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medicalprotection.org
Dr Nick Clements
EDITOR-IN-CHIEF

This edition of Casebook is one of welcomes and farewells. Dr Pardeep Sandhu is the new executive director of your professional services division, where he will be responsible for maintaining and building on the quality of the advice and support that is available to you.

The appointment is a considerable boost to our aim of providing you with world class service. You can read more about Dr Sandhu on page 5, but in summary Dr Sandhu brings with him many years’ experience of working within diverse healthcare environments around the world, and he has also worked extensively with governments to advise on health policy and clinical governance – something that is becoming increasingly important to Medical Protection as we seek to shape the landscape in many countries in which we have members.

This edition of Casebook contains, as ever, our latest collection of case reports. Along with the usual salient learning points – and in this edition there is a general theme on the value of good record-keeping – you will also be interested to note some successful defences. As well as demonstrating the value of the Medical Protection legal expertise available to you, these cases also show how the clinicians involved were able to help their own position, be it through excellent documentation, a robust consent process or an articulate presentation of evidence at trial.

I mentioned at the beginning of this editorial that this edition of Casebook was one of welcomes and farewells. This is my last edition as editor-in-chief of Casebook, as I am moving into a new role within Medical Protection. I have greatly enjoyed my time in the position, especially as it has given me so many opportunities to hear your feedback directly.

I am happy to announce that Dr Marika Davies will be taking on the role, please do get in touch with any comments or suggestions that you wish Dr Davies to take on board.

Dr Nick Clements
Casebook editor-in-chief

Dr Pardeep Sandhu joins us from Aetna International, a global health benefits provider in the USA, where he was medical director and head of business development.

Dr Sandhu spent more than seven years working with governments to create and expand robust healthcare systems. In this international role, Dr Sandhu worked across health policy, clinical governance, business development and strategy, as well as designing and launching Aetna’s international care management programmes in multiple geographies.

Dr Sandhu trained at the University College London and was a GP before serving as a clinical adviser to the UK Department of Health. He also holds a MBA from Kellogg School of Management, Northwestern University, USA.

Simon Kayll, Chief Executive, said: “We are delighted to welcome Dr Pardeep Sandhu.

“With numerous challenges facing the medical and dental professions worldwide, it is vital that we are there for members in the right place, at the right time. As a former practising physician myself, I understand the unique dilemmas clinicians face on a daily basis – and I very much subscribe to the Medical Protection ethos that prevention is better than cure. Ensuring the expertise of my team benefits our membership is a key goal for me.

“Of particular interest to me is the challenge of meeting the needs of our members around the world. With so much variation from country to country, it is imperative that we tailor our services to meet everyone’s requirements as fully as possible. I look forward to working with you and hearing your views on how we can improve even further.”

Dr Sandhu said: “I am very excited to be joining a team of such talented individuals, and look forward to building on their established expertise to deliver an even better service to our members.”

Dr Sandhu’s appointment is one of welcomes and farewells. Dr Pardeep Sandhu is the new executive director of your professional services division. Find out what Dr Sandhu brings to the role and how he plans to further improve your support service.
BE TRUTHFUL WHEN ADVERTISING, SAYS MEDICAL COUNCIL

The New Zealand Medical Council has warned doctors to be honest and balanced when advertising products and services to patients.

In its updated Statement on advertising, published earlier in 2015, the Medical Council outlines a range of guidelines that must be followed in relation to use of titles, discounts and the use of ‘before and after’ photos.

Chairman Andrew Connolly said: “Advertising has a role to play in keeping patients informed, but it also has the potential to mislead.

“Misleading advertising coupled with a lack of consumer knowledge can lead to patients being exploited, medical services being used inappropriately or unnecessarily, and patient harm, or unrealistic expectations.

“Our revised Statement has the sole objective of protecting patients and clearly sets out our expectations of the profession.”

ADVERTISEMENTS

Advertisements must contain truthful and balanced representations. When you choose to make a claim or include scientific information in advertising, it should:

• be valid, evidence based and substantiated
• be readily understood by the audience to whom it is directed
• be from a reputable and verifiable source
• identify clearly the relevant researchers, sponsors and the publication where the results on which any scientific evidence or claims are based appear. (Para 11)

THE USE OF ‘BEFORE AND AFTER’ PHOTOS

Before and after photos:

• Are there solely for the purpose of providing accurate and useful information to patients.
• Show a realistic portrayal of the outcome that can reasonably and typically be expected.
• Only depict patients who have undergone the advertised procedure while under your (or your services’) care.
• Have not been altered in any way.
• Use the same lighting, contrast, background, framing, camera angle, exposure and other photographic techniques in both the ‘before’ and ‘after’ images.
• Ensure consistency in posture, clothing and make-up.
• Are only used when the patient has given his or her fully informed consent. (Para 14)

USE OF TITLES

Mr Connolly said: “The use of titles can be useful in terms of providing patients with information about a doctor’s expertise and experience.

“However, some titles can mislead patients into believing that a doctor is more qualified or experienced than a colleague with the same background and training.

“In regulating the use of titles, the Council’s aim has been to ensure that these provide patients with the clearest and most accurate possible guidance about a doctor’s expertise.”

You must advertise only those titles, qualifications or memberships that have been:

• approved for the purposes of registration and relate to your vocational scope of practice
• conferred or approved by your College, or another training organisation that has been accredited by the Council, or another New Zealand responsible authority. (Para 16)

GIFT CERTIFICATES AND DISCOUNT COUPONS

If you choose to advertise by means of discount coupons or gift certificates, you must ensure that these do not undermine your relationship with the patient and the informed consent process. In particular, you must ensure that your coupon or certificate is clear that:

• purchase of the certificate or coupon does not equate to granting informed consent
• prior to treatment you will discuss treatment options with the patient
• the patient has the right to opt out of treatment at any time
• you will not provide the requested treatment if your assessment indicates that the patient is not a suitable candidate
• you will only use a title with the understanding that you are professionally accountable for the training, ongoing Professional Development and recertification in that area. (Para 19)

Mr Connolly added: “Council has also agreed that it is not appropriate to offer medical treatments as prizes or gifts where this is done to promote a commercial service or for financial gain.”

HOW TO HANDLE COMPLAINTS

Complaints are stressful and time-consuming; often a prompt, well-balanced response to a complaint will be enough to defuse the situation.

