Towards a philosophy of public health

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A few years ago, readers of the journal were invited to discuss the philosophical foundations of public health. This invitation was accompanied—indeed, justified—by the claim that the disclosure of philosophical perspectives is essential to the resolution of complex issues in public health. The authors, Nijhuis and Van der Maesen, argued that debates about the pros and cons of public health approaches are often confined to the methodological scientific level, thereby neglecting implicit ontological notions lying behind and presumably influencing the arguments and decisions flowing from them. Consistent with this view is the idea that public health experts would make better choices if only they would disclose their ontological orientation towards the paired notions of “public” and “health.” Four categories of ontological interpretations of public health were offered: two “public” categories and two “health” categories. “Public” category no 1 emphasises the individual. In this view, the public is primarily comprised of the actions and motives of discrete individuals. “Public” category no 2, on the other hand, emphasises the collective over the individual. In this view, the public is primarily conceived as populations within social, economic, and political systems. “Health” category no 3 is a mechanistic view that emphasises the traditional medical distinction between disease and non-disease in the individual, whereas category no 4 views health as the degree to which an individual reaches an equilibrium state with somatic, psychological, and social influences. Much more could be said about these four categories. Categories no 1 and no 2, for example, could be retrofitted into Rose’s classic paper on sick individuals and sick populations. And Nijhuis and Van der Maesen’s idea that most scientific work in epidemiology emphasises categories no 1 and no 3 whereas most health policy work emphasises categories no 2 and no 4 may reflect the considerable distance between the current practice of epidemiology and the practice of public health that has recently attracted so much attention. In any case, it is reasonable to consider how revealing one’s commitment to these various categories influences practical public health decision making.

Perhaps to encourage further discussion, Nijhuis and Van der Maesen did not illustrate their “disclosure claim” with an example of how a public health decision was made better (or a complex debate resolved) by revealing the ontological orientation(s) of the decision-maker. Yet many potential examples exist. Consider the general situation for public health interventions in which the benefit to the individual is small relative to the benefit to society. Applying a strict interpretation of Nijhuis and Van der Maesen’s claim to this situation would make decision making about the appropriateness of the intervention reasonably straightforward. An intervention is less likely to be advocated if the primary ontological orientation of the decision maker is with the individual—category no 1—than if the decision maker is committed to the population—category no 2.

“Real life” public health decisions are obviously much more complex, no more isolated matters of ontology than they are isolated matters of scientific methodology. Indeed, there is probably more to the philosophical basis of decisions than ontological concerns alone. This disclosure claim, therefore, can and should be expanded to include other types of philosophical commitments. Ethics seems particularly relevant in as much as public health decisions often entail balancing benefits and risks to individuals and to society, an initial condition for the decision described above. Public health decisions may also be affected by the practitioner’s orientations in the shifting philosophical sands of the practice of science.

In this paper, I expand the “disclosure claim” of Nijhuis and Van der Maesen to include not only ontological but also ethical and epistemological perspectives. Brief definitions of these three philosophical categories may assist the reader who is unfamiliar with this territory.

Ontology
Involves the nature or essence of reality, of being and existence.

Ethics
Involves the nature of rightness and the study of what actions are right actions.

Epistemology
Involves the study of how knowledge is gained, and the general validity of claims to knowledge.

Each is a highly developed theoretical discipline in its own right and each can also be connected to (that is, applied to) issues within the theory and practice of public health. For example, Nijhuis and Van der Maesen’s four ontological interpretations relate to the nature of the essential concepts of “public” and “health.” The nature of causation is also an ontological concern, but how knowledge is gained about causation is an epistemological concern. Epistemological commitments to forms of logic (for example, induction or deduction) or to other theoretical approaches may influence how scientific evidence that tests causal hypotheses is examined or interpreted.
Ethics, the study of what constitutes right actions, also has its theoretical side, but may be the most “applied” of these philosophical categories. Making and justifying decisions about what ought to be done in particular situations (that is, cases) is a familiar application of ethics to public health practice.

