**LETTERS**

**The Roman Catholic Church and embryonic stem cells**

Skeene and Parker raise a number of concerns about religious doctrine unduly influencing law and public policy through *amicus curiae* contributions to civil litigations or direct lobbying of politicians. Oakley picks this up in an essay with an emphasis on the Roman Catholic Church’s interest in preventing the destruction of embryos for embryonic stem cell research. Skeene, Parker, and Oakley seem to be concerned mostly with religious views having undue influence on public policy. My concern is the negative effect that such Church influenced public policy may have on the progress of the biomedical research that is itself foundational to the debate. Oakley seems to be particularly incensed that, as he puts it: “Those who support a total ban on embryonic stem cell research sometimes talk as if theirs are the views having undue influence on public policy. My concern is the negative effect that such Church influenced public policy may have on the progress of the biomedical research that is itself foundational to the debate.”

The Roman Catholic Church has historically defended the sanctity and dignity of human life to varying degrees at different times. Human life for much of the past 2000 years was defined by the Church as the presence of the soul, which was thought at different times to appear at various different stages during development. Only recently, with the advent of modern biology, has the Roman Catholic Church shifted its position to claim that the fertilised egg also qualifies as the right sort of human life. It should be noted that this doctrinal change was fundamentally driven by developments in our understanding of embryology and not the process of ensoulment. The Roman Catholic Church tried to prevent Galileo from publishing empirical evidence using his telescope and disseminating his empirical evidence by banning his books. Similarly the Church today has attempted to prevent the gathering of empirical data on the embryo by promoting a ban on all experimentation on early embryos.

The Copernican revolution itself has become a paradigm for the process of theory change in science. Science is not simply a collection of results from experiments (or facts) but perhaps more importantly science is the interpretation of those results and the planning of further experiments. For all its claims of objectivity science is, so the philosophers of science tell us, essentially a theoretical construct. The practical and theoretical sides of science are of course intimately connected. In fact it is well known that a researcher’s actions and observations are most likely guided to some degree by their own hopes and expectations. These same researchers develop the theories that they use to interpret their data. These theories fit the results (or facts) that have been previously observed and predict new experiments to be done. The role of theory at this stage of the process is often underestimated. Theories do not fall out of results. In fact in biology especially theories are often essential to making sense of what constitutes evidence (result) and what is noise (artefact). Theory then is not just a bridge to the next fact or experiment but arguably the very heart and soul of science. Theories that do not fit the facts are of no use and should be discarded. But in biology especially, theories can define what counts as a fact and what does not. Sooner or later a startling new observation is made that cannot be accommodated within the existing theoretical framework. New theories are developed and past observations are re-categorised. What was written off as noise then becomes fact. Thomas Kuhn called this a paradigm shift and his paradigmatic case was the Copernican revolution. One overarching theoretical construct is replaced with another—our understanding of the world is literally changed forever.

A problem arises when an organisation such as the Roman Catholic Church erects its doctrinal structure on the shaky foundations of a specific theoretical construct. Biology and developmental biology in particular are comparatively young sciences that are progressing rapidly and are thus quite theoretically diverse. By lending its support to a certain theory or position within biology the Church may proffer a disordered balance that exists in science whereby theories are valued for their explanatory power or instrumental use, not their doctrinal compatibility. External interest groups with political lobbying power may thus hijack the delicate process of progress in science with dire consequences for future advancement in science and medicine. The Roman Catholic Church’s influence on science is indirect and usually through the medium of public opinion and public policy. As we have seen in the American debate over the status of the embryo with regards to the derivation of embryonic stem cells this influence may be decisive in the formation of public policy. Indeed President Bush’s decision to effectively ban public funding of embryonic stem cell research in America is widely believed to have set back progress in the field worldwide by many years.

The Roman Catholic Church’s input into the embryonic stem cell debate has not been simply moral or ethical as one might assume but has opened defended a particular claim about the biology of the early embryo. Given the basic lack of empirical evidence regarding the embryo and such developments as the unexpected properties of stem cells the Roman Catholic Church’s choice of position on the biology of the embryo seems to be chosen solely as a prop for its doctrinal position. This prop has then been introduced into the secular debate on the status of the embryo as a somehow obvious empirical claim.

