Burden of nonalcoholic fatty liver disease in Canada, 2019–2030: a modelling study

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Abstract

Background: Nonalcoholic fatty liver disease (NAFLD) and nonalcoholic steatohepatitis (NASH) account for a growing proportion of liver disease cases, and there is a need to better understand future disease burden. We used a modelling framework to forecast the burden of disease of NAFLD and NASH for Canada.

Methods: We used a Markov model to forecast fibrosis progression from stage F0 (no fibrosis) to stage F4 (compensated cirrhosis) and subsequent progression to decompensated cirrhosis, hepatocellular carcinoma, liver transplantation and liver-related death among Canadians with NAFLD from 2019 to 2030. We used historical trends for obesity prevalence among adults to estimate longitudinal changes in the number of incident NAFLD cases.

Results: The model projected that the number of NAFLD cases would increase by 20% between 2019 and 2030, from an estimated 7,757,000 cases to 9,305,000 cases. Increases in advanced fibrosis cases were relatively greater, as the number of model-estimated prevalent stage F3 cases would increase by 65%, to 357,000, and that of prevalent stage F4 cases would increase by 95%, to 195,000. Estimated incident cases of hepatocellular carcinoma and decompensated cirrhosis would increase by up to 95%, and the number of annual NAFLD-related deaths would double, to 5,600.

Interpretation: Increasing rates of obesity translate into increasing NAFLD-related cases of cirrhosis and hepatocellular carcinoma and related mortality. Prevention efforts should be aimed at reducing the incidence of NAFLD and slowing fibrosis progression among those already affected.

Nonalcoholic fatty liver disease (NAFLD), defined by the presence of excessive liver fat in the absence of another causative factor, is recognized to cause cirrhosis and hepatocellular carcinoma. Cases of NAFLD can be characterized as simple steatosis or nonalcoholic steatohepatitis (NASH), characterized by additional histologic features. Obesity rates are increasing in Canada, affecting over one-third of Canadians, and obesity is 1 component of metabolic syndrome, which is a risk factor for NAFLD. There is a growing disease burden associated with NAFLD, following the trajectory of increasing obesity in Canada and globally. Even if further increases in obesity in Canada are halted, NAFLD-related morbidity and mortality are projected to increase for decades.

Analyses of the disease and economic burden associated with NAFLD based on data in the existing literature have recently been reported. To our knowledge, there are currently no studies reporting estimates of NAFLD prevalence or disease burden in the general Canadian population, but estimates have been reported in selected populations in other areas. Given the substantial health and socioeconomic burden of NAFLD, there is an increasing need to forecast the burden of disease. A modelling framework can provide a range of outcomes that can help in effective resource use and the development of strategies to prevent further increases in disease burden. The current analysis describes the results of such modelling to forecast the burden of disease of NAFLD and NASH for Canada, from 2019 to 2030.

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Methods

A Markov model (Appendix 1, Supplemental Figure S1, available at www.cmajopen.ca/content/8/2/E429/suppl/DC1) was built with the use of Microsoft Excel to estimate the number of NAFLD cases by disease stage in Canada beginning in 1950.\textsuperscript{15}

Model

The model tracked the Canadian population by age group and gender over time, with adjustment for background mortality at every step. New NAFLD cases entered the model based on longitudinal adult obesity trends. We calculated progression of disease through fibrosis,\textsuperscript{16} decompensated cirrhosis, hepatocellular carcinoma and liver transplantation stages (Appendix 1, Supplemental Figure S1) after accounting for both all-cause mortality (with adjustment for excess cardiovascular and non–liver-cancer mortality) and liver mortality attributable to NAFLD. Fibrosis transition rates were varied by gender (males experience faster progression) and age (Appendix 1, Supplemental Tables S2 and S3).

We implemented a Delphi process in which expert consensus was used to develop key model inputs (Appendix 1, Supplemental Table S1). Experts were invited based on research expertise and represented multiple disease areas including hepatology and diabetology. In addition, the expert panel included representation from provinces representing a majority of the Canadian population.