In the examples that follow, it will become clear that these different philosophical arenas are not easily separated from one another. Independence is more a matter of theoretical than practical interest in everyday decision making. It follows that the future practice of public health decision making may benefit from a mix of ontological, ethical, and epistemological perspectives informed by a general philosophy of public health. This paper will not go so far as to propose such a general theory yet may help set the stage for such an effort. I begin with an example of public health decision making affected by a mix of ontological and ethical perspectives. In the second example, I add epistemological concerns. Clearly there is not room to explore deeply these philosophical roots. My purpose is rather to draw attention to several aspects of practical decision making that, as will be shown below, cannot be easily dismissed.

**Ontology and ethics**

In public health decision making, ontological and ethical perspectives are intertwined. Consider cancer screening as a prototypical example (and assume that the efficacy of the screening test and its side effects are known—that is, not controversial—thereby excluding epistemological concerns from the analysis). Implementation of a mass screening programme, for example, breast cancer screening by mammography or cholesterol screening for heart disease, entails wide public promotion of an intervention. At first glance, two ontological categories proposed by Nijhuis and Van der Maesen undergird the public health decision to go forward with such programmes: the population (as a collective), category no 2, and health promotion, category no 4.

These two perspectives, however, are insufficient to make decisions about the appropriateness of an intervention, even in those circumstances in which efficacy is accepted. There are concerns about the trade off between benefits to the population (the collective) and risks (that is, harms) to the individuals comprising the population. As Rose has argued, individuals typically fare less well than the population for prevention programmes; he dubbed this situation the “prevention paradox.” There are also concerns about the extent to which individual decisions to undergo the intervention are autonomous—that is, not coerced.

Bringing “individuals” into the discussion, however, signals the need to consider ontological category no 1, the gestalt of individuals. Similarly, bringing “risk” into the discussion is a direct reflection of the natural scientific or mechanistic notion of health (category no 3) in as much as cancer screening involves side effects to individuals such as direct injury from the screening test itself, or residual physical trauma from treatment that may be needed if a positive diagnosis is made. In prostate cancer screening, for example, incontinence and impotence are important risks of treatment. Yet “risk” could also reflect a more holistic notion of health (category no 4) because it involves psychological trauma such as anxiety, an increasingly recognised issue in cancer screening.7

It seems, therefore, that there are choices to be made regarding specific public health interventions—here, cancer screening tests—wherein all four ontological interpretations found in Nijhuis and Van der Maesen’s paper are relevant if not often explicitly labelled as philosophical perspectives in themselves. How to make the best decisions in a given situation, especially given the inherent dilemmas such as the prevention paradox, must therefore entail something more than ontology alone, as we suspected from the earlier and simpler example. There is a need to balance the individual’s perspective, emphasising autonomy, self determination, and safety against the perspective of the collective with its responsibility to intervene to increase the overall health of its citizenry5 for the common good. This balancing, however, largely takes place in ethical terms. Rules to “help others” and to “prevent harm to others” are involved and are derived from the general principle of beneficence. Also involved is the principle of respect for persons that is made manifest when information regarding potential risks and benefits10 is provided so that individuals may decide for themselves whether to participate or not in a screening programme. Thus, in practical public health decision making, a combination of ontological categories and ethical constructs are important, consistent with but expanding upon Nijhuis and Van der Maesen’s claim.