This article provides best practice advice on complaints handling.
WHAT TO DO
• Acknowledge the complaint within five working days; offer to discuss with the complainant how the complaint will be handled.
• Undertake your investigation into the complaint.
• Draw up a written response to the complaint.
• You should respond as soon as practicable. At ten working days following acknowledgement of the complaint, you should either have responded, or consider how much more time you will require. If the additional time required is 20 working days or more then you must notify the complainant of the reasons for this.

WRITTEN RESPONSES
• Be mindful when preparing your response that it may be read by more than the complainant, for example passed on to authorities such as the Health and Disability Commissioner.
• Include a sympathetic opening paragraph, placing the complaint in context.
• Include an apology and acknowledgement of distress (condolences) if appropriate.
• Explain how the matter has been investigated and summarise the issues raised in the complaint.
• Make sure you include a clear chronological account of the events in question, with an explanation of what happened and why.
• Answer all the questions raised in the complaint or explain why you cannot answer a point.
• Draw conclusions and advise of any improvements or changes in practice that are required to retain certain complaints, thus ensuring that the matter is recorded in writing and agreed with the complainant. Check who is making the complaint – if it is not the patient, make sure you have consent to contain the patient’s health information in your response, or that consent is not required in the circumstances. Aim to provide a co-ordinated response in multi-doctor/multi-agency complaints.

STORAGE OF PATIENT COMPLAINTS
Where patient complaints are stored will depend on the subject of the complaint, and the extent to which it relates to the health services that have been provided to an individual.

MANAGING HEALTH INFORMATION
The Health Information Privacy Code (HIPPC) sets out a number of rules concerning the management of health information. The definition of health information is wide (including any information about health services that have been provided to the individual). It is arguable that health practitioners are not required to retain certain complaints, such as “the magazines in the waiting room are old” – where the link to the provision of health services is tenuous. The majority of complaints are likely to relate to the provision of health services and therefore be considered health information. Therefore, the information contained in the complaint must be managed in the same way as other collected health information.

DISCLOSING INFORMATION AROUND COMPLAINTS AND STORAGE
Any disclosure of the information contained in a complaint (by virtue of where it is recorded on file) needs to be limited to what is absolutely necessary. This would mean dealing with each complaint on a case-by-case basis, considering the purpose for which the information relating to the complaint was collected, rather than a blanket rule for recording complaints.

For example, if the subject of the complaint is about how care was delivered, eg, the doctor was rude to me, then retaining the complaint in the patient notes (for other doctors to view) would be an unnecessary disclosure and inconsistent with the purpose of collection.

HOW WE CAN HELP
• Copies of all the relevant complaint documentation to date
• Any relevant background information, including the dates on which you interacted with the patient/s (if relevant)
• A draft of your response to the current complaint
• Details of where and how you would like us to reply (including telephone/fax numbers, email addresses etc)

NOTICE
Medical Protection assists members in responding appropriately to a complaint with the aim of resolving the matter quickly, effectively and at the lowest level possible.

Our experienced team of advisers can advise on how to handle a difficult complaint and/or review your draft written response. To speed up our advice to you, it would be useful to send the following information to us:

COMMENTS
Be mindful when preparing your response that it may be read by more than the complainant.

It would, however, be acceptable if the practice manager had access to this information, so as to not book a patient in with that particular doctor in future. In that case, the complaint should be stored in a separate folder but with a reference (or ‘red flag’) on the patient’s file that such a folder exists.

If a patient complained about an adverse reaction to treatment, or the method in which a particular doctor applied a treatment (not voiced during a consultation), then it would be consistent with the purpose of collection to record the medical content of the complaint on the patient’s notes, so other doctors within the practice could avoid repeating similar approaches or treatment. As above, the actual complaint should not be stored within the clinical notes, but separately.

REFERENCE
1. HIPPC clause 4(1)(a).

Any disclosure of the information contained in a complaint (by virtue of where it is recorded on file) needs to be limited to what is absolutely necessary.
As a Medical Protection member, you have access to a wide range of specialist support and advice. Based in our offices in Auckland and Wellington, the Medical Protection team has extensive experience – find out how they are working hard on your behalf.

Dr. Tim Cookson

I have worked as a GP partner since 1987 at City GPs and have also taken on other roles over the years. I spent six years as foundation director and complaints officer for the Wellington Accident and Medical Centre, eight years as foundation director for Matpro (providers of primary maternity care for Wellington) and I am a member of the NZ guidelines development group, which has me involved in a number of national guidelines, including “Dyspepsia, Stroke and Vaginal Birth after Caesarean Section”. I have also served as a senior clinical lecturer at the Wellington School of Medicine for the last 15 years.

I became interested in the complaints process when working at the Wellington AMC and, through this, was invited to apply for the Medical Protection position. I have now spent ten years here. The best parts of the job are being able to assist members through what can be a very stressful process, and dealing with a variety of issues that come across the desk.

Dr. Mark Burns

I am an Auckland medical graduate and trained locally in psychiatry. I obtained fellowship with RANZCP in 2001. I have predominantly worked with young adults in early intervention in psychosis for the last 15 years but I also have experience working in a range of DHBs in general adult community psychiatry, currently in metropolitan Auckland.

I began my legal training because of an interest in the overlap between psychiatry and the law. My part-time studies, however, broadened my interest to a whole range of areas of law, particularly healthcare law and human rights law.

I enjoy the daily interface between law and medicine that Medical Protection brings. In addition, revisiting my knowledge of specialties long forgotten when I talk with colleagues outside psychiatry has been curiously refreshing.

Dr. Samantha King

I am an Auckland born and bred. I graduated from Otago University and have worked as General Practitioner since 1993. I hold a Diploma in Obstetrics and am a Fellow of the Royal College of General Practitioners. I am currently writing my dissertation for a Masters of Healthcare Law and Ethics through the University of Dundee.

Most of my work as a GP has been in South Auckland. I currently work part-time as an associate in a practice in Papatoetoe, where I have been for eight years. I enjoy the cultural diversity of this region and the mix of different socio-economic classes that I meet. I have also worked in the Urgent Care setting on and off for the past 25 years in East Auckland.

I find law fascinating and how this applies to medicine. The medical legal system in New Zealand is very foreign to most doctors. The best part of my work with Medical Protection is being able to support colleagues through the often very stressful process they find themselves in. I also enjoy the teaching side of this role where we get to interact directly with our colleagues.