I believe the Church’s religious fervour for its preferred doctrinal and scientific position of the day is fundamentally at odds with the process and progress of science. Science is an exploration of the physical world that is characterised by continuous change and, historically at least, major shifts in understanding. Over the last 400 years the Roman Catholic Church has been slow to accept that science progresses at all and has proved to maintain its doctrinal position as a matter of faith even when it has been shown to be empirically unsound. My concern here is I think similar to that of Skeene.
and Parker. The Roman Catholic Church’s contributions to public policy are based not only on their moral or ethical principles, but on an effectively arbitrary and dogmatic application of those principles that is backed by the full force of what is effectively a very powerful lobby group in many countries. Like Skene and Parker, I have no answer to the problems they have raised. Historically one thing is certain, in the future the Roman Catholic Church’s current position on the embryo will be judged to have been right or wrong with the wisdom of hindsight. Just as we judge the Church’s persecution of Galileo almost 400 years ago now.

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References


Non-compliance: a side effect of drug information leaflets

The problem of non-compliance with treatment and its repercussions on the clinical evolution of different conditions has been widely investigated.\(^1\) Non-compliance has also been shown to have significant economic implications, not only as a result of product loss but also indirectly through the complication of disease management and its subsequent healthcare and social costs.\(^2\)

Non-compliance as a health problem

The term “non-compliance” might be taken to refer both to the failure to follow a drug regimen and to the failure to adopt other measures that contribute to improvement in health—for example, changes in lifestyle or diet. This letter focuses on the former.

Non-compliance with a drug regimen can be the result of a number of different factors\(^3\) and a variety of techniques have been developed in an attempt to control it.\(^4\) Of these, the few techniques that have been shown to be effective have only managed to solve the problem in specific situations over short periods of time. The use of such techniques to control non-compliance, particularly where these are effective, raises interesting ethical questions about the extent to which their application constitutes an infringement of the patient’s right to decide on how to manage their own health.\(^5\) Here we suggest that in some cases one factor that leads to non-compliance is the tendency to provide extensive and exhaustive information on side effects in patient information leaflets. Consider the following case.

A true story

One morning Dr Smith woke up with a slight cold—muscular aches, headache, chills, and nasal congestion. He decided to take some medicine to counteract its effects. His initial thought was to find something to combat his runny nose, so he chose a product specifically indicated for nasal congestion: “StopSnort”.\(^6\) After reading the product information leaflet, however, Dr Smith felt another kind of chill run down his spine. He was struck cold by the contraindications, warnings, interactions, precautionary considerations listed in the leaflet. If he used this drug, it said, he would run the risk of suffering nausea, anxiety, agitation, insomnia, hallucinations, convulsions, amazement, weariness, arrhythmia, dizziness... Rather than risk all of this, he thought, why not suffer a few bothersome sniffles? For his muscular aches, Dr Smith chose another drug, “Abatache”, but the risks described in the accompanying information leaflet seemed even worse. These included baldness, skin blistering, aseptic meningitis, pneumonitis, fatal hepatitis, gas-trointestinal perforation, blood in the urine, jaundice, kidney disease, peptic ulceration, mouth ulceration, visual abnormality... So in the end, armed with his clinical and pharmacological knowledge, Dr Smith simply opted to continue blowing his nose and suffer a few muscular aches. He had no desire to play Russian roulette with his health.

The principle of autonomy and the right to information

The principle of autonomy in medical ethics places the patient at the centre of medical decision making about his or her care. It places particular emphasis on the importance of informed consent, and suggests that, except in rare situations,\(^7\) no patient should undergo medical treatment or surgical intervention without his or her fully informed authorisation. This is the basis of patient-centred medicine.

