit volume is a maximum. It is clear that this is not an entirely accurate model for cancellous bone, where changes in both pore size and density take place. This may mean that in trabecular bone the attenuation peak occurs at a porosity other than 50%.

As previous workers have shown, the attenuating properties of the os calcis diminish with decreasing density (McCloskey et al 1990, McKelvie and Palmer 1987). Now it can be assumed that in this case density and porosity are inversely proportional and hence the attenuation in the os calcis decreases with increasing porosity. If, therefore, the attenuation coefficients of various porosities of trabecular bone display a similar peak to that given by the phantom, then the porosity of the os calcis should be greater than 50%. Although several studies into the properties of the os calcis have been undertaken, no direct information on its porosity was found. Density values obtained by means of the Archimedes’ principle are available. However, these measurements can be unreliable if great care is not taken to exclude all air from the sample or if there is a degree of uncertainty about the nature of the pore-filling material. For example, McKelvie and Palmer (1987) measured the density of samples of os calcis to lie in the range 940 to 1170 kg m⁻³, despite the density of marrow being (970 ± 60) kg m⁻³ and that of cortical bone (1990 ± 27) kg m⁻³ (ICRP 1975). Nevertheless, in general the densities of the os calcis always give a porosity well in excess of 50%. This is also relevant when considering the BUA values obtained. Several reports (Evans and Tavakoli 1990, Tavakoli and Evans 1991) have concluded that there is a high correlation (0.8 to 0.9) between trabecular bone density and BUA in vivo and in vitro and therefore changes in porosity, and hence also density, over a wide range may not be particularly relevant in the os calcis where there appears to be a relatively small range of values.

The existence of a peak in attenuation as the porosity is changed means that a lower than normal attenuation measurement at a particular skeletal site cannot be interpreted as an indication of osteoporotic bone without knowledge of the porosity of that bone. In fact, if attenuation measurements of a particular bone either fall on the ‘wrong’ side of the peak or straddle the peak this would lead to false conclusions being drawn as to the condition of the bone.

The nature of the ultrasonic attenuating mechanism in porous materials is the subject of debate. Two models have been suggested, one based purely on multiple scatter and the other including losses due to viscoelastic mechanisms first described by Biot (1956, 1962). Biot theory treats attenuation as an effect of the viscous forces that are created as fluid is moved though the porous matrix. The theory therefore requires a fluid filled, permeable structure rather than the closed cell structure filled with semi-solid gelatine which comprises the phantom. Since our inclusions are of the order of 1 mm in diameter and the wavelength is similar, then it is inevitable that scatter will be significant. In fact, it seems probable that it is the dominant mechanism in this phantom material.

The trabecular phantom material displays ultrasonic characteristics which are similar to those of trabecular bone, both velocity and attenuation being highly dependent on porosity; also, the range of velocities, attenuation coefficients and BUA values achievable match those found in real bone.

It seems probable that the dominant attenuating mechanism is scatter due to the material’s closed cell structure. Varying the porosity while using standard pore sizes changes the number of scatterers and hence changes the attenuating characteristics. The number of scatterers could also be altered by varying the pore size whilst keeping the
porosity constant. Hence it is possible that for this material the attenuation and velocity are
independently variable, although further work is required in order to verify this. Another
important potential development is the introduction of anisotropy into the phantom material.
This would be readily achievable by using gelatine strands aligned in some way rather than
the isotropic cubes described here. Such a material would have the advantage of being able
to mimic the directionality of the structures in the os calcis, thereby making the phantom
more realistic. However, in clinical practice, the os calcis is always examined in the same
plane and therefore anisotropic influences should be minimal. Work to investigate this is
now underway.

Acknowledgment

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financial support.

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Comparison of changes in bone mineral in idiopathic and secondary osteoporosis following therapy with cyclical disodium etidronate and high dose calcium supplementation

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Summary

OBJECTIVE Our clinical practice has been to offer treatment with cyclical disodium etidronate and high dose calcium supplements (1500–1600 mg/day) to all female patients with osteoporosis who are unable or unwilling to take hormone replacement therapy (HRT), and male osteoporotics. In a retrospective study we compared the effect of this treatment on measures of bone mineral over a 12-month period in women with post-menopausal and secondary osteoporosis. We also assessed its effects in 10 male osteoporotics.

DESIGN A retrospective analysis of 83 consecutive patients with osteoporosis who completed 12 months of treatment with disodium etidronate and calcium and who had a dual energy X-ray absorptiometry (DEXA) scan at baseline and following 12 months of therapy.

PATIENTS The study included 73 women (45 post-menopausal and 28 secondary osteoporotics) and 10 men with established osteoporosis as shown by spinal and femoral bone mineral densitites (BMD) > 2 standard deviations (SD) below young normals, and radiological evidence of osteoporosis.

MEASUREMENTS Each patient had routine biochemistry at baseline, an X-ray of thoracic and lumbar spine, and a DEXA scan of lumbar spine (L2–L4) and femoral neck. The DEXA scan was repeated following 12 months of therapy.

RESULTS There was no difference between increase in spinal BMD in the post-menopausal (5.7%) versus secondary osteoporotic group (6.7%). There was a significant increase in spinal BMD at 12 months in the 10 male osteoprotics (9.0%, P < 0.01). No overall change in femoral neck BMD was noted.

CONCLUSIONS Cyclical disodium etidronate given with high dose calcium supplements is equally effective in increasing spinal bone mineral density in post-menopausal and secondary osteoporosis. It also results in a significant rise in spinal bone mineral density in male osteoporotics. Whether this produces a reduction in fracture rates is unknown.

Osteoporosis is characterized by an absolute decrease in the amount of bone, leading to increased number of fragility fractures principally of the vertebrae, proximal femur and distal radius. Bone loss occurs because of an imbalance between bone resorption and formation, with the former exceeding the latter (Eastell et al., 1988). Measures of bone mass correlate inversely with fracture rate (Jensen et al., 1983). In post-menopausal women oestrogen deficiency, and consequently negative calcium balance, is a major cause of osteoporosis (Nordin & Morris, 1989; Nordin et al., 1991). In patients with secondary osteoporosis due to thyrotoxicosis or steroid therapy, bone resorption is stimulated and intestinal calcium absorption is inhibited (Mennier et al., 1984; Baran & Braverman, 1991). In steroid induced osteoporosis, bone formation is also inhibited by a direct effect on osteoblasts (Bressot et al., 1979). Anti-resorptive therapy combined with adequate calcium supplementation would therefore be a rational therapy for these forms of osteoporosis.

Bisphosphonates have complex actions on bone. They act mainly as anti-resorptive agents, limiting the accession of osteoclast precursors to bone, the formation of osteoclasts (Boonekamp et al., 1986) and direct bone resorption by mature osteoclasts (Carono et al., 1990). They may also act through osteoblasts affecting their proliferation and inhibiting the secretion of osteoclast stimulating factors (Khoker & Dandona, 1989). They have been shown to provide protection against osteopenia in a rat model (Wronska et al., 1991). In a clinical study, the bisphosphonate (3-amino-1-hydroxypropylidene)-1,1-bisphosphonate (APD)
given continuously with 1 g of calcium supplements per day has been shown to increase vertebral bone mineral density in patients treated with corticosteroids (Reid et al., 1988). Tiludronate has shown a more modest benefit when given continuously without calcium supplements in post-menopausal osteoporosis (Reginster et al., 1989). More recently, well designed clinical trials have reported that 400 mg of disodium etidronate daily, given for 2 weeks, followed by 500 mg of calcium supplements for 11 weeks, in repeated cycles over 2 and 3 years, resulted in an increase in spinal bone mineral and reduced the rate of vertebral fractures in post-menopausal osteoporosis (Storm et al., 1990; Watts et al., 1990).

Our clinical practice has been to treat all patients with established osteoporosis, who are not suitable for, or who refuse, hormone replacement therapy (HRT) (including secondary osteoporosis and male patients) with repeated cycles of disodium etidronate, 400 mg per day for 2 weeks followed by at least 1500 mg of calcium supplements per day for 11 weeks (recommended daily requirement for post-menopausal women, Consensus Development Conference, 1987). In a retrospective study, we have compared the effect of this treatment on spinal and femoral bone mineral in a group of post-menopausal women with that in a group of women with secondary osteoporosis. We have also evaluated its effect on 10 male patients with osteoporosis.

Patients

The study contained 45 women with post-menopausal osteoporosis (mean age 64.4 ± 1.2 years) and 28 women with secondary osteoporosis (mean age 60.5 ± 1.9). Mean Z scores at baseline in these groups were comparable (see Table 1). The group with secondary osteoporosis included patients with previous thyrotoxicosis (8), on oral steroid therapy (7), and with early surgical menopause (13). All other causes of secondary osteoporosis (such as coeliac disease, anorexia nervosa and acromegaly) were excluded because they represented only single cases. We also included 10 males with osteoporosis (see Table 2); the causes were idiopathic osteoporosis (7), hypogonadism (2) and previous gastric surgery (1). Male patients with hypogonadism were on stable treatment with testosterone supplements for at least 1 year prior to treatment with etidronate; their treatment was not altered during the period under study. Osteoporosis was defined as a spinal or femoral Z score of >2 SD below young normals or X-ray evidence of osteoporosis, such as vertebral body collapse. Vertebral collapse was defined as a reduction in the height of the whole or anterior vertebral body of > 20%. In a number of cases spinal Z scores were artefactually increased by vertebral collapse or abdominal aortic calcification, in which case patients were classified as osteoporotic by virtue of X-ray evidence of vertebral collapse.

Patients were excluded from the study for the following reasons: co-treatment with hormone replacement therapy (3), inability to take calcium supplements (2), and treatment with minocycline (which may bind calcium/etidronate in the gut) for acne (1). Treatment was generally well tolerated with only two patients who were unable to take calcium supplements withdrawing.

Methods

All patients had a clinical assessment, routine biochemical tests to exclude metabolic bone disease or other causes of osteopaenia, and standard spinal X-rays at baseline. They

<table>
<thead>
<tr>
<th>Table 1 Details of female patients</th>
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<tbody>
<tr>
<td></td>
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<tr>
<td>Post-menopausal women</td>
</tr>
<tr>
<td>(n = 45)</td>
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<tr>
<td></td>
</tr>
<tr>
<td>Secondary osteoporosis</td>
</tr>
<tr>
<td>(n = 28)</td>
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<tr>
<td></td>
</tr>
<tr>
<td>Age</td>
</tr>
<tr>
<td>64.4 ± 1.2</td>
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<tr>
<td>60.5 ± 1.9</td>
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<td></td>
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<td>Age at menopause</td>
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<tr>
<td>46.2 ± 0.7</td>
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<td>43.0 ± 1.1</td>
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<td>&lt; 0.001</td>
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<td>Years since menopause</td>
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<td>16.7 ± 1.3</td>
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<td>16.5 ± 2.0</td>
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<td>NS</td>
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<tr>
<td></td>
</tr>
<tr>
<td>Height (cm)</td>
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<tr>
<td>155.8 ± 0.9</td>
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<td>155.1 ± 1.5</td>
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<td>Weight (kg)</td>
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<td>57.3 ± 2.3</td>
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<td>NS</td>
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<td>Z score*</td>
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<td>spine</td>
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<td>-3.45 ± 0.20</td>
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<tr>
<td>-3.25 ± 0.28</td>
</tr>
<tr>
<td>NS</td>
</tr>
<tr>
<td>femur</td>
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<tr>
<td>-2.51 ± 0.19</td>
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</table>

* Number of SD below young normals.

<table>
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<tr>
<th>Table 2 Details of male patients</th>
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<tr>
<td></td>
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<tr>
<td>Male osteoporotic</td>
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<tr>
<td>Age</td>
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<td>49.5 ± 5.4</td>
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<tr>
<td>163.0 ± 10.2</td>
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<td>Weight (kg)</td>
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<td>88.3 ± 8.0</td>
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<td>Z score*</td>
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<td>spine</td>
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<tr>
<td>-3.59 ± 0.28</td>
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<tr>
<td>femur</td>
</tr>
<tr>
<td>-1.19 ± 0.52</td>
</tr>
</tbody>
</table>

Mean ± SEM.

* Number of SD below young normals.
each underwent a DEXA scan (Lunar Corporation, Madison, Wisconsin, USA) of the lumbar vertebrae (L2–L4) and femoral neck (reproducibility in normal controls 1·18% in spine and 1·69% in femur) (Truscott et al., 1993). Z score (number of SD below young normals) was derived from a study of 329 normal women (Truscott et al., 1993) and 122 normal men (unpublished data) from the local population. Measurements were performed at baseline and after 12 months treatment in all cases.

Treatment consisted of disodium etidronate (400 mg to be taken in the middle of a 4-hour fast at night for 2 weeks), followed by 1500–1600 mg of calcium supplements per day for 11 weeks. The calcium supplements were given according to patient preference. Each patient started on Sandocal (calcium lactate gluconate/calcium carbonate, Sandoz) 400 mg twice during the day and 800 mg at night initially. Twelve were changed subsequently to Calci chew (calcium carbonate, Shire Pharmaceuticals) 500 mg thrice daily as they found the initial treatment unpalatable. Each cycle was repeated every 3 months.

Statistics

Statistical analyses were performed with paired Student's t-test or Wilcoxon signed rank test. Comparisons of more than two data sets were done by one-way ANOVA with the Bonferroni adjustment when comparing individual groups. All data were expressed as mean ± SEM. Level of significance was set at a P value of < 0.05.

Table 3: Measures of bone mineral in post-menopausal and secondary osteoporotic women at baseline and 12 months

<table>
<thead>
<tr>
<th>Group</th>
<th>Baseline</th>
<th>12 months</th>
<th>Significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Post-menopausal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spinal BMD</td>
<td>0·765 ± 0·025</td>
<td>0·808 ± 0·025</td>
<td>&lt; 0·0001</td>
</tr>
<tr>
<td>Spinal BMC</td>
<td>28·65 ± 1·16</td>
<td>30·21 ± 1·13</td>
<td>&lt; 0·0001</td>
</tr>
<tr>
<td>Secondary osteoporosis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spinal BMD</td>
<td>0·789 ± 0·034</td>
<td>0·842 ± 0·003</td>
<td>&lt; 0·0001</td>
</tr>
<tr>
<td>Spinal BMC</td>
<td>30·20 ± 1·72</td>
<td>33·20 ± 1·65</td>
<td>&lt; 0·005</td>
</tr>
<tr>
<td>Post-menopausal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Femoral BMD</td>
<td>0·695 ± 0·019</td>
<td>0·708 ± 0·017</td>
<td>NS</td>
</tr>
<tr>
<td>Femoral BMC</td>
<td>3·36 ± 0·09</td>
<td>3·43 ± 0·10</td>
<td>NS</td>
</tr>
<tr>
<td>Secondary osteoporosis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Femoral BMD</td>
<td>0·699 ± 0·025</td>
<td>0·714 ± 0·027</td>
<td>NS</td>
</tr>
<tr>
<td>Femoral BMC</td>
<td>3·47 ± 0·164</td>
<td>3·47 ± 0·165</td>
<td>NS</td>
</tr>
</tbody>
</table>

Paired t-test (mean ± SEM.)

BMD, Bone mineral density (g/cm²); BMC, bone mineral content (g hydroxyapatite).

Table 4: Comparison of BMD in sub-groups of patients with secondary osteoporosis at baseline and 12 months

<table>
<thead>
<tr>
<th>Group</th>
<th>Baseline</th>
<th>12 months</th>
<th>Significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Post thyrotoxicosis (n = 8)</td>
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<td></td>
<td></td>
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<tr>
<td>Spinal BMD</td>
<td>0·858 ± 0·036</td>
<td>0·902 ± 0·028</td>
<td>&lt; 0·05</td>
</tr>
<tr>
<td>Femoral BMD</td>
<td>0·679 ± 0·022</td>
<td>0·707 ± 0·025</td>
<td>&lt; 0·05</td>
</tr>
<tr>
<td>Steroid therapy (n = 7)</td>
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<td></td>
<td></td>
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<tr>
<td>Spinal BMD</td>
<td>0·608 ± 0·08</td>
<td>0·684 ± 0·075</td>
<td>&lt; 0·05</td>
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<tr>
<td>Femoral BMD</td>
<td>0·606 ± 0·08</td>
<td>0·583 ± 0·074</td>
<td>NS</td>
</tr>
<tr>
<td>Early surgical menopause (n = 13)</td>
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<tr>
<td>Spinal BMD</td>
<td>0·844 ± 0·04</td>
<td>0·889 ± 0·033</td>
<td>&lt; 0·01</td>
</tr>
<tr>
<td>Femoral BMD</td>
<td>0·756 ± 0·027</td>
<td>0·778 ± 0·034</td>
<td>0·052</td>
</tr>
</tbody>
</table>

Wilcoxon signed rank test (mean ± SEM).

BMD, Bone mineral density (g/cm²).

Results

There was no significant difference between the post-menopausal women (n = 45) and those with secondary osteoporosis (n = 28) in terms of age, years since menopause, height, weight or Z scores at baseline (see Table 1). The secondary osteoporosis group tended to have an earlier menopause, 43·0 ± 1·1 years versus 46·2 ± 7·7 years (P < 0·01). The spinal Z scores in all three groups were not significantly different (ANOVA; F = 0·3033), but the femoral neck Z scores were significantly higher in the male group (Table 2) when compared with the post-menopausal and secondary osteoporotic groups (P < 0·05, P < 0·01 ANOVA; Bonferroni correction) (Table 1).

Spinal BMD and bone mineral content (BMC) rose in all groups (Tables 3 and 5). Percentage changes in BMD and BMC in the post-menopausal women and secondary osteoporotics were not significantly different (Fig. 1). There was no change in spinal area (L2–L4) between baseline and 12 months in either the post-menopausal or
secondary osteoporotic groups (37.25 ± 0.65 vs 37.20 cm$^3$ ±0.61; 38.9 ± 1.04 vs 38.85 cm$^3$ ± 0.98). There was no significant difference between percentage increase in spinal BMD (ANOVA $F=0.906$) or spinal BMC (ANOVA $F=1.557$) between groups (Figs 1 and 3). There was no significant change in femoral neck BMC or BMD in any of the groups studied (Tables 3 and 5). As femoral Z score in the male group was significantly higher than in the female groups they were not compared.

Baseline spinal BMD was significantly lower in the steroid group compared with the post-thyrotoxicosis and post-surgical menopause groups (ANOVA, Bonferroni correction $P<0.05$, $P<0.01$) (Table 4). Femoral neck BMD at baseline was also significantly lower in the steroid group compared with the post-surgical menopause group, but not the post-thyrotoxicosis group (ANOVA, Bonferroni correction $P<0.05$) (Table 4). Spinal BMD increased in all subgroups of secondary osteoporosis (Table 4). Percentage changes in spinal BMD were not significantly different in the sub-groups of secondary osteoporosis (ANOVA $F=2.786$) (Fig. 2). However, there was a significant difference between percentage changes in femoral neck BMD between groups (ANOVA $F=4.18$, $P<0.05$) (Fig. 2) with a significant increase in the post-thyrotoxicosis group ($P<0.05$) and a ‘borderline’ increase in the early surgical menopause group ($P=0.052$) (Fig. 2, Table 4). Both groups were significantly different from the steroid group ($P<0.05$, $P<0.05$ ANOVA, Bonferroni correction). No patient suffered an extra-axial fracture during the study period. We have no data on spinal fracture rate.

**Discussion**

The increase in spinal BMD and BMC in post-menopausal women and secondary osteoporotics were comparable and significantly different from baseline. The mean increases in
BMD and BMC in both groups were similar and the mean spinal areas (L2–L4) were unchanged. This indicates a real increase in bone mineral and not an artefactual increase in BMD produced by further vertebral collapse.

In the two major studies previously referred to (Storm et al., 1990; Watts et al., 1990) the former quoted a 5.3% increase in spinal BMC at 3 years and the latter a 5.2% increase in BMD at 2 years. A recent study (Miller et al., 1991) has shown a 15.7% increase in spinal BMD over 2 years in 47 post-menopausal women treated with a similar regimen to that of Watts et al. (1990) except that they may have received a higher dose of calcium and vitamin D (total daily calcium intake 1500mg/vitamin D 400–800IU). A subsequent similar study (Silberstein et al., 1992) has reported a mean increase of 6.6% in spinal BMD at 12 months, despite removing 15 of 42 patients from analysis due to technical difficulties in measuring BMD in spondylotic spines and 3 patients for not responding to treatment. All crushed vertebrae were excluded from analysis. They reported a mean increase in femoral neck BMC of 4.8% in 28 of 42 patients who ‘responded’ to treatment, with a fall in femoral neck BMD of 3.9% in the 14 ‘non-responders’.

Watts et al. (1990) excluded from analysis any vertebra which suffered a crush fracture either at baseline or during the study. Their results show a greater than 1% increase in spinal BMC in the control group which is contrary to the well recognized decline in spinal BMD in post-menopausal osteoporotics shown in longitudinal studies. This increase in the control group was not found by Storm et al. (1990) who did not exclude any vertebrae from analysis and quoted spinal BMC. Watts’s regimen was similar to Storm’s except for phosphate pretreatment (2g per day for 3 days) to stimulate ‘osteoblast activity’ prior to the commencement of etidronate. Both studies showed evidence of a reduction in the incidence of spinal fractures compared with controls. No consistent increase in femoral BMD was found by Watts et al. (1990) and all three femoral neck fractures which occurred in the study period were in the two treatment groups. Storm et al. (1990) did not report any hip data.

We have found no overall changes in femoral neck BMC or BMD at one year. However, there appeared to be a small increase for patients with previous thyrotoxicosis, with a value of borderline significance for those with an early post-surgical menopause. The lack of any significant change in bone mass at the femoral neck could be interpreted as a beneficial effect in view of the expected bone loss at this site in most of the patients studied. The less marked benefit of anti-resorptive therapy on the femoral neck may result from the lower percentage of trabecular bone and higher percentage of cortical bone in the femoral neck, compared with the vertebral. Evidence suggests that cortical osteoclasts are less sensitive to etidronate than trabecular osteoclasts (Chappard et al., 1991).

In conclusion, cyclical disodium etidronate given with high dose calcium supplements results in a significant increase in spinal bone mineral density and content in post-menopausal osteoporotics, women with secondary osteoporosis and male osteoporotics. The increase may be greater than that produced by a lower daily dose of calcium, but direct comparisons between studies is not possible. The impact of this treatment regimen on fracture rate is not known.

References


A portable system for measuring bone mineral density in the pre-term neonatal forearm

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Department of Paediatrics, Clarendon Wing, The General Infirmary, Leeds LS1 3EX

Abstract
Current systems used to measure bone mineral content (BMC) in the neonate have the major drawback that
the child must be well enough to be moved to the scanner. Consequently, low birth weight pre-term neonates,
a group at particular risk of mineral compromise, cannot be measured. This paper describes a portable
neonatal bone mineral device capable of measuring bone mineral in the incubator. It uses a radiation sensitive,
charge coupled device (CCD) to acquire a bone mineral image enabling bone mineral to be measured at
various sites. It measures bone mineral density (BMD) with a precision of 5.5 mg cm\(^{-2}\) in vivo, reduced to
7.5 mg cm\(^{-2}\) when repositioning between scans is taken into account. The procedure takes under 5 min with
an image acquisition time of 30 s and an absorbed radiation dose to skin of 6 μSv. Calibration has been
undertaken with aluminium foils of differing thickness to confirm the linearity of the system throughout the
intended measurement range. A regression line fitted to the data demonstrated linearity and correlation
between BMD and aluminium thickness with \(r=0.99\) \((p<0.0001)\). Preliminary measurements on pre-term
neonates show values of BMD ranging from 43 to 115 mg cm\(^{-2}\) in babies aged 23-41 weeks post-conception.
These figures are within the linear range of the system.

Improvements in neonatal intensive care have resulted in increased survival rate of very premature babies of
less than 28 weeks gestation. The incidence of osteopenia of prematurity has been shown to be in excess of 50% in
these particular infants [1]. This high incidence may be explained by the fact that the fetal skeleton is most
intensively mineralized during the third trimester of pregnancy when about 70% (30 g) of calcium may be incor-
porated [2, 3]. This means that the very premature infant tends to have low bone mineral content (BMC).
Conventional feeding regimes lead to far lower mineral accretion rates than those achieved in utero [4]. Because
of the need to understand and overcome this problem both calcium metabolism and skeletal mineralization of
the human fetus and neonate have been the subject of intensive study [5] but the lack of a simple, accurate
and safe means of determining BMC has been a limiting factor in progress in the field.

For many years radiography was used but radiographs appear normal until a deficit in BMC of the order of
20% is present [6]. Lack of consistency in sequential radiographs due to variations in both exposure and
development parameters mean that this technique is unsuitable for following infants undergoing different
nutritional regimes. The application of photon absorpti-
ometry to the determination of BMC in infants by Steichen et al [7] in 1978 was a major advance in the
field as it allowed the direct measurement of bone mineral content without the intermediate use of film with all
its problems.

Neonatal bone, even at term, contains only relatively small amounts of mineral, typically 0.2 g cm\(^{-2}\) [8]. To
measure these very small amounts of mineral, we built and validated the first machine designed specifically to
measure the very small bones in the neonatal forearm [9]. Using this instrument we then went on to define
osteopenia of prematurity [10], to describe the pace of mineral incorporation into bone in utero [11] and also
after birth [12]. This system also allowed us to measure mineralization in a number of pathological conditions
[13] and different nutritional regimes [14]. Prior to this
we described the apparent resolution of osteopenia of prematurity in pre-term babies [15]. One major problem
with this study was that the babies had to be well enough to be moved to the measuring instrument. The size of
the samples and sensitivity of the instrument meant that a difference of less than or equal to 15% between normal
term and premature babies at 71 weeks post-conception would not have been statistically significant. This latter
problem could have been addressed by increasing the size of the study groups but the problem of moving the
babies to the scanner proved more intractable. The sickest pre-term babies were not and could not be included in
our studies using the, then, current technology. These babies, typically, fail to thrive and have mineralization
patterns markedly different from those of premature babies who are well at birth and normal for dates.
A system measuring neonatal forearm bone

Lyon et al [16] reported a system for the measurement of bone mineral in babies who were ill enough to warrant intensive care. This system required the observer to obtain two X-ray films of the forearm at 40 kV and 120 kV using a mobile X-ray unit. The films were subsequently digitized using a light box and video camera and a form of dual energy X-ray absorptiometry was carried out. More recently Williams et al [17] have been using this technique to measure premature babies in the incubator. Although offering a method which causes minimal disturbance to the baby, the problems associated with film radiography still affect this system and are reflected in the reproducibility in phantom measurements of between 6% and 9.2% noted by the authors. This technique underestimates the value of bone mineral in phantom sets by between 15.6% and 18.5%. They also give no suggestions as to why this inconsistency in the underestimate occurs, but it is obviously a major problem with this technique. All estimates of reproducibility and accuracy are based on phantom work and one would anticipate that these values would deteriorate in vivo.

This paper describes a new measurement instrument which is: (1) portable; (2) compact enough to be placed in the incubator; (3) fast enough in operation to cause minimal disturbance to the neonate; (4) capable of imaging a section of the forearm in a reproducible manner; (5) low dose enough to allow measurements to be made at frequent intervals. Apart from the obvious advantages in the solution of the problem which made the design desirable, several other advantages accrued. As we can measure a neonate almost immediately after birth we should be able to separate the maternal in utero effects on mineralization, such as placental insufficiency, from the environmental effects experienced post-partum. Such a system would also allow the effects of differing nutritional regimes to be assessed whilst the baby is in the incubator. Similarly the effects of various therapies, on bone, could be monitored at regular intervals.

Apparatus

Equipment design

It is usual for forearm bone mineral scanners to measure bone mineral across a region, often scanning a larger region to identify an appropriately reproducible site. In adults, the point of separation between the distal radius and ulna has often been used. In neonates, little is known about mineralization along the bones of the forearm and so we had no preconceived notion about the optimum site for measurement, both in terms of reproducibility or clinical value. For this reason it was felt that it would be ideal to obtain a map of bone mineral values of a relatively large proportion of the distal forearm.

The equipment was developed using a radiation sensitive charge coupled device (CCD) camera which offered the following advantages:

1. A two-dimensional (2D) image of bone mineral content of the distal and mid-forearm could be obtained.

2. No scanning mechanism and so no moving parts would be involved. Thus the technology could be robust enough to be portable within a neonatal intensive care unit.

3. The device could be sufficiently compact to fit into an incubator.

4. The sensitivity of radiation sensitive CCDs would allow sufficient data to be acquired in 1 or 2 s with a low radiation dose [18].

The block diagram for the system is shown in Figure 1. Photons are produced at 27.5 keV by the decay of an $^{125}$I point source of half life 60 days. A cone beam of photons from the source is passed through the baby's forearm which is held in a water bolus between two parallel sheets of Perspex. Photons are differentially absorbed by bone of varying mineral content and hence the photon beam falling on the front part of the detector has different intensities per unit area. Each of these intensities will be inversely proportional to the bone mineral in the beam.

A bone mineral equivalent image can be calculated on a pixel by pixel basis by the application of the equation:

$$M_b = \left[ \rho_b \rho_t \sigma_b \sigma_t - \rho_t \rho_b \sigma_b \sigma_t \right] \ln\left( I_0 / I \right),$$

where $\rho_b$ and $\rho_t$ are the mass attenuation coefficients for bone mineral $\left( 3.1704 \text{ cm}^2 \text{ g}^{-1} \right)$ and water/soft tissue $\left( 0.48016 \text{ cm}^2 \text{ g}^{-1} \right)$ respectively, $\sigma_b$ and $\sigma_t$ are the density values for bone mineral $\left( 3.225 \text{ g cm}^{-3} \right)$ and water/soft tissue $\left( 1.0 \text{ g cm}^{-3} \right)$ respectively. $I$ is the photon intensity of an image pixel of the sample. $I_0$ is the photon intensity in the same pixel through the water bolus alone. $M_b$ represents bone mass per unit area and is in units of $\text{g cm}^{-2}$ above the soft tissue/background level (which is an arbitrary baseline).

To calculate BMC (grams) the bone image must be integrated within a region of interest (ROI) and an equivalent area in the non-bone background subtracted. The area normalized BMC (BMD) can then be calculated by dividing the BMC by the area of the ROI. Because of the mass attenuation coefficient used these values are in grams of hydroxyapatite.

Most other single photon absorptiometry (SPA) adult forearm BMC techniques do not use two separate acquisitions. In this system the $I_0$ data are acquired mainly to compensate for the different radiation path lengths which results from using a point source with a 2D detector array rather than a single collimated detector. Consequently, with a sample of uniform thickness in the beam, the acquired image of the signal intensity would be maximum at the centre and gradually decrease radially towards the edge of the image. This is due to the increase in radiation path length with distance from the centre (the inverse square law effect) coupled with an increase in sample thickness as the radiation beam passes through with increasing obliquity. The contribution of both these effects could be calculated mathematically from the precise geometry of the apparatus and correction could be made to the image. However, it was felt
that the use of an $I_0$ image would be more reliable for longitudinal studies as the precise equipment geometry could change over a long time period.

**Equipment construction**

A Photonic Science X-ray Imager is used; this is a high performance X-ray sensitive video camera, consisting of a thin polycrystalline layer of gadolinium oxysulphide on a fibre optic substrate, optically coupled to a high resolution, low light level, intensified CCD imager. The active imaging area is a 50 mm diameter circle and the captured image region is a rectangular region 40 mm wide by 30 mm deep. A thin aluminium foil in front of the gadolinium oxysulphide scintillator makes the imager light-tight allowing operation under normal ambient lighting conditions. Using an image intensifier and coherent fibre optic components the image is transferred from the scintillator to a solid state (CCD) image sensor.

The camera electronics produce an output signal in the form of a monochrome composite video signal of 1 volt peak to peak into a 75 Ω load complying with Comité Consultatif Internationale de Radiodiffusion (CCIR) video standards. This video signal provides the input to a Data Translation DT2867 integrated image processor and precision frame grabber with a 16 bit full image buffer (Data Translation, Marlboro, MA, USA). This card, which is controlled by a 80486 personal computer running at 66 MHz, is capable of real time image capture. We use it to capture images of 768 pixels across a video scan line and 512 pixels (lines) in the frame with an effective pixel size of 52 μm square, from which we use a region of 32 mm by 24.5 mm. The digital image may be viewed in real time before data accumulation takes place. When positioning is satisfactory a simple button press will start data collection by adding captured frames to the frame store until an image with sufficient detail is obtained. These data are then stored as a 16 bit image file and archived on a 1.6 GB hard disk drive for later analysis; an image of the water bolus on its own is also captured and stored for use as background in the calculation of a bone mineral image. The specification for this system was determined after a previously reported feasibility study [18].

The internal construction of the system is shown diagrammatically in Figure 2 and illustrated in Figure 3(a). The $^{125}$I source, 2 mm diameter, of initial activity 200 mCi, is held in a shielded container. In front of the container is a simple shutter operated by a self contained subminiature time delay relay unit controlled by a timer dial and push button shutter release. The source, which has a half life of 60 days, is renewed at 120 day intervals. Constant path length is maintained by the Perspex sided forearm and bolus holder located between the source and the camera. Source-to-detector distance is 9 cm and the source-to-sample distance is approximately 6.2 cm with 3.5 cm between the faces of the forearm holder. The bolus is a water filled condom which is wrapped around the forearm. It fills the space between the Perspex sides of the positioning device. An air gap between the positioning device and the detector minimizes the effect of scatter in the image. The video output, camera power and control signals are coupled between computer and instrument via an insulated cable at the rear of the unit.

**Electrical construction and safety**

Electrical construction and safety are consistent with IEC 601/1 and comply with all regulations for safety of the patient and operator. In use, the instrument is ensheathed in a non-conducting Perspex casing giving electrical isolation to both user and subject. The size of the instrument is 52 cm by 20 cm. The controlling computer
A system measuring neonatal forearm bone

is an IPC Porta PC P5-486 with docking station (IPC Corporation (UK), Lancashire, UK) having a footprint of 37 cm by 27 cm working into a Panasonic 5 inch monochrome monitor of size 15 cm by 25 cm (Matsushita Electric Industrial Co. Ltd, Osaka, Japan). This compact controlling system is mounted on a small trolley to improve mobility. The scanner is shown in use in an open incubator in Figure 3b, where it may be seen how compact the instrument is. This figure shows an open incubator in order not to obscure the view of the instrument in use. It is equally functional in a closed incubator.

Data acquisition software, which controls both the camera and the DT2867 frame grabber and frame store, was supplied by Photonic Science, the manufacturer of the X-ray imager. The software allows the user to specify real time viewing of the video image on the monitor, thus permitting the positioning of the forearm before data collection is commenced. Data collected are specified as a number of consecutive “grabbed” video frames added into the frame store. Any number of frames between one and 9999 may be specified but the user should be careful that no pixels overflow (they have a 16 bit limit). Once a satisfactory image is obtained it may be named and saved to disk. Files are later transferred to an archive and analysis system based on a DAN 66 MHz 486 desk top PC system with 1.6 GByte hard drive.

Data analysis

A typical digital image is shown in Figure 4. Working on a pixel by pixel basis a bone mineral image is
calculated and, if required, displayed. Visible bone parts of the image are actually the mineralized section of the radius and ulna. Bone mineral values can be obtained by integrating through an ROI placed in this mineralized section. Currently we are investigating methods for placing the ROI and as a guide have developed a "T" shaped cursor which the operator can place with the bar of the "T" at the end of the mineralized portion of the radius and the tail of the "T" along the axis of the bone as shown in Figure 4. The operator can then opt to position the ROI either at a fixed distance from the end of the bone or at a percentage of the bone length from its end. The length of the radius can be calculated by measuring the palpated distance between the ulnar styloid process and the olecranon and applying the regression equation established by James et al [9]. Results in this paper have been obtained using an ROI 1.5 mm x 5 mm located 1 cm from the mineralized end of the radius axial to the shaft.

Calibration and preliminary validation

The system was checked for linearity in the biological range of interest by measuring aluminium foil. Differing numbers of pieces of 15 μm thick aluminium foil were placed in a water bath in the scanner, measures of its thickness were obtained, and BMD equivalent values were calculated. These are shown plotted against the number of foils in Figure 5. Linear regression analysis applied to these points demonstrated a correlation coefficient of 0.999 (p < 0.0001) indicating a high degree of linearity in the mineral range of interest.

The reproducibility of the system was measured in vivo by two experiments. In the first of these two consecutive measurements were made on a set of 20 subjects without repositioning the forearm between scans. In the second, 16 subjects were scanned twice with the arm being removed from the system between scans. The precision was then calculated for each case using the method outlined by Bland [19]. This gave values of 5.5 mg cm⁻² (8%) and 7.5 mg cm⁻² (11%) respectively. The whole measurement procedure for a single scan took 5 min of which 30 s was patient acquisition time.

The absorbed radiation dose to skin associated with a single acquisition is approximately 6 μSv. This dose was calculated based on an air kerma rate of 30 μGy h⁻¹ GBq⁻¹ at 1 m [20] which allows for both γ and X-radiation. It should be remembered that this dose is given to a small portion of the forearm when comparing it with the typical dose for a paediatric chest X-ray of 40 μGy. As the forearm contains only skin, bone and muscle and the beam impinges on a very small region of the forearm the effective dose to the subject is negligible.

After establishing this linear relationship and the precision of the system we went on to measure pre-term babies at a range of gestational ages from 23 weeks to 41 weeks.

The measurement procedure was well tolerated by the babies studied. All measurements were made at less than 48 h of age and included babies of extreme prematurity undergoing mechanical ventilation. None of them showed any physiological evidence of distress or deterioration in their condition during the image collection. Despite their requirement for intensive care, the most immature babies were often easier to measure than their more mature counterparts as they tended to be less active. More active babies required more patience to
Assessment of Renal Osteodystrophy in Dialysis Patients: Use of Bone Alkaline Phosphatase, Bone Mineral Density and Parathyroid Ultrasound in Comparison with Bone Histology

Abstract
Bone biopsies were studied in 73 patients to determine if a two-site radioimmunometric assay for serum bone alkaline phosphatase (BAP), total serum alkaline phosphatase (ALP), serum intact parathyroid hormone (iPTH), hand X-rays, regional bone mineral density (BMD) measurements and parathyroid enlargement detected by ultrasonography could accurately predict renal osteodystrophy. In the patients studied 57 had hyperparathyroid bone disease, 4 mixed renal osteodystrophy, 3 adynamic bone disease, 1 osteomalacia and 8 normal histology. Serum BAP, ALP and iPTH correlated positively with mineral apposition rate, osteoblastic, osteoid and eroded surface. In the diagnosis of hyperparathyroid bone disease serum iPTH was the most sensitive investigation, detecting 81% of patients at a level >100 pg/ml but with a specificity of only 66%. Serum BAP was more sensitive, 70% at a level of >10 ng/ml, than serum total ALP, 30% at a level of 300 IU/l, with similar specificities, 92 and 100%, respectively. Ultrasound detection of an enlarged parathyroid gland had a sensitivity of 64% and a specificity of 100% for the diagnosis of hyperparathyroid bone disease. Hand X-rays had a poor sensitivity, 47%, but a high specificity, 92%, for the detection of hyperparathyroid bone disease. The majority of patients had regional BMD values within the normal reference range and this test was of poor discriminatory value. The non-invasive markers were unable to distinguish between patients with low turnover, mild hyperparathyroidism and patients with normal histology. In conclusion the measurement of serum iPTH is a useful screening tool for the detection of hyperparathyroid bone disease which can be confirmed by the finding of a raised serum BAP or parathyroid enlargement. For definitive diagnosis, however, the gold standard remains bone biopsy and at present one cannot recommend any non-invasive method as an adequate substitute.

Key Words
Renal osteodystrophy
Bone alkaline phosphatase
Regional bone mineral density
Parathyroid ultrasonography
Introduction

Bone disorders resulting from abnormalities in mineral metabolism are common in patients with renal disease, and their diagnosis and treatment remains a considerable challenge to clinical nephrologists [1]. A reliable diagnosis of metabolic bone disease can be obtained from a transiliac bone biopsy, but this is an invasive procedure unsuited for repeated routine use. Biochemical and radiological investigations are used to try to identify high and low turnover renal osteodystrophy (ROD). Hand X-rays, despite their poor sensitivity, have traditionally been used to detect hyperparathyroid bone disease [2]. Serum intact parathyroid hormone (iPTH), even though not a direct product of bone metabolism, is currently accepted as the most clinically useful marker of high turnover ROD [3, 4]. Dysraphic bone disease, which is being diagnosed with increasing frequency in chronic renal failure, is particularly difficult to establish with traditional markers of bone turnover [5, 6]. The inability to diagnose metabolic bone disease accurately in uraemic patients from serum biochemistry has led to continuing work to develop plasma markers of bone turnover. Serum osteocalcin and collagen breakdown products are renally excreted and although they accurately predict bone turnover in non-renal and pre-dialysis patients, due to the interference of breakdown products their use is limited in dialysis patients [7, 8].

Alkaline phosphatase (ALP), a product of osteoblasts, has a long history as a marker of bone turnover in metabolic bone disease [9]. Common routine measurement of serum total ALP lacks specificity for bone disease due to the inclusion of the activity of all isoenzymes, principally from liver, bone, gut and kidney in biochemical assays [10]. Serum bone-specific alkaline phosphatase (BAP) has shown a greater correlation with abnormal histology in renal failure than serum total ALP [11], but the accurate measurement of specific isoenzymes has required complicated separation techniques based upon electrophoresis, whereas more routine methods of isoenzyme separation, including thermal stability, have provided only a qualitative differentiation of limited clinical value [12-14]. The recent development of a direct radio-immunoassay, using two monoclonal antibodies to BAP, enables a more precise assessment of osteoblastic activity [15].

The aim of the present study is to determine the relative efficacy of certain non-invasive techniques to diagnose renal bone disease using histologically and histomorphometrically defined bone biopsies. In particular, the use of BAP as a plasma biochemical index of bone formation, regional bone mineral density (BMD) measurement, hand X-ray analysis, and ultrasound examination of the neck for parathyroid enlargement was evaluated.

Subjects and Methods

Patients

Seventy-three patients (39 male; 34 female) were recruited, in this cross-sectional study, from the dialysis units of Leeds General Infirmary and St. James’s University Hospital (age range 23-78 years, median 48; time on dialysis 0-14 years, median 3). All patients were medically stable with no changes in therapy for at least 6 months prior to their biopsy. In the 3 months preceding their biopsy there was no change in the dosage or method of administration of vitamin D, although there were minor modifications of dietary phosphate and phosphate binders. Bone biopsies were performed on 39 patients with a known elevated serum iPTH, 32 prior to entering a trial of pulsed calcitriol, and 7 at the time of parathyroidectomy. Thirty-three patients were biopsied when an opportunity arose for a general anaesthetic, 19 at the time of transplantation, 9 when peritoneal dialysis catheters were inserted or repositioned, and 5 at the time of miscellaneous operations unrelated to dialysis. A single patient with clinical features of chronic aluminium overload and a positive deri oxamine stimulation test was biopsied to assess the severity of aluminium deposition prior to treatment. Forty-six patients were on peritoneal dialysis and 28 were on haemodialysis. Table 1 lists the causes of end-stage renal failure in the subjects. All patients gave informed consent and the study was approved by the local Ethics Committees of both hospitals.

Bone Histology

Anterior iliac crest bone biopsies were taken from the 73 patients, 20 of whom had received tetracycline double labelling; 250 mg of oxytetracycline was taken twice daily for 4 days, followed 10 days later by 150 mg of demeclocycline twice daily for 4 days, and bone biopsy 4 days later. Biopsies were taken using a Sheffield trephine with a 7-mm diameter bore (Bolton Surgical Services Ltd., Sheffield, UK). All patients were under general anaesthesia with 2% Maracaine locally infiltrated and a 100 mg diclofenac suppository for analgesia. Patients tolerated the procedure well but there was one case of persistent haemorrhage (which settled when packed) and two wound infec-

<table>
<thead>
<tr>
<th>Disease</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glomerulonephritis</td>
<td>18</td>
</tr>
<tr>
<td>Chronic interstitial nephritis</td>
<td>18</td>
</tr>
<tr>
<td>Diabetes</td>
<td>11</td>
</tr>
<tr>
<td>Hypertension and renovascular</td>
<td>10</td>
</tr>
<tr>
<td>Polycystic</td>
<td>9</td>
</tr>
<tr>
<td>Hereditary (Fabry’s, Alport’s)</td>
<td>4</td>
</tr>
<tr>
<td>Vasculitis</td>
<td>3</td>
</tr>
</tbody>
</table>
ions requiring antibiotics. All biopsies underwent histological examination, 57 were examined histomorphometrically, fragmentation during the collection process making the rest unsuitable for the latter procedure.

The tetracycline-labelled specimens were fixed immediately in 70% alcohol, while the remainder were fixed in 10% phosphate-buffered formalin, pH 7.3. Unfixed calcified specimens were dehydrated in alcohol, cleared in xylene and embedded in methyl methacrylate. Sections, 10 μm thick, were cut on a Jung k heavy duty microtome (Reichert-Jung, Heidelberg) and stained for histomorphometry [16] with 1% toluidine blue stain for 30 min or a modification of the Goldner stain [4]. In addition, sections 5–7 μm thick were stained with 1% solochrome azurine, pH 5, for approximately 18 h for the identification and localisation of aluminium impeding the bone surface [17]. The relative surface extent of aluminium staining was subjectively assigned as <10% (normal-low Al deposition), 10–33% (low-moderate Al deposition), 33–60% (moderate-high Al deposition) and >60% (extensive Al deposition) [17]. Other sections were stained with haematoxylin and eosin for routine histological examination. Histomorphometry was possible on 57 biopsies using a semi-automatic image analysing system OsteoMeasure (OsteoMetrics Inc., Atlanta, USA) and an epifluorescence microscope with ultraviolet light for the analysis of the tetracycline labels. In addition to the relative bone area within each specimen, a comprehensive range of formation and resorption variables was measured without prior knowledge of the corresponding biochemistry. On the basis of the histology, the biopsies were classified according to the following criteria:

1. Normal bone biopsies: no evidence of increased or decreased bone turnover and normal osteoid distribution and thickness;
2. Aplastic bone disease: low bone formation rate with flat osteoblasts, few osteoid seams and rare osteoclasts and resorption cavities;
3. Osteomalacia: low and diffuse uptake of tetracycline, wide and extensive osteoid seams from which cuboidal-shaped osteoblasts are often absent and rare osteoclasts and resorption cavities;
4. Mixed renal osteodystrophy: combination of osteomalacia and hyperparathyroidism with poor mineralisation and wide and extensive osteoid seams in conjunction with increased numbers of cuboidal-shaped osteoblasts, osteoclasts and resorption cavities indicative of hyperparathyroidism;
5. Hyperparathyroidism: a) mild - localised areas of increased osteoblastic and osteoclastic activity, a slight increase in osteoid tissue and resorption cavities but no evidence of periosteal fibrosis; b) moderate - more general increase in osteoblastic and osteoclastic activity, with increased osteoid tissue, resorption cavities with clearly defined but limited peritrabecular fibrosis; c) severe - extensive osteoblastic and osteoclastic activity with increased osteoid tissue, usually of normal thickness, resorption cavities which are increased in both extent and depth, areas of woven bone and extensive periosteal fibrosis.

Biochemical Analysis

The following biochemical measurements were made on stable patients under standard conditions: serum calcium, phosphate, bicarbonate, and total ALP (range 100–300 IU/l); BM Hitachi Autoanalyzer, Boehringer Mannheim, Poole, Dorset, UK), serum iPTH (Nichols Institute, normocalemic range 11–55 pg/ml) and serum BAP (Tandem OSTASE Hybritech, San Diego, Calif., USA, reference range <19 ng/ml), serum aluminium (atomic absorption, Perkin-Elmer, reference range 0.00–1.85 μmol/l). Standard calcium was calculated by the equation of Payne et al. [18]: Std Ca = Ca + (40 – albumin) × 0.025.

The samples were drawn from peritoneal dialysis patients immediately prior to their biopsy and for haemodialysis patients these samples were taken prior to dialysis. There was no significant difference in their serum calcium, phosphate, bicarbonate, total ALP and iPTH between these study samples and the routine pre-dialysis samples taken to monitor the adequacy of dialysis. In the 15 patients who underwent fine needle aspiration of their parathyroid glands, there was no significant difference between the biochemical values drawn prior to their gland aspiration and those taken prior to bone biopsy which was performed after the glands were sampled.

BMD Measurement

BMD of the lumbar spine (L2–L4), right femoral neck and total body were determined in 65 patients prior to biopsy, by dual energy X-ray absorptiometry (DEXA) using a Lunar DPX-L bone densitometer (Lunar Radiation Co., Madison, Wis., USA). Software version 1.3 was used for analysis of spine and femoral neck with extended research mode selected for total body analysis. Results were expressed as Z scores, the number of standard deviations of the measured value from age- and sex-predicted normal values obtained from a local population, reference range ± 2 SD [19]. Single photon absorptiometry of the dominant arm was performed using a Nuclear Data (ND1100A) bone densitometer. Bone mineral measurements were made at sites distal (approximately 25% trabecular bone) and proximal (approximately 75% trabecular bone) to an 8-mm gap between the radius and ulna [20]. Scan analysis was made using VOS software. Z scores were derived by comparison with a local control group.

Ultrasonography

High resolution ultrasound of the neck was performed on 53 patients, prior to biopsy (Ultramark 9 with a L10–5 MHz linear array broadband transducer, Advanced Technology Laboratories, Bothwell, Wash., USA) by 2 radiologists blinded to histology. The detection limit was 0.016 cm³. Since normal parathyroid glands could not be detected, identification of parathyroid tissue was considered a positive scan. Each gland was measured in three dimensions (X, Y, Z) and the volume calculated, assuming it to be an ellipsoid [23]. In 15 patients cytological examination of ultrasound-guided fine needle aspirates confirmed the presence of parathyroid cells whilst elevated aspirate iPTH concentrations indicated a functioning adenoma.

All results are expressed as mean ± SD with Student's t test and Pearson product-moment correlation used as appropriate, using AS-TUTE statistical calculator computer program.
### Table 2. Bone biopsy results with ancillary biological and biochemical data

<table>
<thead>
<tr>
<th>Histology</th>
<th>n</th>
<th>Age years</th>
<th>Time on dialysis years</th>
<th>Standard calcium (2.2-2.4 mmol/l)</th>
<th>Phosphate (0.8-1.2 mmol/l)</th>
<th>Bicarbonate (22-26 mmol/l)</th>
<th>Total ALP (100-500 IU/l)</th>
<th>Bone ALP (0.01-99 ng/ml)</th>
<th>iPTH (1-55 pg/ml)</th>
<th>Serum aluminium (0.01-1.85 µmol/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>8</td>
<td>52.8±12.1</td>
<td>1.5±0.9</td>
<td>2.46±0.23</td>
<td>1.83±0.23</td>
<td>21.7±3.0</td>
<td>147±63</td>
<td>4.24±1.68</td>
<td>99±102</td>
<td>0.36±0.12</td>
</tr>
<tr>
<td>Adynamic</td>
<td>3</td>
<td>54.0±10.6</td>
<td>1.7±1.1</td>
<td>2.43±0.18</td>
<td>1.98±0.31</td>
<td>15.6±2.6</td>
<td>161±53</td>
<td>9.9±3.3</td>
<td>140±187</td>
<td>0.59±0.49</td>
</tr>
<tr>
<td>Osteomalacia</td>
<td>1</td>
<td>48</td>
<td>2.5</td>
<td>1.59</td>
<td>25.6</td>
<td>116</td>
<td>6.9</td>
<td>28</td>
<td>0.28</td>
<td></td>
</tr>
<tr>
<td>Mixed</td>
<td>4</td>
<td>51.5±3.5</td>
<td>1.3±0.5</td>
<td>2.56±0.28</td>
<td>2.0±0.2</td>
<td>21.6±4</td>
<td>141±34</td>
<td>7.0±1.6</td>
<td>68±70</td>
<td>0.38±0.12</td>
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<td>Mild HPT</td>
<td>11</td>
<td>42.7±12.2</td>
<td>3.5±4.6</td>
<td>2.42±0.1</td>
<td>1.89±0.14</td>
<td>19.8±3.2</td>
<td>126±35</td>
<td>5.73±1.7</td>
<td>54±27</td>
<td>0.52±0.27</td>
</tr>
<tr>
<td>Moderate HPT</td>
<td>17</td>
<td>43.6±12.3</td>
<td>3.1±5.6</td>
<td>2.39±0.03</td>
<td>2.01±0.43</td>
<td>20.2±4.8</td>
<td>237±78**</td>
<td>15.8±10.4</td>
<td>405±232**</td>
<td>0.75±0.35</td>
</tr>
<tr>
<td>Severe HPT</td>
<td>29</td>
<td>48.6±13.6</td>
<td>2.5±0.03</td>
<td>2.06±0.53</td>
<td>20.7±4.4</td>
<td>361±190**</td>
<td>43.9±31.5**</td>
<td>780±455**</td>
<td>0.80±0.48</td>
<td></td>
</tr>
</tbody>
</table>

Results are shown as mean ± SD. Reference values for each parameter are indicated. Statistically different values when compared with the patients with normal histology are shown as *p<0.05; **p<0.01; ***p<0.001. HPT = Hyperparathyroidism.

### Table 3. Histomorphometric and histodynamic data for the study population

<table>
<thead>
<tr>
<th>Histology</th>
<th>n</th>
<th>Bone volume BV/TV (23.19±4.37%)</th>
<th>Osteoid volume OV/BV (1.48±0.93%)</th>
<th>Osteoid thickness (10.3±4.21 µm)</th>
<th>Total osteoid surface OV/BS (12.10±4.64%)</th>
<th>Osteoblast surface OB/BS (4.39±1.94%)</th>
<th>Eroded Osteoclast surface ES/BS (4.09±2.53%)</th>
<th>Osteoclast surface Oc/BS (0.69±0.01%)</th>
<th>Mineral appositional rate MAR (0.62±0.2 µm/day)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>6</td>
<td>22.38±7.64</td>
<td>1.63±2.5</td>
<td>8.82±2.09</td>
<td>7.37±10.97</td>
<td>0.51±0.21</td>
<td>3.14±3.47</td>
<td>1.14±1.13</td>
<td>-</td>
</tr>
<tr>
<td>Adynamic</td>
<td>2</td>
<td>20.91</td>
<td>0.45</td>
<td>7.68</td>
<td>3.41</td>
<td>-</td>
<td>7.59</td>
<td>1.62</td>
<td>-</td>
</tr>
<tr>
<td>Osteomalacia</td>
<td>1</td>
<td>23.04</td>
<td>13.19**</td>
<td>19.17**</td>
<td>32.59**</td>
<td>0.36</td>
<td>1.36</td>
<td>0.159</td>
<td>-</td>
</tr>
<tr>
<td>Mixed</td>
<td>3</td>
<td>31.29±15.12</td>
<td>8.57±3.71**</td>
<td>14.89±3.74**</td>
<td>43.12±11.31**</td>
<td>0.24</td>
<td>2.49±2.74</td>
<td>1.62</td>
<td>0.86</td>
</tr>
<tr>
<td>Mild HPT</td>
<td>9</td>
<td>22.24±4.94</td>
<td>1.91±2.00</td>
<td>10.98±2.34</td>
<td>8.1±6.96</td>
<td>0.68±0.25</td>
<td>2.12±2.63</td>
<td>0.43±0.28</td>
<td>0.58±0.14</td>
</tr>
<tr>
<td>Moderate HPT</td>
<td>12</td>
<td>21.77±4.56</td>
<td>6.49±4.34*</td>
<td>11.47±2.98</td>
<td>25.49±16.41</td>
<td>1.35±0.99</td>
<td>6.66±3.23</td>
<td>1.69±1.17</td>
<td>1.21±0.43</td>
</tr>
<tr>
<td>Severe HPT</td>
<td>24</td>
<td>23.37±7.02</td>
<td>12.05±8.69**</td>
<td>16.39±7.35**</td>
<td>37.96±16.66**</td>
<td>2.53±2.51**</td>
<td>9.73±4.64**</td>
<td>2.46±2.46</td>
<td>1.11±0.47</td>
</tr>
</tbody>
</table>

Values are shown as mean ± SD. Where the number of patients is <2 individual values are given. Statistically different values when compared with the patients with normal histology are shown as *p<0.05; **p<0.01; ***p<0.001. HPT = Hyperparathyroidism.

### Results

#### Histology

The histological results classification of the 73 bone biopsies with ancillary biological and biochemical data are shown in table 2. There were 57 patients with predominant hyperparathyroid bone disease which included 1 patient with severe hyperparathyroid bone disease and severe aluminium deposition. There were 4 patients with mixed ROD and 4 patients with low turnover ROD, comprising 3 adynamic and 1 case of osteomalacia. There were 8 patients with normal bone histology. The high incidence of hyperparathyroidism in this study can be accounted for by patient selection. There was no statistical difference in age between histological subgroups. Hyperparathyroid patients tended to be on dialysis longer than other histological subgroups, although this reached significance only between severe hyperparathyroid and normal histology.

The histomorphometry of the 57 undamaged bone biopsies is shown in table 3. The patients with normal and mild hyperparathyroidism on bone biopsy had histomorphometric parameters comparable with the range of the normal population included in the Osteo Measure software package and with our own previously published data. One of the adynamic biopsies had a raised osteoclast surface but the overall histomorphometric and histological picture was that of low bone formation. In the absence of aluminium staining a diagnosis of adynamic bone disease was reached by the 2 observers independently.
The incidence of bone surface aluminium deposition was low; 7 patients (9.6% of the 73 biopsies) had mild deposition, 2 (2.7%) had moderate deposition and the patient with clinical aluminium toxicity had severe deposition. At the time of biopsy 11 (15%) were being prescribed aluminium phosphate binders, although 35 (48%) had been prescribed aluminium hydroxide at some time on dialysis. Mean serum aluminium levels were within the reference range for all histological groups. The patient with severe aluminium deposition had been prescribed aluminium hydroxide for 9 years as a phosphate binder with a serum aluminium within the normal range throughout this time. Persistent bone pain and myopathy raised the clinical suspicion of aluminium toxicity, which was confirmed by a desferrioxamine stimulation test and a bone biopsy. There was no correlation between the severity of bone aluminium deposition and serum aluminium.

Histopathological examination of the marrow revealed 2 unsuspected cases of non-Hodgkin’s lymphoma.

Biochemistry

There was no statistically significant difference in plasma standard calcium, phosphate and bicarbonate concentrations between the subgroups studied, mean levels lying within the adult reference range. Mean serum iPTH levels were raised in all subgroups except the patient with osteomalacia. Mean serum total ALP and BAP levels were within the reference range for all subgroups except those with severe hyperparathyroidism. Moderate and severe hyperparathyroidism showed significant elevated serum iPTH, total ALP and BAP when compared with patients with normal histology (table 2).

Serum BAP, total ALP and iPTH all correlated significantly with mineral apposition rate, osteoid surface, osteoblast surface and eroded surface (table 4). There was a low degree of correlation between the osteoclast surface and the plasma markers of bone turnover. There was a wide variation in the values for osteoclast surface within the histological groups as exemplified by the adynamic patient described previously which the authors are unable to explain.

In the diagnosis of hyperparathyroid bone disease a serum iPTH >100 pg/ml had a specificity of 66% and a sensitivity of 81%. A serum BAP >10 ng/ml had a specificity of 92% and a sensitivity of 70%. A serum total ALP >300 IU/l had a specificity of 100% but a sensitivity of 30%. When the investigations were used in combination a serum iPTH >100 pg/ml and serum BAP >10 ng/ml produced a specificity of 100% and a sensitivity of 66%. A serum iPTH >100 pg/ml and a serum total ALP >300 IU/l had a combined specificity of 100% but a sensitivity of 30%.

Bone Mineral Density

BMD for each histological group is shown in table 5. Mean Z scores for each histological subgroup, except for the spinal measurements for patients with mixed ROD, were within 2 SD of a matched control population. Patients with normal bone histology had a negative mean Z score in the axial skeleton reflecting a loss of mineralisation. There was a negative relation between severity of hyperparathyroid bone disease and BMD in all regions measured. Patients with mixed, mild and moderate hyperparathyroid bone disease had a higher mean axial BMD measurement than patients with normal histology or adynamic bone disease. In the forearm there was an increasing differential loss of BMD between the proximal and distal forearm with increasing severity of hyperparathyroid bone disease which reached significance in patients with severe hyperparathyroidism, p < 0.005.

