Sharing health data: good intentions are not enough
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Abstract
Epidemiologists and public health researchers are moving very slowly in the data sharing revolution, and agencies that maintain global health databases are reluctant to share data too. Once investments in infrastructure have been made, recycling and combining data provide access to maximum knowledge for minimal additional cost. By refusing to share data, researchers are slowing progress towards reducing illness and death and are denying a public good to taxpayers who support most of the research.

Funders of public health research are beginning to call for change and developing data sharing policies. However they are not yet adequately addressing the obstacles that underpin the failure to share data. These include professional structures that reward publication of analysis but not of data, and funding streams and career paths that continue to undervalue critical data management work. Practical issues need to be sorted out too: how and where should data be stored for the long term, who will control access, and who will pay for those services? Existing metadata standards need to be extended to cope with health data.

These obstacles have been known for some time; most can be overcome in the field of public health just as they have been overcome in other fields. However no institution has taken the lead in defining a work plan and carving up the tasks and the bill. In this round table paper, we suggest goals for data sharing and a work plan for reaching them, and challenge respondents to move beyond well intentioned but largely aspirational data sharing plans.

Introduction

As they prepare for careers in science, today’s students doubtless hear the same clichés as we did a generation ago: science advances collaboratively; we reproduce and extend the work of others; we stand on the shoulders of giants. In some fields, such as genomics, these axioms are becoming true. In epidemiology and public health, however, data sharing and collaboration remain more aspirational than real.

Students embark on a career in health research in the spirit of sharing; they want to help improve the well-being of others. For all the talk of collaboration, they will enter a world in which another axiom dominates: “publish or perish”. That system puts the interests of public health researchers in direct conflict with the interests of public health.

Benefits of sharing

The situation was not so different in genomics less than 15 years ago. Then, after years of hoarding their findings in individual laboratories and progressing at an expensive snail’s pace, in 1996 researchers agreed to share all their data openly.1 Now laboratories sequence during the day and post their results that same night; other researchers can begin to stand on their shoulders the very next day. As a result, genetic research is advancing faster than any other area of biomedicine.2

Genomics has taught us that sharing data with other scientists is a way to add value without costing a lot. It allows the same data to be used to answer new questions that may be relevant far beyond the original study. And it allows for meta-analyses that are free from the distortions introduced when only summary results are available.3,4 We could get far more out of public health research if we followed a similar path, if we squeezed more scientific and policy insights out of data that have already been collected.

Routine health and service use statistics can be just as useful for policy analysis as research data. Many countries are reluctant to release detailed service use data because analysis by disinterested outsiders may contradict politically acceptable interpretations. Most countries do, however, contribute aggregate statistics freely to large international databases maintained by multilateral organizations, although they are not always granted free access to those databases when they want to use them. Such restrictions on access, imposed unnecessarily by agencies wanting to protect their institutional mandates, cripple the potential utility of these expensive resources. Researchers and governments are also reluctant to see the data they provide used and manipulated by others in ways they don’t understand because secondary users (including international agencies) do not always publish their methods.

Research data are desperately underused too, in part because of a critical shortage of competent data managers.5 In other fields – genetics, banking and retailing – data management is a valuable skill. People are trained and develop careers in the field. In public health research, data management is the poor cousin of analysis. Undervalued and underfunded, inadequate data management undermines the rest of the scientific enterprise. One review in the United Kingdom of Great Britain and Northern Ireland found that many of the variables collected in epidemiological studies were never cleaned and coded, so they could not be used even by the primary researchers, let alone shared.6 In complex population-based surveys in developing countries, data management and analysis skills are in even shorter supply, so a higher proportion of data probably goes to waste.7

When we’re dealing with public health research, wasted data can translate into shorter, less healthy lives. Improving data management so that data can be shared is a first step to reducing that waste. But it will not be enough. We need to change the in-
centives that pit the interests of individual researchers against the interests of public health, that pit institutional interests against the more rapid advancement of knowledge and understanding. Governments may hold micro-data back from international organizations, but there’s no excuse for international organizations to limit access to the aggregate data that governments do provide.

