As noted in the 2018 clinical practice guidelines,1 Diabetes Canada reported that 3.4 million Canadians were estimated to be living with diabetes in 2016; by 2025, it is expected that the prevalence of diabetes will have increased by 44%. This increase is likely due to a number of individual-level factors such as obesity, aging, ethnicity, lifestyle and socioeconomic status,2–7 environmental factors8 and decreasing mortality rates.9–13 Among First Nations people, the burden of diabetes is disproportionate:14,15 the lifetime risk of diabetes was recently estimated to be 8 in 10 for First Nations adults (> 18 yr), compared to 5 in 10 for non–First Nations people.16 Prevalence rates are 3–5 times higher among First Nations people than other Canadians,17 and diabetes develops at younger ages, progresses more rapidly and has higher complication rates among First Nations people.16,18,19

First Nations people in Canada continue to experience health inequities related to the ongoing impacts of colonization.19,20,21 This history has contributed to poverty and marginalization and has resulted in major disparities in health status and outcomes reported for decades.15,22,23 Despite these persistent health inequities, First Nations people remain resilient in working toward healing.24 To understand the health of their communities and advocate for services within a complex and fragmented health care system,25,26 First Nations leaders need access to population-level data.

**First Nations people with diabetes in Ontario: methods for a longitudinal population-based cohort study**

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**Abstract**

**Background:** To improve diabetes care, First Nations leaders and others need access to population-level health data. We provide details of the collaborative methods we used to describe the prevalence and incidence of diabetes in First Nations people in Ontario and present demographic data for this population compared to the rest of the Ontario population.

**Methods:** To identify the population of First Nations people and other people in Ontario, we created annual cohorts of the Ontario population for each year between Apr. 1, 1995, and Mar. 31, 2015. Through a partnership between First Nations and academic researchers, we linked provincial population-based health administrative data stored at ICES with the Indian Register, which identifies all Status First Nations people. Our collaborative process was guided by the First Nations principles of ownership, control, access and possession (OCAP).

**Results:** Demographic characteristics for the 2014/15 cohort (n = 13 406 684) are presented here. The cohort includes 158 241 Status First Nations people and 13 248 443 other people living in Ontario. Using postal codes, we were able to identify virtually all (99.9%) First Nations people in Ontario as living in (n = 55 311) or outside (n = 102 889) a First Nations community. First Nations people were younger and more likely to live in semiurban or rural areas than the rest of Ontario’s population.

**Interpretation:** The collaborative methodology used in this study is applicable to many jurisdictions working with Indigenous groups who have access to similar data. The Ontario cohort defined here is being used to conduct analyses of health outcomes and use of health care services among First Nations people with diabetes in Ontario.
Little is known about the health care experience of First Nations people with diabetes in Ontario. In 2003, an analysis of prevalence, incidence, mortality and hospital admissions related to diabetes relied on identifying First Nations communities through postal codes, which captured only 7% of First Nations people in Ontario.27 Recent initiatives at ICES, where Ontario’s public health services data sets are stored, linked and used for research, have enabled the identification of Status First Nations people in Ontario, within guidelines established to ensure First Nations data sovereignty.28,29 As a result, we were able to link ICES data sets on disease incidence, demographic characteristics and geography with a data set of registered First Nations people in Canada, which allowed us to collaboratively perform in-depth analyses of health outcomes and use of health care services among First Nations people with diabetes in the province. These analyses are part of a larger, mixed-methods study to describe the landscape of diabetes in First Nations communities across Ontario (Appendix 1, available at www.cmajopen.ca/content/74/4/E680/suppl/DC1).

In this manuscript, we explain the partnership and data-governance principles guiding the analyses of the administrative health data that we are using to compare the prevalence and incidence of diabetes among First Nations people to those among the rest of the Ontario population between 1995/96 and 2014/15. We describe how we identified First Nations people in Ontario and compare their demographic characteristics to those of others living in Ontario. We also describe how we identified First Nations people with diabetes and the analyses we are conducting to compare their disease control, use of health care services and rates of complications to those of the rest of the Ontario population with diabetes. The first such study, focused on cardiovascular disease, has been published.30

Methods

Partnership and collaboration

This study is a result of a partnership between the Chiefs of Ontario, ICES and researchers across Ontario, coordinated by investigators at Queen’s University, Kingston, Ontario, Laurentian University, Sudbury, Ontario, and the University of Minnesota Medical School, Duluth. The collaborative process used for this project has been described elsewhere.31,32 The lead academic investigators have long-standing relationships working with First Nations communities. An Indigenous-led approach, based on principles of Indigenous data sovereignty,29 was used throughout the work, and our partners at the Chiefs of Ontario were central to directing this work. Initial meetings between the academic research team, senior health staff from the Chiefs of Ontario, an elder and 2 First Nations patient navigators established processes for working together (formalized in a research agreement) and determined the research questions for the project. All team members actively participated as coinvestigators in the conception and design of the work and in data analysis and interpretation.31 Our partners at the Chiefs of Ontario are coauthors on all works stemming from this partnership.

