Debate over whether mammography screening saves lives has become front page news again in recent months. The opening salvo came in October 2001 from Gotzsche and Olsen, who argued in the _Lancet_ that five of the seven randomised trials of screening mammography were of poor or flawed quality and that the remaining two fail to show that the benefits outweigh the risks. In the US, scientific experts remain divided in their recommendations regarding the benefits of mammography for women in their 40s, although the practice has been strongly defended by political leaders. This controversy, like many in the realm of disease prevention, raises an important question: when science is uncertain, what else drives an individual investigator’s interpretation of the evidence?

In the midst of a public debate over prevention recommendations, it often seems that even in the best of circumstances (that is, when there is a randomised clinical trial) there is little objective knowledge to be found. Indeed, prevention science is frequently charged with being subjective and value laden. The investigator’s scientific judgment may be polluted by influences other than the evidence itself. The list of suspected polluters is long, including commitment to a favourite theory, type of training (physician, epidemiologist, biologist), personal habits (smoking, diet), concern about the financial costs of screening, and attitudes towards balancing false positives versus false negatives, to name just a few.

Such influences seem to threaten traditional notions of scientific objectivity. At the start of the scientific revolution in the 17th century, the character of a scientist was believed to be as important as any experimental technique. A scientist, it was believed, should exhibit selflessness, disinterestedness, and a willingness to change her views. In other words, the scientific investigator should be a neutral observer with no personal interest in their theories or results. Today, in comparison, we are somewhat jaded. No one claims that scientists can be wholly objective and impervious to outside influences, but, at the same time, few of us are willing to give up on scientific objectivity.

Some epidemiologists have responded to this challenge by insisting that individual investigators try, as much as possible, to insulate their scientific work from the realm of policy making. They urge that investigators should not be active in influencing policy when they are conducting conducting scientific research on a related topic. In other words, they should try to be neutral and disinterested towards their research conclusions. Thus, a researcher who advocates for legislation insuring that women pay for mammography should not conduct a meta-analysis or write a review of clinical trials on the effectiveness of mammography. By separating policy (and morality and politics) from science, the argument goes, objectivity will be enhanced. This idea has previously drawn support in other scientific disciplines that are prone to scientific controversy. For example, in the 1920s, when social scientists were accused of being political and unscientific, sociologist Max Weber responded by warning his colleagues: “It is the duty of the man of science to remain silent . . . on value questions upon which he is so freely encouraged to expound.”

However, others have responded that values cannot be isolated from the conduct of research, because many scientific theories and practices themselves are based on value judgments rather than empirical data and epidemiologists actually carry a professional obligation to participate in policy decisions. Historians and sociologists of science have marshalled much evidence to support this view, demonstrating how implicit individual prejudice can drive scientific debates.

Rigorous methods are frequently touted as the best means of preserving objectivity, but they are no panacea. The threat of the randomised controlled trial as the gold standard for testing new treatments was driven by a need for objective assessments of the claims of drug manufacturers. R A Fisher originally touted randomisation as the only means of generating an objective measurement of the risk of inferential error. However, as the mammography debate vividly illustrates, even a randomised controlled trial (or seven) is not sufficient to enforce objectivity. The methodological benefits of randomisation apply only to a single study and not to the synthesis of evidence from several studies. In contrast, the practice
Science, ethics, and professional public health practice

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Competing values and obligations

PUBLIC HEALTH PRACTICE

At the core of professional public health practice is a promise to help society by preventing disease and promoting health. Public health is a calling, as much an art as it is steeped in scientific theory, method, and evidence. We, the public health professionals, learn theory and practice in the classroom and hone them in experience. We define core values and embrace integrity, prudence, honesty, and trust. We develop standards of excellence and codes of ethics to guide our professional pursuits.1 2 Our practice is a complex blend of acquiring scientific knowledge and participatory policymaking. We study communities and individuals, the healthy as well as those who suffer from disease, injury, malnutrition, and untimely death. We recommend and advocate policies with others, for others, and for ourselves.

Ethics as an academic discipline and as a pragmatic dimension of our daily professional lives offers a conceptual framework and methods for thinking about and improving the practice of public health. Inevitably, we encounter situations marked by tension between competing values and obligations. Of the many problems that require attention, we choose three: evidence to action; the pitfalls and promise of public advocacy, and the balance between individual freedom and the common good.

