In April 2019, the Canadian Institutes of Health Research (CIHR) announced an $81-million funding commitment for an initiative called the Canadian Data Platform under the Strategy for Patient-Oriented Research (SPOR). The funding will enable, among other things, the development of a single portal through which researchers can request access to health administrative, demographic and social data from various federal and provincial sources. This comes 8 years after an international CIHR review panel called for “a Canada-wide effort to harmonize data sets and enable national linkages which would benefit all CIHR institutes and the Canadian research enterprise at large.” The ambitious proposal represents a welcome step forward to capitalize better on the rich population-based data assets in Canada. If successful, it will create a “distributed” network to facilitate multi-jurisdictional research, consolidate efforts to harmonize and validate data definitions, share technical and other expertise, and better support advanced analytics and infrastructure needed to realize the potential of an increasing array of “big data.”

Canada has the opportunity to be an international leader in population-based health and health systems research. The single-payer health systems capture whole populations tracked longitudinally through all publicly funded provincial health services. These administrative data, collected in the day-to-day work of the health system by clinicians (e.g., electronic health records) or government (e.g., physician billings), are the backbone for this research. Many provinces have health services research institutes, which both capitalize on these data and work closely with policy-makers to conduct applied research. In some provinces, health administrative data are linked to a wide variety of other service and demographic data (e.g., immigration, social services, education, child welfare). Manitoba has the most impressive array of linked intersectoral data, which have enabled world-class research on health and social equity. These provincial data repositories are also the “backbone” for linkage with an increasing array of deeper clinical data. Examples include population-based laboratory data, electronic medical records and clinical-trial data, as well as research cohorts, which include imaging, genomic and other “omic” information. Although recent SPOR Support Unit funding has improved data access in provinces without dedicated data institutes, material differences in available data and access remain.

The overarching goal of the national data platform — which will be overseen by a consortium of leaders from provincial data centres, SPOR Support Units, the Canadian Institute for Health Information and Statistics Canada, working under the auspices of the Pan-Canadian Real-World Health Data Network — is to catalyze more multi-jurisdictional research. All provinces and the Northwest Territories are represented, as well as knowledge users from provincial ministries of health and leaders in patient and public engagement and Indigenous health research.

Canada’s 13 health systems provide a natural experiment for policy and program evaluation. Consider, for example, the ability to compare patient outcomes and system costs of differing pharma care eligibility across provinces. To do so, researchers require access to comparable data to create similar patient cohorts and measure drug access or use and outcomes. Increasingly, disease-based quality improvement networks need multi-provincial data to evaluate interventions. As interest grows in both pragmatic trial design and linkage of dormant trials to administrative data for long-term outcomes, access to comparable data is necessary for multi-provincial trials. Similarly, the ability of pan-Canadian cohorts such as the Canadian Partnership for Tomorrow to link to administrative data will enhance their value. Finally, multi-jurisdictional research allows pooling...
of data for the study of uncommon outcomes. The Canadian Net-
work for Observational Drug Effect Studies® is an example of a
distributed data network that uses standardized protocols and
meta-analysis across provinces to study real-world outcomes of
drugs after they are marketed.

How will the national data platform help? At its most basic
level, the platform will provide a single portal through which
researchers will have better access to data from multiple jurisdic-
tions, and dedicated personnel in every province and territory to
help researchers navigate data holdings and processes to access
data and conduct distributed analytics (or comparative analy-
ses). This will overcome the well-known barriers to pooling data
across provinces owing to provincial legislation. The platform
will not change any of the current provincial requirements for
data access, but may be a catalyst for evolution in those jurisdic-
tions with more cumbersome processes for accessing data and
fewer available data sets.

The platform will leverage existing and long-standing provin-
cial and federal investments in health data and data infrastruc-
ture. Some provinces have well-developed capacity for data link-
age and storage, and the platform will allow knowledge sharing with
respect to a number of these technical areas as well as
investment in the technology to create, share and fulfill data
access requests and enable distributed analyses. Data science
approaches to both data handling and analytics, including techni-
quies such as natural language processing and machine learn-
ing or artificial intelligence, are important areas for collective
work, as is the need for data validation.6

The platform proposes to validate disease definitions for a set
of prioritized conditions. Diagnoses in physician billings are
notoriously inaccurate and differences in available data across
provinces pose challenges to standardizing definitions. For
example, drug data (insulin and oral hypoglycemics) or labora-
tory data (glycosylated hemoglobin) can be used in some prov-
inces, but not others, to identify cohorts with diabetes, so the
ideal state of comparable cohorts across a number of conditions
may not be realized. In addition, although electronic medical
record data have the potential to contribute to standardizing dis-
ease definitions and provide rich clinical data, extraction costs
even for structured data elements such as height and weight are
substantial. Apart from cost, the technical challenges are
immense with respect to extracting richer data from, for exam-
ple, clinical notes, and in the context of a multitude of different
electronic medical record vendor systems and data governance
agreements. Finally, apart from disease definitions, researchers
often develop algorithms and other definitions of constructs
(e.g., primary care visits) for studies, and ensuring that research-
ers have access to “local knowledge” of the health system is criti-
cal to data validity and comparability.

Finally, there are many other important areas of leadership
for this platform, including supporting Indigenous data sover-
gignty.7 The platform could build on nascent efforts by some pro-
vincial research institutes to engage the public through public
advisory councils in discussions on the use of large health data
sets to drive research. Although the importance of and support
for involving patients in health research is now well established,
less work has been done in considering how the public should
intersect with research organizations and researchers who use
administrative data. Qualitative work in Ontario suggests the
public is accepting of the use of their data for “the public good,”8
but examples from both Canada9 and England10 indicate that
communication and public engagement are critical to ensuring
this research has public trust.

The SPOR Canadian Data Platform builds on existing CIHR-
SPOR investments and provincial and federal data infrastructure
in an ambitious effort to overcome well-known obstacles to con-
ducting multi-jurisdiction research in Canada. The CIHR’s sub-
stantial investment is a great step forward, but vaulting to inter-
national leadership will likely require additional investments to
ensure the platform goes beyond administrative data to include
other data sets (e.g., detailed clinical data in electronic health
records and “omics” data), as well as to support advanced ana-
lytics including artificial intelligence and machine learning.

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