Big Data and Population Health: Focusing on the Health Impacts of the Social, Physical, and Economic Environment

(Short title: “Big Data and Population Health”)

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Introduction

The growing abundance of data on the factors that produce health, and the capacity to link these to data from individuals, hold extraordinary promise for improving population health. Although public health is built on a long history of creatively using health data going back to the pioneering work of John Snow and William Farr\(^1\),\(^2\), the full potential of contemporary rich data sources is not being fully realised. Instead, the impact of this revolution is being seen mainly in the burgeoning precision medicine agenda, globally, and through the Precision Medicine Initiative (PMI) in the U.S.A.\(^3\) In that context, linkage of omics data to phenotypic information from health records is being exploited as new research platforms, with the hope that this will transform clinical practice. Similar momentum has not yet been achieved for the population health sciences.

We propose that there are two challenges for the population health agenda. First, ensuring that we make better use of existing data, and in some cases, enhance data linkage and methods to do this. Second, in extending our efforts from the individual to the population by exploiting new complex, and sometimes unstructured, data sources.

We recognize that there are major technological challenges when dealing with the massive and rapid flows of data that come from both traditional data sources, such as large administrative databases, and new sources, such as genomics, land-use, neighborhood and climate data, and unstructured social media feeds. However, we argue that the more important challenge is conceptual and perhaps ideological. How do we identify the data that matter most to improve health for the whole population? In some cases, these are existing data and we need to determine how to enrich, link, and analyze these routinely collected data to maximize their value. We also need to think about new sources of big data that can improve our understanding of health, and how we better integrate these in our approaches to studying population health while avoiding the temptation to analyze everything that may
present itself as a new opportunity. Distinct from personalized medicine, the ‘what matters most’ question can be addressed through an understanding of the pressing health challenges of our time and an understanding of how factors across economic, social, behavioral, and biological domains may interact.

The past two decades have shown how factors at multiple ‘levels of influence’ are associated with both individual and population health. Multilevel causal frameworks suggest that policies, features of institutions, characteristics of neighborhoods and communities, living conditions, and social relationships all contribute, together with individual behaviors and individual factors such as genotype poverty and race, to the production of health. For example, quality of the built environment has been shown to be associated with mental ill health and diabetes. Certain social network characteristics are associated with the risk of obesity. It is a logical extension of the goals of big data collection to introduce measures that can capture potential risks at multiple levels of influence. The feature that turns these into “big data” is the coverage and complexity of this information, which enables the population health approach, across geographies, time, and the life course.

Examples of the value of population data linkage and the potential contributions of big data

Two recent papers illustrate the importance of linking health, ethnicity, and personal financial data. Chetty and colleagues linked personal taxation and social security data of the US population to study the independent effects of income and place on life expectancy. This entailed analyses of 1.4 billion person–years of observations (“big data”). Being poor in Detroit is more hazardous to health than being similarly disadvantaged in New York or San Francisco. Case and Deaton documented alarming rises in death rates in middle-aged non-Hispanic white males and females after the global financial crisis. Both studies show that
physical and social factors interact strongly to mitigate or enhance the effects of poverty. Even partial mitigation of these effects could be life-saving on a scale that would compare well with any impact of precision medicine interventions.

Consider how the value of such studies would be enhanced by more complex data that enable longitudinal analyses to determine the effects of migration (e.g., moving from Detroit to New York), the role of behavioral factors, genomics and epigenomic modifications, and access to healthcare, etc. This would enable the study of gene/environment interactions that determine health, the biologic changes induced by stresses associated with poverty and marginalization, the role of health systems, and mediating factors that could potentially afford new opportunities for disease prevention. There are a growing number of centers where such linkages and studies will be possible. One notable example is the Big Data Institute at the University of Oxford, where data from the UK biobank can be linked to administrative data and electronic health records (www.bdi.ox.ac.uk).

