HEPATOLOGY

Nonalcoholic fatty liver disease burden: Australia, 2019–2030
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Abstract
Background and Aim: Nonalcoholic fatty liver disease (NAFLD) and nonalcoholic steatohepatitis (NASH) account for a large and growing proportion of liver disease burden globally. The burden of NAFLD/NASH manifests in increasing levels of advanced liver disease and primary liver cancer in Australia. A Markov model was used to forecast NAFLD burden in Australia through 2030.

Methods: A model was used to estimate fibrosis progression, primary liver cancer, and liver deaths among the Australian NAFLD population, with changes in incident NAFLD cases based on long-term trends for changes in the prevalence of obesity. Published estimates and surveillance data were applied to build and validate the model projections, including surveillance data for the incidence of liver cancer.

Results: Prevalent NAFLD cases were projected to increase 25% from the current burden (5 551 000 [4 748 000–6 306 000] cases in 2019) to 7 024 000 [5 838 000–7 886 000] cases in 2030. The projected increase in the number of NASH cases (40%) was greater than that of NAFLD cases. Incident cases of advanced liver disease are projected to increase up to 85% by 2030, and incident NAFLD liver deaths are estimated to increase 85% from 1900 (1100–3300) deaths in 2019 to 3500 (2100–6100) deaths in 2030.

Conclusions: Restraining growth of the obese and diabetic populations, along with potential therapeutic options, will be essential for mitigating disease burden.

Introduction
Nonalcoholic fatty liver disease (NAFLD) is recognized as a cause of advanced liver disease globally1–3 and particularly in Australia,4 where the increasing burden of metabolic syndrome is reflected in national survey data.5 NAFLD is defined as the presence of excessive liver fat in the absence of other causes such as excess alcohol.6 This analysis tracked NAFLD cases by fibrosis stage and categorized all cases as nonalcoholic steatohepatitis (NASH) or nonalcoholic fatty liver (NAFL) (simple steatosis). NASH is associated with liver fibrosis that may progress to advanced liver disease and related mortality.7,8 Advanced age along with obesity, diabetes, and metabolic syndrome have been identified as predictors for progression to advanced fibrosis.9 Advanced liver disease and NAFLD-related primary liver cancer typically develop after significant fibrosis has occurred; however, primary liver cancer may also occur among non-cirrhotic NASH patients.10

Recent analyses have estimated the disease burden and economic costs related to NASH based on existing literature11 while dynamic modeling techniques can compensate for inherent limitations in estimates of NAFLD disease progression.12 There is an urgent need to understand the future burden of NAFLD-related liver disease in Australia. Using a modeling framework to inform decision making may facilitate development of strategies to ameliorate further increases in disease burden.

Methods
Model. In order to estimate changes over time in NAFLD burden in Australia, a Markov model was constructed in Microsoft Excel to calculate the number of NAFLD cases by disease stage beginning in 1950.12 The total population by age group and gender was tracked over time, with new NAFLD cases entering the model based upon changes in the prevalence of adult obesity in Australia. Afflicted individuals were followed through each stage of the disease including fibrosis and advanced liver disease (Fig. S1) while also accounting for all-cause mortality (including general background mortality and excess cardiovascular and non-liver cancer mortality) as well as NAFLD-related liver mortality. Fibrosis
progression rates increased with advancing age and were varied by gender with male experiencing faster progression (Table S1). Model results were compared with data and studies that estimate the incidence of primary liver cancer attributable to NAFLD in Australia. A Delphi process was used to identify and incorporate key model inputs and review outputs with local experts against available estimates of disease burden (Table S2).

