Use of a catalytic model to estimate hepatitis A incidence in a low endemicity country: Implications for modeling immunization policies

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Use of a catalytic model to estimate hepatitis A incidence in a low endemicity country: Implications for modeling immunization policies

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Running Title: Hepatitis A catalytic model in a low endemic country

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Abstract

Background

Evaluating the cost-effectiveness of vaccine programs with dynamic modeling requires accurate estimates of incidence over time. Since infectious diseases are often underreported, supplementary data and statistical analyses are required to estimate true incidence. We aimed to estimate the true incidence of HAV infection in Canada through a catalytic model.

Methods

Reported HAV seroprevalence and incidence data were obtained. Using a catalytic model, HAV seroprevalence data were reconciled with the corresponding cumulative risk of infection estimated from incidence data. Sensitivity analysis was conducted.

Results

The average annual reported incidence was 6.2 HAV cases per 100,000 (range 4.3-9.5) from 1980-1989 and 7.7/100,000 (5.9-10.8) from 1990-1999, indicating that Canada is a low incidence country. The seroprevalence in Canadian-born individuals (n=7 studies) was approximately 1-8% in ages <20, 1-11% in ages 20-29, 7-29% in ages 30-39, 50% in ages 40-49, and 60-82% in ages ≥50 years. Between 1980 and 1995, the catalytic model estimated an average annual incidence of 60/100,000 (95% confidence interval: 32.75, 523.83); approximately 7.73 (4.21, 67.33) times the average annual reported incidence of 7.78/100,000. For a typical birth cohort of 403,434 Canadians born in 1990, the model predicted 32,750 HAV cases by age 39, with a corresponding seroprevalence of approximately 8.12% by the year 2029.

Implications

Reliable estimates of true incidence of infectious disease are required for cost-effectiveness analysis of infectious disease programs. The catalytic model enables the synthesis of
dispersed data, quantification of data limitations, and reconciliation of these limitations to estimate true incidence for economic evaluations.

Keywords: Hepatitis A, prevalence, incidence, decision support techniques

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Conflict of interest

At the time of study conduct, BP was a full-time employee of GlaxoSmithKline, Canada. MHC, ACT, and CTB were paid to provide consulting services to GlaxoSmithKline, Canada for their vaccine-related research. AMA is a full-time employee of GlaxoSmithKline, Canada. MK has nothing to declare.
Background

Hepatitis A virus (HAV) is prominent in many areas of the world (1-3). In countries with
low endemicity, adults are at risk for infection with HAV due to high-risk activities and
disease importation (4). Safe and effective HAV vaccines have been available since the mid-
1990s (5;6). In the United States, the hepatitis A childhood immunization strategy has
evolved from vaccinating children living in communities with high and medium endemicity
in 1996 and 1999 to routine vaccination of children nationwide in 2005 (7;8). Other low
endemicity countries continue with vaccine programs targeting high-risk groups (9), although
routine vaccination programs for children or adolescents is an alternative option.

In these countries, cost-effectiveness analyses of targeted versus routine vaccination
programs are required to inform immunization policies. The cost-effectiveness ratios
between targeted and routine vaccination programs are expected to be smaller than those
comparing routine vaccinations versus no vaccination (10). Analyses based upon dynamic
models of HAV transmission could increase the sensitivity of detecting such small
differences (11). However, knowledge of the true incidence of HAV infection is required for
such transmission models. Hence, robust approaches to estimate true incidence to be used as
input data for dynamic models of HAV transmission are necessary.

In most countries, the true incidence of HAV is unknown. Reported cases are an
unreliable indicator of the true incidence because the infection is often not clinically
apparent (sub-clinical) and symptomatic cases are not always reported (under-reporting) (4).
Measuring incidence directly with serial seroprevalence surveys is impractical because it is
resource-intensive (4). Data from cross-sectional seroprevalence surveys, if available, cannot
be used to directly infer time-specific incidence without some additional modeling steps (12).
Simple mathematical expressions were first used to model age-specific changes in the force of infection (i.e., the rate at which individuals who have not been exposed to the virus [or susceptibles] acquire infection) (13). These models were described as catalytic because of their structural similarities to equations commonly used in the study of chemical reactions (13). Integrating acute disease surveillance data with seroprevalence data from two national cross-sectional surveys, a catalytic model has been used to estimate the true incidence of HAV infection in the United States (12). According to results from this study, the estimate of true incidence in the United States was approximately 10 times the reported incidence and declined by 4.5% per year over the modeling duration (12). This information was subsequently used in another study to quantify the impact of hepatitis A immunization in the United States in the late 1990’s (14).