To obtain valid informed consent, it is argued that the patient must receive sufficient understandable information to make a fully informed choice. It is practised this means that someone undergoing a specific treatment receives information from at least two sources. First, they are given direct information from their doctor or another health professional about the drug to be taken, recommended lifestyle changes, and perhaps a warning of the hazards related to non-compliance. And they will also be provided with information on some of the side effects attributed to the drug being prescribed. Individual patients will tend to understand this information in a range of different ways, and it is well recognised that they will respond with a variety of known behaviour patterns.\(^8\)

Secondly, the patient will also receive additional information on side effects from the information supplied with the drug itself. These leaflets tend to cite each and every one of the undesirable effects related—note “related”—to the principle active ingredient within the drug. The information can in some cases be so complete or detailed that even any extremely unusual syndrome described in relation to the use of the drug will inevitably be listed in the leaflet as a possible consequence.\(^9\)

This information can sometimes have a significant effect on the likelihood that a patient will take the drug in question and may lead to significant “non-compliance”. When patients with minor ailments read about all the problems that may occur from using the prescribed medication, they may start worrying, to say the least. Some people read the leaflet again and again. They may then consult another source of medical information such as a website and perhaps decide to take only half the dose for half the amount of time prescribed, or simply decide not to take the medicine at all.

In addition to the problem of non-compliance, the so called nocebo effect\(^10\) needs to be considered, whereby the patient’s mindset is often a key element in the appearance of either physical or imaginary side effects, as has been shown in various studies.\(^11\)\(^12\) Such an effect may be caused by information leaflets.

Complete information versus sufficient information

Practically any city dweller would refuse to use transport services, work tools, or recreational facilities if they were supplied with complete, absolute, and extensive information on the hazards these might entail. Precautions and warnings are usually goods things, but they should be kept within reasonable limits to avoid creating outright alarm. Too much information can sometimes undermine autonomy and also lead to significant harms through non-compliance.

It was shown some years ago\(^13\) that information supplied by doctors can generate side effects that cannot subsequently be corroborated by physical examination. As it happens all too often, the information was not as exhaustive or complete as it might be.

In view of this, we believe that the kind of information given in drug descriptions should be reassessed. The information should be true, accurate, and easy to understand in as complete a way as possible, but it should not generate alarm that can lead to deleterious consequences in the healthcare sector or in the economic sphere.

So what did the patient decide?

The patient, shocked and dismayed at the drug’s side effects, finally decides not to follow the doctor’s recommendation. He (or she) will try to relax, perhaps by smoking a cigarette laced with nicotine, tar, and a number of other substances. True enough, doctors recommend giving up smoking. But who will listen to what a doctor says about smoking when they appear to be prescribing drugs truly hazardous to health? After all, a pack of cigarettes only says that cigarette smoking seriously damages your health. There is certainly no leaflet listing each and every one of its possible side effects. Tobacco kills, but it sometimes looks as if medication is worse.

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Response to ‘Patient organisations should also establish databanks on medical complications’”

Gebhardt in his brief report pleads for patient organisations to establish databanks on medical complications. Given the references (for example, an article by Paans, a journalist, entitled ‘Medical errors to be kept secret’), the risk is substantial danger of misinterpretation of the current situation, which in turn may frustrate the process of increased transparency. We would therefore like to respond to this by giving background information and reasons for some of the choices that were made with respect to the registry of complications mentioned by Gebhardt.

First, a distinction needs to be made between error and adverse outcome, which are often confused. From Gebhardt’s reference to the journalist’s article which discusses the same registry of adverse outcomes, but with the title referring to errors, both Gebhardt and the journalist think errors and adverse outcomes are the same thing. However, an error refers to the process in which something has gone wrong, a substandard performance, regardless of the outcome. It has been explained by others that such a judgement may have a degree of subjectivity. An adverse outcome refers to the outcome which is unwanted but does not necessarily imply that an error has been made. This is why the term “adverse outcomes” is used rather than the term “complications”, since the latter term is often confused with an error being made. The registration of medical complications that Gebhardt refers to is a registry of adverse outcomes guided by an unambiguous definition of the term “adverse outcome”, of which only a small percentage is related to errors. Furthermore, some errors will be missed in this registration—that is, errors which have not led to adverse outcomes.

Secondly, with respect to confidentiality, this is relevant in particular for the initial years of such a registry during which it is thoroughly tested and accuracy of the registration may vary widely between participants. Nothing is gained by false positive signals with respect to the high incidence of adverse outcomes in some hospitals, except perhaps by flashing headlines in newspapers. In this respect one may compare the development of such a national registry to the development of a new drug, in which case no one argues about confidentiality and thorough testing until proved safe. Moreover, a pharmaceutical company will probably be sued if it markets a new drug without proper research. It is intended that after this initial period, national adverse outcome data will become available to the public with respect to probability of an adverse outcome given certain types of surgery.