During initial model development, fibrosis progression and NASH status were adjusted to ensure distribution exceeded the number of NASH-related primary liver cancer cases reported in the surveillance system,\(^\text{15}\) with adjustments based on relative rates of overweight (\(25 >\) body mass index [BMI] \(\leq 30\) kg/m\(^2\)) and obesity (\(\text{BMI} \geq 30\) kg/m\(^2\))\(^\text{14,14}\) and published odds of disease progression to advanced fibrosis.\(^\text{15}\) Meta-analysis of reported fibrosis progression rates among NAFLD and NASH cases\(^\text{1}\) has demonstrated a range so broad that modeling projections is not feasible. Therefore, ranges around fibrosis progression rates by age, gender, and fibrosis stage were back-calculated as described in a previous analysis.\(^\text{16}\) Cohort data were used for the relative increase in progression by age and gender\(^\text{17}\) with rates further modified based on results of meta-analysis\(^\text{18,19}\) and historical trends for hepatocellular carcinoma (HCC) incidence by age and gender.\(^\text{19}\) Progression rates to primary liver cancer, decompensated cirrhosis, and liver-related death were based on reported estimates\(^\text{19}\) (Table S1).

**Population and mortality.** The annual Australian population (1950–2050) by age group and gender was based on population data from the Australia Bureau of Statistics.\(^\text{20,21}\) To calculate annual mortality, estimated deaths by age group and gender from the United Nations were divided by population estimates.\(^\text{22}\) Background mortality rates were adjusted to account for incrementally increased mortality related to cardiovascular disease\(^\text{23}\) and non-liver cancers.\(^\text{24}\) Excess non-liver mortality varies by age; among a cohort of adults in the Sydney area with elevated liver enzymes, there was no significant increase in mortality reported for adults aged \(\leq 49\) years as compared with adults with normal enzyme results. Increased risk of mortality was observed among patients aged \(\geq 50\) years, and a significant mortality hazard ratio was observed for adults aged \(\geq 80\) years. Elevated non-liver mortality may also vary significantly by fibrosis stage and NASH status,\(^\text{8,25,26}\) which are heavily correlated with age. Thus, a standard mortality ratio (SMR) of 1.15 [uncertainty range: 1.00–1.30] was applied to background mortality rates for \(\geq F1\) NAFLD cases and the portion of F0 cases that were classified as NASH. F0 cases with simple steatosis were assumed to have no elevated mortality (SMR \(= 1.0\)). Uncertainty analysis was used to account for the potential that NAFLD cases experience no excess mortality with SMR = 1.0 (low) or up to 30% excess background mortality with SMR = 1.3 (high) among non-simple steatosis F0 cases. Model NAFLD-related liver deaths were calculated as a progression rate among prevalent primary liver cancer and decompensated cirrhosis cases.\(^\text{1,27–29}\)

**New nonalcoholic fatty liver disease cases.** In order to estimate relative changes in the number of prevalent NAFLD cases, BMI data were used as a surrogate. While other factors are more strongly predictive of advanced NAFLD-related disease, data for changes in the prevalence of adults in different BMI classes are available for long periods of time, allowing for the estimation of long-term trends. The growth in NAFLD prevalence was assumed to occur simultaneous to changes in the prevalence of adults classified as obese. Because of variations in cutoff levels for obesity by race/ethnicity,\(^\text{30}\) the prevalence of obesity was calculated as a weighted average using a BMI cutoff of \(\geq 25\) kg/m\(^2\) for the population classified in the Australian Census as South-East Asian, North-East Asian, and Southern and Central Asian and a BMI \(\geq 30\) kg/m\(^2\) for the remaining population. The population with an obesity cutoff of \(\geq 25\) kg/m\(^2\) was estimated at 7.5% of the total Australian population in 2006, increasing to 12.9% in 2016. Extrapolating this trend linearly, the selected Asian populations would comprise 20.3% of the Australian population by 2030, meaning that the average BMI cutoff level for adult obesity will continue to decrease over time.