The catalytic model application in the United States was based upon seroprevalence data from two national cross-sectional surveys (12). However, it is less clear whether such a catalytic model can be used to generate reliable estimates of true incidence in countries where national seroprevalence surveys are not available and alternative seroprevalence data are limited in scope and comprehensiveness. We aimed to estimate the true incidence of HAV infection in Canada, a low endemicity country with fragmented seroprevalence data.

Methods

Catalytic Model

Following the notation by Armstrong et al., the force of infection—the percentage of susceptible individuals of age $a$ infected in year $t$—is given by $\lambda(a, t)$ (12). We wish to know the resulting cumulative percentage infected in a cohort that is age $A$ in year $T$. The cumulative exposure $F(A, T)$ experienced by this cohort in the $A$ years since its birth is the
force of infection experienced at age 1 in year $T-A$, plus the force of infection experienced at age 2 in year $T-A+1$, ..., plus the force of infection experienced at age $a$ in year $T-A+a$, ..., plus the force of infection experienced at age $A$ in year $T$. This cumulative exposure experienced from birth ($a=0$) to the present ($a=A$) by individuals in the birth cohort can be expressed as an integral

$$F(A, T) = \int_0^A \lambda(a, t) \, da$$  \hspace{1cm} [1]$$

where $t = T - A + a$ from the arguments above. Since the cumulative exposure in the cohort is $F(A, T)$, the proportion of individuals in the cohort not infected by year $T$ is given by $\exp(-F(A, T))$, hence the proportion of individuals in the cohort infected by year $T$ is simply 1 minus that quantity, or

$$P(A, T) = 1 - \exp(-F(A, T)) = 1 - \exp\left(-\int_0^A \lambda(a, t) \, da\right)$$  \hspace{1cm} [2]$$

To simplify estimation, one can moreover assume that age and time are independent in the force of infection; hence it can be modeled as the product of two functions:

$$\lambda(a, t) = F(a) \times G(t)$$  \hspace{1cm} [3]$$

where $F(a)$ is a function describing the age-specific force of infection (and is distinct from the exposure function $F(A, T)$) and $G(t)$ is a function describing how the force of infection evolves over time.

Here, the cumulative percent infected in a cohort of age $A$ and time $T$, $P(A, T)$, is taken to be identical to the seroprevalence in the Canadian-born cohort of age $A$ at time $T$. Thus $P(A, T)$ is hereafter referred to as seroprevalence. Similarly, the force of infection $\lambda(a, t)$ can be expressed as some function of the incidence of new infections as reflected in case reports. Therefore, equation [2] makes it possible to integrate age-stratified reported
incidence data with seroprevalence data in a birth cohort at a given time in order to estimate epidemiologically important parameters such as the force of infection $\lambda(a, t)$, via model fitting (14;15).

Key steps in fitting the model include: 1) obtaining seroprevalence data; 2) obtaining reported incidence data; 3) adjusting reported incidence to reflect the age-specific and time-specific components of the force of infection $\lambda(a, t)$ in equation [3], 4) fitting the catalytic model to derive parameter estimates and long-term predictions; and 5) conducting sensitivity analyses to address uncertainties in the data (15).

Step 1: Obtaining seroprevalence data

We obtained seroprevalence data from surveys of HAV antibodies in Canadian participants. The included surveys were identified through a systematic review (16). First, the seroprevalence estimates in Canadian-born cohorts that most likely reflected local transmission patterns were derived to provide data for the response variable $P(A, T)$ in equation [2] (Figure 1A). Second, the seroprevalence estimates $P_{all}(a)$ that reflected the total immunity level in the population were derived using data from all Canadians, including foreign-born individuals who might have been infected prior to immigration. The related estimates $(1 - P_{all}(a))$ denoting the proportion of susceptible individuals in age class $a$ in the population were necessary for the adjustment in step 3 (Figure 1B).