The expansion of the disclosure claim, however, does not deny the importance of a basic distinction between the individual and the population in public health. Legislatively mandated public health programmes, for example, highlight this stark contrast. Seat belt laws, immunisation requirements, and mandatory reporting of sexually transmitted diseases are three American examples; each involves considerable restraints on the individual for the benefit of the population. Nevertheless, the justification for these actions—which are enforceable by the power of the state, and which are tax supported and intrusive—involves more than ontology. Cole, for example, argues that commonweal, or “doing the greatest good for the greatest number” is the justification for abrogating the rights of individuals.11 Pellegrino12 provides the view that legislated public health interventions almost always involve a trade off between commitments to general and widely used (if somewhat battered) ethical principles; typically, a paternalistic beneficence is enlisted to limit personal autonomy. Last14 also offers beneficence as a dominant ethical principle in public health but balances it with a respect for the autonomy of people. He notes that it is important to provide them with sufficient information to empower
them to do what they can to promote health rather than coercing them to stop doing what they find pleasurable or to start doing what they find unacceptable.

Ontology, ethics, and epistemology

Epistemological perspectives also lie unrecognised and undisclosed in the philosophical closets of public health decision makers. Consider, for example, the idea that health promotion for the benefit of populations should be based on solid evidence of efficacy. Brought together in this statement are underlying and implicit epistemological commitments, for example, the nature of evidence and its relation to the hypotheses being tested, the meaning of “solid” and “efficacy” as well as ontological concerns (for example, populations) and ethical perspectives (for example, benefit). In keeping with the idea that commitments to any one or a combination of these three philosophical arenas can affect public health decision making, I will show how different published opinions about an important public health issue are at least consistent with different perspectives, some ontological, some ethical, and some epistemological. In most instances, the decision maker’s awareness of the philosophical underpinnings of his individual decision (and whether he would accept Nijhuis and Van der Maesen’s disclosure claim) is a matter for speculation. In a few papers, on the other hand, decision makers have identified philosophical constructs, evidence that Nijhuis and Van der Maesen’s claim has some (perhaps unwitting) advocates in practice.

The example is cancer screening with prostate specific antigen (PSA). Stark differences exist among organisations and among individual commentators regarding whether PSA screening should be recommended. For the purposes of this paper, I examine only published individual decisions, leaving institutional decision making for another time. There is no shortage of examples. In 1993, the British Medical Journal published an editorial in which the author—Schroder—states that “....(PSA) screening should not be recommended....” The same year, Urology published an editorial in which the author—Catalona—presents precisely the opposite opinion, that “....screening with an annual rectal examination and serum PSA measurement should be encouraged....” Four years later, the situation has not changed much. Just one example is a pair of comments appearing in the European Journal of Cancer, one arguing against and another arguing for the “gold standard” status of prostate cancer screening.

What philosophical orientations, whether epistemological, ethical, or ontological, underlie these very different opinions? Answering this question will require a careful examination of the exact wording from decision makers, inferring from them what philosophical commitments are, if not explicitly stated, then are at least consistent with those opinions. I begin with epistemological concerns, because all current published opinions have appeared while randomised trials of PSA screening and a randomised trial of surgery compared with expectant management for localised disease have been undertaken. No trial has been completed. It follows that all published recommendations (to screen or not) either deny or accept the need for a strong (RCT) test of the hypothesis that PSA screening reduces mortality. Put in the language of the expanded “disclosure claim,” none of these decisions are made without at least an implicit epistemological position about the need for randomised trial evidence. Four published opinions follow:

Schroder, the 1993 British Medical Journal editorialist, makes clear his perspective when he writes that “....(PSA) screening should not be recommended as public health policy until clear benefit in terms of reduced mortality from cancer can be shown in prospective screening studies....” Voss, a 1994 editorialist in the Journal of General Internal Medicine provides a similar view when he writes that annual PSA tests are “not warranted by the available evidence.” In both instances, there is a clear epistemologically oriented criterion: no positive trial results means no screening recommendation. Others differ in their evidentiary threshold. Catalona, who wrote the 1993 Urology editorial mentioned earlier, notes that “the National Cancer Institute is conducting a prospective randomised trial to determine whether or not screening reduces the prostate cancer mortality rate, but it will take sixteen years to complete the study. It is estimated that half a million men will die of prostate cancer before this study is completed, and it is unreasonable to expect clinicians to refrain from PSA testing in the meantime.” More recently, an Annals of Internal Medicine editorialist—Middleton—justifies his decision to support screening with the following: “we do not know whether our efforts will ultimately reduce mortality related to prostate cancer, but we can be hopeful.” He cites SEER data showing a decrease in the incidence of new cases of metastatic disease.