Temporal changes in adult obesity were estimated by trending prevalence data from both the Australian National Health Survey\(^\text{3}\) and NCD Risk Factor Collaboration meta-analysis for Australia.\(^\text{31}\) Using the weighted average of adult obesity at different cutoff levels, the National Health Survey reported obesity prevalence at 21.6% in 1995, increasing to 31.8% prevalence in 2015. The NCD Risk Factor Collaboration adjusted obesity was estimated at 11.4% in 1975, increasing to 32.0% in 2014. Trends based on both data sources were considered for uncertainty analysis.\(^\text{3}\)

**Nonalcoholic fatty liver disease prevalence.** Among individuals aged \(\geq 15\) years in 2015, there was an assumed NAFLD prevalence rate of 25% [uncertainty range: 20–30%]. A range of reported NAFLD prevalence exists for adults,\(^\text{32}\) and expert consensus was used to estimate the likeliest prevalence among adults and a conceivable high/low range of prevalence. Adjusting for lower prevalence among persons aged \(< 15\) years, prevalence among all ages was estimated at 20.6% in 2015. The age and gender distribution of prevalent NAFLD cases was based on data from general population studies and is higher in men and those of advanced age.\(^\text{33–35}\) Prevalence among younger people (aged \(< 18\) years) is generally not estimated in large general population-based studies\(^\text{33,36}\) and was assumed to decline with decreasing age. Because of elevated competing mortality risk among NAFLD cases,\(^\text{8}\) it was also assumed that prevalence would naturally decline among the oldest age groups (\(\geq 80\) years), with peak prevalence occurring in late middle age.

**Nonalcoholic steatohepatitis prevalence.** Prevalence of NASH was calculated based on the distribution of NAFLD cases by fibrosis stage given the total NAFLD population, with rates that varied by sex and age group. NASH-related fibrosis can regress in NAFLD patients;\(^\text{5}\) however, there is considerable uncertainty around the presence and staging of NASH due to limitations of liver biopsy results.\(^\text{37}\) The model assumed that 5% of NAFLD cases without NASH could have previously experienced NASH with subsequent regression, including a portion with fibrotic changes. Increasing fibrosis stage was assumed to result in lower probability of experiencing regressed NASH, with each increased fibrosis stage resulting in an exponential decrease in regressed NASH, and the overall number of regressed NASH cases limited to 5%. Overall, 2000 \(\geq F2\) cases were classified as non-NASH NAFLD in 2019, or 0.47% of total NAFLD cases in 2019.
Liver transplants. Total annual liver transplants were reported by the Australia and New Zealand Organ Donation Registry.\textsuperscript{38} Based on expert input and analysis of diagnostic categories for transplant recipients, it was estimated that approximately 15% of current liver transplants are attributable to NAFLD/NASH. Analysis of liver transplant data from Australia and New Zealand has shown that NASH as an indicator grew from 2.0% in 2003 to 10.9% in 2017, making it the third leading indicator for liver transplants.\textsuperscript{39} Given the uncertainties around transplant demand and availability, it was assumed that the annual number of transplants would remain constant through 2030. However, this was a conservative estimate, as data already suggest that the proportion of NAFLD-related transplants is increasing in Western countries and that some portion of transplants indicated for cryptogenic or idiopathic cirrhosis are likely related to NAFLD.\textsuperscript{40} In addition, there are overlapping indications for transplant, as alcoholic liver disease and chronic viral hepatitis can coexist with NAFLD.\textsuperscript{41}

Model validation. Primary liver cancer surveillance data were used to validate the results of the model. The Australian Institute of Health and Welfare estimates that incident primary liver cancer increased from 1076 in 2005 to 2215 in 2018.\textsuperscript{42} These estimates were further adjusted for underreporting, cancer morphology, and the proportion of cancers that could be NAFLD related. For underreporting, it was assumed that 25% of cancers may not be reported to the registry, but based on a Melbourne study, underreporting could historically be as high as 50%.\textsuperscript{4} For cancer morphology, an estimated 90% of incident primary liver cancer were assumed to be classified as HCC. A study of incident HCC cases in the Victorian Cancer Registry from 2012 to 2013 reported risk factors of fatty liver disease (14%), other/unknown (6%), and more than one risk factor (27%).\textsuperscript{4} For modeling purposes, a range of 4.0% to 34.8% was considered for the proportion of HCC that could be NAFLD related.\textsuperscript{43,44} This wide range was utilized because of uncertainty and changes over time in the etiology of liver cancer, with viral hepatitis expected to contribute relatively fewer cases in the future. Cholangiocarcinoma data were incorporated in the validation, with 5% of total primary liver cancers assumed to be classified as cholangiocarcinoma and an estimated 45% of cases potentially NAFLD related. The outcomes of this analysis were compared with model-predicted incident primary liver cancer cases to ensure that the model was predictive based on surveillance data.