The age-specific seroprevalence estimates $P_{all}(a)$ were derived and if necessary, pooled across survey studies using the Dersimonian-Laird random-effects meta-analytical approach (17), for age groups <10 (n=1 study) (18); 10-19 (n= 6 studies) (18-23); 20-29 (n = 6 studies) (19;21-25) 30-39 (n= 5 studies) (19;21;22;24;26); 40-49 (n = 3 studies) (21;22;26); and >50 (n = 4 studies) (19;21;25;26). Across surveys, the reported estimates $P_{all}(a)$ varied substantially with respect to studied populations, timing, and variation in the reported age.
groups (19). The impact of this heterogeneity on modeled estimates and projections was assessed via sensitivity analyses (step 5).

Step 2: Obtaining reported incidence data

Data pertaining to the annual age-specific reported incidence $I(a)$ from 1980 to 2003 were obtained from the National Notifiable Disease Registry of Canada (27). According to the national surveillance program, the case definition is a laboratory confirmation of HAV infection (i.e., positive immunoglobulin M antibody to HAV) with or without symptoms. Hepatitis A incidence is known to vary cyclically, with peaks in outbreak activity every 6-8 years depending on place and time. This cyclicity is apparent in Figure 2 where one observes peaks in 1985 and 1991. Incidence appears to be climbing again after a trough in 1994, but it is clear by the year 2000 that this pattern has been interrupted. The vaccine was first introduced in targeted groups and for outbreak control in the mid 1990s (28). The main analysis was conducted using data from 1980 to 1995 where the reported incidence data were not under any influence HAV vaccination.

Step 3: Adjusting reported incidence

First, the age-specific component of the force of infection $F(a)$ in equation [2], defined as the percent of susceptible individuals of age $a$ infected per year, was estimated as follows:

$$F(a) = \beta \cdot I_{adj}(a) = \frac{\beta \cdot I(a)}{(1 - P_{all}(a)) \cdot P(J | a)}, \quad [4]$$

where the annual reported incidence $I(a)$ represents the percentage of individuals of age group $a$ becoming infected per year. For each age group $a$, $I(a)$ was averaged from 1980-1995 to remove any short-term cyclical effects (29). Similar to the approach proposed by Armstrong and Bell approach (12), we made four adjustments to the reported incidence $I(a)$ in order to capture its contribution to the force of infection.
Adjustment 1: the reported incidence $I(a)$ was divided by $P(J \mid a)$, the probability of developing jaundice during acute infection in age group $a$, to account for sub-clinical infections which were assumed not to have been recognized or reported, thus yielding the true percentage of infections in age group $a$ (4). Here we assumed that HAV infections without symptoms of jaundice were less likely to trigger a laboratory confirmation according to the case definition described in step 2. The estimated age-specific probabilities of jaundice $P(J \mid a)$ came from Armstrong and Bell (12), indicating high sub-clinical infections in children and adolescents. It was approximately 7.2% (95% confidence interval: 4.7-10.9%) in ages 0-4, 37.1% (30.7-43.8%) in ages 5-9, 70.7% (58.8-79.4%) in ages 10-17, and 85.2% (79.9-89.2%) in ages ≥18.

Adjustment 2: The true percentage of infections $I(a) / P(J \mid a)$ was then divided by the proportion susceptible $(1 - P_{all}(a))$ since the force of infection is defined as the percentage of infections in the part of the cohort that is susceptible. This quantity is hereafter referred to as the $I_{adj}(a)$, the adjusted incidence in susceptibles;

Adjustment 3: Finally, $I_{adj}(a)$, was further adjusted by an under-reporting factor $\beta$ ($\beta > 1$), representing any other sources of under-reporting in addition to what results from subclinical infection.

Adjustment 4: A common pattern that was observed in past decades in many advanced countries was a declining risk of infection due to improved sanitation and hygiene, which implies a declining force of infection (8). Hence, in equation [2], the time-specific component of the force of infection was assumed to decline exponentially from 1980 onward at an annual rate $\delta$ such that

$$G(t) = G_0 \exp [-\delta t], \quad [5]$$
where \( G_0 \) is a constant (12). We note that this gradual decline typically occurs over a longer timescale than the 6-8 year timescale of the natural cycling of hepatitis A incidence, which is caused by a separate phenomenon. We also note that this long-term decline is evident in the “S-shaped” age-stratified seroprevalence profile in Canadian-born individuals in Figure 1A. In contrast to the low seroprevalence in recently born cohorts, the high seroprevalence in older cohorts reflect high risk exposure in previous decades. Models that assume a constant force of infection cannot reproduce the observed S-shaped seroprevalence profiles (20;30).

