An estimated 0.64%–0.71% of Canadians (220,000–245,000 people) have chronic hepatitis C virus (HCV) infection,1 and approximately 44%2 of those may be undiagnosed. HCV can be transmitted directly through percutaneous exposure (e.g., through inadequately sterilized medical equipment) or through receipt of contaminated blood products.3 People who inject drugs are at highest risk, but recipients of unscreened blood products, tissues or organs and patients undergoing long-term hemodialysis are also at increased risk.3 Less common modes of transmission include vertical transmission, high-risk sexual contact, unsterilized tattoo or piercing equipment, and occupational exposure.3 Not all people with chronic HCV infection will develop cirrhosis or signs or symptoms indicative of liver disease.4 It is estimated that approximately 84% of people infected with HCV do not develop cirrhosis 20 years after acute infection, and 59% after 30 years.5,6 Progression of liver fibrosis is variable and influenced by factors such as alcohol consumption, age at time of infection, male sex and HIV coinfection.7

Efficacy of treatment for HCV is typically evaluated using sustained virologic response (SVR), a surrogate outcome.8 Sustained virologic response is defined as having undetectable HCV ribonucleic acid (RNA) at 12, 24 or 72 weeks following completion of treatment (SVR12, SVR24 or SVR72). Historically, HCV treatment involved 24–48 weeks of pegylated interferon injections and oral ribavirin (pegylated interferon-ribavirin), which had substantial adverse effects.9 In 2011, Health Canada approved the use of direct-acting antiviral drugs (DAAs) in combination with pegylated interferon-ribavirin, which improved the likelihood of sustained virologic response but increased adverse effects.10,11 In 2016, Health Canada approved the use of pangenotypic interferon-free DAA regimens12 for treatment of HCV. Interferon-free DAA regimens are very costly, but have several advantages: they are administered orally, require shorter treatment duration and have higher likelihood of sustained virologic response, with fewer adverse effects than older regimens.9

There are no organized HCV screening programs for the general population in Canada,13 although the Public Health Agency of Canada (PHAC) and the College of Family Physicians of Canada (CFPC) recommend testing for hepatitis C in people at elevated risk of HCV infection.14

Scope

The Canadian Task Force on Preventive Health Care has not previously developed a recommendation on screening for HCV. Reasons for developing this recommendation include the availability of new treatments for chronic HCV infection and the lack of Canadian guidelines for screening.13 The current recommendations are intended to provide clinicians and policy-makers with guidance on screening asymptomatic Canadian adults for HCV.
Methods

The task force is an independent panel of clinicians and methodologists that makes recommendations about clinical interventions aimed at primary and secondary prevention. These recommendations were developed by a workgroup of seven members of the task force with scientific support from PHAC. The recommendations were informed by two independently conducted systematic reviews that addressed specific aspects of the guideline’s analytic framework (Appendix 1, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.161521/-/DC1). The reviews excluded post-transplant patients, patients with HIV, patients on hemodialysis and patients with occupational exposure; studies on all other population groups were sought, including higher-risk groups (e.g., with history of injection drug use) and higher-prevalence groups (e.g., birth cohort).

The first set of reviews was conducted by the Canadian Agency for Drugs and Technologies in Health (CADTH) and attempted to identify evidence on 1) the effectiveness (benefits and harms) of HCV screening on HCV-related mortality and morbidity, including cirrhosis, hepatocellular carcinoma, rate of liver transplantation, quality of life, sustained virologic response rates, histological improvement, behavioural changes to improve health outcomes, reduced HCV transmission, anxiety, labelling, partner discord, abuse or violence, and overdiagnosis or overtreatment; 2) patient preferences and values in relation to screening; 3) the clinical validity of screening tests; and 4) the cost-effectiveness of screening. The peer-reviewed literature search was conducted in MEDLINE, Embase, PubMed and the Cochrane Library. For research questions on clinical effectiveness (benefits and harms), cost-effectiveness, and patient preferences and values, the search time frame was from January 2000 to March 2016; no date filters were applied to the question on the clinical validity of screening tests, which was run in June 2016. For all review questions, grey literature was sought using CADTH’s Grey Matters checklist up to September 2016.