Population and mortality

We organized the total Canadian population and background mortality rates by year, gender and 5-year age groups.\textsuperscript{17} Increased background mortality among NAFLD cases was based on reported hazard ratios for all-cause mortality by fibrosis stage,\textsuperscript{18} adjusted to remove the impact of increased liver-related mortality,\textsuperscript{19} as this was calculated as a separate progression in the model.\textsuperscript{2,20–22} After adjustment, there was no increased non-liver-related mortality in stages F0–F2; increased mortality ratios of 1.42 and 1.43 were applied to prevalent F3 and F4 cases, respectively (Appendix 1, Supplemental Table S2). For the oldest age groups (age \( \geq 75 \) yr), no excess mortality was assumed given the already high mortality rates in this age group and reported decreases in the incremental impact of cardiovascular disease mortality among people aged 75 years or older.\textsuperscript{23}

New cases of nonalcoholic fatty liver disease

We extrapolated the annual number of new NAFLD cases based on longitudinal changes in the prevalence of adults in different body mass index categories, with the assumption that relative increases in NAFLD prevalence would mirror changes in obesity prevalence\textsuperscript{24–26} (Appendix 1, Incidence [new cases] calculations). Owing to variations in cut-off levels for obesity that vary by race or ethnicity\textsuperscript{27} and differences in health risk by body mass index class,\textsuperscript{28} we calculated the prevalence of obesity as a weighted average using data for body mass index of 25 or greater for the population classified by 1996–2016 census data as South Asian, Chinese, Filipino, Southeast Asian, Korean or Japanese,\textsuperscript{29} and data for body mass index of 30 or greater for the remaining population. We estimated the total population with a body mass index of 25 or greater to be 7.3\% of the total Canadian population in 2006, 14.2\% in 2016 and, with linear trending, 19.0\% in 2030.

Using the weighted average for the body mass index categories as described above, we extrapolated changes in obesity prevalence based on trending of data for the adult prevalence of body mass index of 25 or greater and of 30 or greater,\textsuperscript{24–26} where the adjusted obesity prevalence rate was estimated at 9.1\% in 1975 and 32.7\% in 2014 (Appendix 1, Supplemental Figure S2).

Prevalence of nonalcoholic fatty liver disease

Among people aged 20 years or older in 2018, there was an assumed NAFLD prevalence rate of 25\% (Delphi range 22.5\%–27.5\%) based on expert consensus. The prevalence of NAFLD was assumed to decline among the youngest age groups (\( \leq 18 \) yr), who are often not included in studies based on the general population.\textsuperscript{30,31} We based the age and gender distribution of prevalent NAFLD cases on data from general population studies, in which prevalence increases with increasing age.\textsuperscript{30,32,33} Based on a cohort of patients with NAFLD in Calgary, we assumed that prevalence would be approximately equal between men and women.\textsuperscript{34} Owing to elevated mortality, we assumed that prevalence would naturally decline among the oldest age groups (\( \geq 80 \) yr), with peak prevalence occurring in late middle age.

Liver transplantation

Annual liver transplantation procedures were reported by Canadian Blood Services.\textsuperscript{35} Based on expert input and review of diagnostic categories for transplant recipients,\textsuperscript{36} we estimated that about one-quarter of current liver transplantation procedures could be attributable to NAFLD or NASH.

Statistical analysis

We simulated disease progression by multiplying the total number of cases at a particular stage of the disease by a progression rate to the next stage, with adjustments for mortality at every step (Appendix 1, equation 1). We used Monte Carlo simulation to identify model inputs that accounted for the greatest variation in future disease burden and to produce 95\% uncertainty intervals for selected model outputs (Appendix 1, Uncertainty and sensitivity analysis).

Ethics approval

Since this was a modelling study, no ethics approval was needed.

Results

Nonalcoholic fatty liver disease

Between 2019 and 2030, the number of model-estimated NAFLD cases in Canada would increase by 20\%, to 9 305 000 (uncertainty interval 8 550 000–9 875 000) (Figure 1).

Accounting for population growth, the estimated prevalence would increase from 20.8\% (19.1\%–22.1\%) to 22.9\% (21.1\%–24.3\%), and the age-adjusted prevalence would increase from 20.8\% (19.1\%–22.1\%) to 22.3\% (20.5\%–23.7\%) (Table 1).
In 2019, the largest prevalent NAFLD age group would be 55–59 years, with 995 000 estimated NAFLD cases. By 2030, the largest prevalent age group would be 65–69 years, with 1 079 000 predicted cases (Appendix 1, Supplemental Figure S3).