These examples reveal something about the methodological requirements of these decision
makers but nothing about their epistemological commitments in themselves. According to Vineis, however, different epistemological perspectives may lead to different methodological requirements. He believes that an empiricist philosophy requires randomised trial evidence whereas other philosophical perspectives may only require mechanistic evidence.25 The relations between epistemological commitments and methodological choices or evidentiary thresholds is a rich area for further exploration. What, for example, are the epistemological roots of the hierarchies of study design so often promoted in evidence-based medicine and evidence-based public health? What are the epistemological roots of causal criteria?4 While specific answers to these sorts of questions are beyond the scope of this paper, it is nevertheless reasonable to suppose that the different decisions about PSA screening could be affected by different epistemological perspectives made manifest in different methodological requirements.

Unfortunately, no such epistemological commitments are outlined. And, it is probably too simplistic to argue that methodological commitments (or their roots in different epistemological frameworks if we knew them) completely determine the opinions held about PSA screening. In each example, there are implicit ethical notions and there are also suggestions of ontological commitments along the lines suggested by Nijhuis and Van der Maesen. Schroder, for example, notes that screening should not be recommended “as public health policy.”16 Catalona, as noted above, recommends that clinicians should not refrain from PSA testing but says nothing about public health screening programmes outside the context of the physician-patient relationship.17 Voss23 notes that annual PSA tests are not warranted for asymptomatic patients. Middleton24 subscribes to the ACS recommendation that a man older than 50 years should have a PSA test and that recommendation clearly states that it is not intended as a guideline for public health policy.25

It seems, therefore, that a careful examination of published opinions on the appropriate-ness of PSA screening shows not only subtle commitments to epistemological and ontological frameworks but also somewhat less contentiousness than what was apparent on the surface. These published opinions appear to collect into two camps: those who argue against mass public screening programmes and those who argue for screening as a part of routine clinical practice.

The PSA controversy has also included a few papers in which explicit attention has been paid to philosophical perspectives. Chodak, for example, in a relatively early paper,12 notes that the lack of scientific evidence in support of a mortality reduction (an epistemological claim) does not satisfy the Hippocratic tradition of non-maleficence because screening harms a significant number of men (an ethical claim). He notes that screening asymptomatic men refers to both public programmes and to situa-

tions involving individual patients (an ontological claim). He concludes that the most balanced approach does not involve making strong recommendations for mass screening. For patients, he recommends discussing the nature of the controversy and the potential advantages and disadvantages of screening and treatment, allowing the patient to help make the decision to screen or not. Woolf28 29 as well as Hahn and Roberts30 also support this “informed consent” model for asymptomatic men who present at a physician’s office. Interestingly, a recent study has shown that men informed about PSA testing are less interested in undergoing those tests than controls.31

Conclusion
So it seems that published decisions about the appropriateness of PSA screening have at their heart issues of epistemology, ethics, and ontol-ogy. I doubt that other public health interven-tions are different. Each such decision is a matter of concern (call it respect) for the person or population involved (a combination of ontological and ethical concerns) and it is about the benefits and risks of intervention (a combination of ethical and epistemological concerns, and according to Nijhuis and Van der Maesen, also a matter of ontology). The extent to which decision makers provide their philosophical orientation appears to assist us in understanding their decision; it remains an unresolved and more difficult issue whether decisions consistent with Nijhuis and Van der Maesen’s “disclosure claim” are in fact better decisions. To make matters even more com-plex, decisions about interventions are affected by more than philosophical perspectives. There are powerful economic interests for researchers and for medical practitioners alike. Aronowitz recently argues that extensive investments of research money give researchers as much gain by not recommending PSA screening as physicians have to gain through reimburse-ments by recommending it.32 Along these lines, it is interesting to note that Schroder19 (who does not support public screening) is the prin-cipal investigator of a large trial testing PSA screening. Catalona20 and Middleton24 are both practising urologists and both advocate screening for asymptomatic men. Without frank disclosure on the part of these individuals, it is not clear if these are compelling interests or not. What is clear is that Nijhuis and Van der Maesen’s disclosure claim will require further expansion beyond philosophical foundations to include economic interests, political ideologies, and other social forces. Aronowitz sums it up: “…the best recommendations will be those that take into account the complex, philosophical, social and political forces that shape our response to specific health issues.”32