PHAC conducted a separate systematic review, which focused on the effectiveness of new HCV treatments compared with older treatments, on mortality (all-cause or hepatic), cirrhosis (compensated or decompensated), hepatocellular carcinoma, need for liver transplantation, quality of life, sustained virologic response, improvement in liver histology, reduced HCV transmission, withdrawals due to adverse events, neutropenia, anemia, psychological adverse events, flu-like symptoms and rash. The literature search was an update to a therapeutic review conducted by CADTH in February 2015, which was further expanded by PHAC to include PubMed and ClinicalTrials.gov to November 2015.

Prepublication searches were conducted to update both reviews. The literature search for the screening review and its grey literature search were updated to Dec. 11, 2016. The literature update for the treatment review was updated to Nov. 21, 2016, but was limited to randomized controlled trials (RCTs) published in MEDLINE, Embase and the Cochrane Library.

The Knowledge Translation group of St. Michael’s Hospital (Toronto, Ontario) engaged members of the public on behalf of the task force at two stages of the guideline development process. For phase one, 19 participants from the HCV screening and treatment populations rated outcomes to be included in the systematic reviews, using two online surveys and a focus group. For phase two, 15 asymptomatic, average-risk or increased-risk participants were recruited to provide their perception of the importance of considering harms, benefits and costs in making screening decisions.

A modelling study was also commissioned by the task force and conducted by a team from the Toronto Health Economics and Technology Assessment Collaborative. This modelling study was used to examine the possible impact of screening under certain circumstances on hepatic mortality, hepatocellular carcinoma and decompensated cirrhosis (Appendix 2, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.161521/-/DC1). In the absence of empirical data, expert opinion was sought for four key model input parameters: 1) expected uptake of screening in the general population, 2) uptake of treatment in asymptomatic individuals, 3) genotype distribution, and 4) distribution of hepatic fibrosis scores in a primary care setting. Working group members identified five HCV experts, three of whom provided parameter estimates and ranges and the location of possible supplemental data supporting their estimates. Parameter estimates were established by calculating the mean value of the responses that the experts provided. The latest prevalence estimates from PHAC (2014) were used as inputs for the model for the general population and various subgroups.

The Feasibility, Acceptability, Cost, and Health Equity (FACE) tool was used with organizational stakeholders to gain their perspective on the priority, feasibility, acceptability, cost and equity of the recommendation. The FACE survey was pilot-tested with this guideline as part of a validation exercise. Stakeholder organizations that provided input on the recommendation using the FACE survey are listed in Appendix 3 (available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.161521/-/DC1).

The Grading of Recommendations Assessment, Development and Evaluation (GRADE) system was used to determine the quality of evidence and strength of recommendations (Box 1). The recommendations were revised and approved by the entire task force and underwent external review by academic and clinical experts. More information about task force methods can be found elsewhere. The task force plans to update this recommendation as new evidence or other compelling information about key factors influencing the recommendation becomes available (see Rationale section for details).

Recommendations

We recommend against screening for HCV in adults who are not at elevated risk (strong recommendation, very low-quality evidence).

The recommendation has been graded according to the GRADE system described in Box 1. A summary of the recommendations for clinicians and policy-makers is shown in Box 2.

Screening

The screening systematic review found no studies (RCTs, non-randomized studies with a control group, or disease progression modelling studies) of the effectiveness of HCV screening in the
Box 1: Grading of recommendations
Recommendations are graded according to the Grading of Recommendations Assessment, Development and Evaluation system (GRADE).18 GRADE offers two strengths of recommendation: strong and weak. The strength of recommendations is based on the balance between desirable and undesirable outcomes; the confidence in the magnitude of the estimates of effect of the intervention on outcomes; the confidence in values and preferences and their variability; and whether the intervention represents a wise use of resources.

Strong recommendations are those for which the task force is confident that the desirable effects of an intervention outweigh its undesirable effects (strong recommendation for an intervention) or that the undesirable effects of an intervention outweigh its desirable effects (strong recommendation against an intervention). A strong recommendation implies that most individuals will be best served by the recommended course of action and that the recommendation can be adopted in practice or as policy in most situations.