Results

Nonalcoholic fatty liver disease population. Between 2019 and 2030, NAFLD cases are expected to increase 25% from 5,556,000 (4,754,000–6,312,000) to 7,026,000 (5,842,000–7,890,000) (Fig. 1). Likewise, F0/F1 NAFLD cases are also projected to increase 25% from 5,124,000 (4,212,000–5,994,000) to 6,323,000 (4,946,000–7,393,000). F2 cases are expected to increase more (50%) during the same timespan, from 228,000 (142,000–345,000) cases in 2019 to 347,000 (218,000–504,000) in 2030.
524 000) cases in 2030. The F3 population is predicted to increase by 70% between 2019 and 2030, from 133 000 (79 100–193 000) to 223 000 (134 000–322 000) cases. Compensated cirrhotic cases were forecasted to increase 85% from 62 900 (37 500–105 000) cases in 2019 to 115 000 (68 700–190 000) cases in 2030. Prevalent cases of decompensated cirrhosis, primary liver cancer and liver transplants, are expected to increase concurrently from a combined 8500 (5600–14 500) to 16 000 (11 000–25 700) cases, an increase of 85% during 2019–2030. The prevalence of NAFLD in all ages is predicted to increase from 22.0% (18.8–25.0%) in 2019 to 23.6% (19.6–26.5%) by 2030. Prevalent NAFLD cases by age group and gender were compared with the distribution of the non-NAFLD Australian population (Fig. 2). In 2019, the largest prevalent NAFLD age group was aged 55–59 years, with 595 000 cases. By 2030, peak cases (687 000 cases) were observed in persons aged 60–64 years.

**Nonalcoholic fatty liver population.** The NAFL population was assumed to be cases with simple steatosis that never progressed to NASH, with a relatively small number of cases that were formerly NASH and experienced disease regression. In 2019, the NAFL population is estimated to be 4 239 000 (3 714 000–4 690 000) cases, or 76.3% of all NAFLD cases. By 2030, the
NAFL population is estimated to grow 20% to 5 178 000 (4 403 000–5 634 000) cases (73.7% of total). NAFL cases classified as ≥F2 fibrosis were also estimated to increase from 2000 cases in 2019 to 3200 cases in 2030.

**Nonalcoholic steatohepatitis population.** The number of prevalent NASH cases was projected to increase 40% from 1 317 000 (1 040 000–1 622 000) to 1 848 000 (1 439 000–2 256 000) cases during the 2019–2030 timeframe (Fig. 1). NASH cases were projected to comprise 23.7% of all NAFLD cases in 2019, increasing to 26.3% of cases in 2030. NASH prevalence in the general population (all ages) was estimated as 5.2% (4.1–6.4%) in 2019 and is expected to increase to 6.2% (4.8–7.6%) in 2030 (Table 1).

