Book reviews

Pricing Life–Why It’s
Time for Health Care
Rationing

Peter A Ubel, Cambridge, MA, The

Pricing Life is an extremely timely and
stimulating book. The debate about
health care rationing has now been
running for several decades but there
are still people who publicly deny that
rationing of health care has to take
place, or that health care rationing
could be ethically justifiable. In many
countries we have also had politicians
who have refused to allow the “R”
word to pass their lips. In this well
written, and very direct book Peter
Ubel decisively shows that rationing is
an inevitable feature of modern medi-
cine and that it is not something that
health care administrators and econo-
mists force doctors to perform against
their better judgment. Deciding who
should have access to treatment has
always been a part of the doctor’s
work. Ubel’s analysis of the phenom-
enon of bedside rationing is incisive
and definitive. Anyone who will still
deny that rationing does and must take
place after having read this section can
only be in bad faith.

The book would have been a
valuable contribution to the literature
even if it had only contained the
analysis of the necessity and ubiquity
of rationing in health care. But it also
contains a second part which reviews
and discusses the current knowledge
about how the public view health care
rationing and how they would distrib-
ute health care resources. Ubel is well
known for his own work in this field
and the book gives a very comprehen-
sive and useful critical overview. It is
explained what the main findings are,
but also that these findings are
context-sensitive and sensitive to the
framing of questions. For the health
care ethicist this is perhaps the most
important part of the book. In the
literature one can at times find
categorical statements about the
attitudes of the public–of the type:
“Everyone prefers to give resources to
the young and not the very old”–but
often these statements have no other
basis than the philosopher’s own
musings in his or her armchair, or the
views of his or her friends. Ubel’s
book shows that such categorical
statements are most often wide of the
mark and that the public’s views are
much more nuanced. In doing so he
also shows why empirical studies in
this area are important, as well as their
limitations.

In this part of the book he also dis-
cusses how various forms of prefer-
cences, for example concerning age
groups or justice can be incorporated
into QALY-type measures.

Ubel does not present a new philo-
sophical approach to justice, but he
does show just how much work still
has to be done before our well-known
theories about justice can give deter-
minate answers to the question of how
scarce resources should be rationed in
the health care system.

I can recommend this book without
reservation. It should be read not only
by health care ethicists but by doctors,
health care administrators and, per-
haps especially, by health care politi-
cians.

It is written in an American context
but there is an explicit recognition that
there are many other health care
systems than the American, and that
some of them might even have features
that are superior to the American sys-
tem. Its conclusions are therefore eas-
ily transferable beyond the American
context.

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An Ethics Casebook
for Hospitals:
Practical Approaches
to Everyday Cases

Mark G Kuczewski and Rosa Lynn B
Pnkus, Washington DC, Georgetown
University Press, 1999, 219 pages,
£17.25, $23.95.

In spite of the stress on evidence and
audit, clinicians still talk to each other
mostly about cases, especially when
there are difficulties. So when they
come to consider problems in medical
ethics, the focus is on case presenta-
tion too. However learned the partici-
pants, ethical analysis in medicine
without cases feels like an egg without
salt. Yet the oddity remains that case
discussion is often done very badly,
new tales are thrown in on a “me too”
basis without any apparent question-
ing about why a second one will help
when we haven’t got far down the road
on sorting out the first. It’s not
surprising therefore that some still
dismiss these discussions as “mere
anecdote”, or that there is a continu-
ing critical literature (initiated in part
in this journal) about the use and
value of the genre itself. This ambiva-
lence must be part of the explanation
as to why, among the burgeoning of
books on medical ethics, published
collections of cases well analysed
are still rare. This is therefore a welcome
addition to the landscape.