Dr. Andrew Stacey

I am an Urgent Care Physician. I obtained Fellowship of the Royal New Zealand College of Urgent Care in 2009 and sit on the College’s Executive Committee. I am also a Fellow of the Australasian College of Legal Medicine and an enrolled barrister and solicitor.

I have had a long standing interest in the law, which I developed through studying for a law degree and being admitted to the bar. Medical Protection provided the opportunity to work in the niche area of health law, and to maintain my clinical practice at the same time. The best part of the job is the variety – no two days are the same. I enjoy approaching medicine from another angle and assisting colleagues through difficult times.

Dr. Lucy Gibberd

I trained in the UK where I did undergraduate law papers in the middle of my medical degree. I ended up in Taranaki 23 years ago and have stayed ever since, except for a year back in the UK to do GP training.

I have been a GP partner in a practice in New Plymouth for the last 13 years. I also worked as a medical educator for RNP2GP, running the Taranaki seminar programme from 2007-2014.

I wanted to be an advisor for Medical Protection because I enjoy a new challenge and had always wanted to do medicolegal work. The best part of the job so far is the very supportive work environment and nice people to work with. I enjoy the on-call aspect, hearing members’ issues and giving advice. It is great feeling that Medical Protection can help and that members feel better after talking to us.

Dr. Zarko Kamenica

I am a consultant psychiatrist and previously held roles as the clinical director of Waikarapa DHB Mental Health Services and director of Aotearoa Mental Health Services.

I was interested in working for Medical Protection as I have always been attracted to the legal aspects of practising clinical medicine. I get great satisfaction from being able to minimise, or even avoid, professional and legal consequences for colleagues. I also enjoy the constant intellectual challenge and exposure to the whole of medicine.
FEATURE

LEGAL SERVICES
On all issues requiring external legal assistance, members have access to global law firm DLA Piper’s national healthcare team. They are our nominated legal advisers in New Zealand. DLA Piper have seven dedicated lawyers, all of whom work closely with and are supported by a very experienced barrister panel, which includes Harry Waalkens QC and Catherine Garvey, experienced barrister panel, which includes Harry Waalkens QC and Catherine Garvey, based at Quay Chambers in Auckland and Matthew McClendon QC and Jenny Gibson, based at Harbour Chambers in Wellington.

DID YOU KNOW...

• Medical Protection supports more than 17,000 health professionals across New Zealand
• Over 90% of members in New Zealand would recommend Medical Protection to a colleague (Medical Protection member survey, conducted in 2014)
• In addition to assisting with patient complaints and regulatory proceedings, Medical Protection also offers assistance with ACC inquiries

EDUCATION
Medical Protection members in New Zealand have access to a range of free education opportunities including risk management workshops, which have been developed by our Asia-Pacific Educational Services team. These workshops are presented to members by practising New Zealand clinicians on a regular basis, in locations throughout the country.

Our team of advisers is also available for lectures, presentations and medicolegal talks – members can request a speaker by contacting Medical Protection.

THE WHO CHECKLIST
The World Health Organisation (WHO) has created a checklist to help prevent avoidable complications in surgery. Medical Protection provides a comprehensive education programme to suit the needs of all medical and healthcare professionals.

The Educational Services department includes the Cognitive Institute, a wholly owned subsidiary of Medical Protection, which is committed to providing education that distils complex issues and challenges into relevant practical training. The Cognitive Institute partners with healthcare organisations to provide education that helps clinicians meet the challenges of modern practice. Last year 16,000 clinicians attended a Cognitive Institute designed workshop.

Throat packs are commonly used in oral and maxillofacial surgery for a number of purposes, including the prevention of unwanted material from entering a patient’s oesophagus or trachea. The packs themselves, however, are capable of causing serious injury by obstructing patients’ airways if they are not removed after surgery.

The WHO Surgical Safety Checklist was launched in 2009 to improve teamwork and thus combat avoidable complications in surgery, such as retained swabs and instruments. Two recent Medical Protection cases, however, demonstrate that the problem of retained throat packs persists, notwithstanding the introduction of the WHO Checklist.

CASE 1: MRS A
Mrs A opted to undergo facelift surgery. Dr B was the consultant anaesthetist for the procedure and used a throat pack in order to stabilise Mrs A’s airway.

The WHO Checklist Sign-in was performed and the surgery proceeded uneventfully. However, the WHO Checklist Sign-out did not take place. Dr B reversed muscle paralysis, applied suction to the airway and extubated Mrs A. Dr B would usually perform a laryngoscopy at this point but did not on this occasion, as it was difficult to open the patient’s mouth.

Mrs A was handed over to the recovery staff, where slightly obstructed respiratory movements were noted. Dr B attributed these symptoms to emergence delirium, and therefore inserted a nasopharyngeal airway. On examination around 20 minutes later, Mrs A was awake, the artificial airway had been removed and she indicated to Dr B that she was not in any discomfort.

Around three further hours passed before the throat pack was discovered, during which time she experienced significant respiratory distress. The throat pack was removed and Mrs A made a full recovery.

CASE 2: MISS C
Miss C was admitted to hospital for the routine excision of a benign palatal lump. Dr D inserted the throat pack for the first time she experienced significant respiratory distress. The throat pack was removed and Mrs A made a full recovery.

The surgery proceeded uneventfully. However, immediately after waking up, Miss C experienced some difficulty breathing. The issue of the throat pack was raised by nursing staff and Dr D mistakenly asserted that it had already been removed. The nursing staff therefore removed the sticker that had been placed on Miss C’s head. A laryngeal mask airway (LMA) was inserted, which improved Miss C’s oxygen saturation levels.

On removal of the LMA around 15 minutes later, Miss C coughed up the throat pack. She also made a full recovery.

T • The anaesthetist must be present for all three phases of the list
• All three phases of the list must be signed
• The anaesthetist must be present for all three phases. Best practice is to have all members of the surgical team present for all three phases, although the WHO advises that the Sign-in may take place without the surgeon.

At Sign-in, responsibility for both insertion and removal of throat packs must be assigned.

At Sign-out, removal of the throat pack must be checked, either as part of the swab count exercise, or as a distinct part of the checklist.

The surgical procedure proceeded uneventfully.