These types of data and research can also inform health services planning and delivery. Just as genomic and phenotypic data can predict an individual’s response to treatment, linked socioeconomic, environmental, and health services data can be used to predict population risk for certain diseases. Examples include a population diabetes risk prediction tool for policy makers, and evidence that the development of chronic disease in low socio-economic circumstances identifies individuals who will become very high cost users of the healthcare system.\textsuperscript{14,15}

Linkage with data on education, crime, and occupations can also help elucidate the wider impacts of health status. Such linkage and longitudinal follow-up enabled researchers in Manitoba to document, among teen mothers, the full negative impact of their health status on the educational attainment of their offspring.\textsuperscript{16} In British Columbia researchers used routinely collected data from the Early Development Instrument (EDI) to measure the
negative impact of early childhood vulnerability (a composite of physical health, social competence, emotional maturity, language and cognitive development, communication skills, and general knowledge in the majority language and culture) on school achievement, standardized test scores, and criminality.\textsuperscript{17}

With regards to environmental health, the availability of repeated, increasingly detailed, and geospatially mapped measures of air pollution has allowed studies linking living near major roads and/or exposure to fine particulate matter and other air toxics with cardiovascular disease, diabetes, autism and, very recently, neurodegenerative diseases\textsuperscript{18,19} Linkage with social data, which currently occurs rarely, would help researchers determine, for example, the environmental contribution to the health risks of living in high-poverty areas.

**Prioritizing data from marginalized populations**

Finally, we highlight the current disconnect between big data for health agendas and the health of marginalized populations. Undocumented immigrants, migrant workers, the homeless, and indigenous populations have some of the worst health outcomes in societies.\textsuperscript{20} This is in contrast to documented immigrants who typically have better health than resident populations.\textsuperscript{21} Marginalized groups should benefit from ‘big data’ efforts that identify new targets for prevention and patterns of health care utilization and inform studies of the epigenomic and other biologic impacts of poverty and social exclusion. These populations are rarely included in large cohort studies; however, advances in electronic medical records, data linkage and data security, and socialization, make it possible to track and document the health experience of individuals who may be migratory or homeless while protecting their privacy. As an example with respect to indigenous populations, the administratively linked data on over 13.5 million Ontarians held at the Institute for Clinical Evaluative Sciences which is enabling an accelerated “big data for population health” initiative now includes data
on over 200,000 First Nations individuals. This was made possible by especially close collaboration with and the approval of indigenous communities and their associated governance bodies, a strategy and principle that must extend to research on other disadvantaged groups.

**Recommendations**

At a time when there is so much interest and activity in big data analytics and precision medicine, allowing such efforts to diverge from the interests of population health is unwise. Rather, we have an opportunity to use the momentum that has been created by this movement to do two things: first, to enhance our uses of large population data-sets to gain a better understanding of the determinants of health and health inequities to support policies that will optimize health, and second, to push for more complex data that more closely reflects a person’s context from the social, demographic, economic, and biologic perspectives over time. In so doing we need to be strategic about how we prioritize data efforts and achieve consensus on where to start to build the momentum to a level that is at least as high as the precision medicine realm. Our challenge is not so much employing all types of data and methods, as avoiding the temptation to do everything.

We identify 5 priorities:

1) Data on socio-economic status must be improved and linked to health services and health outcomes data. It is possible to securely link income tax data if revenue agencies can be assured that client privacy will not be breached. In keeping with the big data agenda, as much as possible these data should be longitudinal, reflect the dynamic and contextual nature of socio-economic status and environmental factors.

2) We must strive to include data that identify the most vulnerable members of the population, including indigenous peoples, migrants, refugees, and the homeless. Governance of the use of such data must involve those affected.
3) We should accelerate the shift from ecological studies by securing linkage of data at the level of the individual, which enables analyses at different levels, including micro- and macro-environments, such as occupational and community-level exposures to pollution; neighborhood walkability, “food desert” measures, occupational data, and social isolation among others.

4) Collaborations are needed with: (a) researchers in the fields of education, environment, and social sciences to ensure the validity and accuracy of multi-level data and (b) decision-makers in the public health and healthcare sectors to ensure that research questions and findings are those likely to have maximal impact on health.

5) All reasonable efforts must be made to protect privacy. Analyses should use de-identified data in facilities that provide the necessary combination of policies, staff training, physical, and electronic security.

It is only by addressing these priorities that we can assure that “Big Data” initiatives that pertain to health are not limited to the relatively narrow allopathic goals of the precision medicine agenda. By pursuing, instead, a more expansive “Big Data for Population Health” vision, initiatives so aligned carry the promise of generating insights on the drivers of health that can lead to interventions and policies that promote health on a population scale: the very core mission of public health.
Reference List