It has many features which com-
mand it. These are real cases, culled
from the daily life of ordinary hospi-
tals, not academic centres: so they ring
bells as being important and common
situations. The authors express some
of the discomfort here: such cases
can’t always be neatly categorised, and
they express a “day-to-day moral
struggle”, but they do contain many
of the bread and butter issues of our
discipline, often articulated but still in a
sense unresolved. The cases here are
fleshed out with their particular difficulties, and so (although there are references to key articles), the major signposts of medical ethics seem to be of less use than is often thought in sorting out what to do. As a consequence, the authors speculate on different possible endings. There is a bibliography, readings being helpfully connected to the cases.

But of course as with insurance cover or a long term relationship it's in the details where disappointment may lurk. It's sad, for instance, given the important work published elsewhere, to find that the references are almost entirely from America. But being over-text, over-made and over here is not the main problem. To this reviewer the punchlines are still too flabby, the objectivity not sharp enough. It still reads too often like clinicians speaking to clinician friends. Other voices don't seem properly heard or satisfactorily amplified. Take one case, called “Caring for the indecisive patient”, as an example. This turns out to be about a patient with an ischaemic foot who after admission was found to have widespread and incurable cancer. In spite of an expressed wish for quality of life in her last days, when her ischaemia progresses she is advised to have an amputation and agrees, but then refuses several other interventions and says she wishes to die quietly. Fascinating staff responses are presented but not discussed. The husband has alcohol on his breath, and so it is assumed by nurses and social workers that “he might not have the patient’s best interest at heart”. (His wife who came in with a sore foot is now near death and he’s not supposed to have a drink before he comes in to see her!) The physician is concerned that “reimbursement for hospital days would be denied if no treatment was given” . . . and thinks there was a “significant risk that increased narcotics for pain would ‘do her in’”. He didn’t go to the team meeting to discuss her care. We don’t need to go on. There are well known attitudes represented here that, in part at least, concern the ethics committee. They worry that if they are too critical of a doctor’s care that doctor might not bring future cases to the ethics committee. Understandable, and they all have to meet at PTA gatherings or on the golf course, but that doesn’t surely prevent a robust analysis in a book, where issues not personalities are to the fore. To the reader, this poor woman seems shocked by the discoveries, and caught in transition, but not indecisive. She did a great deal better than Hamlet at making up her mind in difficult circumstances: her tragedy lay in what was beyond her control. This book reminds me of a comment on a close friend’s essay at school on religion, the friend being a parson’s child: “Could we do better, seeing who you are”.

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Promoting Healthy Behavior: How Much Freedom? Whose Responsibility?

Edited by Daniel Callahan, Washington DC, Georgetown University Press, 2000, 186 pages, £32.50.

When my family and I were hiking in northern New York state we got lost. I waved down a passing truck and inside I just could just about through the haze of purple cigarette smoke, make out a couple of very obese people in the car; they looked like mother and son. They very kindly agreed to take us to our car, several miles away, and during the journey I asked them what had brought them to this remote mountain road. “We come here once a week to draw water from a mountain stream”, said the elderly woman. “Most people round here don’t know how to keep healthy”, she and her son opined almost in unison.

This experience, for me, captured many of the attitudes those of us in the West have towards health, health promotion, and disease prevention. This cameo shows how difficult it is to change unhealthy behaviour. Persuading people to give up smoking and reduce weight is more of a challenge than persuading them of the health-giving qualities of mountain spring water. In the West, and in particular in the United States of America, individuals are preoccupied with their health in ways not envisaged fifty years ago. In addition to recognising that we should take greater personal responsibility for our health we are acutely aware, also, of how the behaviour of others can affect our health. And it is in this realm, between the “right” to be free from state interference and lead one’s life as one sees fit and the “right” to be encumbered by unnecessary threats to one’s wellbeing, where tensions arise. These tensions can be seen to play out in the realm of passive smoking, for example. The bans on indoor smoking to prisons in Vermont (and the subsequent lifting of these bans because of a thriving black market) and the US Supreme Court’s decision that exposure of prisoners to tobacco smoke violates the eighth amendment (the right to freedom from cruel and unusual punishment) highlight Beck’s “risk society” in ways which are sometimes difficult to navigate ethically.