Radiology

There was no radiological evidence of hyperparathyroid bone disease in the histologically defined mild subgroup. Four moderate patients and 23 severe patients had radiological evidence of hyperparathyroid bone disease on hand X-rays (table 6). There was no significant difference between the detection rate of subperiosteal erosions and acro-osteolysis in patients with hyperparathyroid bone disease. The specificity of a positive hand X-ray for the diagnosis of hyperparathyroid bone disease was 92% and the sensitivity 47%. When combined with a serum

Table 4. The correlation between the biochemical markers of bone turnover and histomorphometric variables

<table>
<thead>
<tr>
<th></th>
<th>Total ALP</th>
<th>Bone ALP</th>
<th>PTH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total osteoid surface</td>
<td>0.44**</td>
<td>0.55***</td>
<td>0.46**</td>
</tr>
<tr>
<td>Osteoid thickness</td>
<td>0.22</td>
<td>0.35*</td>
<td>0.31*</td>
</tr>
<tr>
<td>Osteoblast surface</td>
<td>0.51***</td>
<td>0.49**</td>
<td>0.37*</td>
</tr>
<tr>
<td>Eroded surface</td>
<td>0.40**</td>
<td>0.36*</td>
<td>0.51***</td>
</tr>
<tr>
<td>Osteoclast surface</td>
<td>0.16</td>
<td>0.16</td>
<td>0.20</td>
</tr>
<tr>
<td>Mineral appositional rate</td>
<td>0.42**</td>
<td>0.54***</td>
<td>0.49*</td>
</tr>
</tbody>
</table>

The data is presented as the r values obtained by Pearson product-moment correlation. Statistically significant values are shown as *p < 0.05; **p < 0.01; ***p < 0.001.
Table 5. Regional BMD expressed as a Z score ± SD

<table>
<thead>
<tr>
<th>Histology</th>
<th>Number</th>
<th>Spine</th>
<th>Femur</th>
<th>Total body</th>
<th>Distal forearm</th>
<th>Proximal forearm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>7</td>
<td>0.38±0.21</td>
<td>-0.87±1.1</td>
<td>-0.93±1.4</td>
<td>-0.87±1.2</td>
<td>-0.83±1.2</td>
</tr>
<tr>
<td>Adynamic</td>
<td>1</td>
<td>1</td>
<td>-0.86</td>
<td>-0.57</td>
<td>-0.23</td>
<td>-1.85</td>
</tr>
<tr>
<td>Mixed</td>
<td>4</td>
<td>2.85±1.98</td>
<td>-0.09±1.1</td>
<td>-0.10±0.78</td>
<td>0.01±1.58</td>
<td>-0.17±2.12</td>
</tr>
<tr>
<td>Mild HPT</td>
<td>9</td>
<td>0.49±1.1</td>
<td>-0.4±1.3</td>
<td>0.05±0.56</td>
<td>0.03±1.0</td>
<td>-0.16±0.9</td>
</tr>
<tr>
<td>Moderate HPT</td>
<td>15</td>
<td>-0.06±1.3</td>
<td>-0.24±0.87</td>
<td>-0.3±1.27</td>
<td>-0.5±1.4</td>
<td>-0.8±1.5</td>
</tr>
<tr>
<td>Severe HPT</td>
<td>25</td>
<td>-0.77±1.6</td>
<td>-0.95±0.9</td>
<td>-1.44±1.5</td>
<td>-1.23±1.5</td>
<td>-1.94±1.9</td>
</tr>
</tbody>
</table>

HPT = Hyperparathyroidism.

Table 6. Radiological evidence of hyperparathyroidism in hand X-rays from the study population

<table>
<thead>
<tr>
<th>Histology</th>
<th>Number</th>
<th>Acro-osteolysis grade I and II</th>
<th>Grade III</th>
<th>Subperiosteal erosion grade I</th>
<th>Grade II and III</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>8</td>
<td>8</td>
<td>0</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>Adynamic</td>
<td>3</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Osteomalacia</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Mixed</td>
<td>4</td>
<td>4</td>
<td>0</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Mild HPT</td>
<td>10</td>
<td>10</td>
<td>0</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>Moderate HPT</td>
<td>17</td>
<td>14</td>
<td>3</td>
<td>13</td>
<td>4</td>
</tr>
<tr>
<td>Severe HPT</td>
<td>29</td>
<td>10</td>
<td>19</td>
<td>6</td>
<td>23</td>
</tr>
</tbody>
</table>

HPT = Hyperparathyroidism.

iPTH > 100 pg/ml the specificity and sensitivity remained the same.

Parathyroid enlargement was only detected in patients with hyperparathyroid bone disease (Table 7). There was no correlation between parathyroid gland volume and serum iPTH levels. The specificity of the detection of an enlarged parathyroid gland in the diagnosis of hyperparathyroid bone disease was 100% with a sensitivity of 68%. When combined with a serum iPTH > 100 pg/ml the specificity and sensitivity remained the same.

Discussion

The histological response of bone to the metabolic disorder of chronic renal failure is modified by age, sex, underlying renal pathology, duration of chronic renal failure, exposure to aluminium, medication and mode of dialysis, leading to considerable individual variation in

Assessment of Renal Osteodystrophy

Nephron 1997;75:412-419
type and severity of ROD [24]. An accurate histological diagnosis may be provided by a bone biopsy but, because bone is a dynamic organ, the nature and severity of any pathology may change with time. Regular, repeated bone biopsies are generally an unacceptable method of monitoring ROD due to the degree of discomfort suffered by the patients. The measurement of plasma biochemical markers of bone metabolism are used frequently to discriminate between high and low turnover metabolic bone disease in non-renal patients. However, ROD is more difficult to accurately diagnose using non-invasive methods, particularly in the early stages, because traditional biochemical markers lack specificity and their metabolism may be altered in the presence of uraemia [25].

In this study BAP, total ALP and serum iPTH correlated significantly with histomorphometric measurements of osteoblastic activity. In patients with normal histology, low turnover and mild hyperparathyroid bone disease there was no significant difference in serum BAP, total ALP and iPTH levels. Markers of bone turnover were unable to differentiate the bone pathologies in these patients. It was surprising that the histomorphometric parameters within these subgroups were all within the reference range, despite the clear diagnostic histological differences. A serum iPTH greater than 55 pg/l, the upper reference limit for normocalcaemic non-renal patients, was the most sensitive investigation for the detection of hyperparathyroid bone disease. A mean serum iPTH of 54 pg/l was observed in the mild group, several patients with clear histological disease having 'normal' iPTH values, i.e. <55 pg/l. However, a mean serum calcium of 2.41 mmol/l for this group suggests that there is a failure of PTH suppression within this group and the selection of an isolated value for iPTH without regard to prevailing calcium levels is inappropriate. We do not have enough data in the present study to identify an appropriate index value which would combine calcium and iPTH values. In any event, lowering the limit below 55 pg/l to increase sensitivity of detection of the mild cases proves unacceptable due to the loss of specificity. An elevated mean serum iPTH was also found in several patients in the adynamic, normal and mixed ROD subgroups, reducing the specificity of a raised serum iPTH for the diagnosis of hyperparathyroid bone disease.

The normal reference range for serum BAP is quoted as less than 19 ng/ml, but in this study all but 1 of the patients with low turnover or normal histology had a serum BAP of less than 10 ng/ml. A serum BAP greater than 10 ng/ml gave a sensitivity similar to serum iPTH but with a greater specificity for the diagnosis of hyperparathyroid bone disease where all the subgroups were included.

BMD measurements were made in only 1 patient with adynamic bone disease and this study cannot comment on the diagnostic value of DEXA scanning to distinguish between high and low turnover ROD. Within the hyperparathyroid group there was a negative correlation between increasing severity of hyperparathyroid bone disease and BMD, although the mean values in all three subgroups were within 2 SD of a matched control population. DEXA scanning has little value in the diagnosis of hyperparathyroid bone disease, but it may have the potential to monitor its progress once it has been histologically diagnosed. Confirmation would require a longitudinal study. Increased bone turnover leads to cancellisation of cortical bone leading to a loss of bone volume principally in the appendicular skeleton [24]. The proximal forearm has a higher proportion of cortical bone than the distal forearm [26]. Increasing severity of hyperparathyroidism caused a greater differential loss of BMD in the proximal than the distal forearm, which was statistically significant in the patients with severe hyperparathyroid bone disease. The forearm, which reflects the changes in bone turnover occurring with hyperparathyroidism, can therefore be used for repeat measurements rather than subjecting the patients to total body scanning.

Using a higher cut-off of iPTH of 100 pg/l gave a specificity of 66% and sensitivity of 81%, the loss of sensitivity being attributable to the number of patients with mild bone disease who had normal or only marginally elevated iPTH values.

Parathyroid volume measured at the time of parathyroidectomy has been shown to correlate with the maximal iPTH secretion, the shift in calcium set point and with the duration and severity of renal failure [27]. Advances in ultrasound technology have improved the sensitivity and specificity of detection of the parathyroid glands. In this study the detection of an enlarged parathyroid gland was more sensitive and specific than for the diagnosis of hyperparathyroid bone disease than the radiological changes on a hand X-ray. Ultrasonography would be the appropriate radiological investigation to confirm the diagnosis of hyperparathyroid bone disease in patients with an elevated iPTH.
Conclusions

At present there are no humeral markers or radiological markers capable of totally replacing a bone biopsy in the diagnosis of ROD. The measurement of serum iPTH was the most sensitive test in detecting hyperparathyroid bone disease but was not highly specific particularly when considering early and mild disease. When raised serum iPTH was combined with an elevated BAP or the detection of an enlarged parathyroid gland, the specificity of the diagnosis was increased. The use of a number of non-invasive investigations, combined with sound clinical judgement, may detect metabolic bone disease when present in the majority of patients, although a minority may still require a bone biopsy.

Acknowledgements

This study was funded by MRC Researchgrant number 9123234. We would like to thank the following: Dr. A. M. Davison and Dr. E. J. Will for allowing their patients to be investigated, Dr. R. C. Fowler, Dr. G. J. S. Parkin for the interpretation of the X-rays and ultrasound, Mr. D. A. Purves for performing the BAP assay, Mrs. P. Shaw and Mrs. B. A. Oakley for the preparation of the bone biopsies for histology and Mrs. W. Barney for help with the preparation of the manuscript.

References

Regional bone mineral density after orthotopic liver transplantation
S. Hyder Hussainia, Brian Oldroydb, Sheena P. Stewarta, Fiona Romanb, Michael A. Smithb, Stephen Pollarda, Peter Lodgea, John G. O'Gradya and Monty S. Losowskya

Objectives Although there is a fall in lumbar spine bone mineral density (BMD) after liver transplantation, little is known about femoral neck or total body BMD. Therefore we determined: (a) the proportion of patients with pre-existing hepatic osteopenia before transplantation and (b) the effects of transplantation on global and regional BMD.

Design Retrospective analysis of BMD measurements of patients before and up to 2 years after liver transplantation.

Methods BMD was assessed by dual energy X-ray absorptiometry in 56 patients, before and at regular intervals after liver transplantation, for up to 24 months, to measure total body, lumbar spine (L2–L4) and femoral neck BMDs.

Results Pre-transplant, 23% of patients had osteoporosis (a negative Z score > 2). Paired data before and after transplantation revealed no change in total body BMD. However, there was a fall in lumbar spine BMD (1.04 ± 0.03 to 1.02 ± 0.03 g/cm²; P < 0.04) at 1 month after transplantation. The reduction in lumbar spine BMD was seen up to 12 months, BMD at 18–24 months being similar to pre-transplant values. Femoral neck BMD also fell (0.96 ± 0.06 to 0.83 ± 0.04 g/cm²; P < 0.03), but only after 6–9 months, thereafter remaining below pre-transplant values until the end of the follow-up period.

Conclusions Although osteopenia is common in patients with liver disease, total bone density does not fall after transplantation. Nonetheless, regional lumbar spine and femoral neck bone density does fall after transplantation with a risk period for femoral neck fracture which may extend for up to 2 years. Eur J Gastroenterol Hepatol 11:157–163 © 1999 Lippincott Williams & Wilkins

Introduction Patients with chronic liver disease suffer from bone disease [1–3]. Osteomalacia occurs particularly in patients with cholestatic disease and is secondary to vitamin D deficiency [4,5]. Reduced bone mineral density (BMD), known as osteopenia, is present in up to 40% of patients with chronic liver disease [6,7]. Osteopenia results in a two-fold increase in the rate of axial spinal and appendicular peripheral spontaneous bone fracture in patients with chronic liver disease compared to healthy controls [6].

Orthotopic liver transplantation (OLT) is an established treatment for patients with end-stage chronic liver disease with 1 year survival rates for elective patients of over 90% [8]. In some patients undergoing liver transplantation, pre-existing bone disease may deteriorate due to factors such as prolonged bed rest, immobilization [9] and immunosuppressive therapy [10–12]. Nonetheless, good graft function after liver transplantation might be expected to overcome the osteoporotic effects of the underlying liver disease. The initial studies of bone disease following liver transplantation have found a high rate of atraumatic bone fracture [13–15] and a fall in BMD immediately after transplantation [14,16,17]. However, long-term studies suggest an overall improvement in BMD following transplantation [14,17]. Immunosuppressive regimens utilizing relatively high doses of steroids, especially in the early post-operative period, may in part have been responsible for these high fracture rates post-transplantation [13,17]. Osteopenia in the non-transplant population is a heterogeneous disorder with marked anatomical regional variations in bone density [18–20]. The initial studies of BMD after transplantation focused on the changes of lumbar spine BMD [14,16,17], which consists of predominantly trabecular bone. Only one small series has examined the effects of
OLT on regional bone density [21]. Therefore, in a cohort of patients undergoing OLT for end-stage chronic liver disease we determined the following: (a) the proportion of patients with pre-existing hepatic osteopenia before transplantation and (b) the effects of transplantation on global and regional BMD. Furthermore, we assessed whether duration of hospital stay, dose of steroids, number of rejection episodes, presence of cholestatic liver disease, age and sex were potential risk factors for post-transplant osteopenia.

Methods
Patients
We retrospectively identified all patients who underwent dual energy X-ray absorptiometry (DEXA) scanning before and after liver transplantation between September 1989 and December 1994. Fifty-six patients (33 female) with end-stage chronic liver disease were identified, with a mean age of 51 ± SEM 1.5 y (range 24–69 y). Some of the patients studied have been reported in a separate, but related study concerning the changes in body composition after liver transplantation [22]. Thirty-three patients had cholestatic liver disease, 24 with primary biliary cirrhosis (18 female), seven with primary sclerosing cholangitis (three female), and two with secondary biliary cirrhosis (two female). A total of nine patients had alcohol-related cirrhosis (three female) and five had chronic autoimmune hepatitis (four female). Of the remaining 10 patients, three had cryptogenic cirrhosis (two female), four had neoplastic disease (one female), one patient had hepatitis B cirrhosis and one suffered from alpha-1 antitrypsin deficiency.

All patients were diagnosed on the basis of biochemical, immunological or serological markers, together with liver histology when appropriate. The diagnosis of cryptogenic cirrhosis was made if a liver biopsy demonstrated cirrhosis with no diagnostic features, with the exclusion of excess alcohol intake and following laboratory findings of: (a) no autoantibodies and normal immunoglobulins (to exclude autoimmune liver disease), (b) negative viral serology for hepatitis B and C and, (c) normal studies for iron, copper and alpha-1 antitrypsin. All patients with primary biliary cirrhosis received 100 000 units intramuscular of vitamin D (calciferol) per month.

Transplantation details
Clinical data were available in 43 patients with regard to disease severity before transplantation. Six of the 43 patients were Child–Pugh [23] class A, 33 were class B and 4 were class C. All patients were given triple immunosuppressant therapy consisting of cyclosporin (or tacrolimus in seven patients), azathioprine and prednisolone. In 41 patients details of steroid dosage, duration of hospital stay and number of rejection episodes were available. Patients received a median dose of prednisolone of 20, 12.5, 10, 7.5, and 7.5 mg at time intervals of 1, 2–5, 6–9, 12 and 18–24 months, respectively. The mean duration of hospital stay after transplantation was 32 ± 7 days (range 14–96 days; median 28 days). Ten (25%) patients had an episode of cholestasis after transplantation due to a bile duct stricture or cholestatic hepatitis. Only five (8%) patients experienced an episode of post-transplant hepatitis. One patient who was re-transplanted for recurrent hepatitis B did not receive any prednisolone postoperatively, to minimize the risk of hepatitis B recurrence.

Controls
A group of 329 women (age range 20–81 y) and 115 men (age range 20–78 y) were recruited locally as controls [24]. Subjects with a history of prolonged immobilization, fractures affecting the hip, spine or wrist, medical conditions or on medication known to affect BMD were excluded as control subjects.

Bone mineral density measurements
Total body, lumbar spine (L2–L4) and right femoral neck BMD were measured by dual energy X-ray absorptiometry using a Lunar DPX Bone Densitometer (Lunar Radiation Corporation, Wisconsin, USA). Analysis of the bone scans was made using the manufacturer's software (version 3.6) which calculated bone mineral content (g) and bone area scanned (cm²). Hence BMD was calculated (g/cm²) and Z scores derived subsequently, corrected for age and sex, were used in data analysis. The Z score for BMD is the number of standard deviations above or below the mean BMD for an age and sex-matched normal population. Patients were assessed before transplantation, at 1, 2–5, 6–9, 12 and 18–24 months after transplantation.

Statistical analysis
The number of patients studied at each of these time periods was variable. Thus, bone mineral content was analysed as paired data for each time period compared to pre-transplant values, and tested for statistical significance with the paired t-test. P values of < 0.05 were considered to be statistically significant.

To determine possible risk factors for osteopenia, patients were stratified into those with negative Z scores ≥ 1. The World Health Organization defines osteopenia as a BMD of > 1 SD below the mean for a healthy young population [25]. Since we wished to compare patients with liver disease with controls of the same sex and similar age, we defined osteopenia as BMD > 1 SD below that for an age and sex-matched control population, that is a negative Z score of > 1. Osteoporosis was defined as a negative Z score of > 2. Risk factors for osteopenia, i.e. severity of liver disease, age, gender and cholestatic chronic liver disease, were
compared before and after transplantation with the non-parametric heteroscedastic t-test. Further possible risk factors for osteopenia, namely duration of hospital stay, dose of steroids and numbers of rejection episodes were then compared in those with and without osteopenia after transplantation using the non-parametric analysis. Since multiple analyses, to determine risk factors for bone disease, were performed on the same dataset before and after transplantation, only \( P \) values of \(< 0.01\) were taken as demonstrating a significant difference. The results were expressed as mean values \pm \text{standard error of the mean (SEM)}, unless otherwise stated. Excel software version 5.0 (Microsoft Corporation, 1 Microsoft Way, Redmond, WA 98052-6399) was used to analyse data.

**Ethical considerations**
The study was approved by the Ethics Committee of St James's University Hospital.

**Results**

**Pre-transplant bone mineral density**
The data for total body, lumbar spine and femoral neck BMD for controls and patients in all subjects and stratified for gender are shown in Fig. 1. The total body BMD of \(1.15 \pm 0.01\text{ g/cm}^2\) in controls was significantly greater than the BMD of \(0.62 \pm 0.15\text{ g/cm}^2\) in patients (\( P < 0.001\)). Similar differences were seen for lumbar spine (\(1.16 \pm 0.02\text{ vs } 0.49 \pm 0.15\text{ g/cm}^2; \ P < 0.001\)) and femoral neck (\(0.94 \pm 0.02\text{ vs } 0.52 \pm 0.13\text{ g/cm}^2; \ P < 0.002\)) BMD in controls and patients, respectively. As can be seen from Fig. 1, the pattern of osteopenia was unchanged when stratified for gender, although the difference in BMD was more marked between women and their controls.

The number of patients with osteoporosis defined as a negative \( Z \) score of \(> 2\) or osteopenia defined as a negative \( Z \) score of \(> 1\) is shown in Fig. 2. A total of 13 (23%) of 56 patients studied had osteoporosis, i.e. either total body, lumbar spine or femoral neck negative \( Z \) scores of \(> 2\). However, the numbers of patients with total body, lumbar spine or femoral neck osteoporosis was variable, highlighting the fact that the osteoporosis was rather heterogeneous. Whole body DEXA results detected only an extra two patients to those with osteoporosis at either femoral or lumbar spine sites.

The prevalence of osteopenia was far greater than that of osteoporosis, with 32 (57%) of 56 patients with either total body, lumbar spine or femoral neck negative \( Z \) scores of \(> 1\). No additional patients with osteopenia were detected with whole body DEXA scanning.

**Pre-transplant determinants of osteopenia**
Although total body BMD for patients with cholestatic liver disease was lower (\(1.04 \pm 0.03\text{ g/cm}^2\)) than for patients with non-cholestatic liver disease (\(1.15 \pm 0.02; \ P < 0.002\)), this difference was not statistically significant when corrected for age and sex as \( Z \) scores (\(-0.90 \pm 0.26 \text{ vs } -0.70 \pm 0.96; \ P = 0.07\)). Similarly no significant difference was seen in lumbar spine (\(-0.96 \pm 0.22 \text{ vs } -0.63 \pm 0.19\)) or femoral neck (\(-0.57 \pm 0.30 \text{ vs } -0.32 \pm 0.23\)) \( Z \) scores from patients with cholestatic and non-cholestatic liver disease, respectively.

No significant differences in \( Z \) scores were observed between patients with a Child–Pugh (CP) score of \(< 7\) (range 5–7; median 7) and those with a CP score \(> 7\) (8–13; 9.5)) for: (a) total body (\(-0.52 \pm 0.25 \text{ vs } -0.48 \pm 0.34\)); (b) lumbar spine (\(-0.72 \pm 0.20 \text{ vs } -0.75 \pm 0.29\));
The regional variation in bone density is depicted as the number of patients with osteoporosis (negative Z score of > 2; a) or osteopenia (negative Z score of > 1; b). The histogram shows the number of abnormal Z scores at total body (TB), lumbar spine (LS) or femoral neck (FN) sites; total body alone or combined lumbar spine and femoral neck measurements and as combined total body, lumbar spine or femoral neck measurements.

or (c) femoral neck (−0.63 ± 0.25 vs −0.17 ± 0.47) BMD.

Bone mineral density after liver transplantation
The paired data for BMD after transplantation are given in Fig. 3. The numbers of patients studied at 1, 2–5, 6–9, 12 and 18–24 months were 30, 23, 26, 17 and 11 patients, respectively. Total body BMD fell slightly at 2–5 months from 1.12 ± 0.03 to 1.11 ± 0.03 g/cm², although this just failed to reach statistical significance (P = 0.053). Thereafter total BMD remained stable.

Immediately after transplantation, there was a fall in lumbar spine BMD, which persisted over the next 12 months. However in the 11 patients studied before and 18–24 months after transplantation no difference in BMD was observed. In contrast to lumbar spine BMD, there was no fall in femoral neck BMD immediately after transplantation. Nonetheless, significant reductions in femoral neck BMD were observed after 6–9 months which were sustained at 18–24 months after OLT.

Symptomatic non-traumatic fractures
The patients included in this study did not undergo systematic axial and spinal screening radiology to determine the rate of asymptomatic fractures. However, the rate of symptomatic bone fractures was 11% (six of 56 patients) at 2 years. The details of the fractures with BMD before and at the time of fracture are given in Table 1. Only one patient with primary biliary cirrhosis (PBC) experienced a fracture of the lumbar spine, the remaining five patients sustaining femoral neck fractures. All the patients were female, four with PBC, five over the age of 50 years. Four patients had negative Z scores > 0.5 pre-transplantation and all had a negative Z score > 0.5 at the time of fracture.

Post-transplant determinants of osteopenia
No differences in BMD were found between patients stratified for age, gender, disease severity or disease type. Neither the dose of steroids administered at each
Osteopenia after liver transplantation

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age (y)</th>
<th>Disease</th>
<th>Site of fracture</th>
<th>Time post-transplantation (mo)</th>
<th>Pre-transplantation</th>
<th>Z score at site of fracture</th>
<th>At time of fracture</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>58</td>
<td>PBC</td>
<td>LS</td>
<td>4</td>
<td>-0.56</td>
<td>-0.56</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>44</td>
<td>CC</td>
<td>FN</td>
<td>12</td>
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<td>0.40</td>
<td>-0.77</td>
</tr>
<tr>
<td>3</td>
<td>56</td>
<td>PBC</td>
<td>FN</td>
<td>24</td>
<td>-2.12</td>
<td>-2.12</td>
<td>-1.98</td>
</tr>
<tr>
<td>4</td>
<td>52</td>
<td>PBC</td>
<td>FN</td>
<td>2</td>
<td>-0.88</td>
<td>-0.88</td>
<td>-1.82</td>
</tr>
<tr>
<td>5</td>
<td>59</td>
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<td>FN</td>
<td>8</td>
<td>0.20</td>
<td>0.20</td>
<td>-0.98</td>
</tr>
<tr>
<td>6</td>
<td>59</td>
<td>PBC</td>
<td>FN</td>
<td>9</td>
<td>-2.86</td>
<td>-2.86</td>
<td>-2.86</td>
</tr>
</tbody>
</table>

Demographic details of the six female patients who experienced asymptomatic atraumatic bone fractures after liver transplantation at either femoral neck (FN) or lumbar spine (LS) sites with Z scores corresponding to the site of fracture before transplantation and at a time period nearest to the time that the bone fracture occurred. PBC, primary biliary cirrhosis; ALD, alcohol-related liver disease; CC, cryptogenic cirrhosis.

**Discussion**

Our results confirm that there is a loss of BMD after liver transplantation [14,16,17,26]. However, we have reported the effects of transplantation on both trabecular and cortical bone, rather than trabecular bone alone [14,16,17,26]. We consider this to be important since osteopenia is a heterogeneous disorder with variable bone density depending on the site examined [21].

The paired data did show a significant, though small, fall in lumbar spine bone density, confirming the findings of other groups [14,16,17,26]. Thereafter there was no improvement over pre-transplant values in lumbar spine BMD up to 24 months post-transplantation. In contrast to our data, Eastell et al. [17] found that lumbar spine BMD returned to the pre-operative values by about 12 months and was convincingly greater than pre-transplantation BMD by 2 years. The delay in recovery of lumbar bone density that we found confirms the observations of Arnold et al. [14], who documented an early fall in BMD within 3–6 months, which thereafter remained unchanged in the majority of patients up to 24 months after liver transplantation. However, long-term studies of bone density are required to determine the degree of reversibility of post-transplant osteopenia, especially in transplant programmes which use long-term corticosteroid therapy.

The timing of the changes bone density in the femoral neck was different to that in the lumbar spine, with a fall in femoral BMD after transplantation, confirming regional differences in another, smaller, study which examined both femoral and lumbar BMD after transplantation [21]. A possible explanation for the discordance in the fall of femoral and lumbar spine BMD may have been the large time intervals used in the early time periods. Thus a subtle fall in femoral BMD in the 2-5 month period may have been missed. Nonetheless, using this time scale we were able to demonstrate a significant fall in lumbar spine BMD in the early post-transplant period. Femoral bone density 2 years after OLT was still significantly lower than pre-transplant bone density. Thus sixty-four patients may extend beyond 1 year and perhaps still exists 2 years following first transplantation.

The regional variation in osteopenia before transplantation, observed in the current study, has been documented previously in studies of bone loss in patients with osteoporosis [18–20]. Lumbar spine is predominantly made up of trabecular bone, whilst proximal femur consists of mainly cortical bone. Trabecular bone has a higher turnover rate compared to cortical bone [27] which may account for greater variation in bone loss.

The regional loss of bone density without a total BMD has been demonstrated previously in renal transplantation, but has not been reported following liver transplantation. Following transplantation these changes have been attributed to differing regional skeletal responses to immobilization [29,30] or a redistribution of bone content from trabecular to cortical bone sites [30]. The findings of an initial fall of lumbar spine bone density and subsequent fall in femoral neck bone density little change in total body BMD support the hypothesis that there is a redistribution rather than absolute bone loss following transplantation. However, as is practice, the measurement of total BMD, in a cortical and trabecular femoral neck and lumbar spine BMD, is included in this model.
were detected before transplantation and no overall changes in BMD were seen after transplantation.

The regional variations in bone density shown in the current study, together with the definitions used for osteopenia and osteoporosis, demonstrate the difficulty in defining the exact prevalence of bone disease patients with chronic liver disorders. In the current series, 23% of our patients had osteoporosis and 37% were osteopenic, although the exact figures would vary, depending on which region of the body was studied. Although these prevalence rates are comparable to earlier data [6,7], we would suggest that both lumbar spine and femoral neck BMD need to be assessed in order to accurately determine the prevalence of osteopenia in patients with chronic liver disease.

The small numbers of patients with cholestatic liver disease in our study, together with the fact that these patients received early transplantation and vitamin D supplementation prior to transplantation, may account for the absence of a relationship between cholestatic disease and bone density found by some [31], but not all, authors [17,32]. Furthermore, the short duration of hospital stay and relatively low dose of steroids administered were probably responsible for these factors not being implicated in the development of post-transplant bone disease. The median CP scores for the group of patients with mild (7) and those with moderate severity (9.5) liver disease were similar, with only four patients with class C disease. This skew in the distribution of disease severity towards milder disease may account for the similar BMD in patients with mild or moderate severity chronic liver disease before transplantation.

Since patients were not investigated systematically for evidence of asymptomatic fractures, we believe that the low fracture rate of 11% in the current series underestimates the true asymptomatic fracture rate. Others have reported a fracture rate of 20–65% in the first year of transplantation [13,15,17,33]. However, an alternative explanation for the low fracture rate is the relatively small magnitude of the reduction in BMD after transplantation. The quantitative fall in lumbar spine BMD was lower than some reports [13,15,17], although similar to others [14]. Moreover a recent study [34] reported no reduction in bone density in 82 patients undergoing transplantation. We speculate that the reason for the diminished reduction in BMD after transplantation, compared to a decade ago, is a consequence of earlier liver transplantation, shorter hospital stays and the use of steroids in lower dosages (with steroid withdrawal in some programmes).

In conclusion, although osteopenia is common in patients with chronic liver disease, total bone density does not fall after transplantation. Nonetheless regional lumbar spine and femoral neck bone density does fall after transplantation with a risk period for femoral neck fracture which may extend for up to 2 years. However, the magnitude of the fall in bone density after transplantation has declined over the last decade. This suggests that the prevalence rate of asymptomatic fracture should be reassessed with the use of modern transplantation techniques.

References
Osteopenia after liver transplantation

The comparison of neutron activation analysis and photon absorptiometry at the same part-body site

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Abstract. Two non-invasive methods of bone measurement were, for the first time, compared at the same part-body site. Bone calcium, using in vivo neutron activation analysis, and bone mineral, using photon absorptiometry, were measured in the forearm in a group of renal patients. A highly significant correlation was obtained from absolute values \( r = 0.91 \) which was reduced after normalisation \( r = 0.75 \). Percentage changes, over periods totalling 31.3 patient years, were also highly significantly correlated \( r = 0.61, P < 0.001 \). The precision of the techniques accounts for the spread of the bivariate data and the estimated line of the true relationship was not significantly different from the line described if the percentage changes measured by the two techniques were equal. The two basic techniques were shown to be equally sensitive in vivo methods for monitoring calcified bone in patients suffering from metabolic bone disease.

1. Introduction

In metabolic bone disorders such as renal osteodystrophy, osteomalacia and osteoporosis, the loss of calcified bone will increase the tendency to fracture. It is therefore necessary to establish a means of quantifying any such bone loss, both for the routine management of patients and also to evaluate possible therapeutic regimes.

The two most precise non-invasive techniques of investigating the bone status are neutron activation analysis (NAA) and photon absorptiometry. NAA has been used clinically to measure both whole-body calcium and part-body calcium whilst photon absorptiometry has been used principally to measure the bone mineral content (BMC) of the forearm. The two techniques measure similar, though not identical, parameters. NAA measures purely calcium; photon absorptiometry, on the other hand, reflects total mineral content, usually in a small volume of bone.

Comparisons have been made in clinical studies between absolute values, normalised values (a correction being made to eliminate the variation in patient size) and fractional changes over a time period using the two techniques. Highly significant correlations have been found between absolute photon absorptiometry and both whole-body and part-body NAA (Chesnut et al 1973, Cohn et al 1974, 1975, 1976, Harrison et al 1974, Manzke et al 1975, Naik et al 1977, Zanzi et al 1978, Maziere et al 1979). As both techniques give values that are dependent on the size of the patient, these correlations are to be expected. The correlation coefficients were reduced when normalisation procedures were adopted in an attempt to correct for the size of the patient (Cohn et al 1974, 1975, Manzke et al 1975). The significant correlations that exist after normalisation may reflect the inadequacies of the normalisation procedures rather than demonstrating any relationship between the techniques themselves.
The most important comparison is therefore between changes measured over a period by the two methods. All such comparisons reported so far have failed to demonstrate any significant correlation (Harrison et al 1974, Aloia et al 1975, Cohn et al 1975, 1976, Dubek et al 1977) though 'reasonable agreement' was found in one study (Harrison et al 1974). It must be noted that different body sites were used for the two techniques so the lack of correlation could therefore reflect the different rates of change in bone at different sites in the body rather than differences between the two methods.

This study seeks to obtain a valid comparison of the two methods by comparing absolute, normalised and percentage changes measured by photon absorptiometry and NAA at the same part-body site, measurements of the forearm being correlated in a long-term clinical study.

2. Methods and patients

The main purpose of our research work has been the development and evaluation of NAA to measure forearm calcium. Care was taken in the design and construction of the apparatus to obtain a high level of precision. Photon absorptiometric measurements however were performed using available apparatus which had been constructed many years earlier in our workshop with the aim of monitoring bone mineral content in renal patients.

2.1. Part-body NAA

The calcium in the forearm was activated using two $^{252}$Cf neutron sources positioned one either side of the forearm. The induced $^{49}$Ca activity was then measured by two 15 cm × 10 cm NaI detectors, also in a bilateral arrangement (Smith and MacPherson 1977, Smith and Tothill 1979). The irradiation apparatus had initially been designed to measure only changes in forearm calcium. Three repeated measurements were performed at each visit producing a precision of the calcium measurement of 1.8%. At the final visit an additional measurement was performed using the apparatus which was more suitable for absolute calcium measurements. Immersion of the forearm in a water bath (Smith and Tothill 1979) eliminated the variation in the sensitivity of activation due to soft tissue around the bone. The precision of a single measurement of absolute forearm calcium was 2.6%. To normalise for body stature the absolute calcium values were divided by the cube of the patient's height, this factor giving the smallest spread of the data from a series of normal forearm measurements (Tothill et al 1979).

The bilateral geometry effectively eliminated errors due to minor patient movement. The design of the apparatus, with the arm always held rigidly in the same arm rest, eliminated any subjective choice of the measurement site and hence inter-operator error. The efficiency of activation and detection are not uniform along the length of the forearm, the combination approximating to a normal distribution of full width at half maximum of 14 cm centred 19 cm from the centre of the fist. It is therefore not appropriate to express the results in grams of calcium. The measurements are 'absolute' in the sense that the $^{49}$Ca counts recorded depend only on the mass of calcium and its distribution along the sensitive volume, and not on such factors as the thickness or composition of overlying tissue. The results are expressed as 'calcium counts', corrections being derived for such factors as source decay by measurements on a calcium standard.
2.2. Photon absorptiometry

The bone mineral content of the radius was measured using the techniques described by Cameron and Sorenson (1963). Calipers were used to define the measurement site 5 cm from the styloid process after which the forearm was positioned horizontally with the inside of the forearm uppermost in a water bath. A 1.665 GBq (45 mCi) $^{241}$Am source and NaI detector with a collimation of 0.5 cm $\times$ 1.2 cm, scanned across the radius, counting for 5 s every millimetre (approximate count rate, $6 \times 10^3$ cps). Three repeated scans across the radius were performed and the data were stored on punched tape for computer analysis. The precision of the BMC measurement on normal volunteers was approximately 3.5%. The width of the radius, obtained by plotting the BMC values across the forearm, was used as the normalisation factor.

It is recognised that neither the site nor the source were ideal for BMC measurements. The former was used to be consistent with previous studies whilst the latter was used for financial reasons. Our method of photon absorptiometry was more prone to patient movement and repositioning errors than NAA; in addition, varying amounts of overlying fat would affect the BMC measurements.

Only about 10% of the bone at the 5 cm site is trabecular (Melsen et al. 1979) so both NAA and photon absorptiometry are measuring predominantly cortical bone.

2.3. Patient group

Nineteen renal patients undergoing chronic haemodialysis at home were followed for 22 months in a study to evaluate the effect of dialysate calcium levels and treatment with 1α-hydroxycholecalciferol. The same NAA apparatus was used to measure the forearm calcium at times 0, 8 and 22 months to determine the percentage changes over the first 8-month and then subsequent 14-month period. Changes in bone mineral content by photon absorptiometry were also measured over the same periods. Of the 19 patients at the start of the study, 18 were measured at eight months and the surviving 16 at 22 months. In addition absolute forearm calcium was measured at 22 months.

Absolute and normalised part body calcium and bone mineral content were correlated in 16 patients and percentage changes were correlated from 34 sets of data totalling 31.3 patient years. Regression analysis of the second kind (Documenta Geigy 1970) was performed on the data and $P > 0.05$ was taken to be non-significant. Having performed the regression analysis on the percentage changes measured by NAA and photon absorptiometry a $\chi^2$ value was calculated which established whether the known errors of the two methods could account for any observed spread of the bivariate data about a straight line (see Appendix 1).

3. Results

The results of the absolute measurements are displayed in figure 1 and the results of the percentage changes in figure 2. As it is the percentage difference between the two sets of measurements which is plotted in figure 2, the error becomes the precision $\times \sqrt{2}$, i.e. 5.0% and 2.6% for BMC and NAA respectively. The estimated line of the true relationship (see Appendix 1) is displayed. This would only correspond to the major axis of the ellipse (figure 2) if the precisions of the two techniques were identical. Tolerance ellipses ($P = 0.05$) are displayed with the data and the results of the regression analysis are given in table 1. The result of the $\chi^2$ analysis of the data in figure 2 was $\chi^2 = 36.2$ ($P = 0.30$).
Figure 1. Correlation of absolute bone mineral content (BMC) measured by photon absorptiometry against forearm calcium measured by NAA.

Figure 2. Correlation of percentage changes in bone mineral content (BMC) by photon absorptiometry against percentage changes in forearm calcium by NAA. Estimated line of true relationship is displayed.

<table>
<thead>
<tr>
<th></th>
<th>n</th>
<th>r</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Absolute values</td>
<td>16</td>
<td>0.91</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Normalised values</td>
<td>14</td>
<td>0.75</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Percent changes</td>
<td>34</td>
<td>0.61</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

n = number of sets of patient measurements.
4. Discussion

This group of renal patients proved to be an ideal one to correlate NAA with photon absorptiometry because not only did they have a wide variation in absolute NAA values— but also some large percentage changes over the period of the study. The NAA results are supported by clinical findings; a highly significant inverse correlation \( r = -0.84, P < 0.001 \) was found between the normalised forearm calcium and the duration of dialysis and the patients receiving 1α-hydroxycholecalciferol were shown to increase significantly their forearm calcium during the study (Winney et al 1977, Smith et al 1979, 1981).

The correlation coefficient \( r = 0.91 \) for absolute values falling to \( r = 0.75 \) after normalisation agrees very well with the values found in comparisons between whole body NAA and forearm photon absorptiometry (Cohn et al 1975, Manzke et al 1975) and is better than values obtained by Maziere et al (1979) and Harrison et al (1974) who compared forearm photon absorptiometry with part-body NAA of the hand and trunk respectively. The decrease in the correlation coefficient from 0.91 to 0.75 after normalisation suggests that the high correlation between absolute values is due to a large extent to the size of the patient. As neither of the normalisation factors are perfect, the true correlation is probably lower still. This inability to eliminate completely the effect of stature means that the most important information can only be obtained by comparing the changes measured by the two techniques.

There are two important questions that need to be answered from the analysis of the percentage changes. Firstly, are changes in calcified bone measured by NAA and photon absorptiometry related and, secondly, if this is so, which is the more sensitive method.

This study is the first to show a highly significant correlation \( \rho = 0.61, P < 0.001 \) between changes measured by NAA and photon absorptiometry. In addition the \( \chi^2 \) value obtained showed that the errors of the techniques could account for the spread of the bivariate data. This is further demonstrated by the fact that 80% of the data points in figure 2 are within one standard deviation of the estimated line of the true relationship, suggesting that the scatter about the line is due to the known measurement errors. Consequently, if a perfect correlation between the methods did exist, the data would not be distributed along a straight line but in an ellipse of comparable width to the one obtained in this study.

Information about which of the two methods is the more sensitive is obtained from the estimated line of the true relationship of the data (figure 2). This line crosses the axes extremely close to the origin and the slope of the line appears to suggest that NAA is the more sensitive, as would be expected due to the better precision of the method. However, the 95% confidence interval of the line contains the line \( Y = X \) so the line through the origin with slope unity is not significantly different from the estimated line of the true relationship of the data.

The discussion must now be extended to consider how the results from our specific methods of NAA and photon absorptiometry can be used to draw more general conclusions about the two techniques. In our hands NAA has a better precision, although figures of 1% have been quoted for some photon absorptiometry apparatus (Christiansen and Rodbro 1977). If, in figure 2, the BMC had the same precision as NAA then the line of relationship would be the major axis of the ellipse, which has a slope of 0.95. Thus NAA and photon absorptiometry, generally speaking, are equally sensitive in vivo techniques for monitoring bone status in the forearm in patients suffering from metabolic bone disease. An implication of the findings is that the lack of correlation of
changes found by others is due to real physiological differences between parts of the body and not to the use of alternative measurement techniques.

**Appendix 1.**

The equation to calculate \( \chi^2 \) to test whether the specified error variances of the NAA and photon absorptiometry measurements are large enough to account for the observed spread of the bivariate data about a straight line (Sprent 1966) is as follows:

Let

\[
(S_x)^2 = \text{error variance of } x\text{-axis}
\]

where

\[ S_x = \text{precision of NAA measurement } \times \sqrt{2}. \]

Similarly for the \( y \)-axis for photon absorptiometry.

Then

\[
\chi^2 = \frac{S_{xy} - 2bS_{xy} + b^2S_{xx}}{(S_x)^2 + b^2(S_y)^2} (n - 1 \text{ degrees of freedom})
\]

where

\[
b = \frac{S_{xy} - \lambda S_{xx} + [(S_{xy} - \lambda S_{xx})^2 + 4\lambda S_{yy}^2]^{1/2}}{2S_{xy}}
\]

and

\[
\lambda = \frac{(S_x/S_y)^2}{S_{xy}}
\]

\[ S_{xx} = \sum (x_i - \bar{x})^2, \quad S_{yy} = \sum (y_i - \bar{y})^2 \]

\[ S_{xy} = \sum (x_i - \bar{x})(y_i - \bar{y}). \]

\( b \) is the slope of the estimated line of the true relationship between \( x \) and \( y \).

**Acknowledgments**

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**Résumé**

Comparaison en une même partie du corps de l'analyse par activation neutronique et de la mesure par absorption des photons.

Pour la première fois, une étude comparative de deux méthodes non-invasives de mesure de l'os ont été comparées, en une même partie du corps. Nous avons mesuré, au niveau de l'avant-bras, dans un groupe de patients ayant une atteinte rénale, le calcium osseux, en utilisant l'analyse par activation neutronique in vivo et l'os minéral, en utilisant la mesure par absorption des photons. Nous avons obtenu une correlation hautement significative pour les valeurs absolues \((r = 0.91)\), celle-ci diminue après normalisation \((r = 0.75)\). Les changements en pourcentage, sur une période totale de 31,3 années-malades, sont également corrélés de manière hautement significative \((r = 0.61), P < 0.001\). La distribution de ces données bivariates est
Activation analysis and absorptiometry correlation

explained by the precision of techniques. The line of regression estimated to part of the true values shows no difference significantly from that obtained by supposing that the changes in percentage measured by the two techniques are equal. We have shown that these two techniques of base, of surveillance in vivo of bone, even if patients suffering from maladies osseuses metaboliques, have a similar precision.

Zusammenfassung

Ein Vergleich zwischen Neutronenaktivierungsanalyse und Photonenabsorptiometrie bei gleicher Teilkörperlage.

Zwei nicht-invasive Methoden zur Messung am Knochen werden bei gleicher Teilkörperlage verglichen. Mit der in vivo Neutronenaktivierungsanalyse wurde das Kalkzium im Knochen, mit der Photonenabsorptiometrie der Mineralgehalt des Knochens im Vorderarm von Nierenpatienten gemessen. Aus den Absolutwerten ($r = 0.91$) erhielt man eine hochsignifikante Korrelation, die nach der Normalisierung ($r = 0.75$) reduziert wurde. Prozentuale Änderungen waren ebenfalls hochsignifikant korreliert ($r = 0.61$, $P = 0.001$). Die Genauigkeit, die man erhält durch das Spreizen der zweidimensionalen Werte und die geschätzte Linie der wahren Beziehung, war nicht sehr verschieden von der Linie, wenn die gemessenen prozentualen Änderungen bei beiden Techniken gleich waren. Es wird gezeigt, daß beide Techniken gleich empfindliche in vivo Methoden zur Darstellung verkalkter Knochen sind bei Patienten, die an einer Stoffwechselkrankheit leiden.

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Sir,—Your recent leading article (4 April, p 1096) draws conclusions with which we cannot entirely agree. Certainly patients who undergo surgery for peptic ulcer are at increased risk of dying prematurely. This was the conclusion in our article to which you refer, and was re-emphasised in an extended study (on 779 males aged 30-59 years operated on from 1947 to 1965) recently presented to the British Society of Gastroenterology. Over a 15-32-year follow-up period highly significant increases in mortality occurred in all age groups. In both studies we found that the major contribution to this excess mortality was provided by diseases which have close associations with cigarette smoking (carcinoma of the lung and oesophagus, ischaemic heart disease, and chronic bronchitis and emphysema). In our experience gastric carcinoma was no more common than predicted for an age-matched Scottish population. Eighty-four per cent of the patients studied were smokers prior to operation and this proportion of smokers remains unchanged among our patients now undergoing highly selective vagotomy.

Surgically treated ulcer patients as a group are heavy cigarette smokers. The crucial question therefore is not whether 5% or more of the survivors of the "gastrectomy bonanza" will subsequently develop carcinoma of the stomach but whether patients coming to surgery can be persuaded to cease smoking. Otherwise the trend towards premature death will not diminish with the abandonment of gastrectomy and patients undergoing highly selective vagotomy will die as prematurely as their gastrectomised predecessors.

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LATE MORTALITY AFTER SURGERY FOR PEPTIC ULCER

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Abstract

To examine the claim that life expectancy is reduced after curative peptic-ulcer surgery, we studied mortality and its causes in 779 men with peptic ulcer treated surgically between 1947 and 1965. The minimum follow-up period of survivors was 15 years.

There was an excess mortality in the three major age groups (30 to 39, 40 to 49, and 50 to 59 years at operation), as compared with the general population (P<0.001), with a mean shift of 9.1 years in the survival curve. Surgery was not the direct cause of death. In particular, the eight deaths from carcinoma of the stomach were no more than expected. Excess mortality was due to smoking-associated disease, which accounted for 200 deaths. Eighty-three per cent of the patients were cigarette smokers.

We conclude that substituting highly selective vagotomy for gastrectomy will not lead to improved survival, because 80 per cent of patients now having operations for ulcer are heavy cigarette smokers. (N Engl J Med. 1982; 307:519-522.)

The long-term survival of patients who undergo curative surgery for peptic ulcer has attracted relatively little recent attention.1 In 1958 Krause showed that patients tended to die prematurely in the years after Polya gastrectomy;2 the major contributors to this excess mortality were tuberculosis, carcinoma of the gastric remnant, suicide, and alcoholism. In the same year Balint confirmed that the number of deaths from pulmonary tuberculosis was significantly increased after gastric surgery; this report also added ischemic heart disease and carcinoma of the lung to the list.3 Other diseases since linked with gastric surgery include carcinoma of the esophagus,4 certain neurologic disorders,5 and gallstones.6

The general increase in life expectancy since the reports of Krause and Balint, together with earlier claims that patients who had undergone surgery for ulcer were bad risks in life-insurance terms,7 prompted us to review the mortality trends in men undergoing curative peptic-ulcer surgery.

The aims of our study were twofold: to determine whether there was still a tendency for patients to die prematurely after a successful peptic-ulcer operation, and if so, to determine which factors contributed to the excess mortality.

Methods

The records of all men between 30 and 59 years old who underwent peptic-ulcer surgery in the Western General Hospital between 1947 and 1965 were reviewed. These records are complete at the time of surgery and updated at regular intervals thereafter by standardized questionnaires or personal interviews.

Three categories of patients were excluded from the analysis: those who died within one year of surgery, those who underwent simple closure of a perforated ulcer, and those in whom peptic ulcer subsequently recurred.

The patient's age at operation, the date of operation, the type of operation, the smoking status at operation, the age at death, and the cause of death were recorded. The cause of death in patients who died at home during the study period was taken as the underlying cause registered on the death certificate.

Of the 898 patients in the series, 77 (9 per cent) were excluded either because their records were incomplete or because they were lost to follow-up by the end of 1979. The study population therefore comprises 779 patients. By the end of 1979, 360 patients had died.

Six hundred seventy-six (86 per cent) of the 779 patients were treated by gastrectomy, and 101 (13 per cent) by drainage with or without vagotomy.

Method of Analysis

Actuarial survival curves were constructed to assess the mortality rate for each of three 10-year age groups: 30 to 39, 40 to 49, and 50 to 59 years at operation. These curves were compared with curves of expected survival calculated for the general population from life-table figures obtained from the Faculty of Actuaries of Edinburgh and appropriate to the period and the age groups under study. All patients were included in the calculation of the percentage surviving at 5, 10, and 15 years after surgery. In order to calculate the percentage surviving 20 and 25 years after surgery, subgroups of patients who underwent surgery before 1955 were used. The numbers of patients in the subgroups from 30 to 39, 40 to 49, and 50 to 59 years old were 83, 157, and 132, respectively.

The expected numbers of deaths from individual diseases were obtained from figures published by the Registrar General for Scotland (RGS).6 For this purpose, the study group was not divided into the three groups used for the survival analysis. Two methods of calculating the expected number of deaths were used. When available from the RGS statistics, the death rate (expressed as the number of deaths per year per 100,000 population) was used. Data were analyzed in three five-year periods from 1963 (when input into the study group had ceased and the size of the group was 701 patients) until 1980. The age spectrum of the study group for each such period was calculated in 10-year intervals and combined with data from the RGS for the same age decades. The RGS data were used those for the middle year of each period (1967, 1972, and 1977). The three causes of death analyzed by this method were ischemic heart disease, chronic bronchitis and emphysema, and carcinoma of the lung.

For the causes of death for which the death rate per 100,000 population was not published, a second method was used. The study group was split into five periods from 1955 to 1980. As before, the RGS figures used were those for the middle year of each period, considered in 10-year age intervals. For each death, the number of deaths per year recorded in the RGS figures was expressed as a percentage of the total number of deaths from all causes in the same age group. This was then combined with the total number of deaths in the corresponding age group of the study population for the particular five-year period, resulting in the expected number of deaths from a particular cause. This method of analysis alone was applied to six causes of death: carcinoma of the stomach, carcinoma of the pancreas, carcinoma of the esophagus, carcinoma of the colon and rectum, suicide, and cirrhosis of the liver. It was also employed for ischemic heart disease, chronic bronchitis, and carcinoma of the lung for the period from 1955 to 1965, when death rates expressed as deaths per year per 100,000 population were not published. Both methods of analysis took account of variations in causes of death with age and time.

When appropriate, the chi-square test was used to assess significance. When the expected number of deaths was less than five and the chi-square method was consequently invalid, the probability was calculated directly from the equation of the binomial distribution.8
RESULTS

When the life-table survival curves for each of the three age groups studied are compared with the survival curves predicted for the general population, it is seen that the study population had a significant increase in mortality rate (Fig. 1) (P<0.001 in each age group). Figure 2 shows the death rate per year for each age group at operation over the first 15 years after the operation. The curve for the study group (which includes a figure for 20 to 29-year-old patients who underwent surgery for peptic ulcer during the same period) is shifted to the left, indicating premature mortality; the mean shift of the curve is 9.1±0.1 years.

Table 1 shows the number of deaths from individual diseases by age group during the study.

Deaths from diseases that have been identified as having close associations with smoking,210 diseases that have been related to the sequelae of gastric surgery,2 and other diseases with significant variations from the predicted values are detailed in Table 2.

Eighty-three per cent of the patients were cigarette smokers at the time of operation.

There were eight deaths from carcinoma of the stomach; the average time from gastrectomy to death from carcinoma was 18.9 years.

There were significant increases in the numbers of deaths from cirrhosis of the liver, suicide, carcinoma of the colon and rectum, and carcinoma of the pancreas.

DISCUSSION

Since Krause2 first identified the increased risk of premature death in the years after peptic-ulcer surgery, this finding has been confirmed by others.3,11,13

Earlier, the nature of the link between the increased incidences of ischemic heart disease, of tuberculosis,13 of carcinoma of the lung, esophagus, or stomach, of suicide, and of alcoholism after peptic-ulcer surgery was not always clear. However, the growing realization of the part played by smoking in the causation of a number of these diseases clarifies the relation of gastric surgery to shortening life expectancy.10,14,15

Our study confirms a tendency for patients who have undergone a curative ulcer operation to die prematurely in the years after the operation. The shortening of life expectancy was approximately nine years—a figure similar to that calculated several years ago with some of the population data presented here.11

The trend toward premature death occurred in each age group at operation that we studied.

Thus, persons who have had surgery for ulcer are indeed bad risks in life-insurance terms. The question is why.

Of the four causes originally shown to be related to gastrectomy, pulmonary tuberculosis has now been virtually eliminated. We could attribute only two deaths to this disease.

An increased incidence of gastric carcinoma after gastrectomy has been reported,16-18 but we found that it was no more common than predicted for an age-matched Scottish population. Stalsberg19 suggested...
that the increase in incidence occurs only 15 years or more after surgery; before then, the incidence may be decreased.20 The average interval between gastrectomy and death from gastric carcinoma was 18.9 years in our study, so that insufficient time may have elapsed. If a subsequent increase does occur, its contribution to the overall excess mortality will be small.

Thus, of Krause's original four contributors, only alcoholism (as measured by liver cirrhosis) and suicide have in our experience contributed to the excess mortality. But numerically the excess was small. The major contribution was from the diseases that are closely associated with smoking. In men these diseases are identified as carcinoma of the lung and other respiratory sites, carcinoma of the esophagus, tuberculosis, chronic bronchitis and emphysema, pulmonary heart disease, aortic aneurysm, hernia, and ischemic heart disease.10 In our study tuberculosis, hernia, and aortic aneurysm accounted for only three deaths. Overall, deaths due to the other five diseases were significantly more common than predicted for an age-matched Scottish population (P = 0.025), although individually these increases over the predicted values did not reach statistical significance. In particular, the number of deaths from ischemic heart disease, though increased, was not significantly different from the predicted value. Doll and Peto have shown that in men the link between ischemic heart disease and smoking is strongest under the age of 45.10 In patients over this age, the connection with smoking becomes weaker. Since the majority of our patients had the operation in their 40s, this may explain the insignificance of the rise over the predicted values.

Confirming the findings of others,19,21 we found no significant increase in carcinoma of the esophagus. Although suicide and alcoholism are both traditionally regarded as potential sequelae of gastric surgery, both have established connections with smoking status. Possibly, they are related to the patient's psychological background as well as the need for peptic-ulcer surgery.

Carcinomas of the colorectum and of the pancreas were significantly increased in our study population, and these are also related to smoking status, although in the case of colorectal carcinoma the association is limited to carcinoma of the rectum.10 No connection between these diseases and gastric surgery has been described, to our knowledge.

Thus, we found that diseases known to be related to smoking contributed more toward mortality than diseases traditionally linked with the sequelae or gastric surgery.

Eighty-three per cent of our patients were habitual cigarette smokers at the time of operation, and few ceased to smoke after surgery. The trend continues into the 1980s. Eighty per cent of our male patients presently undergoing highly selective vagotomy for chronic peptic ulceration are cigarette smokers, and this is occurring in a population in which the proportion of male smokers is diminishing. Clearly, the men who come to surgery for duodenal ulcer represent a select group.

Table 1. Causes of Death According to Age Group.*

<table>
<thead>
<tr>
<th>Cause of Death</th>
<th>Age Group at Death (Yr)</th>
<th>No. of Deaths</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total 30-39 40-49 50-59 60-69 70-79 &gt;80</td>
<td></td>
</tr>
<tr>
<td>Ischemic heart disease</td>
<td>128 - 4 - 36 - 19 - 3</td>
<td></td>
</tr>
<tr>
<td>Carcinoma of the lung</td>
<td>47 - 1 - 8 - 28 - 10</td>
<td></td>
</tr>
<tr>
<td>Cerebrovascular accident</td>
<td>29 - 2 - 5 - 13 - 9</td>
<td></td>
</tr>
<tr>
<td>Chronic bronchitis and emphysema</td>
<td>22 - 4 - 14 - 4</td>
<td></td>
</tr>
<tr>
<td>Bronchopneumonia</td>
<td>21 - 5 - 7 - 6 - 3</td>
<td></td>
</tr>
<tr>
<td>Carcinoma of the colon and rectum</td>
<td>16 - 1 - 2 - 11 - 2</td>
<td></td>
</tr>
<tr>
<td>Carcinoma of the pancreas</td>
<td>11 - 3 - 2 - 5 - 1</td>
<td></td>
</tr>
<tr>
<td>Suicide</td>
<td>10 - 4 - 3 - 12</td>
<td></td>
</tr>
<tr>
<td>Carcinoma of the stomach</td>
<td>8 - 1 - 4 - 1</td>
<td></td>
</tr>
<tr>
<td>Accident</td>
<td>8 - 2 - 4</td>
<td></td>
</tr>
<tr>
<td>Hepatic cirrhosis</td>
<td>7 - 2 - 3</td>
<td></td>
</tr>
<tr>
<td>Peripheral vascular disease</td>
<td>6 - 2 - 2</td>
<td></td>
</tr>
<tr>
<td>Carcinoma of the prostate</td>
<td>5 - 3</td>
<td></td>
</tr>
<tr>
<td>Pulmonary embolism</td>
<td>5 - 2</td>
<td></td>
</tr>
<tr>
<td>Other malignant diseases</td>
<td>21 - 12 - 5 - 3 - 1</td>
<td></td>
</tr>
<tr>
<td>Other gastrointestinal diseases</td>
<td>4 - 1 - 2 - 1</td>
<td></td>
</tr>
<tr>
<td>Other respiratory diseases</td>
<td>3 - 2 - 1</td>
<td></td>
</tr>
<tr>
<td>Other vascular diseases</td>
<td>3 - 1</td>
<td></td>
</tr>
<tr>
<td>Miscellaneous</td>
<td>6 - 1</td>
<td></td>
</tr>
</tbody>
</table>

* Diseases that caused five or more deaths in the series are listed individually.