Miss C was admitted to hospital for the routine excision of a benign palatal lump. Dr D inserted the throat pack, he was not under the impression that its removal was his responsibility.

Further, this throat pack had been obtained from the anaesthetic room, and as such did not form part of the scrub nurse’s swab count. Dr D did, however, place a sticker on Miss C’s head notifying that a throat pack had been used.

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MEDICAL PROTECTION

Medical Protection provides protection for doctors in all stages of their career. The organisation has a dedicated team of legal, risk management and educational specialists who can provide you with advice on any medical matter.

We have a team of specialist medi-cal legal advisers, who work with members to provide the best possible advice in relation to legal and medicolegal issues. They also provide support and advice to the Educational Services team on medical matters.

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THROAT PACKS

Medicolegal advisers Dr Helen Hartley and Professor Carol Seymour examine two recent Medical Protection cases, which demonstrate that the risk of retained throat packs has survived the introduction of the WHO checklist.

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The WHO Surgical Safety Checklist was launched in 2009 to improve teamwork and thus combat avoidable complications in surgery, such as retained swabs and instruments. Two recent Medical Protection cases, however, demonstrate that the problem of retained throat packs persists, notwithstanding the introduction of the WHO Checklist.
Dr Richard Stacey, senior medicolegal adviser, introduces this edition’s collection of case reports and reminds readers of the importance of good note-keeping.

Before joining Medical Protection in 2003, I was a GP and always enjoyed reading the cases in Casebook, irrespective of whether they related to primary or secondary care cases. In my role at MPS I meet many doctors from different specialties and when I introduce myself, invariably the first thing they say is that they enjoy reading the cases in Casebook – with the caveat that it often causes them to reflect on their own practice (which, of course, is one of the reasons why the particular cases are chosen).

In this edition of Casebook there is the usual array of thought-provoking cases, with varying outcomes and learning points. A common issue is that of record-keeping; in the case “Poor notes, fatal consequences”, Dr A is criticised for not documenting a thorough history or the fact that Mrs Y was reluctant to be admitted to hospital; and in the case “Elbow arthroscopy – radial nerve injury”, the operation note was not deemed to be of an acceptable standard. Conversely, in the case “Alleged anticoagulation failure”, the fact that the consultant cardiologist had specifically stated that anticoagulation was not indicated on the advice slip to Dr B was an important feature in defending the claim.

There is a real tension in the context of a busy surgery or outpatient clinic, and other clinical settings, in that patients can perceive that the making of records intrudes into the consultation – yet the records provide the basis of your defence in the event of an adverse outcome. I have often heard it said by patients “the doctor did not pay attention to me as they were far too busy tapping into their computer”. The likelihood is that, in fact, the doctor was making a thorough contemporaneous record, hence there is a real art to being able to take thorough and contemporaneous notes without appearing to disengage from the consultation (or without missing what could be very important non-verbal clues).

What’s it worth?

Since precise settlement figures can be affected by issues that are not directly relevant to the learning points of the case (such as the claimant’s job or the number of children they have) this figure can sometimes be misleading. For case reports in Casebook, we simply give a broad indication of the settlement figure, based on the following scale:

- HIGH NZ$1,000,000+
- SUBSTANTIAL NZ$100,000+
- MODERATE NZ$10,000+
- LOW NZ$1,000+
- NEGLIGIBLE <NZ$1,000

I hope that you find the cases thought-provoking and that they provide you with an opportunity to reflect (amongst other things) on your approach to record-keeping.
M rs S was a 51-year-old teacher. At the start of term Mrs S developed a troublesome cough and went to see her GP, Dr B, about it. Dr B diagnosed a chest infection and prescribed antibiotics but also noted that she had an irregular pulse. An ECG was performed at the surgery the same day, which showed that Mrs S was in atrial fibrillation. Dr B sent Mrs S to the medical assessment unit for urgent review.

The hospital doctors confirmed the diagnosis of atrial fibrillation and prescribed warfarin to reduce her risk of thromboembolic stroke and bisoprolol to slow her heart rate. They put Mrs S on the waiting list for a cardioversion procedure and discharged her home.

Mrs S attended for her cardioversion procedure but was found to be in sinus rhythm. The cardiologist (Dr T) advised Mrs S to stop taking her warfarin and to reduce her bisoprolol dose further. Dr B arranged a further ECG for the following week and reduced her bisoprolol dose further. Dr B documented that Mrs S was “awaiting cardiology follow-up” and that she had had a chest infection when the atrial fibrillation was initially diagnosed.

Dr B saw Mrs S again with the cardiologist’s advice slip. Dr B documented that her pulse was regular now (although she was slightly bradycardic). Dr B arranged a further ECG for the following week and reduced her bisoprolol dose further. Dr B documented that Mrs S was “awaiting cardiology follow-up” and that she had had a chest infection when the atrial fibrillation was initially diagnosed.

Dr B circled the response “no longer requires anticoagulation”.

A month later, Mrs S suffered a stroke. There were no other risk factors for stroke identified other than atrial fibrillation, thus the likely cause of Mrs S’s stroke was an embolic event arising as a consequence of thrombus formation within the atrium.

As a result of the stroke, Mrs S felt unsteady and hesitant every time she walked. Despite rehabilitation, her writing was slow and clumsy and she slurred her words. Sadly, teaching was no longer possible and Mrs S had to retire early on grounds of ill health.

Mrs S was devastated. She felt that her stroke could have been prevented if she had been anticoagulated. Mrs S made a claim in negligence against Dr B. It was alleged that Dr B should have prescribed some form of anticoagulation and that he should have contacted the hospital to query the medication position, especially in light of the non-attendance letter from the anticoagulation clinic.

Medical Protection served a letter of response denying liability and Mrs S did not pursue the claim any further.

**Learning points**

- **NICE, Atrial fibrillation: the management of atrial fibrillation (June 2014)** state that doctors should consider anticoagulation for men with a CHA2DS2-VASc score of 1 and to offer anticoagulation to people with a CHA2DS2-VASc score of 2 or above, taking bleeding risk into account.

- Documentation of the reasons behind the decision-making was invaluable in defending this case.