Strong recommendations are normally based on high-quality evidence (i.e., high confidence in the estimate of the effect of an intervention). Strong recommendations may recommend in favour of an intervention (when there is high confidence of benefit) or against an intervention (when there is high confidence of harm). However, there are five circumstances in which the Task Force may consider a strong recommendation based on low- or very low-quality evidence:8

1) When low-quality evidence suggests benefit in a life-threatening situation (evidence regarding harms can be low or high)
2) When low-quality evidence suggests benefit and high-quality evidence suggests harm or a very high cost
3) When low-quality evidence suggests equivalence of two alternatives, but high-quality evidence of less harm for one of the competing alternatives
4) When high-quality evidence suggests equivalence of two alternatives and low-quality evidence suggests harm in one alternative
5) When high-quality evidence suggests modest benefits and low-/very low-quality evidence suggests possibility of catastrophic harm

Weak recommendations are those for which the desirable effects probably outweigh the undesirable effects (weak recommendation for an intervention) or undesirable effects probably outweigh the desirable effects (weak recommendation against an intervention), but appreciable uncertainty exists. Weak recommendations result when the balance between desirable and undesirable effects is small, the quality of evidence is lower, or there is more variability in the values and preferences of patients. In cases where the balance of cost and benefits is ambiguous, key stakeholders differ about the acceptability or feasibility of implementation, and the effects on health equity are unclear are likely to result in a weak recommendation. A weak recommendation implies that most people would want the recommended course of action, but that many would not. For clinicians, this means that they must recognize that different choices will be appropriate for each individual, and that they must help each person arrive at a management decision consistent with his/her values and preferences. Policy-making will require substantial debate and involvement of various stakeholders.

Evidence is graded as high, moderate, low or very low quality, based on how likely further research is to change our confidence in the estimate of effect.

Box 2: Summary of recommendations for clinicians and policy-makers
We recommend against screening for HCV in adults who are not at elevated risk (strong recommendation, very low-quality evidence).

This recommendation applies only to adults who are not at elevated risk for HCV. It does not apply to pregnant women14 or adults who are at elevated risk for hepatitis C,14 such as:

- Individuals with current or history of injection drug use
- Individuals who have been incarcerated
- Individuals who were born, travelled or resided in HCV-endemic countries (Appendix 6)
- Individuals who have received health care where there is a lack of universal precautions
- Recipients of blood transfusions, blood products or organ transplant before 1992 in Canada
- Patients on hemodialysis
- Individuals who have had needle-stick injuries
- Individuals who have engaged in other risks sometimes associated with HCV exposure, such as high-risk sexual behaviours, homelessness, intranasal and inhalation drug use, tattooing, body piercing or sharing sharp instruments or personal hygiene materials with someone who is HCV positive
- Anyone with clinical clues suspicious for HCV infection (and above risk factors)

general population or in any other higher-risk or higher-prevalence subgroup, including the 1950–1975 birth cohort. Only one uncontrolled retrospective study22 was found that reported on harms. That study22 was a retrospective review of records from a large urban US Department of Veterans Affairs hospital; the study reported that of 12 485 people screened for HCV in 2001, only one patient experienced serious harm (hospitalization for pain control following liver biopsy).

The modelling study commissioned by the task force21 suggested that under the assumptions of the model, one-time screening of 100 000 individuals not at elevated risk of HCV (0.2% prevalence) could result in 199 new diagnoses of chronic HCV infection, compared with 91 persons identified through case finding (testing for HCV in individuals who show signs or symptoms or who are suspected of exposure), over a lifetime horizon (i.e., the model assumes that cases are ascertained across the lifespan of all 100 000 simulated individuals).21 Assuming 89% of identified cases would receive treatment, screening was estimated to prevent 26 cases of decompensated cirrhosis and 20 cases of hepatocellular carcinoma, contributing to about 40 lives saved over a lifetime horizon per 100 000 screened.21 Given the natural history of chronic HCV infection, this model showed that the expected benefit from screening would not be realized for 20–30 years from time of initial infection, with only 3 HCV-related deaths prevented at 5 years and 6 deaths at 10 years24 after screening 100 000 individuals not at elevated risk.

The model21 has several important limitations that contribute to the uncertainty in the estimate of the effect: 1) an inability to consider potential differences in long-term outcomes between initiating treatment at earlier versus later stages of liver fibrosis,
2) the possibility that the baseline risk of disease progression used for the model (based on nonscreened populations) is higher than that for the asymptomatic screened population, which would translate into an overestimation of the benefit of screening, 3) an inability to take into consideration the harms of overdiagnosis and overtreatment that result from screening, and 4) the inherent inability of modelling to account for unknown factors that may influence screening outcomes.