Table 2. Actual and Predicted Numbers of Deaths from Individual Diseases, 1955-1979.*

<table>
<thead>
<tr>
<th>Cause of Death</th>
<th>No. of Deaths</th>
<th>P Value</th>
<th>Predicted</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ischemic heart disease and pulmonary heart disease</td>
<td>128 - 112.5</td>
<td>NS</td>
<td></td>
</tr>
<tr>
<td>Chronic bronchitis and emphysema</td>
<td>22 - 19.9</td>
<td>NS</td>
<td></td>
</tr>
<tr>
<td>Carcinoma of the lung and other respiratory sites</td>
<td>47 - 35.7</td>
<td>0.05</td>
<td></td>
</tr>
<tr>
<td>Carcinoma of the esophagus</td>
<td>3 - 2.7</td>
<td>NS</td>
<td></td>
</tr>
<tr>
<td>All smoking-associated diseases</td>
<td>200 - 170.8</td>
<td>&lt;0.025</td>
<td></td>
</tr>
<tr>
<td>Carcinoma of the stomach</td>
<td>3 - 10.4</td>
<td>NS</td>
<td></td>
</tr>
<tr>
<td>Suicide</td>
<td>10 - 3.0</td>
<td>&lt;0.05</td>
<td></td>
</tr>
<tr>
<td>Cirrhosis</td>
<td>7 - 2.2</td>
<td>&lt;0.05</td>
<td></td>
</tr>
<tr>
<td>Carcinoma of the colon and rectum</td>
<td>16 - 8.9</td>
<td>&lt;0.01</td>
<td></td>
</tr>
<tr>
<td>Carcinoma of the pancreas</td>
<td>11 - 3.9</td>
<td>&lt;0.01</td>
<td></td>
</tr>
</tbody>
</table>

* Diseases with established links with smoking, diseases traditionally linked with gastric surgery, and other diseases with significant differences from the predicted number of deaths are listed.

NS denotes not significant.

† Smoking-associated disease.
Recently it has been suggested that the crucial question determining the fate of patients undergoing surgery for peptic ulcer is whether 5 per cent or more will subsequently have gastric-remnant carcinoma. Should this occur, it will have relatively little impact on the problem of premature death after peptic-ulcer surgery. Of much greater numerical importance is the problem of premature death caused by diseases associated with cigarette smoking. Despite reductions in the number of men in the general population who smoke, no such reduction is apparent in the group of patients with chronic peptic ulceration who require surgical treatment. The increase in premature death after surgery that cures ulcer will therefore continue, and since this increase is unrelated to the operation, the move from gastrectomy toward more conservative physiologic operations will not be followed by a significant improvement in life expectancy.

We are indebted to the staff of the Gastric Follow-Up Clinic and the Department of Medical Physics and Engineering for their invaluable assistance during this study, and to Prof. Sir Richard Doll for his helpful comments.

REFERENCES

Late mortality after vagotomy and drainage for duodenal ulcer

Sir,—In their report (5 May, p 1335) on the survival of 711 patients who underwent truncal vagotomy and drainage for peptic ulcer Mr P C H Watt and others refer to our previously published study of 779 patients undergoing surgery for peptic ulcer mainly of gastrectomy. Direct comparison is not possible. The Belfast series included men and women patients, our series was restricted to men. The inclusion of women may explain the discrepancy between actual/predicted ratios (APR) for lung cancer (Belfast 3-3%, Edinburgh 1-32). The actuarial curves show a greater reduction in survival for women than men. To compare the two studies it would be interesting to know the significance levels for survival and for individual disease incidence for the group of men only.

Whatever the reason for any discrepancies between the calculated APRs in Edinburgh and Belfast, there is clearly a real difference in the number of deaths due to gastric carcinoma, which accounts for 7.1% of all deaths in the Belfast study, compared with only 2.2% in the Edinburgh study. As all patients had vagotomy and drainage in the Belfast study compared with less than 13% in the Edinburgh study, the type of operation may explain the difference, though longer follow up may be required before increases are seen after gastrectomy.

None the less, of greater importance is the confirmation by Watt et al. of the much larger contribution to excess mortality from disease having close associations with smoking (ischaemic heart disease, lung and oesophageal carcinoma, and chronic bronchitis). One hundred and forty one of their patients died of these conditions while only 16 died of gastric carcinoma, emphasising the need to influence smoking habits if late mortality is to fall appreciably.

Finally, they confirm our finding of a statistically significant excess mortality from colorectal carcinoma following gastric surgery (APR 1-8, p = 0.01). Although not a major contributor to excess mortality, the increased incidence of colorectal carcinoma is academically stimulating in the search for factors of aetiological importance in the development of such tumours.

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Metabolism of Sodium Pentosan Polysulphate in Man - Catabolism of Iodinated Derivatives

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Key words
Heparin analogue - SP54 - Metabolism - Iodinated derivatives - Organ distribution

Summary
An iodinated derivative of the heparin analogue SP54 has been prepared and used in conjunction with unlabelled SP54 to study the catabolism and organ distribution of this potential antithrombotic agent in healthy human volunteers. As observed previously with 125I-heparin, we found that the 125I-SP54 was rapidly cleared from the circulation, returning later in a desulphated form. Organ distribution studies with 125I-SP54 suggested that the liver and spleen were major sites of desulphation. Gel filtration and Polybrene binding showed the presence of sulphated macromolecular SP54 and desulphated macromolecular and depolymerised SP54 in post-injection urines. No depolymerised material was present in plasma suggesting depolymerisation occurs in the kidney.

Introduction
Studies have demonstrated that heparin analogues which are relatively inactive in vitro may be potent anticoagulants in vivo and have led to a reappraisal of such properties (1, 2, 3). Pentosan polysulphate (SP54) is a semisynthetic linear polymer of (1→4) linked β-D-xylpyranose residues which contains ≈ 1.8 sulphate groups per monosaccharide unit and has a mean molecular weight of 6,000 daltons. It is less active than heparin in vitro but exhibits marked anticoagulant activity after injection in man. Several workers have shown that the inhibition of factor Xa activity in vivo is greater than that obtained with a similar weight of injected heparin while thrombin inhibition is less (4, 5, 6, 7). Fischer et al. have also demonstrated that inhibition of factor Xa generation may be an important mechanism of SP54’s activity (8). Such results have prompted clinical trials of the antithrombotic efficacy of SP54 which suggest that it can prevent postoperative thromboembolic complications (9, 10).

We have, therefore, investigated the catabolism of SP54 in man in view of its potential clinical use as an antithrombotic drug. Studies on the catabolism of heparin and its analogues have been previously hampered by the lack of a labelled molecule combining high specific activity with retention of biological activity but we have described the preparation and use of an iodinated derivatised heparin which possessed both properties (11). We have used a similar approach with SP54 by preparing a derivative containing tyrosyl residues via which the molecule can be iodinated using the conventional chloramine T method.

The labelled SP54 has been used to study the catabolism of SP54 after intravenous and subcutaneous injection in human volunteers and to determine its organ distribution. The results indicate that the catabolism of SP54 follows similar general pathways to those previously described for heparin (11).

Materials and Methods

Derivatisation and Iodination of SP54

Gel filtered fractions of SP54 (Bene-Chemie, W. Germany) were provided by Drs. Tangen and Granath, Pharmacia. Their mean molecular weights were 3,000, 6,000 and 10,000 daltons as determined by low angle light scattering.

Fractionated SP54 was converted from the sodium salt to the tetrahydro-2-lammonium salt, freeze dried and dissolved in 0.5 ml dimethylsulphoxide to a concentration of 40 mg/ml. It was then reacted with 14 mg of carbonyldimidazole at 56°C for 45 min following which 36 mg tyrosyl hydrazide was added and allowed to react overnight at 20°C. The tyrosyl-SP54 derivative was then converted to the Na+ salt by adding excess sodium chloride, gel filtered on Sephadex LH-20 (Pharmacia, U.K.) in distilled water, and finally freeze dried. Tyrosyl incorporation was determined from the dry weight and absorption using a molar extinction coefficient ε_{190}^{max} of 4.55 x 10^4.

50 µg aliquots of derivatised SP54 were iodinated with 125I or 123I (Na125I and Na123I, carrier free, Amersham, U.K.) using the chloramine T method of Greenwood et al. (12). The fraction of mean molecular weight 6,000 was used in all metabolism experiments.

Characterisation of 125I-SP54

The molecular size distribution of 125I-SP54 was compared with that of unlabelled SP54 by gel filtration on a 95 x 3.6 cm column of Sephadex G-50 (Pharmacia, U.K.). The column was developed with 13 mM trisodium citrate, 150 mM NaCl, pH 6.7 and 8.3 ml fractions were collected. The elution position of unlabelled SP54 was determined by measuring metachromasia with azure A (13), while that of the 125I-SP54 was measured by gamma counting (NE-1600 γ counter, Nuclear Enterprises, U.K.).

The labelled and unlabelled SP54 preparations were chromatographed on a 7 x 5.6 cm column of proteamine agarose using an exponential gradient of 13 mM trisodium citrate, pH 6.7 containing from 0 to 3 M NaCl and were detected as described above. Salt concentration was measured conductometrically.

Biological activity of derivatised SP54 was measured in an activated partial thromboplastin time assay (APTT) in vitro by adding known concentrations to a normal human citrated plasma pool (14). The APTT calibration curve was compared with that obtained using unlabelled SP54.

Injection of 125I-SP54 in Human Volunteers

Five healthy volunteers were fully informed of the procedures and aims of the study. Their weights ranged from 52-77 kg (mean 68) and their ages from 28-44 years (mean 34). Four subjects were injected intravenously with either 0, 0.1, 1, 7 or 50 mg SP54 containing 370 KBq 125I-SP54, one subject receiving tracer alone and tracer plus 50 mg SP54 at an interval of 3 weeks. The fifth volunteer was injected subcutaneously with 50 mg SP54.
containing 370 kBq I125I-SP54. The actual amount of injected radioactivity was measured by counting a known fraction of the total volume of injection. 30 mg potassium iodide was given orally each day to the subjects up to one week after injection. The unlabelled SP54 was an unfraccionated sample used clinically.

Blood samples were taken at various times after injection by clean venepuncture from the antecubital veins and processed as described later. Urine samples of measured volume were collected at known times after injection.

**Clearance of I125I-SP54**

Whole blood, platelet poor plasma (PPP), packed blood cells and urine obtained at various times after injection were counted for radioactivity in a gamma counter with the counts being corrected for background counts. The binding capacity of I125I-SP54 for the polybasic chemical Polybrene (Aldrich, U. K.) was determined by end over end mixing of 5 ml plasma or urine with 0.2 ml of a slurry of Polybrene immobilised by linkage to epoxy agarose 6B (Pharmacl, U. K.). The molecular size distribution of I125I-SP54 in plasma or urine was estimated by gel filtration on Sephadex G-50 as described above.

**Organ Distribution of I125I-SP54**

Prior to injection, the syringe containing 10.0 MBq I125I-labelled SP54 and 1 mg unlabelled SP54 was measured against an I125 standard of approximate activity 30 MBq. The subject was then injected in front of a Scintag-Berthold gamma camera and a series of nine anterior views at 5 min intervals starting 7.5 min after injection, were stored on a Technicare MCS 560 computer processor. Immediately following this a single 5 min posterior view was recorded. The net detected I125I counts in the liver and spleen were obtained for each image using variable regions of interest on the computer. The absolute uptake, as a percent of the administered activity, was obtained by relating the geometric mean of the 45 min and 50 min anterior and posterior image to the I125 standard which was also counted using the gamma camera. The uptake in the liver plus spleen was also measured at 3 hr and 43 hr after injection using a profile scan on the whole body counter. The separate retention in the liver and spleen at 3 hr and 43 hr could not be obtained because of the absence of lateral resolution in the whole body counter profile. In all calculations of the I125I retention, allowances were made for radioactive decay.

**Results**

**Characterisation of Iodinated SP54**

In initial labelling experiments with molecular weight fractions of SP54, I123I was preferentially incorporated, on a weight basis, into the lowest molecular weight fraction of 3,000 daltons. This was thought to be due to derivatisation proceeding via the

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**Fig. 1** Clearance of I125I-SP54 from the circulation. Radioactivity was monitored in blood samples taken at various times after intravenous injection of (nominal) 370 kBq I125I-SP54, with 50 mg SP54, ▲ ▼; with 7 mg SP54, ○ - ○; with 1 mg SP54, △ - △; with 0.1 mg SP54, ● - ●; and without unlabelled SP54 □ - □. 5 ml sample volumes were counted throughout.

**Fig. 2** Absorption and clearance of I125I-SP54 in the circulation after subcutaneous injection. Radioactivity was monitored in samples taken at various times after subcutaneous injection of (nominal) 370 kBq I125I-SP54 with 50 mg unlabelled SP54. Radioactivity in whole blood, ▲ ▼; in plasma, ○ - ○; and in packed cells, ● - ●. 5 ml sample volumes were counted throughout.

**Fig. 3** Elution profile of gel-filtered radioactivity in plasma and urine obtained after i.v. injection of I125I-SP54. Radioactivity was measured in fractions obtained after gel filtration of 5 ml plasma obtained at 320 min after injection of 370 kBq I125I-SP54, ● - ●; or 5 ml urine at 90 min after injection, ○ - ○; on a 95 × 3.6 cm column of Sephadex G-50 equilibrated with 13 mM citrate, 150 mM NaCl, pH 6.7. V₀ = void volume and Vₗ = solvent volume of the column.
reducing end of the molecule, rather than at free hydroxyl groups throughout the molecule, since the charge density of each labelled fraction was similar. Accordingly, a fraction of SP54 whose mean molecular weight was 6,000 was used for derivatisation and iodination and this exhibited a major elution peak coincident with that of unlabelled SP54 upon gel filtration. A second peak of radioactivity, which averaged 10% of the applied load, eluted in the solvent volume of the Sephadex G-50 column. Tyrosine was typically incorporated at 0.5 mole per mole of SP54.

Labelled and native SP54 eluted from protamine agarose at a similar salt concentration of 1.7 M NaCl. Up to 14% of the labelled material did not bind to the solid phase in 15 mM citrate.

Biological activity of the derivatised SP54 preparation of mean molecular weight 6,000 daltons was assessed by prolongation of the APTT of normal human plasma containing from 0–10 µg/ml derivatised SP54 or unmodified SP54. The activity of the derivatised SP54 averaged 69% of the activity of unmodified SP54, tested at three different concentrations.

**Clearance of Injected \( {\text{\(^{125}\text{I}}\)SP54} \)**

Radioactivity was initially cleared from the blood with a half-life of between 13–18 min at all i.v. doses except 50 mg where the half-life was 45 min, assuming zero order kinetics (Fig. 1). 90% of the radioactivity had been removed within 80 min of injection for the lower doses and within 240 min for the highest dose. Most remaining radioactivity was cleared in a second phase over the next 24–96 hr. The clearance of \( {\text{\(^{125}\text{I}}\)SP54} \) from plasma and blood was initially similar, but the radioactivity in plasma decreased more rapidly than in whole blood, due to the progressive association of \( {\text{\(^{125}\text{I}}\)SP54} \) with the packed cell fraction as shown in Table 1.

After s.c. injection of 50 mg SP54 with 370 kBq \( {\text{\(^{125}\text{I}}\)SP54} \) radioactivity became detectable at 5 min post-injection and the peak radioactivity in the blood occurred at 80 min as shown in Fig. 2.

**Appearance of \( {\text{\(^{125}\text{I}}\)SP54} \) in Urine**

Radioactivity was detectable in urine within an hour of i.v. injection. The recovery of radioactivity in the urine within 24 hr following both i.v. and s.c. injection averaged 31% of the injected dose (range 22–43%) and was not related to the dose of unlabelled SP54.

### Table 1  \( {\text{\(^{125}\text{I}}\)SP54} \) associated with plasma or packed cells

<table>
<thead>
<tr>
<th>Time after injection of 7 mg SP54 plus 370 kBq ( {\text{(^{125}\text{I}})SP54} ) (min)</th>
<th>Radioactivity expressed as % of counts in whole blood</th>
<th>Plasma</th>
<th>Packed cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>100</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>10</td>
<td>98</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>20</td>
<td>101</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>40</td>
<td>97</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>80</td>
<td>93</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>160</td>
<td>70</td>
<td>32</td>
<td>0</td>
</tr>
<tr>
<td>320</td>
<td>67</td>
<td>34</td>
<td>0</td>
</tr>
<tr>
<td>640</td>
<td>67</td>
<td>35</td>
<td>0</td>
</tr>
<tr>
<td>1440</td>
<td>61</td>
<td>38</td>
<td>0</td>
</tr>
</tbody>
</table>

Radioactivity due to plasma trapped in the packed cell fraction was estimated and subtracted to allow the calculation of cell associated \( {\text{\(^{125}\text{I}}\)SP54} \).

#### Characterisation of \( {\text{\(^{125}\text{I}}\)SP54} \) in Plasma and Urine

a) **Affinity for immobilised Polybrene**. The percentage of radioactivity in plasma which bound to immobilised Polybrene decreased with time after injection and the rate of decrease in binding was less after 50 mg unlabelled SP54 than after the lower doses (Table 2). \( {\text{\(^{125}\text{I}}\)SP54} \) with affinity for immobilised Polybrene was present in urine samples obtained soon after injection but not in later samples. Early urine samples contained more Polybrene-binding material after injection of \( {\text{\(^{125}\text{I}}\)SP54} \) with 50 mg unlabelled SP54 than for the lower doses of cold SP54. Thus in the initial urine sample 95 min after injection of 50 mg SP54, 78% of the radioactivity bound to Polybrene agarose while in the urines at 48 and 90 min after injection of 0.1 mg SP54 the binding was 43 and 9% respectively. With all doses the binding in urine was less than 10% by 6 hr after injection.
Table 2  Relationship between binding of 125I-SP54 in plasma to immobilised Polybrene and time after injection showing the influence of accompanying dose of unlabelled SP54

<table>
<thead>
<tr>
<th>Dosage of unlabelled SP54 (mg) injected together with 370 kBq 125I-SP54</th>
<th>Time after injection (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>5</td>
</tr>
<tr>
<td>0.1</td>
<td>81</td>
</tr>
<tr>
<td>1</td>
<td>89</td>
</tr>
<tr>
<td>7</td>
<td>75</td>
</tr>
</tbody>
</table>

Binding shown as % of total counts added to binding system

b) Molecular size of 125I-SP54 in plasma and urine assessed by gel filtration. Figure 3 shows the elution profiles of gel filtered plasma and urine obtained after injection of 370 kBq 125I-SP54 in the absence of any unlabelled SP54. Radioactivity in the plasma sample obtained at 320 min after injection, eluted in a major peak prior to the solvent volume demonstrating its macromolecular nature. In contrast, radioactivity in a urine sample obtained 90 min after injection eluted almost entirely in the solvent volume of the column, indicating that it was associated with low molecular weight material (<1,000 daltons). Samples of urine obtained at various times after injection of 125I-SP54 plus 7 mg unlabelled SP54 were also gel filtered. Figure 4 shows the elution profile of urines sampled at 125 and 425 min after injection. Radioactivity in the first sample was associated both with macromolecular and degraded material while that in the later sample was entirely associated with low molecular weight material in the solvent volume of the column.

Radioactivity bound to Polybrene-agarose after adsorption of post-SP54 plasma (as described in section [a]) was eluted with 5 M NaCl and gel filtered on Sephadex G-50 using 13 mM citrate containing 3 M NaCl. The radioactivity eluted entirely before the solvent volume indicating the absence of any low molecular weight degraded material (results not shown).

Organ Distribution of 125I-SP54

125I-SP54 was prepared immediately before injection and coeluted with unlabelled SP54 when filtered on Sephadex G-50. 88% of the labelled SP54 had high affinity for immobilised Polybrene. After i.v. administration of 10.0 MBq 125I-SP54 with 1 mg unlabelled SP54 images taken at 5 min intervals from 7.5 to 47.5 min showed progressive uptake of 125I by the liver and spleen (Fig. 5), such that at 50 min 60% and 7.5% of the dose respectively was associated with these organs. A profile scan at 3 hr post-injection showed 60% of the activity in the liver plus spleen and 13% in the bladder. By 43 hr post-injection the liver plus spleen retention was 37%. Urine collected over 18 hr post-injection contained 35% of the administered activity while stools passed at 18 and 42 hr post-injection contained 0.13% and 0.07% respectively.

Discussion

Radioactively labelled heparin has been used by several groups to study the metabolism of heparin in a number of species (15-18). The most commonly used labels have been 35S and H both of which are rather unsuitable for use in man. The former label is quickly detached from the polysaccharide chain during the desulphation reactions and cannot be used to follow the subsequent metabolism of desulphated material. The latter label is hazardous and it is consequently difficult to administer sufficient labelled material for accurate counting. We have previously shown that iodination of a derivatised heparin resulted in a labelled material of high specific activity which retained several of the biological activities characteristic of heparin (11) and which was suitable for injection in man.

A similar approach was chosen with SP54 and the following criteria led us to conclude that the labelled molecule could be successfully used as a tracer of SP54 metabolism in human volunteers. Firstly, the derivatisation of SP54 involves activation of ring hydroxyls or reducing end groups with carbonyldimidazole to give an imidazolyl carbamate derivative which reacts with tyrosyl hydrazide to yield tyrosine coupled to SP54 via a urethane linkage (19). This linkage is very stable under the conditions used in the chromatographic experiments and those existing in plasma. Secondly, the labelled SP54 exhibited similar affinities for protamine and Polybrene to native SP54 and the molecular size distribution taking into account the effect of substitution of the derivatising groups was also similar. Thirdly, the labelled SP54 retained biological activity as assessed by the APTT assay.

The kinetics of clearance of 125I-SP54 could not be accurately determined due to probable redistribution following the bolus injection. Even so it was clear that 125I-SP54 was removed from the circulation more slowly when injected together with 50 mg of unlabelled SP54. These results again point to the similar properties of 125I-SP54 and native SP54, in this case competing for the same mechanism of uptake from the blood-stream. A point of inflection in the plasma clearance curves which was also previously observed after injection of 125I-heparin (11) represents the reappearance of 125I-SP54 or its degradation products from extravascular sites.

Peak absorption of 125I-SP54 occurred after subcutaneous injection at 80 min post-injection, and followed a similar time course to that observed for biological activity in other studies (5, 6, 7, 8). These results demonstrate the suitability of the tracer for following uptake and clearance of SP54 by the subcutaneous route.

The redistribution of 35S-labelled SP54 from the plasma to the packed red cells has been noted in the rat. In the present experiments 30-40% of labelled material in later samples was associated with cells (shown to be leucocytes by cell fractionation), but this total radioactivity only represented about 2% of the injected dose. Binding to cellular elements of the blood is therefore not a major metabolic pathway for SP54 clearance which, like heparin, is probably mainly removed by the reticuloendothelial system (20).

Gel filtration and Polybrene-binding experiments showed that only macromolecular 125I-SP54 was present in plasma and that the proportion with affinity for Polybrene decreased with time after injection. We conclude that the material without affinity is macromolecular desulphated SP54 and that the probable sites of desulphation are the liver and spleen which are rich sources of sulphatases (21). The 125I-SP54 imaging data supports this conclusion since a major portion of the label became localised in these organs.

From the absence of depolymerised 125I-SP54 in plasma it appears that the kidney is the site of depolymerisation. Only depolymerised label was found in the urine after injection of 125I-SP54 alone while after injection of 125I-SP54 with high doses of unlabelled SP54 both macromolecular label (some with affinity for Polybrene) and depolymerised label was detected. These results demonstrate the saturation of the desulphation and depolymerisation mechanisms at pharmacologically relevant doses of SP54. Such saturation has also been observed with clinical doses of heparin (22), although the enzymic pathways for
SP54 catabolism must differ from that of heparin due to structural differences and remain to be defined.

Acknowledgements

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References


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Correction Factors for Gravimetric Measurement of Peritumoural Oedema in Man

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Abstract
The water content of samples of normal and oedematous brain in lobectomy specimens from 16 patients with cerebral tumours has been measured by gravimetry and by wet and dry weighing. Uncorrected gravimetry underestimated the water content of oedematous peritumoural cortex by a mean of 1.17%, and of oedematous peritumoural white matter by a mean of 2.52%. Gravimetric correction equations calculated theoretically and from an animal model of serum infusion white matter oedema overestimate peritumoural white matter oedema in man, and empirical gravimetric error correction factors for oedematous peritumoural human white matter and cortex have therefore been derived. These enable gravimetry to be used to accurately determine peritumoural oedema in man.

Key words: Brain oedema, specific gravity, gravimetry.

Introduction
Interest in gravimetry of the brain began 135 years ago in Devon, with Sir Charles Bucknill's measurements on the brains of deceased lunatics. A century after Bucknill's studies, Lowry used gravimetry to study changes in brain volume produced by freezing, and in 1971 Nelson and his colleagues introduced microgravimetry for the measurement of cerebral oedema induced by water intoxication in mice. Nelson's technique was subsequently refined in the cat, and has been widely applied to other animals and to man.

The estimation of tissue water content, and hence oedema, from gravimetric measurements requires the specific gravity of the tissue solids to be known. This is a constant in oedema produced by water intoxication, but varies when the oedema fluid contains protein, and when the blood volume in the tissue changes. Marmarou and his colleagues derived a correction for the white matter of the cat brain made oedematous by the direct infusion of serum, which introduced an error in uncorrected gravimetric water content measurements that approached 4.5%. This serum infusion oedema model is designed to simulate vasogenic oedema in which vascular permeability to plasma proteins is abnormal. Whilst it is suggested that the correction factors derived from these experiments might allow gravimetry to accurately measure water content in cryogenically induced oedema, the possibility that a different experimental model and different species may introduce new errors is discussed, and no correction to account for blood volume changes is made.

An assessment of the accuracy of gravimetry
to measure clinical brain oedema in man has not been made, and this is what has been done in the study reported here.

**Patients and Methods**

Neurosurgical operative treatment offered the opportunity to measure brain water in 16 patients with intrinsic supratentorial tumours of glial or metastatic origin. There were 14 men and two women, with an age range of 39–73 years (mean 58 years), and all were investigated with preoperative CT (computed X-ray transmission tomography) and NMR (nuclear magnetic resonance) imaging of their tumour. Surgery followed MRI (magnetic resonance imaging) within 24 h, and 7 patients had a frontal lobectomy, 4 a temporal lobectomy, and 5 an occipital lobectomy. The lobectomy specimens were sealed in copolymer film (Ethicon Ltd, Edinburgh, Scotland) to prevent evaporative water loss, and taken from the operating theatre to an adjacent laboratory.

Eleven samples of oedematous peritumoural cortex were obtained from the specimens, and 13 samples of oedematous peritumoural white matter. Thirteen samples of normal cortex and 8 of normal white matter were obtained from the poles of the lobectomy specimens in those patients whose MRI showed a normal longitudinal relaxation time ($T_1$) at the poles of the lobe to be resected.

Each tissue sample was divided, and part of it was weighed in its fresh state on a small piece of dry aluminium foil that had an accurately predetermined weight. The sample was then dried in an oven at 80°C, and re-weighed at 24 h intervals until it reached a constant weight, which was then recorded. The wet tissue weight ($W$) was obtained by subtraction of the foil weight from the fresh weight, and the dry tissue weight ($D$) by subtraction of the foil weight from the dried weight. The wet tissue weights ranged from 8.9 to 484.1 mg, with a mean of 102.8 mg. Water content was expressed as a percentage by weight of the fresh tissue using the formula:

\[
\% \text{ water} = \frac{W-D}{W} \times 100
\]

The remainder of each fresh tissue sample was then divided into three portions weighing approximately 5 mg each, and they were placed in a calibrated layered oil density column for specific gravity measurements\(^3\).\(^4\). Tissue water content was calculated from the specific gravity of fresh tissue ($SG$) using the equation:

\[
\frac{m}{SG} = \frac{m - b}{SG - 1}
\]

where $m = \frac{SG\text{solids}}{SG\text{solids} - 1}$

and $b = \frac{1}{SG\text{solids} - 1}$ (Marmarou\(^4\))

and $SG\text{solids} = \frac{1 - (SG-1) \times W}{D \times SG}$ (Nelson\(^2\))

($SG\text{solids}$ = derived specific gravity of the solid component of normal brain tissue)

For our samples of normal brain, $m = 4.688$ and $b = 3.688$ for cortex, and $m = 7.430$ and $b = 6.430$ for white matter. These values are similar to those derived for man by Takagi and his associates\(^6\). Modified formulae were also derived for man using Marmarou's theoretical equations\(^2\) for both oedematous cortex, where $m = 8.145$ and $b = 7.005$, and for oedematous white matter, where $m = 15.905$ and $b = 14.566$. (For the derivation of these values of $m$ and $b$ please refer to the appendix).

**Results**

For the 13 samples of normal cortex, the mean (± standard deviation) water content measured by wet and dry weighing was 80.96% (±0.86%), and derived from specific gravity measurements was 81.01% (±0.92%). The specific gravity of the solid component of normal human cortex was 1.2711 (±0.0230).
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For the 8 samples of normal white matter, the mean water content measured by wet and dry weighing was 70.23% (± 0.88%), and derived from specific gravity measurements was 70.25% (± 1.18%). The specific gravity of the solid component of normal human white matter was 1.1555 (± 0.0086).

The specific gravity of the solid component of the brain increased as peritumoural oedema increased, for both cortex and white matter. The rise in specific gravity of the tissue solids was greater in cortex, where it averaged 0.0181 per 1% oedematous water increase (Fig. 1), compared to an average rise of 0.0044 per 1% oedematous water increase for white matter (Fig. 2).

For the 11 samples of oedematous peritumoural cortex, the mean water content measured by wet and dry weighing was 83.54%, and derived from uncorrected specific gravity measurements using Nelson's formula3 was 82.37%. The cortex results are presented graphically in Fig. 3, and the progressive underestimate of water content with increasing oedema is apparent from the slope of the regression line of the data (0.3841) compared with the line of identity for the two different measuring techniques which has a slope of unity. For the 13 samples of oedematous peritumoural white matter, the mean water content measured by wet and dry weighing was 77.74%, and derived from uncorrected specific gravity measurements using Nelson's formula3 was 75.22%. The white matter results are presented graphically in Fig. 4, and also show a
progressive underestimate of water content with increasing oedema, although with less error than in cortex, as the slope of the regression line for the data (0.6068) is closer to unity.

Discussion

Many workers who quantify cerebral oedema with gravimetry have recognised the potential for the errors highlighted by Marmarou and his colleagues and have avoided calculating water content from specific gravity measurements. They express their results directly as specific gravity figures, and take a fall in specific gravity to indicate cerebral oedema. This may not always be a valid interpretation. Taking as an example a 1 ml sample of normal human white matter:

**Example 1**

<table>
<thead>
<tr>
<th>Property</th>
<th>Formula</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fresh weight</td>
<td>$1.0417,g$</td>
<td></td>
</tr>
<tr>
<td>Specific gravity</td>
<td>$1.0417,g/1,ml$</td>
<td>$1.0417$</td>
</tr>
<tr>
<td>Dry weight</td>
<td>$0.3104,g$</td>
<td></td>
</tr>
<tr>
<td>Water content from wet</td>
<td></td>
<td>$70.21%$</td>
</tr>
<tr>
<td>and dry weights</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Water content from $SG$ (Nelson)</td>
<td></td>
<td>$70.21%$</td>
</tr>
</tbody>
</table>

For the same 1 ml sample made oedematous by adding 0.5 ml of pure water to it:

**Example 2**

<table>
<thead>
<tr>
<th>Property</th>
<th>Formula</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fresh weight</td>
<td>$1.0417+0.5000,g$</td>
<td>$1.5417,g$</td>
</tr>
<tr>
<td>Specific gravity</td>
<td>$1.5417,g/1.5,ml$</td>
<td>$1.0278$</td>
</tr>
<tr>
<td>Dry weight</td>
<td>$0.3104,g$</td>
<td></td>
</tr>
<tr>
<td>Water content from wet</td>
<td></td>
<td>$79.87%$</td>
</tr>
<tr>
<td>and dry weights</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Water content from $SG$ (Nelson)</td>
<td></td>
<td>$79.87%$</td>
</tr>
</tbody>
</table>

If 0.5 ml of normal human plasma containing 0.036 g of plasma protein is substituted for water as the oedema fluid:

**Example 3**

<table>
<thead>
<tr>
<th>Property</th>
<th>Formula</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fresh weight</td>
<td>$1.5417+0.0360,g$</td>
<td>$1.5777,g$</td>
</tr>
<tr>
<td>Specific gravity</td>
<td>$1.5777,g/1.5,ml$</td>
<td>$1.0518$</td>
</tr>
<tr>
<td>Dry weight</td>
<td>$0.3104+0.0360,g$</td>
<td>$0.3464,g$</td>
</tr>
<tr>
<td>Water content from wet</td>
<td></td>
<td></td>
</tr>
<tr>
<td>and dry weights</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Water content from $SG$ (Nelson)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

In these examples it has been assumed that none of the protein from the plasma oedema has been cleared from the oedematous tissue, and the approximation that 1 ml water weighs 1 g at the temperature at which weighing was performed has been accepted. With these limitations, the extent of the error introduced by plasma protein in oedema fluid is apparent in Example 3, with an erroneous fall in water content calculated using the uncorrected equation. This erroneous fall in water content would also be construed from the measured rise in specific gravity to 1.0518. Specific gravity and water content derived by Nelson's formula are directly mathematically related, and the acceptance that calculated water content figures are inaccurate means that the underlying specific gravity figures are an equally inaccurate indicator of oedema.

Specific gravity figures alone cannot therefore be quoted when quantifying oedema, and the derivation of a correction factor to allow water content to be calculated from specific gravity, as advocated by Marmarou and his colleagues, is essential for each particular experimental protocol, or clinical aetiology. To test whether Marmarou's corrections derived from the serum infusion white matter oedema model in the cat are applicable to clinical peritumoural oedema in man, we have compared white matter water content measured by wet and dry weighing, with water content calculated from the specific gravity of the same samples by Marmarou's derived equation corrected for man (Fig. 5). Marmarou's equation progressively overestimates water content for this type of oedema in man, as the slope of the regression line for the data (1.3044) exceeds unity. Deriving a corrected equation for human cortex in accordance with Marmarou, a closer
estimate of water content is obtained from specific gravity measurements of oedematous peritumoural cortex (Fig. 6).

![Graph 1](image1)

**Fig. 5.** Oedematous white matter water content derived by Marmarou's equation.

![Graph 2](image2)

**Fig. 6.** Oedematous cortex water content derived by Marmarou's equation.

The reason why Marmarou's corrected equation overestimates oedema in peritumoural white matter relates to the degree of the increase in the specific gravity of tissue solids that occurs with increasing white matter oedema. Marmarou assumes in his equations that the protein content of vasogenic oedema fluid equals that of serum. This latter assumption will lead to an overcorrection if in fact the oedema fluid contains less protein than plasma. This is highly likely in the case of peritumoural white matter oedema in man, where the leak in the blood brain barrier may be partially selective and exclude some plasma proteins from the oedema fluid. Some of the proteins that are extravasated into the oedema fluid may also be cleared from the tissue by reabsorption into the vascular compartment, further reducing the equilibrium protein content of the fluid. The measured rise in the specific gravity of tissue solids for a 1% increase in oedematous water content is less for human peritumoural white matter (0.0044 from Fig. 2) than for peritumoural cortex (0.0181 from Fig. 1), and this disparity between the two tissues is likely to be accounted for by a lower protein content of white matter oedema fluid compared to cortex oedema fluid.

Empirical correction factors can be derived from the regression line equations of our data in Figs. 3 and 4. The error for oedematous cortex shown in Fig. 3, can be corrected by subtracting the constant (50.28) from the uncorrected specific gravity derived water content and multiplying the result by 2.603 (the reciprocal of the slope of the regression line). The error for oedematous white matter shown in Fig. 4, can be corrected by subtracting the constant (28.05) from the uncorrected specific gravity derived water content and multiplying the result by 1.648 (the reciprocal of the slope of the regression line).

It must be stressed that these correction factors are for uncorrected water content derived from specific gravity measurements by Nelson's formula, and apply only to peritumoural oedema in man. They cannot be applied to animals, or where oedema in man has been produced by pathological processes other than a tumour. All measurements of brain oedema, where the oedema fluid is not pure water, must initially use wet and dry weighing as a reference standard to allow the derivation of correction factors for subsequent gravimetric estimations. Gravimetry alone in these circumstances cannot measure brain oedema accurately, and the error is in the specific gravity values themselves (Example 3) as well as in the water content figures derived from them by uncorrected equations.
Appendix

Derivation of \( m \) and \( b \) for human brain, where the oedema fluid is pure water, from Marmarou's equation:

\[
m = \frac{SG_{\text{solids}}}{SG_{\text{solids}} - 1}
\]

\[
b = \frac{1}{SG_{\text{solids}} - 1}
\]

For our samples of normal human cortex, mean \( SG_{\text{solids}} = 1.2711 \), therefore:

\[
m = \frac{1.2711}{1.2711 - 1} = 4.688
\]

\[
b = \frac{1}{1.2711 - 1} = 3.688
\]

For our samples of normal human white matter, mean \( SG_{\text{solids}} = 1.1555 \), therefore:

\[
m = \frac{1.1555}{1.1555 - 1} = 7.430
\]

\[
b = \frac{1}{1.1555 - 1} = 6.430
\]

Derivation of \( m \) and \( b \) for human brain, assuming the oedema fluid contains the same amount of protein as plasma, from Marmarou:

Equation A10

\[
m = \frac{(Brain \: Water - Plasma \: Water) \times (SG_{\text{brain}} \times SG_{\text{plasma}})}{(SG_{\text{plasma}} - SG_{\text{brain}})}
\]

Equation A11

\[
b = \frac{(Plasma \: Water \times SG_{\text{plasma}}) - (Brain \: Water \times SG_{\text{brain}})}{(SG_{\text{plasma}} - SG_{\text{brain}})}
\]

For normal man, \( SG_{\text{plasma}} = 1.0264 \) and Plasma Protein = 0.0720 g protein/ml plasma.

\[
\begin{align*}
\text{Plasma Protein} & = 0.0720 \\
SG_{\text{plasma}} & = 1.0264 \\
\end{align*}
\]

For our samples of normal human cortex, mean Brain Water = 0.8096 g water/g brain (by wet/dry weighing), and mean \( SG = 1.0422 \).

Therefore:

\[
\begin{align*}
m & = \frac{(0.8096 - 0.9299) \times (1.0422 \times 1.0264)}{1.0264 - 1.0422} \\
& = 8.145
\end{align*}
\]

and

\[
\begin{align*}
b & = \frac{(0.9299 - 1.0264) \times (0.8096 - 1.0422)}{1.0264 - 1.0422} \\
& = 7.005
\end{align*}
\]

For our samples of normal human white matter, mean Brain Water = 0.7023 g water/g brain (by wet/dry weighing), and mean \( SG = 1.0417 \).

Therefore:

\[
\begin{align*}
m & = \frac{(0.7023 - 0.9299) \times (1.0417 \times 1.0264)}{1.0264 - 1.0417} \\
& = 15.905
\end{align*}
\]

and

\[
\begin{align*}
b & = \frac{(0.9299 - 1.0264) \times (0.7023 \times 1.0417)}{1.0264 - 1.0417} \\
& = 14.566
\end{align*}
\]

Editorial comment

The gravimetric method has been shown by many authors to be a useful technique for measurement of brain tissue water. It allows us to measure water content of small samples rapidly and accurately and has considerable advantages over conventional oven drying methods. This article by Dr Bell and his colleagues, emphasises an important point which is crucial to the correct application of the gravimetric methodology. Specifically, one must determine the nature of the edema fluid before using the appropriate equations, which convert the gravimetric measures directly to water content. In this study, it has been shown that the fluid, which exudes from the tumorous site in man, differs from that of serum. Thus, new conversion factors are necessary in order to accurately measure water content in adjacent tissue. Accordingly, it would seem proper that studies of tumorous edema in man utilise the equations described in this report. Complexities arise when compressive effects of the tumor volume result in barrier compromise or ischemia, in which case the nature of the extracellular fluid is difficult, if not impossible, to define. Under these circumstances, conventional wet/dry techniques share the same errors as gravimetry and given a choice, the features of small sampling and rapid determination offered by gravimetry remain attractive.

ANTHONY MARMAROU
Interindividual differences in the pituitary-thyroid axis influence the interpretation of thyroid function tests

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Summary

OBJECTIVE We investigated interindividual differences in the shape, slope and setpoint of the pituitary-thyroid axis (PTA) in normal persons. Based on these physiological data we propose a novel bivariate concept for the interpretation of thyroid function tests which is less biased by interindividual differences in the PTA than the currently used univariate approach.

DESIGN In two separate trials (A and B), healthy volunteers were given small, increasing doses of T3 (7–45 μg/day orally) over 5 days. The regulation characteristics of the individual PTAs and the effects of age and gender were assessed by general linear regression models. In addition, serum samples were collected from normal persons to establish the proposed bivariate approach for the interpretation of thyroid function tests.

SUBJECTS The regulatory characteristics of the PTAs were determined in a total of 21 normal volunteers (eight females, 13 males; age 24–49 years). Single blood samples were collected from 257 normal volunteers. The participants had no history of pituitary or thyroid disease.

MEASUREMENTS Free and total thyroid hormone and TSH concentrations were determined in the serum. All samples from one person were analysed in the same assay in duplicate.

RESULTS A log-linear relationship between T3 and TSH was found to describe best the individual PTAs (multiple r=0.96). However, significant differences were found in the setpoint (P<0.001) and to a lesser degree in the slope (P=0.05) of the PTAs; this variability was not dependent on age or gender. Since these findings invalidate the assumptions on which the current univariate interpretation of thyroid function tests is based, we propose a novel model for the evaluation of thyroid function tests derived from the experimentally determined shape and average slope of the PTA.

CONCLUSIONS The presence of significant age and gender-independent interindividual variations in the set-point of the pituitary-thyroid axis raises conceptual problems with the current approach for interpreting thyroid function tests. An easy to use graphical bivariate representation of the normal ranges for thyroid function tests is presented and exemplified by the thyroid hormone and TSH measurements in a large reference population. This concept should improve the diagnostic accuracy in the borderline-normal, and particularly subclinical hypothyroid region of these hormone measurements.

To assess the thyroid function of a patient, the thyroid hormone and TSH concentrations are measured in the serum. The results are formally interpreted independently of each other, i.e. by applying separate normal ranges for T3, T4 and TSH levels (Caldwell et al., 1985; Klee & Hay, 1987; Helfand & Crapo, 1990; Liewendahl, 1990; Sukrs et al., 1990). Nevertheless, many clinicians intuitively reason that, for example, a low-normal T4 level is more likely to be indicative of early hypothyroidism when accompanied by a high-normal rather than a low-normal TSH level. Presently, however, no method exists which permits this knowledge to be exploited quantitatively. The categorization of primary thyroid disorders into subclinical and overt hypo- and hyperthyroidism is a direct consequence of this univariate approach as shown in Fig. 1. Two fundamental problems are raised by such a classification.

(1) When interpreting the thyroid function tests in a patient, the thyroid hormone and TSH levels are implicitly assumed to be uncorrelated in an individual normal person within the normal range of the hormone concentrations. If a correlation between these parameters exists, the individual shape, slope and setpoint of this relationship influence the extension of the individual normal range as represented by the maximally allowed variations of thyroid hormone and TSH levels within the normal range derived from a population (Fig. 1, see legend) thus introducing an unpredictable bias.
measure the FT4 serum levels within a range of 0–93 pmol/l, the normal range being 9–24.5 pmol/l (intra-assay CVs 9 and 4%, respectively). The concentration of TSH was determined with the ultrasensitive RIA-gnost hTSH assay using a normal range of 0.3–3.0 mIU/l. The intra-assay CVs for 0–13 mIU/l were 20 and 3% for 5 mIU/l.

Study B. The FT3 serum concentrations were measured using the Amerlex-M RIA kit (Amersham Inc., UK) covering a range of 0–40 pmol/l with a normal range of 2.9–8.9 pmol/l. The intra-assay CV was 3% at 5.0 pmol/l. The Amerlex-M Free T4 RIA was utilized to measure the FT4 serum levels within a range of 0–130 pmol/l, the normal range being 9.4–25 pmol/l. The intra-assay CV was 4% at 14.8 pmol/l. The concentrations of TSH in the sera were determined with the ultrasensitive Amerwell TSH immunoradiometric assay (monoclonal) using a normal range of 0.36–3.25 mIU/l. The sensitivity was better than 0.06 mIU/l and the intra-assay CV for 0.8 mIU/l was 4–4%.

All samples from one person were analysed in the same assay in duplicate. The normal ranges given above are those currently in use by the respective laboratories and were utilized in Fig. 2.

Statistical methods

The data transformations, summary statistics, generalized linear regression modelling, Wilcoxon signed rank test, Kolmogorov-Smirnov and chi-squared goodness-of-fit tests were carried out on an IBM-PC with SYSTAT 5.03 (Systat Inc., Evanston IL, USA). Analysis of covariance with a repeated measures design was used to compare the slope and intercept of multiple regression lines (Neter et al., 1990). Similar multiple regression models were specified to test the influence of sex and age on the interindividual differences in the pituitary–thyroid axis. Unless stated otherwise the results were expressed as mean ± one standard deviation (± SD).

Results

Table 1 summarizes the hormone levels of the 21 persons from studies A and B after the administration of the T3 doses indicated. In study A, the TSH and FT4 concentrations had already declined significantly on day 2. Until and including day 4 the TT3 levels remained within the normal range accompanied by steadily declining TSH concentrations, demonstrating the continuous regulation of TSH by T3 within the normal range of the hormone concentrations. Because a small threshold in the pituitary–thyroid regulation could not be excluded from these data, this possibility has been investigated by the measurement of the TSH levels every 20 minutes after a single oral dose of 6 µg T3 was given to a healthy person. This resulted in a decline of TSH levels by 0.6 mIU/l within 4 hours (duplicate experiment in one person, controlled for circadian rhythmicity; data not shown). Taking into account the slightly different dosage regimen, study B showed a similar time course of the relative hormone level changes.

The regulation characteristics of the individual feedback axes were investigated by fitting a separate line through the T3 vs TSH data points of each person using multiple regression with a repeated measures factor. The residual mean squares were significantly lower when a regression line was fitted through the log-transformed rather than the linear TSH levels (P < 0.0005 in studies A and B); hence a logarithmic curve described the individual relationship between TT3/TSH in study A (multiple r = 0.97) and FT3/TSH in study B (multiple r = 0.95) well. However, a signifi-
Fig. 3 Representation of the proposed modified bivariate concept for the interpretation of thyroid function tests. The FT3 and TSH levels of 257 normal individuals were plotted and a univariate normal range box (dashed lines) based on the 2.5th and 97.5th percentiles was calculated. Three elastic bivariate confidence ellipses (for 95-95, 99-99 and 99-99%) corresponding to the probabilities of ±2 SD, ±3 SD and ±4 SD, respectively) are shown in solid lines. The zones (stippled lines) for ±2 SD, ±3 SD and ±4 SD derived from the proposed modified bivariate model have been calculated as described in the Appendix. Essentially, for each person the FT3 and TSH levels are standardized and subtracted to yield a new parameter ZD, which is representative of the distance moved along the average pituitary-thyroid axis. The normal range for ZD is between ±2 SD, while the other lines (±3 SD, ±4 SD) may represent an empirical measure of the probability to suffer from thyroid disease. Contrary to the univariate and classical bivariate approach, this concept allows for interindividual differences in the setpoint of the feedback axis.

feedback axis and its parallels as depicted in Fig. 3 and described in detail in the Appendix. The average axis is defined by the experimentally found logarithmic shape and the average slope from study B. These zones provide several advantages over the univariate normal range matrix as well as over the bivariate confidence ellipses.

1. They allow for differences in the setpoint of the pituitary-thyroid axis and are less sensitive to interindividual variations in the slope of the axis than is the conventional approach. It is apparent from the statistical analysis and Fig. 2 that the variability in the setpoint is an important factor for the interindividual differences in the regulation axes. Although an ideal model would take into account the slope of the axis in a given patient, this information is obviously not generally available.

2. Not only can the binary question, whether a pair of FT3 and TSH levels is within or outside the normal range, be answered but an empirical measure of the probability that a patient with borderline abnormal thyroid function tests suffers from early thyroid dysfunction may be derived in prospective studies. This could prove particularly useful in a more refined classification of subclinical thyroid disease, particularly for research purposes (O'Malley et al., 1989; Ross et al., 1989; Tenerez et al., 1990; Roden et al., 1991; Wartofsky, 1991).

3. The different stages of thyroid disease (e.g. normal—subclinical hypothyroidism—overt hypothyroidism) are conceptually better represented in adjacent zones rather than by the univariate matrix (Wehmann & Nisula, 1984) or concentric confidence ellipses.

The clinical usefulness of this zonal concept and the assignment of probabilities for thyroid dysfunction to the different zones needs to be established in future studies. Since thyroid function tests are used simultaneously as the tools for defining and diagnosing thyroid disease, only a large prospective study will establish a pattern of borderline abnormal T3 and TSH levels which are associated with an increased risk for thyroid disease. Nonetheless, in addition to the theoretical considerations outlined above, indirect evidence suggests that the proposed concept will prove clinically useful. When the serum T4 levels from a large population of patients with and without thyroid disease are plotted against the logarithm of the TSH concentrations, the data points form a distinct band of constant width along a linear regression line representing the average regulation axis (Spencer et al., 1990). It is evident that in such a situation zones orthogonal to the normalized regression line would represent a more adequate normal range than a univariate box with no direct relationship to the regulation axis. The clinical relevance of the modified bivariate model as depicted...


A whole body NMR imager has been installed in the Royal Infirmary, Edinburgh, for clinical evaluation. This paper gives details of the installation of the first commercial NMR imager to be used in a hospital environment in the United Kingdom.

**NMR IMAGER**

The NMR imager is the first production machine manufactured by an Aberdeen-based company.* The design of the imager is based on the "Mark II NMR Imager" developed at Aberdeen Royal Infirmary. Though the physical principles are the same, the computer control circuits and much of the electronics have been redesigned.

The water-cooled, resistive magnet consists of four circular coils, horizontally mounted, producing a field of 0.08T (1 Tesla = 10,000 Gauss) in the vertical direction. The patient enters between the two larger coils, which have a separation of 0.35 m, and lies inside the patient tube which is 0.56 m diameter and 1.2 m long. Wound round the outside of the patient tube are the windings which produce the three orthogonal magnetic field gradients. The RF coil has a slightly larger diameter and is in the form of two linked solenoids in the central region of the patient tube. The overall dimensions of the imager, including the radiofrequency (RF) shield are 2.35 m wide, 2.03 m high and it weighs about 2000 kg. The patient couch extends a further 2.65 m and it weighs 350 kg.

The RF and gradient amplifiers, temperature control, power supply control and NMR lock are contained in one cabinet with the power supply itself in another cabinet. The power supply cabinet is also water-cooled in parallel with the magnet. The imager is controlled by a PDP 11/23+ computer which drives the RF and gradient amplifiers and also analyses the NMR signals. This computer has two RL02 discs for storage and is part of the operator's console, which in addition contains a black and white monitor and a VDU with keyboard. Images in the transverse, sagittal or coronal plane are obtained using a two-dimensional Fourier transform, the "spin warp" technique (Edelstein et al, 1980). Saturation recovery and inversion recovery pulse sequences are used with a suitable time between pulses to give proton density and T1 images. The computer of the operator's console is linked to a second PDP 11/23, working on a serial line, which acts as an independent diagnostic console. The diagnostic console contains a Winchester disc, an 8-inch floppy disc and a magnetic tape unit for patient data storage, plus black and white and colour monitors and a VDU with keyboard.

**SITE PREPARATION**

The NMR imager is installed in a room which previously had been a laboratory in the University Department of Medicine. The room is part of a single-storey, metal framed building, with girders of the metal frame in the centre of the room. One adjacent room is a store which contains some ferrous material, most of which is rarely moved. In the room where the imager is sited there had been a large treadmill which could be lowered below floor level when not in use. Though this contains a large amount of ferrous material it was not removed but permanently floored over. The specifications from M&D Technology Ltd. required that there should be no ferrous material within 3 m of the centre of the magnet. The material in the storeroom, the vertical support girders and the treadmill are sited at different positions around the magnet site but all are just outside the 3 m limit (Fig. 1). The only ferrous material within the 3 m limit which was removed was a short length of steel trunking sunk into the floor and two central heating radiators along the wall. The laboratory cupboards at the end of the room nearest the magnet are wooden with a metal frame but though they are within the 3 m limit they were not removed.

A separate three-phase supply has been installed for the NMR imager, computers and refrigerator unit. Plastic trunking is used for the electricity cables within the NMR room. Although a water supply was already in the room, a larger 22 mm mains supply has been installed from the adjacent laboratory to increase the flow for the run-to-waste system. A 15 mm mains supply was connected which ran outside the room to the position of the refrigerator unit.

Other modifications made to the room, such as the construction of a partition to create office space, are included in our financial costs for the installation of the NMR imager, though they are not specific for NMR.
The four magnet coils and two sets of shimming coils. In addition, small amounts of ferrous material were attached to the magnet to make fine adjustments to the uniformity. Unfortunately, at the end of the first shimming procedure, a fault developed on the upper large coil which had to be replaced. The upper coil was lifted from the magnet using a winch and a frame constructed around the magnet, then tipped onto its side so that it could be rolled out of the room. The replacement coil was installed in the same way. When the magnet was shimmed once more, a uniformity of 126 ppm in a 44 cm diameter disc was obtained. After the magnet had been shimmed, the RF shield was fitted and the couch was fixed into position.

**Operation of the imager**

The orientation of the coils of the magnet produces a vertical magnetic field, so the fringe field is greatest above and below the magnet, which in our case presents no problem. Magnetic field contours, as low as 0.0005 T, were plotted using a gaussmeter (Fig. 2). It

**Machine installation**

Before the magnet was delivered, the refrigeration unit was placed in position on a concrete plinth outside the NMR room using a large crane. The suppliers of the unit installed the plumbing necessary for the magnet and power cooling system. A closed-loop water system using 22 mm copper pipe was installed, linking the magnet, power supply and refrigerator with junctions in the flow pipe connected to the 22 mm mains supply, and in the return pipe connected to a pipe which ran to waste. Stopcocks can either isolate the refrigeration unit or the mains and waste pipes, so that it is possible to switch easily from one cooling system to the other.

The magnet was delivered in one piece with the four coils on their supporting frame and the patient tube in position. The magnet was lifted from the lorry, then manhandled along a corridor and into the NMR room. Part of the wall of the room was removed to allow access. The base of the RF shield had already been fixed into position and after the magnet had been pushed into place the castors at the base of each leg of the frame were removed. The magnet stands directly on a concrete floor which forms the foundation of the building.

The magnet operates at 332 A and 100 V when producing a field strength of 0.08 T. To obtain a uniform magnetic field at the centre of the magnet it had to be shimmed. Before this was done, all the cabinets, computers and couch motor assembly, in fact everything which contained ferrous metal, were sited in their permanent positions as shown in Fig. 1. The process of shimming the magnet involved making small adjustments to the relative positions and orientations of...
can be seen that there is circular symmetry of the low field contours even though there is ferrous material in the vicinity of the magnet. The 0.0005 T (5 gauss) contour is often quoted to signify the "limit" of the fringe field though this value is far lower than the field strength that might be hazardous to patients with pacemakers or aneurysm clips.

The refrigeration plant successfully maintains the magnet temperature between 23°C and 25°C. The run-to-waste cooling system also works successfully, keeping the magnet at a temperature of about 26°C with a constant water flow of 20 l/min.

The imager is designed with a "standby" mode of operation. This can be used at night to control the current supplied to maintain the magnet at any desired temperature with no water cooling. This is designed to ensure that the core of the magnet is at the correct temperature first thing in the morning for imaging. There is also an NMR feedback device which uses a small NMR probe in the imager to adjust the current and voltage and so maintain a stable field. It is our experience, however, that everything could be switched off at night and that the NMR imager could be used half an hour after switching on in the morning, even with the NMR feedback device disconnected.

**Cost of Installation**

The cost of the NMR imager with the independent diagnostic console was £939,000, including three years' warranty. Edinburgh received the first imager manufactured by the company and as a consequence, it was expected that there would be "teething troubles" with the imager and that any new developments would be evaluated in Edinburgh. The price paid for the system may not therefore reflect the price of subsequent systems.

The cost of site preparation was £6,500. The largest item of the cost (£5,000) was for the installation of a three phase electricity supply to the NMR Unit and the associated wiring for the electricity supply to the power racks and computers. The cost incurred was less than if an X-ray CT body scanner had been installed rather than an NMR imager, expensive radiation protection measures not being required for the latter.

**Summary**

It has been demonstrated that a relatively low-cost resistive NMR imager can be installed in a normal hospital environment without many major or expensive modifications. The magnet can be adjusted to give adequate uniformity and there is sufficient RF shielding to give good quality clinical images (Fig. 3). The fringe field of the magnet of this system, which operates at the lowest field strength of any commercial NMR imager, does not present a problem to imaging unit staff (NRPB, 1984).

The long term reliability and detailed specifications with regard to image quality have yet to be determined. These will be determined whilst the imager is being used.

Images of the normal head and body using the two types of RF coils: (a) proton density, (b) inversion recovery, (c) calculated $T_1$. 

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for clinical studies as part of the national clinical evaluation of NMR imaging supported by the Medical Research Council.

Acknowledgments
The authors would like to thank Professor Robson and members of staff in the University Department of Medicine for their support and forebearance with the disruption associated with the installation. We also thank Mr. A. Lough, Mr. A. Brown, Mr. J. Walker and Dr. H. Brash for their assistance with the installation together with the staff of M & D Technology Ltd. We thank Miss C. N. Taylor for secretarial assistance. We are indebted to the Lothian Health Board, M&V Trust, MRC, SHHD, University of Edinburgh and the Wellcome Trust for their generous financial support.

References

Book review
Environmental Radioactivity, NCRP Proceedings No. 5, pp. vi + 284, 1983 (National Council for Radiation Protection and Measurements, Bethesda, Md. 20814 USA), $17.00.
ISSN 0195–7740; No. 5
This publication contains the scientific and briefing sessions of the 19th annual meeting of NCRP and the text of the 7th Lauriston Taylor Lecture given by Merrill Eisenbud.
A wide range of topics was covered in the scientific session, including papers on the role of the environment and of natural and man-made sources within it, assessments of radioactivity at the Nevada test-site, in the atolls of the Northern Marshall Islands and at the site of the Three-Mile-Island incident. There were also contributions on environmental modelling, the particular problems associated with long-lived radionuclides, and the ever-present problem of waste disposal. The concluding paper of this section summarised and discussed the current criteria for setting dose-limits for the public.
Merrill Eisenbud’s lecture, “The human environment—past, present and future” was an interesting and stimulating account of environmental pollution, taken in a fairly broad sense. In contrast to many recent papers on this topic, his treatment was not gloomy, nor were his conclusions pessimistic. It is a pity that his time limit only allowed him to touch on relatively few aspects.
The scientific briefing session contained five papers: mostly reviews of published reports, including that of the NRPB on the Windscale release, the OECD report on the biological behaviour of plutonium and allied transuranic elements, the 1982 UNSCEAR document and two upon USA legislation in the fields of protection and radioactive waste. The proceedings closed with some selected task-group reports.
As usual, the papers are well presented, and continue to maintain the high standard we have come to expect from this Council’s publications. There is much in this volume that will be of interest to those engaged in radiation protection and to environmental health physicists.

John R. K. Savage
THE LANCET, June 1985

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BRAIN WATER IN CHRONIC ALCOHOLIC PATIENTS MEASURED BY MAGNETIC RESONANCE IMAGING

Sir,—In magnetic resonance imaging (MRI) the T1 relaxation time is related to the state of water in the tissue. The T1 values, measured in milliseconds, increase as the proportion of free to bound water increases. MRI is therefore a potentially valuable technique in the study of brain water during withdrawal of alcohol in chronic alcoholic patients. It has been used previously in such a study1 which suggested that there was "decreased free water during intoxication and an increase in brain water during alcohol withdrawal", a somewhat surprising result that the authors admitted was "at variance with current clinical practice which emphasizes the role of dehydration in the syndrome and encourages fluid replacement". We have repeated the study in a larger group of patients, since the result could have implications for the treatment of alcoholism.

The subjects were 9 chronic alcoholic patients (7 male, 2 female, aged 20–60 years, mean 46) voluntarily admitted to hospital for detoxification. They had been drinking at least 130 g ethanol per day (mean 180 g per day) before admission and had a 5–15 year history of alcoholism. None was clinically malnourished, or had physical disorders other than raised liver enzymes in serum. Over the first 5 days each patient received a normal diet, intramuscular vitamin B supplement, and decreasing doses of diazoxide. 9 normal controls, sex and age matched (range 26–62 years, mean 45), were chosen from hospital staff; their alcohol consumption ranged from total abstinence to the equivalent of 80 g ethanol per week.