**Medical Protection**

- The likely cause of Mrs S’s stroke was thromboembolic. Professor G pointed out that some patients develop atrial fibrillation secondary to other illness such as chest disease. In such a setting, if the atrial fibrillation resolves when the underlying cause has been treated, and the clinician feels that there is a low risk of it recurring, then it is reasonable not to anticoagulate. Mrs S would have had a CHA2DS2-VASc score of 1 because of her sex but an absence of congestive heart failure, hypertension, diabetes, stroke or vascular disease and age below 75 years. Professor G felt that it would have been quite reasonable not to anticoagulate in this context.

**EXPERT OPINION**

Medical Protection sought the advice of an expert GP, Dr H. Dr H felt that the care given by Dr B was of a reasonable standard. Dr H did not consider that Dr B had a mandatory duty to prescribe anticoagulation or that he should have contacted the hospital to query the medication position. Dr H noted that the decision to stop anticoagulation had been clearly relayed on an advice slip from a cardiologist. Mrs S had also told Dr B that she was waiting for cardiology review and her subsequent ECG had shown sinus rhythm.

The opinion of a professor in stroke medicine (Professor G) was also obtained by Medical Protection. Professor G confirmed that the cause of Mrs S’s stroke was probably an embolic event arising as a consequence of thrombus formation within the atrium.
**Miss F** was an 18-year-old university student, had been taking the combined oral contraceptive pill microgynon for 18 months for dysmenorrhea. When she presented to her GP, Dr K, worried about acne on her back, Miss F had heard from her flatmate that dienette is a better pill to take for acne than microgynon and wanted to give it a try. Dr K recorded that Miss F was a non-smoker with a normal BMI and BP and switched her pill to dienette, advising her to start it when her microgynon cycle finished in another fortnight.

Two weeks after commencing the dienette, Miss F was rushed into hospital with sudden onset chest pain and shortness of breath. Miss F was diagnosed with a pulmonary embolism and went on to have a cardiac arrest in the emergency department. Dr K was thrombolysed, which resulted in return of spontaneous circulation, and she was transferred to intensive care. On waking she reported reduced vision and was found to have a left homonymous hemianopia.

Imaging of Miss F's brain revealed oedema suggestive of a cerebral infarction and a small subdural haemorrhage. Miss F's long-term warfarin was initiated and she was transferred to rehabilitation to have her stroke assessed. After two days of treatment Mrs B refused to take any more tablets because her nausea was so severe and she was commenced on intravenous ciprofloxacin.

The following day Mrs B had a cardiac arrest and despite adrenaline and DC cardioversion she died. A postmortem report showed she had died of a gram negative sepsis and gastroenteritis with salmonella parapharynx A. Mrs B's family were devastated and made a claim against Dr A. They felt that her death could have been avoided if Dr A had investigated and treated her diarrhoea earlier.

**EXPERT OPINION**

Medical Protection commissioned a report from a GP expert, Dr D. Dr D was not critical of Dr A's first consultation with Mrs B. At that time Mrs B had a three-day history of diarrhoea. Dr D explained that viral gastroenteritis is the commonest cause of diarrhoea and that traveller's diarrhoea is an extremely common presenting complaint. Even in cases of bacterial infection, antibiotic treatment is not usually required. As traveller's diarrhoea is self-limiting in the majority of cases, Dr D felt that few GPs would have requested a stool sample on that occasion.

Dr S was, however, critical of Dr A's second consultation. At that time Mrs B had complained of significant diarrhoea for ten days. Dr S felt the clinical records were very brief and did not include a record of the presence or absence of blood in the stool or abdominal pain.

Dr S thought that the patient's ongoing symptoms at this consultation required the identification of a causative organism and that a stool culture should have been arranged. It was his view that the failure to do so represented an unreasonable standard of care. He postulated that if a stool sample had been taken, this would have led to the causative organism being known within four to seven days.

The case was settled for a moderate sum.
Mr K was a 36-year-old man who ran a pub. Mr K smoked and drank heavily. Mr K's dentist had noticed a painless swelling on the right side of his neck during a routine check-up and asked him to see his GP. Mr K was seen by Dr A, one of the GPs at his surgery, who noted that Mr K was unsure as to how long the lump had been there, and referred him to the ENT outpatient department.

A letter came back to the practice confirming the presence of a lymph node in the anterior triangle of Mr K's neck, which was felt to be innocuous. The plan was for Mr K to be reviewed in six weeks' time and significant X-ray investigations to be pursued if the node was still present. Mr K was busy at work and did not feel too concerned about the lump because it was not painful. He did not attend his follow-up appointment, and a letter stating this was sent from the hospital to his GP.

Eight months later, Mr K began to get some discomfort in the neck swelling so decided to see his GP again. This time he was seen by Dr B at his surgery. Dr B notified his pain swelling and also a history of chronic idiopathic meningeal pain. Dr B did not document his previous referral to the ENT department regarding the same lump or the intended follow-up. Dr B's brief examination notes detailed the tender, swollen lymph node but did not include an examination of the mouth, tongue or throat. Dr B prescribed ibuprofen to help with the discomfort and did not arrange any follow-up.

Over a year later, Mr K was still struggling with his symptoms and went again to see Dr B. This time Dr B made a referral to head and neck surgery. His referral letter stated “intermittent chronic right sided neck swelling in the pre-auricular and submandibular area”. There was no mention of any previous referral in his letter. Dr B documented a differential diagnosis of a possible parotid lesion or salivary gland disorder.

Mr K's neck lump subsequently proved to be malignant. As a result he had to have neck surgery and resection of a primary in his tonsil. He had a course of radiotherapy and since has not had recurrence of his disease. Unfortunately he was left with shoulder weakness and a dry mouth, which he found difficult to cope with. Mr K was angry with Dr B and felt that he caused a delay in his diagnosis. He brought a claim of negligence against Dr B because he felt the delay had necessitated more radical surgery, leaving him with debilitating symptoms.

EXPERT OPINION

Medical Protection sought the advice of an expert GP (Dr F). Dr F felt that Dr B bore liability for the delayed diagnosis. He was critical of Dr B's history-taking and record-keeping. Dr F commented that Dr B had responsibility for establishing the history of his previous referral to the surgical assessment unit. Had Dr B known of that referral, then the duration and the continuing nature of the lymph node would have necessitated immediate re-referral back to that team. Dr F also criticised Dr B's inadequate examinations, stating that he should have documented an examination of the patient's neck, mouth, tongue and throat.