Over the same period, the unadjusted number of stage F0 NAFLD cases would increase by 15%, to 7 400 000 (6 577 000–8 048 000). The number of prevalent stage F1 cases would increase by 30%, to 807 000 (542 000–1 123 000).

Figure 1: Model-estimated prevalent cases of nonalcoholic fatty liver disease (NAFLD) (A), nonalcoholic steatohepatitis (B), stage F0 NAFLD (C), stage F1 NAFLD (D), stage F2 NAFLD (E), stage F3 NAFLD (F), compensated cirrhosis NAFLD (G) and decompensated cirrhosis, hepatocellular carcinoma and liver transplantation related to NAFLD (H) for Canada, 2015–2030. Shaded areas represent 95% uncertainty interval.
Even greater relative increases are projected in later disease stages: the estimated number of stage F2 cases would increase by 45%, the estimated number of stage F3 cases would increase by 65%, and the estimated number of compensated cirrhosis cases would increase by 95%.

The number of estimated prevalent cases classified as hepatocellular carcinoma, decompensated cirrhosis or liver transplantation related to NAFLD would increase the most, by 100%, to 28 200 (17 700–44 700).

Nonalcoholic steatohepatitis

The number of prevalent NASH cases would increase by 35% between 2019 and 2030, to 2 630 000 (2 107 000–3 136 000) (Figure 1). Among the total NAFLD population, 25.2% of cases were projected to have NASH in 2019, increasing to 28.3% in 2030. The crude all-ages prevalence of NASH would increase from 5.2% (4.2%–6.2%) to 6.5% (5.2%–7.7%), and the age-adjusted prevalence would increase from 5.2% (4.2%–6.2%) to 6.1% (4.9%–7.2%) (Table 1).

Of NASH cases in 2019, 349 000 would be estimated to be stage F3–F4 fibrosis, decompensated cirrhosis, hepatocellular carcinoma or liver transplantation, encompassing 17% of estimated NASH cases and 0.9% of the Canadian population (all ages). By 2030, this number would be expected to increase by 65%, to 578 000 cases, and would account for 22% of predicted NASH cases and 1.0% of the total population.

Decompensated cirrhosis and hepatocellular carcinoma related to nonalcoholic fatty liver disease

Estimated incident cases of decompensated cirrhosis would almost double between 2019 and 2030, from 3 400 (1 900–5 800) in 2019 to 6 700 (3 800–10 900) in 2030 (Figure 2). The modelled cumulative incidence during the period would number 59 800 cases. The estimated number of incident hepatocellular carcinoma cases would increase by 80%, from 660 (440–990) to 1 200 (770–1 800), and the modelled cumulative incidence would number 11 000 cases. Estimated incident hepatocellular carcinoma was compared with reported estimates for 1992–2010, and model results were within reported ranges (Appendix 1, Supplemental Figure S4).