In addition, a patient advisory group made up of 7 First Nations people with lived experience of diabetes as a patient or family member provided additional insight. As described elsewhere,41 we worked with the Health Coordination Unit, which includes members from various First Nations organizations in the province, to invite members from different communities, acknowledging the diversity of First Nations people in Ontario. The advisory group met twice yearly, providing constructive and meaningful feedback. All members of the advisory group were offered authorship on all publications.

Data governance

The Chiefs of Ontario and ICES established a data-governance agreement to ensure First Nations governance of the Indian Register data held at ICES, in accordance with OCAP (ownership, control, access and possession) principles.33 In addition, this project was approved by the First Nations Data Governance Committee, which has 7 members from across the province appointed by the Ontario Chiefs Committee on Health.28 The committee developed an application process for researchers interested in accessing the Indian Register to identify First Nations people in Ontario or using the Regional Health Survey data. The current project served as a pilot for this application process. The project was also reviewed and approved by the Health Coordination Unit. We provided regular updates and presented findings to the patient advisory group, the Health Coordination Unit and the annual First Nations Health Forum in Ontario before releasing the results elsewhere.

Data sources

This project used the Ontario population-level health administrative data housed at ICES.34 Key to these analyses is the ability to link provincial population-based health information at the individual level with the Indian Register. The Indian Register is maintained by Crown–Indigenous Relations and Northern Affairs Canada and lists all people who are registered as members of a First Nation that is recognized under the Indian Act (i.e., all “Status Indians” or “Registered Indians”).35 These data sets (listed in Appendix 2, available at www.cmajopen.ca/content/74/4/E680/suppl/DC1) were linked by means of unique encoded identifiers and analyzed at ICES.

Study cohort

We defined 2 study cohorts. The first cohort is the population of First Nations people and other people in Ontario; we report on their characteristics in this manuscript. Second, we identified people with diabetes from within these cohorts. They will be used to estimate the annual prevalence and incidence of diabetes, as well as describe measures of disease control, use of health care services and rates of complications of diabetes.30,32

Incidence and prevalence cohorts

To identify the population of First Nations people and other people in Ontario, we created annual cohorts of the Ontario population for each year between Apr. 1, 1995, and Mar. 31,
2015. All people from the Registered Persons Database who were Ontario residents eligible for a health card, were alive for the entire year of the cohort (Apr. 1–Mar. 31 of the following year), had had contact with the health care system within the previous 7 years and were aged 105 years or younger were included in the population cohorts. We included all people who were alive at the end of each fiscal year to ensure that all new cases of diabetes were captured.

We used the Indian Register to identify all Status First Nations people in Ontario; all other Ontario residents were considered to be the population of “other people in Ontario.” We used a combination of residence codes and postal codes to identify whether a person lived in a First Nations community (i.e., “Indian Reserves”35). Residence codes are unique to each municipality or First Nations community and are recorded in the National Ambulatory Care Reporting System and the Canadian Institute for Health Information Discharge Abstract Database when people are admitted to emergency departments or hospitals. For those who did not have a recorded residence code, we used their postal code from the Registered Persons Database, which includes the address associated with a person’s health card, to map their home address to a census subdivision. Most census subdivisions are clearly and exclusively within a First Nations community or outside a First Nations community. We considered First Nations people who lived in the few census subdivisions that included both First Nations communities and other communities to be living in a First Nations community. Through this process, we classified the residence for First Nations people as being in a First Nations community, outside a First Nations community, out-of-province (based on a postal code outside Ontario) or unknown (there was no available residence code or postal code for the person) each year. As the geographic information available in the 1990s has limited reliability, we determined the classification of living in or outside a First Nations community from 2001 onward.

Within each yearly cohort, we established the presence of diabetes through the Ontario Diabetes Database. This data set has been validated against primary care health records and has a reported sensitivity of 86% and specificity of 97%.36

Diabetes-specific cohorts
To estimate diabetes-specific outcomes, we created annual cohorts of the population with diabetes for each year between Apr. 1, 1995, and Apr. 1, 2014. All people from the Registered Persons Database who were Ontario residents eligible for a health card, were alive at the beginning of each year, had a prior diagnosis of diabetes, had had contact with the health care system within the previous 7 years and were aged 105 years or younger were included in the diabetes cohorts.