The opinion of a professor of otolaryngology (Professor Y) and head and neck surgery was also obtained. Professor Y commented that there was a significant delay between the initial presentation and the final treatment. Professor Y thought that had Mr K been examined at diagnosis, there may have been less radical neck dissection and it may have been possible to spare the accessory nerve, which controls the functions of the trapezius and sternocleidomastoid muscle. This would have resulted in less dysfunction to the shoulder and neck.

In addition, Professor Y considered that it may have been possible to spare radiotherapy if he had been treated earlier. The need for radiotherapy in this case was due to the size of the lymph node in the final specimen and the positive margins, which was evident following removal of the tonsil primary.

Due to expert-opinion finding Dr B to be in breach of his duty, the claim was settled for a high amount.

Mr P, a right-handed project manager, developed a stiff right elbow following a previous injury, and had reached the limit of his progress with physiotherapy. X-rays showed degenerative changes and he was referred to an orthopaedic consultant, Mr A, who diagnosed osteoarthritis of his elbow. He advised Mr P that as he had significant anterior and posterior osteophytes he may need multiple arthroscopic debride-ments to achieve a good outcome.

After an arthroscopic anterior debridement, there was minimal improvement and further surgery was planned. There were another two debride-ments, the third one being more than six months after the initial procedure, before Mr P was happy with the result.

Two months later Mr P returned with a reduced range of movement in his elbow. X-rays confirmed the presence of massive heteroetopic ossification (new bone growth), which was confirmed on CT. Mr A planned a fourth arthroscopic debride-ment two months later. No discussion relating to the possible risks and complications of surgery was documented. The limited operation note for this complex arthroscopic debride-ment described significant bone removal and a full range of movement at the end of the procedure.

In clinic two days later Mr P was noted to have a radial nerve palsy, but Mr A felt that some nerve conduction was present and that this was a neurapraxic nerve injury, which should recover completely. He commented that the procedure had been lengthy at over an hour and ten minutes. Mr P returned ten days later as there was no change in his symptoms, but Mr A was reassured by the presence of a positive Tinel's test and felt the nerve palsy would recover. He planned for review in six weeks, which was three months post-surgery, but again there was little improvement. Mr A commented that the positive Tinel's could now be felt up to the fingertips. An appointment for three months later was made, but still there was no improvement.

Six months post-surgery, Mr A now requested nerve conduction studies, which were performed within five days, and reported the presence of a severe radial nerve injury. Plans were then made for surgical exploration of the nerve with possible repair, grafting or neurolysis as necessary. Mr P made a claim against Mr A, stating that his nerve injury had left him with a permanent disability including reduced grip and manual dexterity, plus an inability to extend his fingers. He believed that the surgery should have been an open procedure rather than arthroscopic, and that had his injury been diagnosed sooner, and not presumed to be a neurapraxia, then he would have had a better outcome.

On review of the case, an expert felt that as long as Mr A had the necessary experience it was not negligent to carry out the surgery arthroscopically. There is still a risk of radial nerve injury when carrying out this surgery with an open technique. However, Mr A was found to be negligent in causing the nerve injury, keeping poor documentation, and delaying nerve conduction studies. The lack of any documented discussions about the risks of the surgery was also a factor in the outcome of the case. The case was settled for a substantial sum.
Dr A, expert in forensic medicine, stated that had Dr A been made aware of the test from Romania, it would have been a breach of duty to discount it. Assuming that Mrs S would have accepted the offer of amniocentesis, based on the timings, the test had been done. A month later, Mrs S was given the results of her Romanian triple test, which allegedly gave a risk of Down’s Syndrome of 1 in 67. Her combined test in the UK gave a much lower risk of 1:835. Based on the triple test, Mrs S was offered a termination of pregnancy. Mrs S made a claim against Dr A, stating that she had been given false reassurance regarding her test results, which had also failed to be documented adequately in her notes. It was alleged that had she been referred to an obstetrician for amniocentesis, then she would have chosen to undergo a termination of pregnancy.

EXPERT OPINION
Expert GP Dr D maintained that Dr A’s standard of care did not fall below that expected of a GP. Dr D felt that Dr A was entitled to rely on the screening performed in the local secondary care setting, which indicated a low risk of Down’s Syndrome and had no need for further investigations. Dr D’s account was that he was not told of the Romanian result, so was unable to take this into consideration. Dr D maintained that it would have been prudent to refer if this result had been available, given that it was carried out at 16 weeks – at a time when it would be less sensitive – it would have been reasonable for Dr A to have advised the patient to undergo a termination of pregnancy.

LEARNING POINTS

- Consultation with patients who do not speak the same language present a significant challenge for all healthcare professionals. If you cannot understand and communicate with a patient, you may be putting both yourself and the patient at risk. It is preferable to try to use an interpreter rather than a family member if possible, unless a patient presents acutely.

- Patients who undergo investigations overseas often return home for ongoing care and may present a challenge to GPs. Questioning an account of events, if possible, should be the best course of action.

Mrs S, a 27-year-old Romanian woman who lived with her husband in the UK, became pregnant and presented to her local GP surgery to commence antenatal care. Mrs S did not feel well and usually brought a family member with her to interpret. Mrs S presented to the emergency department at six weeks’ gestation and since she had previously suffered with a haemorrhage, an early scan was carried out, which confirmed a viable pregnancy. Mrs S received IV hydration and was discharged with oral cyclizine to use if the vomiting persisted.

A month later, she was feeling better. The vomiting had resolved and she was no longer using the progestogens. She visited GP Dr A, who noted “had Down’s scan, family member interpreter present, review at 16 weeks”.

Mrs S visited Romania for a holiday to see her family. While she was there she presented to hospital complaining of possible kidney problems with a secondary concern over reduced foetal movements. Mrs S underwent a pelvic ultrasound scan, which appeared to have shown a growth on her right kidney. Mrs S also claimed she underwent a triple test at this point.

After returning to the UK, Mrs S attended her routine 16-week check with Dr A. The practice antenatal template was completed and Dr A ticked that the Down’s screening test had been done. A month later, Mrs S visited Dr A, and the results of her Romanian triple test, which allegedly gave a risk of Down’s Syndrome of 1 in 67. Her combined test in the UK gave a much lower risk of 1:835. Based on her age, Mrs S had a background risk of 1.800 – therefore a risk of 1.67 would represent a significantly increased risk.