Step 4: Fitting the catalytic model

Combining equations [2]-[5] yields

\[
P(A,T) = 1 - \exp \left( \int_0^A \beta * I_{adj}(a) * \exp[\delta * (M - T + A - a)] da \right), \quad [6]
\]

where \( M \) denotes the mid-point of the years in which the reported incidence data were used and accounts for the constant \( G_0 \) in equation [5]. The paired parameter \((\beta, \delta)\) was estimated by minimizing the difference between the prevalence \( P(A, T) \), which uses the seroprevalence data while the corresponding right side of equation uses the reported incidence data [6].

A smoothing function was fitted to the relationship between \( I_{adj}(a) \) and age \( a \) to obtain a continuous approximation to the discrete age-structure in the reported incidence data (Appendix). The catalytic model was fitted in two steps. First, \( \delta \) was varied by increments of 0.01 from 0.00001 – 1.0 (12). Given a fixed value of the parameter \( \delta \), the expression \( \log(1-P(A,T)) \) according to equation [6] becomes a linear function of the parameter \( \beta \). A least-square estimate of \( \beta \) was then derived from a univariate linear regression of \( \log(1-P(A,T)) = \hat{\beta} * X \). Model goodness-of-fit was assessed using the fitted statistic \( R^2 \) and residual plots (31). Fitted values of the paired parameter \((\hat{\delta}, \hat{\beta})\) were selected to maximize the fitted
statistic $R^2$. The 95% confidence intervals for estimates of $(\hat{\delta}, \hat{\beta})$ were derived by varying estimates of the age-specific sub-clinical infection probability $P(J | a)$ from their lower to upper 95% confidence intervals (12). Age-specific estimates of the adjusted incidence $I_{adj}(a)$ were derived according to equation [4] using the fitted values $\hat{\beta}$. Estimates of the force of infection $\lambda(a,t)$ for a hypothetical birth cohort of 1990 ($n=403,434$) were derived via equation [2] using the fitted values $(\hat{\delta}, \hat{\beta})$. Corresponding estimates of icteric or symptomatic infection incidence were calculated by applying the age-specific probability of jaundice $P(J | a)$ to estimates of the true incidence $F(a)$.

Step 5: Sensitivity analysis

In the main analysis, reported incidence data from 1980-1995 and seroprevalence data from studies with data collection in 1995 or before were used. Incidence data from 1995 to 2003 was in part influenced by targeted HAV vaccination programs, as the vaccines were first used around 1995 in Canada (5). Including this late time period, however allowed the inclusion of prevalence data from recent studies (Figure 2). In a sensitivity analysis, data from 1980 to 2003 were used.

Age-specific estimates of HAV prevalence among all Canadians varied substantially (Figure 2). In the main analysis, the pooled estimates from random-effects models for $P_{all}(a)$ were used. Sensitivity analyses were also conducted by using the lower and upper bounds of the 95% confidence intervals of the pooled estimates.

Results

The systematic search yielded 7 studies (18;19;21;22;24-26) reporting 20 point estimates for seroprevalence among Canadian-born participants (Figure 1A) (16). Seroprevalence samples
collected from these studies occurred between 1980 and 2003 (Figure 1). The seroprevalence in Canadian-born individuals was approximately 1-8% in ages <20, 1-11% in ages 20-29, 7-29% in ages 30-39, approximately 50% in ages 40-49, and 60-82% in ages ≥50 years (Figure 1A). A similar pattern was observed for all Canadians including foreign-born individuals, with large variation in each age group (Figure 1B). Point estimates in the age groups >40 were high and most likely influenced by the cohort effect; meaning that the high seroprevalence reflected life-long immunity from HAV infection experienced by birth cohorts in past decades (Figure 1A) (25;32).

In Canada, the average annual reported incidence was 6.2 cases per 100,000 (range 4.3-9.5) from 1980-1989, and 7.7 (5.9-0.8) from 1990-1999 (Figure 2). Coinciding with the use of HAV vaccines in targeted groups and outbreak control, there appeared to be a decreasing trend from 1995 to 2003, reaching below 2 cases per 100,000 from 2000-2003.