Measurements were performed using a 0.15T resistive MRI system. All patients were investigated on two occasions, at 24 h and at 7–21 days after cessation of alcohol consumption; controls were investigated once. Measurements of T1 were made from a calculated T1 image of a 12 mm thick transverse section 10 mm above the maximum diameter of the lateral ventricles. To define the T1 in grey and white matter in the frontal, parietal, and occipital regions small regions of interest were defined—for grey matter 20 mm3 and for white 69 mm3—and the mean of similar regions in the left and right hemisphere was noted. In addition the T1 value over the whole brain, excluding the cerebrospinal fluid (CSF), was measured. The precision of T1 estimation was estimated from four repeated measurements on 3 normal volunteers over a period of a month. The T1 precision of grey matter was 2.8%, grey matter 4.9%, and the whole brain, excluding CSF, 2.6%.

At 24 h T1 over whole brain was higher in alcoholic patients than in controls (see figure). It then falls over the next 7 to 21 days. The change in T1 in the whole brain in alcoholic patients correlated with the time between abstinence and the second measurement (r = −0.81, p<0.01). The mean decrease in T1 in alcoholic patients
in grey matter was 10.1 ms (SE = 6.9 ms) whereas the decrease in white matter was only 0.6 ms (SE = 4.0 ms). The decrease in T1 in grey matter correlated with the duration of abstinence (r = 0.76, p < 0.05). Though the decreases in T1 over whole brain and in grey matter in alcoholic patients were not significantly different from zero, the reduction is significantly different (p < 0.05) from the increase previously reported by Besson et al.2

The preliminary results of this study suggest that chronic alcoholic patients have a higher T1 value in the brain than do normal controls. This raised T1 probably reflects an increase in free water within the brain. During withdrawal from alcohol the T1 drops, probably because of a decrease in the brain water content. Our findings contradict those of Besson et al2 but are consistent with the generally accepted hypothesis that the brain becomes excessively hydrated during chronic alcohol consumption and that abstinence results in dehydrolysis of the brain.

We intend to determine in more detail the differences between T1 in alcoholic patients and normal volunteers, whether the subsequent reduction in T1 is significant, and if so whether it returns to normal values with prolonged abstinence. A larger number of patients will enable the apparent difference in response between grey and white matter to be explored.

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ENDOCRINE ADJUVANT THERAPY FOR BREAST CANCER

SIT,—The six-year follow-up analysis by the Nolvadex Adjuvant Trial Organisation (NATO) of their comparison of tamoxifen with no adjuvant therapy after surgery for early breast cancer (April 13, p 856) focused on oestrogen receptor (ER) content of the primary tumours and on patterns of first relapse. Data from the Ludwig Breast Cancer Trials III and IV,1 in which one year of daily prednisone 7.5 mg and tamoxifen 20 mg was compared with no adjuvant therapy after surgery for node-positive premenopausal breast cancer, also cast light on these two issues.

In the Ludwig trials, 32% of patients whose ER status was known had ER values below 10 fmol/mg (taken as the threshold of ER-negative in previous publications) and 23% had ER values below 5 fmol/mg, compared with 37% in the NATO trial. The higher frequency of ER-negative tumours in the NATO trial might be due to the (few) premenopausal patients included, the limited tumour material available from node-negative patients, or the loss of receptor content during harvesting and transport of the specimens to the laboratory. The NATO investigators suggest that receptor loss during handling would have yielded a higher percentage of patients whose tumours contained less than 30 fmol/mg than that observed. The higher percentage of patients in the NATO trial with ER measurements below 5 fmol/mg cytosol protein almost certainly included some whose tumours in fact contained higher levels of receptor than that measured and who therefore benefited from adjuvant endocrine treatment.

The investigators provide an analysis of events by first site of recurrence and report that tamoxifen appeared to prevent both local/regional and distant disease. This observation is in contrast to the findings in the Ludwig trials III and IV, which indicated a significant reduction of local and regional recurrences but not of distant metastases.12 The raw recurrence data presented by NATO show that the frequency of distant events was reduced by 17% (15-1% in the tamoxifen-treated group and 18-5% in the controls). On the other hand, the incidence of local/regional relapses (also including second primaries which are reported together) was reduced by 46% (7-5% in the tamoxifen group and 13-8% in the controls). The table analyses type of recurrence by time in the Ludwig trials. The reduction in mastectomy scar recurrences appeared primarily within the first two years after randomisation, while significant control of regional recurrence and some impact on distant spread was only apparent beyond the first two years. The two-year duration of the tamoxifen treatment in the NATO trial—in contrast to only one year in the Ludwig trial—could therefore account for the early evidence of a survival benefit in the adjuvant-treated group. We do feel, along with our NATO colleagues, that long-term follow-up is needed for a better comprehension of the real effect of adjuvant endocrine treatment in early breast cancer.

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OSTEOGEN RECEPTOR STATUS AND CELLULAR COMPOSITION OF BREAST CANCERS

SIT,—In the continuing debate (April 27, p 876) about the predictive value of oestrogen receptor (ER) concentrations in the management of breast cancer13 we risk losing sight of a fundamental flaw in conventional ER assay methods. ER concentrations can now be measured precisely: regrettably the name “prediction” has not always been applied to the assessment of the predictability of the assay tumour sample. Indeed, the tissue homogenisation required for conventional assays prohibits any assessment of cellularity or even verification that visible tumour is present in the sampled tissue.

Variations in tumour epithelial cellularity (clinically significant) is assumed to reside in the epithelial compartment of breast

<table>
<thead>
<tr>
<th>SITE OF FAILURE</th>
<th>Failure at &lt;2yr</th>
<th>Failure at &gt;2yr</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>O/E</td>
<td>O/E</td>
</tr>
<tr>
<td>Mastectomy scar alone</td>
<td>24</td>
<td>0.64</td>
</tr>
<tr>
<td>Contralateral breast alone</td>
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<td>0.21</td>
</tr>
<tr>
<td>Regional lymph nodes</td>
<td>22</td>
<td>0.61</td>
</tr>
<tr>
<td>Distal</td>
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<td>0.92</td>
</tr>
<tr>
<td>2nd primary (not breast)</td>
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<td>1.00</td>
</tr>
<tr>
<td>Death without recurrence</td>
<td>15</td>
<td>1.39</td>
</tr>
<tr>
<td>Total failures</td>
<td>152</td>
<td>0.81</td>
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</tbody>
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P + T | = adjuvant endocrine therapy (prednisone + tamoxifen) (n = 320); control = observation only (n = 309); O/E = observed number of events; O/E = ratio of observed to expected number of events.
Nuclear magnetic resonance in hypertrophic cardiomyopathy

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SUMMARY The large differences in the spin lattice relaxation times ($T_1$) of blood and myocardium (measured by nuclear magnetic resonance) allow the heart to be visualised without the use of contrast media. The findings using nuclear magnetic resonance in 11 unselected patients with hypertrophic cardiomyopathy were compared with those in equal numbers of normal subjects and patients with electrocardiographic features of left ventricular hypertrophy. In patients with hypertrophic cardiomyopathy characteristic septal hypertrophy was noted together with variable and sometimes pronounced hypertrophy of the left ventricular free wall, which is consistent with the heterogeneous nature of this disease. The mean (SD) ratio of septal to free wall thickness was 1-5(0-8) for patients with hypertrophic cardiomyopathy, 0.8(0-2) for those with left ventricular hypertrophy, and 0-9(0-2) for normal subjects. Although septal measurements by nuclear magnetic resonance were greater than those obtained by echocardiography there was a significant correlation between the two. Septal and free wall area were significantly smaller in normal subjects. There were no differences in septal or free wall $T_1$ values between the three groups.

Non-gated nuclear magnetic resonance can detect septal and free wall hypertrophy. With the addition of multiple slice acquisition, rapid estimation of myocardial mass will be possible allowing the potentially important assessment of progression or regression of myocardial hypertrophy.

Hypertrophic cardiomyopathy is characterised by hypertrophy of myocardium, particularly of the interventricular septum, without apparent cause. Distinct histological appearances—such as cardiac muscle cell disorganisation, again mainly of the septum—have been reported. The diagnosis may be confirmed by echocardiography if the typical features (systolic anterior motion of the mitral valve, early aortic valve closure, and disproportionate thickening of the septum relative to the left ventricular free wall) are present. When few of the features are present or if echocardiography is technically difficult than doubt remains. In 1982 Goodwin suggested that angiocardiography was still the most useful investigation.

Nuclear magnetic resonance can be used for both non-invasive visualisation of cardiac structures and tissue characterisation; the second can be achieved by using the spin lattice relaxation time ($T_1$). The large differences in the $T_1$ values of myocardium and blood allow clear separation between cardiac muscle and cavity without the use of contrast medium.

Using nuclear magnetic resonance we therefore examined the appearances of hypertrophic cardiomyopathy by measuring septal and free wall thickness and contrasting $T_1$ values with those found in concentric left ventricular hypertrophy and normal hearts.

Patients and methods

SUBJECTS Three groups were examined. The first consisted of 11 unselected patients attending a cardiac review clinic with an established diagnosis of hypertrophic cardiomyopathy based on clinical, echocardiographic, and in two cases angiographic findings (Table 1). The second group consisted of 11 patients with left ventricular hypertrophy based on the electrocardiographic criteria of Romhilt and Estes,4 of whom nine had World Health Organisation stage III hypertension, one aortic stenosis, and one aortic coarctation. The third group consisted of 11 age matched healthy

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Nuclear magnetic resonance in hypertrophic cardiomyopathy

Table 1  Echocardiographic and nuclear magnetic resonance measurements in 11 patients with hypertrophic cardiomyopathy

<table>
<thead>
<tr>
<th>Case No</th>
<th>Age (yr)</th>
<th>M Mode echocardiography</th>
<th>Nuclear magnetic resonance</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Septal thickness (mm)</td>
<td>Posterior LV wall thickness (mm)</td>
</tr>
<tr>
<td>1*</td>
<td>50</td>
<td>14</td>
<td>16</td>
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<tr>
<td>2</td>
<td>55</td>
<td>24</td>
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<td>12</td>
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<tr>
<td>11</td>
<td>45</td>
<td>14</td>
<td>12</td>
</tr>
</tbody>
</table>

*Smiles.
LV, left ventricular; SAM, systolic anterior motion.

volunteers drawn mainly from hospital staff. In all cases consent was obtained and the study approved by our institute's ethical committee.

Nuclear magnetic resonance was carried out using a commercial (M&K Technology) low field (0.08T) resistive imaging system. Cardiac gating was not used. Three transverse sections with a slice of 16 mm were obtained from each subject with a separation of 20 mm between each section. The pulse sequence used was an interlaced saturation recovery and inversion recovery sequence each with a repetition rate of 1 s and a tau value of 200 ms for the latter. For each section four images were obtained at the end of the 4 min 16 s acquisition: proton density, calculated T1, inversion recovery, and a “difference” image (proton density-inversion recovery). The inversion recovery and difference images contain both T1 and proton density information. Of the four images, the calculated T1 image gave the best contrast between blood and myocardium. The long term precision of T1 estimation using this particular technique has been found to be 2-7% for a T1 value of 300 ms.

The section giving best definition between septum and left ventricular cavity was selected for each patient and studied in more detail. Initially, an estimate of the mean and range of T1 values for myocardium and ventricular blood in the 33 subjects. The mean (SD) T1 value for myocardium was 345(28) and for blood 635(124). The window on the T1 images was set so that all pixels with a T1 value >350 ms appeared uniformly bright enabling myocardium and blood to be clearly separated. Irregular regions of interest were drawn around the interventricular septum and free wall of the left ventricle using the computer. Computer derived measurements of septal and free wall thickness and area were made together with total cardiac diameter.

Echocardiography
M mode and real time echocardiography were carried out in the left oblique position within two weeks of the nuclear magnetic study. Septal and free wall diameters were measured from M mode recordings using an Irex-3 phased array system with a 2-5 MHz transducer in the groups with hypertrophic cardiomyopathy and left ventricular hypertrophy. Recordings were made at end diastole at a level just above the mitral valve. Septal and free wall thickness were measured in three consecutive cycles and the mean values noted for each subject. Five of the patients with hypertrophic cardiomyopathy showed systolic anterior motion of the anterior mitral leaflet and seven some degree of early closure of the aortic valve. None of these abnormalities was seen in patients with left ventricular hypertrophy. Two patients did not have echocardiographic features typical of hypertrophic cardiomyopathy, the diagnosis having been established at angiography (cases 1 and 5).

Statistical analysis
A non-paired Wilcoxon rank test was used for comparison between groups. Student's t test was used for comparison of T1 values between septum and left ventricular free wall with all groups combined.

Results
In eight patients with hypertrophic cardiomyopathy nuclear magnetic resonance showed characteristic appearances with pronounced septal hypertrophy and a variable degree of left ventricular free wall hypertrophy (Figs. 1 and 2). Table 1 shows the results and patient details for the group with hypertrophic car-
Table 2  Septal and left ventricular free wall spin lattice relaxation times ($T_1$) in three groups of patients. Values are mean (SD)

<table>
<thead>
<tr>
<th></th>
<th>Septal $T_1$ (s)</th>
<th>Free wall $T_1$ (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal subjects</td>
<td>374 (18)</td>
<td>331 (29)</td>
</tr>
<tr>
<td>Patients with ventricular hypertrophy</td>
<td>355 (31)</td>
<td>333 (22)</td>
</tr>
<tr>
<td>Patients with hypertrophic cardiomyopathy</td>
<td>352 (27)</td>
<td>342 (24)</td>
</tr>
</tbody>
</table>

Cardiomyopathy. In those patients with secondary left ventricular hypertrophy the free wall and septum appeared of equal thickness, but the muscle mass was greater than in normal subjects. The differences in area and dimension between patients with hypertrophic cardiomyopathy, left ventricular hypertrophy, and normal hearts (Figs. 3 and 4) are summarised below.

SEPTAL AND FREE WALL THICKNESS

There was considerable variation in the thickness of both septum and left ventricular free wall measured by nuclear magnetic resonance in the group with hypertrophic cardiomyopathy (Table 1). Although septal measurements were greater than those obtained with echocardiography there was a significant correlation between the two ($r=0.71, p<0.001$). The mean ratio of septal to free wall thickness measured by nuclear magnetic resonance and M mode echocardiography was similar at 1.5 (0.8) and 1.6 (0.6) respectively. The mean ratio of septal to free wall thickness as measured by magnetic resonance imaging was 0.8 (0.2) for patients with left ventricular hypertrophy and 0.9 (0.3) for normal subjects. In view of the variability of this ratio in patients with hypertrophic cardiomyopathy, there was no statistically significant difference detected between the three groups.

Fig. 1  Typical appearance of hypertrophic cardiomyopathy by nuclear magnetic resonance showing pronounced septal hypertrophy (arrow 1) (a) with mild left ventricular free wall hypertrophy (arrow 2) (case 10) and (b) with pronounced left ventricular free wall hypertrophy (arrow 2) (case 8).

Fig. 2  (a) Septal and free wall area and (b) ratio of septal to free wall area in individual patients with hypertrophic cardiomyopathy, those with left ventricular hypertrophy, and normal subjects.
Nuclear magnetic resonance in hypertrophic cardiomyopathy

SEPTAL AND FREE WALL AREA
There was a significant difference in septal area between normal subjects and both patient groups. Measured area in normal subjects was 2.7 cm² compared with 4.6 cm² (p<0.05) and 6.1 cm² (p<0.05) for patients with left ventricular hypertrophy and hypertrophic cardiomyopathy respectively. Corresponding values for left ventricular free wall area were 3.7 cm², 6.3 cm² (p<0.05), and 6.1 cm² (p<0.05).

TOTAL CARDIAC DIAMETER
The diameter as measured from the right ventricular border to the mid portion of the left ventricular free wall was greater in both patient groups at 138(10) mm and 130(14) mm for patients with hypertrophic cardiomyopathy and left ventricular hypertrophy respectively compared with 120(12) mm for normal subjects. The difference did not reach statistical significance.

SPIN LATTICE RELAXATION TIME (T₁)
Table 2 shows the mean values for T₁. There was no significant difference between patient groups, although septal T₁ was consistently higher than free wall T₁ for all groups (p<0.01).

Discussion
This study shows the ability of nuclear magnetic resonance to outline the interventricular septum and left ventricular free wall and to detect differences in myocardial thickness in patients with increased afterload and hypertrophic cardiomyopathy when contrasted with normal subjects. The pronounced contrast between flowing blood, which gives a high signal intensity, and the chamber walls allows hypertrophy to be demonstrated since there is also contrast between myocardium and surrounding tissues.

Within the group of patients with hypertrophic cardiomyopathy large variations in septal area were found. In those cases in which a diagnosis of hypertrophic cardiomyopathy could confidently be made by nuclear magnetic resonance the findings on echocardiography were more impressive (Table 1). Our unselected population of patients with hypertrophic cardiomyopathy represented the full spectrum of the disease, which is known to be heterogeneous. This variability in septal and free wall size was detected both by echocardiography and by nuclear magnetic resonance.

Steiner, in outlining the applications and potential of nuclear magnetic resonance in cardiac imaging, described the concentric nature of left ventricular hypertrophy in patients with hypertension and aortic stenosis and the non-uniform septal and free wall hypertrophy of hypertrophic cardiomyopathy. Although these patterns may occur in typical examples, it is important to recognise that both the extent and position of myocardial hypertrophy are variable as illustrated by nuclear magnetic resonance (Figs. 1 and 2).

Stone et al recently reported the appearances of hypertrophic cardiomyopathy obtained with non-gated computed tomography using contrast medium in nine patients. Visualisation of the septum was good and showed the variability of the position of maximal septal hypertrophy. Definition of the left ventricular free wall was, however, poor and inadequate for measurements. By using the differences in the relaxation index T₁ between flowing blood and ventricular muscle, nuclear magnetic resonance can allow clear visualisation of both septum and free wall without the use of contrast agents.

It was not the purpose of this study to compare nuclear magnetic resonance with echocardiography as there is a large difference in their cost and stage of development. There is also a wide variation in purchase and installation costs between different nuclear magnetic resonance systems and the variables they measure. The system we used is a low cost low magnetic field system. Echocardiography has contributed to the understanding of hypertrophic cardiomyopathy (for example, in determining the cause and level of obstruction when present) and is of value in making the diagnosis. Echocardiographic features are not, however, specific individually. Mode echocardiography gives diagnostic information in over 85% of patients, but precise measurements of left ventricular dimensions are possible in only 60-65% of patients. Real time echocardiography improves diagnostic ability, and because of its rapidity and widespread availability echocardiography will remain the commonest and most practical technique for confirming hypertrophic cardiomyopathy. Nevertheless, nuclear magnetic resonance is possible in most patients and may be helpful in those in whom echocardiographic quality is poor.

The differences observed between our three groups were from images taken without cardiac gating. The extent by which gating improves resolution in nuclear magnetic resonance is not yet established. Some reports have emphasised the need for cardiac gating to obtain satisfactory anatomical information. Choyke et al, however, have shown that images of diagnostic quality were obtained without gating using single spin echo sequences with a very short echo time (10-20 ms). In practice it gives a very similar echo time to that used in our own system. They had noted pronounced degradation of the image with longer echo times and concluded that the previous unfavourable reports using non-gated techniques may have been due to longer echo times used in other centres.
The ability to obtain measurements of myocardial area and thus by multiple slice acquisition to estimate myocardial mass may prove valuable in the follow up of patients with myocardial hypertrophy. Drug treatment in both patients with hypertension and those with hypertrophic cardiomyopathy should ideally result in a reduction in muscle mass. The non-invasive nature of nuclear magnetic resonance makes this an attractive method for long term assessment of such patients.

This work was funded in part by the Medical Research Council, Department of Health and Social Security, the Scottish Home and Health Department, University of Edinburgh, Melville, and the Wellcome Trusts. MB was in receipt of a Squibb Cardiovascular Fellowship.

References

Two Examples of CNS Lipomas Demonstrated by Computed Tomography and Low Field (0.08 T) MR Imaging

D. M. Kean, M. A. Smith, R. H. B. Douglas, C. N. Martyn, and J. J. K. Best

Abstract: The low field (0.08 T) magnetic resonance (MR) findings in two lipomas of the central nervous system (CNS), one adjacent to the corpus callosum, the other in the cervical cord, are described. The characteristic short T1 of fat and the image acquisition in the sagittal plane permit accurate localisation of these lipomas. Low field MR can provide images of diagnostic quality in a number of CNS lesions. Index Terms: Central nervous system, neoplasms—Nuclear magnetic resonance, techniques—Computed tomography.

Whereas the general trend seems to be toward higher field magnetic resonance (MR) imaging, low field imaging is being performed with satisfactory results in a number of centers. We report here on the usage of low field (0.08 T) imaging in the cases of fatty tumors of the central nervous system.

MATERIALS AND METHODS

Magnetic resonance investigations were performed using a commercial low field (0.08 T) resistive imaging system, manufactured by M & D Technology Ltd., which has been installed in the Royal Infirmary, Edinburgh. A section 12 mm thick was used and the pulse sequence was an interlaced saturation recovery and inversion recovery, each with a repetition time of 1,000 ms and inversion time of 200 ms for the latter. For each section a proton density image, inversion recovery image, and a calculated T1 image were obtained in a total acquisition time of 4 min 16 s.

CASE REPORTS

Case 1

A 32-year-old man was admitted after a seizure that had occurred while he was asleep. He had a previous history of seizures but had never received anticonvulsant therapy. Plain posteroanterior skull radiography showed a faint curvilinear calcification, deep in the left frontal region, parasagittally. Computed tomography (EMI CT 5005) (Fig. 1) showed a large lesion of low attenuation values in the region of the corpus callosum with a rim of calcification anteriorly. With MR the sagittal inversion recovery image (Fig. 2) showed a region of high signal intensity closely related to the corpus callosum. The axial calculated T1 image (Fig. 3) demonstrated the very short T1 of this abnormality. All features seen in both CT and
FIG. 2. Case 1. Sagittal inversion recovery MR image shows high signal intensity from lipoma.

FIG. 3. Case 1. Calculated T1 image shows short T1 of lipoma.

FIG. 4. Case 2. CT scan of cervical cord lesion.

FIG. 5. Case 2. Sagittal inversion recovery MR image shows high signal intensity from cervical lipoma.

FIG. 6. Case 2. Sagittal calculated T1 image shows short T1 of lipoma.
MR are in keeping with the diagnosis of lipoma adjacent to the corpus callosum.

**Case 2**

A 57-year-old man had a 6 year history of increasing spasticity of the right leg and unsteadiness of gait associated with more recent onset of clumsiness and loss of sensation in both upper limbs. Myelography demonstrated an intradural, extramedullary mass lesion in the cervical region (C2-C7). Figure 4 is a CT scan with intrathecal metrizamide, at the level of the C4 vertebral body. This showed the spinal cord compressed and displaced anteriorly by a large mass with low attenuation values. The sagittal inversion recovery MR image (Fig. 5) showed a large abnormality extending from C2 to C7. The T1 values, as demonstrated by a calculated T1 (Fig. 6), were very short, strongly suggesting that the lesion contained fat tissue. The CT and MR findings were in keeping with the diagnosis of lipoma. At surgery a lipoma that was both adherent to and incorporated within the cord was identified, biopsied, and subsequently shown to be a lipoma.

**DISCUSSION**

These two cases illustrate that images of good technical quality can be obtained using a low field MR system. Further, the advantages of MR are again demonstrated. In particular we may emphasize the ability to accurately delineate midline lesions by the use of sagittal plane imaging, and the ability to image spinal cord pathology without the injection of intrathecal contrast material. Finally, we have shown that measurements of magnetic relaxation times, in this case T1, may be useful in characterising lesions such as the two lipomas reported here.

**Acknowledgment:** We thank all our clinical colleagues for their continuing support and in particular Professor J. D. Miller, Mr. J. F. Shaw, and Dr. E. H. Jellinek whose cases are presented. We also thank Miss C. N. Taylor for typing this manuscript. We are indebted to the Department of Health and Social Security, Lothian Health Board, Medical Research Council, Melville Trust, Scottish Home and Health Department, University of Edinburgh, and Wellcome Trust for their generous financial support.

**REFERENCES**


CHARACTERISATION OF ACUTE MYOCARDIAL INFARCTION BY GATED MAGNETIC RESONANCE IMAGING

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R. H. B. DOUGLAS D. M. KEAN
J. J. K. BEST A. L. MUIR

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Summary

In studies with gated nuclear magnetic resonance imaging, patients with recent transmural myocardial infarction showed significantly longer spin-lattice relaxation times (T₁) in the infarct area than did healthy volunteers or patients with non-ischaemic or chronic ischaemic heart disease. 10 of the 13 patients had a T₁ longer than that found in any healthy subject or in any patient with other heart disease. Changes in T₁ should prove useful in the assessment of interventions designed to limit infarct size.

Introduction

Proton magnetic resonance imaging is a non-invasive technique with which body structure can be visualised in multiple planes. It relies on the principle that protons in a static magnetic field will tend to recover to their original position after deflection by a radiofrequency pulse. Anatomical information may be obtained by measuring proton density, which will differentiate between tissues such as muscle and lung.

Measurements of differences in the spin-lattice relaxation time (T₁) and of the spin-spin relaxation time (T₂) allow detailed tissue characterisation. These relaxation parameters are influenced by the ratio of "free" to "bound" water within the tissue. Oedematous tissue would thus have a longer T₁ and T₂ than normal tissue.

In dogs T₁ and T₂ are longer in infarcted than in normal myocardium in vitro. Differences in T₁ relaxation times in acute myocardial infarction in man have not been recorded, although an increase in T₂ after acute myocardial infarction has been reported. The prognosis after myocardial infarction depends on the extent of myocardial damage and residual left ventricular function. Current emphasis on intervention in myocardial infarction designed to limit infarct size highlights the need for techniques that can localise and define ischaemic and infarcted myocardium. Since current methods of assessing myocardial infarct size are imprecise, require repeated blood sampling, or involve the use of radioisotopes, we have explored the use of gated T₁ images to identify the position and size of myocardial infarcts in man.

Patients and Methods

Patients

13 patients with a recent transmural myocardial infarction diagnosed from electrocardiographic (ECG) evidence and the finding of raised levels of heart muscle enzymes were studied 2–12 days after the onset of infarction. 8 patients had ECG evidence of anterior, 3 of inferior, and 1 of predominantly lateral infarction, and in 1 patient the site could not be determined from the electrocardiogram. 6 patients received standard therapy in the coronary-care unit for the first 48 h, and 7 received intravenous thrombolytic therapy. A further 2 patients with subendocardial infarction were also studied. The age range of patients was 37 to 68 years.

Measurement of T₁

The T₁ image was transferred to a diagnostic console and displayed in colour corresponding to the range of T₁ values. The overall visual appearance was noted, and any obvious abnormalities such as wall thinning, pericardial fluid, or obvious area of increased myocardial T₁, were recorded. For all studies the left ventricular wall was divided into six areas for both transverse and coronal scans (fig 1), and the T₁ for each region was measured.

Results

The appearance of normal myocardium was of a uniform T₁ (fig 2), which was seen in both the healthy volunteers and patients with non-ischaemic cardiac disease. The mean T₁
was 305±25 ms for normal volunteers, which was not significantly different from the T₁ values for the patients with non-ischaemic cardiac disease or old infarction but was lower than for recent infarcts where the mean T₁ was 329±60 ms (p<0.01).

In the group with recent transmural infarction the most dramatic finding was of distinct areas of increased T₁ in the left ventricular wall (table I). The precise area of myocardium involved varied from a localised region of prolonged T₁ (fig 2) to a more diffuse area of patchy increase in T₁. Differences in maximum T₁ values between the groups were greater than differences in mean T₁ (table II). For patients with recent infarction the maximum T₁ was 426±67 ms. This compares with a maximum T₁ of 352±30 ms for old infarct (more than 2 months) (p<0.05), 316±9 for non-ischaemic cardiac disease (p<0.01), and 339±17 for normal volunteers (p<0.02). 10 of the 13 patients with recent transmural infarction had a region of myocardium with a T₁ longer than 390 ms, whereas none of the other groups contained any areas of myocardium with a T₁ longer than 390 ms. There was no significant difference in the prolongation of T₁ between those treated with and without thrombolytic therapy. Of the 3 patients with recent infarction who did not show areas of increased T₁, all had only one or two sections scanned and it is possible that the region of infarction may not have been included. No regions of high T₁ were seen in the 2 patients with subendocardial infarction. Unsuspected pericardial fluid was noted in 2 patients, pleural effusion in 1, and suspected early aneurysm formation in 1. Patients with previous myocardial infarction had changes such as ventricular dilatation and wall thinning but no areas of increased T₁. A degree of wall thinning was also seen in 3 patients with recent infarction. In patients with inferior myocardial infarction affecting the inferior border of the left ventricle, the area of increased T₁ was often best appreciated on the coronal scan, although it was present on the lowest transverse section as a very broad area of raised T₁.

**Discussion**

Using a magnetic resonance imager that gives accurate measurements of the relaxation parameter T₁, we have demonstrated that recent myocardial infarction can be visualised in man. We found an average 40% increase in T₁ in areas of suspected infarction in patients imaged between 2 and 12 days after the onset of infarction. A smaller percentage increase in T₁ was reported by Williams et al. in biopsy specimens of ischaemic tissue from dogs within 30 min of experimental coronary occlusion. Our findings of increased T₁ with recent but not with old infarction are in keeping with the in-vitro measurements of Brown et al., who showed prolongation of T₁ in canine myocardium 3 h and 4 days after
coronary occlusion, but at 21 days $T_2$ was more variable, and by 56 days it was shortened.

Higgins et al. found a 14% prolongation of $T_2$ and a good correlation with the percent water content in an experimental model of myocardial infarction and have confirmed these findings in man. In dogs, Westby et al., having demonstrated a change in spin echo signal intensity in vivo, reported 69% prolongation of $T_2$ in vitro. It is not yet possible to say whether $T_1$ or $T_2$ is superior either in vitro or in vivo. Most magnetic resonance imaging systems are designed so that images of high spatial resolution can be obtained, less emphasis being placed on precise estimation of $T_1$ or $T_2$. In such systems the spin echo pulse sequence is generally used for cardiac imaging because it is technically easier to gate and because it produces images of good spatial resolution. Using a system such as this, $T_1$-weighted images can be obtained and $T_1$ and $T_2$ values can be calculated, $T_1$ with generally poorer accuracy and precision than $T_2$. However, our method, using inversion recovery, provides a more precise value for $T_1$.

Whether $T_1$ or $T_2$ will in the end be the more useful remains to be determined.

The ability to obtain serial sections through the heart suggests that it should be possible to obtain an accurate measure of infarct size with gated images in man. This is of particular importance in assessing interventions designed to limit infarct size. Thrombolytic therapy causes recanalisation and improves blood flow and ventricular function but may cause reperfusion damage or haemorrhagic infarction. Our studies have not revealed a significant difference in $T_2$ values between patients treated conventionally and those with documented coronary reperfusion, but the varying time of imaging from onset of infarction make a more detailed study necessary. With nuclear magnetic spectroscopic analysis Slutsky et al. found an increase in $T_2$ in excised dogs' hearts after 3 hr of ischaemia and a slight further prolongation with reperfusion. Brown et al. found a differential effect of varying lengths of occlusion and reperfusion on $T_2$ values in canine hearts. With measurement of $T_1$ it should be possible to follow the natural history of myocardial infarction and to determine the effect of intervention therapies, including coronary reperfusion.

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REFERENCES


NINDIUM-111 LABELLED MONOCLONAL ANTIBODY TO PLACENTAL ALKALINE PHOSPHATASE IN THE DETECTION OF NEOPLASMS OF TESTIS, OVARY, AND CERVIX

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Summary

A monoclonal antibody (H17E2) against placental alkaline phosphatase (PLAP) and testicular placental-like alkaline phosphatase was labelled with indium-111 and used in radioimmunoassay of 15 patients known or suspected to have germ-cell carcinoma of the testis or carcinoma of the ovary or cervix. Good images of neoplastic lesions were obtained in most patients with active disease. In 1 patient with testicular teratoma and elevated human chorionic gonadotrophin who had a normal computed tomography scan, the labelled antibody located microscopic disease in a lymph node, which was then removed. No false positive localisation was seen in patients with PLAP-negative tumours or sites of inflammation. This method may be helpful in the diagnosis, staging, and monitoring of PLAP-positive neoplasms of testicular, ovarian, and cervical origin.

Introduction

RADIOIODINATED polyclonal1,2 and monoclonal antibodies3,4 have been shown to localise successfully in a wide range of neoplasms. In-vivo dehalogenation and antibody cross-reactivity with healthy tissues are limitations of this method and may explain the low levels of radiiodine detectable in resected tumours.5,6

We selected a new monoclonal antibody (H17E2)7 against placental alkaline phosphatase (PLAP) and testicular placental-like alkaline phosphatase which shows high reactivity against PLAP-positive tumours and very low, if any, reactivity against healthy tissues.7,8 For radiolabelling we used indium-111, which has been shown to produce better tumour-to-healthy-organ (except for liver) ratios than those with radiiodine.9-16

Patients and Methods

15 patients, aged 21-70 years (mean 40 years), with testicular (n=8), ovarian (n=5), and cervical (n=1) neoplasms were studied. Written, informed consent was obtained from all patients, and they were skin tested for allergy to mouse antibodies before injection.

Monoclonal Antibody H17E2

This mouse IgG1 reacts with placental alkaline phosphatase as well as the leucine inhibitable form of alkaline phosphatase found at low levels in the healthy testis and which cross-reacts with the

M. BEEN AND OTHERS: REFERENCES—continued


References continued on next column...
BRAIN WATER MEASURED IN VOLUNTEERS AFTER ALCOHOL AND VASOPRESSIN

Sir,—We have reported that brain water levels in chronic alcoholics are raised during the acute phase of withdrawal and then decrease over 7–21 days of abstinence. We used magnetic resonance imaging (MRI), the T₁ relaxation time being related to the amount of free water in tissues.² These findings are consistent with the hypothesis that the brain, in the initial stage of withdrawal, is overhydrated. Vasopressin may have a role in mediating these changes.³ We have attempted to reproduce these changes in healthy volunteers.

Four male and one female volunteers, mean age 32 (range 26–40), with an average weekly consumption of 16 units of alcohol (range 5–42) had an NMR scan before and 45 min after the consumption of half a bottle of spirits (14 units). The alcohol was consumed over 1½ hours. This produced blood alcohol levels of 143–197 mg/l at the time of the second scan and caused a mean increase in plasma osmolarity of 12 mmol/kg (after correction for alcohol). We used a 0.8T Siemens benchtop MRI system.

Measurements of T₁ were made from a calculated T₁ image of a, scan thick transverse section 10 mm above the maximum of the lateral ventricles. Measurements were made for the left and right hemispheres of the whole section, excluding CSF, and for individual small areas of grey and white matter in the frontal and occipital regions. Statistical significance of results was calculated using Wilcoxon's paired rank test.

There was a significant decrease in T₁ of the whole section (mean 4.7 ms, p<0.05) (table). In white matter there was a significant decrease when frontal and occipital measurements were analysed together (t=−3.9 ms, p<0.05) and there was a similar trend in grey matter (t=−2.6 ms, NS). The precise variation of T₁ in grey matter is poorer than that of white matter or of the whole brain.² The change in T₁ is consistent with the observation that alcohol inhibits vasopressin and this promotes diuresis and dehydration in the short term.

Three further volunteers (all male, aged 30, 31, and 32 yr) were then scanned daily at 0900 hr for 4 days. Day 1 acted as a baseline and the volunteers took vasopressin 20 µl twice daily for the next 3 days. This produced a fall in mean sodium from 140 to 135 mmol/l and in mean plasma osmolarity from 292 to 281 mmol/kg. The T₁ of frontal and occipital white matter increased by a mean of 4.1 ms (consistent with an increase in free water). This increase just failed to reach significance at the 5% level. No general trend was noted in the grey matter or whole slice measurements.

Thus we have demonstrated that we can alter T₁ acutely and in the direction we would expect if the change is due to alterations in the amount of free water in the brain tissue, although the magnitude of the change is about half of that seen in chronic alcoholics.³ This supports the view that changes in T₁ during withdrawal are due to dynamic changes in water balance.

Hydroptic overhydration has been noted in alcoholics and so have decreased levels of vasopressin in plasma and CSF.⁴⁻⁶ Eisenhofer and Johnson⁷ found that in normal volunteers early suppression of vasopressin is followed by raised levels as the increasing plasma osmolarity overcomes the direct suppression. Perhaps the mechanism of vasopressin release becomes reset in chronic alcoholics. These and other studies⁸ suggest that water retention, induced by raised vasopressin levels, may be responsible for some of the clinical features of alcohol withdrawal.

NMR is a safe, non-invasive tool for monitoring changes in tissue hydration, and we plan to look next at changes in total body water, especially after consumption of 6 units of alcohol.

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CYSTIC LEUCOMALACIA IN PRETERM INFANT: SITE OF LESION IN RELATION TO PROGNOSIS

Sir,—In your July issue (p 137) we reported the very poor outcome in infants with periventricular leucomalacia (PVL) compared with those who had large intraventricral haemorrhages. We did not distinguish between lesions confined to the periventricular area and those extending further into the white matter. We have since studied another 7 cases and have become aware of the prognostic importance of the distribution of these lesions (figure). We suggest that the term "periventricular" should be restricted to lesions adjacent to the ventricles in the area of the centrum semiovale, trigone, and optic radiation. Lesions further from the ventricles, at the depth of the sulci in the subcortical region, should be classed as "subcortical leucomalacia" (SCL), on the basis of the post-mortem angiographic studies of Takashima et al. Lesions are prominent in both the subcortical and periventricular regions would be classed as "mixed".

Of the 17 infants studied so far (including the 10 previously reported) 9 had extensive lesions in the periventricular area, 4 in the subcortical region, and 4 were mixed. Although serial ultrasound studies showed the same basic pattern for all lesions, in SCL the areas of increased echogenicity tend to occur earlier in the insult and were especially marked around the interhemispheric fissure, and cysts occurred sooner and persisted far beyond 40 weeks' postnatal age. PVL was seen in preterm infants below 32 weeks' gestation while SCL tended to occur in older preterm infants. Takashima et al suggested that the distribution of the cystic lesions is related to the maturity of the vascular supply. Another suggestion is that the different energy metabolism in the various parts of the brain at

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ECG-gated $T_1$ images of the heart

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1. Introduction

Magnetic resonance imaging (mri) has been used to investigate heart disease, either studying abnormal anatomy or attempting to identify pathological alterations of myocardial tissue. The anatomical studies have used cardiac-gated saturation recovery or spin-echo pulse sequences with short time to echo (te), which produce reasonably high spatial resolution images containing mainly proton density information (Longmore et al 1985) or real-time echo planar imaging (Rzedzian et al 1983). For information concerning tissue, particularly important for the investigation of myocardial infarction (mi), it may be possible to identify abnormal pathology within the cardiac tissue using the relaxation times $T_1$ or $T_2$.

In vitro measurements of excised dog hearts have shown that both $T_1$ and $T_2$ increase following recent myocardial infarction (Williams et al 1980, Higgins et al 1983). The changes in $T_1$ correlate with changes in the water content of the tissue (Higgins et al 1983, Brown et al 1984, 1985) demonstrating that the initial increase in $T_1$ and $T_2$ is due to tissue oedema. The increase in $T_1$ is then reversed as the region of the heart wall is replaced by fibrous tissue (Brown et al 1984). Increases in $T_2$ in vivo following mi have been demonstrated in dogs (Wesbey et al 1984, Pflugfelder et al 1985) and in man (Higgins et al 1984, McNamara et al 1985). In one of these studies $T_1$ weighted images were also investigated but no increases in signal were detected (Pflugfelder et al 1985).

There is a wide variation in the results of the magnitude of the changes in $T_1$ and $T_2$ following infarction and also some disagreement concerning the relative merits of $T_1$ or $T_2$ in the assessment of mi. We feel that the suggested superiority of $T_2$ over $T_1$ is likely to be influenced by the fact that the mri systems used were not designed to give accurate or precise $T_1$ values (Pflugfelder et al 1985) or that the pulse sequence combination used was more appropriate for $T_2$ measurements than for $T_1$ (Higgins et al 1983). In order to assess the true value of in vivo $T_1$ measurements of mi in man a pulse sequence specifically designed for $T_1$ measurement with high precision has been gated with the cardiac cycle to produce gated $T_1$ images.

2. Methods

A commercial 0.08 T resistive system, installed in a hospital environment for clinical studies (Smith et al 1984), has been modified to produce cardiac gated $T_1$ images. The
The imager generally uses a fixed pulse sequence which produces a saturation recovery image, an inversion recovery image and a calculated $T1$ image for each section in a 4 min 16 s acquisition time. The imager uses an interlaced saturation recovery and inversion recovery pulse sequence with $TR = 1000 ms$ for both and $TR = 200 ms$ for the latter. One novel feature of the system, which makes it particularly suited to gated cardiac imaging, is the use of an adiabatic fast passage (AFP) pulse to invert the nuclei in the inversion recovery sequence (Hutchinson and Smith 1983). This pulse is very efficient at inverting the nuclei throughout the section imaged and it also has the advantage that it is not slice-selective in the inversion part of the sequence. This latter feature of the inverting AFP pulse is important because of the necessarily long delay between it and the subsequent 90° pulse, which is slice-selective, and which therefore must be synchronised with the cardiac cycle.

The timing of the pulse sequence is controlled by two timing integrated circuits, each with ten channels. In the normal mode of operation one of the timing channels controls the $TR$ interval, which can be fixed at a preset value. This counter works in a free running mode and is automatically re-armed. For the gating sequence, this channel is changed to one shot mode. A single pulse sequence can then be started under direct software control by re-arming the appropriate channels. The computer used in the MRI system is a Digital PDP 11/23+. The software was programmed to respond to a pulse occurring on one of the status lines and a socket was fitted so that the system could be triggered externally from either a cardiac or respiratory gating pulse.

To obtain an ECG pulse from a subject inside the NMR imager, care was taken to reduce interference on the ECG signal from the RF pulses and also to minimise the noise introduced into the system from any patient connections. It was decided to monitor the ECG and use the R wave to determine the position of systole and diastole in the cardiac cycle rather than the less precise method of using a peripheral pulse from a region such as the finger. It was found that two electrodes placed one on either wrist produced severe interference on the ECG signal from the RF pulse. ECG detection in a perpendicular direction, with one electrode placed on the right shoulder and the other electrode just above the right hip produced no measureable interference from the RF pulses on the ECG signal. Conventional ECG electrodes were used in this position as illustrated in figure 1, the leads being taped to the patient with Micropore to ensure that there were no loops to pick up RF interference. The cables from the two electrodes were twisted together and connected to a rechargeable battery-powered unit 1.2 m from the centre of the magnet which converted the ECG to optical pulses so that it could be transmitted along a 12 m optical fibre. The distance of the battery-powered optical fibre unit from the central region of the magnet was found to be important. If the unit were nearer the centre of the magnet the ferromagnetic material within the unit would degrade the image; if the unit were further away then the electrode leads would act as an aerial for the external RF interference. The optical fibre is connected to a unit to convert the ECG signal into a form that can be fed into a conventional ECG monitor (Kontron Micromon 7141), both of which are sited 4 m from the magnet. The monitor has been modified to produce a pulse immediately after the detection of the QRS complex.

The trigger pulse from the modified ECG monitor is fed into a microcomputer (BBC model B) which in turn produces a pulse which is fed into the MRI system computer to trigger a single NMR signal acquisition. The microcomputer controls the delay between the patient’s R wave and the part of the cardiac cycle to be imaged. A window setting can be used such that only R–R intervals within a certain range produce a
ECG-gated $T_1$ images of the heart

Figure 1. Schematic plan view of subject in MRI system. The positions of the ECG electrodes and leads and the battery powered box, converting the ECG to optical pulses, are shown in relation to the subject and the largest ring magnet.

Signal for the MRI system: thus ectopic beats are excluded. This is particularly important for end diastolic images as the section is not imaged immediately after the R wave, as is usually the case, but immediately before the succeeding R wave.

The timing of the $T_1$ gating sequence is illustrated in figure 2. There is a delay of 20 ms between the patient’s R wave and the trigger from the ECG monitor due to the QRS complex detection. The delay from this pulse to the start of the AFP inverting

Figure 2. Timing diagram for a gated $T_1$ image showing inversion recovery and saturation recovery pulses during the interleaved sequence. The trigger from the ECG monitor occurs 20 ms after the upstroke of the R wave. The delay between the trigger pulse and the AFP is controlled by the microprocessor and will be different for each subject. A time from inversion (ti) of 200 ms is shown. The NMR signal is a field echo occurring 10 ms after the $90^\circ$ pulse.
pulse is controlled by the microcomputer. The time from inversion (TR) can be altered in the main MR acquisition program but is generally set at 200 ms. The time from the trigger pulse and the 90° pulse is identical in both the saturation recovery and inversion recovery parts of the pulse sequence. Thus in both types of pulse sequence resonance occurs in the section at the same part of the cardiac cycle. The field echo occurs approximately 10 ms after the 90° pulse. The shortest delay available between the patient's R wave and the part of the cardiac cycle to be imaged is 230 ms, falling within systole even in patients with heart beat rates as fast as 120 min⁻¹. Signal acquisition can be triggered from every heart beat, though for accurate T₁ estimation every other heart beat is used in subjects with a heart beat rate faster than 70 min⁻¹.

A Perspex phantom was used to check the gated T₁ images. It contained three layers of CuSO₄ solution, the central layer having a T₁ of 330 ms and the outer layers a T₁ of 528 ms, and moved up and down with a maximum distance of travel of 5 cm. It was driven by a stepping motor controlled by a microprocessor which enabled either regular motion or motion which simulated an irregular R-R interval to be used. The microprocessor also produced a trigger pulse at the start of each cycle. The frequency of movement studied was 38–70 min⁻¹, equivalent to the frequency of trigger pulses obtained from human subjects, remembering that for heart beat rates greater than 70 min⁻¹ a trigger pulse is produced on every second beat.

3. Results

Results obtained from the moving phantom are given in table 1. It can be seen that the precision of the calculated T₁, as demonstrated by the standard deviation of pixel values within the region of interest used, is not degraded by movement. There is a slight variation in T₁ value with TR, as would be expected. However the variation is within the long-term reproducibility (coefficient of variation) of T₁ measurement which is 2.9 and 6.5% for T₁ values 330 and 528 ms respectively.

An example of a cardiac-gated T₁ image of a normal subject at end systole is illustrated in figure 3 with an ungated image of the identical section for comparison. The time to end-systole from the R wave was determined using echocardiography. The

<table>
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<tr>
<th>Frequency (cycles/min)</th>
<th>TR (ms)</th>
<th>T₁ + 2SD (ms)</th>
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<td></td>
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<td>Solution 1</td>
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<tr>
<td>38</td>
<td>1580</td>
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<td>47</td>
<td>1280</td>
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<td>56</td>
<td>1070</td>
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<td>65</td>
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<td>55–67</td>
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<td>50–75</td>
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<td>326 ± 32</td>
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<tr>
<td>Static Phantom</td>
<td>1000</td>
<td>330 ± 30</td>
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Figure 3. (a) Gated and (b) ungated transverse T₁ images of a normal human subject. The linear grey scale covers the range 50 to 650 ms.

ECG-gated T₁ images of the heart

section thickness used was 16 mm and the acquisition time for the gated image just under 4 min. The absence of signal from the anterior chest wall on the right-hand side is caused by the ferromagnetic material in the screening of the cable of the ECG connection. In the gated T₁ image the region of the left ventricular wall can be seen quite clearly with a uniform T₁ value. In the ungated image there is no such region of uniform T₁ corresponding to the ventricular wall; the T₁ value steadily increases from the outer edge of the heart to the centre. Eight small regions of interest were drawn encompassing the whole of the left ventricular free wall and the coefficient of variation of the T₁ values was found to be 3% in the gated image and 10% in the ungated image.

At end-systole the T₁ of the blood is fairly uniform. This is not the case in images later in the cardiac cycle, particularly during early diastole, where the T₁ of blood varies considerably. The cause of this is illustrated in figure 4 which shows the saturation recovery image at different stages during the heart cycle. It can be seen that the image acquired at the beginning of diastole contains a region of very low signal, caused presumably by the turbulence of the blood flowing into the ventricles. As the signal level from this image is the denominator in the equation to calculate T₁, this results in indeterminate values of T₁.

To date the technique has been used on 12 normal volunteers, many of these on more than one occasion, 13 patients with hypertrophic cardiomyopathy and 32 patients with my 7 within 4 days and the rest within 12 days of infarction. Occasionally the
position of the two electrodes makes it impossible to produce a trigger pulse from the cardiac monitor. On such occasions the electrode on the right hip is moved to the left side of the patient and the cable to the electrode on the right shoulder is supported away from the chest wall by foam pads so that no signal loss occurs in the region of the heart. The use of this arrangement has improved the ECG signal sufficiently to allow a trigger pulse to be produced in all but one patient. However, the position of the electrode on the left-hand side of the patient causes interference on the ECG from the RF or gradient pulses, depending on the size of the patient and consequent proximity to the gradient windings or RF coil, which can cause unwanted trigger pulses. This can be overcome by gating the signal acquisition on every second, third or fourth pulse, depending on the type of interference. In some instances this can result in a lengthening of the time for an acquisition of a single section.

Figure 4. Gated saturation recovery images of a normal subject (a) at end-systole, (b) during diastole, showing a region of low signal in the left ventricle as it is filling, and (c) towards the end of diastole.

Figure 5. Gated coronal $T_1$ image of a patient with recent MI. The region of increased $T_1$, corresponding to infarcted tissue, is indicated.
Patients with recent infarction find the procedure acceptable, a complete investigation of four transverse and one coronal section taking between 20 and 30 min. Regions of increased $T_1$ have been found in the ventricular wall in patients with a recent infarct (Been et al 1985) and an example is shown in figure 5.

4. Discussion

The ability to obtain ECG-gated $T_1$ images of sufficient quality and in a sufficiently short time for clinical use has been demonstrated. The commercial system which has been modified has been shown to be particularly suitable for obtaining such images. This technique will be of value in assessing the use of in vivo $T_1$ measurement to identify abnormal pathology within the ventricular wall, particularly in patients with recent infarction. Most other MRI methods of investigating cardiac disease rely on the identification of wall thinning or abnormal motion using gated spin-echo images, often presented in cine mode. Such techniques can take a long time, particularly if the complete left ventricle is to be studied, which may prove unacceptable for patients with recent infarction. Moreover, echocardiography is already a well established and readily available technique producing similar information.

Many patients with cardiac disease will have an irregular heart beat; even if ectopic beats are not present then the R-R interval will vary during an investigation. Such a variation can produce a significant error in the $T_1$ estimation when a gated spin-echo sequence with short $TR$ is used. In contrast, our use of a gated inversion recovery sequence with an inverting pulse that is not slice-selective ensures that the typical R-R variations obtained from patients have little effect on the $T_1$ values obtained.

It would be possible to obtain better linear spatial resolution by increasing the section width, thereby increasing the signal size, or by signal averaging, which would result in a considerably longer acquisition time. However, thicker sections have not been used as these would increase the partial volume effect, which increases with section thickness, as the heart is imaged in the transverse plane which itself is not perpendicular to the ventricular wall. To minimise the time that the patient is inside the MRI system signal averaging is not performed.

Future improvements to the system will be the availability of oblique sections so that the left ventricle can be imaged in a more appropriate plane, either perpendicular to the ventricular wall or along the length of the ventricle through the apex.

Acknowledgments

We are grateful to Mr Robert Chesser of M&D Technology Ltd and Mr David Norris of the Department of Biomedical Physics and Bioengineering, Aberdeen University, for valuable discussion. We would also like to thank Mrs C N Rowan for secretarial services. Financial support is acknowledged from the Lothian Health Board, Medical Research Council, Melville Trust, Scottish Home and Health Department, University of Edinburgh and Wellcome Trust.

References

In view of the absence of conclusive evidence to support the existence of non-thermal effects we retain our hypothesis that the applied heat dose, which is dependent upon both the temperature obtained and the exposure time, is the critical parameter. In this case extensive thermometry is required to ensure that the patient receives an effective adjunct treatment to radiotherapy.

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REFERENCES


The absence of tissue specificity in MRI using in vivo TI or T2 determination: true biological variation or technical artefact?

The Editor—Sir,
The discovery by Damadian (1971) that malignant tissue measured in vivo has a higher T1 than normal tissue acted as the stimulus for the development of NMR imaging techniques. In the beginning it was hoped that in vivo measurements of relaxation parameters may uniquely identify different tissues or the same tissue in different pathological states, perhaps enabling the differentiation between normal, benign or malignant tissue. However, by the time commercial magnetic resonance imaging (MRI) systems became available in about 1982, the emphasis in MRI system design was generally not on precise or accurate in vivo TI or T2 determination but rather higher resolution image production, the images being T1 or T2 weighted. It is generally accepted now that neither T1 nor T2 are disease-specific, there being too great an overlap for diagnostic purposes.

We feel that this conclusion may be erroneous. The spread of T1 or T2 values from a particular tissue may not be due solely to the biological variation but instead may be significantly expanded due to technical errors in their calculation. There are a number of possible sources of error in the in vivo relaxation parameter determination.

(a) The calculation of in vivo T1 from a saturation recovery and an inversion recovery pulse sequence will have an error if there is not efficient inversion in the latter sequence. This will occur in all imaging systems when a spatially selective 180° pulse is used or when there is significant radiofrequency inhomogeneity across the image plane. An exception is the system which uses an adiabatic fast passage RF pulse (Edelstein et al., 1980), which efficiently inverts all the nuclei.

(b) Some two-point methods of relaxation parameter calculation are known to be imprecise; for example, two saturation-recovery or two spin–echo sequences with different TR, or two inversion-recovery sequences with different TI to calculate T1 and two spin–echo sequences with different TE to calculate T2. More precise estimations using such sequences require a larger number of points each with a different TI or TE value to be used.

(c) The spin–spin and spin–lattice relaxation processes may not be characterised by a single exponential.

(d) The width of the tomographic sections used in MRI, particularly when values of T1, or T2 are being calculated, can be as thick as 20 mm. As a consequence there can be a significant partial volume effect. The partial volume effect can produce a considerably reduced estimate of the T1, or T2 in a pathological lesion which has a higher value than surrounding normal tissues unless the lesion was extremely large. Much thinner section widths must, therefore, be used for accurate in vivo T1, or T2 determination; the thickness of the section should be about half the diameter of any tissue under investigation and care must be taken to position the section through the centre of the tissue.

(e) It is recognised that cardiac and respiratory movement degrade the resolution of images; this can be avoided by the use of gating techniques. However, not only does it degrade image resolution, it will also produce an error in the calculation of T1 or T2 if the tissue is moving.

(f) There can be an error introduced due to subjective operator variation in the choice of region for T1 or T2 calculation.

Experience has shown that, as the techniques of in vivo T1 and T2 determination are improved, so the spread in their values from a particular type of tissue is reduced. Only when the technical errors of T1 and T2 measurement are eliminated can we hope to draw any firm conclusions concerning their specificity and consequent clinical use.
We would recognise that in many MRI systems it is not possible to eliminate all the sources of technical error, particularly as there is often limited time available for patient investigation. In the centres which adopt a pragmatic approach to T1 and T2 calculation and are investigating its use in the clinical environment, knowing that, for technical reasons, the values obtained may not be entirely accurate, we would suggest that such quantitative values should be defined as $T_1^{**}$ and $T_2^{**}$. The term $T_1$ and $T_2$ would be reserved for what is believed to be true biological relaxation times.

Yours, etc.,

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REFERENCES


Radioprotection by ascorbate in vitro: temperature-dependent uptake

THE EDITOR—Sir,

The role of glutathione in radioprotection has been much investigated, but relatively little attention has been paid to ascorbate, which is present at high concentrations in human tumours (Kakar & Wilson, 1974; Moriarty et al, 1977). Both glutathione and ascorbate can act as H-atom donors, although ascorbate is more efficient in electron transfer reactions, and both these reactions may be important in DNA repair processes (Redpath & Wilson, 1973; O'Neil, 1983; Wilson, 1983). There have been some reports in the literature of radioprotection by ascorbate in vitro (Redpath & Wilson, 1973; O'Connor et al, 1977; Baverstock, 1979) but other workers have found no effect of ascorbate on radiosensitisation by misonidazole (Koch et al, 1979).

As a part of an investigation into the radioprotective role of ascorbate in vitro, we have measured the uptake of ascorbate by V79 379A Chinese hamster cells incubated in Eagles' minimum essential medium +10% fetal calf serum at various temperatures. Figure 1 shows the concentration of ascorbate in cells plotted against time of incubation under N2 at 4°C, 21.5°C and 36°C. The point we wish to emphasise is the marked temperature-dependence and slow time course of the uptake, with essentially no uptake at 4°C, and intracellular concentrations increasing for several hours at the higher temperatures. In some radiobiological experiments the pre-irradiation incubations are carried out at 4°C in order to reduce toxicity. If the agent of interest is actively transported, or is taken up slowly, both of which appear to apply to ascorbate, a lack of effect could arise from poor intracellular uptake rather than inactivity per se. Thus, Koch et al (1979) showed that ascorbate at 5 mM had no effect on radiosensitivity when the pre-gassing was carried out at 4°C. We have recently shown that if pre-incubation is at room temperature, then ascorbate can afford significant protection in both control and GSH-depleted cells (Fig. 2). Full details will be published elsewhere. We would, therefore, like to take this opportunity to emphasise the importance of measuring the uptake of any
A Description of a Low Field Resistive Magnetic Resonance Imaging System and its Application in Imaging Midline Central Nervous System Pathology

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A commercial low field magnetic resonance imaging (MRI) system installed in the Royal Infirmary, Edinburgh at the end of 1983 has since been used for the clinical investigation of over 1000 patients. This system uses an interlaced saturation recovery and inversion recovery pulse sequence which yields four types of clinical image. A T₁ weighted image has been found to be the most sensitive for the detection of pathology. Several examples of pathology with negative X-ray computed tomography examinations including three examples of a syrinx and two cases of tonsillar herniation have been demonstrated using this system. It is suggested that the system could provide a routine clinical service for imaging the central nervous system.

The past four years have seen considerable progress in magnetic resonance imaging (MRI). In the majority of this work imaging systems with field strengths of 0.15 T and above have been used, though some clinical work has been performed at 0.08 T (Hawkes et al., 1980; Bydder and Steiner 1982; Bydder et al., 1982; Brant-Zawadski et al., 1983; Bydder et al., 1983; Gamsu et al., 1983; Herfkens et al., 1983; Hricak et al., 1983a; Hricak et al., 1983b; Smith 1983; Worthington et al., 1983; Kean et al., 1985). Since the major application of MRI appears to be in the investigation of the central nervous system it is necessary to produce satisfactory images of this system with sufficient resolution and pathological contrast within an acceptable patient investigation time.

THE MAGNETIC RESONANCE UNIT

The magnetic resonance machine is a low field 0.08 T system manufactured by M&D Technology Ltd, installed in the Royal Infirmary, Edinburgh (Smith et al., 1984). It employs a resistive magnet whose magnetic field is directed along the vertical axis. The design of the system is closely based on the Aberdeen magnetic resonance imager (Hutchison and Smith, 1983) and uses an interlaced saturation-recovery and inversion-recovery pulse sequence (Fig. 1). The repetition rate (TR) and time to inversion (TI) are normally 1000 ms and 200 ms respectively resulting in a total data acquisition time of 4 min 16 s for each section.

Four types of clinical image are processed for each section (Fig. 2). The so-called 'P' or 'proton density' image (Fig. 3) is obtained from the saturation-recovery pulse sequence which utilises a field echo with an echodeay (TE) of 10 ms. This image has the best signal to noise ratio and consequently the best spatial resolution of the four images, but since there is very little difference between proton density in different normal tissue types and between different pathological tissues little pathological information is contained in this image.