At 20 weeks, Mrs S presented to Dr A – her husband was present to translate but no interpreter was available. The baby was born with Down’s Syndrome and patent ductus arteriosis and developed septicaemia and pulmonary hypertension.

Mrs S made a claim against Dr A, stating that she had been given false reassurance regarding her test results, which had also failed to be documented adequately in her notes. It was alleged that had she been referred to an obstetrician for amniocentesis, then she would have chosen to undergo a termination of pregnancy.

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Mrs S, a 39-year-old chef, opted to see consultant obstetrician Mr B for private antenatal care. It was her first pregnancy and other than a BMI of 30 she had no pre-existing medical problems. She was reviewed regularly throughout pregnancy and noted to have elevated blood pressure through the first trimester, between 126/83 – 157/90. Methyldopa was introduced and this was well tolerated. Mrs Y gave birth to a healthy son. She was discharged from hospital with half hourly BP and hourly urine output measurements, which Mr B should have initiated.

A week following delivery Mrs Y continued to have elevated BP with proteinuria in spite of ongoing antihypertensive therapy. Mr B was contacted by the ward team and provided telephone advice to continue anti-hypertensives. The following morning the decision was made to deliver by caesarean section due to a raised blood pressure and patient request. Mrs Y gave birth to a healthy son. She was discharged on propranolol and irbesartan.

A week later Mrs Y was admitted to hospital with no relief of her meningeal irritation from a small bleed versus haemorrhage. Expert neurosurgeon Mr G commented that although based on the clinical signs there was a strong suspicion of an intracranial bleed, a scan confirmed a cerebral haemorrhage. Mrs Y died four days later.

EXPERT OPINION
Experts were critical of Mr B, commenting that it was unacceptable for him to fail to visit Mrs Y when called by the ward team regarding her symptoms; Mrs Y’s consistently elevated BP warranted high dependency management with half hourly BP and hourly urine output measurements, which Mr B should have initiated.

Mr B was criticised for not reviewing Mrs Y early enough when the ward team was calling. He was incorrect in the belief that a patient when well could be discharged with the education that he had provided. The patient was displaying red flag symptoms where a scan confirmed a cerebral haemorrhage. She died four days later.

Learning points

- It is easy to attribute any new symptoms a woman may develop during pregnancy to the pregnancy itself, but you should not require urgent assessment.

- Always documentation is essential. Mr B had commented that the patient was well and could be discharged, and his note may have been consistent with Mr G’s comment that a timely intervention could have made a difference.

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Mrs L, a teacher, was first prescribed the oral contraceptive pill microgynon by her GP, Dr G, when she was 22. Her blood pressure was taken and recorded as normal. At this time, no other mention was made in the records of her risk profile or family history. Later, Mrs L’s medical records showed that she was changed to ovran and then ovranette, but there was no explanation why these changes were made. Mrs L was changed again to levonorgestrel. The reasoning for this was due to “excessive bleeding on ovranette”. At her review consultation, Mrs L’s blood pressure was taken and recorded as normal.

When she was 26, Mrs L was seen by her GP as she was under significant stress. Her records showed that she had increased her smoking to 35 cigarettes per day and had not exercised. Counselling was offered but she admitted that she smoked 50 cigarettes per day. 50mg was prescribed and exercise was advised. In addition, a prescription for nicotine patches was also issued.

For the next six years, Mrs L was given repeated prescriptions of the microgynon pill without any record of her blood pressure being taken or her risk factors being assessed. Mrs L was now 35, but her GP, Dr F, did not say whether she was still smoking, under a lot of stress, or whether or not she was still exercising.

Four months after her last repeat script, aged 35, Mrs L presented to the same practice with central chest pain and was seen by another GP, Dr G. She had been under a lot of stress at work and had been smoking a cigarette a day. Her blood pressure was recorded as normal. After her first child had been born, Mrs L was prescribed amitriptyline, before she changed to the combined pill.

Three years later, Mrs L consulted her GP as she was under significant stress. Her records showed that she had increased her smoking to 25 cigarettes per day and did not exercise. Counselling was again offered, but she admitted that she was smoking 50mg prescribed and exercise was advised. In addition, a prescription for nicotine patches was also issued.

Mrs L was seen on a number of occasions in the practice for a repeat prescription for microgynon and other matters, including further chest pain, collapse and migraine.

At age 41, Mrs L collapsed and was admitted to the Emergency Department, where investigations found that she had a stroke. She was unable to return to work due to paralysis affecting her left side.

Mrs L made a claim against Dr F. She alleged that he had been negligent in continuing to prescribe microgynon after she was 35 years old, when she had three risk factors: a family history of heart attack, smoking and being over the age of 35.

**EXPERT OPINION**

**Expert opinion** found that a reasonably competent GP would have stopped prescribing microgynon from the age of 35 onwards and changed Mrs L to a progestogen-only pill (at least have warned Mrs L of the increased risks in order that she could have considered the alternative options). Mrs L’s notes show that the practice knew of Mrs L’s family history and her smoking, but despite these risks continued to prescribe the pill.

The case was settled for a substantial sum.

Mrs S was a 36-year-old patient diagnosed with a benign giant cell tumour of the sacrum. She was seen by Mr A, consultant in orthopaedic oncology, and listed for resection of the lesion. Prior to surgery Mrs S underwent preoperative embolisation.

Mrs S was also reviewed by Mr B, consultant vascular surgeon, who planned to introduce an aortic balloon through the femoral artery prior to the tumour resection if required, the balloon to be inflated and deflated during the surgical resection in order to reduce blood loss. Mr B sought consent for aortic balloon occlusion and documented that the risks included “femoral artery injury, limb ischaemia and bleeding from rupture”. Separate consent was obtained by the orthopaedic team.

Surgery was initially planned in the supine position to allow access to the femoral vessels. The right common femoral artery was cannulated and a 6Fr sheath inserted. This was exchanged for a 14Fr sheath under radiological control. A 40mm aortic balloon was introduced to the level of L3, its position being confirmed under fluoroscopy.

Mrs S was then turned to the prone position to allow tumour resection. The balloon position was re-imaged and found to be unchanged. Mr B left the operating theatre.

After two hours, Mr B was called back to the theatre to inflate the aortic balloon as haemostasis was required. The balloon was inflated by Mr B using an inflation device. Haemostasis was improved and the blood pressure stable. No further imaging was required, the aortic balloon could be inflated during the orthopaedic procedure.