It’s easier to understand why individual researchers are reluctant to share data they have collected. That reluctance will certainly remain entrenched as long as their employers – research councils, foundations and universities – regard publication of research papers in peer-reviewed biomedical journals as the main yardstick of success. If, however, “publish [papers] or perish” were to be replaced by “publish [data] or perish,” the picture might change rapidly, as it did in genomics.

What did that experience teach us? That a change in the culture of science requires the buy-in of key research teams, yes, but that it also requires considerable and very concrete commitments from funders. The two largest funders of the Human Genome Project, the Wellcome Trust and the National Institutes for Health, invested massively in the infrastructure needed to share data on a large scale for the long term. They also changed funding mechanisms to emphasize team work and the value of roles such as data management, rather than just looking at publication and citation records. Inevitably the rapid change of culture raised some tensions, but those have now largely been resolved. It would be perfectly feasible for research funders to take similar steps in other fields so that personal and professional incentives are aligned rather than in conflict.

Genomics and the social sciences (which have a dramatically better record of sharing data than most biomedical sciences) have developed techniques to deal with two of the other main obstacles to sharing of public health research data – confidentiality and consent. In part because of the development of research tissue banks (biobanks), broad consent procedures are increasingly becoming a norm. Anonymization removes some of the obstacles associated with consent, and techniques for protecting identities are improving constantly. Despite concerns about the theoretical possibility of identifying individuals in shared data sets, no breaches of confidentiality have yet been recorded in anonymized data sets. Social and economic sciences have also gone further in making the sharing of data sets easy through standard metadata, both for aggregate data through Statistical Data and Metadata Exchange (SDMX) standards and for individual data using Data Documentation Initiative (DDI 3.0) standards. A further lesson from other fields: it is possible to make data widely available to the research community while still safeguarding integrity, through the use of standardized data use agreements and licences. These define who may use data and how, and may require secondary analysts to contribute both derived data and a record of their analytic methods back to the database, so that primary and other users can both verify and benefit from their work.

The data that we collect and don’t make full use of do not come free. The collection of routine health statistics is paid for by our tax money. Most research aiming to reduce ill-health in the developing world is also funded either from the public purse or by charitable foundations. It is irrational to invest so much in collecting data and yet so little in ensuring that we make the best use of it. It is also ethically unsound: people who participate in research have a right to expect that the results will be used to improve life for them and/or for their communities.

Funders and standard-setters have been aware of this for some time. Gradually, they are urging or adopting policies that aim to increase the use and recycling of data. Although they don’t all yet practice what they preach, several international organizations, including the Organisation for Economic Co-operation and Development and the World Health Organization, have issued statements calling for increased access to routine statistics and other publicly-funded data. Many biomedical journals have recently addressed the importance of data sharing in editorials and commentary articles. A few biomedical journals expect researchers to make the data that underlie research articles available to others on request. An even smaller number of journals have followed the lead of Annals of Internal Medicine and now require authors to state whether and how they will make protocols, analysis tools and data available to others. But even Annals stops short of requiring authors to publish data sets along with their articles. “If we did that, we’d have a very thin journal,” commented editor Christine Laine at a recent conference on biomedical publication.

There are indications that public and foundation funders of public health research wish to strengthen data sharing policies, shepherding epidemiologists down the road already travelled by geneticists. Many field researchers who have battled difficult climates, erratic electricity supplies, fuel shortages and recalcitrant local authorities will doubtless resent increasing pressure to “give data away”. Some are also apprehensive that people looking at the data in the comfort of some distant, well resourced office will spot the errors that are the inevitable by-product of research in the real world.

Governments are equally reluctant to expose their data to interpretations other than those published by their official statisticians. There is a fear, too, that data may be used by others not just for professional but for economic gain. This is sometimes cast as a “north–south” divide; one spectre raised is of pharmaceutical companies exploiting data from developing countries to develop products that those countries then can’t afford.