The screening review found low-quality evidence from 26 cross-sectional studies on the clinical validity of HCV screening tests (anti-HCV antibody and HCV-antigen tests) in predicting active infection (measured with a confirmatory HCV RNA test). Anti-HCV antibody tests detect both active and past infection; therefore, individuals with past and resolved infections screen positive although they do not have active or chronic HCV infection. The proportion of confirmed active infections among positive screening test results varies widely across studies and test types. The proportion of RNA-positive, active HCV infection cases ranged from 0% to 89.7% among positive antibody-based assays and from 0% to 100% among antigen-based assays. Only three studies conducted confirmatory polymerase chain reaction on samples that tested negative for HCV antibody or antigen. The proportion of negative anti-HCV assays that were confirmed with polymerase chain reaction to be RNA negative ranged from 73.7% to 99.7% for two antibody assays and 89.7% in one antigen assay. The wide variation in results may be as a result of actual differences in test performance, varying study sample sizes or underlying variation in population characteristics, including population prevalence.

**Treatment**

Indirect evidence on treatment of HCV from the treatment review examined the benefits and harms of pegylated interferon-ribavirin treatment regimens compared with newer DAAs. This evidence is indirect, because the population examined is not a screened population and includes symptomatic individuals, who (by definition) are not the target of a screening program. Moderate-quality evidence from seven RCTs showed that treatment with DAA-based regimens achieved higher sustained virologic response rates than PR at 48 weeks following treatment (SVR48) and reduced the frequency of harms (Appendix 4, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.161521/-/DC1). Very low-quality evidence showed no difference in quality of life (three RCTs, n = 1342) or all-cause mortality from 36 to 72 weeks following treatment (five RCTs, n = 1853) (Appendix 4). No studies reported on hepatic mortality, cirrhosis, hepatocellular carcinoma, need for liver transplant, and improvement in liver histology or reduced HCV transmission. Three RCTs suggested that there may be higher rates of SVR12 when treatment is given at an earlier fibrosis stage (F0 to F2) versus a later stage (F3 to F4). However, this was a post hoc analysis in which no statistical tests were done and that may serve as hypothesis generating only.

The evidence from the treatment review has several key limitations that reduce its applicability to this guideline. 1) Study participants were not identified by screening and could differ from people detected by population-based screening. It is uncertain whether the effects of treatment would be similar for symptomatic and screened, asymptomatic patients. 2) The identified studies did not include a comparison to an untreated control group. 3) Evidence of benefit associated with newer treatments was restricted to surrogate outcomes assessed following a relatively short interval, given the chronic nature of HCV infection, and did not consider the potential risk of occurrence of hepatocellular carcinoma after achieving sustained virologic response.

**Patient values and preferences**

The screening review found moderate- to low-quality evidence from 12 observational studies, which reported on patient preferences and values related to the decision to be screened for HCV. Findings were highly variable in terms of patient preferences related to screening and there was a high level of uncertainty about the value that patients place on the clinical outcomes considered in the review. Concerns related to stigma and to access to care were reported as important factors for decision-making. A survey of 15 patients commissioned by the task force generally reinforced these findings and found that participants placed equal value on the benefits and harms of screening. Reduced mortality was perceived as a very important benefit, and concerns were noted about stigma and psychological adverse events from positive results of screening tests.

**Resource use**

A recent study examined the potential budget impact of HCV screening in Alberta. Assuming a 0.69% HCV prevalence, 70% screening uptake and 24% treatment uptake (reflects uptake common with older treatments, which led to more severe adverse events) and hypothetical negotiated drug costs of about 50% off list price, screening and treating the general population of Alberta was estimated to cost $253 million over one year. This estimate was driven primarily by drug costs. If treatment uptake were to increase from 24% to 75% based on the use of DAA-based regimens with fewer adverse effects, the cost of screening and treatment would increase to $501 million. Extrapolating these findings to the Canadian population not at elevated risk for HCV (n = 19,855,629), and assuming an HCV prevalence of 0.2%, a screening uptake of 70%, treatment uptake of 75%, and a hypothetical negotiated drug cost of about 50% off list price, would yield an estimated cost of more than $844 million for screening only and approximately $1.5 billion to screen and treat with DAA-based regimens. For the Canadian extrapolation, only 75% treatment uptake is used because it is more likely to reflect the situation going forward, when DAA-based regimens will be primarily used.