The image is not a true proton density image as substances with a very long T₁ relaxation time like cerebrospinal fluid will give very little signal due to the incomplete relaxation of magnetisation between successive pulses with a TR of 1 s.

The inversion-recovery pulse sequence employs an adiabatic fast passage (AFP) radiofrequency (RF) pulse to ensure complete 180° inversion of magnetisation throughout the section of data collection. This differs from more conventional 180° RF pulses where it may be

![Fig. 1](https://example.com/image1.png)

**Fig. 1** - The NMR pulse sequence used in the M&D 800 imager.

![Fig. 2](https://example.com/image2.png)

**Fig. 2** - The four different types of image produced by the imager.
uncertain whether complete inversion through 180° has been achieved. The inversion-recovery image can be displayed by either a non-modulus or a modulus form of display (Fig. 4). The inversion recovery image is \( T_1 \) weighted, allowing for discrimination between tissues of different \( T_1 \) values such as grey and white matter. The contrast, that is the difference in signal between two different tissues, will vary with the time allowed for longitudinal relaxation before a 90° pulse is applied (TI).

The ‘D’ or ‘difference’ image is obtained by subtracting the numerical data of the inversion-recovery pulse sequence from the saturation recovery pulse sequence. This image (Fig. 5) contains much of the spatial resolution of the saturation-recovery image but its \( T_1 \) weighting provides much more pathological information. This \( T_1 \) weighting will result in regions of prolonged \( T_1 \) giving a higher signal intensity.

A calculated \( T_1 \) image is obtained from the following equation:

\[
T_1 = \frac{200}{\ln \left( \frac{2 \times P}{D} \right)}
\]

where ‘P’ and ‘D’ represent the pixel values in the proton density and difference images respectively. This image is often useful in confirming the presence of pathology which is suspected on the ‘difference’ or inversion-recovery image. The \( T_1 \) calculation is precise due to the complete inversion achieved by the AFP pulse. The long-term in vitro precision over six months varies from 2.3% (coefficient of variation) to 4.1% for \( T_1 \) values of between 163 ms and 416 ms respectively. The in vivo precision is 4.9% for grey matter and 2.8% for white matter (Smith et al., 1985). An example of a \( T_1 \),
image is shown in Fig. 6. The $T_1$ values can be displayed using a linear grey scale or colour-coded display. The pixel values on the grey scale and the colours on the colour-coded display are directly related to the $T_1$ value in milliseconds.

THE MACHINE IN CLINICAL PRACTICE

The patient is positioned by means of a light source and is transported into the magnet system on a motor-driven couch. A smaller head coil is fitted into the larger body coil for cranial examination. A 16 s 'fast' scan is used to check the anatomical level before the 4 min 16 s full data collection is begun. There is some $T_1$ information in the 'fast' scan and this can show pathology such as the subdural collection shown in Fig. 7. This is useful if the patient is unable to cooperate or elderly when chance of movement during the longer scan increases. Fast scans, due to the short $TR$ are also more flow dependent and may be used to confirm the presence of flowing blood within a lesion like an intracranial aneurysm (Fig. 8). Subsequent transverse images above
or below the first image are then obtained by controlled couch movement. Sagittal and coronal images can also be obtained but at present these are only available through the mid-sagittal and mid-coronal planes of the imaging volume. Slight adjustment to the position of this plane in the midline of the imaging volume may be made by alterations to the static magnetic field. These alterations however are limited and only permit about 1 cm latitude. It is also possible, by moving the patient slightly, to alter the position of the region of interest being imaged.

As with other MRI systems claustrophobia may be a problem, particularly with cranial examinations where the head is more enclosed within the smaller head coil. Gentle persuasion overcomes this problem in all but the most severely claustrophobic. The overall total rejection rate due to claustrophobia is 2%.

To date 49 children have been examined, eight of these under general anaesthesia. Several problems are posed in the administration of general anaesthesia in a magnetic environment but these have been successfully overcome. The problems are minimised by the low mag-

![Image](a)  
![Image](b)

Fig. 11 - (a) A midline sagittal proton density image showing central low signal in the cervical cord due to a syrinx cavity. (b) 'Normal' CT scan of the same patient (L+100, W250).

![Image](a)  
![Image](b)

Fig. 12 - (a) A midline sagittal inversion-recovery image showing high signal from a large lipoma in the cervical spinal cord. (b) Midline sagittal calculated \( T_1 \) image showing the very short \( T_1 \) of this lipoma compatible with fat. The displacement of the spinal cord is also shown. (From Kean et al., 1985.)
nentic field. It is important that no metal is used in the anaesthetic tubing, particularly at the mouthpiece. Metal stethoscopes cannot be used but an oesophageal stethoscope has proved ideal. A standard anaesthetic trolley causes distortion of the main magnetic field when it is in close proximity to the magnet and adjacent to the RF shield next to the patient couch. Once an anaesthetic trolley is in position however this distortion can be compensated for by adjusting the static magnetic field; and provided it is not moved, the quality of the image is unaffected (Fig. 9). A pulse monitor with a light-emitting diode detector around a digit may be used to monitor the pulse. It is important that all anaesthetic staff are warned of the possibility of danger from projectile metal objects in the proximity of the static field. This hazard however is not so great a problem with a lower field system than with higher field systems. In common with all other MR systems, patients with pacemakers are not allowed within the imaging suite and at present patients with aneurysm clips are not imaged.

CLINICAL RESULTS

Craniovertebral Junction

To date 44 patients with suspected pathology in this region have been examined. The assessment of suspected pathology in this location is made particularly easy by the use of the sagittal imaging plane. This allows the position of the cerebellar tonsils to be accurately defined and any downward displacement of these structures to be easily assessed. Since cortical bone is not visualised by MR the anterior and posterior margins of the foramen magnum are not directly visualised but can be inferred. The clivus, which contains fat, points to the anterior margin of the foramen magnum and the fat in the occipital diploe points to the posterior margin. When the cerebellar tonsils are displaced inferiorly (Fig. 10) they are seen to herniate through the foramen magnum and in this example are seen to approach the posterior arch of the second cervical vertebra.

The presence of a syrinx can be rapidly demonstrated by low-field MRI. Figure 11(a) demonstrates a low signal from a syrinx cavity within the cervical cord. This cavity contains CSF which on a proton density image gives a low signal due to its very long relaxation time when examined with a TR of 1 s. A moderate degree of tonsillar herniation is also seen. The computed tomography (CT) examination, with intrathecal iopamidol (Fig. 11b), shows a normal sized cord and no evidence of a syrinx.

A lipoma in the cervical cord is dramatically shown on inversion recovery and calculated $T_1$ images (Fig. 12). The short $T_1$ of this lesion allows a diagnosis of lipoma to be made. The anterior displacement of the cervical spinal cord is also well seen.

Brain Stem

To date 50 patients with suspected pathology in this region have been examined. Lesions in the brain stem cause a non-specific rise in relaxation times. Figure 13a
demonstrates an expanding lesion in the pons, which is effacing the 4th ventricle. The coronal image (Fig. 13b) confirms the intra-axial location of this pathology. Vascular lesions such as brain-stem infarcts are easily demonstrated. Figure 14 demonstrates a region of low signal on an inversion-recovery image corresponding to prolonged $T_1$ in a large brain-stem infarct. Figure 19 demonstrates a region of prolonged $T_1$ in the pons in a patient with clinical signs of brain-stem demyelination. It must be pointed out that this $T_1$ abnormality is not specific and may also be seen in a small area of ischaemia.

**Corpus Callosum**

This is another structure which is easily assessed by MRI. Figure 16a demonstrates agenesis of the corpus callosum with an associated lipoma. The transverse section (Fig. 16b) shows the separation of the lateral ventricles and the presence of a rounded short $T_1$ lesion anteriorly.

**Pituitary Gland**

To date 60 patients with suspected pathology in this region have been examined. MRI does not demonstrate the destruction of cortical bone by pituitary pathology but the presence of most pituitary pathology is easily demonstrated. A very large pituitary macroadenoma is demonstrated in Fig. 17a; although demonstrated well by X-ray CT (Fig. 17b), it should be pointed out that this examination required over 45 1.5 mm sections together with intravenous contrast medium to provide such an image. The spatial resolution of the MRI machine is at present insufficient to demonstrate small pituitary lesions like microadenomas.

**DISCUSSION**

The advantages of MRI in the investigation of the CNS are now well known and many of these can be obtained from a low-field system. There have now been numerous publications demonstrating the use of MRI in the central nervous system. All of these have shown that MRI is the modality of choice in this region due to the following features:

1. The variability of the pulse sequences give MRI flexibility and allows the examination to be tailored to suit the particular clinical problem since these sequences may be altered to provide more contrast between normal and abnormal tissues which makes MRI a more sensitive technique.

2. The lack of artefact, particularly streaking in the posterior fossa, permits the demonstration of this area along with the craniovertebral junction much better than X-ray CT.

3. The direct acquisition of sagittal and coronal images permits the rapid and accurate localisation of pathology particularly at the cranio-vertebral junction and in the brain stem.

Fig. 15 - Transverse 'D' image showing high signal in the posterior aspect of the pons due to an area of demyelination.

Fig. 16 - (a) Midline sagittal inversion-recovery image showing high signal from a lipoma of the corpus callosum. (b) Transverse calculated $T_1$ image showing low signal from the same lipoma. The separation of the lateral ventricles is also seen.
These advantages have been demonstrated on systems with a higher field strength than the one described. We believe that the low-field system described is capable of producing excellent clinical results. The most obvious disadvantage is the inferior spatial resolution compared with high-field systems. This drawback may only be a problem in examining regions such as the pituitary and the internal auditory meatus. Signal averaging may overcome this but this would mean a slightly increased imaging time. Another drawback to the system at present is the lack of offset slices in the sagittal and coronal planes. It is planned to upgrade the system with this facility together with the addition of a facility for multiple simultaneous slice acquisition.

In conclusion, we feel that this imaging system, at least when applied to the central nervous system, can provide almost all the advantages inherent to MRI but without many of the problems associated with higher field, higher cost systems; and with the additions of offset slices and multiple simultaneous slice acquisition this system will be capable of providing a routine clinical service in imaging the CNS.

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REFERENCES


Technical note
A technique for velocity imaging using magnetic resonance imaging
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Magnetic resonance imaging (MRI) is becoming well established as a technique which both shows anatomy and allows tissue characterisation. A further development of the technique allows the imaging of regular movement such as blood flow and cardiac wall motion. In particular, much effort has been put into developing techniques for imaging flowing blood and deducing such parameters as velocity and acceleration.

Early work by Singer (1959) described a technique of blood flow velocity measurement using a "time-of-flight" method. Velocity images have been obtained by this method using a modified version of a spin-echo pulse sequence (Feinberg et al, 1984). Normally, a slice is excited using a selective 90° radio-frequency (RF) pulse and an echo is produced at a later time by applying a selective 180° pulse to the same slice. If moving hydrogen nuclei have, in the meantime, moved out of the slice, no signal will be returned from them, resulting in a reduction in signal intensity from blood vessels in the image. This is the spin washout effect. If, however, the 180° RF pulse is applied to a slice downstream, an echo signal will be returned from excited nuclei having a velocity such that they coincide with the second slice selection. In this way, a series of images can be obtained, each corresponding to a certain range of velocities.

The standard spin-echo technique can also be exploited to produce flow-dependent signals in vessels (Axel, 1984; Bradley et al, 1984; Waluch & Bradley, 1984). The effect, known as paradoxical enhancement, results in vessels appearing to have a much higher intensity than the surrounding tissue. This effect becomes particularly noticeable for shorter repetition times since freshly saturated nuclei moving into the slice will give a stronger signal relative to the surrounding stationary tissue, which is only partially saturated.

In a multi-echo sequence a phenomenon known as even-echo rephasing is also present (Waluch & Bradley, 1984). The magnetic field gradient (subsequently referred to as the gradient) used for slice selection and the gradient used for readout are applied twice in a spin-echo sequence. The first time they are applied, nuclei dephase along the direction of the gradient; the second time, they are applied with opposite polarity and nuclei move back into phase along that direction. If the nuclei are stationary then they will move back exactly into phase with one another; however, if nuclei are moving along a gradient direction they will have a phase shift at the first echo, having only partially rephased. This phase shift will be linearly dependent on their velocity. All nuclei having the same velocity will have the same phase shift on odd echoes but will have zero phase shift on even echoes, the effect of the gradients having been reversed provided the velocity is constant. Thus, flow can be identified by comparing first and second echo images. On the first echo, flowing blood will appear to have a reduced signal while, on the second echo, the signal will be of higher intensity. We are, of course, neglecting dephasing effects caused by spin-spin relaxation.

Since the non-zero phase shift on the first echo is linearly dependent on velocity, it can be used to measure velocity directly (Bryant et al, 1984; Moran & Moran, 1984; Redpath et al, 1984; Van Dijk, 1984; Moran et al, 1985; Wedeen et al, 1985). In most MRI systems the signal is collected from two channels in quadrature, yielding a real and an imaginary data set. These are usually combined to calculate a modulus image used for diagnosis. These two data sets can, however, be used separately to produce a real or an imaginary image, or can be combined to calculate a phase image. All three of these contain phase information which is normally discarded in the calculation of the modulus image and it is in this information which provides a way of imaging velocity.

Although real and imaginary images have been used to demonstrate imaging of velocity, they also contain spin-density information (Moran et al, 1985). The phase image is easiest to interpret and quantify since it contains only motion information and represents a direct linear map of a single velocity component. In order to quantify the velocity measurements accurately, standard spin-echo and field-echo sequences have been modified by adding extra gradient pulses in the direction of required velocity measurement.

To date, this has been done in two ways (Moran et al, 1985); either two identical gradient pulses separated by a 180° RF pulse are inserted, or a bipolar pulse consisting of two successive gradient pulses equal in magnitude and duration but of opposite sign is inserted into the pulse sequence between the 90° RF pulse and the readout gradient. The first of these
Technical note

gradient pulses dephases the nuclei in the applied direction and the second pulse refocuses them. Stationary nuclei are all brought back into phase, whereas for nuclei moving along the gradient the refocusing is not complete, leaving them with a phase shift proportional to velocity.

The accuracy of the technique is dependent on the two gradient pulses having exactly the same duration and magnitude over the imaging volume and on the uniformity of the RF, gradient and static fields. A non-uniform field will introduce a spatially dependent phase shift over the image (McVeigh et al. 1985; Norris, 1985b). A technique has been developed (Redpath et al., 1984) to overcome these difficulties by varying the size of the flow-encoding pulses for a number of data acquisitions and applying a 3D Fourier transform technique to the whole set of data. The resultant set of images will be coded in intensity, each corresponding to a certain velocity range, and may be used to reconstruct a simple velocity-coded image. A further limitation of the techniques is that velocities corresponding to a phase shift of greater than +180° or less than −180° will lead to an ambiguity of velocity measurement. This can be avoided by adjusting the magnitude of the gradient so that the resultant phase shifts will lie within the range of ±180°. This paper presents a technique which uses the calculated phase image as a direct measurement of velocity. The problem of field non-uniformity is avoided by using two separate pulse sequences and subtracting the resultant phase images.

METHOD

The technique has been integrated into the standard spin-warp imaging sequence (Edelstein et al., 1980) used on an M&D 0.08 T resistive MRI system (Smith et al., 1984). The slice selection gradients, which are effectively of bipolar form, were initially used to provide the phase encoding for velocity components perpendicular to the imaging slice (Fig. 1). In order to avoid the appearance of phase shifts due to the spatial non-uniformities of the fields, two pulse sequences were used to produce two phase images which were subtracted to produce the final velocity image. The two pulse sequences were identical except that the timing of the rephasing gradient was later in one than in the other. Phase errors due to the time-independent inhomogeneities in the main field and the RF field would be the same in each case and would, thus, cancel in the final subtracted image. The only remaining phase differences are due to motion, such that for a given velocity component, v, in the direction of the applied slice selection gradient, G, the phase difference is:

\[ \phi_{\text{diff}} = -\gamma G t_v \Delta \phi \]

where \( t_v \) is the duration of the rephasing pulse, \( \Delta \phi \) is the difference in pulse separation for the two pulse sequences, and \( \gamma \) is the gyromagnetic ratio for protons (Fig. 1).

All the computer processing is performed within the acquisition program so that the subtracted phase images are displayed 8 s after acquisition. An ungated velocity image can be obtained in 64 s using a repetition time (TR) of 250 ms. For in-vivo imaging, however, the pulse sequences are gated (Been et al., 1985) in order to look at an instantaneous velocity, and consequently the imaging time is dependent on the heart rate. Typically, one section will take about 4 min. The two pulse sequences are interlaced so as to reduce the effects of gross patient movement between scans. The magnitude of the flow pulse gradient, G, is 1.25 mT m\(^{-1}\). For the studies described here, the flow pulse timing was such that \( t_{\text{diff}} \) = 2 ms and \( t_v \) = 2.4 ms. The velocity sensitivity can be increased simply by increasing \( t_{\text{diff}} \).

![Fig. 1.](image-url) A simple adaptation of the Aberdeen spin-warp pulse sequence for imaging of velocity through the slice by varying the position of the rephasing gradient (broken line) between positions 1 and 2. Two corresponding phase images are produced and then subtracted to produce the velocity-coded image.
Velocity measurements have been calibrated using a simple flow phantom consisting of a 1 cm-inside diameter polythene tube surrounded by a bath of 2.5 mm solution of copper sulphate. The same solution of copper sulphate was pumped through the tube at various known constant rates. The mean flow velocity for the tube was calculated and calibrated against the mean phase shift found within the tube. A number of normal volunteers have been imaged in the neck, thorax and abdomen using the interlaced gated sequence. An initial comparison with continuous wave Doppler ultrasound at 2.5 MHz has been carried out in the abdominal aorta using a Hewlett Packard 77020 Doppler imaging system. Using MRI, a number of delays were used from the electrocardiogram R-wave in order to plot the velocity in the abdominal aorta over the systolic period of the heart cycle. The sample volume for this measurement was taken at the centre of the aortic diameter.

**RESULTS**

The calibration of the velocity measurement shows a good linear relationship between velocity and phase shift (Fig. 2). Three examples of the volunteer studies are shown in Fig. 3. These images display zero phase shift as mid-grey, positive phase shift of 180° as white and negative phase shift of 180° as black. Where the values in the modulus image are below a certain threshold the phase values are not calculated.

Figure 4 shows the velocity measurements taken in the abdominal aorta compared with those taken from continuous-wave Doppler ultrasound measurements. There is a good agreement in profile shape and timing, although the velocities appear to be lower in the case of the Doppler measurements.

**DISCUSSION**

It has been shown that velocity measurements are possible on the low field imaging system at Edinburgh with changes only to the pulse sequence and software and without any improvements required on the field homogeneity.

This technique, using phase as a direct measurement of velocity, has advantages over the other techniques.
mentioned in the introduction due to its relative simplicity in interpretation, processing and display. The time-of-flight method is restricted to imaging of one particular velocity range per image. Paradoxical enhancement and even echo rephasing are not quantitative measures of flow and can only be used to identify whether flow is present. The 3D Fourier transform method requires many flow encoding stages, whereas the technique described here requires only two sections to produce the final velocity image.

Initially, high flow velocities are being examined since they occur in the larger, better defined vessels. Lower velocities of the order of 1 cm s\(^{-1}\) have been imaged by the technique simply by extending the time \(T_{im}\). However, if this becomes too large then inhomogeneities in the field lead to loss of signal from the imaging slice, so it would be necessary instead to increase the magnitude of the flow gradient.

While peak velocity in the abdominal aorta can easily be imaged, the peak velocity in the ascending aorta is rather more difficult to image due to loss of signal around high velocity gradients over the vessel cross-section (Moran & Moran, 1984; Moran et al., 1985; Wedeen et al., 1985). These large velocity gradients result in a large range of velocities and, hence, a large range of phase shifts being present in the same pixel volume. This phase dispersion causes the signal from the nuclei in the same pixel volume to cancel, resulting in loss of signal, and is often seen as a dark line along the inner surface of the vessel. This effect can be seen even in standard pulse sequences, as the slice selection and signal readout gradients induce accidental phase shifts for nuclei moving in their direction. Of particular importance is the case of mitral-valve stenosis where high-velocity jets are present.

Methods have been suggested to overcome the phase dispersion which causes this loss of signal (Le Roux & Floch, 1985; Pattany & Naylor, 1985). It is proposed to use an adaption of the compensating bipolar pulse technique suggested by Le Roux and Floch, modifying it for use with the basic spin-warp pulse sequence. The compensating bipolar pulse is of opposite polarity to that already in the sequence, so that the dephasing effect is reversed. Figure 5 shows the suggested pulse sequence for use in further work, incorporating compensating bipolar pulses into the technique in all three directions. At present, this is not possible to do due to hardware limitations, although it is hoped to implement it in the near future.

It has already been shown (Norris, 1985a) that double bipolar pulses can be used to image acceleration. The greater the separation between the bipolar pulses, the greater the phase shift for a given acceleration. Thus, phase shifts due to acceleration can be minimised by merging the two middle pulses of the double bipolar flow pulse into one. The velocity measurement is achieved by varying the position in time.
of the broken line pulses in Fig. 5 between positions 1 and 2. Using the previous notation, the expression for phase shift is given by:
\[ \phi_{\text{det}} = \gamma G v_{\text{det}} t_{\text{det}} + \gamma G a (7t_{\text{det}}^2 + t_{\text{det}}^2) \]
where \( a \) is the constant acceleration.

Thus, for conditions where maximum acceleration occurs in vivo (200 cm s\(^{-2}\)) this would produce at most a 10% error in velocity measurement, given that typical velocities at this point are around 50 cm s\(^{-1}\).

It is proposed to extend the technique to the measurement of flow in all directions in all phases and to pursue further volunteer studies and initial clinical trials. The technique will be compared directly with results from Doppler ultrasound measurements where possible.

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Correlation of human NMR $T_1$ values measured in vivo and brain water content

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ABSTRACT

A relationship has been demonstrated between nuclear magnetic resonance (NMR) longitudinal relaxation times ($T_1$ values) obtained in vivo in both normal and oedematous (peritumoral) brain tissue, and measurements of brain water obtained by gravimetric analysis of operative samples. Significant correlations were found in seven patients in both cortex ($r = 0.97$; $P < 0.001$) and white matter ($r = 0.92$, $P < 0.001$). These findings suggest that NMR may prove a useful technique for monitoring brain oedema.

Nuclear magnetic resonance (NMR) $T_1$ values in body tissue are considered to be related to total water content and to the ratio between “free” and “bound” water (Go & Edzes, 1975; Mathur-De Vre, 1984). Longitudinal relaxation time ($T_1$) lengthens in oedematous tissue and in a variety of pathological conditions including malignancy. Previous authors (Go & Edzes, 1975; Naruse et al., 1982) have shown a linear correlation between $T_1$ of experimentally induced cerebral oedema, measured by in vitro NMR spectrometry, and brain water, calculated from the weight of wet and dry tissues. The present study was performed in order to examine the relationship between $T_1$ values, obtained from clinical NMR images of patients about to undergo surgery for cerebral tumour, and brain water measurements obtained by gravimetric analysis of the excised specimens, and also to establish whether $T_1$ images can be used to predict the degree of cerebral oedema.

METHOD

Patients

All seven patients examined had histologically verified solitary tumours within the brain. Six were gliomas and the seventh a metastasis from an adenocarcinoma of lung. All patients were receiving 4 mg dexamethasone four times a day at the time of imaging and surgery. Four of the patients received 200–500 ml 20% mannitol at the time of dural opening, within 30 min of tissue excision. Six patients underwent lobectomy and one patient had multiple needle biopsies of tumour and peritumoral tissue. The excised specimens were rapidly enclosed in “cling film” and transported to the adjacent laboratory for measurement of brain water content.

Brain water measurements

Brain water content was measured by microgravimetry using small tissue samples from the lobectomy specimens. The specific gravity (s.g.) of each sample was determined by placing it in a layered gradient column of a mixture of kerosene and bromobenzene oil, that had a high specific gravity at the bottom and a low specific gravity at the top (Marmarou et al., 1978). Two minutes after insertion of the sample into the column, its depth in the oil was measured, and the column was subsequently calibrated by measuring the depth of potassium-sulphate droplets of known specific gravity. The water content of each sample was then calculated from its specific gravity using the formula:

\[
g = \frac{1}{(1 - \frac{\text{s.g.}}{\text{solids s.g.}}) \times \text{sample s.g.}} - 1
\]

derived by Nelson et al. (1971). The s.g. of the normal human cortex and white matter solids was determined by wet-weighing 500 mg samples from five patients with normal brain in their operative specimens, and then
reweighing the same samples after drying to constant weight at 80°C (Nelson et al., 1971).

The lobectomy specimens were sectioned in the transverse plane. Three samples of peritumoral white matter, each weighing about 10 mg, were taken from each site. Each sample was measured in the layered kerosene/bromobenzene density column (Mamarou et al., 1978), and samples of normal cortex and white matter, when present in the specimen, were similarly measured. In two patients, macroscopically oedematous cortex was also sampled. All gravimetric measurements were completed within 15 min of excision of the specimen, to minimise water loss from the tissues. The lobectomy specimen was then fixed for subsequent histological examination.

Analysis

The NMR images were reviewed by the same radiologist and surgeon without reference to the gravimetric data. The sites of tissue sampling had been noted by the surgeon with reference to known landmarks. T1 values for abnormal tissue were obtained from those sites using the region of interest facility on the diagnostic console. Values for normal tissue were taken by the same method, wherever possible, within the lobe which had been resected.

RESULTS

A graph of T1 values plotted against brain water readings for normal and oedematous tissue is shown in Fig. 1. A significant correlation exists between the two measurements for white matter (r = 0.92, P < 0.001), and cortex (r = 0.97, P < 0.001). Statistical analysis of the data (Smith et al., 1981) demonstrates that the errors of the techniques account for the spread of the bivariate data about the regression line (\( r^2 = 36.9, P < 0.001 \)).

DISCUSSION

Brain tumours are often surrounded by a zone of oedema. This form of oedema is regarded by Klatzo (1972) as vasogenic, where oedema fluid emerges through a defective blood-brain barrier and spreads in the extracellular space of the white matter. This preliminary in-vivo study confirms in patients with brain tumours the linear correlation between brain water content and NMR T1 values, previously reported in experimental animal models. Our results, however, highlight differences between white matter, where T1 increases rapidly as water content rises, and cortex, where the increase is slower. In normal tissue the water content of cortex is greater than that of white matter, our finding of 10% agreeing closely with the 10–15% reported by other authors (Mathur-De Vrë, 1984). The T1 of normal cortex is correspondingly longer than that of normal white matter. Oedema fluid accumulating in cortical tissue is taken up by the cells and is, therefore, influenced by the intracellular macromolecules. The water protons are “bound” in the hydration layer around the macromolecules and this alteration in their dynamic state shortens their relaxation time (Mathur-De Vrë, 1984). In oedematous white matter more water remains in the extracellular compartment. A lower rate of collisions between molecules in this “free” state results in a longer relaxation time. Previous authors, working with rat brain in vitro, have attributed the linear increase in T1 with water content to increase in the “free” water (Go & Edzes, 1975). Although results from our small sample must be interpreted with caution, we have shown a less marked rise in cortex, in keeping with the smaller rise in “free” as opposed to “bound” water. It appears that NMR T1 values can be used to predict the degree of cerebral oedema in white matter, and possibly in cortex.

Four of our patients received intra-operative mannitol. Measured water content in their peritumoral white matter is lower than might be expected from preoperative T1 readings. Magnetic resonance imaging may prove a useful tool in assessing the effect of mannitol and other agents on cerebral oedema.

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Correlation of human NMR $T_1$ values and brain water content


Original Contribution

A SURFACE COIL DESIGN FOR A VERTICAL FIELD MRI SYSTEM AND ITS APPLICATION IN IMAGING THE BREAST

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The general design of a flat surface coil for use with a vertical field MRI system is discussed. A development of the design for imaging the breast is used as a specific illustration. The phenomenon of "sympathetic resonance" is described in which transmission and reception is performed on the standard body coil with the surface coil disconnected, its presence amplifying the signal to the body coil from the region of interest. Results show that better quality images can be obtained with the surface coil than with the standard body coil and that the sympathetic resonance mode overcomes to a large extent the limited field of view problem normally associated with a surface coil. A threefold improvement in signal to noise ratio was obtained using the surface coil.

Keywords: Magnetic resonance imaging; Surface coils; Vertical main field; Sympathetic resonance; Breast T₁ imaging.

INTRODUCTION

Tomographic images through human subjects are obtained by magnetic resonance imaging (MRI) using a combination of magnetism and radio-frequency (RF) radiation. The RF coil or coils are either in the form of a combined transmitter and receiver coil, sometimes known as a transceiver, or separate transmitter and receiver coils. It is possible to obtain a significant improvement in the signal to noise ratio (S/N) of the NMR signal by placing a smaller RF coil, generally referred to as a surface coil, close to the patient at the region of interest. Such a coil is generally only a receiver coil, a much larger coil being used as a transmitter. The improvement in S/N produces images of a higher quality but over a restricted region of the patient.

The simplest type of surface coil is a circular loop of conducting material. As the RF field (B₁) must be perpendicular to the main magnetic field (B₀), there is a restriction in the orientation of these surface coils. Most MRI systems have a horizontal magnetic field B₀, which means that a circular surface coil can be used in planes parallel to the patient couch. This is very convenient for imaging regions such as the orbit or breast.¹

When the main magnetic field B₀ is in the vertical direction, circular surface coils cannot be used in the plane parallel to the patient couch. To image a specific region of the body, different coil configurations must be developed which produce the RF field B₁ in a horizontal plane.

An additional criterion to be considered in the design of a surface coil when a vertical field MRI system is used is that the access for the patient in the vertical direction is generally more restricted than in horizontal field systems. The form of coil for imaging the breast or spine, for instance, must take up the minimum amount of space in the vertical direction. In particular, a flat spine coil would allow imaging with the patient lying supine over it. A further criterion which applies to all breast coils, irrespective of the main field orientation, is that the axillary tail must be imaged as well as the breast.

Although the general design discussed here is applicable to imaging various parts of the body using a vertical field system, and has been tested by us for

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imaging both breast and spine, there is particular clinical interest in Edinburgh in the former, and so the technique is specifically illustrated here with the design of a breast coil.

METHODS

A breast coil was developed for the M&D 0.08-T vertical field resistive MRI system sited in the Royal Infirmary, Edinburgh. The patient enters between the faces of the two larger horizontal ring magnets; the distance between the patient couch and the lower face of the upper magnet is 33 cm. One advantage of the vertical field orientation for the main magnetic field is that solenoidal RF coils can be used, resulting in a higher S/N than would be obtained from a similar size saddle coil. The RF coil used for conventional body imaging is a split solenoid, which is situated on the outside of the patient tube containing the gradient windings. The coil is electronically switched to transmit and receive mode. Prior to the development of surface coils, body images were obtained using a 16-mm section width. Calculated $T_1$ maps were obtained using an interleaved saturation recovery and inversion recovery pulse sequenced with $TR = 1000$ ms for both and $TI = 200$ ms for the latter.

Coil design

At the operating frequency of 3.4 MHz the skin depth in copper is less than 0.1 mm. As a consequence it is possible to construct RF coils using copper tube or ribbon. To reduce the resistance of the coil as much as possible, surface coils were constructed using copper pipe with an outside diameter of 8 mm and a wall thickness of 0.6 mm. To construct a surface coil, the appropriate length of copper pipe was straightened and then filled with fine dry sand. The pipe could then be shaped by hand, the sand enabling fairly sharp bends to be obtained without distorting the pipe. Having shaped the coil, the sand was removed. To tune the coil, fixed polystyrene capacitors and nonmagnetic variable capacitors were used. Polystyrene or polyethylene capacitors suffer less RF loss than other types of capacitors. 300-Ω twin feed cable connected the breast coil to the standard M&D preamplifier. When rigid support was required, polycarbonate was used rather than perspex because there is less RF loss with the former.

The surface coils developed for our MRI system used the principle illustrated in Fig. 1. The parallel sections of conductor in the central region of the coil carry current in the same direction resulting in a magnetic field with a significant component perpendicular to the parallel conductors. The return paths of the coil are positioned far enough away from the central region so that the RF field from them does not interfere with the central region. This type of coil can be flat, and such surface coils have been constructed for the spine and mediastinum.

The breast coil design was an extension of this principle and is illustrated in Fig. 2. The breast coil was connected as two loops in parallel and mounted on an expanded polystyrene block in which a trough

Fig. 1. The transmitter coil produces an RF field $B_T$ parallel to and along the long axis of the patient couch. The central conductors of the flat surface coil produce an RF field $B_f$ which is perpendicular to both $B_0$ and $B_T$. The field $B_f$ is produced in the region of the central conductors of the flat surface coil; the rest of the flat surface coil is oval in shape, the conductor being sufficiently far from the central region not to interfere with the RF field.
was cut. A piece of high-density foam, 1 cm thick, was then placed on top of the coil and the polystyrene block to reduce capacitive coupling between the breast coil and the patient. The subject lies prone with the centre of the breast of interest positioned in the middle of the parallel conductors.

The surface coil was not used to transmit RF radiation; the conventional solenoidal body RF coil was used as a transmitter resulting in uniform spin excitation throughout the tomographic section. The orthogonal orientation of the RF fields associated with the transmitter coil and breast coil, which acted as the receiver, reduced the degree of cross-coupling. The presence of the breast coil did not significantly change the resonant frequency of the body coil, nor was there any interference on the breast coil from transmitted RF radiation. The breast coil was tuned inside the imager at the central position and connected to the preamplifier which had a capacitive loading effect of 60 pF. The pixel size was reduced from 3.5 mm square for body imaging to 2 mm square and slice thicknesses of 8–12 mm were used.

Coil performance assessment

To assess the performance of the breast coil, the Q factor and the S/N were measured. Two methods were used to determine the Q of the breast coil. The first method to measure Q was performed during the tuning of the breast coil. Two 15-cm-diameter circular copper search coils were placed 20 cm apart, one either side of the breast coil. One coil was connected to a signal generator and frequency counter, the former being set to the exact reference frequency of the MRI system. The second coil was connected to an oscilloscope and the capacitors of the surface coil adjusted until maximum signal was obtained from the oscilloscope. The Q of the coil was obtained from the equation

$$Q = \sqrt{3} \frac{f_{\text{max}}}{f_1 - f_2} ,$$

where $f_{\text{max}}$ equals frequency of maximum signal, $f_1$ equals frequency above $f_{\text{max}}$ at which the signal amplitude is reduced by a half and $f_2$ equals frequency below $f_{\text{max}}$ at which signal is reduced by half. The second method of measuring Q employed a vector impedance meter connected across the surface coil terminals to trace its impedance variation with frequency about 3.4 MHz. This technique was useful for making measurements of Q for both unloaded and patient loaded conditions, search coils being difficult to apply in the latter. All Q values were measured with the breast coil in the imaging position inside the magnet.

A major aspect of our research in Edinburgh into the use of MRI in the breast is the use of $T_1$ in the investigation of breast disease. As a consequence the image S/N was obtained from the calculated $T_1$ map for specific section widths and at various distances from the breast coil. The definition of S/N used for evaluation and comparison of surface coils was
$S/N = \text{mean } T_1/\text{SD}$,

where the mean $T_1$ was obtained from a region of interest containing at least 80 pixels, and SD equals the standard deviation of individual pixel $T_1$ values within that region.

Coil performance was assessed from images on normal volunteers and also from images of a 7.5-cm-diameter bottle containing CuSO$_4$ with a $T_1$ of about 200 ms. The bottle was placed in the trough of the breast coil and imaged in the same way as a human subject.

**Sympathetic resonance**

The phenomenon which we have called "sympathetic resonance" was identified while testing the breast coil. In the conventional surface coil mode the breast coil was tuned to the imager resonant frequency (3.4 MHz) and acted as the RF receiver. However, similar improvements to those obtained using the conventional surface coil mode could be obtained if the large body coil used for transmission was also used for RF detection, with the breast coil still tuned to the resonant frequency but not connected to the preamplifier nor directly coupled to the imager electronics in any way. Images were obtained using the breast coil in this sympathetic resonance mode and also, for comparison, with its connectors shorted so that the coil did not resonate near 3.4 MHz. The principle of a sympathetic resonance coil is illustrated in Fig. 3.

**RESULTS**

The $Q$ of the patient breast coil for different situations is listed in Table 1. As expected, there is a reduction in $Q$ when the coil is inside the imager and a further reduction when the coil is loaded with a human subject. There is an increase in $Q$ when operating in the sympathetic resonance mode, because the coil is no longer damped by connection to the preamplifier input.

As expected, there is a variation in sensitivity with distance from the breast coil; this variation, however, is less in the sympathetic resonance mode. The calculated $T_1$ map is obtained from a saturation recovery and inversion recovery image in the usual way, both images having the same signal depth variation. As a consequence, when they are combined, the $T_1$ map is uniform with depth. However, the decreasing signal in the original images results in a gradual decrease in $T_1$ $S/N$ with distance from the breast coil. The variation in $S/N$ with depth is shown in Fig. 4 with the $S/N$ for the conventional body coil and head coil shown for comparison. The section width used with the breast coil is narrower than the 16 mm used for routine patient body imaging. The $S/N$ values at different

<table>
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<tr>
<td>Breast coil:</td>
<td>153</td>
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<tr>
<td>Unloaded</td>
<td>109</td>
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<td>With patient</td>
<td>295</td>
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<td>Sympathetic resonance:</td>
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<td>With patient</td>
<td>316</td>
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<td>Body coil:</td>
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<tr>
<td>Unloaded</td>
<td>475</td>
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<td>With patient</td>
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Table 1. $Q$ values of the breast coil and normal body coil unloaded and loaded with a patient in position

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**Fig. 3.** Schematic diagram demonstrating the coil used as a sympathetic resonator. The larger RF coil is used for both transmit and receive. The surface coil is tuned to the resonant frequency but not directly coupled to the imager electronics in any way.

**Fig. 4.** Variation in $S/N$ in the $T_1$ image with distance from the surface coil at different section thicknesses. For comparison the signal to noise using the head coil and the large body coil at different section thicknesses is shown.
Fig. 5. $T_1$ images of the same normal breast using a slice thickness of 12 mm with (a) breast coil connected in the conventional manner, (b) breast coil connected in the sympathetic resonance mode and (c) conventional body coil.
depths for the breast coil connected in the conventional surface coil mode and also in the sympathetic resonance mode are given in Table 2.

An indication of the improvement obtained using the breast coil on a normal volunteer is illustrated in Fig. 5. For comparison an image is shown using the standard body coil but with the magnet gradients and section width that were used for breast coil imaging. The smaller pixel size and section width used with the conventional body coil produce an apparently noisier image, due to a 75% reduction in signal per pixel, than would be obtained if it were used in its conventional mode. The improvement in image quality compared with the conventional body coil can be seen clearly. This improvement is present when the surface coil is connected in the two different modes though in the sympathetic resonance mode the field of view is larger. This is because the variation in sensitivity with distance from the coil is less in the sympathetic resonance mode and so the image S/N does not deteriorate so quickly.

**DISCUSSION**

It has been demonstrated that a surface breast coil can be used with a vertical field MRI system to obtain significant improvements in breast image quality. It has also been shown that the depth response of a breast coil can be improved when connected in the sympathetic resonance mode. This is particularly important when investigating regions of abnormal tissue near the chest wall or in the axilla.

It is hoped that the use of this breast coil will enable $T_1$ maps to be obtained in patients suffering from breast disease with a much thinner section width than could be obtained using the conventional body coil. This will result in a much smaller partial volume error in the $T_1$ values obtained which may in the past have been partly responsible for the degree of overlap that was found in $T_1$ values between normal and abnormal tissue.

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Cardiac gating in nuclear magnetic resonance imaging

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In order to obtain NMR images of the heart, and to measure blood flow by NMR, the signal acquisition must be synchronized with the patient's cardiac cycle. Some of the problems of detecting the cardiac cycle within a strong uniform magnetic field without introducing RF interference into the imaging system are discussed. Some general applications of cardiac gated NMR imaging are presented.

KEYWORDS: NUCLEAR MAGNETIC RESONANCE IMAGING, CARDIAC GATING, HEART, FLOW.

Introduction

The basic principles of nuclear magnetic resonance (NMR) have been explained in detail both in this Journal [1] and elsewhere [2 and 3]. Essentially the technique involves placing a sample, in our case a human subject, in a strong constant magnetic field. Radiofrequency (RF) radiation is applied at a specific frequency which causes the proton nuclei within the body to resonate; the source of radiofrequency radiation is then switched off and the nuclei continue to resonate emitting RF radiation which can be detected. The size of the NMR signal will be influenced by the amount of protons present, i.e., the proton density. There are two time constants which relate to the exponential decay of two components of the resonance and these are referred to as the relaxation parameters $T_1$ and $T_2$. During the transmission and subsequent detection of the RF radiation additional magnetic fields are briefly applied to encode spatial information within the NMR signal. These additional magnetic fields are known as 'magnetic gradients'.

NMR images are produced using a series of RF pulses and the timing and type of pulses used will result in the amplitude of the detection NMR signal containing information not only of the proton density of the sample but also the $T_1$ and $T_2$. Thus different pulse sequences can be used in magnetic resonance imaging (MRI) which produce images of the body which contain either mainly proton density information, $T_1$ information, $T_2$ information or a combination of two or even three of these parameters. The term ‘weighted image’ is often used to indicate that the signal intensity of the image is mainly influenced by one particular variable. For example in a $T_2$ weighted image there will be a lot of information concerning the proton density but the image intensity will also be influenced by the $T_2$ of the tissue. A $T_1$ image on the other hand contains purely information related to the $T_1$ of the tissue. Such an image is sometimes referred to as a 'T1 map'.

Cardiac gating is important in MRI, not only to eliminate cardiac motion and phase smearing (figure 1) when imaging the heart, but also to image velocity and flow within the body. It is possible to store velocity information in the phase of the NMR signal [4 and 5] resulting in the production of a phase image the intensity of which is related to velocity. Measurements of pulsatile flow in vessels within the body can only be achieved using cardiac gating.

The amount of $T_1$ or $T_2$ weighting in an image using a particular pulse sequence will be influenced by the separation between the pulses in the sequence. As the RF pulses in these sequences are often slice selective, difficulties can arise when attempts are made to synchronize the pulses with the cardiac cycle. Not only can the amount of $T_1$ or $T_2$ weighting vary during a single scan, thus distorting the final image, but the pulses may not all occur at exactly the same part of the cardiac cycle and so the motion of the heart may not be completely ‘frozen’. In addition, the motion of the heart may cause dephasing of the NMR signal resulting in a loss of signal amplitude and low intensity regions in the image.

Cardiac gating is something of a misnomer in MRI. In nuclear cardiology signals are emitted from the patient all the time and the patients' ECG is used to define a ‘window’ or ‘gate’ in time during which signals are recorded. Hence the image obtained is cardiac gated. In MRI the image is built up from a number of different NMR signals, usually 128 or 256. These signals are
produced after a short burst of RF and magnetic gradient pulses which are ‘triggered’ by the patient’s ECG so the NMR signal is obtained in the same part of the cardiac cycle. It is sometimes argued therefore that the term cardiac triggered image is more appropriate.

### Detecting the position of the cardiac cycle

There are a number of problems which must be overcome to synchronize the NMR image with the cardiac cycle:

1. Care must be taken not to introduce an excessive amount of ferromagnetic material into the MRI system. The presence of ferromagnetic material will disturb the magnetic field uniformity and hence can degrade the image.

2. Both the RF and magnetic gradient pulses can produce interference on any system used to detect the cardiac cycle which is situated inside the MRI system.

3. The MRI is very sensitive to RF noise and any leads attached to the patient can act as aerials and introduce unwanted noise into the system.

4. Any system which is developed must not just work for normal volunteers but must be able to tolerate a wide range of abnormal ECGs, compensate for variations in the patient’s R-R interval and detect and reject ectopic beats.

A peripheral pulse can be used to identify the cardiac cycle. This has two advantages: the first is that there need be no electrical contact with the patient, and, secondly, any device attached to the patient need be no nearer than 0.5m to the section imaged. The main disadvantage of this technique is that unlike an ECG it does not provide such an accurate determination of the cardiac cycle [6]. Often in cardiac imaging it is required to trigger each signal at end diastole but the delay which exists between end diastole and a peripheral pulse can prevent such images being obtained.

A patient’s cardiac cycle is most accurately specified using their ECG. It is possible to detect a patient’s ECG using two electrodes, one on either wrist. The advantage of such an arrangement is that the electrodes and cables are away from the region being imaged. The main...
disadvantage is that the ECG signal is low but also interference can be detected from RF transmitter and the gradient coils. In our vertical field MRI system it is possible to detect a patient's ECG using two electrodes, one placed on the right shoulder and one placed on the right hip, which produces a larger signal and less interference [7].

It is possible to use some conventional electrodes, which do not contain sufficient ferromagnetic material to distort the image, at low field strengths but in high field systems the amount of ferromagnetic material may produce significant distortion and carbon electrodes may be necessary. Signal loss can occur around the electrode leads (see figure 1) but as the lead is positioned away from the heart, this is quite acceptable. The small local loss of signal is probably due to eddy currents in the screening of the cable; removal of the screening eliminates this signal loss but increases the RF noise introduced into the system.

**Transfer of the ECG signal**

It is sometimes possible to use a simple filter on the ECG leads to prevent interference being introduced into the MRI system. More common is the use of an optical fibre to transmit the ECG signal. The two ECG leads from the patient are connected to a device which converts the ECG to optical pulses. In Edinburgh a battery-powered module is used with rechargeable nickel-cadmium batteries. This rests at the side of the patient approximately 1.2m from the imaging section and level with the entrance to the imager.

The optical fibre is then connected to a device, this time several meters from the main part of the NMR imaging system, which reconverts the optical pulses to an ECG signal. The signal can then be connected to a modified ECG monitor which produces a trigger pulse at the R wave. A visual display of the ECG has been found to be important to obtain the best position of the electrodes on the patient and also to confirm that the R wave is being detected. In addition the ability to monitor many cardiac patients during the investigation is essential. It must be remembered however that the ECG cannot be considered as diagnostic because of the possible presence of RF and gradient pulses.

**Triggering the image acquisition**

A NMR image is produced from a number of NMR signals, each obtained with a different phase encoded gradient. In non-gated images the time between each of these signals will be constant for a particular acquisition. However, the time will vary for different types of acquisition typically from about 50ms to 4000ms.

In cardiac gated MRI the trigger pulse initiates a sequence of RF and gradient pulses which produces an NMR signal. The next trigger pulse initiates the production of the next signal and so on until an image is produced. The time from the R wave to the start of each NMR signal production can be altered so that the heart is imaged at any point within the cardiac cycle. RF and gradient interference can produce sharp spikes on the ECG which can be detected as R waves in the ECG monitor producing a trigger pulse. This can be overcome in the software of the MRI system as any such pulses will occur at exact times after the trigger pulse which initiated the signal production. As a consequence pulses detected at such times are ignored.

As in other cardiac gated imaging techniques trigger pulses produced by ectopic beats or from R waves following R–R intervals outwith a predefined range must be ignored. This can be done using a microcomputer or within the main MRI computer itself.

**Applications of cardiac gated NMR imaging**

In other cardiac imaging techniques there is usually only one parameter which influences the image intensity. In nuclear cardiology it is the isotope concentration, with ultrasound it is tissue reflectivity, and in computerized tomography it is X-ray absorption. In MRI the situation is quite different as we have three separate parameters which can influence the image produced. It is generally true to say that images that are essentially images of proton density can be used to demonstrate anatomy. Pathology, on the other hand, can be imaged using the $T_1$ or $T_2$ relaxation parameters.

A cardiac gated image of the heart containing mainly proton density information is shown in figure 2. A spin echo sequence has been used which produces good contrast between blood and cardiac tissue due to the motion of the blood. Such an image would be used to identify abnormal anatomy within the heart or to calculate ventricular volumes and parameters such as ejection fraction and stroke volume [8]. The trigger pulse after the R wave can initiate a number of NMR signals occurring at different parts of the cardiac cycle. The sequence of images so produced, usually 8 or 16, can be displayed in a cine loop mode at the end of acquisition.

![Figure 3. A cardiac gated $T_1$ map showing mainly the left ventricle at end systole. The scale on the right-hand side of the image is calibrated in milliseconds. Generally such images are viewed in colour to highlight increases in the $T_1$ parameter following myocardial infarction.](image_url)
This can be used to demonstrate abnormalities in ventricular wall motion.

Cardiac pathology can be visualised using gated T1 maps as illustrated in figure 3. Two sequences are used to calculate the T1 maps, both of which are gated. An important example is the identification of myocardial infarction which results in a local increase in tissue water due to oedema. The anatomy of the heart may appear quite normal even though the pathology is abnormal. The relaxation parameters T1 and T2 are both prolonged in regions of oedema and consequently will identify regions of recent infarction that would not be shown on a proton density image [9 and 10]. This time the blood has a high signal corresponding to a high T1.

The application of cardiac gating in flow measurements is illustrated in figure 4. Moving tissue will influence the phase of the NMR signal; phase maps can therefore be produced in which the image intensity is directly related to velocity and direction. Fairly accurate estimates of flow can be obtained from these velocity measurements and the technique may rival Doppler ultrasound for some applications in the future. To measure pulsatile flow, whether in blood or the CSF, the sequence must be synchronized with the cardiac cycle.

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References

BRAIN WATER MEASURED BY MAGNETIC RESONANCE IMAGING

Correlation with Direct Estimation and Changes After Mannitol and Dexamethasone

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Summary

Brain water content was measured in tissue samples taken at operation from 19 patients with intrinsic cerebral tumours imaged preoperatively by magnetic resonance. A high correlation (r = 0.94, p < 0.0001) between white matter content and the longitudinal relaxation time (T1) enabled water content to be estimated from T1 to within 4%, 11 patients received dexamethasone and improved clinically but their T1, and thus brain water content, was unchanged an average of 6 days after infusion of 20% mannitol in 11 patients significantly reduced T1 in oedematous white matter and tumour within 15 min of administration, and by 30 min the T1 of oedematous white matter had fallen to a mean of 32.4 (SEM 7.1) ms, corresponding to a reduction in water content of 1-4 (0.3)%.

Introduction

In 1919 Weed and McKibben showed that the brain bulk of cats could be increased by intravenous infusion of distilled water and decreased by hyperosmotic (30%) saline or saturated sodium bicarbonate solution. They reported that the cortical capillaries were distended and the intercellular spaces were diminished, but did not examine changes in water content. What are the physical effects of osmotic agents on the brain? Some workers argue that mannitol does not decrease brain water content, but rather changes intracranial vascular caliber and reduces ventricular cerebrospinal fluid (CSF) pressure.

Others report measurements suggesting that mannitol can reduce brain water content.

In cerebral neoplasms there is relief of symptoms after administration of dexamethasone, and clinical improvement is seen within 24 h. Initially it was thought that steroids produced immediate clinical improvement by reducing the oedema around the tumour, but papilloedema was noted to be slow to resolve. The water content of oedematous brain biopsy samples from untreated tumour patients is higher than in those treated with steroids for 4 days, but a rapid fall in water content in an individual patient coincident with clinical improvement has not been shown. Dexamethasone and other glucocorticoids can reduce high intracranial pressure in tumour patients, but the reduction lags behind steroid administration and clinical improvement by days, although there is evidence that the viscoelastic properties of brain tissue are changed within 24 h of the start of steroid therapy. In acute head injury, no such changes are seen.

Computed X-ray transmission tomography (CT) has been used to assay brain water content changes after dexamethasone, and although some investigators have seen changes of about 3 Hounsfield units in oedematous white matter 48 h after starting dexamethasone, changes in density of this order are close to the resolution limits of CT and are influenced by other factors besides water content. Magnetic resonance imaging (MRI) is a better technique than CT for the investigation of brain water. The longitudinal relaxation time (T1) is mainly determined by the ratio of "free" to "bound" water within tissue. The expected T1 changes have been shown after alcohol and vasopressin intake, and T1 correlates with brain water content in vivo.

This study was undertaken to confirm the correlation between T1 and brain oedema in patients with intrinsic tumours, to investigate water changes in the brain following mannitol administration, and to examine with MRI the effects of dexamethasone.

Patients and Methods

Brain Water Estimations

Brain water was measured in 16 men and 3 women aged 39-73 years (mean 58) having neurosurgery for intrinsic supratentorial tumours of glial (18 patients) or metastatic (1 patient) origin. The site of the tumour allowed a frontal (7 patients), temporal (5 patients), or occipital (6 patients) lobectomy to be done, and biopsies of peritumoural tissue were taken in 1 patient. The lobectomy specimens were sealed immediately in cosympore sheet (Ethicon, Ltd, Edinburgh) to prevent evaporative water loss, were taken to a laboratory adjacent to the operating theatre, and were cut transversely as close as possible to the plane of a preoperative MRI of the brain. Peritumoural white matter and, where possible, peritumoural cortex, normal white matter, and normal cortex were sampled from the lobe, and the positions of the samples were marked.

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The samples were then placed in a calibrated layered oil density column for specific gravity measurement and the calculation of water content. In 16 patients half of each sample was placed in its fresh state on a small piece of dry aluminium foil that had an accurately predetermined weight. The sample was then dried in an oven at 80°C and reweighed at 24 h intervals until it reached a constant weight, which was then recorded. The wet tissue weight (W) was obtained by subtraction of the foil weight from the fresh weight and the dry tissue weight (D) by subtraction of the foil weight from the dried weight. Wet tissue weights ranged from 9.9 to 484.1 mg, mean 102.8. Water content was expressed as a percentage by weight of the fresh tissue with the formula: % water = (W - D)/W × 100. Wet and dry weighing of normal samples of brain allowed the derivation of the specific gravity of the solids of human grey and white matter. The specific gravity of the tissue solids is used to calculate water content from the wet tissue specific gravity measurements, and wet and dry weighing also allowed correction of the error introduced by protein in the oedema fluid. The interval between operative lobectomy and completion of the specific gravity measurements was kept to less than 20 min in all cases, so as to limit evaporative water loss from the tissue specimens.

**Correlation of** $T_1$ **with Brain Water**

The patients had MRI the day before surgery by a 0.8 T vertical field resistive MRI system (M & D Technology, Aberdeen). Calculated $T_1$ maps were obtained by means of an interleaved saturation recovery and inversion recovery pulse sequence with a sequence repetition time of 1000 ms and a time from inversion of 200 ms. Transverse sections through the tumour were obtained, each with a section thickness of 12 mm and an acquisition time of 256 s. Interpretation of the digitally stored $T_1$ map enabled $T_1$ values to be measured in any region of a tomographic section. Experiments on normal volunteers showed that the precision of $T_1$ measurements in the cortex and white matter was 5% and 3%, respectively.

All patients had their operations within 24 h of MRI, and in 6 patients mannitol was given peroperatively to control intracranial pressure. 13 patients did not receive mannitol during surgery. After operation, measurements of $T_1$ were made from the stored preoperative MRI at sites corresponding as closely as possible to those of brain tissue sampling. The correlation between the measured $T_1$ and the water content of the samples was assessed by least squares regression analysis, and confidence limits were calculated for the regression line for white matter.

**Changes in** $T_1$ **with Dexamethasone and Mannitol**

11 patients with glial tumours were imaged before steroid administration and were scanned on dexamethasone 1 mg 6-hourly by mouth immediately after MRI. They had a second MRI 2–11 days later (mean interval 6), and $T_1$ values for normal cortex, normal white matter, peritumoural white matter, and tumour were measured from the same positions on equivalent sections of the pretreatment and post-treatment scans. All the dexamethasone group had improved clinically by the time the second MRI was done.

17 patients with glial tumours were imaged while having an intravenous infusion of 0.9% saline, after which the infusion fluid was changed to 20% mannitol in 11 patients. Over the next 15 min, 200 ml of infusion fluid was administered to each patient; further MRIs were taken 15 min and 30 min after the control image. On equivalent sections of the control, 15 min, and 30 min images, $T_1$ values for normal cortex, normal white matter, peritumoural white matter, and tumour were measured at fixed sites by two observers who did not know whether the patient had received 20% mannitol or placebo (0.9% saline), or the timing of the scan.

Changes in $T_1$ after dexamethasone or mannitol were assessed by non-parametric statistics. Changes from zero were assessed by the Wilcoxon signed rank test and, in the case of mannitol, from patients receiving a placebo infusion by the Mann Whitney test. Results were regarded as significant if $p<0.05$.

**Results**

**Comparison of Brain Water Estimations**

In the 16 patients for whom there were wet and dry weight brain samples in addition to specific gravity estimations of brain water, gravimetry underestimated tissue water content of oedematous white matter by a mean of 2.4% and of oedematous cortex by a mean of 1.2%.

**Correlation of** $T_1$ **with Brain Water**

In 19 patients there were nine samples of white matter from normal areas of brain and twenty-one from oedematous peritumoural areas. The nine samples of normal white matter had a mean $T_1$ of 298.0 (SD 15.1) ms and water content of 69.7% (0.7%), and these are the lowest 9 points on the graph in fig 1A. There was a high correlation ($r=0.94$, $p<0.0001$) between $T_1$ and water content with narrow 95% confidence limits. From the image of a single patient the $T_1$ of white matter allows estimation of water content to within 4%.

In 17 patients there were twelve samples of cortex from normal areas of brain and eight from oedematous peritumoural areas. The twelve samples of normal cortex had a mean $T_1$ of 464.8 (SD 25.9) ms, and water content of 80.5 (0.8%), and these are the lowest 12 points on the graph in fig 1B. Oedematous cortex was hard to identify in the lobectomy specimens, since the distribution of peritumoural oedema is largely within white matter, as evident on CT as

![Fig 1—Correlation of white matter (A) and cortex (B) water content with $T_1$ values.](image-url)

Open circles, patients who had peroperative mannitol; regression line --- 95% confidence limits ---- (A) linear regression line slope = 0.043, constant term = 57.13; (B) linear regression line slope = 0.031, constant term = 66.98.)
well as NMR images. In addition the $T_1$ of cortex was less easy to determine than that of white matter because of the proximity of the CSF in the subarachnoid space, with a consequent partial volume effect, and the smaller area of tissue for measurement on the NMR images. Nonetheless, there was a correlation ($r=0.65$, $p<0.002$) between $T_1$ and water content of cortex; but confidence limits were not calculated because there were insufficient data from abnormal tissue.

Changes in $T_1$ with Dexamethasone

Dexamethasone had no significant effect on the $T_1$ of normal cortex and white matter, oedematous peritumoural white matter, or the tumour tissue (fig 2A). We therefore concluded that the water content of these tissues was unchanged by dexamethasone and could not explain the clinical improvement (seen in all patients) by any effect of dexamethasone on the extent of brain oedema.

Changes in $T_1$ with Mannitol

Mannitol did have an effect in reducing the $T_1$ of oedematous peritumoural white matter and the $T_1$ of the tumour tissue, but did not have a significant effect on normal white matter or cortex. After 15 min (fig 2B) the change in the $T_1$ of the peritumoural white matter and tumour tissue was significantly less than zero ($-19.6$, SEM 4.0 ms, $p<0.01$; and $-26.9$, SEM 11.3 ms, $p<0.05$, respectively) and for peritumoural white matter it was significantly less than the values obtained during the placebo infusion ($p<0.01$). 30 min after the intravenous infusion of 20% mannitol was started (fig 3) the mean reduction in $T_1$ of the peritumoural white matter was 32.4, SEM 7.1 ms ($p<0.01$) and for tumour was 57.0, SEM 15.6 ms ($p<0.01$). In both peritumoural white matter and tumour the reduction in $T_1$ was significantly greater ($p<0.005$) in patients given mannitol than in those given placebo. The fall of 32.4 ms corresponds to a reduction in water content of 1-4% within the oedematous white matter.

Discussion

This study partly confirms the simple linear correlation between brain water and $T_1$ reported in animal models of cytotoxic and cryogenic oedema. Within a given tissue (cortex or subcortical white matter) brain water is linearly related to $T_1$, but the relaxation behaviour of the water molecules in white matter seems to be different from the behaviour of the water in cortex. The water within normal cortical tissue is largely intracellular, the extracellular fluid space being small. When white matter becomes oedematous, water accumulates outside the myelin sheaths, and although it can permeate between the myelin lamellae, it remains largely extracellular. This extracellular oedema fluid in white matter probably has a high free to bound ratio, because of a low concentration of macromolecules in the extracellular space of white matter.

