Diet and physical activity are of immense importance to primary and secondary prevention of coronary heart disease (CHD). However, information relating to these activities at the individual and population level remains scant, with opportunities to alter the situation overlooked continuously. A recent independent review in the UK was highly critical of existing information systems to assess needs and plan service provision. Others are critical too in relation to CHD specifically. We argue that the existing poor information infrastructure relating to diet, physical activity and other variables associated with a person’s way of life, known to influence CHD, could be improved by utilising disease registers. We put forward key arguments why this may not be happening and encourage practitioners and researchers to help overcome such barriers.

Disease-specific registers are increasingly being developed in countries such as the UK in response to national quality frameworks. General practitioners in the UK are given financial incentives to establish administrative systems to identify patients with and at significant risk for CHD, with the purpose of facilitating the systematic delivery of care and regular patient follow-up. Earlier work suggests that disease-specific registers can provide good-quality information relating to individual patients as well as for monitoring and service planning. It has been argued that well-administered CHD registers enable patient profiling and patterns of care to be examined within individuals or groups of general practices. Yet it remains to be seen if CHD and other registers lead to enhanced patient care.

The possible benefits of CHD registers extend beyond individual patient care and that delivered by a single care provider. Advances in technology make it feasible to develop single registers within a particular administrative district, for example, with primary and secondary care providers linking ‘real-time’ information, regardless of their place of work. Not only would this allow patterns of care to be monitored between providers and over time, it would also create a unified longitudinal dataset with which to examine the often complex relationships between patient and clinical/treatment variables and changes in outcomes over time. This analysis can be missing from clinical trials because of short follow-up times, focus on a single primary outcome measure, or because trial participants are not always representative of the general population. Information on variations in inputs and their effect on patient care also provides analysis from ‘natural experiments’, again argued to be an untapped resource.

Concerns remain about using data from CHD registers for research purposes, alongside missed opportunities for individual patient care. Many CHD registers focus on optimal prescription of CHD drugs, which are underpinned by national government quality indicators. However, these indicators provide no incentive to adequately record variables associated with CHD and a person’s way of life, which is usually limited to assessing smoking behaviour. This is at odds with the evidence base showing a strong impact of diet and physical activity on the total population burden of CHD and with guidelines for preventing and treating this disease.

Thus clinicians are currently denied systematic methods for recording and reviewing patient information relating to diet and physical activity during a patient consultation, despite its potential contribution to reducing their risk of future events or their recovery following an event. Why are clinicians expected to measure and record blood pressures over time in patients with or at risk of CHD, yet not do the same for diet and physical activity? This absence of information also poses problems to the research community. Failing to measure and account for potential confounding variables, of which diet and physical activity are, will seriously (and rightly so) reduce the validity of research findings based on CHD registry data. It is, in effect, as serious as a longitudinal analysis of β-blockers ignoring blood pressure in its analysis.

Why does the problem remain?

If diet and physical activity are so important to the primary and secondary prevention of CHD, then why does this information remain absent from CHD registers? We put forward two related arguments. The first rests with clinicians. We suggest that some clinicians, following a clinically dominated training, remain unaware of the true strength of the impact of diet and physical activity on the course of CHD. While attention will probably be given to cholesterol for example, this still fails to consider wider dietary components and their impact on CHD.
This scenario is perhaps reinforced through pharmaceutical marketing techniques, diverting attention from interventions to change a patient’s way of life where this impacts on their illness. Others may feel lacking in skills to address such issues, and some could be daunted by what they see as an impossible task in supporting appropriate dietary changes and increases in physical activity by their patients.

A second but related issue refers to the availability (or lack) of suitable methods or instruments to measure diet and physical activity systematically within the constraints of a general practice consultation and then to record this easily on the patient register. Few simple instruments exist to measure dietary intake, with most designed to examine nutrient–disease relationships. We argue this level of detail is not necessary within the context of CHD registers. However, information would be of value to clinicians, service planners and researchers alike if it identified the extent that patients followed key dietary recommendations in relation to CHD protection. Short-form tools are available to assess physical activity, although not validated in this context, but still remain absent from CHD and other disease registers14.

We accept that complex issues need to be resolved in developing a short instrument to collect and record this type of information within the confines of a patient consultation. However, it is not without such possibilities15 although existing tools are not suitable for this task16. We do not accept that complexity is a good enough argument to continue to ignore diet and physical activity in CHD registers, given their importance in the primary and secondary prevention of CHD. No doubt many sceptics prevailed when short-form mental health instruments were mooted. Yet now a number of short instruments are available to collect robust information on dimensions of mental health and are used successfully in patient consultations17.

CHD registers offer hope in improving individual patient care and in addressing local and national information gaps about this dominant chronic disease. However, registers that continue to ignore the contribution of diet and physical activity to the primary and secondary prevention of CHD will only reinforce traditional health-care practice, and do nothing to encourage clinicians to pay attention to patients’ wider way of life and its impact on their illness. Moreover, doubt will remain about the validity of research findings from CHD registers. This diminishes the potential of the registers as tools of population health surveillance until the effect confounding from diet and physical activity can be assessed alongside other routinely collected data. The challenge to the research community is to now develop such short-form assessment methods, and to clinicians, to incorporate these as part of the patient consultation. It then remains to be seen if a CHD register, capable of prompting and storing all information appropriate to the care of individual patients, leads to significant improvements in outcomes over time. Without this, the cost and effort involved in managing any register is futile.

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References
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