We read with interest the papers on informed consent published in a recent issue of the Journal of Medical Ethics.1 Whatever their differences, and however much they question some aspects of the duty to respect autonomy through attempting to obtain informed consent for therapeutic interventions, there was general agreement that competent adult patients are entitled to a core of basic information about their treatment options. There was also consensus that training in the process of obtaining consent is important. In our experience, two dimensions of such training are of particular interest. On the one hand, students require good theoretical understanding of the ethical and legal background to the professional emphasis now placed on informed consent. On the other hand, they need practical training in the relevant communication skills and how to apply them to obtain consent for specific clinical procedures. To do so, doctors must obviously also have a good understanding of these procedures. We recently encountered serious problems in cases such understanding in a study among junior doctors in England (Schildmann J, Cushing A, Doyal L, Vollmann J). The ethics and law of informed consent: knowledge, views and practice of pre-registration medical students. Submitted (subsequent publication). No matter how good their philosophical and legal knowledge, preregistration house officers (PRHOs) will not be able to deliver the minimal standards of informed consent outlined by O’Neill unless, suffice it to say, they know what—practically speaking—they are talking about. In contrast to Bravo et al’s results (in the same issue of the journal), almost all the PRHOs who took part in our survey had good legal understanding of the differences between competent and incompetent patients. This may be interpreted as a positive result of the change in the curricula at their particular medical school, which includes extensive sessions about informed consent. These embrace ethics, law, and communication skills. However, despite their understanding, the junior doctors in our study still experienced problems about their role in the consent process. The problems pertained to pressure of time and lack of support by senior doctors, as well as pressure on them at times to obtain consent in circumstances where they had been taught that they should not. This gap between the standards of informed consent currently taught to medical students and the clinical realities they face, and into which they are thrust, is an ongoing problem.

If informed consent is to fulfill the purpose of respecting the autonomy and dignity of patients, sufficient resources are required to train young doctors to do the job properly, especially as regards their understanding of procedures for which they are providing information and their competence as communicators. One thing is clear: if they cannot complete the task in accordance with the guidance issued by the General Medical Council and the Department of Health, they should not be doing it at all.1 Trusts and colleges should ensure that all supervisory staff are aware of their responsibilities in this regard.

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Patients need information to make a well-informed choice

Who is a good doctor and what is a good hospital? This simple question is not easy to answer. For some, it involves a recent diagnosis and the best treatment. The NPCF (Dutch Federation of Patients and Consumer Organisations) and its member organisations have published several consumer guides for specific diseases to help patients find their way in the labyrinth of the healthcare system. Patients experience many difficulties in getting access to relevant information from doctors’ organisations and insurance companies. Therefore, the NPCF wants to cooperate with these organisations to create consumer information based on the important and relevant data that are available. A joint project for a databank on best practices started in September 2003.

Patients are not interested in black lists of doctors and malpractices; they prefer to know about good and best practices to make a well-informed choice for a doctor or hospital. They need consumer information on objective measures such as the risk of infection in a hospital, the specific skills of a doctor, how many patients with this specific disease a doctor treats a year, etc. Patients would also like to receive subjective information on a specific hospital or doctor: How is the communication between a doctor and his or her patients? Does the team give enough information and support when needed?, etc. This experience based information is often available from patient organisations.

The NPCF has chosen to work together with organisations of healthcare providers and insurance companies to use parts of their databanks as a basis for consumer information. One task of the NPCF is to translate the data into consumer information that meets the needs of the patients based on research and experiences of patients. Joint efforts are needed to make this important information accessible for doctors and patients.