Dexamethasone does not seem to change $T_1$ in intrinsic tumours and the surrounding oedematous brain, or in the normal brain remote from the tumour, within an average of 6 days of administration. The initial effect of dexamethasone must therefore be mediated by some mechanism other than a change in the water content of the brain. Although resting intracranial pressure is not reduced by steroid therapy for 48 to 72 h, patients are clinically better within 24 h and plateau waves become less frequent. The lumped intracranial elastance, as measured by pressure volume tests, shows a return towards normal over 72 h, but the biggest change is within the first 24 h. In patients with posterior fossa tumours, the same clinical response follows steroid therapy over the same time scale, with the disappearance of plateau waves, and yet in these patients the cause of the raised intracranial pressure is predominantly obstructive hydrocephalus rather than brain oedema. This suggests that the first change is an alteration in the viscoelastic properties of brain tissue. The present study shows that brain water is not reduced at this early stage, and to explain the elastance changes some other mechanism, such as a change in cerebral blood volume or alteration in the structural properties of cell membranes, must be invoked.

The intravenous infusion of mannitol reduces $T_1$, but only in abnormal tissues (tumour and oedematous peritumoural white matter). After rapid intravenous infusion in normal cats, mannitol has been shown to produce considerable pial arteriolar vasoconstriction within 5
minutes of injection, with subsequent vasodilatation at 1 hour. The time course of this vasodilatation closely matches the reduction in intracranial pressure and blood viscosity that mannitol is known to produce. Some workers argue that the blood viscosity reduction is the primary change and an autoregulatory mechanism is responsible for the vasodilatation. When autoregulation is impaired, mannitol does not reduce intracranial pressure.

Mannitol lowers brain oedema by drawing water osmotically from tissues other than normal brain, because of the high hydraulic resistance of the normal cerebral capillary wall to water movement (thirty times higher than muscle capillaries and three thousand times higher than mesenteric capillaries). This would explain our finding of no change in water content in the normal brain regions of our patients, while in tumour and oedematous peritumoural white matter the water content was reduced. In these abnormal regions, the brain-blood barrier is known to be disrupted by the appearance on contrast enhanced CT and pertechnetate radionuclide scans. The hydraulic resistance of the capillaries in these areas will be low, and mannitol would seem from our findings to withdraw water osmotically from these abnormal tissues.

Our results show that it is possible to estimate tissue water from the total water content of the cerebrospinal fluid and the brain. Patients with brain tumours are not treated by reduction of cerebral oedema, and that mannitol does reduce water content of oedematous brain, but not of normal brain.

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MECHANISM OF EXACERBATION OF RHEUMATOID SYNOVITIS BY TOTAL-DOSE IRON-DEXTROX INFUSION: IN-VIVO DEMONSTRATION OF IRON-PROMOTED OXIDATIVE STRESS

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Summary

The mechanism by which a synovial flare occurred in a patient with rheumatoid arthritis after intravenous infusion of iron-dextran was investigated. After the infusion, serum and synovial fluid iron-binding capacity became saturated, giving rise to low-molecular-mass iron chelates with the capacity to cause oxidative damage ("bloomycin-iron"). At the same time lipoperoxidation and the concentration of oxidised ascorbic acid (dehydroascorbate) increased in both serum and synovial fluid, and red-cell glutathione fell. These changes corresponded closely to an exacerbation of rheumatoid synovitis. Hepatic function was transiently disturbed 7 days after the infusion, reflecting hepatic oxidative stress on the iron-loaded liver. Such changes provide clear evidence that iron-catalysed oxidative reactions influence the inflammatory process in human beings.

Introduction

The observation that the rheumatoid synovial membrane contains greater than normal deposits of iron1 led to speculation2-5 that iron-catalysed generation of toxic free radicals may promote rheumatoid synovitis. The superoxide anion radical and hydrogen peroxide, derived from polymorphonuclear leucocytes (PMN) within the synovial cavity or activated macrophages within the synovial membrane, react together in the presence of transition-metal catalysts (eg, iron) to form the highly reactive hydroxyl radical.4,6 This radical is thought to oxidise lipids,7 proteins,8 ascorbic acid,9 and other biomolecules within the joint and thus perpetuate inflammation. The oxidative reaction depends on the presence of iron in a suitable form, such as a low-molecular-mass complex (eg, ADP·Fe).10 As long as iron in synovial fluid or serum is bound to transferrin (serum transferrin is normally about 30% saturated) catalysis of hydroxyl radical formation will not take place.14 Iron with the capacity to cause oxidative damage can be detected by the bloomycin assay.16

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Demonstration of pulsatile cerebrospinal-fluid flow using magnetic resonance phase imaging

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Abstract

The study of pulsatile cerebrospinal-fluid (CSF) flow may be useful in diagnosis of certain forms of intracranial disease. Previous techniques used to study CSF flow either are invasive or do not allow accurate measurement. Magnetic resonance imaging (MRI) offers a non-invasive method of studying the CSF pathways. Our technique uses MR phase images and allows quantitative measurement of flow velocities and volume-flow rates. Four volunteers were studied at the level of the second cervical vertebra (C2). The MRI pulse sequence was gated from the R-wave of the subject's electrocardiogram and 12 scans were taken, corresponding to different times in the cardiac cycle. The variation in flow velocity throughout the cycle was plotted, and maximum caudal and cephalad flow velocities and flow rates were calculated. Good agreement was found between three of the four volunteers. The mean maximum caudal velocity was 2.91 cm s⁻¹ occurring at a mean time of 190 ms after the R-wave. This corresponds to a mean maximum flow rate of 4.13 ml s⁻¹. The total imaging time for each study was about 1 h. Technical developments, allowing simultaneous acquisition of several images throughout the cardiac cycle, will reduce this time significantly.

The cerebrospinal-fluid (CSF) pathways are particularly difficult to study, and to obtain reliable information the haemodynamic conditions must not be altered, for example, by surgical intervention. Radiologists have, for a long time, been aware of the pulsatile nature of CSF flow at fluoroscopy during Pantopaque myelography and in cineradiography of myelography and encephalography (du Boulay, 1966; du Boulay et al., 1972). Estimations of CSF flow using these techniques, which were derived from observation of a television monitor, were inevitably imprecise. The appearance of flowing fluid using magnetic resonance imaging (MRI) has been the subject of much research, and various techniques have been developed to demonstrate the velocity of blood flow. The effects of blood flow on magnetic resonance (MR) images are dependent on the pulse sequence used, and on the velocity profile of the flow. They can be interpreted as either an increase or decrease in signal compared with stationary material (Axel, 1984; George et al., 1984). The precise variation in signal intensity as a function of velocity is dependent on many factors and is thus difficult to predict. The nuclear magnetic resonance signal has two components which are referred to as the real and imaginary parts of the signal. These two components can be used to calculate both the modulus and the phase of the signal for each pixel. It is the modulus value that is normally used to obtain the standard clinical MR images. It has been shown, however, that phase images can also be used to quantify flow velocities (Moran et al., 1985). Whereas the modulus MR image contains information on proton density, relaxation time parameters (T₁ or T₂) and flow effects, the phase image contains information on the velocity and direction of flow, as well as phase shifts due to magnetic-field inhomogeneity, chemical shifts, and changes in magnetic susceptibility. The velocity information is usually encoded by adding extra gradient pulses into the pulse sequence (Bryant et al., 1984; Van Dijk, 1984). Alternatively, the bipolar form of gradient pulses which are part of the standard imaging sequence, such as those used for slice selection, can be exploited to provide velocity encoding (Ridgway & Smith, 1986). The range of velocities to be measured can be altered by changing the temporal separation of the two components of the bipolar pulse. Thus, peak velocities in the ascending aorta, and slower venous flow, can both be imaged using the same technique.

Pulsatile CSF flow has been demonstrated in the cerebral aqueduct and pontine cistern using MRI by showing a variation in signal intensity over the cardiac cycle (Bergstrand et al., 1985). An inversion-recovery pulse sequence was used in this study and though a cyclical variation in intensity was shown, no direct quantification of velocity or direction of flow was possible. Phase imaging has not previously been used to image pulsatile CSF flow.

The study describes the extension of a phase imaging technique which was originally developed to measure arterial flow in the body. The technique was modified to measure slow flow in the head and neck.

Method

This study was performed using a 0.08 T resistive MRI system manufactured by M&D Technology Ltd.
The imaging technique acquires two phase images, each having different velocity encoding, achieved by varying the temporal separation of the dephase and rephase components of the slice selection gradient (Ridgway & Smith, 1986). The final velocity image is obtained by subtracting these two images, thus eliminating any phase shifts due to field inhomogeneity, chemical shifts or changes in magnetic susceptibility. The technique was modified to image at lower velocity ranges by extending the separation of the bipolar slice selection gradients in one of the velocity-encoding pulse sequences. The difference in separation for the two pulse sequences was thus extended from 2 ms to 12 ms. The maximum measurable velocity, corresponding to a phase shift range of ±180°, was thus decreased from 120 cm s⁻¹ to 20 cm s⁻¹ making finer velocity resolution possible. Further modifications involved the inclusion of stronger gradients and the use of a smaller-diameter RF coil, used routinely for head imaging.

The flow measurement was calibrated for mean flow velocities in the range 1–11 cm s⁻¹. A simple continuous-flow phantom was used, incorporating a tube of inside diameter 10 mm, containing copper sulphate solution with \( T_1 = 250 \) ms. The phantom was fed from a constant head tank and mean flow rates were calculated by finding the time taken for a known volume to pass through the phantom. The MRI pulse sequence was triggered from an external source at a repetition rate corresponding to a heart rate of 75 beats min⁻¹. Each scan was repeated three times and the flow rate recorded before and after each scan.

Four volunteers were imaged, gating the pulse sequence with the subject's electrocardiogram (ECG) (Smith et al, 1986). Heart rates were monitored and images were obtained with delays of 110, 140, 165, 190, 215, 240, 290, 390, 490, 590, 690 and 790 ms after the subject's R-wave. More images were obtained during the systolic period where the velocity variation appeared to be more rapid. The total imaging time for each subject was about one hour.

Prior to this study, one volunteer was imaged at 12 levels in the head and neck at the time during the cardiac cycle when the peak velocity was attained. The result images suggested that the peak velocity was higher in the neck than in the head. Imaging too low in the neck resulted in increasing loss of sensitivity with distance from the head coil. As a compromise the level chosen for the four volunteers was at the level of the second cervical vertebra (C2).

The mean phase shift within the ring-shaped subarachnoid space was calculated using two regions of interest: the outer represented the boundaries of the spinal canal and the inner, the spinal cord. From these two regions the mean phase shift within the subarachnoid space and its area were determined. The calibration curve was used to determine the velocity from the phase shift, and the area of the subarachnoid space was used to calculate flow.

In order to assess the reproducibility of the technique, expressed as a coefficient of variation, one volunteer was scanned six times at the same level at the time of peak velocity.

**RESULTS**

The results of the velocity calibration are plotted in Fig. 1, showing 95% confidence limits at ±0.92 cm s⁻¹. There was a significant correlation (\( r = 0.998, p < 0.001 \)) between the phase shift (degrees) and the flow velocity (cm s⁻¹).

A representative transverse section from one of the normal volunteers is illustrated in Fig. 2, showing both the modulus and the phase images. The CSF flow in the subarachnoid space can be seen on the phase image as a ring-shaped area. The displayed intensity of the phase image is related to velocity. Velocity in the caudal direction is represented by lighter shades of grey which correspond to a positive phase shift (Fig. 3a). Similarly, velocity in the cephalad direction is represented by darker shades of grey which correspond to a negative phase shift (Fig. 3b). Mid-grey represents zero velocity, corresponding to zero phase shift.

All volunteers demonstrated pulsatile flow within the subarachnoid space during the cardiac cycle and a representative flow-velocity curve is shown in Fig. 4.
Pulsatile CSF flow demonstrated by MR phase imaging

Representative section of those used in the volunteer study showing (a) the modulus image and (b) the phase image. The CSF flow is seen on the phase image as a ring of increased intensity (arrows). Also seen are left and right internal jugular veins.

Maximum flow rates, maximum flow velocities and time to peak flow for all subjects are listed in Table I. At intervening times during the cardiac cycle, flow velocities were relatively small and their corresponding phase shifts were not significantly different from phase shifts measured in adjacent soft tissue. The maximum variation in heart rate in any one volunteer was from 65 to 83 beats min^{-1}.

The reproducibility of the repeated scan, analysed once, was 14.4%. However, this figure includes the error associated with the estimation of phase information from a single image which was found to be 8.7%. The reproducibility of the technique itself is therefore 11.5%.

**DISCUSSION**

This study not only confirms the ability of MRI to demonstrate pulsatile CSF flow, but also demonstrates

Two representative phase images demonstrating the pulsatile nature of the CSF flow: (a) shows caudal flow (positive phase shift); (b) shows cephalad flow (negative phase shift).

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that phase imaging enables quantitative measurement of flow velocity to be obtained with reasonable accuracy. In addition, the direction of flow can be assessed. The technique therefore has advantages over methods which rely on signal intensity (Bergstrand et al, 1985). However, unlike Bergstrand, we have been unable to demonstrate flow within the cerebral aqueduct, though this may be due, in part, to the lower signal-to-noise ratio that is obtained with the lower-field system used in this study.

There is good agreement between the flow rates and velocities obtained in three out of our four normal subjects. Shorter sampling intervals would enable these values to be established more accurately; currently the "peak" caudad and cephalad velocities could be underestimated. The variation in the time to peak CSF flow velocity may be due to many factors including the sampling time, i.e. time after R-wave, respiratory rate or variation in heart rate.

At present, the total imaging time for each subject is probably too long for routine clinical use. Simultaneous multislice acquisition would substantially reduce total imaging time but the spatial location of each section would be different. Alternatively, the acquisition of multiple images of a single section throughout the cardiac cycle, as has been done in cardiac imaging (Waterton et al, 1985) would also reduce imaging time.

This technique offers potential for the investigation of disorders involving the CSF circulation. In the case of complete obstruction, for example, pulsation distal to the obstructing lesion will stop, but it will persist on the cranial side. In addition, du Boulay (1966), using myelographic techniques, observed that when the subarachnoid space was narrowed but not obliterated the amplitude of movement could be considerably exaggerated. Magnetic resonance imaging offers a non-invasive method of studying the direction and velocity of CSF flow in such patients.

**Acknowledgments**

This research is generously funded by a grant from the Scottish Home and Health Department. We would like to

**TABLE 1**

<table>
<thead>
<tr>
<th>Subject</th>
<th>Max. caudad flow velocity (cm s⁻¹)</th>
<th>Max. cephalad flow velocity (cm s⁻¹)</th>
<th>Time of max. cephalad flow after R-wave (ms)</th>
<th>Max. caudad flow rate (ml s⁻¹)</th>
<th>Max. cephalad flow rate (ml s⁻¹)</th>
<th>Time of max. cephalad flow after R-wave (ms)</th>
<th>Approximate mean R-R interval (ms)</th>
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<td>3.17</td>
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<tr>
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<td>1.74</td>
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<td>4.96</td>
<td>2.99</td>
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<td>0.55</td>
</tr>
<tr>
<td>Mean</td>
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<td>1.47</td>
<td>190</td>
<td>4.13</td>
<td>2.24</td>
<td>190</td>
<td>—</td>
</tr>
<tr>
<td>SD</td>
<td>0.78</td>
<td>0.31</td>
<td>20</td>
<td>1.15</td>
<td>0.55</td>
<td>20</td>
<td>29</td>
</tr>
</tbody>
</table>

**FIG. 4.**

Variation of CSF flow velocity (cm s⁻¹) plotted against time after the R-wave (ms). The set of measured points are shown as filled circles. The same set of points was plotted (open circles) in adjacent ECG cycles to show the continuity of the curve. In this case the average R-R interval was 800 ms (corresponding to a heart rate of 75 beats min⁻¹).
Pulsatile CSF flow demonstrated by MR phase imaging

thank Mrs C. N. Rowan for secretarial services and Mr I. Lennox of the Department of Medical Illustration. We would also like to thank Professor J. J. K. Best and Dr D. M. Kean for their helpful comments and support.

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Correspondence
(The Editors do not hold themselves responsible for opinions expressed by correspondents)

Magnetic resonance flow imaging: a possible method for distinguishing communicating syringomyelia from cystic intraspinal lesions

The Editor—Sir,

The important radiological differential diagnosis of syringomyelia is an intramedullary tumour. Magnetic resonance (MR) is now the modality of choice to image the spinal cord and canal (Han et al, 1983; Yeates et al, 1983; Norman et al, 1983). However, differentiation between the cerebrospinal fluid (CSF) contained within a syrinx and a cystic tumour is a potential problem, as there may be little difference in the relaxation parameters ($T_1$ and $T_2$) or in the morphology. One possible means of differentiating these conditions is to study CSF flow using a MR phase imaging technique (Ridgway & Smith, 1986; Ridgway et al, 1987), which has recently been developed to allow quantitative measurement of flow velocities, including low values found in CSF flow.

We recently studied a 38-year-old lady who presented with a history of mild spinal-cord disturbance, suggestive of a demyelinating disease. The MR scan showed a probable syrinx extending from the level of the first cervical vertebra to at least the second thoracic vertebra, with some associated tonsillar herniation (Fig. 1). Although in this case the demonstration of a coexisting Chiari malformation established the diagnosis, phase images were obtained in order to detect CSF flow within the syrinx. These were obtained in the transverse plane at the level of the second cervical vertebra; the pulse sequence was gated from the R-wave of the subject's electrocardiogram and images obtained at 50, 225, 300 and 325 ms following the R-wave. Both the time intervals and choice of level corresponded to those used in a previous study carried out on normal volunteers (Ridgway et al, 1987). The cross-sectional area of the syrinx was measured from the images and maximum caudal and cephalad flow rates and velocities were calculated.

Pulsatile CSF flow was demonstrated within the syrinx during the cardiac cycle (Fig. 2). A maximum caudal flow rate of 1.27 ml s$^{-1}$ was achieved at 300 ms following the R-wave. This corresponds to a maximum velocity of 1.63 cm s$^{-1}$, which was calculated from the area of the syrinx (0.87 cm$^2$).

Cephalad flow was maximum at 30 ms with a flow rate of 0.64 ml s$^{-1}$ and a velocity of 0.86 cm s$^{-1}$. Due to cord expansion, the subarachnoid space at the level imaged was too small for accurate measurement of CSF flow outside the syrinx to be obtained.

Magnetic resonance phase imaging offers a means of studying the velocity and direction of flow in both normal and pathological CSF circulation. It may prove to be useful non-invasive technique to distinguish communicating and non-communicating forms of syringomyelia, including those associated with spinal-cord tumours, posterior-fossa tumours or cysts, and spinal-cord trauma.

We would like to thank Dr G. Venables for valuable clinical assistance.

Yours, etc.

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Fig. 1.

Sagittal proton density image showing fusiform enlargement of the spinal cord. The syrinx cavity is identified as a low-intensity area, vertically orientated, running throughout the length of the cord.

Fig. 2.

Transverse phase image at the level of the second cervical vertebra. The flow within the syrinx is seen as an oval area of increased intensity (arrow). Blood flow within the right deep cervical vein is noted.
Correspondence

Avoidance of ring artefacts in lumbar spine computed tomography in obese patients

THE EDITOR—Sir,

Central ring artefacts (Kowalski & Wagner, 1977) are sometimes unavoidable on computed tomography (CT) in very obese patients, even when meticulous care has been taken with calibration (Seeram, 1982). This is because of low photon transmission and consequent low signal-to-noise ratio (Haaga et al., 1981). The spinal canal is often at the precise centre of the axis of tomographic rotation and, thus, especially liable to be obscured by such artefacts (Fig. 1). This can be avoided by purposely positioning the patient slightly off centre within the gantry aperture so that the spinal canal is removed from the axis of rotation (Fig. 2).

Quantum mottle on the initial lateral scanogram is indicative of low photon transmission and provides a clue that deliberate off-centring may be beneficial.

Yours, etc.,

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Fig. 1. Computed tomogram of the L5/S1 disc of an extremely obese patient (W = 1024, L = +45). A central field of reconstruction has been used with a magnification factor of 4.2. The central ring artefacts obscure detail of the disc.

Fig. 2. The patient has been moved 3.5 cm to the right. The ring artefact is now over the left sacral ala. The degenerate L5/S1 disc is seen more clearly.
Cerebral and Brain Stem Changes After ECT Revealed by Nuclear Magnetic Resonance Imaging


Nuclear magnetic resonance images of the brain were obtained in fourteen patients with major depression during a course of ECT. The \( T_1 \) relaxation time rose immediately after the fit, reaching a maximum 4–6 h later. The \( T_1 \) values then returned to their original level; no long-term increase occurred over the course of treatment. These results are consistent with an extensive but temporary breakdown of the blood-brain barrier during ECT.

Nuclear magnetic resonance (NMR) is a long established technique in physics and chemistry, but relatively new in medicine. Nuclei in a constant magnetic field can be made to resonate by the application of an oscillating magnetic field in the radiofrequency range. As the resonance decays to zero the radiofrequency radiation emitted by the nuclei can be detected; this is the basis of the NMR signal. The time taken for the nuclei to return to their original equilibrium position is the spin lattice relaxation time, \( T_1 \).

In medical NMR imaging, the signal is usually obtained from hydrogen nuclei (protons) within the body. Since most biological systems consist of 70–90% water, it is these water protons that cause the major component of the NMR signal. Solid tissues have short \( T_1 \) values, while relatively fluid tissues have longer ones. The \( T_1 \) of grey matter is significantly longer than that of white matter because of their different fat:water ratios. \( T_1 \) times can be altered by changes in the ratio of 'free' to 'bound' water in a tissue (Mathur De Vre, 1984) or in its total water content (MacDonald et al., 1985). NMR is therefore a useful tool for investigating possible changes in the brain during a course of ECT, particularly as there is no ethical objection to scanning patients repeatedly. Information obtained in this way about changes in the brain induced by ECT may also shed light on the intracerebral changes occurring after spontaneous epileptic fits.

Method

Eleven patients (4 male, 7 female, mean age 49.6 years) had an NMR scan before and after the administration of the first treatment (scans 1 and 2) and again before and after a later treatment (scans 3 and 4) of a course of ECT. A further three patients (all female, mean age 46.3 years) had a scan before and after their last treatment and then six further scans during the 6 h following the treatment. One of these had a final scan after 25 h. Of the total group of fourteen patients, eleven had bilateral and three had unilateral electrode placings; all of them satisfied Research Diagnostic Criteria for a major depressive episode (Spitzer et al., 1978). Patients were excluded if there was a history of serious head injury, brain damage, alcohol abuse or previous radiological abnormality. None had received ECT in the previous 6 months. A standard anaesthetic procedure was used: methohexisone (60–100 mg) and suxamethonium (30–50 mg). Four patients also received diazepam (2.5–10 mg i.v.) before scan 2. The treatment was delivered using an Ectron CC-100 machine, except in one case where a Mecta machine was used. The post-ECT scan was performed as soon as possible after the patient had recovered from the effects of the anaesthetic.

An M&D Technology Ltd 0.08 T resistive magnetic resonance imaging system was used. For each scan, calculated \( T_1 \) images were obtained in three planes: a transverse section 10 mm above the maximum diameter of the lateral ventricles, a midline sagittal section of the brain stem, and a coronal section through the parietal and temporal lobes. All sections were 12 mm thick, and the pixel size was 2 mm\(^2\). Measurement of mean \( T_1 \) of the entire area of the transverse section (the grey and white matter together but excluding the cerebrospinal fluid (CSF)) was made. The area of each hemisphere to be sampled was delineated separately using an irregular region of interest; a cursor was used to draw the boundary of the area to be measured directly onto the image of the section as it was displayed on the monitor. The mean \( T_1 \) within the region of interest was calculated and displayed. CSF has a high \( T_1 \), and is white on the coloured display. It is therefore possible to exclude sulcal CSF by ensuring that it is not included within the boundary of the area to be sampled. In addition, the computer was programmed to present the mean \( T_1 \) of the tissues in this area that were within the range 60–600 ms. This range includes grey and white matter, but not CSF, which has a much higher value. This latter method also ensures that in the small regions of interest sulcal CSF is not included in the samples of grey matter that were analysed. Individual small areas of \( T_1 \) in the pons, mid-brain (at the level of the
optic chiasma and posterior corpus callosum (sagittal section), frontal and occipital lobes (transverse section) and parietal and temporal lobes (coronal section) were also measured. \( T_1 \) was measured at the mid-point of each area. In each case the sample measured had a volume of 1.41 cm\(^3\), except the mid-brain and posterior corpus callosum (0.42 cm\(^3\)).

To control for the effects of the general anaesthetic, five volunteers (three male, two female, mean age 32.6 years) had a transverse section scan before and 30 minutes after an identical anaesthetic procedure.

### Results

Patients had a mean of 5.4 treatments given at a mean frequency of 2.4 per week prior to scan 3. As those having unilateral ECT did not show any lateralisation of change in \( T_1 \), unilateral and bilateral treatments have been analysed together. There was no significant change in \( T_1 \) in any of the areas examined between the beginning and the end of the course of ECT, i.e. when scan 1 was compared with scan 3 and scan 2 with scan 4.

In the transverse section there were significant differences in the mean \( T_1 \) of the whole section before and after the individual treatments, i.e. between scans 1 and 2 and between scans 3 and 4 (Table I). The mean change between scans 1 and 2 was 6 ms (\( P < 0.01 \)). The rise after bilateral ECT was larger (7.3 ms) than after unilateral ECT (4.0 ms), although this was not significant, probably because of the small numbers in the study. There was a smaller mean increase in \( T_1 \) of 4 ms between scans 3 and 4 (\( P < 0.05 \)); bilateral ECT produced a larger increase (6.7 ms) than unilateral ECT (1.3 ms), although again this difference was not significant. Patients were scanned sooner after the last ECT (15.8 ± 4 min, range 13–28 mins) than after the first (30.9 ± 16 min, range 15–62 mins), probably because by then they were less depressed and also familiar with the technique (\( t = 2.69, \) d.f. = 19, \( P < 0.02 \)); this suggested that the maximum rise in \( T_1 \) may not have occurred within 15 min of treatment. This was shown to be so in the three patients scanned six times during the six hours following ECT (Fig. 1). In the patient scanned after 25 h the \( T_1 \) time had returned to below its starting value.

The same anaesthetic sequence in the five controls was associated with a non-significant decrease in mean \( T_1 \) from 340 ms (s.d. 9 ms) to 336 ms (s.d. 8 ms).

Of the nineteen individual small regions in the grey and white matter, \( T_1 \) was found to increase in ten between scans 1 and 2; there was a decrease in seven, and no change in the other two. This increase was statistically significant in the frontal and occipital white matter (\( P < 0.05 \)) and in the left temporal grey matter (\( P < 0.01 \)). For the 8 patients having bilateral ECT only, there was a significantly decreased \( T_1 \) in the right occipital grey matter (\( P < 0.01 \)). Between scans 3 and 4 the \( T_1 \) in eleven regions increased; there was a decrease in six and no change in the other two.

### Discussion

The mean \( T_1 \) of the transverse section showed a significant increase after treatment, both at the start and end of a course of ECT, reaching a maximum 4–6 h after the fit. This was shortlived, and no cumulative effects of ECT were noted. The increases reported in this study are likely to be underestimates.
of the changes produced by ECT itself, since anaesthesia alone was associated with a fall in $T_1$.

Measurement of the 19 small regions described above added little further information. Most of the $T_1$ values in the transverse and coronal sections (20 of 32) increased following treatment. Because there were 64 comparisons involved, it is likely that some of the significant differences arose by chance.

The brain stem showed a non-significant fall in $T_1$ in five of the six measurements. This may be due to the anaesthetic having a greater effect in this area, and is especially interesting since the brain stem reticular formation is involved in arousal.

Interpretation of changes in $T_1$ is difficult. There is no good evidence that ECT causes brain damage (Meldrum et al., 1974; Galloway et al., 1981), and recently necroscopy could reveal no pathology attributable to ECT in a patient who had received over 1250 treatments (Lippmann et al., 1985). Similarly, it is unlikely that these $T_1$ changes represent inflammation or brain damage, since they are reversed so quickly. It is more likely that they are the result of an increase in the total water content of the brain. It is known that ECT produces a temporary breakdown of the blood-brain barrier (BBB) and that this is probably due to the concomitant increase in blood pressure (Bolwig et al., 1977a) and cerebral blood flow (Bolwig et al., 1977b). It is also known that repeated convulsions at short intervals produce cerebral oedema (Laursen et al., 1985). It has been shown that macromolecules leak into cerebral tissue during the temporary breakdown of the BBB induced by ECT (Bolwig et al., 1977a). This would cause an increase in the relative osmotic pressure of the brain.

After restoration of the BBB, movement of water from blood vessels into the tissues would therefore occur, and this would provide an explanation for the rise in $T_1$ over time. It is likely that similar changes occur after spontaneous epileptic fits, although the possibility that the observed changes in $T_1$ are induced by the electrical stimulus rather than by the subsequent convulsive activity in the brain cannot be excluded.

More work is needed to determine the duration and magnitude of the change in $T_1$ and to find out if it is related to clinical improvement or the side-effects of treatment.

Acknowledgements
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References


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Serial changes in the $T_1$ magnetic relaxation parameter after myocardial infarction in man

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D P BONO,* J J K BEST,‡; A L MUIR*

From the Nuclear Magnetic Resonance Imaging Unit (Departments of Medicine; Medical Radiology, and Medical Physics and Medical Engineering, University of Edinburgh), Royal Infirmary, Edinburgh

Summary A low field resistive nuclear magnetic resonance imaging system (0.08 Tesla) was used to study the in vivo changes in the relaxation parameter $T_1$ of the left ventricular myocardium from the first day to six months after acute myocardial infarction in 41 consecutive patients admitted to a coronary care unit. $T_1$ maps were constructed from transverse and coronal images at various times after infarction. Thrombolytic treatment had been successful in 28 patients. Thirty three of the 34 patients studied within two weeks of infarction had a significantly increased $T_1$ value but this developed only after the third day in four. At day 1–3 the mean (SD) maximum $T_1$ was 413 (29) ms (n = 23) compared with 430 (41) ms (n = 22) at day 4–7, 433 (35) ms (n = 24) at day 8–14, 420 (34) ms at one month (n = 22), 388 (39) ms (n = 20) at three months, and 361 (24) ms (n = 14) at six months. The number of regions of interest with an increased $T_1$ followed a similar time course. Although the increase in $T_1$ measured at three months correlated with the initial maximum creatine kinase and with the left ventricular ejection fraction measured at one month, the number of regions with abnormal $T_1$ from day 4 through to one month correlated best with left ventricular ejection fraction. There was no significant difference in $T_1$ between patients with or without reperfusion. The rise in $T_1$ over the first few days together with the prolonged time course of $T_1$ increase suggests that the increase in $T_1$ may reflect cellular infiltration as much or more than tissue oedema.

It has been suggested that the magnetic relaxation parameters $T_1$ and $T_2$, which are associated with an increase in free water content may be used to demonstrate tissue abnormalities.1 After experimental canine myocardial infarction in vitro $T_1$ and $T_2$ were prolonged, and these changes correlated well with increases in tissue water content.2,3 In vivo prolongation of both $T_1$ and $T_2$ after myocardial infarction has been confirmed in dogs4,5 and recently in human beings.6,7 In dogs the changes begin within 30 minutes of coronary occlusion8 and become more pronounced with occlusions of one, two, and three hours.2,3 Recently, Plughfelder et al described the serial changes in the intensity of the in vivo signal in canine myocardial infarction5 but similar studies in human beings have not been reported.

The purpose of the present study was primarily to define the time course of changes in $T_1$ after acute myocardial infarction in human beings.

Patients and methods

We studied 41 patients with transmural myocardial infarction that was confirmed by both electrocardiographic and creatine kinase changes. Whenever possible, imaging was carried out on day 1–3, day 4–7, day 8–14 and one month, three months, and six months after infarction. Serial imaging was performed on 2–6 occasions (mean 3.4 per patient); however, in two patients adequate images were obtained on one occasion only. Consecutive patients were selected provided that the imager was available, the patient consented, and none of the following exclusion criteria was present: current symptomatic pulmonary oedema, continuing chest pain, major

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arrhythmia within 12 hours, previous infarction in the same electrocardiographic territory, or any myocardial infarct in the preceding year.

The study was approved by the local ethics review committee and all patients gave written informed consent.

Ten patients did not have thrombolytic treatment and in a further three thrombolysis was found to be unsuccessful at subsequent coronary angiography (group 1). The remaining 28 patients (group 2) had successful coronary thrombolysis with intravenous anisoylated plasminogen streptokinase complex (n = 23) or recombinant tissue type plasminogen activator (n = 5) (table 1).

The study population was heterogeneous in terms of the site of infarct, rise in creatine kinase, and the use and success of thrombolytic treatment (table 1). Patients on thrombolytic treatment underwent early coronary angiography to determine whether the infarct related vessel was patent, and left ventricular function was assessed by gated blood pool scan.10 In all patients creatine kinase activity was measured twice a day during their stay in the coronary care unit.

The number of patients in group 1 (n = 13) was smaller than that in group 2 (n = 28) because the use of thrombolytic treatment increased during the study and because non-reperfused patients tended to be less well and were therefore more likely to be excluded from this study.

MAGNETIC RESONANCE IMAGING TECHNIQUE

Patients were imaged with a 0-08 T vertical field resistive magnetic resonance imaging system, manufactured by M & D technology, installed in the Edinburgh Royal Infirmary,11 and situated between the coronary care unit and the angiography laboratory. Calculated T₁ maps were obtained by an interleaved saturation-recovery and inversion-recovery pulse sequence with a time from inversion (TI) of 200 ms. Inversion was obtained with an adiabatic fast passage inverting pulse rather than the usual 180° pulse; this is particularly suitable for precise T₁ calculation because it efficiently inverts all the nuclei within the section. In addition, it is not slice selective which means that it inverts all the nuclei within the heart and therefore does not have to be synchronised with the cardiac cycle.

The patient’s electrocardiogram was used to trigger the imager such that the 90° pulse in both the saturation-recovery and inversion-recovery sequences occurred in the same part of the cardiac cycle.12 A field gradient echo of 20 ms was used. The minimum time from the R wave trigger at which imaging can be performed is 230 ms, principally because of the time from inversion (TI) values used. Purdy as a consequence of this, but also to obtain tomographic sections of maximum myocardial thickness, images were obtained towards end systole. The pixel size was 3-5 mm × 3-5 mm.

In an attempt to obtain comparable sections on serial studies for individual patients we used the following routine. After attachment of the electrocardiographic leads we used an alignment marker to position the patient on the couch. The lower edge of the xiphisternum was used as the reference level. The distance of subsequent couch movements was recorded from the digital readout provided by the system. Initial images were obtained with a 16 s ungated saturation-recovery sequence as an anatomical guide for the sections to be scanned. Having optimised the image position and recorded its distance from the xiphisternum we performed a gated scan with a slice thickness of 10 mm. To encompass the whole left ventricle, further scans were performed at two or three 15 mm levels cranial or caudal to this or both, and the position was noted to allow equivalent sections to be obtained at subsequent imaging.

A single coronal section through the left ventricle was obtained. The most suitable imaging slice was again selected from rapid ungated images, but in this case the position was adjusted by slightly altering the strength of the magnetic field. Despite the measures described above, it is likely that changing left ventricular geometry over the six month period prevented serial imaging of exactly the same portion of myocardium. For this reason we used the area of highest T₁ found in any region of interest in the infarct territory for analysis.

The imager was triggered by each R wave for heart rates <75/min and by alternate R waves for faster heart rates. Patients who were scanned during the first 48 hours while in the coronary care unit were moved on a trolley equipped with a defibrillator, and electrocardiographic monitoring was continued throughout.

Table 1 Data (mean (1 SD)) on the study population

<table>
<thead>
<tr>
<th></th>
<th>Group 1</th>
<th>Group 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coronary occlusion (proven or assumed)</td>
<td>Coronary reperfusion (proven vessel patency)</td>
<td></td>
</tr>
<tr>
<td>No of patients</td>
<td>13*</td>
<td>28</td>
</tr>
<tr>
<td>Anterior infarction</td>
<td>5</td>
<td>19</td>
</tr>
<tr>
<td>Inferior infarction</td>
<td>8</td>
<td>9</td>
</tr>
<tr>
<td>Age (years)</td>
<td>57 (10)</td>
<td>53 (10)</td>
</tr>
<tr>
<td>Peak creatine kinase (U/l)</td>
<td>1889 (800)</td>
<td>1947 (1334)</td>
</tr>
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</table>

*Confirmed at angiography1 or assumed because thrombolysis was not used.
Magnetic resonance $T_1$ changes after myocardial infarction

**Data Analysis**

For analysis of $T_1$ values, the myocardium was divided into six sections for each image containing the left ventricular myocardium and cavity. Figure 1 shows the areas into which the myocardium was divided. $T_1$ values for each of the six sections of every image were obtained by using the computer to draw circular regions of interest within the myocardium. The largest circle that could be placed clearly within the myocardium was used. The area of interest selected was either 0.26 or 0.51 cm$^2$, but where wall thinning was considerable the smallest available region of interest (0.08 cm$^2$) was used. This gives a $T_1$ value of the full thickness of the ventricular wall but with a greater contribution from the central portion of the myocardium. In nine patients there was considerable thinning of the myocardium in the infarct region (eight anterior, one inferior). To avoid the possibility that measurements of myocardial $T_1$, from this region might be rendered inaccurate by inclusion of intracavitary signal (from blood or thrombus) or extramyocardial signal (from pericardial fat or connective tissue) the region of interest was modified where necessary to encompass the closest lateral position which lay clearly within the myocardium. Attempts at obtaining a $T_1$ value from the narrowed segment illustrated the necessity for this modification since minor inward or outward adjustment of position resulted in major alterations in $T_1$.

The maximum $T_1$ value for each patient investigation was noted as was the number of areas with a $T_1$ value > 390 ms, the value previously identified as the cut off between normal and infarcted tissue. Values for non-affected myocardium were then obtained as follows. On the first image showing an increased $T_1$ value (> 390 ms) we selected the $T_1$ value from an area anatomically remote from the site of infarction to control for and give information about the variability of $T_1$ measurements with our system. $T_1$ values for this selected non-affected region were noted for subsequent examinations.

The data were analysed in both a cross sectional and longitudinal manner. In the cross sectional analysis the maximum $T_1$, for all patients at each time period after infarction is presented as a mean and 1 SD. The variation in $T_1$ with time since infarction was studied by fitting a gamma variate curve to the mean maximum $T_1$ values at each time period and calculating the regression coefficient. The maximum $T_1$ at day 1–3 was compared with the value obtained from the non-affected region on the same scan by means of a paired Student’s $t$ test. Linear regression analysis was used to correlate the maximum $T_1$ values and the number of areas of increased $T_1$ with left ventricular ejection fraction and peak creatine kinase activity.

Longitudinal analysis of the data was performed by noting the change in $T_1$ at each study interval.

**Results**

**$T_1$ Values in the Non-AFFECTED Area**

For the group as a whole the mean $T_1$ of the non-affected myocardium at day 1–3 was 320 (19) ms. There was no significant difference in the measured $T_1$ at any time in the non-affected region, the highest mean value being 326 (18) ms at one month. The range of $T_1$ values in the 41 patients over all these periods was 274–382 ms, confirming our previous observation that a cut off of 390 ms differentiates between normal and infarcted myocardium.

**$T_1$ Values in the AFFECTED Area**

**Maximum $T_1$ values**

Table 2 shows the results for each patient at the six times after infarction. Maximum $T_1$ values always occurred in an anatomical position consistent with the electrocardiographic changes (fig 1a and b). But the highest $T_1$ was not always seen in the same region on serial imaging.

Cross sectional analysis of the data (fig 2) showed that at day 1–3 $T_1$ was significantly higher in the infarct region than in the non-affected area.
Table 2  Patient details, maximum creatine kinase (CK) and T1 values at intervals shown

<table>
<thead>
<tr>
<th>Patient No</th>
<th>Age</th>
<th>Site of infarct</th>
<th>Successful thrombolysis</th>
<th>Max CK</th>
<th>T1 maximum</th>
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<td></td>
<td></td>
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<td>480</td>
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<tr>
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<td>1261</td>
<td>439</td>
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<tr>
<td>3</td>
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<td>1395</td>
<td>415</td>
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<td>371</td>
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<td>371</td>
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<td>371</td>
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</table>

The number of areas with significantly increased T1 varied greatly both between patients and at different times. The same general pattern to that of the peak T1 values was found. By six months only two patients had any areas of T1 > 390 ms.

**Comparison of reperfused and non-reperfused groups**

The maximum T1 and number of areas of increased T1 (above 390 ms) were not significantly different between the two groups at any time period. Neither was there any significant difference in T1 values between those patients in the reperfused group who were treated early (within two hours) and those who were treated later, although there was a trend towards a more rapid fall in the number of areas of raised T1 in the group treated early.

**Identification of infarct site**

For inferior infarcts the maximum T1 was most commonly seen on the coronal image (fig 1a) but...
**Magnetic resonance T₁ changes after myocardial infarction**

Fig 2  Mean maximum T₁ values (2 SE) after myocardial infarction. Values were greater than the normal mean maximum T₁ (SE) at first imaging (day 1-3) and tended to increase until the third imaging sequence (day 8-14); values only returned to the mean maximum T₁ values seen in normal subjects six months after infarction.

maximum T₁ values were more commonly seen on the transverse views when there was an anterior infarct (fig 1b).

**CORRELATION OF T₁ WITH OTHER MEASURED VARIABLES**

**Creatine kinase**

Because inferior infarcts tended to have a smaller peak activity of creatine kinase (1471 (581) vs 2162 (428) U/l) and left ventricular function was better preserved (52 (7)% vs 32 (11)%), they were first considered separately from anterior infarcts. There was no significant difference in maximum T₁ values between anterior and inferior infarction, however. There was a correlation (r = 0.59, p < 0.01) between the peak activity of creatine kinase and the maximum T₁ found at three months in the combined groups.

**Left ventricular ejection fraction**

There was a correlation between ejection fraction and peak T₁ values at one month (r = -0.57, p < 0.05). The number of areas of increased T₁ (>390 ms) correlated with ejection fraction at 4-7 days (r = -0.63, p < 0.01), 8-14 days (r = -0.53, p < 0.05), and at one month (r = -0.6, p < 0.02).

**Discussion**

After myocardial infarction T₁ rises in the area of infarction in almost all patients. There is no significant change in T₁ in the unaffected myocardium. This confirms reports of in vitro ²³ and in vivo ⁵⁶ experimental studies and those in human infarction. ⁷⁸ The changes may not appear or be appreciable before four days after the onset of infarction, and they reach a peak by the end of the first week. While it is likely that there was some increase in T₁ during the early stages of infarction in all our patients, the magnitude of this increase was not sufficient to permit reliable identification of abnormal myocardium in every case. The values in the five patients who did not have a T₁ > 390 ms on their initial scan ranged from 366 to 378 ms compared with a mean 320 (9) ms in the normal myocardium and they were within our previously reported range for normal myocardium.

Our finding that maximum T₁ values occurred at about two weeks was unexpected. The reported close correlation between T₁ and water content ² suggests that a more uniform and rapid increase in T₁ would be expected, because myocardial interstitial oedema and mitochondrial swelling occur early after the onset of infarction. ¹³ The dissociation between the timing of the peak T₁ and the expected peak of myocardial oedema formation may partly be explained by the fact that T₁ depends not simply on the total myocardial water content but also on the binding or interaction of water with biological macromolecules. ¹⁴

The difference between our findings and previous in vitro studies of early changes in T₁ is likely to reflect the major differences between ischaemic human myocardium and excised heart muscle. Not only will the quantity of blood in the intramyocardial vessels differ, as does the coronary anatomy, but in addition there are differences in the state of binding of water within cells. In living systems metabolic processes continuously produce water and its biological state is regulated by cellular processes. That such differences affect T₁ has been demonstrated by comparative in vivo and in vitro T₁ measurements in rabbits. ¹⁴

Our results accord with the in vivo experimental work of Pflugfelder et al who studied the serial changes in relaxation parameters and signal intensity in eight dogs after myocardial infarction. ⁹ In one animal the increase in signal intensity was only seen 4–6 days after infarction. The timing of peak signal intensity was dependent on the pulse sequence used but occurred between the fourth and thirteenth day. Furthermore, these workers frequently found an area of increased signal intensity on day 20 (the final image) and they commented that the affected area seemed to get smaller. Similarly in our study T₁ values remained raised at one month and even in some patients at three and occasionally six months, but the number of areas of increased T₁ fell more rapidly.
The presence of $T_1$ prolongation for up to three months confirms that factors other than myocardial oedema are responsible. The classic histological findings after myocardial infarction indicate that neutrophils appear after about six hours and increase rapidly after 24 hours, reaching a peak between the third and fifth day. Thereafter the number of large mononuclear macrophages increases, reaching a maximum at six weeks and subsequently declining. Other cellular infiltrates also occur, as does proliferation of capillaries which reaches a peak between 3-6 weeks. The histological changes may vary in the same patient at various times. These cellular changes may influence magnetic resonance variables but, as yet, precise histological correlations are not possible. The possibility that $T_1$ and $T_2$ changes may provide a method of obtaining in vivo information about pathogenesis is exciting and has received some support from the recent finding that $T_2$ correlates with the histological severity of rejection in

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Fig 3  Sequential coronal images in a patient with inferior infarction: (a) 3 days, (b) 14 days, (c) 1 month, and (d) 3 months after infarction.
heterotopic cardiac transplantation in dogs.\textsuperscript{18} Some studies have suggested that \( T_2 \) is better than \( T_1 \) for the identification of infarcted tissue, but this may reflect different imaging sequences where the pulse sequence combination used was more appropriate for \( T_2 \) measurements than for \( T_1 \). Spin echo sequences with a short time to recovery do not contain as much \( T_1 \) weighting as an inversion-recovery sequence. In addition, variation in a patient's RR interval can influence time to recovery in the spin echo sequence causing a variable \( T_1 \) weighting and, as a consequence, error in the \( T_1 \) value. The constant time for inversion used in our technique, coupled with a non-slice selective efficient inverting pulse, minimises the error in \( T_1 \) caused by variation in the RR interval.\textsuperscript{12}

Ratner \textit{et al} measured endocardial and epicardial \( T_1 \) and \( T_2 \) and correlates these variables with perfusion measured after 30 and 60 minutes of coronary occlusion.\textsuperscript{19} They suggested that the magnitude of change in \( T_1 \) and \( T_2 \) correlates with the extent of the ischaemic insult. The magnitude of the ischaemic insult is dependent not only on the duration of vessel occlusion but also on vessel size, collateral flow, and the rate of oxygen consumption which all modify the response and determine the infarct size. In this clinical study our means of assessing infarct size were limited. We used the peak creatine kinase activity as a crude assessment of infarct size. There was a correlation with maximum \( T_1 \), but only at one month. Thus the maximum level of \( T_1 \) seems to correlate poorly with infarct size in clinical practice. Because the kinetics of creatine kinase release are altered by reperfusion we also measured left ventricular ejection fraction at 8-10 days. All patients included in this analysis presented with a first infarct and so the reduction in left ventricular function provides some information about infarct size. The number of areas of increased \( T_1 \) did correlate with the ejection fraction at 4-7 days, 8-14 days, and at one month. No correlation was expected at day 1-3 or at three and six months because some patients did not develop changes until after the third day and by three months the area of increased \( T_1 \) was considerably reduced. This correlation presumably reflects the amount of myocardium affected by the ischaemic event. Thus the extent of the area with abnormal \( T_1 \) values seems a better guide to infarct size than the magnitude of changes in \( T_1 \). When this study was undertaken we were unable to perform multiple slice acquisitions but with this capability a precise determination of the volume of infarcted tissue should be possible.

Studies of changes in \( T_1 \) in dogs after coronary occlusion and reperfusion have suggested that \( T_1 \) is further prolonged by reperfusion,\textsuperscript{20} and an increase in cellular swelling after reperfusion has been reported.\textsuperscript{13} None the less, we found no significant difference in maximum \( T_1 \) between our reperfused and non-reperfused patients. This may reflect a balance between beneficial and harmful\textsuperscript{21} effects of reperfusion; however, further analysis of our data in which those given early and late thrombolytic treatment were compared gives no evidence of this. Because time to reperfusion is critically important\textsuperscript{22} to the effect of reperfusion, it might be expected that the \( T_1 \) values would be different in these groups. There was no significant difference in maximum \( T_1 \) or the number of areas showing raised \( T_1 \) between those treated within two hours and those treated more than two hours after the onset of symptoms. We noted a trend towards a more rapid fall in the number of areas with a raised \( T_1 \) in those treated early. The small numbers in these subgroups and the heterogeneity of the patients (in terms of the vessel affected, the position of the lesion, rate of reperfusion, extent of residual stenosis, collateral flow, disease of other vessels and therefore of infarct size and degree of continuing ischaemia) may be masking real differences.

The factors which produce these alterations in \( T_1 \) are complex and as yet we do not know whether changes in \( T_1 \) values reflect specific histological
findings. If the demonstrated changes reflect healing then alterations in $T_1$ may provide an important method for studying a neglected area in our understanding of myocardial infarction.

This work was supported by the Medical Research Council, The Wellcome Trust, and the Scottish Home & Health Department. MB was a Squibb cardiovascular research fellow.

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1 Mather-De Vre R. Biomedical implications of the relaxation behaviour of water related to NMR imaging. Br J Radiol 1984;57:935-76.
Myocardial involvement in systemic lupus erythematosus detected by magnetic resonance imaging


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KEY WORDS: Relaxation parameter T1, Lupus activity criteria count.

Myocardial involvement in systemic lupus erythematosus is commonly found at autopsy but seldom recognized clinically or by routine cardiological investigations. As the magnetic resonance relaxation parameter, T1, is altered by changes in tissue cellularity, we carried out magnetic resonance imaging in 10 patients with systemic lupus erythematosus. Five had active systemic lupus erythematosus when assessed using the lupus activity criteria count.

The mean (±SD) T1 was 319±12 in normal volunteers and 321±10 in a second control group with hypertrophic cardiomyopathy. In the group with systemic lupus erythematosus, there was a higher mean value of 336 ms with a wider scatter of individual results (SD±22 ms). In the subgroup of patients with active disease, T1 was significantly higher (349±24) than in either of the two control groups.

In addition, there was an inverse correlation between serum complement and myocardial T1 in patients with systemic lupus erythematosus. Myocardial abnormalities in systemic lupus erythematosus were demonstrated by magnetic resonance imaging even where other non-invasive cardiac investigations were negative. We conclude that T1 calculated from magnetic resonance imaging is often abnormal in systemic lupus erythematosus and probably indicates myocardial involvement.

Introduction

Myocardial abnormalities are frequently found in post-mortem studies of patients with systemic lupus erythematosus[1-3] and discovery of inflammatory infiltration of the myocardial interstitium led to the first description of systemic lupus erythematosus as a 'connective tissue disease'[4]. The diagnosis of myocarditis in vivo, however, has remained difficult to establish. The electrocardiogram and chest X-ray are often abnormal in systemic lupus erythematosus but do not clearly differentiate myocardial and pericardial disease[5,6]. Echocardiography may be helpful particularly in the assessment of myocardial function[7,8], but cannot differentiate ischaemic from non-ischaemic dysfunction and is of limited use in determining tissue changes. Thallium imaging reveals perfusion defects in patients with both active and inactive disease[9], but little is known about the underlying mechanism. The lack of an adequate method for non-invasive assessment of myocardial abnormalities has limited the understanding and recognition of the myocardial component of cardiac involvement in systemic lupus erythematosus.

Magnetic resonance imaging may allow myocardial tissue characterisation. Although we reported that the T1 was not altered in hypertrophic cardiomyopathy[10], these observations were made before we had developed a satisfactory gating technique for magnetic resonance imaging. Increases in the magnetic resonance relaxation parameters, T1 and T2, have been noted after experimental myocardial infarction and relate to myocardial tissue water content[11,12]. We demonstrated an increase in T1 in end-systolic images after myocardial infarction in man[13] but the time of maximum change was at about 14 days after infarction, suggesting that alterations in T1 may reflect cellular repair[14] rather than myocardial oedema alone. This concept is further supported by recent studies of the time course of magnetic-resonance parameter changes after coronary occlusion in both dogs[15] and rabbits[16].

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The purpose of the present study was to determine whether the tissue characterisation provided by magnetic resonance imaging could indicate the presence of myocardial involvement in patients with systemic lupus erythematosus and, thereby, provide an index of myocarditis.

Methods

SUBJECTS

Ten female patients with established systemic lupus erythematosus fulfilling the revised American Rheumatism Association criteria11 were studied. Clinical details of the patients are illustrated in Table 4. Nine patients were free of cardiac symptoms at the time of study, although one of these had a bioprosthetic mitral valve. The tenth patient was in congestive cardiac failure and suffered from recurrent supraventricular dysrhythmias.

Two control groups were also studied. The first was an age- and gender-matched group of eight normal volunteers drawn mainly from hospital staff. The second group consisted of 10 age-matched patients with hypertrophic cardiomyopathy, seven of whom were female.

All patients and volunteers gave written, informed consent prior to magnetic resonance imaging.

INVESTIGATIONS

A clinical history was taken from all patients with systemic lupus erythematosus and a physical examination was performed by one physician (BJT). Blood was taken for estimation of the full blood count, immuno-reactive complement proteins C3 and C4, antibody to double stranded DNA and circulating immune complexes. Disease activity was then assessed using the lupus activity criteria count, a validated index of disease activity which comprises seven primarily clinical variables10,19. The presence of two or more positive variables at the time of study indicates active disease. Each subject had a 12-lead electrocardiogram, M-mode and cross-sectional echocardiography on the same day as magnetic resonance imaging. The results of the cardiac imaging were assessed by a second clinician (MB) who was unaware of the clinical assessment.

MAGNETIC RESONANCE IMAGING

Subjects were imaged in a low field (0.08 Tesla) resistive imaging system (M&D Technology). An interleaved saturation recovery and inversion recovery pulse sequence was used to allow accurate measurement of the magnetic relaxation parameter T1. Electrocardiographically triggered cardiac gating was used as described previously20. Images were obtained at end-systole with acquisition of each image taking about 5 min. Four contiguous transverse slices, each of 16 mm thickness were obtained, encompassing the left ventricle. A single coronal scan was also obtained at a level that bisected the left ventricle.

The T1 maps which make up the images were

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age (years)</th>
<th>Duration of SLE (years)</th>
<th>Clinical manifestations</th>
<th>Blood pressure (mmHg)</th>
<th>Treatment (mg day⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LC</td>
<td>21</td>
<td>4</td>
<td>Alopecia</td>
<td>140/86</td>
<td>Pred. 12.5 Aza.</td>
</tr>
<tr>
<td>MJ</td>
<td>37</td>
<td>2</td>
<td>Arthritis</td>
<td>154/80</td>
<td>25 150</td>
</tr>
<tr>
<td>EG</td>
<td>48</td>
<td>2</td>
<td>Alopecia</td>
<td>140/88</td>
<td>15</td>
</tr>
<tr>
<td>JB</td>
<td>38</td>
<td>6</td>
<td>Arthritis</td>
<td>160/96</td>
<td>15</td>
</tr>
<tr>
<td>JW</td>
<td>57</td>
<td>14</td>
<td>Arthritis</td>
<td>138/80</td>
<td>2.5</td>
</tr>
<tr>
<td>CMC&amp;D</td>
<td>68</td>
<td>20</td>
<td>—</td>
<td>175/102</td>
<td>10</td>
</tr>
<tr>
<td>GD</td>
<td>33</td>
<td>4</td>
<td>—</td>
<td>150/80</td>
<td>—</td>
</tr>
<tr>
<td>JM</td>
<td>51</td>
<td>10</td>
<td>—</td>
<td>160/80</td>
<td>12.5</td>
</tr>
<tr>
<td>AR</td>
<td>31</td>
<td>7</td>
<td>Cardiac failure</td>
<td>150/96</td>
<td>60 50</td>
</tr>
<tr>
<td>AL</td>
<td>26</td>
<td>4</td>
<td>—</td>
<td>140/90</td>
<td>—</td>
</tr>
</tbody>
</table>

Pred., prednisolone; Aza., azathioprine; SLE, systemic lupus erythematosus
then analysed by dividing each image of the left ventricular myocardium into six regions of interest to test for homogeneity of signal. From the T1 values obtained, a mean T1 of all regions of interest for each subject was obtained and the maximum T1 in any section noted.

**ECHOCARDIOGRAPHY**

M-mode and cross-sectional echocardiography were performed using a Hewlett-Packard phased array system with a 3-5-MHz transducer. Left ventricular function was assessed from cross-sectional images and measurement of myocardial thickness and cavity size made from M-mode tracings.

**STATISTICAL ANALYSIS**

In addition to analysis between three clinical groups, the group with systemic lupus erythematosus was further subdivided into those with and without active disease based on the disease activity.

**Table 2**  
ECG and echocardiographic findings in patients with systemic lupus erythematosus

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age (years)</th>
<th>ECG</th>
<th>SEPTUM</th>
<th>LV FREE WALL</th>
<th>EDD</th>
<th>ESD</th>
</tr>
</thead>
<tbody>
<tr>
<td>LC</td>
<td>21</td>
<td>Normal</td>
<td>11</td>
<td>14</td>
<td>50</td>
<td>36</td>
</tr>
<tr>
<td>MJ</td>
<td>37</td>
<td>Normal</td>
<td>13</td>
<td>14</td>
<td>33</td>
<td>20</td>
</tr>
<tr>
<td>EG</td>
<td>48</td>
<td>Normal</td>
<td>11</td>
<td>12</td>
<td>56</td>
<td>30</td>
</tr>
<tr>
<td>JB</td>
<td>38</td>
<td>LBBB</td>
<td>12</td>
<td>12</td>
<td>64</td>
<td>56</td>
</tr>
<tr>
<td>JW</td>
<td>57</td>
<td>Normal</td>
<td>16</td>
<td>16</td>
<td>50</td>
<td>35</td>
</tr>
<tr>
<td>CMcD</td>
<td>68</td>
<td>Non-specific</td>
<td>13</td>
<td>20</td>
<td>44</td>
<td>24</td>
</tr>
<tr>
<td>GD</td>
<td>33</td>
<td>Normal</td>
<td>12</td>
<td>14</td>
<td>50</td>
<td>36</td>
</tr>
<tr>
<td>JM</td>
<td>51</td>
<td>Left axis deviation</td>
<td>13</td>
<td>16</td>
<td>63</td>
<td>45</td>
</tr>
<tr>
<td>AR</td>
<td>38</td>
<td>AF, widespread T-wave changes</td>
<td>12</td>
<td>11</td>
<td>64</td>
<td>48</td>
</tr>
<tr>
<td>AL</td>
<td>26</td>
<td>Normal</td>
<td>11</td>
<td>11</td>
<td>52</td>
<td>33</td>
</tr>
</tbody>
</table>

SLE, systemic lupus erythematosus; LBBB, left bundle branch block; AF, atrial fibrillation.

**Table 3**  
Magnetic resonance T1 values in volunteers, patients with hypertrophic cardiomyopathy and SLE (complement values are shown for patients with SLE)

<table>
<thead>
<tr>
<th>Mean T1 (ms)</th>
<th>Complement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vol.</td>
<td>HCM</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>321</td>
<td>311</td>
</tr>
<tr>
<td>318</td>
<td>320</td>
</tr>
<tr>
<td>334</td>
<td>333</td>
</tr>
<tr>
<td>322</td>
<td>324</td>
</tr>
<tr>
<td>333</td>
<td>333</td>
</tr>
<tr>
<td>297</td>
<td>320</td>
</tr>
<tr>
<td>322</td>
<td>321</td>
</tr>
<tr>
<td>314</td>
<td>318</td>
</tr>
<tr>
<td>314</td>
<td>305</td>
</tr>
</tbody>
</table>

Vol., volunteers; HCM, hypertrophic cardiomyopathy; SLE, systemic lupus erythematosus; adsDNA, antibody to double stranded DNA; LACC, lupus activity criteria count.