Feelings of ownership over hard-won data, viscerally held even by researchers who support the idea of data sharing in principle, are understandable. And peer reviewers, mostly researchers themselves, are reluctant to approve funding for data management if it cuts into budgets for data collection. But funders of science are themselves under pressure to get the most out of expensive research studies. They have to wrestle with two important questions: how much data sharing is desirable and how much is feasible?

Researchers sometimes argue that interpretation of their data is so dependent on understanding local conditions that the data would be worthless to other scientists. This is often a reflection of inadequate documentation, but also a necessary failure of imagination. Sailors keeping log books on whaling boats in the 1600s could not have predicted that, centuries later, the data would be an important source of information for climate change scientists. Most funders have stringent peer-review procedures; few invest in research that they believe is of only very localized importance, and few wish to support research that produces data of such poor quality that it has no further value. Publicly-funded data can also be invaluable to students learning data management and analysis skills. It thus seems
fair to expect that almost all public health research funded by taxpayers or charities might be useful to secondary analysts. If a piece of research is considered worthy of publication in a peer-reviewed journal, the underlying data should also be worth publishing.

How feasible would it be to make these data available to the scientific community? Technically, the challenges are not trivial, but they have been overcome in several other fields; they are broken down here into manageable parts. We maintain that the major constraints to feasibility are a cultural resistance to change from within our own scientific community, and a reluctance of any institution to take leadership of the data sharing agenda. We also believe, however, that the imperative to share data will only grow stronger. The research community should look at this pressure from funders as an opportunity rather than an imposition.

Goals for funders and researchers

Here we propose several goals to which funders and researchers can jointly aspire and towards which progress can be measured: (i) all data of potential public health importance funded by taxpayers or foundations will be appropriately documented and archived in formats accessible to the wider scientific community; (ii) all data provided by governments to databases developed by publicly-funded organizations will be freely available to any user, at the level of detail at which it was provided; (iii) the publication of a research article in a biomedical journal will be accompanied by the publication of the data set upon which the analysis is based; (iv) funders and employers of researchers will consider publication of well-managed data sets as an important indicator of success in research, and will reward researchers professionally for sharing data; and (v) all planned research will budget and be funded to manage data professionally to a quality adequate for archiving and sharing.

Plan of work

These goals can only be achieved with considerable investment in several practical areas. We propose the following plan of work, necessary to underpin progress towards our stated goals.

Fill the gaps in data management

There is a need to develop metadata standards, which will lead to improved documentation and allow data to be combined more easily across time, locations and sources. This will probably require the extension of DDI and SDMX standards to encompass areas of public health interest. Agreement is also required on standards for anonymization and safeguarding of confidentiality.

We need to develop a search portal that will allow data to be discovered across a range of repositories, and standards for repositories similar to those used for registries of clinical trials. We also need to invest in training in data management for public health, especially in developing countries, and the development of career paths in bioinformatics.

Increase incentives to share data

We need to further develop and adopt reliable citation standards for data sets, such as those proposed by DataCite collaboration, and ensure they are indexed in databases such as PubMed. Standards and procedures for peer review or quality control of data sets are also needed. Digital fingerprinting of data would allow tracing of secondary use and we should develop methods and measures to track the value that sharing data adds to the work of both primary researchers and funders of research. There is a need to agree on norms and standards governing fair use periods for primary researchers, data access policies and data use agreements.

Data libraries

To underpin the long-term viability of data libraries, we need to invest in expanding existing infrastructure to cover curative and access of data of public health importance. This calls for a business or funding model that assures the long-term viability of data archives.

Conclusion

All of these areas have already been identified as critical to promoting data sharing, often repeatedly so. Funders, governments, publishers and many researchers want these things to happen, it seems. Some of the organizations calling for greater sharing of public health research data have expressed willingness to pay for parts of the work. But none are willing to take charge of the agenda, committing themselves to orchestrating the dull, messy but essential work of developing the norms and standards that will allow data sharing to revolutionize public health research.

It is time to move beyond expressions of good intentions and to get on with the practical work that will allow data to be shared. The first thing that is needed is leadership. We challenge other participants in this round table to commit to coordinating, funding or carrying out the work described in this paper. Only after someone takes the lead in tackling these issues will today’s students of public health be able to climb onto the shoulders of the current giants in our field.