Modeled estimates of seroprevalence fitted the observed seroprevalence reported in survey studies for age <40 (Figure 3). Limited seroprevalence data were available for modeled estimates in ages ≥ 40 (n=3 data points, Figure 3). The annual rate of decline in the force of infection over time was estimated to be $\delta = 0.12$ (95% confidence interval: 0.08, 0.15; Table 1). The under-reporting factor estimated from the linear regression involving the response $\log(1-P(A,T))$ derived from equation [6] was $\beta = 7.73$ (4.21, 67.33), with a goodness-of-fit statistic $R^2 = 0.86$. This under-reporting estimate suggests that for every reported HAV case, approximately 8 cases were not reported. In the sensitivity analysis, estimates of the annual declining rate $\delta$ and under-reporting factor $\beta$ exhibited large confidence intervals. The maximum boundaries across different confidence intervals for $\delta$ ranged from 0.05 to 0.15, and for $\beta$ from 4.21 to 76.83 (Table 1).
The reported incidence was adjusted to better reflect estimates of the force of infection (Figure 4). In ages 0 to 9, reported incidence substantially underestimated the true incidence, most possibly due to sub-clinical infection. For age groups ≥30, reported incidence also underestimated true incidence. Note that the seroprevalence estimates among all Canadians increased with age, leading to a decrease in the relative size of susceptibles in the older age groups and an increase in the gap between reported and adjusted incidence according to equation [4].

The model estimated an average annual incidence of 60.14 HAV cases per 100,000 in Canada (95% confidence interval: 32.75, 523.83); approximately 7.73 times the average annual reported incidence of 7.78 cases per 100,000 from 1980 to 1995. For a typical birth cohort of 403,434 Canadians born in 1990, the model predicted 32,750 HAV cases by age 39, with a corresponding prevalence of approximately 8.12% by the year 2029 (Table 2). The cumulative number of cases in older age groups was predicted taking both the cohort effect and susceptibility into account (Table 2).

Discussion

This paper provides an example of how catalytic modeling can be employed to bridge the gap between reported incidence and seroprevalence data in order to estimate the true incidence pattern over time in a population. This information can be used to understand transmission dynamics of infectious diseases over time by distinguishing age and cohort effects on the force of infection. This type of modeling is particularly salient for epidemiologists and economic modelers attempting to estimate the future effects of infectious disease control programs, particularly vaccines, on a population’s health.
Cost-effectiveness analyses of disease control programs often rely on cohort models, which essentially consider the individual in isolation from the population and do not take population-level mechanisms of disease transmission into account (10). They implicitly assume that the force of infection does not change when control programs are implemented, which is rarely the case. Hence, they can either under-estimate or over-estimate cost-effectiveness of the intervention (11). In contrast, dynamic models take into account mechanisms that govern the time evolution of epidemiological measures, such as the number of infected individuals, mortality rates, and force of infection. In principle, they are more accurate than corresponding cohort models for cost-effectiveness analysis of control programs (10;11). However, application of dynamic models requires knowledge of the true incidence of infection; hence catalytic modeling or a similar approach is necessary.

The catalytic model described in this research was first proposed by Armstrong and Bell (12), where they reconciled the reported incidence of HAV in the United States with the observed prevalence of HAV antibody from two nation-wide surveys (12). In the current application of the model, fragmented and heterogeneous seroprevalence data identified through a systematic view was used (16). The data was of low quality (16), substantially contributing to large confidence intervals around key parameter estimates. Consequently, the model was not robust as it is constrained by the lack of reliable data.

An advantage of catalytic modeling is that it allows the identification of cohort effects (i.e., changes in the force of infection on generational timescales) (2). Cohort effects are reflected in S-shaped age-structured seroprevalence profiles (e.g., Figure 2). Age-specific data from a prevalence survey at one point in time can not distinguish between age- and time-dependence (33). Data from multiple studies are under the influence of both time-specific and age-specific effects. Note that HAV transmission depends on hygiene, food
preparation, or contact with infective individuals. These factors change over time and risk-related behaviors could change across generations. Estimates of true incidence are biased unless both age- and time-specific effects in seroprevalence data are modeled (33). The current model fulfilled this requirement. Failure to account for cohort effects has wide implications for interpreting seroprevalence data and predicting the impact of vaccination programs with dynamic models (30).