*Active disease.
Myocardial involvement in systemic lupus erythematosus

Hypertrophic cardiomyopathy

Mean T1 values in normal volunteers, patients with hypertrophic cardiomyopathy and patients with systemic lupus erythematosus. O, Active disease.

Analysis of variance was used to establish whether there was a significant difference between the means of the initial three groups and also between the means of the further subdivision into four groups. Subsequent analysis was performed using a non-paired t-test to decide which group means differed significantly.

Correlation between T1 values and complement levels was by linear regression analysis. Significance was accepted at $P < 0.05$.

Results

CLINICAL FINDINGS

The clinical findings in the group with systemic lupus erythematosus are summarised in Table 1. Active disease, as indicated by a lupus activity criteria count of 2 or more, was present in five of the 10 patients. Eight patients were on prednisolone and two on azathiaprine. Clinical evidence of cardiac disease was only detectable in the one patient with heart failure.

ELECTROCARDIOGRAPHIC AND ECHOCARDIOGRAPHIC FINDINGS

Four of the group with systemic lupus erythematosus had non-specific electrocardiographic abnormalities (Table 2). The presence of an abnormal electrocardiogram did not relate to disease activity. On echocardiography, mild hypertrophy of the interventricular septum and left ventricular free wall was common (Table 2). Poor left ventricular function was confirmed in the patient with cardiac failure and a dilated left ventricle with moderate contraction was noted in the patient with a mitral-valve replacement.

All patients with hypertrophic cardiomyopathy had evidence of septal hypertrophy, usually with some free-wall hypertrophy.

PERICARDIAL INVOLVEMENT

A small amount of pericardial fluid was noted on the magnetic resonance image in two patients with systemic lupus erythematosus, but was also seen in two normal volunteers and in one patient with hypertrophic cardiomyopathy. The pericardial fluid was not detectable by echocardiography in the normal volunteers or patients with systemic lupus erythematosus but was noted in the patient with hypertrophic cardiomyopathy (in whom the effusion was larger).
Figure 3. Transverse sections illustrating T1 values in (a) a normal subject and (b) a patient with active lupus erythematosus. The colour bar shows the T1 values with the normal subject having T1 values of around 300 ms, but the patient with lupus having a diffuse increase in T1 of around 360 ms.

MAGNETIC RESONANCE IMAGING

The individual mean left ventricular myocardial T1 for each group is shown in Table 3. The mean T1 (1 standard deviation) for normal volunteers was 319 ± 12 ms, similar to the group with hypertrophic cardiomyopathy (321 ± 10 ms). For the group with systemic lupus erythematosus the mean T1 was higher at 336 ± 22 ms, this difference being significant ($P < 0.05$) when compared to the group with hypertrophic cardiomyopathy ($N = 10$), but not when compared to the normal volunteers ($N = 8$). However, there was marked variability in T1 values within the systemic lupus erythematosus group (Fig. 1). Values were particularly elevated in those with active disease as indicated by the lupus activity criteria count. For this subgroup the mean T1 was 349 ± 24 as compared to 324 ± 12 in the subgroup with inactive disease (Table 3). Analysis of variance demonstrated a significant difference between the means at the 0.99 level.
However, the increases in systemic lupus erythematosus (SLE) wall thickness were prolonged myocardial involvement. Inactive patients with systemic lupus erythematosus and raised T1 values show a more diffuse pattern of T1 elevation, with values usually not exceeding 400 ms (Fig. 3).

No abnormalities were seen in the electrocardiogram or echocardiographs of the normal volunteers.

**Discussion**

This study confirms our previous findings that there is little variation in T1 in normal subjects and the T1 is not abnormal in patients with hypertrophic cardiomyopathy. Furthermore, it demonstrates that the magnetic relaxation parameter is frequently, but not invariably, raised in patients with active systemic lupus erythematosus. The patients with systemic lupus erythematosus were recruited from those attending an outpatient clinic. They thus tended to have severe disease, reflected by the fact that eight were on steroid therapy (Table 1). However, of the five with active disease at the time of the study, only two had clinical, radiological, electrocardiographic or echocardiographic evidence suggestive of myocardial involvement.

Badui et al. found evidence of myocarditis in only 14% of cases using routine non-invasive investigations. Despite the selective nature of the patient group, the fact that four out of the five with active disease had a raised T1 suggests that abnormalities in the myocardium in systemic lupus erythematosus are common. It is likely that commonly available cardiac investigations are insensitive to the detection of such myocardial involvement. In two patients with markedly prolonged myocardial T1 values the electrocardiogram was entirely normal.

The possibility that the elevated T1 values might be related to the slight increase in left ventricular wall thickness found in the group with systemic lupus erythematosus is discounted by the normal values in the patients with hypertrophic cardiomyopathy who had marked myocardial hypertrophy.

Increases in myocardial T1 are not specific to systemic lupus erythematosus. We have previously reported increases in T1 following myocardial infarction. However, the pattern of increase is very different in the two conditions. In myocardial infarction there is a localised area of increased T1, with values in the infarcted area rising above 390 ms. There is often associated wall thinning, while the unaffected myocardium has a normal appearance. In contrast, those patients with systemic lupus erythematosus and raised T1 values show a more diffuse pattern of T1 elevation, with values usually not exceeding 400 ms (Fig. 3).

After acute myocardial infarction the time course of the changes is for a localised abnormal area to increase in size up to about 14 days after infarction. There is a gradual return to normal values by six months after the acute event. We have not yet been able to make serial measurements in the patients with lupus erythematosus.

The underlying mechanism for this increase in T1 is not yet clear. Histological findings reported in the myocardium of patients dying of systemic lupus erythematosus include lymphocytic infiltration, sometimes accompanied by myocardial oedema. In addition to the changes following acute myocardial infarction, T1 also rises in acute rejection following heterotopic cardiac transplantation.

In both these conditions, as in systemic lupus erythematosus, cellular infiltration is a prominent histological feature and may directly or indirectly be responsible for the increase in T1.

Although prolonged myocardial T1 values appear to reflect overall disease activity, one patient with active disease had a normal T1. Although we have no histological proof, increases in T1 may specifically reflect myocardial involvement. It is notable that the one patient with acute cardiac failure had the highest T1.

The likelihood is that increases in T1 documented by this interleaved saturation recovery and inversion recovery pulse sequence reflect altered cellularity of the tissue being examined. The relaxation parameters T1 and T2 depend on the precise pulse sequence used in magnetic resonance imaging. The pulse sequence used in our system is the direct spin-echo pulse sequence, so some caution has to be taken before applying these results to measurements from other magnetic resonance imaging systems. However, estimates of T1 values from magnetic resonance imaging using inversion recovery sequences may aid in the investigation of other myocardial diseases such as myocarditis, congestive cardiomyopathy, connective tissue diseases other than systemic lupus erythematosus and cardiac rejection after transplantation where myocardial inflammation is suspected.
References


Brain hydration during alcohol withdrawal in alcoholics measured by magnetic resonance imaging


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(Received October 28th 1987)

Twenty-seven patients had a first Magnetic Resonance Imaging (MRI) scan 1–3 days after stopping drinking and a second approximately 2 weeks later with no change in whole brain $T_1$ or $T_2$ in selected brain areas. Six patients whose first scan was over 36 h after the last drink underwent an increase in whole brain $T_1$ in the interval to the second scan. The later the first scan was performed the greater was the increase in $T_1$. These results are compatible with a very early fall in brain water immediately on cessation of drinking (perhaps due to a rebound increase of vasopressin activity) with a return to 'baseline' after two weeks. A third scan after discharge from hospital in 23 individuals who had abstained from alcohol or drank very little did not reveal any further significant change in brain $T_1$.

Key words: brain hydration; alcohol withdrawal; magnetic resonance imaging

Introduction

Water retention within the brain has been postulated as a contributant to some of the clinical features of alcohol withdrawal in chronic alcoholics [1]. In magnetic resonance imaging (MRI), $T_1$ relaxation time is related to the state of water in the tissue [2]. The $T_1$ value, measured in milliseconds (ms) increases as the proportion of free to bound water within the tissue increases. In vivo $T_1$ values have been shown to correlate with the level of brain water [3–5] and so may enable a study of brain water during withdrawal of alcohol in chronic alcoholics.

MRI was first used for the investigation of chronic alcoholics by Besson and colleagues [6]. These authors suggested that brain water is diminished during intoxication and increased during alcohol withdrawal. Acute intoxication in occasional drinkers results in a fall in whole brain $T_1$, compatible with the dehydration which follows acute intake of alcohol [7]. Eisenhoffer and colleagues [1] suggest that such an effect, if chronic, might result on alcohol withdrawal in a rebound increase of vasopressin activity, followed by a return to baseline.

It was therefore hypothesised that whole brain $T_1$ in alcoholics would be seen to fall from the level at 24–48 h following cessation of drinking to a lower level ('baseline') some 2 weeks later. We were unable to perform $T_1$ measurements in the present study in intoxicated alcoholics immediately prior to admission and thus unable to study the early part of the postulated pattern of changes (that is, the putative rise in brain water and $T_1$).

Patients and Methods

Chronic alcoholic patients who were about to be voluntarily admitted as inpatients for detoxification to the Alcohol Problems Clinic, Royal Edinburgh Hospital, were fully informed about the study and invited to participate.
Twenty-seven patients (18 men, 9 women), were admitted to this study. Their mean age was 44 years (range 26–63, S.E.M. 5.2). They were to be scanned within 48 h of admission (day 1) and again approximately on day 15. However, the timing of scans varied according to the availability of the scanner and the requirements of the patients' participation in the ward programme.

In connection with a separate study [8] of cognitive impairment and MRI parameters, results of a scan performed at approximately day 15 were available in a further 42 patients. These patients had not been scanned at or near admission.

Patients had been drinking at least 130 g ethanol/day before admission and had a 5–15 year history of alcoholism. No patients were included if they had a history of drug abuse, or were clinically malnourished, or had physical disorders other than raised serum liver enzymes. Over the first 5 days each patient received a normal diet, intramuscular multivitamin supplements and decreasing doses of chlordiazepoxide to control withdrawal symptoms.

Our procedures for measuring whole brain $T_1$ and $T_2$ in regions of interest in the brain have been described [9]. The inter-operator reliability of our $T_1$ measurements has been reported [8]. Differences in $T_1$ values between groups of patients at different times were

![Graph](image-url)

**Fig. 1.** Whole Brain $T_1$ measurements with means according to length of time since last drink.
Table I. Mean change in $T_1$ from Scan 1 to Scan 2 ($\Delta = T_1$ (Scan 2) - $T_1$ (Scan 1)).

<table>
<thead>
<tr>
<th>All subjects</th>
<th>Scan 1 within 36 h of last drink</th>
<th>Scan 1 36–84 h after last drink</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta$ Frontal White</td>
<td>$1.83 \pm 9.46$</td>
<td>$-0.39 \pm 2.04$</td>
</tr>
<tr>
<td>$\Delta$ Frontal Grey</td>
<td>$0.50 \pm 17.51$</td>
<td>$-1.833 \pm 4.26$</td>
</tr>
<tr>
<td>$\Delta$ Parietal White</td>
<td>$0.83 \pm 9.17$</td>
<td>$-0.30 \pm 2.12$</td>
</tr>
<tr>
<td>$\Delta$ Parietal Grey</td>
<td>$-6.75 \pm 32.21$</td>
<td>$-11.17 \pm 5.09$</td>
</tr>
<tr>
<td>$\Delta$ Whole brain</td>
<td>$0.75 \pm 8.89$</td>
<td>$-1.78 \pm 2.01$</td>
</tr>
</tbody>
</table>

n: 24 18 6

*Significantly greater ($P = 0.012$) than in those whose first scan was within 36 h of last drink.

Results

Three patients were unavailable for their second scan. The mean whole brain $T_1$ at the first scan in the 24 patients who had a second scan was 314 ms (range 283–398, S.E. 2.71). At the second scan the mean $T_1$ was also 314 ms (range 283–334, S.E. 2.69). $T_1$ measurements in small areas of interest likewise showed no significant change between scans. Figure 1 illustrates all our available data on whole brain $T_1$ in the withdrawal period. It includes the subjects on whom only the first scan was done, and subjects from our other study scanned only at day 15 approximately. No trend suggestive of a fall in whole brain $T_1$ is visible.

Lest the hypothesised fall in $T_1$ had been early and had been followed by an early 'return to baseline', an analysis of the change in $T_1$ between the first and second scan was made, comparing those patients who had their first scan within 36 h of the last drink to those whose first scan was later. The 18 patients scanned within 36 h had their second scan at a mean of 13.6 days later (range 9–16 days, S.E. 0.43) and the six whose first scan was up to 48 h later had their second scan at a mean interval of 12.3 days since the first (range 9–16, S.E. 1.15). This interval was not significantly different ($P = 0.33$).

In those whose first scan had been within 36 h of the last drink, the difference between whole brain $T_1$ on the first and second scans was insignificant (mean difference $-1.78$ ms — a fall — range $-15 - +9$, S.E. 2.01). A significant increase occurred, however, in those whose first scan had been up to 48 h later (mean $8.33$ ms, range $+2 - +14$, S.E. 2.04; t = 4.08, d.f. 5 P < 0.01). The difference between the changes in these two groups of patients, (between those scanned within 36 h of admission and those scanned later), is

Table II. Correlation of TIME OF SCAN (post admission) with CHANGE IN $T_1$ from Scan 1 to Scan 2.

<table>
<thead>
<tr>
<th>TIME OF</th>
<th>$\Delta$ WB</th>
<th>$\Delta$ FW</th>
<th>$\Delta$ PW</th>
<th>$\Delta$ FG</th>
<th>$\Delta$ PG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scan 1</td>
<td>0.4048*</td>
<td>0.2434*</td>
<td>0.1080*</td>
<td>0.4076*</td>
<td>0.4242*</td>
</tr>
</tbody>
</table>

R-values (* $P < 0.01$) n = 24
significant ($t = 2.73$, d.f. 22, $P = 0.012$) (Table I). The trend was visible but not significant in each of the $T_1$ measurements made in areas of interest. The later the first scan was performed after the last drink, the greater the change between the first and second scans, as shown by the correlation coefficients given in Table II.

In case our timing of the scans had resulted in our missing any later changes in $T_1$, another comparison was made. Four patients had the second scan on the 9th or 10th day and 20 patients beyond the 10th day. There was no significant difference when these groups were compared in the mean change between the first and second scans (2.25 ms, S.E. 5.28, range $-11 - +13$ and 0.45 ms, range $-15 - +14$, S.E. 1.97, respectively; $t = 0.36$, d.f. 22, $P = 0.72$).

We had some further data after patients had been discharged from the 2-week ward programme. Seventeen patients from the present study had a further scan 2 months after discharge. There was also a scan at that point in 38 patients in the collateral study already mentioned. The difficulty in using this data is that we were not always certain whether or not an individual had continued to abstain from alcohol during that period. We examined the $T_1$ values in 23 individuals who had been scanned at about day 15 and had a repeat scan at a mean of 102 days and who as far as we could tell had abstained during that period or drunk very little. No significant change in $T_1$ in any of the regions of interest or in the whole brain was noted. However, the trend in these apparent abstainers in all our $T_1$ measures was for a slight increase, but with considerable individual variation (for example, the mean increase in whole brain $T_1$ was 2.39 ms, range $-11 - +19$, S.E. 1.55 ms, $n = 23$).

**Discussion and Conclusions**

The only possible way in which our data would support the notion that there is a fall in brain water following withdrawal from alcohol is using the sparse data comparing the changes in those scanned within 36 h of the last drink with those whose first scan was later. Our data would support a fall in $T_1$ between 36 h and 84 h, with a subsequent return to 'baseline' during the next 10 days approximately. It would have to be postulated, however, that individual differences in 'baseline' $T_1$ prevented this from emerging in the data on the whole sample as illustrated in Fig. 1.

Our data do not uphold the provisional result which we obtained on our first 9 patients [9]. In that paper, we reported a fall in whole brain $T_1$ which correlated positively with the length of abstinence prior to the second scan. In our present sample, the correlation between change in whole brain $T_1$ and interval to the second scan was insignificant ($-0.09$). Neither were there significant correlations between the change in $T_1$ in the areas of interest and interval to the second scan.

Further studies should comprise daily measurements beginning on the day prior to admission, when the patient is still drinking, and continuing for 2 weeks.

**Acknowledgements**

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**References**

Magnetic resonance spectroscopy in vivo

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ABSTRACT

Nuclear magnetic resonance (NMR) has become an important non-invasive investigative technique in medicine and biology. The most recent development has been the ability to perform magnetic resonance spectroscopy (MRS) in selected regions within the human body. Such volumes can be selected by techniques which fall into the following broad categories: surface coil methods, surface coils with depth selection, volume selection and chemical shift mapping. The latter two methods use magnetic field gradients, present in magnetic resonance imaging systems, to select the volume. MRS can be used to measure phosphorus and proton metabolites and hence study tissue biochemistry in-vivo.

Keywords: Medical imaging, magnetic resonance spectroscopy, biochemical analysis.

INTRODUCTION

There has been a great deal of interest in the application of nuclear magnetic resonance (NMR) in the field of medicine and biology over the last 15 years. The initial interest was fueled by the development of techniques of magnetic resonance imaging (MRI) so that now commercial MRI systems are present in many hospitals and research centres. More recently, developments in magnetic resonance spectroscopy (MRS) have provided a non-invasive technique for monitoring biochemical activity from selected regions within the body.

Following the first observations of NMR in 1946 by Bloch1 and Purcell2, it has long been possible to obtain in-vitro NMR spectra from small samples in liquid or solid form. In 1974 it was demonstrated that NMR can be used to study the major phosphorus containing metabolites in isolated intact tissues3. With the development of larger bore high field superconducting magnets came the possibility of acquiring NMR spectra from animal and human tissues in-vivo. Nuclei that can be studied by MRS include phosphorus, protons and carbon, all of which play an important role in human biochemistry.

One important requirement for in-vivo studies is spatial selection, so that spectra can be collected from a well defined region within the body without contamination by signals originating from tissues not under investigation. After a short account of the basic principles, this article reviews the techniques used to meet the in-vivo objectives and presents examples of human in-vivo NMR spectra.

PRINCIPLES OF MRS

In order to obtain NMR spectra, the sample to be studied is placed in a strong magnetic field of high homogeneity. The strength of this field is denoted by B0 and its direction is defined as the z axis. Interaction between the magnetic moment of the nuclei and the static field B0 results in the preferential alignment of the nuclear spins along the field direction. In this way, a nuclear magnetization is produced along the z axis. An individual nucleus, whose spin is inclined at an angle to B0, will precess about the field direction at a characteristic frequency:

$$\omega = \gamma B_0$$

where $\omega$ is known as the Larmor frequency and $\gamma$ is the gyromagnetic ratio for the particular nuclear species.

This behaviour is analogous to the precession of the axis of a gyroscope about the Earth's gravitational field. A radio frequency (RF) magnetic field at the same frequency can be employed to tilt the magnetization away from the z axis. This time varying field, denoted by $B_1$, is applied in a direction perpendicular to $B_0$. The magnitude and duration of this RF excitation pulse determines the angle of tilt, $\theta$, away from the z axis as follows:

$$\theta = \gamma \int B_1(t) \, dt$$

where $t_0$ is the pulse duration.

A 180° pulse leaves the nuclear magnetization along the $-z$ axis and a 90° pulse leaves it in the $xy$ plane. The continued precession of the nuclei about $B_0$ means that any $xy$ component of magnetization present after the RF pulse also precesses about $B_0$. It is this rotating magnetization that is detected by the induction of an electromotive force (EMF) in a radio frequency receiver coil. The decay of this signal is characterized by two time constants $T_1$ and $T_2$. $T_1$ is known as the longitudinal or spin–lattice relaxation time and describes the relaxation of the system to equilibrium with the nuclear magnetization along the z axis. $T_2$ is the transverse or spin–spin relaxation time and is a measure of the time during which the individual nuclear magnetic moments that contribute to the rotating $xy$ magnetization remain in phase with each other.
The magnetic field experienced by the nuclei is modified by aspects of its local environment such as the distribution of electrons. As a result, different compounds of the same nucleus will experience marginally different $B_0$ fields and consequently produce resonances at slightly different frequencies. This change in resonant frequency is known as a chemical shift. Formally, the chemical shift for a compound with resonant frequency, $\omega$, is given by:

$$\text{chemical shift (p.p.m.) } = \frac{\omega - \omega_0}{\omega_0} \times 10^6$$

$\omega_0$ is the resonant frequency of a reference compound which is standard for a given nuclear type. Providing the bandwidth of the RF excitation pulse is sufficient to encompass the entire range of chemical shift for a given nuclear type, the time signal produced by the rotating $\gamma$ magnetization contains information about the relative signal intensity as a function of chemical shift. This information is retrieved by Fourier transformation of the time signal and allows relative concentrations of different compounds of the same nucleus to be inferred. In order to resolve the different chemical shifts, a high degree of magnetic field homogeneity is required. If this condition is not met the resonant frequency of a nucleus with a given chemical shift will vary significantly across the sample and degrade resolution. Before data can be acquired in an NMR experiment the magnetic field $B_0$ is adjusted until the required homogeneity is produced. The field homogeneity can be monitored by measuring the line width of a single resonance. The resonance of proton nuclei in water molecules are frequently used for this purpose. This process of $B_0$ field adjustment is known as shimming. For imaging purposes a uniformity of 1 in 10⁶ is sufficient whereas for spectroscopy a uniformity of 1 in 10⁷ is required.

**SURFACE COIL TECHNIQUES**

In 1980 the first in-vivo NMR spectra were collected from phosphorus compounds in skeletal muscle and brain of intact rats, using a single turn circular RF coil placed close to the skin. The coil, which is tuned to the appropriate resonant frequency, is positioned so that the coil axis is perpendicular to the static magnetic field $B_0$. The coil is used both to send the RF excitation pulse and to receive the subsequent RF signal from the irradiated tissues. In conventional in-vivo NMR spectroscopy a saddle-shaped or solenoidal coil is used to provide a uniform $B_1$ field over the sample. Under these conditions, different regions within the sample contribute evenly to the detected signal. The inhomogeneous magnetic field associated with a surface coil is used to confine the NMR signal to a region close to the surface of the body. The relative signal strength as a function of spatial coordinates is determined by two factors. Firstly, as a result of the principle of reciprocity, the EMF induced in the coil by the rotating magnetization at a given point in the sample is proportional to the magnetic field produced at that same point by unit current flowing through the coil. Secondly, the pulse angle following an RF excitation pulse varies across the sample according to equation (1).

It therefore follows that the relative contribution to the detected signal from a given point in the sample, is proportional to $z \sin \varphi$, where $z$ is the pulse angle at that point. This behaviour can be used to a limited extent to select regions at different distances from the coil. Figure 1 shows the relative signal received from points in a homogeneous non-conducting sample as a function of distance along the coil axis when a 90° and 180° pulse are generated at the coil centre. Frequency dependent attenuation of the RF signals in conducting biological samples must be considered for a more careful analysis. The behaviour of the surface coil $B_1$ field away from the coil axis means that regions of high sensitivity selected by adjustment of pulse length have a complex shape. It is not possible to achieve efficient depth selection using simple radio frequency excitation pulses as a result of surface contamination. This arises due to the high $B_1$ flux near the coil wire, where a large range of pulse angles exist and where the inherent sensitivity is high.

An example of a phosphorus spectrum from a human calf muscle using a 6 cm diameter surface coil is shown in Figure 2.

**DEPTH SELECTION**

A number of techniques have been developed to improve the ability of surface coil methods to select regions of tissue at different distances from the coil.

One approach has been to make the signal received from a given point a stronger function of the pulse angle at that point. In this way the signal is more effectively localized to regions where the pulse angle is close to 90°. This has been achieved by using a series of multiple RF pulses to excite the nuclear magnetization away from its equilibrium direction along the $z$ axis. This contrasts with the simple NMR experiment, outlined above, where a single RF pulse is applied for this purpose.
A number of different RF pulse sequences have been used. An example of one of the less complex is a follows: an initial RF pulse of duration $t$ is followed by a second RF pulse of duration $2t$ before signal acquisition. Four separate signal acquisitions are performed in the same way, but the phase difference between the two RF pulses is increased by $90^\circ$ for each acquisition; $180^\circ$ phase adjustment of the receiver is also required for alternate acquisitions. Analysis of the motion of the nuclear magnetization during this sequence shows that summation of the four signals produces a signal whose magnitude is proportional to $x \sin x$. Figure 3 compares this dependence on pulse angle with that obtained from the use of a single RF pulse for excitation.

Other multiple pulse sequences produce signals that are proportional to higher order trigonometrical functions of the pulse length. This greater selectivity to pulse length is accompanied by greater complexity of the sequence and an increase in the number of signal acquisitions required. The problem of avoiding contamination by signals from regions close to the coil has also been addressed by multiple pulse techniques.

The introduction of magnetic field gradient techniques provided a more efficient method of selectively receiving a signal from tissue at a given distance below a surface coil. One technique which utilizes a magnetic field gradient in conjunction with a surface coil to produce spatial localization is DRESS (Depth Resolved Surface Coil Spectroscopy). Superposition of a linear magnetic field gradient upon the $B_0$ static magnetic field has the effect of making the resonant frequency of the nuclei a linear function of position along the field gradient. Hence, the application of a narrow-bandwidth RF excitation pulse in the presence of the magnetic field gradient only excites spatial regions of nuclei within the appropriate range of resonant frequency. In DRESS a magnetic field gradient is applied perpendicular to the surface coil plane. The use of a narrow-bandwidth RF excitation pulse, such as an amplitude-modulated sine function, then has the effect of confining the signal to a plane perpendicular to the field direction and parallel to the surface coil plane. Localization of the signal within the plane is determined by the geometry of the surface coil. The thickness of the selected plane is dependent upon the magnitude of the magnetic field gradient and the bandwidth of the RF pulse.

In this technique the effects of chemical shift must be examined. Consider two compounds of the same nucleus whose resonances are separated by $\Delta \omega$ in the field $B_0$. The range of frequencies contained within the narrow-bandwidth RF pulse corresponds to slightly different spatial regions for each of the two compounds. The spatial offset between the two regions is given by:

$$\Delta x = \frac{\Delta \omega}{\gamma G_x}$$

where $G_x$ is the field gradient in the $x$ direction. Clearly, the offset is minimized by using as large a field gradient as possible.

The DRESS technique makes one further modification to the surface coil method of MRS. Separate RF coils are used for excitation and receiving. The coil used for excitation is chosen to have a uniform $B_1$ field over the sensitive volume of the surface receiver coil. A larger surface coil can be used for the excitation. It is then possible to optimize the signal strength received from the selected plane by applying a $90^\circ$ pulse to the sample. Pulse length variation, and associated signal reduction, is inevitable if the smaller surface coil is used for the excitation. The Johnson noise received from the sample is determined by the receiver coil dimensions.
Hence, this coil arrangement detects no greater noise than when a single send and receive coil is used.

VOLUME SELECTION

The first technique of volume selection was christened topical magnetic resonance (TMR) and was used to collect the first human in-vivo NMR spectrum in 1981 from forearm muscle. Static high order magnetic field gradients are superimposed upon the magnetic field \( B_0 \) to produce high field homogeneity in one region only. The signal detected from this region contains highly resolved frequency information, characterized by narrow lines in the spectrum. The detector, which may be a surface coil, also receives signals from the inhomogeneous parts of the body elsewhere. This low resolution information can be filtered out from the signal mathematically to recover the signal originating from the region of high field homogeneity only. The complex nature of the spatial variation of magnetic field means that the volume selected in this manner is irregular and not well defined. The technique is further limited by the requirement that the homogeneous region can only be produced at the centre of the main magnetic field. This makes positioning of the localized volume a difficult task. This method of spatial localization has been superseded by others and is only of historical interest.

More recently, techniques have been developed that achieve localization of the detected signal to a well-defined volume, such as a cube, deep within the body. The position and size of the volume is very flexible and changes can be made without repositioning of the subject under investigation. Such methods employ pulsed field gradients that are used in MRI. As in the DRESS technique, described above, the combination of magnetic field gradients and narrow-bandwidth pulses allows the excitation of selected regions of nuclear spins. In DRESS a field gradient is applied in one dimension only, confining the signal to a plane. Field gradients applied in all three dimensions can be used to confine the signal further to a cube. NMR images of the body provide a means of identifying the region from which a spectrum is to be obtained. Then appropriate field gradients and RF pulses are used to acquire localized information from the desired location. The choice of position of the volume is, however, restricted by the sensitive volume of the receiver coil. A number of methods for achieving volume selection have been developed at MIT and SPARS.

ISIS (Image Selected In-vivo Spectroscopy) makes use of slice selection in three orthogonal planes. A slice selective 180° pulse applied to a sample leaves the nuclear magnetization with its direction along the \(-z\) axis in the selected slice and unperturbed in the \(+z\) axis equilibrium direction elsewhere. Application of a non-selective 90° pulse, or detection pulse, rotates the magnetization into the \(xy\) plane across the whole sample. The magnetization then rotates in the \(xy\) plane as described previously. However, the magnetization in the selected slice points in the opposite direction to that elsewhere in the sample. As a result there is a 180° phase difference between signals from the two regions so that signal from the selected slice contributes negatively to the total signal. A signal acquired from the same total volume, omitting the 180° slice selective pulse has positive signal contributions from the entire sample. Hence, subtraction of the signal acquired with the slice selection from that acquired without it results in the summation of signal from the selected slice and cancellation elsewhere.

In the ISIS technique this principle is extended to three dimensions to achieve cancellation in all regions except a selected cube. Three orthogonal field gradients are used to produce three selected slices whose intersection is the required cube. The three 180° slice selective pulses are applied successively before the detection pulse. The slice selective pulses can be applied in any combination, ranging from the omission of all three to the use of all three. When no slice selective pulses are applied during the preparation period, before the detection pulse, all regions within the sample contribute positively to the signal. If a single selective pulse is used, a single slice perpendicular to the appropriate field gradient contributes negative signal as above. When two or three slice selections are used, regions experiencing zero or two 180° pulses contribute positively to the signal and regions where one or three 180° pulses have been applied contribute negatively. Signals received using different permutations of the slice selective pulses can be arithmetically combined to cancel signal from everywhere except for the cube formed by the intersection of the three planes, where the signal is summed. Eight separate signal acquisitions are required to achieve this.

As with all techniques that incorporate selective excitation using magnetic field gradients, the spatial localization is a function of chemical shift as described by equation (2). In our case, a maximum magnetic field gradient of 3 mT/m means that \( \Delta x \) is about 1 cm for the range of phosphorus compounds at 1.5 T. Although selective 180° pulses give the optimum signal, the method of cancellation does not depend on the pulse angles used. As a result ISIS can be used in conjunction with RF coils, such as surface coils, with an inhomogeneous \( B_1 \) field.

SPARS (SPAtially Resolved Spectroscopy) uses a more complex sequence of magnetic field gradient and RF pulses to produce signals from the selected volume in each signal acquisition. In this case the method does rely on accurate pulse angles and a homogeneous \( B_1 \) field is required. Firstly, a non-selective 90° pulse is applied, rotating the magnetization into the \(xy\) plane. A combination of pulsed magnetic field gradients and a 180° RF pulse are then used to produce a phase coherence of the nuclear spins in the \(xy\) plane. It is then possible to apply a slice selective 90° pulse to rotate the spins within a given slice back along the \(z\) axis. The remaining spins in the \(xy\) plane become dephased as a result of the position dependent resonant frequency created by the magnetic field applied during the 90° selective pulse. The sequence is repeated and slice selections are performed in the other two dimensions. The
signals used to identify the acquisition of Figure 5. The proton coil is used to magnetize the phosphorus head coil of the human subject, positioned close to the head to optimize sensitivity. The subject is then placed inside the magnet with the head inside the proton imaging coil. Figure 4 shows the position of the two RF coils during the investigation. Firstly, an image is acquired, from which the position and size of the volume to be studied by MRS is determined (see Figure 6). The proton coil is then used to collect localized proton signals from the volume of interest using SPARS. The static magnetic field, $B_0$, is shimmed by varying the current through a series of resistive coils around the magnet bore until the required magnetic field homogeneity of 0.2 to 0.3 p.p.m. is obtained. ISIS is then used to collect a localized phosphorus spectrum; the phosphorus coil is used for both excitation and receiving. A phosphorus spectrum obtained from a 5 x 5 x 5 cm cube in the brain of a normal volunteer is shown in Figure 7.

Figure 4 The 1.5 T Gyroscan Magnetic Resonance Imaging and Spectroscopy system manufactured by Philips Medical Systems. In the foreground the operator can be seen in front of the control console on top of which is the box used to adjust the magnet shim. Through the window can be seen the patient couch and magnet.

Figure 5 Positioning of the RF coils about the head during acquisition of a phosphorus spectrum from the brain. The proton coil is used both to collect magnetic resonance images to identify the region of interest and to acquire localized proton signals used to shim the static magnetic field $B_0$.

Figure 6 A magnetic resonance image is used to select a volume in the brain from which a phosphorus NMR spectrum is obtained.

Figure 7 In vivo phosphorus n.m.r. spectrum collected from the brain of a healthy volunteer. The resonances arising from the different phosphorus compounds are labelled as follows: PME, phosphomonoesters; Pi, inorganic phosphate; PDE, phosphodiesters; PCr, phosphocreatine; g, a, b ATP, gamma, alpha and beta adenosine triphosphate.
CHEMICAL SHIFT MAPPING

In chemical shift mapping, instead of collecting a single NMR spectrum from a selected region, a map is produced of the spatial distribution of each of the compound's chemical shifts. The method has been modified to enable one-dimensional chemical shift images to be obtained from the human body. A one-dimensional $B_1$ field gradient, of duration $t_1$, is followed by a period, $t_2$, of free precession in the homogeneous static field $B_0$. A series of acquisitions are performed, varying $t_1$ each time. During the period $t_1$, the nuclear spins rotate in a plane containing the $z$ axis. The volume angle of the rotation during $t_1$ is given by equation (1). The larger the value of $B_1$, the higher the frequency of this rotation. Hence, for the linear $B_1$ field gradient, Fourier transformation with respect to $t_1$ yields a frequency variable $f_1$ which linearly describes position along the $B_1$ field gradient. Fourier transformation with respect to $t_2$ provides the chemical shift information for each of the spatial regions defined by $f_1$.

A second method for obtaining chemical shift images uses pulsed magnetic field gradients. After the nuclear spins have been excited into the $xy$ plane, an applied linear magnetic field gradient makes the frequency of precession of the nuclei a linear function of position. When the field gradient is removed the frequencies of precession are equal, although a small variation exists dependent on the homogeneity of the static field $B_0$. However, a phase variation exists in the direction of the applied field gradient, so that the spins are phase encoded with respect to their positions. The signal is acquired during a period of free precession $t_2$, after removal of the field gradient. Subsequent signals are collected using different field gradient strengths, so that the resulting phase variation has a different spatial frequency each time. If $N$ signal acquisitions are performed, incrementing the field gradient by the same amount each time, a two-dimensional data set is produced comprising the $N$ sets of acquisition times $t_2$. Two dimensional Fourier transformation of this data produces both chemical shift and positional information. $N$ spectra are produced associated with $N$ slices perpendicular to the applied field gradient. One dimensional chemical shift imaging using phase encoding techniques was first applied to in vivo animal studies in 1983 and in vivo human phosphorus chemical shift images were obtained in 1987.

By using three orthogonal field gradients for phase encoding, three dimensional chemical shift images of the human body have been produced. All three gradients are applied simultaneously to produce phase encoding in all three directions. The number of acquisitions performed is given by $N_x \times N_y \times N_z$, where $N_i$ is the number of voxels in dimension $i$. As above the field gradients are removed before signal acquisition. Four dimensional Fourier transformation produces information in three spatial directions along with the chemical shift information. The technique has been used in conjunction with a surface coil to produce chemical shift images of phosphorus in human liver with a voxel size of $2 \times 2 \times 2 \text{ cm}^3$.

SUMMARY

MRI has many applications in medicine and would be the investigation technique of choice in a number of clinical conditions. However, the role of MRS in routine clinical practice is much less clear. Many of the above spectroscopic techniques are currently undergoing evaluation of their potential clinical applications. Of particular interest is the non-invasive monitoring of pathological conditions and their response to different therapeutic regimes rather than the use of MRS as a diagnostic tool.

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REFERENCES


Blood flow imaging by magnetic resonance

Atherosclerosis of major arteries is a very common disease in the Western world, and the biggest single killer. It also accounts for a considerable morbidity resulting from the ischaemic effects on a wide variety of organs, leading to angina, transient ischaemic attacks, claudication, rest pain etc. Investigation of this disease has, for many years, relied upon X-ray angiography which produces high quality images of the anatomy of the arteries under study and enables interventional therapeutic procedures, most notably angioplasty, to be performed. However, X-ray angiography has several important drawbacks. First, it is necessary to inject a contrast medium into the vascular system and, even with the new low osmolar agents, this is uncomfortable and not without a significant morbidity and mortality. Secondly, the patient is exposed to X-radiation and, thirdly, as the femoral artery is punctured, most angiograms necessitate an overnight hospital stay with the associated financial implications.

X-ray angiography has been greatly improved by digital subtraction techniques which allow angiography to be performed as an out-patient procedure using small catheters, a lower contrast medium dose and a possible reduction in X-radiation dose. Nevertheless, for all this, it still remains an unpleasant, expensive and potentially hazardous investigation. Duplex ultrasound offers a non-invasive alternative with many attractions. It is a truly non-invasive technique, as no significant harmful effects are known. However, it too suffers from many drawbacks. Ultrasound is attenuated by gas and bone, so that certain areas of the body are less accessible and, although B-scan ultrasound produces images of vessels, the resolution is poor compared to X-ray angiography. In order to study a long vessel, a picture has to be built up from several images. Such scans are highly operator-dependent and time consuming, and it is sometimes difficult to identify which vessel one is imaging. The volume of blood flowing per unit of time through named arteries and to individual organs is a very basic physiological parameter. A non-invasive accurate in-vivo method for measuring blood flow would be of great value in both clinical and research situations. The only non-invasive in-vivo technique at present applied to measure blood flow through individual arteries is Doppler ultrasound. This has proved itself accurate in those vessels which are accessible to ultrasound. However, an inherent problem in the Doppler technique is that it does not measure flow directly. Values for instantaneous velocity are obtained, which are then integrated throughout the cardiac cycle to produce a figure for time-averaged velocity. It is then necessary to measure the cross-sectional area of the vessel under study in order to calculate flow. Thus, this method involves several stages at which errors may occur. The need for accurate measurement of blood vessel cross-sectional area is a particular drawback as many vessels, not just those within the head, are difficult to image.

Other techniques have also been used for in-vivo blood flow measurement. For example, a limited degree of success has been achieved using radioactive isotopes, and quantitative measurements from digital X-ray angiography. However, these techniques are invasive. There is, therefore, a need for a rapid non-invasive and accurate technique for imaging the vascular system and for performing quantitative blood flow measurements.

MR Flow techniques
Flow effects on magnetic resonance (MR) signals have been known for over 30 years, and no sooner had the first magnetic resonance images of the human body been produced than workers in several centres were studying ways of exploiting these effects to produce quantitative and qualitative information on blood flow.

MR angiography
One of the attractive features of conventional X-ray angiography is the production of a projection image of the contrast-filled blood vessels within the beam. However tortuous the vessel may be, it will be shown throughout its length (albeit foreshortened in places). Attempts to provide images of the blood vessels using MR have generally
concentrated on simulating such a projection image with a section thick enough to contain all the vessels of interest. These techniques have been collectively termed MR angiography. The first technique of MR angiography relied on the different degrees of phase dispersion within voxels in different parts of the cardiac cycle. Van Wedeen used a conventional 2D FT spin-echo sequence. Two cardiac-triggered images were acquired: one at systole and the other at diastole. In diastole the blood flow is relatively slow, so there was no significant phase dispersion, resulting in preservation of the signal from the slow-flowing blood. In the image acquired during systole, however, significant phase dispersion caused a reduction in the signal intensity from the more rapidly flowing blood. The signal from stationary tissue was the same in both images. Subtraction of the systolic from the diastolic image resulted in an MR angiogram.

A disadvantage of this technique is that it will only work where the flow is relatively strongly pulsatile. Vessels in which there is a constant velocity cannot be imaged. Probably a more important disadvantage is the change in position of some vessels in different parts of the cardiac cycle. Such movement of vessels will produce registration errors when the images are subtracted. A further disadvantage is the variability in the phase dispersion at a fixed point in the cardiac cycle in a patient with vascular disease or with a variable heart rate.

The general technique of MR angiography, which overcomes the problems mentioned above, produces different velocity-dependent phase shifts in two images which can be at the same phase of the cardiac cycle. Stationary tissue will have the same phase shift in each image resulting in zero signal after subtraction. Careful choice of the imaging parameters to produce a phase shift of 180° at the peak velocity likely to be found in the region to be studied will give a maximum signal within the vessel following subtraction.

Van Wedeen et al. use a subtraction of two SE sequences with an unpulsed frequency encoded gradient and different echo times. This variation of TE produces the velocity-dependent phase shift in the subtracted image. Dumoulin et al. inserted an additional bi-polar pulse during the evolution period of a gradient echo sequence. Data for two images were acquired, the polarity of the additional bi-polar pulse being reversed for the second image. This reversal modulates the phase of the moving spins but has no effect on the stationary spins.

Groen et al. developed a sequence with the acronym FLAG (Flow Adjusted Gradients); this is the technique we use, and we shall discuss it in detail below.

**MR velocity-phase maps**

The technique of using MR images to
2. MR angiogram of upper legs of normal volunteer. Note good visualization of deep and superficial femoral arteries, but motion artifact blurring of signal within the pelvis.

3. MR angiogram of lower legs of normal volunteer. This is a composite image taken by summing 6 images taken throughout the cycle.

antitate velocity was developed by Bryant et al.1, Moran1, Moran7 and Van Dijk11. These early methods had limitations for clinical application, but there have been a number of significant advances which make clinical use practical. Nayler et al., Ridgway and Smith9 and Young et al.15 utilized a field echo to increase the range of velocities that could be measured. Methods to correct the phase dispersion, which can produce a loss signal, were also developed8-9. The technique has also been extended to measure very slow flow10.

- **FLAG sequence**

The FLAG sequence additional bi-polar gradient pulses added to a field echo sequence to compensate for the phase-dependent phase shifts. Two images are acquired, where the bi-polar pulse compensates for velocity, giving zero phase change in flowing blood, and the second the added bi-polar gradient changed in magnitude a velocity-dependent phase shift. For MR angiography a thick section is selected and an additional flow gradient is applied in the ‘slice select’ direction to phase the signal from stationary tissue. Complex traction of the two images gives the required high signal from flowing blood and zero signal from the background. For velocity-phase maps, thin sections are selected and the separate phase images are subtracted. The FLAG sequence is illustrated in Fig 1. In addition to the standard sequence parameters such as slice thickness, matrix size, TE flip angle etc., the following additional operator-adjustable parameters are available: cardiac triggered or continuous acquisition, direction of flow sensitivity, velocity to give the maximum signal or phase difference, and slice dephasing gradient.

Recently there have been a number of developments of the FLAG sequence. The minimum echo time has been reduced to enable thin-section phase maps of the aorta to be acquired with less signal loss; this has also improved the signal-to-noise ratio in MR angiograms. In a further refinement of the FLAG sequence the flow-sensitive and compensated acquisitions are interleaved within the same cardiac cycle instead of at the same point in adjacent cycles. Consequently the separation is reduced from perhaps 800 ms to 50 ms, reducing the effect of extravascular motion. This technique is referred to as Rapid Sequential Excitation5.

Our work has been carried out on a Philips Gyroscan MR system operating at 1.5T using the original FLAG sequence, the short echo modification and also the rapid sequential excitation version.
Fig. 4. Comparison of X-ray and MR angiograms of a patient with a block at the origin of the left peroneal artery.

a. X-ray angiogram.
b. MR angiogram demonstrating absence of flow in the left peroneal artery.

Fig. 5. MR angiogram of a patient with patent bilateral femoro-popliteal grafts.

MR angiography

Methods

In MR angiography, the acquisition sequence is normally synchronized to the cardiac cycle in order to obtain images at different times within the cycle. This is important in regions such as the lower limb where the flow is very pulsatile, but is not necessary in the carotid arteries where the flow is continuously present throughout the cardiac cycle. Using the standard FLAG sequence to assess flow in the lower limb arteries, we normally collect images at 6 points throughout the cycle, at 75 ms intervals starting 50 ms after the R-wave. We have found a velocity setting of 0.5 m/s to be optimal for the lower leg and cerebral circulation, and 1.0 m/s for the upper leg.

The slice thickness is selected at the minimum which ensures inclusion of all the vessels of interest. A maximum thickness of 10 cm is satisfactory in most circumstances. The direction of flow sensitivity can be set in either of the two co-ordinate axes within the slice. One direction usually suffices, but two directions are sometimes necessary if the vessels are very tortuous. The thick slice under study may be angled if necessary, which is especially useful for examinations of the carotid bifurcation. We generally use
Fig. 5. Comparison of X-ray and MR angiograms in a patient with aortobifemoral graft, and a block of the left superficial femoral artery at its origin with reconstitution via collaterals from the deep femoral artery to the popliteal.

a, b. X-ray angiograms.

c, d. MR angiograms.
Fig. 7. Comparison of X-ray and MR angiograms in a patient with a block of the right superficial femoral artery at its origin, and reconstitution of the popliteal. There is also a short block of the left superficial femoral artery at the adductor hiatus.

a. X-ray angiogram.
b. MR angiogram.

Fig. 8. MR angiogram, with sagittal view of the right side of head and neck of a normal volunteer. This demonstrates the common, external and internal carotid arteries, the siphon, the vertebral and basilar arteries and the posterior superior sagittal sinus and transverse sinus.

Fig. 9. MR angiogram of the carotid bifurcation in a normal volunteer. Note the loss of signal in the bulb due to recirculation effect.
Fig. 10. Comparison of 3 cm thick midline sagittal MR angiograms in a normal volunteer and a patient with occlusion of the superior sagittal sinus.

a. MR angiogram in a normal volunteer, using velocity encoding of 25 cm per second. Note good visualization of flow in the superior sagittal sinus.

b. Same scan as Fig. 10a in a patient with occlusion of the superior sagittal sinus. Note absence of flow signal from within the sinus.

Matrix size of 256 x 256, a TE of 25 ms, a flip angle of 5° and four measurements. A slice dephasing gradient is used to reduce signal from stationary tissue; the magnitude of this gradient depends on the thickness of the slice.

Investigation of the lower limb arteries from aorta to bifurcation using the standard FLAG sequence takes 25 minutes for each position, the whole lower limb being imaged in two positions making a total imaging time of under one hour. Untriggered angiograms of the head and neck can be obtained in under 10 minutes.

Results in the lower limb
Using the original FLAG sequence, high quality images of femoral, popliteal and lower leg vessels were obtained in all normal volunteers (Figs 2, 3). The veins could be easily distinguished from the arteries by their relative intensity at different phases through the cycle and there were very few artifacts. However, as was also found by Laat3, studies in patients were less satisfactory. The quality of the images was lower in older patients and those with more generalized disease. This is probably due to patient motion.

Despite these difficulties we have obtained many diagnostic studies, such as the example shown in Fig. 4. Particularly high quality images were obtained in a series of 10 patients with femoro-popliteal grafts. All the patent grafts were clearly shown on the MR angiograms, as illustrated in Figure 5.

The quality of the angiograms in the thigh and lower leg was improved with the use of the new sequence with a shorter TE. However, a major problem still remained, in that the abdominal aorta and iliac arteries were impossible to image adequately due to bowel and respiratory motion (Fig. 2). Rapid sequential excitation showed a dramatic improvement in the ability to image the iliac and aortic bifurcation and, for the first time, we were able to routinely visualize the lower aorta and the iliac arteries.

We have now used this sequence to study several patients, with excellent results (Figs 6, 7).

Results in the head and neck
Nowadays, most carotid angiography is performed for the detection of atherosclerosis (bifurcation and siphon) or intracranial aneurysm. Non-invasive images with Doppler can compete very strongly with X-ray angiography for carotid bifurcation lesions, but cannot show concomitant
Fig. 11. Velocity-phase mapping.

a. Spin echo transaxial scan through the ascending and descending aorta, used to plan for phase maps.
b. MR phase map corresponding to Fig. 11a. This shows a fairly uniform white signal indicating flow into the slice in the ascending aorta, and a black signal indicating flow out of the slice in the descending aorta. The background and the pulmonary artery show a mid-grey signal indicating zero flow vector within the slice.
c. Sagittal MR phase map of ascending and descending aorta planned from Fig. 11b.
d. Colour coded MR phase map as in Fig. 11c, superimposed on the correspondingly orientated spin echo image.
disease of the siphon or intracranial aneurysms.

MR angiography could potentially image both areas (Fig. 8) and has been applied to imaging the carotid bifurcation (Fig 9). Studies in this area, however, have been disappointing for several reasons. First, there is confusing overlap of the signal from the jugular vein. Secondly, the spatial resolution is not sufficient to detect small but nevertheless important lesions and, thirdly, flow disturbances near the carotid bulb are common, causing signal loss even in normal volunteers.

The ability of MR to image within the skull and to obtain low information in two directions should allow it to be used to produce angiograms of the intracranial vessels. Unfortunately, however, the spatial resolution of this system is not yet sufficient to cope adequately with the small size of these vessels or to detect small but clinically significant aneurysms. On the other hand, the cerebral venous drainage is a large-volume, low-velocity system which can be usefully imaged with MR angiography. We have obtained high quality angiograms of the superior sagittal sinus in normal volunteers (Fig. 10a). In two patients with proven thrombosis of the superior sagittal sinus we have demonstrated absence of flow (Fig 10b), confirming the diagnosis which is very difficult to make clinically, and normally requires high quality rebral X-ray angiography.

- R velocity-phase maps

Methods
The same pulse sequences used to produce the MR angiograms can also be utilized to produce quantitative phase images whereby the grey scale is proportional to the velocity of the blood, black representing maximum flow in one direction, and white representing maximum flow in the opposite direction. Mid-grey represents stationary tissue.

It is important to realize that although both the angiograms and the velocity-phase maps are produced using the same pulse sequence, the variable parameters and image processing are very different so that it is not, in fact, possible to quantitate flow from an angiographic phase. Velocity-phase maps have potentially more clinical use in the MR angiographic technique described above. Unlike Doppler methods, time-averaged flow can be determined directly by analysing a region of interest physically larger than the cross section of the vessel. It is not necessary to measure the exact area of the vessel, but only the velocity profile within it. This can then be used to measure blood flow to individual organs. Time-

averaged or instantaneous velocity can be measured if desired. Phantom experiments and comparison of our technique with Doppler ultrasound studies in normal volunteers have shown it to be accurate for quantifying both time-averaged velocity and flow. Since clinically, and in most research situations, the primary interest is in time-averaged flow or velocity, there is no need to synchronize the sequence to the cardiac cycle, and continuous acquisition sequences are used in many vessels. This significantly reduces the time needed for data acquisition and processing – a typical study taking about 8 minutes to acquire.

Results

Typical examples of cardiac-gated phase maps of the thoracic aorta are shown in Figure 11b and 11c. Phase maps can be colour coded to give more immediate visual impact of the direction of flow and, if desired, superimposed on standard spin echo images (Fig. 11d). This is likely to have its most important application within the heart to study flow disturbances arising from congenital defects.

We have used the technique to measure blood flow in the common and internal carotid arteries (Fig. 12), middle cerebral artery (Fig. 13b), renal artery and femoral artery.
Fig. 13. Velocity-phase mapping of the left middle cerebral artery.

a. Sagittal field echo image perpendicular to the left middle cerebral artery before its division. This image is planned from a series of routinely orientated transaxial spin echo head scans which, in almost all cases, adequately demonstrate the course of the middle cerebral artery for planning purposes. Note the high signal in the centre of the vessel at the peak of the parabolic flow profile. Nearer the edges of the vessel there is a large variation of velocities within each voxel which produces dephasing and, therefore, loss of signal.

b. Corresponding phase map from which flow measurements are obtained.

Results from the carotid and femoral arteries were compared with Doppler ultrasound and good agreement was achieved.

The sensitivity of the technique to velocity changes can be of great help as an adjunct to the interpretation of MR angiographic defects. An area of signal loss on an MR angiogram can be due to a stenosis (the high velocity leading to signal loss) or an occlusion. Phase mapping can more clearly define the problem in difficult situations. (Fig. 14).

Conclusions
We now have a technique which, totally non-invasively and with no contrast medium, provides a high quality ‘projective’ angiogram of all the major vessels of the lower limbs in under one hour. Currently, the technique has clinical applications in the diagnosis of peripheral vascular disease and assessment of femoro-popliteal grafts. X-ray angiography will continue, however, to offer the potential for angioplastic intervention.

MR angiography has immediate application in the investigation of cerebral venous sinus disease and long term potential in the imaging of the intracranial arteries. It is unlikely to ever compete with ultrasound in the diagnosis of disease of the carotid bifurcation.

MR phase maps can measure flow and velocity accurately and non-invasively through large and medium size vessels. There is a huge potential clinical and research application for the technique.

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Fig. 14. Comparison of X-ray angiography and MR velocity-phase mapping in a patient with tight stenosis of the left external iliac artery.

a. X-ray angiogram.

b. MR thin section sagittal phase map along the left external iliac artery demonstrating very high signal at the stenosis where the velocity is very high. Note the dark signal in the iliac vein lying anterior to the artery.

- bibliography


Magnetic resonance imaging of the infant heart at 1.5 T

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Magnetic resonance (MR) has been found to be of value in many fields of medicine but has yet to find an established place in the study of diseases of the heart. Published studies of the heart have demonstrated its ability to identify areas of acute myocardial infarction (Been et al, 1985b) and old infarcts (Higgins et al, 1984b), to recognize hypertrophic cardiomyopathies (Been et al, 1985a), pericardial effusions (Stark et al, 1984) and constrictive pericarditis (Soulen et al, 1985). It is, in addition, possible to measure ventricular volumes (Longmore et al, 1985) and images containing blood-flow-related data can be acquired (Ridgway & Smith, 1986; Underwood et al, 1987).

Magnetic resonance imaging (MRI) has been suggested to have a particularly valuable role in the study of the anatomy of the congenitally malformed heart (Higgins et al, 1984a) and imaging studies of a limited number of cases have been published (Jacobstein et al, 1985a,b; Boxer et al, 1986; Didier & Higgins, 1986; Didier et al, 1986; Fletcher & Jacobstein, 1986; von Shultess et al, 1986). However, in these studies from a small number of research centres there is very little experience in the use of MRI in infants. Since 90% of patients with congenital heart disease present in the first year of life (Hoffman & Christianson, 1978), if MRI is to play a role in the investigation of congenital cardiovascular abnormalities it is essential to be able to image infants within the first year.

Imaging of the heart has generally been performed using low- and medium-field strength magnetic resonance systems with a field strength of 0.6 T or less (Higgins et al, 1985; Didier et al, 1986; Smith et al, 1986; Underwood et al, 1987). Cardiac triggering poses few problems at these field strengths but it has been thought that difficulties may arise with higher-field systems at 1.5 T and above. Despite this, however, the high-field MR systems have a potentially important advantage for the study of congenital heart disease in children: the signal-to-noise ratio from these systems is higher than from low- and mid-field systems resulting in improved spatial resolution or shorter imaging times. Both these advantages are particularly valuable in the study of infants.

An alternative approach for paediatric imaging is to use the echo-planar imaging technique (Rzedzian, 1983). With this method the images can be obtained in 60–120 ms, so that patient and cardiac motion are frozen. A drawback with echo-planar imaging is that, as yet, the image quality is poorer than that obtained using triggered conventional imaging techniques at the same field strength. In addition, the technique is not generally available.

The research described in this paper set out to determine the feasibility of investigating cardiovascular abnormalities in young infants making use of the improved spatial resolution that can be obtained with a high-field MR system.

Imaging technique

The subjects were imaged using a 1.5 T whole-body MR system manufactured by Philips Medical Systems (Gyroscan). To image the heart or the great vessels, the acquisition of the MR image must be synchronized with the patient's heartbeat. To achieve this with a high-field MR system, care must be taken over the positioning of the electrodes. If the three electrodes are placed in the conventional position used for triggering in other imaging modalities, or in MRI at low- and medium-field strengths, unacceptable distortion of the electrocardiogram (ECG) is produced. The R wave is sharply defined as required but the T-wave amplitude is greatly increased and may be the same size as the R wave, thus...

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introducing the possibility of incorrect triggering from the T wave. This distortion does not indicate a risk to the patient because it is not caused by the magnetic field disturbing the electrical conduction within the heart but by the motion of conducting material, particularly blood in the great vessels, in the high magnetic field. The distortion is reduced to an insignificant level by placing the three electrodes close together around the left nipple. Pre-gelled disposable silver/silver chloride electrodes (medicostat Q-10-A) were used and the ECG was transmitted using a telemetry system. Specialized electrodes, sometimes proposed for MRI, were not required.

The standard proton radiofrequency (RF) transmit and receive head coil was used. This is circular in cross-section with an internal diameter of 32 cm. The size and design of the coil enabled the infant to be positioned inside without difficulty and visual contact was possible if necessary throughout the investigation.

The infant was wrapped in a blanket and placed on a thin, firm mattress, which was strapped in a semicircle round the patient to restrict movement. The majority of subjects were imaged supine. The first 12 babies to be imaged were fed immediately prior to the MRI investigation and the investigation was undertaken without sedation. However, although this procedure was successful in most cases, in some, time was needed for them to settle during the investigation and this difficulty increased as infants with more severe abnormalities were studied. Consequently, most subsequent infants were given sedation of 75 mg/kg chloral.

Simultaneous 5 mm sections were acquired using a spin–echo sequence with echo time (TE) of 30 ms and a field of view of 200–250 mm. A matrix size of 256 × 256 was used but with each image being constructed from 180 signals instead of the usual 256. This resulted in a reduction in scan time with only a slight reduction in

Figure 1. Aortic coarctation in a 6-week-old infant. This section shows clearly the relationship of the left subclavian artery (LSA) to the stenosis.

Figure 2. Aortic hypoplasia in a 12-week-old infant. The MR scan clearly demonstrates severe hypoplasia of the whole aortic arch and descending aorta without any localized narrowing. The patient's weakened medical condition and reduced lumen made her a poor candidate for X-ray angiography.
spatial resolution in the phase-encoded direction. The MR system was triggered on each heartbeat, even for heart rates as high as 140 beats per minute. Once in position, a series of sections was acquired in the transverse direction from which the angles of obliquity could be obtained. Double-oblique imaging was then used to obtain the appropriate angulation, for example, through the arch and descending section of the aorta. Two signal averages were performed resulting in a typical scan time for a set of seven sections of about 6 min. The total investigation time per patient varies with the complexity of the congenital abnormality; typically, it is currently between 60 and 90 min.

Measurements were performed on a high-contrast resolution phantom and a line pair separation of 1.5 mm and 0.95 mm was obtained in the phase-encoded and frequency-encoded directions, respectively. The RF power absorption for children of different age imaged using the procedure are all below the minimum recommended specific absorption rate (SAR) with a typical value for a 6-week infant of 0.08 W/kg.

Results

The technique has been used to image over 40 infants under the age of 12 months. Electrocardiogram triggering was successfully achieved in all cases and respiratory motion artefacts were not found to be a problem. Constant observation of the patient from the end of the magnet and monitoring of the ECG ensured patient safety. In one instance, an ill, oxygen-dependent infant was imaged with the attendant clinician inside the magnet.

The technique has been successful in imaging a range of congenital disorders including coarctation of the aorta \( n = 22 \), atrial and/or ventricular septal defects \( n = 14 \), transposition \( n = 3 \), Fallot’s tetralogy \( n = 3 \) and multiple cardiac defects. Images were obtained successfully from all patients and examples showing the quality of images obtained are given in Figs 1 to 5.