**Feasibility, acceptability, cost and equity**

The majority of individuals identified by screening are anticipated to be asymptomatic and in early stages of HCV (F0 to F1). At the time of writing this guideline, certain provinces had negotiated a price reduction with pharmaceutical companies that
produce DAA-based treatment regimens. However, more advanced fibrosis (F2 to F4) and presentation with comorbidities is still needed to qualify for treatment in Canada. As a result, it is likely that most individuals who would be identified by screening would not qualify for drug coverage under their provincial plan. Given the high listed cost of DAA-based regimens (based on 2015 list prices in Alberta for a course of treatment: sofosbuvir + pegylated interferon-ribavirin, $104,429; ledipasvir + sofosbuvir $95,000), few people with HCV are likely to have the funds to pay out of pocket for a course of treatment. Therefore, a recommendation in favour of screening would increase the number of people with known HCV (and who are potentially susceptible to harms of stigma and anxiety) who could not access treatment, thus deriving no clear benefit despite the potential for harm from a diagnosis combined with treatment ineligibility. To the extent that wealthier individuals with screen-detected HCV could access treatment, a recommendation for screening has potential to increase health inequity.

Presently, the lack of health system resources required to successfully roll out a treatment strategy that includes all individuals with HCV means that population-based screening is unlikely to be acceptable to funders such as provincial and territorial governments, especially given the uncertain benefits of a screening program. The results of the FACE survey are presented in Appendix 3.

Data from 38 countries show that the price of DAA medication varies substantially between lower- and higher-income countries and even across high-income countries. For example, one bottle of sofosbuvir is reported to cost (in USD) $161 to $312 in India, $500 in Ivory Coast, $14,000 in Spain and $20,590 in Switzerland. These set costs are not correlated with gross national income or with manufacturing costs. The task force encourages policymakers to continue to work with pharmaceutical companies to offer affordable prices for new HCV treatments, which might justify the allocation of the resources required for a successful roll-out of a treatment strategy for all individuals identified with HCV, including those with no comorbidities in the early stages of liver disease (F0 and F1). If pharmaceutical companies were to decrease substantially the cost of new HCV drugs for all people diagnosed with early stages of fibrosis (i.e., F0 and F1) regardless of comorbidity, and policy-makers were to confirm that health system resources are in place to ensure a successful rollout of a treatment strategy, it is possible that under such circumstances, the resources spent on screening individuals not at elevated risk for HCV could be worth the expected net benefit, even if that benefit were small.

Rationale

The screening review did not identify any evidence for the benefit of screening for HCV on critical outcomes. The modelling study estimated possible results for three critical outcomes (hepatic mortality, hepatocellular carcinoma and decompensated cirrhosis), but this model was rated as very low-quality evidence. Given the lack of direct evidence on screening for HCV and the many assumptions required by the model, the task force considers the overall quality of evidence supporting this recommendation to be very low (i.e., highly uncertain). Given the prevalence of HCV in Canadians not at elevated risk for HCV, the lack of direct evidence on the benefits and harms of screening in all groups of the population, and the very low quality of the indirect evidence produced by modelling, substantial uncertainty remains about the effectiveness of screening among adults in Canada.

This recommendation places a relatively lower value on 1) very low-quality indirect evidence suggesting a potentially small benefit from screening, 2) the low risk of household and sexual transmission of HCV among individuals not at elevated risk, as well as the low risk of transmission through blood products given routine screening of blood and organs, and 3) the potential risk of the individual developing end-stage liver disease and transmitting the infection despite being asymptomatic.

On the other hand, this recommendation places a relatively higher value on 1) the anticipated increase in harm resulting from diagnosing and treating individuals who screen positive but would have never developed HCV-related disease during their lifetime; 2) false positives and false negatives, which could lead to unnecessary anxiety and/or false reassurance; 3) the potential for screening to increase inequity, given that among those who do not meet current eligibility criteria (e.g., specific comorbidities), only wealthier individuals or those with private insurance would obtain earlier access to treatment not currently funded by government; 4) the unknown magnitude of benefit of treatment on reducing risk of transmission; and 5) the very large impact that screening and treatment would have on health care budgets, and associated opportunity costs (i.e., the limit this would place on the ability to provide other health care interventions that would have to be forgone for lack of funds, despite being supported by better evidence).