<table>
<thead>
<tr>
<th>Year</th>
<th>Country population</th>
<th>Prevalent cases</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2015</td>
<td>2020</td>
</tr>
<tr>
<td></td>
<td>23 816 000</td>
<td>25 710 000</td>
</tr>
<tr>
<td>NAFLD cases</td>
<td>4 915 000 (4 220 000–5 605 000)</td>
<td>5 710 000 (4 879 000–6 424 000)</td>
</tr>
<tr>
<td>NAFLD prevalence rate (all ages)</td>
<td>20.6% (17.7–23.5%)</td>
<td>22.2% (19.0–25.2%)</td>
</tr>
<tr>
<td>F0</td>
<td>4 211 000 (3 541 000–4 840 000)</td>
<td>4 818 000 (4 009 000–5 515 000)</td>
</tr>
<tr>
<td>F1</td>
<td>360 000 (244 000–509 000)</td>
<td>438 000 (295 000–513 000)</td>
</tr>
<tr>
<td>F2</td>
<td>186 000 (116 000–282 000)</td>
<td>238 000 (149 000–361 000)</td>
</tr>
<tr>
<td>F3</td>
<td>105 000 (62 000–153 000)</td>
<td>140 000 (83 600–204 000)</td>
</tr>
<tr>
<td>Compensated cirrhosis</td>
<td>47 900 (28 600–80 200)</td>
<td>67 000 (39 900–111 000)</td>
</tr>
<tr>
<td>Decompensated cirrhosis, HCC, and liver transplant</td>
<td>9100 (6000–15 500)</td>
<td>12 200 (8300–20 100)</td>
</tr>
<tr>
<td>NASH cases</td>
<td>470 000 (330 000–620 000)</td>
<td>620 000 (400 000–840 000)</td>
</tr>
<tr>
<td>NASH prevalence rate (all ages)</td>
<td>4.7% (3.7–5.8%)</td>
<td>5.3% (4.2–6.5%)</td>
</tr>
<tr>
<td>Incident cases</td>
<td>1 119 000 (886 000–1 380 000)</td>
<td>1 366 000 (1 078 000–1 612 000)</td>
</tr>
<tr>
<td>HCC</td>
<td>330 (220–520)</td>
<td>360 (250–550)</td>
</tr>
<tr>
<td>Liver death</td>
<td>1300 (760–2300)</td>
<td>1800 (1100–3200)</td>
</tr>
</tbody>
</table>

HCC, hepatocellular carcinoma; NAFLD, nonalcoholic fatty liver disease; NASH, nonalcoholic steatohepatitis.

![Figure 3](https://wileyonlinelibrary.com)
Nonalcoholic fatty liver disease burden

LA Adams et al.

NASH cases and 0.8% of the total population (all ages). By 2030, this number was expected to increase 75% to 354,000 cases, accounting for 19.1% of all NASH cases and 1.2% of the total Australian population.

**Nonalcoholic fatty liver disease-related decompensated cirrhosis and primary liver cancer.** Incident decompensated cirrhosis cases were projected to increase 85% from 2100 (1100–3800) cases in 2019 to 3900 (2300–6900) cases in 2030 (Fig. 1). Cumulative incidence of decompensated cirrhosis during the same time period was estimated to be 35,800 (20,800–63,200) cases. Cases of incident primary liver cancer were forecasted to increase by 75% from 420 (280–660) cases in 2019 to 730 (480–1100) cases in 2030; cumulative incident cases of primary liver cancer between 2019 and 2030 were estimated to be 6800 (4500–10,700). Modeled incident liver cancer was compared with reported estimates for 2005–2015, for which the most reliable data were available (Fig. 3). During this timespan, model incident liver cancer cases were within the expected range.

**Mortality.** In the NAFLD population, annual liver-related deaths are estimated to increase 90% from 1700 (1000–3000) to 3200 (1900–5700) cases during 2019–2030 (Fig. 1). Cumulative liver deaths during the same time period were estimated to be 29,300 (17,300–51,500). Total deaths in the NAFLD population in 2019 are estimated at 54,800 deaths increasing 55% to 85,500 deaths in 2030. In 2019, modeled liver mortality accounted for 3.1% of annual deaths in Australia’s NAFLD population, increasing to 3.8% of deaths in 2030. Among the NASH population, annual deaths are estimated to be 19,800 in 2019, increasing 75% to 34,800 deaths in 2030. Modeled liver deaths in the NASH population were estimated to comprise 8.7% (1700 deaths) of total deaths in 2019, increasing to 9.3% (3200 deaths) in 2030.

**Discussion**

The results of NAFLD modeling suggest that NAFLD presents an expanding burden such that the Australian population is expected to experience substantial increases in NAFLD-related disease burden in the coming decades. In the coming decade, NASH may become a leading indication for liver transplantation in Australia, in tandem with a reduced burden of viral hepatitis.