Finally, what does the patient want? (see box 1). International research has shown that patients do not use public information on performance of hospitals or doctors for making a choice of treatment or hospital because, among other reasons, they do not understand and do not trust these data. This also applies to adverse outcomes data. For interpreting the incidence of hospital specific adverse outcomes it is important to know the context—for example, since older, sicker, and more complex patients have higher probabilities of adverse outcomes. It is therefore vital to establish a reliable registry which can be trusted and understood both by medical professionals and the public. For this reason, the Association of Surgeons of the Netherlands and the Dutch Federation of Patients and Consumer Organisations (NPCF) are collaborating with respect to the national surgical adverse outcome registry, in particular, to produce information that is relevant for patients about treatment and hospital choices. Supported by the international literature, the NPCF holds the view that patients are not primarily interested in data on adverse outcomes, since they are aware that these data need to be interpreted in the right context. Patients are more interested in the experience of doctors or hospitals to treat certain diseases or to perform certain operations, since the question they want answered is “What is the best place to go to for this type of problem?”. That this doctor or hospital probably has a high adverse outcome record is not relevant, since this may well be explained by the complex patients who are referred to more experienced doctors. As argued in a previous paper, it is essential that there is an increased mutual trust between the medical profession and patients’ organisations that supports a combined effort to improve the quality and availability of patient information. Such initiatives, such as the database on best practices started in September 2003, are needed.

In the last paragraphs of their article, Joffe et al write: “We do not recommend that patients’ perspectives should unilaterally determine ethical frameworks. We do, however, believe that data such as those presented here can contribute to the search for reflectieve equilibrium in bioethics”. The term “reflective equilibrium”, as the authors note, was introduced by John Rawls. At least in its first instance, it refers to a way of constructing a moral theory by balancing one’s considered moral judgements against one’s moral principles, until one’s judgements and principles form a consistent set—that is, a moral theory (p 288). Joffe et al’s idea seems to be that by surveying patients’ perspectives they will be able to capture one side of this equilibrium, considered moral judgements, or moral principles (they do not specify which), and in so doing contribute to the desired end: a consistent ethical framework to govern medical encounters, built (at least in part) from the principles and moral judgments by the patient community. Whatever the merits of this goal, however, Joffe et al fail to capture either the considered moral judgements or the moral principles of those they survey and so fail to contribute to the moral theory they seek to construct.

Rawls defines considered moral judgements as those judgements in which our moral capacities, which he considers analogous to our linguistic capacities, are not likely to be displayed without distortion—for example, those offered without hesitation, given without strong emotions like fear, and made in the absence of conflicting interests (p 47). The discussion focused on the considered judgements and judgements generally is important. When constructing a moral theory for a particular community—our community, the patient community—we want to use only those judgements that reflect the respondents’ real moral sensibilities, and not those stemming from superficial prejudices or their mood on the day they happen to respond. For instance, the authors ask: (1) how can a researcher know whether he or she is collecting them—that is, what survey method, if any, is appropriate for the task?
Although it is difficult to give a positive answer to these questions (and I will not attempt to do so here), some survey methods, such as the mailed questionnaires that Joffe et al used, seem particularly inadequate. Rawls suggests that certain external conditions favour the formation of considered judgements: ‘the person making the [considered] judgment is presumed to have the ability, the opportunity and the desire to reach a correct decision (or at least, not the desire not to)’ (p 48). Very likely, however, many of Joffe et al’s respondents lacked the necessary ability, opportunity, or desire to consider their moral judgements when responding to the questionnaire they received in the mail. Furthermore, even if a number of patients did offer legitimate considered judgements, there is no way to distinguish these from those made by respondents who lacked the requisite ability or desire. Although the size of Joffe et al’s study is of value for its ability more accurately to reflect a population’s response to its survey questions, because of the practical limitations that come with its size, the study falls short of capturing patients’ considered moral judgements.

Any empirical approach using reflective equilibrium, as Joffe et al’s, faces a second challenge: why do we want people’s considered moral judgements to influence our theories of ethics in the first place? In his influential critique of reflective equilibrium, D W Haslett writes:

… given the wide differences between people’s considered moral judgments, and given that these differences are, as we know, largely just a reflection of differences in upbringing, culture, religion, and so on, it would appear that, far from having a reason for giving people’s considered moral judgments initial credibility, we have instead a reason for initial skepticism (p 309).4

If moral judgements are liable to reflect superficial prejudices, one could argue, considered moral judgements are liable to reflect deep seated ones. Surely this prejudice is something a health care provider would like to overcome, not codify. While I do not think this challenge is insurmountable, it does demand that researchers justify the inclusion of considered judgements in ethical theory before using the method of reflective equilibrium. Joffe et al have failed to do this.