Indirect evidence showing that new DAA-based treatment is much more effective in achieving sustained virologic response than older pegylated interferon-ribavirin treatment regimens was also considered in developing this recommendation. As such, emerging evidence on new treatments with similar effectiveness would not change this recommendation. Our recommendation is also informed by limited evidence on patient preferences: although some patients would want to be screened, many would forgo treatment if it were not funded by provincial and territorial drug plans. The rationale for this recommendation is outlined in the GRADE Evidence to Decision framework (Appendix 5, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.161521/-/DC1).

The recommendation is strong because the task force is confident of the potential for harm resulting from screening and treatment for HCV, and substantial uncertainty remains about the benefits of screening. Considering the totality of evidence, this means that the estimate of beneficial effect could be substantially different from the true effect of screening. The task force is confident that implementing any recommendation to screen and treat those identified as HCV positive would require substantial health system resources. This burden would limit the ability of clinicians to deliver other interventions of proven value.

Further, because of the degree of uncertainty related to any benefit of screening, it would be difficult to define a set of circumstances under which a practitioner should offer screening to an
individual who is not at increased risk for HCV. The strong recommendation implies that the majority of such patients will be better served by not being screened for HCV. Considering that funding for treatment is presently limited to individuals with identified HCV and more advanced liver fibrosis (F2 to F4), the task force believes our recommendation should allow clinicians to focus on providing treatment to individuals who have already been diagnosed.

This guideline may be re-evaluated as evidence on the benefit and/or harm of screening emerges, such as evidence examining the real-life consequences of screening and treating those who would have otherwise never developed complications or died from liver disease; or as other factors influencing the recommendations change, such as improved access to treatment. This could occur, for instance, if there were a substantial reduction in the price of treatment, to an extent that would permit all individuals with HCV to be treated. The task force notes that given the potential impact of treatment on budget, even very large reductions in price might not be sufficient to recommend screening. Even if markedly lower drug prices were available, changes to models of care may also be required before population-based screening could be warranted, such as changes in health system policies to support a successful rollout of a treatment strategy that would include all individuals identified as having chronic HCV infection, regardless of fibrosis stage or comorbidity. Better access to DAA-based treatment may require extending management of HCV to clinicians in primary care.

Considerations for implementation

The task force recommendation applies to individuals who are not pregnant or at elevated risk for HCV. Subgroups of the population who are at increased risk for HCV (and not included in this recommendation) may require special attention from clinicians. A joint 2009 recommendation from CFPC and PHAC,14 although not based on a systematic review of the evidence, addressed those individuals who are at increased risk. That guidance suggests testing for HCV in “anyone with risk behaviours for HCV, with potential exposure to HCV, and/or with clinical clues suspicious for HCV.” Populations targeted in the CFPC/PHAC guideline14 include people who inject drugs (currently or in the past); individuals who have been incarcerated; individuals who may have been exposed to contaminated blood, blood products or medical equipment; and those who have travelled or resided in endemic regions.13,14

Some immigrants are at increased risk for HCV because they are from countries where HCV infection is common. Unlike the nonimmigrant population, these persons are at increased risk for HCV because of iatrogenic exposure in their country of origin (e.g., lack of standard precautions, or as a result of medical or dental procedures with contaminated equipment) and not necessarily from injection drug use or other higher-risk behaviours.83 The CFPC/PHAC guidance14 recommends testing for HCV in individuals who were “born, traveled or resided in a region in which HCV infection is more common.” A list of endemic countries and a related map are provided in Appendix 6 (available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.161521/-/DC1).