Simple catalytic models have been used previously to describe the age-specific prevalence of susceptibility to rubella virus (34;35), to estimate HAV prevalence rates over times as part of an assessment of the temporal relationship between childhood acute lymphoblastic leukemia and hygiene conditions (36), and to estimate true incidence of primary toxoplasmosis infection (33), HCV infection (37), and HSV-2 infection (38). Catalytic modeling was also used to estimate age-related rates of measles infection (14), and to study the age distribution of attack of measles (29).

Early catalytic models focused on the functional form of the age-specific force of infection $\lambda(a)$ (13). Other functional forms have been used: constant since birth (39), constant over the range of data, but not necessarily since birth (13;29;40); higher order polynomials (14;41); and an exponentially damped linear function (42). Nonparametric functions have also been introduced (43). Ades and Nokes suggest that flexible functional forms might have the disadvantage of occasionally generating extreme curvatures at the edges of the data, precisely where prediction of future trends is desired (33). Griffiths provided an approach to graphically presenting observed age distributions to reveal the nature of dependence of the true incidence upon age (29).

In the current model, the time-dependent component of the force of infection (i.e., $G(t)$) was modeled using an exponential function to allow for the cohort effect. However,
recent research suggests that the temporal evolution of the force of infection in Canada in the past century can be well-described by a sigmoidal function, where transmissibility declines from a high constant level to a low constant level with a relatively rapid period of transition in between (30). The period of rapid transition may correspond to significant post-war improvements in sanitation and hygiene, for example. The use of an exponential function may partly explain the shape of the curve in Figure 3, where seroprevalence appears to be increasing exponentially with age in the model but not in the data. A sigmoidal function might produce a better fit to the data, and because it is not piecewise, yields an easier analysis. Future research may explore the use of a sigmoidal function in catalytic modeling.

In addition to offering an example of the use of catalytic modeling with fragmented and limited data, our results also highlight the potential biases associated with the use of readily-available epidemiological data in cost-effectiveness evaluation of infectious disease control programs. Data from notifiable disease reporting substantially under-estimates true incidence due to sub-clinical symptoms in HAV infection (4) and under-reporting of clinical cases (12). Prevalence data from cross-sectional surveys was fragmented and suffered from a cohort effect; the high prevalence in the older age groups most likely reflected infection risk of past decades (20;30). Reported prevalence from surveys including foreign-born subjects might not exclusively reflect local transmissibility. The uncertainty induced by these potential sources of bias in the input data should be recognized, quantified, and assessed in cost-effectiveness analyses of infectious disease control programs.

A number of key assumptions were made in the catalytic model we used. First, we assumed that HAV infection leads to life-long immunity. The age-dependent force of infection was assumed to be constant and the susceptible population homogeneous with
respect to infection exposure. A population with stable age distribution was assumed, as it was convenient to ignore deaths and migration to and from the population (29). With respect to input data to the model, seroprevalence estimates from small studies and heterogeneous study populations were generalized to represent Canadian estimates.

In summary, catalytic models integrate reported incidence and seroprevalence data providing reasonably reliable estimates of past, current, and projected force of infection. They provide a logical structure to link highly dispersed data, quantify limitations in the data sources, and reconcile these limitations to derive estimates of true incidence. They are useful to epidemiologic and economic modelers attempting to estimate the cost-effectiveness of disease control programs through dynamic modeling.
A 2-parameter exponential function form was used for the relationship between $I_{adj}(a)$ and age $a$, $I_{adj}(a) = \exp(-\alpha a^\tau)$. Fitted values for $(\alpha, \tau)$ were obtained through fitting the univariate linear regression $\log(-\log(I_{adj}(a))) = \alpha + \tau \log(a)$. This resulted in $\alpha = 5.40$ (95% confidence interval: 5.10, 5.87), $\tau = 0.15$ (0.13, 0.18), and a goodness-of-fit statistic $R^2$ of 0.92.
### Figure 1: Age-specific prevalence (%) abstracted from clinical studies identified through a systematic review (16)

#### Notes:
The column “Author” displays the first author and year of publication of the included studies; “Population” the major age groups, participant characteristics, and year of data collection in the included studies; “Age” the age groups according to the original study reports; “n” sample size. Plotted lines are percent estimates of participants with positive HAV antibodies and their corresponding 95% confidence intervals. A total of 7 studies contributed 20 data points in Panel 1A and 10 studies contributed 30 data points to Panel 1B.