Joffe et al’s study is susceptible to a second line of critique. Even if the study’s use of mailed surveys is appropriate, it fails to capture either patients’ considered judgements or principles, because, put simply, it does not ask for considered judgements or principles. Instead, it asks patients whether providers respected their person or respected their autonomy, and then tests patients’ responses to questions against whether they report being satisfied with their care. If a provider’s acting with respect for persons is a better predictor of patient satisfaction than him or her acting with respect for autonomy, Joffe et al conclude that the principle of respect for persons should be assigned as much importance, ethically speaking, as the principle of respect for autonomy. As should be clear, this conclusion does not follow from Rawls’s conception of how one constructs a moral theory. In a Rawlsian view,4 a moral theory requires knowing which principles patients hold, not whether those principles are associated with patient satisfaction. Joffe et al seem to be operating with an underlying utilitarian assumption to the effect that what we ought to do is whatever will lead to the greatest patient satisfaction. Although there may be reasons for accepting this utilitarian assumption (which Joffe et al do not provide), certainly there are others for rejecting it. For instance, although patient satisfaction may give a hospital a very good reason to change a policy, we probably do not want to say this reason is a good ethical reason. It is just good business sense. This is an especially important point given the principles that Joffe et al evaluate. Respect for autonomy and respect for persons are traditionally viewed deontologically—that is, it terms of duties or rights, which are valued for their own sake and not for the consequences (such as patient satisfaction) that they produce. In any case, these utility considerations take us far from patients’ actual moral views, the very things Joffe et al, by invoking Rawls’s reflective equilibrium, propose to capture.

Lastly, there is a question of their instrument’s validity. As I have been arguing, Joffe et al claim to assess whether patients are treated according to principles of respect for autonomy and respect for persons. Yet, their single item assessing respect for autonomy—the question, ‘do you feel you had your say?’—does not do the principle justice. The principle of autonomy not only requires that the health care provider asks the patient for his or her opinion, but also that the provider acts on the patient’s opinion. Their instruments are similarly inadequate for the principle of respect for persons, which, they suggest, includes ‘assertion, fidelity, veracity, avoiding killing, and justice’, as well as ‘respect for the body, respect for family, respect for community, respect for culture, respect for the moral value (dignity of the individual), and respect for the personal narrative’ (p 104). How are we to know whether patients had all or any of these in mind when they answered the question: ‘Did you feel like you were treated with respect and dignity while you were in the hospital?’ Joffe et al acknowledge that these ethical concepts are a bit unwieldy for a survey of manageable length. However, these practical considerations should be used not only to excuse the study but also to question its ability to clarify the ethical concepts it claims to assess. They should prod us to ask, regardless of the survey’s scale and the limitations that its size produces: does this survey really address what we mean by the principles of respect for autonomy and respect for persons?

With any empirical study in bioethics, there is a gap between the empirical hypotheses the study confirms and the normative conclusions its authors would like to draw from it. In their article Joffe et al hoped to bridge this gap by invoking Rawls’s notion of the reflective equilibrium. As I have explored, however, the study does not contribute to either side of the reflective equilibrium they imply, and, thus, they fail to demonstrate how their findings challenge the centrality of autonomy and shared decision making in bioethics.

Joffe et al’s failures are instructive, however, insofar as they suggest how we could better bridge the gap between research and theory. The use of the reflective equilibrium in empirical research has promise, provided researchers are clear about: (1) how to define considered moral judgements and/or principles; (2) how their methods capture these judgements and/or principles reliably; (3) how the inclusion of considered moral judgements strengthens rather than weakens bioethical theory; and (4) how their findings are valid for the judgements or principles they mean to assess. In empirical research can contribute to bioethics by questioning the assumptions implicit or explicit in our normative views. Joffe et al try to do just this when they argue in the introduction to their article (p 103) that patients’ desire to delegate decision making challenges the mandatory autonomy view. However, if empirical findings are to defeat a particular normative principle, the assumption that those findings challenge must be logically necessary for our holding that principle. For instance, without showing that patients’ desire for autonomy is necessary for our holding the mandatory autonomy view, the studies that Joffe et al cite, even if valid, can be interpreted variously as devaluing the mandatory autonomy view or as recommending that we better consider the value of autonomy. This normative question cannot be settled empirically.