More persons in subgroups such as the Indigenous population (3% prevalence)1 and the cohort born from 1950 to 1975 (0.8% prevalence) are diagnosed with chronic HCV;2 these populations have a higher proportion of individuals at higher risk for HCV because of identifiable risk factors.1,2,7,9,24,45 For example, removing from the Indigenous population people who inject drugs would reduce the HCV prevalence from 3% to 0.5%.1,83 Individuals from the Indigenous population who are not otherwise at increased risk are, therefore, included in the present task force guidance, which recommends against screening adults who are not at elevated risk. Similarly, the increased reported prevalence in the cohort born between 1950 and 1975 is likely driven by an increased prevalence of risk behaviours or potential exposures, rather than birth year per se.1,24 In the judgment of the task force, neither Indigenous people nor members of the 1950–1975 birth cohort should be screened for HCV in the absence of other characteristics that would place them at increased risk for HCV.

The task force considered the possibility of screening a birth cohort; that is, one-time testing of all people born, for example, between 1950 and 1975. Most individuals in the birth cohort who are at elevated risk are included in the joint CFPC/PHAC guideline.14 Following this risk-based guideline will likely increase the identification of those who will benefit most from testing. Those born from 1950 to 1975, who are not otherwise at increased risk, are included in the present task force guidance, which recommends against screening adults who are not at elevated risk. More evidence would be needed before making a recommendation about birth cohort testing, separate from adults in the general population.

The task force developed knowledge translation tools to help clinicians assess their patients’ risk for HCV, so that testing can be offered to those at increased risk. These are available at http://canadiantaskforce.ca/tools-resources/hepatitis-c-2/.

Monitoring and evaluation

Given that the task force has recommended against screening adults who are not at elevated risk of HCV, a clear indicator of the uptake of this guideline would be decreased screening of individuals who do not present with risk factors.

Other guidelines

Population-based screening for HCV is not recommended by the task force. The task force recommendation aligns with recently published clinical guidelines from the World Health Organization,46,56 the National Institute for Health and Care Excellence,46 the Scottish Intercollegiate Guidelines Network;45 Immigration, Refugees and Citizenship Canada;40 UK National Screening Committee;91 and the Gastroenterological Society of Australia.83 It partly aligns with guidelines from the Canadian Collaboration for Immigrant and Refugee Health,93 the US Preventive Services Task Force (USPSTF)94 and the US Centers for Disease Control and Prevention (CDC) (1998)95 (Table 1). Although there is variation in definitions, most jurisdictions recommend some sort of risk-based testing.

The more recent guideline from CDC (2012)96 and the USPSTF94 guideline recommend one-time screening for those born between
1945 and 1965. This recommendation relies on indirect evidence such as prevalence (estimated to be 3.25% in the US,\textsuperscript{94} which is four times higher than in Canada at 0.8%\textsuperscript{95}), attainment of sustained virologic response (a surrogate outcome) and the ability of practitioners to influence screening uptake. The CDC and USPSTF recommendations acknowledge the lack of direct evidence on effectiveness of screening in this cohort and the potential for screening to increase overall harms in this population\textsuperscript{97,98} related to overdiagnosis and overtreatment.