Covariates
We described multimorbidity using the Johns Hopkins ACG System (version 7) Aggregated Diagnosis Groups (ADGs)37 applied to diagnostic codes associated with hospital admissions and physician services data sources. We classified each person as having a low (0–4 ADGs), medium3–9 or high (≥ 10 ADGs) degree of comorbidity. We assigned urban–rural location using the Rurality Index of Ontario,18 a broad measure of rurality based on the dissemination area of a person’s postal code. A score of 0–9 indicates a major urban centre, a score of 10–39 specifies a nonmajor urban centre (referred to as semiurban), and a score of 40 or greater is considered rural. People with a missing Rurality Index of Ontario score were categorized as “Remote/missing.”

Analytic approach
We calculated annual diabetes prevalence rates using each annual population cohort as the denominator for that year. We calculated the numerator as people in the denominator who received a diagnosis of diabetes on or before the end of the year according to the Ontario Diabetes Database. We calculated diabetes incidence rates in a similar fashion using all newly diagnosed cases in a particular year as the numerator and all people in the annual population cohort with no previous diagnosis of diabetes as the denominator.

To estimate diabetes-specific outcomes, we calculated each outcome measure on an annual basis using each annual diabetes cohort as the denominator for that year. The numerator was people in the denominator who experienced the outcome during the 1 year of follow-up. For rare events, we calculated outcome measures on a 2- or 5-year basis. Definitions for these specific outcomes will be described in detail in each forthcoming paper.

We estimated crude and age- and sex-standardized prevalence and incidence rates and corresponding 95% confidence intervals among First Nations people and compared them to those among other people in Ontario. We calculated age- and sex-standardized rates, along with $\gamma$-distributed 95% confidence intervals,39 using a direct standardization based on the 2001 Ontario census population to adjust for the differences in population distribution between populations and over time. Among First Nations people, we stratified estimates by those living in or outside First Nations communities, rurality and comorbidity.

Analyses were conducted with SAS Enterprise Guide version 7.1 (SAS Institute).

Ethics approval
This project was reviewed and approved by the research ethics boards of Queen’s University and Laurentian University.

Results
Description of population of First Nations people and other people in Ontario
Registered First Nations people accounted for 1.2% of the overall Ontario population ($n = 13\,406\,684$) included in our study cohort. We were able to determine the location of residence (living in or outside a First Nations community) for almost all First Nations people; the 41 people for whom we were unable to determine location of residence are excluded from any analysis stratifying data by First Nations people living in or outside First Nations communities.
The proportion of First Nations people living in First Nations communities decreased from 36.4% (52,348/143,702) in 2001/02 to 35.0% (55,311/158,241) in 2014/15. Compared to other people in Ontario, First Nations people were younger and slightly more likely to be male (Figure 1). Those living in First Nations communities were more likely to be male and were younger than First Nations people living outside these communities (Figure 2).

First Nations people living outside First Nations communities had a higher burden of comorbidity than those living in First Nations communities (19.1% v. 13.1%) (Table 1). Significantly fewer First Nations people than the rest of the Ontario population lived in urban areas (33.1% v. 73.2%). Overall, 27.5% of First Nations people had a Rurality Index of Ontario score of Remote/missing; the values for those living in and outside First Nations communities were 59.9% and 10.0%, respectively. Almost half (49.7%) of First Nations people lived in northern areas of the province, compared to 5.5% of the rest of the Ontario population (data not shown). The proportion of First Nations people across the province is shown in Figure 3.

**Figure 1:** Age distribution of First Nations people (A) and other people (B) in Ontario, 2014/15.
The age and sex distribution of First Nations people differed from that of other Ontarians. First Nations people tended to be younger and were more likely to live in northern regions of the province and in semiurban, rural or remote areas. Those living in First Nations communities had a lower burden of comorbid disease than those living outside First Nations communities, possibly because they were younger. Also, people may move outside First Nations communities as they age or experience illness to be closer to health care services. First Nations people with diabetes have noted a lack of health care providers in their communities, which results in the need to travel to receive care.19

Our collaborative approach to identifying First Nations people with diabetes in Ontario and describing their access to care and health outcomes is potentially applicable to those working with other Indigenous groups who have access to similar data and need to feel confident that they will be central in deciding how the data are used and how conclusions will be drawn. Although there are provincial and regional differences across Canada, the methods used in

Figure 2: Age distribution of First Nations people living in (A) and outside (B) First Nations communities, 2014/15.
our administrative data analyses are likely applicable to jurisdictions with access to similar data sources.