EVIDENCE TO ACTION

The scientific knowledge that matters to public health interventions extends from the physical and biological sciences to epidemiology and on to the environmental, social and behavioural sciences. The problem of deciding when to act on the basis of that knowledge is as much synthetic as it is analytic. We collect evidence and use methods, both qualitative and quantitative, for its interpretation. We recommend actions in a climate buffeted about by politics, economics, and religious beliefs.

We cannot act in a vacuum. For primary prevention, we need to know something about how people are exposed, some semblance of a mechanism of action, how well the factor explains disease occurrence, and how that factor is connected to other determinants that make up the complex tapestry of causation. We need to know something about the expected changes in incidence, morbidity or mortality if the factor is removed, how much such interventions cost, the trade offs in risks and benefits, and how well such changes are welcomed by the public and their cultural institutions.

How much do we need to know? We rarely have the luxury of waiting for a complete understanding of causation. With every new shred of evidence we ask the question: now is it time to act? Sometimes the answer is obvious. Other times we swing back and forth on the pendulum of uncertainty. The scholarship of ethics suggests that such judgments are a product of circumstances—including the current scientific evidence—and ethical principles, obligations, guidelines, and maxims. The principles that guide such decisions are multiplying by the hour. Bioethics gave us four: non-maleficence, beneficence, respect for persons, and justice. Twelve so-called principles of public health ethics recently appeared.3 The precautionary principle suggests that actions should be taken when the evidence is somewhere below that of the unachievable levels of certainty or proof.4 But what is the least amount of evidence needed to warrant action to reduce risk, minimise harm, respect the autonomy of others, achieve justice, and maintain the public trust in our profession?

ADVOCACY, OBJECTIVITY, AND VALUES

As public health professionals we debate the pitfalls and promise of public advocacy. There are those in the profession who warn us away from advocacy in the hope that we can maintain an objective scientific neutrality. But science alone will not get the work of public health done and objectivity is less a characteristic of the scientist than it is the property of scientific methods. Besides, we are obliged to come to the aid of communities.5 Thoughtful, just, and reasoned advocacy is as much a part of our practice as is science.

Ultimately we seek balance between the dispassionate description of scientific findings and a persistent plea to use those findings for public health action. Call it finding a balance between the pursuit of truth for its own sake and solidarity with others for whom we advocate. Call it the balance between realism and pragmatism6 or between objectivity and subjectivity. Mix in the values that cut across science and its application. That is the second problem for public health professionals.

INDIVIDUAL FREEDOM AND THE COMMON GOOD

In mission and means, public health strives for healthy communities and for healthy individuals in communities.7 Historically, even when public health activities were directed to individuals (for example, immunisation), there was a
SUMMARY

Public health is a multidimensional entity: a complex of concepts and concrete institutions, both quest and practice, a desired goal and a present vocation. Its domain is extensive, stretching horizontally from providing preventive services as a safety net for individuals to promoting the health of communities, and vertically spanning policies, interventions, and research ranging from fundamental physiological processes to the social forces that change society.

It is inevitable that public health professionals will encounter, even engage, tension between competing values and obligations. We have suggested only three areas where tensions seem particularly pressing. The first is deciding when to act in public health, given a synthesis of the current knowledge gained from applying scientific methods to cells, individuals, communities, and society at large. It is a balancing act between what needs to be known and what needs to be done. The pursuit of scientific knowledge (value laden as it is) and the dedicated application of what we know to achieve ends we value. Both are mutually reinforcing and potentially in conflict. Discerning and maintaining the proper balance, especially in the face of diverse personal and public values and political adversity, is the second of our challenges.

The third problem requires us to determine when and whether the presumed goods of promoting health and preventing disease justify constraints on fundamental rights, and to balance closely held individual values of self-determination, privacy, and freedom with community values and wellbeing.

We have not proposed how these three areas of ethical tension are resolved. Indeed their resolution in specific cases is the very stuff of ethical reasoning.

What we have hoped to show is that our fundamental commitments as public health professionals impose upon us ethical dilemmas unique to our calling.

REFERENCES

place of the “patient”, who requires the proximity of his/her pair—the “physician”—to be defined as such. Curiously, with the increasing distance in the patient-physician relationship, the patient’s place has received new designations. In our opinion, some of these terms are appropriate, including “medical ends user.” Other terms are clearly up for discussion because of their premises, like “e-health consumer” and “consumer health informatics”.