| Table 1: Model-estimated burden of nonalcoholic fatty liver disease and nonalcoholic steatohepatitis in Canada, 2019–2030 |
|---------------------------------------------------------------|
| **Variable** | **2019** | **2020** | **2025** | **2030** |
| Population of Canada | 37 280 000 | 37 603 000 | 39 173 000 | 40 618 000 |
| Model-estimated prevalent cases | | | | |
| NAFLD | 7 757 000 (7 138 000–8 323 000) | 7 930 000 (7 298 000–8 414 000) | 8 712 000 (8 012 000–9 244 000) | 9 305 000 (8 550 000–9 875 000) |
| Crude all-ages NAFLD prevalence rate, % | 20.8 (19.1–22.1) | 21.1 (19.4–22.4) | 22.2 (20.5–23.6) | 22.9 (21.1–24.3) |
| Adjusted all-ages NAFLD prevalence rate, %† | 20.8 (19.1–22.1) | 21.0 (19.4–22.3) | 22.0 (20.2–23.4) | 22.3 (20.5–23.7) |
| Stage F0 | 6 436 000 (5 784 000–6 956 000) | 6 553 000 (5 884 000–7 086 000) | 7 060 000 (6 300 000–7 655 000) | 7 400 000 (6 577 000–8 048 000) |
| Stage F1 | 630 000 (425 000–872 000) | 649 000 (437 000–899 000) | 736 000 (496 000–1 022 000) | 807 000 (542 000–1 123 000) |
| Stage F2 | 360 000 (235 000–508 000) | 376 000 (245 000–530 000) | 451 000 (298 000–636 000) | 518 000 (341 000–730 000) |
| Stage F3 | 216 000 (131 000–316 000) | 228 000 (139 000–334 000) | 293 000 (179 000–424 000) | 357 000 (219 000–511 000) |
| Compensated cirrhosis | 101 000 (60 400–167 000) | 108 000 (65 200–179 000) | 150 000 (91 300–242 000) | 195 000 (120 000–309 000) |
| Decompensated cirrhosis, hepatocellular carcinoma and liver transplantation | 14 000 (8 600–21 700) | 15 100 (9 300–23 500) | 21 300 (13 300–33 700) | 28 200 (17 700–44 700) |
| NASH | 1 953 000 (1 582 000–2 320 000) | 2 020 000 (1 635 000–2 401 000) | 2 345 000 (1 888 000–2 792 000) | 2 630 000 (2 107 000–3 136 000) |
| Crude all-ages NASH prevalence rate, % | 5.2 (4.2–6.2) | 5.4 (4.3–6.4) | 6.0 (4.8–7.1) | 6.5 (5.2–7.7) |
| Adjusted all-ages NASH prevalence rate, %† | 5.2 (4.2–6.2) | 5.3 (4.3–6.3) | 5.8 (4.7–6.9) | 6.1 (4.9–7.2) |
| Model-estimated incident cases | | | | |
| Decompensated cirrhosis | 3 400 (1 900–5 800) | 3 700 (2 100–6 200) | 5 100 (2 900–8 500) | 6 700 (3 800–10 900) |
| Hepatocellular carcinoma | 660 (440–990) | 710 (470–1 100) | 940 (620–1 400) | 1 200 (770–1 800) |
| Liver death | 2 700 (1 500–4 500) | 2 900 (1 700–4 800) | 4 200 (2 400–6 800) | 5 600 (3 200–9 000) |

Note: NAFLD = nonalcoholic fatty liver disease, NASH = nonalcoholic steatohepatitis.
*Except where noted otherwise.
†Adjusted to 2019 Canadian population.
Mortality

Over the study period, the estimated number of incident liver-related deaths in the total NAFLD population would double, from 2700 (1500–4500) to 5600 (3200–9000) (Figure 2), and the cumulative number of liver-related deaths would number 48 700. The total estimated number of deaths in the NAFLD population would increase by 75%, from 66 100 to 115 000. Estimated liver-related mortality would account for 4.0% of deaths in 2019 and 4.8% of deaths in 2030. Among the NASH population, the number of annual deaths was estimated to be 24 400 in 2019, doubling to 49 100 in 2030. Estimated liver-related death in the NASH population was estimated to account for 10.9% of total deaths in 2019 and 11.3% in 2030.

Uncertainty and sensitivity analysis

The leading drivers of uncertainty for the number of estimated NASH cases in 2030 included ranges around the rate of transition from stage F0 to stage F1, the starting prevalence among adults aged 20 years or more in 2018, and the standard mortality ratios applied to background mortality rates (Appendix 1, Supplemental Figure S5).

Interpretation

The current analysis shows that NAFLD may represent a growing burden on the Canadian health care system over the next decade. The population of Canada is aging and will be prone to increased disease progression to advanced fibrosis in the coming years.

Model estimates from this analysis are similar to projections for other countries where the same modelling was applied. Differences between countries in changes in disease burden are a result of different length of model study period, changes in the background population (population change and aging), and the timing and magnitude of the obesity epidemic in each country.

Other investigators have applied a modelling framework to consider the economic impact of NAFLD, including quality-adjusted life-years. A large portion of estimated costs were among patients with simple steatosis, who were the largest estimated group in the current analysis. However, it is unknown what proportion of patients with simple steatosis are diagnosed, under care and incurring costs, given the burden of occult liver disease, including compensated cirrhosis.

Estimation of quality-adjusted life-years is an important measure for estimating disease burden, but challenges exist for understanding NAFLD in this context. Given the common comorbidities among patients with NAFLD, decreased quality of life may be negligible, as utility measures are not additive. Future research should focus on the burden of NAFLD in terms of quality-adjusted life-years, with attention to advanced cases and adjustment for comorbidities.