### Abbreviations:
NS Nova Scotia, BC British Columbia, G6 Students Grade 6 students, and CF Canadian Forces.
Figure 2: Age-adjusted reported incidence of hepatitis A in Canada from 1980 to 2003
Figure 3: Observed seroprevalence reported in seven clinical studies identified from the systematic review (i.e., 20 data points are represented by the dots) and corresponding predicted prevalence denoted by the fitted line.
Figure 4: Reported and adjusted average annual incidence of hepatitis A in Canada, 1980-1995.

Notes: Reported incidence data was available for each decade of age except for the age group 40-59 and 60+. The blue bars show the unadjusted reported incidence $I(a)$. The red bars show incidence after adjustment for sub-clinical infection using the age-specific probability of jaundice $P(I|a)$. The yellow bars show incidence after further adjusting the population denominator for the relative size of susceptibles $[I(a)/(1-\mu(a))]$. Additional adjustment for under-reporting was possible according to expression [3] (data not shown in the graph).
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<tr>
<td>Main analysis with pooled estimates¹</td>
<td>0.12 (0.08, 0.15)</td>
<td>7.73 (4.21, 67.33)</td>
</tr>
<tr>
<td>Sensitivity analysis with lower bound²</td>
<td>0.09 (0.07, 0.13)</td>
<td>16.56 (6.89, 76.83)</td>
</tr>
<tr>
<td>Sensitivity analysis with upper bound³</td>
<td>0.13 (0.05, 0.14)</td>
<td>3.88 (2.69, 48.23)</td>
</tr>
</tbody>
</table>

Table 1: Parameter estimates from the main analysis and corresponding sensitivity analyses.

Notes: ¹Pooled estimates from random-effects model of HAV prevalence in all Canadians (data from Figure 2 – Panel B) were used to adjust reported incidence data for the proportion susceptibles (i.e., in expression [3]). ²Lower bounds from 95% confidence intervals of the pooled estimates from footnote 1. ³Upper bounds from 95% confidence intervals of the pooled estimates from footnote 1.

Abbreviation: CI confidence interval.
<table>
<thead>
<tr>
<th>Age Group</th>
<th>Year</th>
<th>HAV cases*</th>
<th>Icteric cases*</th>
<th>HAV cases / 100,000</th>
<th>Icteric cases / 100,000</th>
<th>Estimated Prevalence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-9</td>
<td>1990-1999</td>
<td>7,261</td>
<td>362</td>
<td>1,800</td>
<td>90</td>
<td>1.80</td>
</tr>
<tr>
<td>10-19</td>
<td>2000-2009</td>
<td>3,954</td>
<td>492</td>
<td>980</td>
<td>122</td>
<td>2.78</td>
</tr>
<tr>
<td>20-29</td>
<td>2010-2019</td>
<td>12,346</td>
<td>599</td>
<td>3,060</td>
<td>148</td>
<td>5.84</td>
</tr>
<tr>
<td>30-39</td>
<td>2020-2029</td>
<td>9,198</td>
<td>238</td>
<td>2,280</td>
<td>59</td>
<td>8.12</td>
</tr>
</tbody>
</table>

**Table 2:** Long-term projection from the catalytic model for true incidence of HAV infection in a Canadian birth cohort of 1990 (n=403,434).

**Note:** *Cumulative number of cases in each age group (e.g., 7,261 cases from birth to age 9 in the first age group).
Reference List


(5) Scheifele DW, Oehnio J. Hepatitis A vaccine: is it being used to best advantage? CMAJ 2002 Jul 9;167(1):44-5.


(22) Cook D, Wilton L, Patrick D, Zou S, Sherman M, Krajden M. Prevalence of antibodies to hepatitis A virus in a cohort of women of child-bearing age. 68th Canadian Association for Clinical Microbiology and Infectious Disease, Nov 5-8, 2000 - Ottawa . 11-5-2000.

Ref Type: Generic


Ref Type: Abstract