Empirical researchers have the potential to contribute substantially to bioethics, but their work needs the kind of purity and empirical rigor that comes from truly interdisciplinary collaboration and must be informed by a careful reflection on the proper relationship between descriptive and normative ethics. Joffe et al take us part of the way down that path. An exciting research itinerary lies ahead.

D P Narendra

References

How to be a ‘good’ medical student
The public revelation in 2003 that medical students perform intimate examinations without patient consent has engendered much debate in the press and scientific journals. Using this case as a springboard for discussion, I will argue that medical schools should encourage students to raise their ethical concerns and call for a change of policy making it easier for students to do so. I will also address the question of medical students’ moral obligations towards their patients, and conclude that medical students ought to express their discontent when faced with unethical practices or attitudes.

In early January 2003, a study appeared in the British Medical Journal revealing that nearly a quarter of rectal and vaginal examinations on anaesthetised patients were performed by medical students without patient consent. Although the study did not generate the firestorm of controversy many expected, it engendered much discussion on ethical issues surrounding informed consent and patient autonomy, as well as stressing the need for greater ethics training for medical students. As an ethical problem, the need for greater ethics training for schools should encourage students to raise their ethical concerns and call for a change of policy making it easier for students to do so. I will also address the question of medical students’ moral obligations towards their patients, and conclude that medical students ought to express their discontent when faced with unethical practices or attitudes.

sphere of intimate examinations. It concerns the moral obligations of medical students faced with ethically dubious situations. In short, what should a “good” medical student do?

In an article on the scope of medical ethics, Professor Raanain Gillon recounts two experiences from his days as a medical student. The first describes his colleague’s refusal to grant an abortion to a 14 year old girl on the grounds that she was “a slut”; the second his own refusal to examine a scrotal lump on a patient whose testicles had already been examined by five other students. Gillon’s objections were very much the exception. When these events took place in the 1960s, medical students were simply expected to follow their teachers’ orders and to absorb their evident wisdom without question. Since then, medical ethics has developed from an ill-defined embryonic subject to an academic discipline in its own right, with specific journals and associations, and a place in the medical curriculum.

Judging from some of the comments students at Bristol, however, the growing emergence of medical ethics has not dispelled the awkward climate of unquestioned reverence towards teachers. Many of the students felt uneasy about the examinations, but were too intimidated to voice their concerns: “You couldn’t refuse comfortably. It would be very awkward, and you’d be made to feel inadequate and stupid”, commented a fourth year student who participated in the study. It seems clear that medical schools should strive to foster a climate more conducive to open discussion on ethical issues between students and teachers. Students should not have to perform heroic acts of courage to raise ethical concerns. In light of medical ethics’ place in the curriculum, the situation is deeply paradoxical. Students may be taught the importance of respecting the patient’s autonomy one day, but witness an obvious violation of this principle by their teachers the next. For the subject to be of any relevance to the medical students, jointly with their super-

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2 Coldicott Y. “To respect the patient’s integrity is the key.” BMJ 2003;326:100.

NOTICES

The JME editorial office has now moved
The JME editorial office has now moved to BMA House. The new contact details are: Journal of Medical Ethics, BMA House, Tavistock Square, London WC1H 9JR. Tel: +44 (0) 207 383 6439. Fax: +44 (0) 207 383 6668. The point of contact is Nayahna Siva, Editorial Assistant.

Institute of Medical Ethics Student Electives
The JME wishes to award 10 bursaries of up to £500 each to support Medical Student Electives, or exceptionally Special Study Modules, on issues in medical ethics.

Medical students, jointly with their supervisor, are invited to apply by 28th February 2005. Application is to be done via email, explaining the project’s relevance to medical ethics and the reasons why a bursary is requested. An outline study protocol and project budget should be included or attached.

Applications should be sent to Mrs M Bannatyne, JME Bursaries Administrator, daniel.sokol@imperial.ac.uk

SUCCESSFUL APPLICANTS

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An error has been pointed out in the affiliation for R Andorno, author of The right not to know: an autonomy approach (J Med Ethics 2004;30:435–439). The correct affiliation is Interdepartmental Center for Ethics in the Sciences and Humanities (IZEW), University of Tubingen, Tubingen, Germany. The journal apologises for this error.

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