Challenges exist for modelling based on health records. Chronic liver disease, even cirrhosis, may go unrecognized in primary care, but algorithms exist for identifying advanced cases. Recorded hospital admissions for NAFLD increased sharply during 2007–2014 in the United States; however, varied diagnostic modalities with different sensitivity levels, along with increased awareness, likely contributed to an unknown degree. Mortality data are subject to unusable or insufficient coding of cause of death, and cause of death in NAFLD cases may be coded as complications of cirrhosis or other complications of metabolic syndrome. Rates of mortality related to cardiovascular disease and nonliver cancers are elevated in NAFLD cases, and the relation between these comorbid conditions and elevated mortality risk has not been fully quantified.

The time frame of our analysis aligns with 2030 targets to reduce mortality from noncommunicable disease by one-third. Interventions for obesity will reduce NAFLD disease burden. They should follow a multipronged approach, including...
“1) health services and clinical interventions that target individuals, 2) community-level intervention to influence behaviours, and 3) public policies that target broad social or environmental determinants.” Lifestyle modification interventions that result in weight loss of more than 10% can result in resolution of NAFLD and regression of fibrosis.

There are opportunities for intervention among Canadians aged 6–17 years, as one-third were classified as overweight or obese based on the Canada Health Measures Survey, and this rate remained stable between 2004 and 2014. Without intervention, these people will contribute to an ever-growing disease burden in future decades and may experience progression to advanced disease at an earlier age.

Limitations
Challenges to understanding NAFLD disease burden include the lack of consistent definitions for measuring prevalence in general populations and the potential failure to identify a substantial portion of the NAFLD population owing to the lack of simple and widely available diagnostic techniques with high positive predictive value. Estimates of disease burden are usually based on historical data, which may not represent the current situation in an epidemic of increasing obesity, especially among younger people. A lack of consistent diagnostic measures means that reported NAFLD prevalence rates vary between studies, with NAFLD diagnosis often considered an incidental diagnosis when steatosis is found on ultrasonography. Longitudinal studies of disease using non-invasive measures in general populations should inform modelling, as in such studies more data are reported for longer periods and noninvasive measures are refined.

We calculated the number of new NAFLD cases over time using changes in obesity, classified on the basis of body mass index. Although waist circumference has been shown to be a better measure of visceral fat, which is central to the development of NAFLD, measures at the national level are available for limited time points, which makes comparisons problematic. We ultimately selected body mass index because of the availability of nationally representative longitudinal data over several decades. However, it is important to note that trends using body mass index may not capture changes in the burden of disease in people with normal body mass index who are metabolically unhealthy, among whom the prevalence of NAFLD has been estimated to be 7% in the US and 19% in Asia.

We compared liver cancer incidence to model outputs. A limitation of this comparison is that overall liver cancer incidence may be underestimated, and levels of underreporting are continuously changing, so that application of historical estimates may result in overestimation when applied to the latest data. There is further uncertainty and changes over time with regard to the cause of liver cancer, with viral hepatitis expected to contribute relatively fewer cases in the future.

There are several uncertainties when considering future liver transplantation, including the overall demand for and availability of donor organs, and changes in disease burden for competing indications, such as viral hepatitis. We assumed that the total annual number of transplantation procedures would remain constant in the future, but there is evidence that the proportion of NAFLD-related transplantation procedures is growing in Western countries. Nonalcoholic steatohepatitis is also a growing cause of transplantation among patients with hepatocellular carcinoma. Furthermore, the fact that there are overlapping indications for transplantation means that some transplantation procedures for other primary indications may also be related to NAFLD but not identified as such.

Our model excluded data on comorbidities that are confounded with NAFLD, including diabetes and dyslipidemia, to adjust model inputs. However, we applied excess background mortality to adjust for competing mortality risks among NAFLD cases. Finally, we did not adjust base prevalence and disease progression for ethnicity, although this factor may be important for understanding disease burden.

Conclusion
Nonalcoholic fatty liver disease may represent a growing burden on the Canadian health care system over the next decade. Increasing rates of obesity, in combination with an aging population, translate into increasing NAFLD-related cases of cirrhosis, hepatocellular carcinoma and related mortality. Prevention efforts should be aimed at reducing the incidence of NAFLD and slowing fibrosis progression among those already affected.

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