Discussion

The superior signal-to-noise ratio obtained with a high-field MR system enables good-quality, thin sections to be obtained in a reasonably short time period, which we feel is particularly suitable for the investigation of cardiac disease in the infant. Imaging soon after feeding can eliminate the need for any sedation, but for reliable, rapid throughputs imaging, mild sedation is required. A series of diagnostic quality images can be obtained in a sufficiently short time to enable MRI to be considered as a possible routine technique for cardiac investigation in these children. Although a rigorous comparative study with ultrasound has not yet been undertaken, our experience suggests that the amount of anatomical detail is at least equal to and generally greater than that achieved using echocardiography. In a number of cases MRI eliminated the need for an X-ray angiogram.

A study is currently in progress to determine the clinical value of MRI in congenital heart disease in infants.

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References


Magnetic resonance imaging at a high field strength of ventricular septal defects in infants

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SUMMARY Magnetic resonance imaging at a high field strength has potential benefits for the study of the heart in infants, which is when most congenital heart disease presents. Seventeen infants with various anatomical types of ventricular septal defect were studied by this technique. Good quality, high resolution, images were obtained in every case. There were no major practical problems. The morphology of the defects in all 17 hearts was displayed in great detail. In some instances, the interpretation of the images resembled that of equivalent images from cross sectional echocardiography. But this new technique allowed imaging in planes that cannot be obtained by echocardiography. One particularly valuable plane gave a face on view of the inlet and trabecular components of the septum. This allowed very precise localisation of defects in these areas. The relation between the defects and the atrioventricular and arterial valves was exceptionally well shown in various different imaging planes. One patient in the series had multiple trabecular defects that were clearly shown.

Magnetic resonance imaging gives detailed morphological information about ventricular septal defects.

High resolution cross sectional imaging can provide a precise description of the morphology of defects in the ventricular septum, and echocardiography is a useful investigation in patients with such defects. But newer cross sectional imaging methods have not been evaluated in the study of ventricular septal defects. Gated cardiac magnetic resonance imaging, with image systems of low field strength, can detect large defects in older patients. The structural diagnosis of congenital heart defects, however, is usually made in infants. This is the age group in which new imaging techniques need to be evaluated.

We studied an unselected series of infants with various types of ventricular septal defect by a magnetic resonance imaging system that operates at high field strength.

Patients and methods

Patients

We studied 17 infants with ventricular septal defects undergoing clinical assessment at Guy's Hospital (table). The selection criteria were the presence of a known ventricular septal defect and the availability of the infant for imaging. They were aged between three weeks and six months. All had a normal atrial arrangement. Nine had normal cardiac connections. In three there was a discordant ventriculoarterial connection, and one of these infants also had a discordant atrioventricular connection. Two cases had double outlet from the morphologically right ventricle, two others had pulmonary atresia, while the final patient had a common arterial trunk. One patient has since died and the heart has been examined (patient 5). Another has undergone surgical closure of the defect (patient 17).

Methods

The imaging protocol was described in detail elsewhere. In all patients an initial series of sections was acquired in the transverse plane of the body, encompassing the whole heart. Typically, this required seven sections. Subsequent imaging planes were chosen in the light of the suspected cardiac abnormality and the position and orientation of the cardiac structures as shown on these images. Imaging planes...
Morphology demonstrated by magnetic resonance imaging at high field strength

Table

<table>
<thead>
<tr>
<th>Case No</th>
<th>Age</th>
<th>Morphology demonstrated by magnetic resonance</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>3 mnth</td>
<td>Large muscular inlet defect, multiple small muscular trabecular defects</td>
</tr>
<tr>
<td>2</td>
<td>4 mnth</td>
<td>Double outlet of the morphologically right ventricle, anterior aorta, posterior pulmonary trunk overriding a perimembranous confluent defect</td>
</tr>
<tr>
<td>3</td>
<td>3 wk</td>
<td>Pulmonary atresia, aorta arising anteriorly from the right ventricle, no central pulmonary arteries, muscular outlet defect</td>
</tr>
<tr>
<td>4</td>
<td>4 mth</td>
<td>Perimembranous confluent defect with overriding by the tricuspid valve and malalignment between the atrial septum and the trabecular ventricular septum</td>
</tr>
<tr>
<td>5</td>
<td>3 mth</td>
<td>Double outlet of the right ventricle with the great arteries arising, side by side, the aorta to the right, complete muscular infundibulum beneath both great arteries, defect with entirely muscular margins</td>
</tr>
<tr>
<td>6</td>
<td>6 wk</td>
<td>Double committed juxta-arterial defect with muscular posteroinferior border, coarctation</td>
</tr>
<tr>
<td>7</td>
<td>4 mth</td>
<td>Pulmonary atresia with central confluent pulmonary arteries; aortic valve overrides a perimembranous outlet defect</td>
</tr>
<tr>
<td>8</td>
<td>6 wk</td>
<td>Perimembranous inlet defect with accessory tricuspid valve tissue partially closing it</td>
</tr>
<tr>
<td>9</td>
<td>3 mth</td>
<td>Small perimembranous defect with accessory tricuspid valve tissue</td>
</tr>
<tr>
<td>10</td>
<td>1 mth</td>
<td>Muscular outlet defect, complete muscular subaortic infundibulum coarctation</td>
</tr>
<tr>
<td>11</td>
<td>5 mth</td>
<td>Discordant ventriculoarterial connection, muscular outlet defect with posterior deviation of the outlet septum and subpulmonary stenosis</td>
</tr>
<tr>
<td>12</td>
<td>1 mth</td>
<td>Perimembranous confluent defect with overriding tricuspid valve, overriding aortic valve, and infundibular pulmonary stenosis</td>
</tr>
<tr>
<td>13</td>
<td>3 wk</td>
<td>Perimembranous defect with overriding common arterial trunk</td>
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<tr>
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<td>2 mth</td>
<td>Small perimembranous defect, coarctation</td>
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<td>15</td>
<td>5 mth</td>
<td>Discordant ventriculoarterial connection, muscular outlet defect</td>
</tr>
<tr>
<td>16</td>
<td>1 mth</td>
<td>Discordant anteroseptal and ventriculoarterial connections, perimembranous outlet defect with overriding pulmonary valve</td>
</tr>
<tr>
<td>17</td>
<td>6 mth</td>
<td>Single large muscular inlet defect</td>
</tr>
</tbody>
</table>

Results

Images of adequate quality for morphological study were obtained in every patient. The electrocardiographic gating was also satisfactory in all instances.

Fig 1  Magnetic resonance image of oblique coronal section in patient 1. The section includes the ventricular septum and shows multiple muscular defects (arrows), the largest being in the inlet septum. There is no defect in the outlet septum; it appears to be defective because it curves backwards out of the plane of the section.

Fig 2  Magnetic resonance image of oblique coronal section in patient 17. There is a single large muscular defect (arrow). The membranous septum was shown to be intact in sections orthogonal to this, because it is thin it appears to be defective in this plane.
Magnetic resonance imaging of ventricular septal defects

The table lists the descriptions of the morphology that were discernible from the magnetic resonance images. The morphological criteria for the description of ventricular septal defects have been described in detail elsewhere. In the two patients in whom direct observation of the morphology of the defect was possible at necropsy or operation the magnetic resonance description of the defects was confirmed. Although the imaging planes used in each patient were different, some general principles emerged. For
The use of oblique imaging planes individually tailored to the morphology seen in each case was essential (figs 4, 5, and 6). The morphology of the ventricular septal defects studied in this series was very varied. There was no standard imaging protocol that would have provided the best images in every patient.

Discussion

This series contains examples of a wide variety of anatomical types of ventricular septal defects. In every patient, the magnetic resonance study provided the morphological details to define the anatomy of the defects present. The high resolution of the images obtained, even in these small infants, compared favourably with those from cross sectional echocardiography. A unique feature of magnetic resonance imaging is its capability to image in any plane through the heart. There should in theory, therefore, be no feature of the morphology of any ventricular septal defect that cannot be determined within the limits of resolution of the images. Further experience will indicate whether this is indeed true. It is already
The role of magnetic resonance imaging in the diagnosis of ventricular septal defects

Clear to us that considerable experience is required in choosing the best imaging plane in individual patients. We believe that the standard series of imaging planes that have been described by others, have no place in the imaging of congenital malformations of the heart. The protocol in each patient has to be matched to the observed anatomy. This is already well established in echocardiography. The same general principles about the interpretation of cross sectional images of the heart apply to magnetic resonance images.

Relation of the Defect to Arterial Valves
Examples of muscular, perimembranous, and doubly committed aorta-septal defects all opening between the subarterial outlets are included in this series. The outlet septum cannot be shown face on in the same way as the inlet and trabecular components of the septum. This is because, in hearts with normal connections, the septum curves posteriorly out of the imaging plane. Despite this, extension of defects to space between the subarterial outlets was exceptionally well shown. For example, the fibrous continuity between aortic and pulmonary valve in a patient with a doubly committed defect was very clear (fig 5). A comparison of the necropsy specimen with the magnetic resonance images in a patient with a bilateral infundibulum shows how well this aspect of the anatomy was shown (fig 6). Features such as overlapping of arterial valves and deviation of the outlet septum were also well shown.

Relation of the Defect to the Central Fibrous Body and Atrioventricular Valves
The relation of defects to the central fibrous body was shown in various planes. So perimembranous defects were readily distinguished from other defects. Perimembranous defects extending into the inlet septum and roofed by the atriointricular valves were well shown in a four chamber imaging plane. In two patients in this series such defects were associated with overriding of the tricuspid valve. This was clearly shown (fig 4). In one of these, however, there was echocardiographic evidence of additional straddling of the tension apparatus of the valve that was not evident on the magnetic resonance images.

Multiple Defects
In one patient in this series, the magnetic resonance images clearly showed multiple small trabecular defects in association with a large muscular defect of the inlet septum (fig 1). Only one trabecular defect was shown in this patient on both Doppler colour flow mapping and cineangiography. It is, therefore, possible that magnetic resonance imaging will prove of special value in detecting the presence of multiple defects. Great care is needed to differentiate trabeculations on the right ventricular surface of the septum from true small trabecular defects. By following the defects in contiguous sections from the cavity of the left ventricle to that of the right we were confident that in this patient they were true septal defects. If this is not always done, trabeculations will be falsely identified as septal defects.

The Place of Magnetic Resonance Imaging
There are few practical problems with this imaging technique. The patient must be prevented from moving. We found that light sedation was sufficient for this. The infant does not suffer any discomfort during the imaging procedure and, if well prepared, will generally sleep soundly. The heart rate is constantly monitored and the patient's colour and respiratory motion can be observed from outside the magnet. Cardiac gating was a potential problem because of possible distortion of the electrocardiogram. But we have not found any major difficulty with this. Occasionally the position of the electrodes has to be adjusted to give the best signal, but we have always been able to obtain satisfactorily gating.

Magnetic resonance imaging can, therefore, provide excellent images of ventricular septal defects. Much of the information they contain is comparable with that from cross sectional echocardiography. The images, however, are of high resolution, and new imaging planes that are not possible with echocardiography can be used. This new technique enables a very detailed assessment of the morphology of ventricular septal defects to be made non-invasively. This calls into question the role of
cineangiography in the assessment of infants with ventricular septum defects.

This research was supported by the British Heart Foundation. The imaging system was purchased with the assistance of Sir Philip and Lady Harris and the Special Trustees of Guy’s Hospital.

References
Magnetic resonance imaging of coarctation of the aorta in infants: use of a high field strength

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SUMMARY Nineteen infants with suspected coarctation of the aorta were studied with electrocardiographically gated magnetic resonance imaging on a 1·5 T whole body imaging system. In all cases imaging was successful and produced diagnostic images of high resolution. Coarctation was shown in 12 cases. The position and shape of the coarctation were well displayed by the magnetic resonance images. In addition, they clearly showed the relation of the coarctation to arteries arising from the aortic arch and to the length and diameter of the aortic isthmus and the distal aortic arch. The anatomy was confirmed at operation in all 12 patients, except for two with a small ductus arteriosus (arterial duct), which was not seen in the magnetic resonance images. In the seven remaining patients, coarctation was excluded.

Magnetic resonance imaging produced high quality images that showed the anatomy better than other non-invasive methods. It provided all the anatomical information required for surgical correction.

Magnetic resonance has been promoted as an important advance in the imaging of congenital malformations of the heart and great vessels. As yet, however, there is little evidence that it will either supplant or supplement established imaging techniques. There is no experience in very young children, the age group in which most cases of coarctation now present. We report our experience of imaging the aorta in a group of infants with suspected coarctation with a high field strength magnetic resonance imaging system. The superior spatial resolution of such a system makes it particularly suitable for this age group.

Patients and methods

PATIENTS

Nineteen infants, aged between three days and 10 months, were imaged (table). In every case there was clinical or echocardiographic suspicion of coarctation of the aorta. Infants who were in severe heart failure that required intravenous inotropic support or ventilation were not included; otherwise there were no selection criteria. Four of the patients studied (patients 4, 5, 6, and 10) subsequently had aortograms. Surgical correction was performed in all the cases in which coarctation was confirmed by magnetic resonance imaging.

METHODS

The infants were fed just before the imaging procedure and, if necessary, sedated with 75 mg/kg of chloral hydrate. They were imaged by a 1-5 T whole body imaging system (Gyroscan) manufactured by Philips Medical Systems. Electrocardiogram electrodes were placed closely around the left nipple, an arrangement that minimises the distortion of the electrocardiogram produced by the movement of blood within the high magnetic field. The electrocardiogram was transmitted by a telemetry system. The infants were wrapped in a blanket and their movement was restricted by a firm foam mattress curved in a semicircle around them. They were imaged while supine in a 32 cm diameter proton head coil.

Electrocardiographically gated sections (5 mm thick), separated by 0-5 mm, were simultaneously acquired by a spin echo sequence with an echo time of
Results

Images of diagnostic quality were obtained in every patient. There were no imaging failures. In all patients the electrocardiographic gating was satisfac-

30 ms. The matrix used was 256 × 256, with a typical pixel size of 0.75 mm. To reduce acquisition time we used 180 phase encoding gradients instead of the usual 256 to produce the image. Two acquisitions of each phase encoding step were averaged. This resulted in a typical scan time for one set of seven sections of about six minutes.

In every case, the initial series of sections was in the transverse plane of the body, encompassing the structures between the arch of the aorta and the diaphragm. This, typically, required seven to nine sections. Subsequent imaging planes, oblique to the standard planes of the body, were chosen to show the area of suspected coarctation. The angles for these planes, chosen individually in each patient, were determined directly from the initial series of transverse images. The most valuable imaging plane for the aortic arch was found to be an oblique sagittal, rotated around a vertical axis, to show the aortic arch (figs 1–5). Occasionally, the imaging plane had to be additionally rotated around an anterioposterior axis to show the aortic arch optimally. A coronal plane, which showed the descending aorta, was also useful in some cases.

In all the patients with coarctation, the area of narrowing was shown in the oblique sagittal plane and at least one other plane, either the transverse or coronal. In most patients it was imaged in all three planes.

Table  Details of patients studied

<table>
<thead>
<tr>
<th>Number</th>
<th>Age</th>
<th>Anatomy shown by magnetic resonance</th>
<th>Surgical findings where different</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3 wk</td>
<td>Discrete shelf, mild isthmal hypoplasia</td>
<td>Small arterial duct</td>
</tr>
<tr>
<td>2</td>
<td>3 wk</td>
<td>Waist lesion, no isthmus, mild distal arch hypoplasia</td>
<td>Agree</td>
</tr>
<tr>
<td>3</td>
<td>8 days</td>
<td>Discrete shelf, severe isthmal hypoplasia, patent arterial duct</td>
<td>Agree</td>
</tr>
<tr>
<td>4</td>
<td>17 days</td>
<td>Mild isthmal narrowing, no coarctation</td>
<td>Agree</td>
</tr>
<tr>
<td>5</td>
<td>6 days</td>
<td>Discrete shelf, anomalous right subclavian artery proximal to coarctation</td>
<td>Agree</td>
</tr>
<tr>
<td>6</td>
<td>2 month</td>
<td>Anomalous pulmonary veins to coronary sinus, coarctation excluded</td>
<td>Agree</td>
</tr>
<tr>
<td>7</td>
<td>5 days</td>
<td>No coarctation</td>
<td>Agree</td>
</tr>
<tr>
<td>8</td>
<td>6 wk</td>
<td>Discrete shelf, mild isthmal hypoplasia, ventricular septal defect</td>
<td>Agree</td>
</tr>
<tr>
<td>9</td>
<td>10 month</td>
<td>Waist lesion, no isthmus</td>
<td>Agree</td>
</tr>
<tr>
<td>10</td>
<td>3 month</td>
<td>Hypoplasia of abdominal aorta, no coarctation</td>
<td>Agree</td>
</tr>
<tr>
<td>11</td>
<td>3 days</td>
<td>Normal aortic arch</td>
<td>Agree</td>
</tr>
<tr>
<td>12</td>
<td>3 month</td>
<td>Discrete shelf, mild isthmal hypoplasia</td>
<td>Agree</td>
</tr>
<tr>
<td>13</td>
<td>1 month</td>
<td>No isthmus, hypoplasia of distal arch, ventricular septal defect, subaortic infundibulum</td>
<td>Small arterial duct</td>
</tr>
<tr>
<td>14</td>
<td>2 month</td>
<td>Mild isthmal hypoplasia, discrete shelf, small ventricular septal defect</td>
<td>Agree</td>
</tr>
<tr>
<td>15</td>
<td>10 days</td>
<td>Mild isthmal hypoplasia, no coarctation</td>
<td>Agree</td>
</tr>
<tr>
<td>16</td>
<td>2 month</td>
<td>Severe isthmal hypoplasia, discrete shelf</td>
<td>Agree</td>
</tr>
<tr>
<td>17</td>
<td>12 days</td>
<td>Normal aortic arch</td>
<td>Agree</td>
</tr>
<tr>
<td>18</td>
<td>7 wk</td>
<td>Waist lesion, hypoplasia of distal arch, no isthmus</td>
<td>Agree</td>
</tr>
<tr>
<td>19</td>
<td>6 wk</td>
<td>Discrete shelf, distal arch and isthmal hypoplasia</td>
<td>Agree</td>
</tr>
</tbody>
</table>

Fig 1  A normal aortic arch in a three day old infant (patient 11). This image is in an oblique sagittal plane rotated around a vertical axis. There is some mild isthmal narrowing but no coarctation. Cross sectional echocardiography did not rule out the presence of coarctation.
Magnetic resonance imaging of coarctation

Fig 2. An oblique sagittal section in a three month old infant with coarctation. There is a discrete shelf across the descending aorta. The length of the aortic isthmus and the relation of the coarctation to the left subclavian artery is clearly seen. There is only very mild isthmal hypoplasia.

Fig 3. An oblique sagittal section in another patient aged 10 months with coarctation. In contrast with the anatomy in figure 2, there is a waist-like narrowing of the descending aorta. The origin of the left subclavian artery arises at the level of the coarctation and so there is no isthmus.

Fig 4. An oblique sagittal section in an infant with a very long, hypoplastic isthmus (between the arrows). The coarctation forms a discrete shelf rather than a waist.

tory. The diagnosis of coarctation was confirmed in 12 infants. It was excluded in seven. In the first case in which it was excluded (patient 4), we lacked experience in interpreting the magnetic resonance images and so the clinical picture prompted investigation by angiography. The aortogram confirmed normal aortic anatomy. In all the subsequent cases the diagnosis based on magnetic resonance imaging was accepted. Angiography was performed in two other negative cases (patients 6 and 10) to define aspects of the cardiac anatomy other than the aortic arch. In one positive case the arrangement of the branches of the aortic arch was unclear. An aortogram was performed before surgical correction to clarify this. The right subclavian artery arose anomalously just proximal to the coarctation and, in retrospect, this could be seen in the coronal images of the coarctation in this patient. In all the remaining positive cases, surgical correction was performed without further investigation.

Surgical correction confirmed the anatomy in all the patients with coarctation. But in two patients small arterial ducts (ductus arteriosus) were found at
operation that were not seen in the magnetic resonance images. In practice this was not of clinical importance. In one patient a large arterial duct was shown by magnetic resonance imaging (case 3).

The presence or absence of obstruction in the thoracic aorta could invariably be shown in the initial transverse series of images. Further oblique sagittal images were, however, obtained in all cases to display the anatomy of the aortic arch clearly. From such images the diameter and length of the aortic isthmus were evident. The coarctation shelf, itself, gave a very intense magnetic resonance signal in these T1 weighted images. Images obtained in the coronal plane were valuable in defining the relation of the coarctation to the vessels arising from the aortic arch, particularly the left subclavian artery.

Discussion

To date magnetic resonance imaging of the heart has generally been performed with systems that use low and medium field strengths, < 0.6 T. It was thought that it would be difficult to achieve cardiac gating in strong magnetic fields such as 1.5 T. This is because the electrocardiographic signal is distorted by the flow of blood in the high magnetic field. But the newer, high field strength imaging systems have a potentially important advantage for the study of congenital heart disease. The signal to noise ratio from these systems is considerably better than from those that use low and medium strength fields and so a more favourable compromise between spatial resolution and imaging time can be achieved. These advantages are especially valuable when infants are studied. A major criticism of published reports is that there is almost no experience in the use of gated magnetic resonance cardiac imaging in infants. Indeed, most workers specifically avoided imaging in this age group. Because almost all patients with congenital heart disease present in the first year of life, the anatomical diagnosis of cardiac defects is now generally performed in infancy. An imaging technique, to be of value, must be capable of producing high quality images in this age group.

As we have shown, magnetic resonance images can define the position of the coarctation in the aortic arch, its relation to the arterial duct and the arteries arising from the arch, the length and diameter of the aortic isthmus, and whether the coarctation is a discrete shelf or a waist lesion. Identification of the coarctation is greatly helped by the intensity of the signal from the shelf itself. In infants, this is usually made up of tissue originating from the arterial duct (ductus arteriosus). We always made certain that the shelf could be seen in images in at least two different planes, thus ensuring it was not an artefact produced by the oblique transection of the aorta. In practice, this does not seem to be a problem. In none of our normal patients were the appearances of the aortic arch ambiguous.

We believe that our experience shows that high field strength magnetic resonance imaging can consistently produce high resolution cardiac images in infants. In this series, we have shown that it is possible, with great clarity, to image the aortic arch. The images were of much higher resolution than those obtained with cross sectional echocardiography. In addition, magnetic resonance imaging is non-invasive and safe. It does not use any ionising radiation. It provides all the anatomical information required for surgical repair of the anomaly. It thus has many advantages for studying this age group. Its clinical role requires further evaluation.

This research was supported by the British Heart Foundation. The imaging system was purchased with the assistance of Sir Philip and Lady Harris and the Special Trustees of Guy’s Hospital.

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Magnetic resonance imaging of coarctation


Magnetic Resonance Imaging of the Brain in Alcoholics: Cerebral Atrophy, Lifetime Alcohol Consumption, and Cognitive Deficits

J. D. Chick, M. A. Smith, H. M. Engleman, D. M. Kean, A. J. Mander, R. H. B. Douglas, and J. J. K. Best

Magnetic resonance imaging of the brain in 69 detoxified alcoholics revealed that relaxation time ($T_1$) in whole brain and in grey matter and parietal white matter was greater than in age-matched controls. In 48 patients, data on cognitive function and lifetime alcohol consumption were available. With age-controlled, lifetime consumption, and impairment in performance in the cognitive test (a Category Sorting Test) correlated positively with $T_1$ in whole brain and in selected regions. Impairment in the cognitive test correlated with $T_1$ in whole brain and white matter independently of cerebral atrophy. Alcohol consumption patterns in the following 8 months were unrelated to changes in $T_1$. The excess water implied by the elevated $T_1$ values may be intra- or extracellular. It is uncertain whether or not $T_1$ elevation in alcoholics is a marker of neuronal damage. $T_1$ elevation appears to be a marker of one type of alcohol-related cognitive impairment.

EXCESSIVE DRINKING is associated with impaired cognition.\(^1\) There is evidence in some drinkers that a deficit in cognition predated the heavy drinking.\(^2\) However, improvements in cognition occur in some alcoholics who become abstinent.\(^1,3\) Also indicators of predrinking intelligence such as early academic attainment and retained verbal skills suggest that the excessive drinking is at least as much a cause of certain types of cognitive impairment as a consequence thereof.\(^1,2\)

Brain shrinkage is also well established,\(^4,5\) and modest correlations between measures of shrinkage (especially central shrinkage) and cognitive impairment have been found.\(^1,3\) The cause of this shrinkage is unknown. However, the cerebellar atrophy of alcoholism is due at least in part to neuronal death.\(^6\) Neuronal death may contribute to the cerebral shrinkage of the alcoholic, a possibility that was raised over 80 years ago.\(^7\) Animal studies have shown that ethanol consumption can cause nerve cell dysfunction and death.\(^8,9,11\)

A potential marker of nerve cell death in alcoholic patients is, of course, the shrinkage visible in computer tomographic (CT) scans. However, a search for another marker seems justified since the correlations between cognitive impairment indices and CT measurements is, as mentioned, modest at best. An alternative marker might be provided by magnetic resonance imaging (MRI) of the brain. Furthermore, the observation that the histology of neuronal death may show osteomatous cells\(^12\) suggests a particular importance for MRI.

In magnetic resonance imaging of living tissue the parameter known as $T_1$, relaxation time is related to the state of water in the tissue.\(^12\) The $T_1$ value, measured in milliseconds, increases as the proportion of free to bound water within the tissue increases. MRI scanning of brain in the intact patient before neurosurgery has shown that in vivo $T_1$ measurements predict the degree of brain hydration.\(^14\) Besson et al.\(^15\) were the first to report brain $T_1$ measurements in a small group of alcoholics but reported only on change in the phase immediately after cessation of alcohol intake.

The present study had four hypotheses: (a) that the detoxified alcoholic brain some 2 weeks after the last drink would be overhydrated compared to age and sex matched non-alcoholics; (b) that in detoxified alcoholics the degree of overhydration in the brain would be related to the total lifetime consumption of alcohol; (c) that this overhydration would predict cognitive impairment; (d) that extended abstinence in the posttreatment phase would be associated with reduction of overhydration and improvement in cognitive functioning.

SUBJECTS AND METHODS

Alcoholic patients admitted for detoxification to the Alcohol Problems Clinic at the Royal Edinburgh Hospital were fully informed about the purpose and procedures of the study and invited to participate. Signed consent was obtained. The project was approved by the appropriate Ethics of Research Committee.

Hypothesis 1

Sixty-nine patients (46 males, 23 females) were scanned at the end of detoxification—a mean interval of 14 days after the last alcoholic drink. These 69 patients had a minimum of 5 years of recent heavy drinking. Their mean age was 44 years (range, 22-70; st. 1.16). None had the clinical features of Wernicke-Korsakoff syndrome, had a history of drug abuse, were clinically malnourished, or had physical disorders other than raised serum liver enzymes. They had all been drinking at least 130 g of ethanol per day during drinking episodes. In Britain, a “standard drink,” i.e., a half pint of beer, a glass of wine, or a single measure of spirits, contains 8.0 g of ethanol. They spanned the range of socioeconomic
backgrounds, and though half were unemployed none were without a fixed address.

Hypotheses 2, 3, and 4

A subgroup of 48 of the 32 males (16 females) was invited to have cognitive testing and to attend for repeat MRI scans and cognitive testing at 6 months, (though in the event only 26 patients kept the following appointments for both scans and testing). These patients were selected on the basis of their willingness and likely ability to attend for further testing. Their mean age was 44 years (range, 22–70; ss, 1.36). They had mean red cell volumes ranging from 88 to 111 fl (mean, 98.9; ss, 0.98; n = 42) and serum y-glutamyl transpeptidase (GGT) values ranging from 5 to 1000 units (median, 100; mean, 217; ss, 42; n = 43). The consumption pattern of this group was studied in more detail. They were asked to give anchor dates throughout their adult lives and commencing with the year in which they first drank alcohol, to reconstruct their typical pattern of drinking month by month during each of the succeeding years. The method is based on that of Skinner and Sheu. Thus an estimate was obtained of cumulative total lifetime alcohol consumption. Inter-rater reliability of this method in 10 patients whose responses were rated by two independent assessors showed an average agreement between assessors of 87%. (Taking the past year’s consumption enabled some of the relationships of interest to be analyzed using a recent, rather than a lifetime consumption measurement. However, to reduce the total number of tests of significance made, it was decided to concentrate on our cumulative lifetime measure of consumption).

Normal MRI values were available from a group (n = 26) recruited from hospital staff and patients’ relatives. Their usual alcohol consumption ranged from nil to 80 g ethanol per week. Their ages ranged from 23 to 74 years with a mean of 42 years, and a similar distribution to that of the patients. The ratio of women to men in the normal sample was 14 males to 12 females, that is, a higher proportion of women than in the patients. However a two-tail t test of the differences in the females and males of the control group found that there were no significant differences in their whole brain T1, measurements (p = 0.88) so the differences in the proportions of the sexes in the control and the alcoholic samples are not considered to be of importance. They may have been of slightly higher mean socioeconomic status than the patients but this was not thought likely to bias the results, though no study of MRI and socioeconomic status in a knowledge has been reported.

Our index of cognitive function was the computer-administered Maudsley Category Sorting Test. This provides a measure of abstract reasoning ability and rigidity of thought processes and it is of interest because rigidity of thinking in alcoholics is a major impediment in the psychological treatment of the condition. The subject is required to deduce the categorization system being used by the computer program and to shift his or her set when the program changes. The program offers six different methods and the number of times out of six in which the subject correctly arrives at the solution is counted (CATEGORIES). The number of trials needed to achieve that number of correct solutions is counted (SORTS). The total number of errors made is summed (ERRORS). The number of perseverative errors (errors repeated despite feedback from the program) is totalled (PERSEVERATIONS). We have concentrated on the Category Sorting Test because it has been shown that deficits in this type of function correlate well with CT parameters of atrophy, and are common to both older and younger alcoholics. A number of tasks on tests of memory and general intellect tend to be seen only in older patients.

Alcohol consumption in the follow-up period was estimated from the clinic treatment records, and interviews with the patient conducted by a research assistant at 3 months and 6 months. This information was corroborated where possible by an additional interview with a relative and by changes in the serum GGT in those patients who had an elevation of this marker at intake to the study. For some patients only a qualitative report on consumption was available. Patients were categorized as follows for each of the 3-month period: total abstinent; "intermediate" drinking (1–40 g per day); "heavy" drinking (over 40 g per day).

MRI measurements were performed using a 0.3 Tesla-resistive MRI system. Three or four transverse sections were obtained through the brain centered to the maximum diameter of the lateral ventricles. The slice thickness was 12 mm and pixel size was 2 x 2 mm. An interleaved saturation recovery and inversion recovery pulse sequence, with a repetition time of 1000 ms for both and a time from inversion of 200 ms for the latter, was used to obtain a calculated T1 map of each section. Measurements of the patient’s brain T1 were made from a section approximately 10 mm above the maximum diameter of the lateral ventricles. The T1 in grey and white matter in the frontal and parietal regions was measured from the T1 map using small regions of interest generated by the computer. Different size regions of interest were used for grey matter (20 mm2) and for white matter (69 mm2) and the mean of similar regions in the left and right hemisphere was noted. Larger irregular regions of interest were also measured to attain mean T1 values of each cerebral hemisphere. To achieve this an electronic cursor was used to delineate the border of each hemisphere directly on the computer-reconstructed image of the brain as displayed on the monitor. Obvious areas of CSF (which have a very high T1, and appear white on the coloured image) were avoided. As a further precaution to exclude CSF the computer was programmed to display the mean T1 for each hemisphere only for values within the range 60–600 ms. An example of a T1 map in a patient including the regions of interest used for analysis is shown in Fig. 1.

The precision of the T1 measurements was estimated from four repeated measurements on three normal volunteers over a period of a month. The T1 precision of white matter was 2.8%, grey matter was 4.9%, and the whole brain 2.6%. The poorer precision of the measurements of grey matter are due to the fact that it is generally of small volume and adjacent to CSF resulting in partial volume errors. In additional smaller regions of interest are used resulting in a larger error. The interoperator variation of our T1 measurements was assessed in two independent operators in 20 sections from five different patients. The percentage differences between raters were: whole brain 1.9%; frontal, occipital, and parietal white matter 2.5%; temporal white matter 5.2%; grey matter (all areas) 5.5%. All ratings were blind to the clinical state of the patient, consumption measures, and cognitive test scores. However, control scans were identified as such, and dates of all scans were also known to the raters. T1 maps were analyzed by three operators; results from the same patient were always analyzed by the same operator.

Atrophy Measurements

Commonly used indices of cerebral atrophy were made by caliper measurement on MRI scans: (a) Third ventricle; widest diameter; (b)
The Ventricular Index, the distance between the choroid plexuses divided by the maximum width between the frontal horns; (c) The Huckmann Index, the sum of the maximum and minimum widths of the frontal horns; and (d) The mean width of four cortical sulci was measured on a slice 16 mm above the upper limit of the lateral ventricles, choosing the two widest sulci on each side.

As an attempt at assessing the reliability of these measures in our hands, we checked the correlations between the indices of internal atrophy. The Huckmann index correlated \( r = 0.33 \) (\( p < 0.05 \)) with the ventricular index and \( r = 0.47 \) (\( p < 0.01 \)) with "3rd ventricle" (\( n = 46 \)).

Differences in \( T_1 \) values within groups of patients at different times were compared using Student's t test. Changes in \( T_1 \) were assessed using a paired Student's t test. The relationship between changes in \( T_1 \) values, drinking and changes in cognitive function were assessed using linear regression analysis. Partial correlation analysis was used to investigate the relationship between cognitive function and \( T_1 \) value and cognitive function and previous drinking history to control for the effect of age and atrophy.

RESULTS

The \( T_1 \) values obtained from the patients at 2 weeks and 6 months are given in Table 1 with the results from normal controls for comparison. The most pronounced differences between normals and alcoholic patients are seen in the whole brain and in grey matter \( T_1 \) values.

Relationship between \( T_1 \), Age, and Alcohol Consumption

In the subgroup studied intensively the mean lifetime drinking total was \( 775 \times 10^3 \) g (se, \( 104 \times 10^3 \); range, \( 71 \times 10^3-3329 \times 10^3 \)). The \( T_1 \) values at the time of the first scan correlated positively with the patient's age, significantly so in frontal white matter (\( r = 0.49 \), \( p < 0.01 \)) and in posterior white matter (\( r = 0.41 \), \( p < 0.05 \)). Eliminating the effect of age using partial correlation, a significant positive correlation was found between whole brain \( T_1 \) in the detoxified patient and the patient's cumulative total lifetime consumption \( r = 0.36 \), (\( p < 0.01 \)). The past year's consumption, which not unexpectedly correlated significantly with lifetime consumption, correlated positively with parietal white matter \( T_1 \) but not significantly with whole brain \( T_1 \). (\( n = 47 \)).

Relationship between \( T_1 \) and Cognitive Impairment

In the alcoholics given cognitive testing, impairment of each of the scores on the Category Sorting Test was related to an increase in \( T_1 \) in all regions of the brain. The findings are shown in Table 2. The effect of age has been controlled for. Correlations of each of the test measures with whole brain \( T_1 \) and with frontal white \( T_1 \) were significant for either the 99% or 95% levels. The relationships remained significant when the sexes were examined separately. Both lifetime consumption and past year's consumption correlated significantly with each of the test measures with age controlled.

Role of Atrophy

Table 3 shows that internal atrophy as measured by the Huckmann index appears to be associated with increases in \( T_1 \) in white matter, and with impairment in the number of categories achieved in the sorting test. However, by chance, in the 36 correlations shown in Table 3, two would be significant at the 5% level.

There is little evidence of a relationship between other measures of atrophy and either \( T_1 \) values or cognitive test impairment. However, the correlation of 0.26 between sulcal width and \( T_1 \) in frontal grey matter is probably meaningful in that when patients are divided about the median for sulcal widths, the \( T_1 \) (frontal grey) is 10 points higher for those with sulcal width above the median than for those below the median (\( t = 3.3, p < 0.01 \)). This supports the possibility that CSF in the subarachnoid space, increased in cortical atrophy, might tend to increase \( T_1 \) measurements in grey matter.

Table 1. Mean ± se (Range) of \( T_1 \) in Whole Brain and Regions of Interest and in the Whole Sample of Alcoholics at a Mean of 15.0 Days after Admission

<table>
<thead>
<tr>
<th></th>
<th>Whole brain</th>
<th>Frontal white</th>
<th>Parietal white</th>
<th>Frontal grey</th>
<th>Parietal grey</th>
<th>Timing (days after admission)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control group (( n = 26 ))</td>
<td>306 ± 1.69</td>
<td>268 ± 1.58</td>
<td>269 ± 1.97</td>
<td>341 ± 2.09</td>
<td>334 ± 1.50</td>
<td>( 15.0 ± 0.06 )</td>
</tr>
<tr>
<td>(220-324)</td>
<td>(252-283)</td>
<td>(248-296)</td>
<td>(326-371)</td>
<td>(310-381)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patients—whole sample (( n = 69 ))</td>
<td>321 ± 1.67*</td>
<td>285 ± 1.49</td>
<td>279 ± 1.45</td>
<td>367 ± 2.28*</td>
<td>360 ± 1.76*</td>
<td>( 15.0 ± 0.06 )</td>
</tr>
<tr>
<td>(223-388)</td>
<td>(252-326)</td>
<td>(250-266)</td>
<td>(314-402)</td>
<td>(321-382)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patients—subgroup with cognitive test (( n = 48 ))</td>
<td>325 ± 1.91*</td>
<td>256 ± 1.98</td>
<td>287 ± 1.11*</td>
<td>373 ± 2.13*</td>
<td>361 ± 2.25</td>
<td>( 15.7 ± 0.08 )</td>
</tr>
<tr>
<td>(504-383)</td>
<td>(230-265)</td>
<td>(250-265)</td>
<td>(315-402)</td>
<td>(326-382)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patients—subgroup attending for follow-up (( n = 26 ))</td>
<td>322 ± 1.74*</td>
<td>266 ± 1.69</td>
<td>276 ± 1.62</td>
<td>372 ± 2.99*</td>
<td>366 ± 2.99*</td>
<td>( 14.0 ± 0.09 )</td>
</tr>
<tr>
<td>First scan</td>
<td>(304-333)</td>
<td>(249-284)</td>
<td>(254-268)</td>
<td>(336-400)</td>
<td>(335-392)</td>
<td>( 14.0 ± 0.09 )</td>
</tr>
<tr>
<td>Follow-up scan</td>
<td>(&lt; 40 ) g ethanol per day</td>
<td>324 ± 1.37*</td>
<td>266 ± 1.85</td>
<td>286 ± 2.71*</td>
<td>373 ± 4.17*</td>
<td>365 ± 2.86*</td>
</tr>
<tr>
<td>(312-333)</td>
<td>(254-285)</td>
<td>(262-301)</td>
<td>(349-402)</td>
<td>(348-390)</td>
<td>( 147-258 )</td>
<td></td>
</tr>
<tr>
<td>( \geq 40 ) g ethanol per day</td>
<td>330 ± 5.35*</td>
<td>270 ± 3.84</td>
<td>277 ± 3.18</td>
<td>374 ± 4.31*</td>
<td>363 ± 4.04</td>
<td>( 147-258 )</td>
</tr>
</tbody>
</table>
| (311-373) | (251-298) | (258-263) | (346-393) | (342-392) | * Patients greater than controls, \( p < 0.01 \); \( \ddag p < 0.05 \).
MAGNETIC RESONANCE IMAGING IN THE BRAIN OF ALCOHOLICS

Table 2. Age Controlled Correlation Coefficients of T
1 Values and Estimated Lifetime Consumption with Scores on Components of the Category Sorting Test in Alcoholics at a Mean of 15.7 Days after Admission (*p<0.01; **p<0.05). (Impairment Results in a Lower Score on Categories, and a Higher Score on Sorts, Errors, and Perseverations.)

<table>
<thead>
<tr>
<th>Category Sorting Test scores [mean ± se (range)]</th>
<th>Whole brain</th>
<th>Frontal white</th>
<th>Parietal white</th>
<th>Frontal grey</th>
<th>Parietal grey</th>
<th>Lifetime drinking total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Categories</td>
<td>5.42 ± 0.16</td>
<td>-0.44*</td>
<td>-0.50*</td>
<td>-0.42*</td>
<td>-0.24*</td>
<td>-0.17</td>
</tr>
<tr>
<td>(37-136)</td>
<td></td>
<td>(37-136)</td>
<td></td>
<td></td>
<td></td>
<td>(37-136)</td>
</tr>
<tr>
<td>Sorts</td>
<td>60.90 ± 0.42</td>
<td>0.23*</td>
<td>0.30*</td>
<td>0.19</td>
<td>0.21</td>
<td>0.14</td>
</tr>
<tr>
<td>(37-136)</td>
<td>(37-136)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(37-136)</td>
</tr>
<tr>
<td>Errors</td>
<td>21.00 ± 3.02</td>
<td>0.37*</td>
<td>0.34*</td>
<td>0.23</td>
<td>0.20</td>
<td>0.40*</td>
</tr>
<tr>
<td>(1-90)</td>
<td></td>
<td>(1-90)</td>
<td></td>
<td></td>
<td></td>
<td>(1-90)</td>
</tr>
<tr>
<td>Perseverations</td>
<td>6.33 ± 1.25</td>
<td>0.41*</td>
<td>0.34*</td>
<td>0.29*</td>
<td>0.21</td>
<td>0.25</td>
</tr>
<tr>
<td>(0-35)</td>
<td>n = 48</td>
<td>n = 48</td>
<td>n = 48</td>
<td>n = 48</td>
<td>n = 48</td>
<td>(0-35)</td>
</tr>
</tbody>
</table>

Changes in T
1, and Cognitive Function in Relation to Degree of Abstinence

There were 26 patients who reattended for the repeat scan and on whom we had data on consumption of alcohol in the interval (Table 1).

Whether or not patients were deemed to have resumed drinking heavily, the trend was for whole brain T
1, to rise slightly over the 6-month period. Whole brain T
1, measurements at 6 months in those 15 patients who had been abstinent or who had drunk on average less than 40 g of ethanol per day had risen by a mean of 2.33 ms (range, -11 to 29; se, 2.87) (see also Fig. 2). The mean serum GGT of this group was within the normal range (21 IU/liter, n = 14). In those 11 patients who had drunk more than 40 g per day on average during the 6-month period the mean rise in whole brain T
1, was 6.27 ms (range, -15 to 60; se, 5.94) (see also Fig. 2). This was not a significantly greater mean rise (p = 0.56) than in those drinking less than 40 g/day. The mean serum GGT of this group was elevated (178 IU/liter, n = 10). The two patients with the highest T
1, values at follow-up were in this group. They were men who had severe long-standing dependence on alcohol for many years and were once again drinking in excess of a bottle of spirits per day (280 g ethanol per day) up to the time of the scan.

Table 4 shows that internal atrophy did not account for the relationship between raised T
1, in white matter and cognitive impairment. Table 2 showed, as already mentioned, correlations in the expected direction though in general not significant between T
1, in grey matter and cognitive impairment. Table 5 shows little alteration in the size of these correlations when cortical atrophy as measured by sulcal width is controlled for. T
1, in whole brain also remains significantly related to cognitive impairment despite the possible confounding effect of atrophy.

The relative lack of importance of atrophy in this study may partly reflect our measurements. We used indices previously used in CT scans, and by comparison MRI scans lack definition. Our measurements were made by caliper rather than by computer.

Table 4. Age-controlled Correlation Coefficients of Atrophy Indices, T
1, Values (Whole Matter and Whole Brain) with Cognitive Test Scores, Controlling for Age and Huckmann Index (n = 48)

<table>
<thead>
<tr>
<th></th>
<th>Whole brain</th>
<th>Frontal white</th>
<th>Parietal white</th>
</tr>
</thead>
<tbody>
<tr>
<td>Categories</td>
<td>-0.42*</td>
<td>-0.55*</td>
<td>-0.44</td>
</tr>
<tr>
<td>Sorts</td>
<td>0.32*</td>
<td>0.40*</td>
<td>0.30*</td>
</tr>
<tr>
<td>Errors</td>
<td>0.36*</td>
<td>0.47*</td>
<td>0.40*</td>
</tr>
<tr>
<td>Perseverations</td>
<td>0.36*</td>
<td>0.46*</td>
<td>0.40*</td>
</tr>
</tbody>
</table>

* p < 0.05; ** p < 0.01.

Table 5. Correlation Coefficients of T
1, Values in Whole Brain and Grey Matter with Cognitive Test Scores, Controlling for Sulcal Width and Age (n = 48)

<table>
<thead>
<tr>
<th></th>
<th>Whole brain</th>
<th>Frontal grey</th>
<th>Parietal grey</th>
</tr>
</thead>
<tbody>
<tr>
<td>Categories</td>
<td>-0.43*</td>
<td>-0.21</td>
<td>-0.20</td>
</tr>
<tr>
<td>Sorts</td>
<td>0.33*</td>
<td>0.20</td>
<td>0.17</td>
</tr>
<tr>
<td>Errors</td>
<td>0.37*</td>
<td>0.21</td>
<td>0.23</td>
</tr>
<tr>
<td>Perseverations</td>
<td>0.40*</td>
<td>0.19</td>
<td>0.27</td>
</tr>
</tbody>
</table>

* p < 0.05; ** p < 0.01.

Changes in T
1, and Cognitive Function in Relation to Degree of Abstinence

There were 26 patients who reattended for the repeat scan and on whom we had data on consumption of alcohol in the interval (Table 1).

Whether or not patients were deemed to have resumed drinking heavily, the trend was for whole brain T
1, to rise slightly over the 6-month period. Whole brain T
1, measurements at 6 months in those 15 patients who had been abstinent or who had drunk on average less than 40 g of ethanol per day had risen by a mean of 2.33 ms (range, -11 to 29; se, 2.87) (see also Fig. 2). The mean serum GGT of this group was within the normal range (21 IU/liter, n = 14). In those 11 patients who had drunk more than 40 g per day on average during the 6-month period the mean rise in whole brain T
1, was 6.27 ms (range, -15 to 60; se, 5.94) (see also Fig. 2). This was not a significantly greater mean rise (p = 0.56) than in those drinking less than 40 g/day. The mean serum GGT of this group was elevated (178 IU/liter, n = 10). The two patients with the highest T
1, values at follow-up were in this group. They were men who had severe long-standing dependence on alcohol for many years and were once again drinking in excess of a bottle of spirits per day (280 g ethanol per day) up to the time of the scan.

There were only nine patients in whom we could be sure that abstinence or near abstinence had been achieved for the whole of the 6-month period. The mean change in whole brain T
1, in these patients was -0.78 ms (range, -11 to 10; se 2.85). Figure 2 illustrates the pattern of whole brain T
1, changes in individual subjects from the initial scan to the 6-month scan showing abstainers, “intermediate” and “heavy” drinkers separately. Analysis of variance to compare the changes in T
1, values in whole brain and regions of interest over the 6-month period across these three groups defined by increasing posttreatment consumption did not reveal a significant trend for any of the T
1, measurements (frontal white, F = 1.85, p = 0.21; frontal grey F = 1.06, p = 0.36; parietal white, F = 1.92, p = 0.17; parietal grey, F = 0.26, p = 0.77; whole brain, F = 0.67, p = 0.52; in each test df = 2/23).

There were 25 patients who repeated the Category Sorting Test in a sober state at the time of the 6-month scan. The mean changes in these individuals' scores were: CAT-
EGORIES, +0.44 (range, −1 to +3; SE, 0.17); SORTS, −5.2 (range, −44 to +18; SE, 3.4); ERRORS, −5.2 (range, −32 to +16; SE, 2.5); PERSEVERATIONS, −0.8 (range, −15 to +7; SE, 1.0). However despite this trend towards amelioration, no significant correlations were found between these changes and changes in T₁.

DISCUSSION

The findings of a raised whole brain T₁ in newly detoxified patients and of a significant relationship between T₁ measurements and both cognitive impairment and the possible cause of that impairment, total lifetime consumption, suggest that T₁ measurements have both potential clinical utility and a neuropathological significance. Cognitive deficits were also linked to increased T₁ in white matter, but not grey matter. The finding that is not corroborative is the lack of difference between controls and patients in frontal white matter T₁, given that cognitive deficits appeared to relate to that measure. Grey matter T₁ measures, while they did distinguish alcoholics from controls did not relate (significantly) to cognitive impairment. This raises the question that our T₁ findings are artefacts due to cerebral atrophy (which would be expected to affect grey rather than white matter T₁ measurements). However, the only T₁ measurement which we made which was significantly related to atrophy was that in frontal grey matter. Thus, while noting that frontal white matter T₁ measurements did not distinguish alcoholics from controls, we feel we have found a relationship between T₁ in whole brain and in white matter that relates to alcohol consumption and to cognitive impairment, and that this is independent of atrophy. We note that Harper et al. have stressed that it is loss of white matter, rather than grey matter, which is most important in alcoholics.

Previous workers have attempted to estimate total lifetime consumption but its reliability remains in some doubt. Although we have little confidence in the absolute accuracy of the estimates, the rank ordering that the method yields is possibly more reliable. This is supported by our finding a correlation of this estimate with T₁ values, and with cognitive test scores (Table 2). Measures of more recent consumption might be thought to be more reliable. However, while more reliable because the memory is fresher, patients' recent consumption often poorly reflects longer standing patterns of drinking: for example, the patient coming to a clinic has often been trying recently to abstain for periods; or, he has presented for treatment because of one or two especially heavy recent binges. Thus in the general working population, where patterns of drinking are fairly stable over time, a marker such as the serum GGT may correlate well with a measure of recent drinking, while in patients admitted to clinics no apparent relationship is found.

In the "Subjects and Methods" section we explained some of the reasons for our choice of the Category Sorting Test. In addition, its microcomputer-assisted administration makes it an inexpensive and convenient test to perform and reduces tester-induced bias; it tests a psychological function of clinical relevance; previous (unpublished) work in some 150 patients had led us to conclude that among a range of tests it was sensitive to increasing severity of alcoholism. It is not a widely used test in the literature, but it is closely based on the better known Wisconsin Card Sorting Test. Category Sorting Tests are believed to reflect frontal lobe functioning. It is notable therefore that the largest correlations between test scores and T₁ in the region of interest measurements were found in frontal tissue. Unfortunately, over a period of 6 months, we believe that a subject who had mastered the test could later recall his method of doing it and so we hesitate to
interpret changes in scores in this test during the follow-up period.

The associations which have been found by other authors between CT changes and cognitive tests suggest that it is central atrophy (i.e., loss of white matter), rather than cortical atrophy (loss of grey matter) which best predicts cognitive impairment, albeit only explaining less than 20% of the variance in cognitive performance overall. Central changes appear to regress less quickly with abstinence than cortical changes. Thus, our failure to find significant correlation between T1 in grey matter and cognitive impairment is consistent with Bergman and also Ron.

Only slightly over half of the patients invited for a 6-month scan attended, despite repeated arrangements being made in some cases. Only very occasionally did the patient give as a reason that he or she found the enclosed space of the scanner unpleasant. Nonattenders were mostly patients who had resumed drinking and wished to break off contact with the clinic. It is difficult to obtain reliable information on drinking in the follow-up phase without, for example, incorporating at the outset specific monitoring by relatives, or urine, blood, or sweat testing at frequent intervals. It would be premature to conclude that our lack of T1 reduction in our small number of supposedly abstinent patients indicates that elevated values are permanent in the established alcoholic. However, other follow-up studies of abstinent alcoholics have failed to show improvements in cognitive function and the classic alcohol-related deficit, Korsakoff's syndrome, is marked by persisting permanent alterations in brain cells and persistent cognitive impairment despite abstinence. On the other hand, cortical atrophy as identified by cerebral tomography sometimes resolves with abstinence.2 Also, cerebral perfusion, diminished in the newly detoxified alcoholic by comparison to controls, appears to improve with abstinence.

A number of neuropathological processes may occur simultaneously in the alcoholic. An increase in the measured T1 is a marker of overhydration or oedema of either intra- or extracellular tissue. It remains speculative whether an elevated T1 is a marker of damaged neurones. Christie et al.28 have shown that frontal and parietal grey and white matter T1 values are not elevated in patients with presenile Alzheimer-type dementia when compared to age-matched controls whereas patients with Korsakoff psychosis, abstinent for 1–14 years, (mean, 5 years), have raised cortical T1 values when compared to age-matched normal controls and to frontal grey and white matter in patients with Alzheimer's disease. Christie et al. were able to distinguish increased T1 measurements from atrophy in this study of dementia, supporting our conclusion in the present study that our findings are independent of atrophy). Measurements of T1, like cortical atrophy, increase with age as also found in a previous study as well as the present study.

In conclusion we believe we have shown that MRI T1 measurements, especially in white matter, are elevated in alcoholics proportionate to their cumulative alcohol intake and to the degree of impairment on a Category Sorting Test of cognitive function. These findings are analogous to those of ageing. The precise location of the extra water which is presumed to underly this change, and its cause, is unknown. Potentially, MRI T1 is of clinical value as a marker of alcohol-related brain damage.

ACKNOWLEDGMENTS

We are grateful to the staff of the Alcohol Problems Clinic for their assistance; to Iris Cansdale without whose perseverance and skill the study would not have been possible; to Colette Rowan and Chris Stirling who helped prepare the manuscript; to Professor R. E. Kendell; to the Medical Research Council and the Alcohol Education and Research Council for funding; to colleagues at the Department of Neuropathology, Institute of Psychiatry, London University; and above all to our subjects for their generous cooperation.

REFERENCES


Phospholipids are visible in $^{31}$P n.m.r. spectra of human breast tumours

MARTIN LOWRY,* DAVID A. PORTER,* CHRISTOPHER J. TWELVES,* PHILIPPA E. HEASLEY,* PAMELA B. GARLICK,* MICHAEL A. SMITH,† ROBERT D. RUBENS,* MICHAEL N. MAISEY,* and MICHAEL A. RICHARDS*†

*L.C.R.F. Clinical Oncology Unit and †Division of Radiological Sciences, Guy's Hospital, London SE1 9RT, U.K.

$^{31}$P magnetic resonance spectroscopy provides a non-invasive method for studying human tumour biochemistry. In vivo, spectral resolution is, however, inferior to that achievable in vitro. Furthermore, analysis in vitro may assist in determining the chemical composition of the peaks observed in vivo. In particular, the peaks commonly known as phosphomonoesters (PME) and phosphodiesters (PDE) may arise from several different compounds. As a preliminary to studies of breast cancer patients, we have characterized the spectra obtained both in vivo and in vitro from this tumour type.

Tumours (>6 cm) from 15 patients with breast cancer were studied by spectroscopy in vivo alone, in vitro only and both in vivo and in vitro. Spectra were obtained in vivo with a Philips Gyroscan 1.5 T system using a 5 cm surface coil for volume localization. In 11 patients a biopsy was obtained and rapidly frozen in liquid nitrogen. The specimen was divided and two extracts prepared by homogenization in 6% (v/v) perchloric acid (PCA) and chloroform/methanol (CM), respectively. Spectra from these extracts were obtained with a 9.4 T Bruker AM400 spectrometer.

The spectra collected from PCA extracts of breast tumour specimens consisted of approximately 12 peaks each of which was identified and quantified. These results are shown in Table 1. P1 was present in high concentrations while phosphocreatine (PCr) was present in only 6 of the 11 tumours at very low concentrations. Other compounds identified include phosphoethanolamine,-phosphocholine, glycerophosphoethanolamine (GPE), glycero-phosphocholine (GPC) and nucleotide di- and tri-phosphates. Similar observations have been made on several other tumour types [1, 2, 3].

Comparison of spectra obtained in vivo with corresponding spectral regions in the PCA-extract spectra revealed the following major differences. The PCA-extract spectra had greater relative proportions of P1 and PME and smaller PDE than the spectra in vivo. In contrast the CM extract showed a large peak in the PDE position ($5340 \pm 1235$ nmol/g wet wt. of tissue), i.e. about 20-fold greater than the combined concentrations of GPE and GPC. The differences in P1 and PME observed between in vivo and in the PCA-extract spectra can be explained by saturation effects and n.m.r.-invisible pools. The low PDE in PCA-extract spectra cannot be explained in this way and suggests that some n.m.r.-visible phosphorus-containing material has been lost during the extraction procedure. As PCA only extracts the water-soluble metabolites from tissue, one candidate for the missing PDE is phospholipids in cell membranes. The results imply that the PDE peak observed in vivo arises primarily from membrane phospholipids. However, phospholipids are generally considered to be n.m.r. invisible owing to their low mobility giving rise to very broad signals. The solution to this apparent paradox lies in the fact that the line-width of the peak is proportional to the magnetic field strength [4]. At the low field strengths used for human spectroscopy in vivo, the line-width is small enough for the phospholipid peak to make a significant contribution to the spectrum.

The results of these study examine have major implications with respect to the interpretation of spectra collected from patients in vivo. The finding that the PDE peak is pri-

---

Table 1. Concentrations of phosphorus-containing metabolites in human breast tumours

<table>
<thead>
<tr>
<th>Peak</th>
<th>Component metabolites</th>
<th>Concentration (nmol/g wet wt.)</th>
<th>Contribution to total area (%)</th>
<th>Extracts</th>
<th>in vivo</th>
</tr>
</thead>
<tbody>
<tr>
<td>PME</td>
<td>Phosphoethanolamine</td>
<td>$1295 \pm 491$</td>
<td>$29 \pm 9$</td>
<td>$14 \pm 8$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Phosphocholine</td>
<td>$554 \pm 478$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P1</td>
<td>Inorganic phosphate</td>
<td>$2543 \pm 1871$</td>
<td>$36 \pm 17$</td>
<td>$10 \pm 3$</td>
<td></td>
</tr>
<tr>
<td>PDE</td>
<td>Glycerophosphoethanolamine</td>
<td>$125 \pm 80$</td>
<td>$5 \pm 2$</td>
<td>$33 \pm 8$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Glycerophosphocholine</td>
<td>$160 \pm 65$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PCr</td>
<td>Phosphocreatine</td>
<td>$47 \pm 57$</td>
<td>$1 \pm 1$</td>
<td>$3 \pm 6$</td>
<td></td>
</tr>
<tr>
<td>$\gamma$-ATP</td>
<td>$[\gamma-P]$Nucleotide triphosphate</td>
<td>$467 \pm 230$</td>
<td>$11 \pm 4$</td>
<td>$16 \pm 6$</td>
<td></td>
</tr>
<tr>
<td>$\alpha$-ATP</td>
<td>$[\alpha-P]$Nucleotide diphosphate</td>
<td>$223 \pm 83$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\beta$-ATP</td>
<td>$[\beta-P]$Nucleotide diphosphate</td>
<td>$188 \pm 72$</td>
<td>$13 \pm 4$</td>
<td>$18 \pm 3$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$[\alpha-P]$Nucleotide triphosphate</td>
<td>$430 \pm 212$</td>
<td>$6 \pm 2$</td>
<td>$11 \pm 5$</td>
<td></td>
</tr>
</tbody>
</table>

Vol. 17
Molecular graphic displays of antibody–antigen interactions

MALCOLM M. FRASER and E. JAMES MILNER-WHITE
Department of Biochemistry, University of Glasgow, Glasgow G12 8QQ, U.K.

The three-dimensional structure of the complex between chicken egg white lysozyme and the mouse antibody HyHEL-5 has been determined by X-ray crystallography at 2.5 Å resolution [1]. In this communication, we present photographs of models of the complex displayed on Gemini Challenger graphics workstations. The displays make use of new software based on the CHEMMOD molecular graphics package (U-Microcomputers Ltd, Calver Rd., Warrington, Cheshire WA2 8RF, U.K.) designed for investigating interfaces between macromolecules.

![Image of antibody-antigen interaction](image-url)

Fig. 1. The interface between the antibody (two variable domains, VL and VH) and antigen

Residues that lie within 13 Å of any residue in the other interface are drawn as pale spheres, shaded according to depth. The program originally selects a view along the line joining their centres of gravity. To view the two interacting interfaces simultaneously, the lysozyme molecule is rotated by 180° so that it is placed next to the antibody. Geometrically this is like opening a book. In the original view the interacting interfaces correspond to middle pages of the book. A simultaneous view of both interfaces is produced by opening the book.

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