Mr B received a telephone call to inform him the operation was finishing and he should return in order to remove the sheath and deflate the aortic balloon. Prior to him arriving at the operating theatre, the patient suffered a cardiac arrest and CPR was commenced.

Mrs S had an unrecordable blood pressure and at laparotomy a large retro-peritoneal haematoma was discovered secondary to a 2.5cm tear in the anterior aorta. The aorta was surgically repaired but after release of the clamps, Mrs S’s blood pressure was unable to stabilise and she suffered a further cardiac arrest and died.

Mrs S’s family made a claim against Mr B. It was alleged that consent was inadequate as the risks of death was not specifically mentioned. It was also alleged that the aortic balloon used was inappropriate and that it was inappropriate to inflate the balloon without radiological guidance. In addition, it was alleged that delegation of the deflation of the balloon to the orthopaedic team was unacceptable.

**EXPERT OPINION**

Medical Protection sought an expert vascular surgery opinion from Professor T. Although the risk of vessel rupture and bleeding was discussed, he was critical of the failure to warn of the small risk of death from aortic balloon inflation.

Whilst acknowledging that re-inflating the aortic balloon without guidance may have been acceptable as a last-ditch effort to save the patient’s life under extreme circumstances, the decision to initially inflate the balloon without radiological guidance and to delegate deflation to the orthopaedic team was also criticised.

The case was settled for a high sum.
TOO MUCH OXYGEN

I read with interest your case report of an extremely preterm baby with high oxygen saturations, who was not screened for retinopathy of prematurity (ROP) and who subsequently developed severe ROP, causing blindness.

However, the learning point that safe levels of oxygen saturation in low birth weight infants are between 86–92% is incorrect. In two large, multi-centre trials a targeted oxygen saturation level of 85–89% increased infant mortality compared with an oxygen saturation target level of 91–95%.

While the incidence of ROP was lower with lower oxygen saturation target levels, this does not outweigh the increased risk of babies dying. It is recommended that extremely preterm babies should have target oxygen saturations levels between 91–95%.

Dr Jane Alswieker
Neonatal paediatrician
Auckland
New Zealand

Response

Thank you for your email. We have discussed your comments with the author of the case report in question.

He has confirmed that the oxygen range quoted was from guidelines issued in 2010 and that a more recent meta-analysis has found that the lower range of oxygen saturations are associated with higher mortality at a later stage.

We are happy to correct this point and would like to thank you for your helpful comments.

REFERENCES
ESTABLISHING, MANAGING AND PROTECTING YOUR ONLINE REPUTATION – A SOCIAL MEDIA GUIDE FOR PHYSICIANS AND MEDICAL PRACTICES

by Kevin Pho and Susan Gay

Dr Aidan O’Donnell, consultant anaesthetist, New Zealand

How social media savvy are you? If you are a medical student, the chances are that you are online more or less permanently. If, like me, you are a practising doctor who qualified in the last century (read ‘dinosaur’), you might be a bit less comfortable. I’ve been using computers since you could measure the pixels with a ruler, and I carry my smartphone as if it were grafted onto my hand, but even I admit I am feeling a little left behind by the social media tsunami that has arisen around us. Social media is becoming increasingly popular among doctors and patients alike.

Where clear ethical and behavioural boundaries are well established in traditional face-to-face relationships, the online community has developed so rapidly that the medical profession is finding itself in uncharted waters. How do you respond when a patient wants to “friend” you on Facebook? Or when someone harshly criticises your doctoring on a public forum?

My organisation has released guidelines about how to behave online, but they are a series of don’ts. Don’t publish pictures of yourself drunkenly incapacitated on your Facebook page, where employers and patients can see them.

Into this environment come Kevin Pho and Susan Gay, with their book, Establishing, Managing and Protecting your Online Reputation. Pho is himself a doctor, writing for doctors, which gives him immediate authority. His blog, www.kevinmd.com, is well-known and successful.

The central theme of the book is that doctors’ online reputation is just as important as their real-life one. Whether we like it or not, our basic information is already out there, but we usually don’t take any ownership of it. Done properly, we can establish and cultivate an online reputation, which can be professionally and personally rewarding. In short, we can use social media to our professional advantage. To quote: “First, do no harm; second, get an online profile.” Rather than don’ts, this book is full of dos.

The book is informal and readable, and covers the absolute basics well: techno-novices need have no fear. My main criticism is the book’s overwhelmingly American perspective. Patterns of work and ethos of practice are very different where I work, and I don’t need to build myself – or my practice – as a brand, or attract my paying customers. Social media is here to stay, and need not be a threat. We can ignore it, or use it to our advantage, and this book goes a long way toward telling us how.

I’LL SEE MYSELF OUT, THANK YOU: THIRTY PERSONAL VIEWS IN SUPPORT OF ASSISTED SUICIDE

Edited by Colin Brewer and Michael Irwin

Reviewed by Dr Ellen Welch – GP, London

Following the recent rejection of the Assisted Dying Bill in the UK House of Commons by an overwhelming majority of 330 against to 118 in favour, this collection of essays in support of the issue provides the reader with some of the key arguments in the debate for the legalisation of what the authors term medically assisted rational suicide (MARS).

The book has been compiled by former psychiatrist Colin Brewer and former medical director of the United Nations Michael Irwin, with essays contributed by doctors, priests, politicians, philosophers and, most poignantly, from people suffering with terminal illness.

The writers discuss the facts and the law surrounding the subject in both the UK and overseas, with both ethical and religious perspective offered. Dignitas writes a chapter on their experiences in Switzerland over the last 16 years of their existence. And a chapter is dedicated to palliative care – both its promises and its limitations.

Perhaps the most thought-provoking stories come from people who have been faced with the reality of a painful, undignified death. They tell of their struggle, their pain, the frustration that they feel in a life they no longer want to live, but are unable to end. Several quotes are given from the 2014 House of Lords debate which sum up some of the main arguments.

A major limitation of this book is that it only presents one side of the argument on the debate and it would certainly provide more of a balanced read if there had been contributors from those who oppose assisted dying. Whatever your view may be, it does provide an interesting and comprehensive read in support of the issue.
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In the interests of confidentiality please do not include information in any email that would allow a patient to be identified.

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