**Limitations**

There are a number of limitations with the administrative health data we used. Population denominators based on the Registered Persons Database do not include people who have had a lapse in their health care coverage, which may lead to overestimation of the rates of diabetes in some regions or communities. Postal codes in the Registered Persons Database are associated with a person’s health card and are updated when a health card is renewed (usually every 5 yr) or when a person interacts with the health care system. Therefore, the Registered Persons Database does not have up-to-date postal codes for all Ontario residents, and there is a high likelihood that we misclassified the place of residence of some people with high mobility. First Nations people are more geographically mobile than other Canadians, and these inaccuracies in postal codes may have resulted in incorrectly ascribing First Nations people to living in or outside First Nations communities. Last, the Registered Persons Database contains a small number of people who are deceased or no longer living in Ontario, which may slightly underestimate mortality rates.

Our cohort includes only Status First Nations people and excludes those who are not registered with the federal government or are members of a First Nation that is not recognized by the government of Canada. In addition, we were not able to capture care delivered by community health centres or Aboriginal Health Access Centres, which provide care to 25% of Indigenous populations in Ontario. This will result in underestimation of measures of access to primary care.

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<th>Table 1: Demographic characteristics of First Nations people, including those living in and outside First Nations communities, and other people in Ontario, 2014/15</th>
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Note: IQR = interquartile range, SD = standard deviation.

*Except where noted otherwise.
†This includes all First Nations people included in the Indian Register, regardless of whether we could determine whether they lived within a First Nations community.
‡We were unable to determine location of residence for 41 people.
care and outcomes that rely on billing claims by primary care physician (e.g., measures of diabetes control such as glycosylated hemoglobin and lipid monitoring) among patients who used these facilities. In addition, those living in First Nations communities who were treated by nurses working at federally supported nursing stations and health centres will have no physician visit billing claims, which will result in underestimates of access to primary care. This is most relevant in northern Ontario. People living in northwestern Ontario may receive specialist and hospital-based care in Manitoba and, to a lesser extent, those living in southeastern Ontario may receive such care in Quebec. Our data include claims only from Ontario physicians and facilities, so out-of-province care has been missed. Many First Nations people live in northwestern Ontario, and this limitation may be important for some diabetes outcome measures, including hospital admissions and visits to endocrinologists. In addition, our measure of multimorbidity relies on diagnostic codes associated with hospital admissions and physician visits, and thus may be underestimated in people who accessed care outside the Ontario system, or in nursing stations or from other nonphysician providers.

Although the algorithm used by the Ontario Diabetes Database to identify people with diabetes has high sensitivity and specificity, the diagnostic accuracy of the algorithm may be different in First Nations populations. In addition, the database does not capture gestational diabetes or distinguish between type 1 and type 2 diabetes. Although the majority of people with diabetes have type 2, type 1 accounts for a greater proportion of the diabetes burden among younger people. This is particularly relevant in First Nations populations, who are younger than other people in Canada.

Income quintiles are a commonly used proxy for individual socioeconomic status. However, this measure is less accurate in rural regions than in urban regions. Prior work and preliminary analyses (data not shown) suggest that this proxy method is not sufficiently accurate for use in the Status First Nations population, so we did not include socioeconomic status in these analyses. We also found that the Rurality Index of Ontario score does not perform as well in First Nations populations compared to the rest of the Ontario population. Because the Rurality Index of Ontario is partially based on travel times to the nearest health care centre via roads and highways, fly-in communities are excluded.

Figure 3: Proportion of the Ontario population identified as Status First Nations, by Local Health Integration Network, 2014/15. Adapted from reference 32 with permission from ICES.
from this calculation, as are populations with less than 500 and other centres that do not belong to a municipality. Thus, we included the category “Remote/missing” as a fourth category.

Last, administrative health data do not capture information on other important risk factors for diabetes, such as food security, diet, body mass index, smoking status and physical activity levels, which also influence the outcomes of people with the disease. These factors are incorporated in our analysis of the First Nations Regional Health Survey (Dr. Laura Rosella, University of Toronto: unpublished data, 2019).

Conclusion
We identified First Nations people in Ontario using administrative health data and created 2 cohorts with diabetes, First Nations people and other people in Ontario. These cohorts are being used as the basis of a series of studies describing the landscape of diabetes among First Nations people in Ontario. We hope this will help First Nations communities prevent and manage diabetes and cope with the long-term complications of the disease. Because reports often focus on negative findings regarding Indigenous health, our advisory group communicated the need to ground the findings of this work in the context of the history of First Nations people in Ontario and their experiences with colonization. This is important as we consider how diabetes affects First Nations people compared to other people in Ontario.

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Contributors: Michael Green, Baiju Shah, Carmen Jones, Roseanne Sutherland, Kristen Jacklin and Jennifer Walker conceived and designed the study. Shahriar Khan analyzed the data. Morgan Slater drafted the manuscript. All of the authors contributed to data interpretation, revised the manuscript critically for important intellectual content, approved the final version to be published and agreed to be accountable for all aspects of the work.

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