The current analysis utilized both literature review and expert interviews to design the model and validate model outputs. Longitudinal trends in adult obesity levels, including estimates from both the Australian National Health Survey and NCD Risk Factor Collaboration, were used. Trending of obesity estimates from both data sources shows that the period of fastest growth in obesity has passed, while future growth rates in obesity remain uncertain.

Even if obesity prevalence in Australia stabilizes, NAFLD-related morbidity and mortality are projected to rise. The Australian population aged ≥ 65 years is projected to increase by over 1.4 million persons between 2019 and 2030. With increasing average age, the population will experience greater risk for advanced liver disease. The number of primary liver cancer cases identified as NAFLD-related has already been increasing over time; however, patients with NAFLD face high rates of non-surveillance for liver cancer. Our results further confirm the continued growth in NAFLD-related liver cancer in Australia and the pressing need to better identify NAFLD cases, especially persons with significant fibrosis. Persons with diabetes represent a substantial portion of the NASH population, and physicians should consider diabetics a high-risk group. There were 1.2 million diagnosed diabetics in Australia in 2015, and approximately 20% of cases may be undiagnosed. By 2030, the diabetic population is projected to grow to 2.2–3.0 million cases.

Modeling projected disease burden is subject to multiple limitations. Some limitations apply to all disease models, such as uncertainties around the future growth of the total Australian population and the impact of immigration. Other limitations are present when modeling the growth in chronic lifestyle conditions. Trends projecting growth in the prevalence of obesity, diabetes, and other facets of the metabolic syndrome are informed by current and historical data, but such trends may change in the future.

Rates of childhood/adolescent obesity are relatively high in Australia but may be stabilizing. In the future, a larger portion of NAFLD patients will have experienced a longer duration of obesity and a potentially earlier onset of NAFLD with disease progression occurring at younger ages. In contrast, another uncertainty is the potential availability of new therapies targeted at different facets of metabolic syndrome (e.g., NASH, diabetes, and cardiovascular disease) that would impact the natural history of disease by reducing the rate of disease progression.

This analysis differs from previous work, in that assumptions for model inputs, as well as low and high ranges for sensitivity analysis, varied. This analysis considered the impact of changing obesity at two different cutoff levels (≥ 25 and ≥ 30 kg/m²) to account for changes in demographics of the population and the impact on NAFLD prevalence. In addition, this analysis included assumptions for excess background mortality rates that were adjusted for the proportion of non-NASH F0 cases, assuming that these cases did not experience elevated mortality. Given the long natural history of the metabolic syndrome and NASH, the impact of competing mortality will play a large role in future disease burden (Fig. S3).

The current analysis was calibrated and validated using surveillance data for advanced liver disease, primarily NAFLD-related liver cancer. The historical incidence of liver cancer may be underreported to a greater degree than assumed in this analysis. The relative contribution of NAFLD to liver cancer changes over time as competing risk factors for liver cancer (e.g., viral hepatitis) vary greatly over time, and others such as alcoholic liver disease remain relatively constant.

Results of analyses demonstrate growing disease burden associated with NAFLD and NASH, following the trajectory of increasing obesity in Australia. Over one quarter of Australians aged 5–17 years are overweight or obese. This may translate to increasing rates of NAFLD in younger age groups and continued high levels of NAFLD-related disease burden in the coming decades. Intervention is needed to slow the growth in obesity and metabolic syndrome. Both lifestyle modifications and other therapeutic options must be considered to avert the coming epidemic of NAFLD-related liver disease.
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### Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

### Data S1. Supplementary Information.

#### Table S1.

Fibrosis Transition Probabilities by Disease Stage, Sex and Age Group.

#### Table S2. Delphi Process.

#### Figure S1. NAFLD Disease Progression Model.

#### Figure S2. Reported Prevalence of Adult Obesity – Australia, 1975–2015.

#### Figure S3. Key Drivers of Uncertainty for Prevalent NAFLD and NASH Cases – Australia, 2030.