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Although we cannot clearly demonstrate that decisions about public health interventions would have been better had philosophical per-spectives been made explicit, the claim is con-sistent with research that has shown that, for ethics, the more you know the more likely you
are to make ethically appropriate decisions.32–34. It follows that in order for these philosophical perspectives to be recognised by public health decision makers, to be made explicit, and to make a difference in practical decision making, they should be incorporated into the formal training and career development programmes of the public health professions.4

Not everyone agrees that philosophy can be taught nor that it is relevant to practice. Schlesinger, for example, writes that scientists’ problem solving skills are not likely to be improved by studying the philosophy of science. He suggests that inspiration and a type of mental exhalation are the primary benefits of philosophical inquiry.39 And perhaps the interest epidemiologists and other public health practitioners have expressed in philosophical issues in the past two decades can be explained in such terms, although I find more satisfying the notion that by describing the ontological nature of causal (and other types of) hypotheses, the epistemological framework for testing those hypotheses, and the ethical foundation for applying that knowledge we will be rewarded with a better understanding and perhaps even justifications for the difficult decisions we make in the practice of public health. In the absence of such an effort, we are left with the important but poorly characterised constructs of common sense and judgement, a host of contradictory decisions in daily practice,16 and the interesting but slippery problem of inferring from those decisions the corresponding philosophical commitments. Although we may conclude that differences in public health decision making can be explained as a matter of different (and sometimes non-negotiable) values, some scientific and some extrascientific,36 disclosure of philosophical commitments or “values” remains the key concern, as some philosophers who see science as social knowledge have emphasised.24

If public health professionals embrace the need to examine and proclaim their philosophical foundations, then a guide encompassing ethics, ontology, and epistemology is needed. Call it a guide to the philosophy of public health.

No such document exists. More work has been done on the ethics of public health44 including the ethics of screening45–48 than on ontology, a fact that may have spurred Nijhuis and Van der Maesen’s appeal to that part of philosophy.1 There has, however, been a recent discussion of the role of systems theory in conceptualising the nature of epistemological studies.49 Epistemological concerns, at least in epidemiology, have been discussed for 20 years.14

How might such a philosophy of public health arise? If efforts in the philosophy of medicine are a reasonable template, then we should expect a gradual transition through three phases.49 In the first phase, what could be called philosophy and public health, practitioners would use a philosophical idea to illuminate an issue in the practice of public health. Philosophers in turn would use a public health problem to illustrate some aspect of philosophy. The two disciplines would remain distinct and the analyses superficial. In the second phase, called philosophy in public health, more formal philosophical analyses would be applied to problems that comprise the “matter” of public health. This phase best represents some efforts in public health and epidemiology; the two decades long debate about the utility of Popperian philosophy comes to mind as well as the extensive efforts to examine the ethics of public health and epidemiology. In the final phase, a philosophy of public health would consist of a general theory of public health within which the problems examined and solutions proposed in the previous phases would be incorporated and synthesised. The types of problems could be ontological, ethical, and epistemological as described in this paper. These problem-solutions in turn would provide the public health practitioner with a foundation for philosophical perspectives that presumably underlie and influence daily public health decision making.

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