**Table 1 (part 1 of 2): National and international guidelines on testing and screening for hepatitis C virus**

<table>
<thead>
<tr>
<th>Organization</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>Canadian Task Force on Preventive Health Care (current guideline, 2017)</td>
<td>We recommend against screening for HCV in adults who are not at elevated risk (strong recommendation, very low-quality evidence). This recommendation applies only to adults who are not at elevated risk for HCV. It does not apply to pregnant women or adults who are at elevated risk for hepatitis C, such as individuals with current or history of injection drug use; individuals who have been incarcerated; individuals who were born, travelled or resided in HCV endemic countries; individuals who have received health care where there is a lack of universal precautions; recipients of blood transfusions, blood products or organ transplant before 1992 in Canada; patients on hemodialysis; individuals who have had needle-stick injuries; individuals who have engaged in other risks sometimes associated with HCV exposure, such as high-risk sexual behaviours, homelessness, intranasal and inhalation drug use, tattooing, body piercing or sharing sharp instruments or personal hygiene materials with someone who is HCV positive; and anyone with clinical clues suspicious for HCV infection.</td>
</tr>
<tr>
<td>US Centers for Disease Control and Prevention (1998)\textsuperscript{96}</td>
<td>The Centers for Disease Control and Prevention recommends routine testing of those at high risk, defined as history of injection drug use, those at risk of health care associated transmission, individuals with HIV, those with a recognized exposure, or those who are concerned. This is an unrated risk-based recommendation, based on expert opinion related to prevalence in risk groups</td>
</tr>
<tr>
<td>College of Family Physicians of Canada and the Public Health Agency of Canada (2009)\textsuperscript{14}</td>
<td>The College of Family Physicians of Canada and Public Health Agency of Canada guideline recommends screening anyone with risk behaviours or potential exposures, defined as injection drug use, incarceration, exposure in a high-prevalence region (born, travelled or resided), health care associated transmission, higher-risk sexual behaviour, tattoos or body piercing, or ceremonial rituals requiring skin piercing done with suspect infection control, etc. Individuals with any of the above risk factors and the following should also be tested: abnormal ALT, HIV, hepatitis B, non-Hodgkin lymphoma, signs of chronic liver, etc. These are unrated risk-based recommendations, based on expert opinion.</td>
</tr>
<tr>
<td>UK National Screening Committee (2011)\textsuperscript{91}</td>
<td>The National Screening Committee recommended against a national screening program for HCV among people of ethnic minorities who are born outside the UK. This is an unrated recommendation based on a lack of randomized controlled trial data on the effectiveness of screening programs to reduce morbidity and mortality in addition to other areas related to program initiation.</td>
</tr>
<tr>
<td>Canadian Collaboration for Immigrant and Refugee Health (2011)\textsuperscript{91}</td>
<td>The Canadian Collaboration for Immigrant and Refugee Health recommends screening all immigrants and refugees from regions with prevalence of disease ≥ 3% (this excludes South Asia, Western Europe, North America, Central America and South America). This prevalence-based recommendation is based on moderate-quality evidence factoring in increased risk of death from viral hepatitis and hepatocellular carcinoma in immigrants and refugees compared with the general population.</td>
</tr>
<tr>
<td>US Centers for Disease Control and Prevention (2012)\textsuperscript{96}</td>
<td>The Centers for Disease Control and Prevention augmented its 1998 guideline by recommending one-time testing without prior ascertainment of HCV risk for adults born between 1945 and 1965. This is a strong cohort-based recommendation based on moderate-quality evidence that these benefits outweigh harms: attainment of SVR, and SVR’s association with reduced risk of hepatocellular carcinoma and all-cause mortality. Ancillary evidence was also considered: undergoing liver biopsy, false-positive tests, anxiety, treatment access and effect of HCV status on insurance and employment. Cohort chosen was based on prevalence estimates.</td>
</tr>
<tr>
<td>Immigration, Refugees and Citizenship Canada (2013)\textsuperscript{95}</td>
<td>Immigration, Refugees and Citizenship Canada recommends one-time screening of those with risk factors such as injection drug use; health care associated transmission, including occupational exposure; higher-risk sexual behaviour; tattoos or body piercing; former incarceration; children born to mothers with chronic hepatitis C infection; those with signs or symptoms of liver disease, active tuberculosis, HIV or syphilis, etc., with particular emphasis on those from endemic countries: Egypt, Pakistan and China are specifically mentioned. These are unrated risk-based recommendations, based on expert opinion and provincial recommendations (British Columbia) and the World Health Organization.</td>
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**Gaps in knowledge**

An important gap is the lack of studies that examine the benefits, harms and other potential consequences of screening asymptomatic populations. RCTs that compare treatment at earlier versus later stages of liver fibrosis are needed. A large population-based prevalence study of chronic HCV in Canada is also lacking. Small studies, primarily in lower- or higher-risk groups and using modelled data, are available, but the confidence in the certainty of those
estimates is low. Although there is some evidence on the natural history of HCV infection, there is uncertainty about the factors that influence the progression of liver disease and how these factors may affect the proportion of people who go on to develop end-stage liver disease, in some cases, despite achieving sustained virologic response. Although there are observational data examining the association of sustained virologic response with long-term outcomes important to patients, additional studies are needed to ascertain whether sustained virologic response associated with newer agents (DAAs) yields similar outcomes.

**Conclusion**

The HCV prevalence in most adults in the general Canadian population is low and direct evidence examining the benefits and harms of screening for HCV is not available. Thus, the task force recommends against screening adults who are not at elevated risk for HCV. Not screening for HCV will focus our limited health care resources to test (and treat) individuals at elevated risk for HCV and to provide other medical interventions that are proven to be of benefit.


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