Competing interests: None declared.
Sharing health data

Résumé
Partage des données sur la santé : les bonnes intentions ne suffisent pas

Les épidémiologistes et les chercheurs en santé publique s’engagent très lentement dans la révolution que subit le partage des données et les agences chargées d’entretenir les bases de données mondiales sur la santé sont réticentes à ce partage. Une fois certains investissements consentis dans les infrastructures, le recyclage et la combinaison des données peuvent donner accès à un maximum de connaissances pour un coût additionnel minimal. En refusant le partage des données, les chercheurs ralentissent les progrès vers la réduction de la morbidité et de la mortalité et interdisent l’accès à l’information à un public tout juste bon à payer les impôts qui financent la plupart de leurs recherches.

Les apporteurs de fonds pour la recherche en santé publique commencent à appeler au changement et à développer des politiques de partage des données. Cependant, ils n’ont pas encore trouvé de moyens adéquats pour aplatiser les obstacles responsables de l’échec de ce partage. Il s’agit notamment de structures professionnelles qui récompensent la publication d’une analyse, mais pas celle des données, et de flux de financement et d’évolutions de carrière qui continuent de sous-évaluer le travail essentiel de gestion des données. Il convient aussi de sérifier les problèmes pratiques : où et comment les données doivent-elles être stockées sur le long terme, qui exercera un contrôle sur les accès et qui paiera pour ces services ? Les normes existantes pour les métadonnées doivent être étendues pour couvrir les données relatives à la santé.

Ces obstacles sont connus depuis un certain temps ; la plupart d’entre eux peuvent être surmontés dans le domaine de la santé publique tout comme ils l’ont été dans d’autres secteurs. Néanmoins, aucune institution n’a pris la direction des opérations pour définir un plan de travail et répartir les tâches et la facture. Dans cet article destiné à une table ronde, nous proposons des objectifs pour le partage des données et un plan de travail pour les atteindre et nous sollicitons des réponses pour aller au-delà de plans de partage des données bien intentionnés, mais largement utopistes.

Resumen
Intercambio de datos sanitarios: las buenas intenciones no son suficientes

Los epidemiólogos e investigadores en salud pública están avanzando muy lentamente en la revolución del intercambio de datos, y además los organismos que mantienen las bases de datos mundiales sobre salud se muestran reacios a compartir su información. Una vez realizadas las inversiones en infraestructuras, la reutilización y combinación de datos brindan acceso a un máximo de conocimientos con un costo adicional mínimo. Al negarse a compartir los datos, los investigadores están frenando los progresos hacia la reducción de la morbilidad y la mortalidad y están negando un bien público a contribuyentes que apoyan la mayor parte de las investigaciones.

Los agentes de financiación de las investigaciones en salud pública están empezando a exigir cambios y a elaborar políticas de intercambio de datos. Sin embargo, aún no están abordando adecuadamente los obstáculos que impiden compartir esos datos. Entre ellos cabe citar unas estructuras profesionales que recompensan la publicación de análisis, pero no de datos, y unas fuentes de financiación y unas perspectivas de carrera que siguen sin reconocer el carácter crucial de la gestión de datos. Es preciso esclarecer también algunos aspectos prácticos: cómo y dónde deben conservarse los datos a largo plazo, quién controlará el acceso y quién pagará esos servicios. Las normas existentes sobre metadatos deben ampliarse para poder manejar los datos sanitarios.

Estas dificultades son conocidas desde hace ya algún tiempo, pero la mayoría pueden ser superadas en el campo de la salud pública al igual que han sido superadas en otros campos. Sin embargo, ninguna institución ha tomado la iniciativa para definir un plan de trabajo y repartir las tareas y los costos asociados. En este artículo de la mesa redonda proponemos metas para el intercambio de datos y un plan de trabajo para su consecución, y alentamos a los encuestados a trazar algo más que unos planes de intercambio de datos bienintencionados pero demasiado ambiciosos.

References