The contemporary world imposes a difficult test on all those who intend to join it, by demonstrating adherence and the economic capacity to accompany the unceasing, whimsical trends of market consumerism. Those who fail the test and cannot afford or are denied access to the necessary commodities become non-consumers or consumer failures. They are incapable of becoming “free”, autonomous individuals. Huge contingents of the world’s population live and die in such conditions.

Still, the dominant perspective in the sociopolitical and economic road to be trod by the peoples of the planet seems to be towards globalisation, pursued by monopolistic capitalism in its various facets (especially in the so called post-industrial society, where the services and knowledge production sectors now enjoy economic hegemony).

Health promotion policies place a clear emphasis on autonomy. In general, when dealing with the notion of personal autonomy, we should be clear about the individualist premises underlying the prevailing concept of “person” in the West. Such premises become manifest in self-care proposals. There are currently various forms of “healthy self building” through physical activities aimed at risk avoidance (for example, cardiovascular risk), aesthetics (production of a personal appearance with standardised forms of physical beauty and bodily attractiveness), rehabilitation (for the very obese or the middle aged who seek to “feel more fit”), and performance (athletic contests, both amateur and temporary times).

Certain aspects pertaining to the “fitness” concept deserve special attention. Without going into the distinctions and imbrications between this concept and that of good health, let us highlight how pertinent it is that health care is also linked to the idea of feeling fit, or feeling physically well disposed.

Subjective feelings of possessing fitness by way of disciplined training are especially important. The quest for fitness imposes a state of frequent and endless training, with ever present self scrutiny and a nagging anxiety over the possibility of “getting out of shape”.

Countless men and women currently attempt to keep their bodies ready/adjusted/in shape to deal with overly hazardous events like rallies, resistance trials in wilderness areas, x-games, etc. And they also attempt to “keep in shape” for the potentially unexpected events provided by multiple possibilities from a veritable menu of adventures in contemporary life. Yet they must also obviously have the wherewithal to consume the goods and services needed to engage in such activities.

The spread of ideas involving self care and health promotion behaviour is in full swing. A rapid search on Medline will identify a deluge of articles on this issue under the various (and not always clear) terminology that goes with it. This perception was highlighted by Kulbok and associates: they referred to the reigning confusion in the self care/health promotion field. After a critical analysis of the terminology, these authors highlighted the different meanings in the ways by which specialist use ideas and concepts inherent to the area: “health promotion”, “health promotion behaviour”, “health protection behaviour”, “disease prevention behaviour”, “preventive health behaviour”, “healthy behaviour”, and “healthy life style”.

New ethical issues and sociocultural repercussions have emerged in both the health care and self care fields through mediations between the electronic media, health professionals, and users, who in turn become agents of consumption. Still, what is to be done with the vast multitude of non-consumers surviving precariously on this planet? How does one deal with issues at the societal level: poverty, inequality, and nutritional and housing deficiencies and their repercussions on illness, violence, crime, and exclusion? Such problems have the same roots as our inability to establish realistic values that are pertinent to living in contemporary times.

Critics of bioethics contend that it is impossible to apply the principles of freedom and autonomy from classic principlist bioethics as instruments for moral conflict mediation in societal settings with great socioeconomic inequality, where non-consumers also tend to be more vulnerable in terms of health. In this sense the agenda of a global bioethics should also include the analysis, discussion, and development of strategies to deal with social inequalities and the vulnerability of populations that are excluded as agents of consumption. This implies not allowing health related decisions to be made primarily at the private level.

Faced with the question of whether traditional principlist bioethics is sufficient to deal not only with moral dilemmas but also with public health problems, unfortunately our answer cannot be categorically affirmative. We face the important challenge of taking lack of autonomy into consideration as a crucial aspect for building a “public health ethic”. Although it may sound like a truism, such an ethic should also include old public health issues on its agenda. It should provide the basic conditions for autonomy (and citizenship) to populations excluded from the markets and consumption through a struggle against the contemporary causes of destitution and ignorance. In particular, it should seek ways to reduce the vulnerability and deep inequalities in the distribution of means for protection, treatment, and rehabilitation of the vast multitudes within the context of the heavy side effects of socioeconomic inequities in the “globalising” economic trends dictated by economically stronger nations. The moral arguments emerging from the confrontation between the large pharmaceutical industry and representatives of economically weaker nations (whether from the public sector or civil society) that resulted in breaking patents on antiretroviral drugs to make them more accessible to people with HIV undeniably belongs to the field of public health ethics.