This excellent book, the result of a two-year research project conducted by the Hastings Center and the Stanford University Center for Bio-medical Ethics, collects together a series of essays that reflect upon ethical, political and cultural issues surrounding the promotion of healthy behaviour. Perhaps not surprisingly, therefore, given the authors’ North American origins, the book takes a US perspective. Although health promotion and disease prevention terms are used on occasion interchangeably with the promotion of healthy behaviour it is the latter, as the book’s title suggests, that the collection focuses upon. This is interesting in itself and illustrates something of the strictures placed upon those in the US who ally social justice to public health. Ronald Labonte and Ann Robertson, the two non-US-based contributors, each provide papers which add breadth to the book and are superb. Meredith Min- kler and Beverly Ovrebo have also produced impressive chapters, which highlight with clarity the ethical dilemmas which arise when trying to balance individual and societal needs and responsibilities. Ovrebo notes that: “While the debates are old, what is new is the context within which these debates are occurring” and it is these changing circumstances which make the book so interesting.

The collection has, in my opinion, failed in two areas. Firstly, although the ethical and social tensions in changing behaviour are described well, little space is given to potential solutions. These, of course, are political. But if collections such as this are to have a lasting impact then they must provide argued political solutions. This is a lot to ask of a book but if an ocean liner of the magnitude of the American body politic is to change course it needs all the help it can get. Secondly, the health promotion consequences of the unravelling of the human genetic code are insufficiently covered. In the recent years or so the knowledge which arises from this project will transform the ethical
debates regarding the promotion of healthy behaviour.

More than a quarter of a century after Marc Lalonde drew attention to the individual as an important focus for intervention to control risk factors for ill health, this book provides a timely reminder that changing behaviour continues to be politically, culturally and ethically challenging.

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**Bioethics: an Anthology**


It has often been remarked that developments in science and technology proceed at a pace which far outstrips efforts to understand or control them. This clearly applies in the field of bioethics. Scarcely a day goes by without the newspapers being filled with the latest genetics scare story or medical controversy. So Blackwell’s latest anthology is certainly a book whose time has come. The question really is not whether it will find a market, for there are many, both students and professionals, who will find this tome an invaluable reference, but whether the editors have managed to strike a suitable balance between their own, avowedly utilitarian approach, and approaches of those who place a greater emphasis on notions of rights and duties.

Kuhse and Singer introduce their admirably comprehensive collection by noting that we humans owe a duty not just to other humans–but to the “biosphere”, and that applied ethics is rooted in a need (as argued by R M Hare) to find general rules and principles which can then be applied universally. The anthology follows the same pattern as their earlier and popular *Companion to Bioethics*, yet allows additional space for papers on topics where the “Companion” was weak, particularly debates surrounding new reproductive technologies.

The anthology is grouped into eleven main categories. Part one, Before birth, deals with abortion, including Judith Jarvis Thomsons famous violinist in need of a loan of a woman’s life-support system. Part two, Issues in reproduction, looks at IVF, “Dolly” the sheep’s experience and fertility treatment. A very brief piece by Hope, Lockwood and Lockwood, uses the rather fatuous claim that it is (almost) always better to exist than not to exist, to argue that IVF treatment should be made available without judgment as to the suitability of the would-be parents. Jennifer Jackson argues that because “a century ago” life expectancy was 47 for women and no one objected to them, and because “myth” that letting die is not equivalent to intentionally causing death, her article on “Why killing is not always worse–and sometimes better–than letting die”, and Singer’s piece, Is the sanctity of life ethic terminally ill?, which cites the case of Tony Bland.

Parts five and six look more briefly at resources and organ donation, whilst parts seven and eight offer some thoughts on experimentation with, in a significant choice of sequence, human subjects and animals.

Finally, contributions on issues in the practice of health care such as truth-telling and consent, conclude the volume along with four papers on ethicists and ethics committees, to the effect that, as Dan Brock puts it, professional philosophers of health care ethics are a bad thing, whilst occasional academic visits into the arena, by those whose primary interests remain outside, are best.

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