Evidence-based clinical guidelines for immigrants and refugees

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Conditions covered in systematic reviews

Infectious diseases
- Measles, mumps, rubella
- Diphtheria, tetanus, polio, pertussis
- Varicella
- Hepatitis B
- Tuberculosis
- HIV
- Hepatitis C
- Intestinal parasites (Strongyloides and Schistosoma)
- Malaria

Chronic and noncommunicable diseases
- Diabetes mellitus
- Iron-deficiency anemia
- Dental disease
- Vision health

Women’s health
- Contraception
- Cervical cancer
- Pregnancy

Mental health and maltreatment
- Depression
- Post-traumatic stress disorder
- Child maltreatment
- Intimate partner violence

Key points
- Clinical preventive care should be informed by the person’s region or country of origin and migration history (e.g., forced versus voluntary migration).
- Forced migration, low income and limited proficiency in English or French increase the risk of a decline in health and should be considered in the assessment and delivery of preventive care.
- Vaccination (against measles, mumps, rubella, diphtheria, tetanus, pertussis, polio, varicella, hepatitis B and human papillomavirus) and screening (for hepatitis B, tuberculosis, HIV, hepatitis C, intestinal parasites, iron deficiency, dental pain, loss of vision and cervical cancer) should be routinely provided to at-risk immigrants.
- Detecting and addressing malaria, depression, post-traumatic stress disorder, child maltreatment, intimate partner violence, diabetes mellitus and unmet contraceptive needs should be individualized to improve detection, adherence and treatment outcomes.
1. Overview: evidence-based clinical guidelines for immigrants and refugees

There are more than 200 million international migrants worldwide, and this movement of people has implications for individual and population health. The 2009 United Nations Human Development Report suggested that migration benefits people who move, through increased economic and education opportunities, but migrants frequently face barriers to local health and social services. In Canada, international migrants are a growing and economically important segment of the population (Table 1A).

Immigrants to Canada are a heterogeneous group. Upon arrival, new immigrants are healthier than the Canadian-born population, both because of immigrant-selection processes and policies and because of sociocultural aspects of diet and health behaviours. However, there is a decline in this “healthy immigrant effect” after arrival. In addition, compared with the Canadian-born population, subgroups of immigrants are at increased risk of disease-specific mortality; for example, Southeast Asians from stroke (odds ratio [OR] 1.46, 95% confidence interval [CI] 1.00–1.91), Caribbeans from diabetes mellitus (OR 1.67, 95% CI 1.03–2.32) and infectious diseases (e.g., for AIDS, OR 4.23, 95% CI 2.72–5.74), and immigrant men from liver cancer (OR 4.89, 95% CI 3.29–6.49).

The health needs of newly arriving immigrants and refugees often differ from those of Canadian-born men, women and children. The prevalence of diseases differs with exposure to disease, migration trajectories, living conditions and genetic predispositions. Language and cultural differences, along with lack of familiarity with preventive care and fear and distrust of a new health care system, can impair access to appropriate health care services. Additionally, patients may present with conditions or concerns that are unfamiliar to practitioners.

Many source countries have limited resources and differing health care systems, and these differences may also contribute to health inequalities among migrants. In these guidelines, we refer to low- and middle-income countries as “developing.”

Why are clinical guidelines for immigrants needed?

Canadian immigration legislation requires that all permanent residents, including refugees, refugee claimants and some temporary residents, undergo an immigration medical examination. Screening is undertaken to assess the potential burden of illness and a limited number of public health risks. The examination is not designed to provide clinical preventive screening, as is routinely performed in Canadian primary care practice, and it is linked to ongoing surveillance or clinical actions only for tuberculosis, syphilis and HIV infection.

The Canadian Task Force on Preventive Health Care and the US Preventive Services Task Force have produced many high-quality clinical prevention recommendations, but these statements have not explicitly considered the unique preventive needs and implementation issues for special populations such as immigrants and refugees. Evidence-based recommendations can improve uptake and health outcomes related to preventive services, even more so when they are tailored for specific populations.

How are these guidelines different?

Use of evidence-based methods has yet to substantially affect the field of migration medicine. The Canadian Collaboration for Immigrant and Refugee Health explicitly aims to improve patients’ health using an evidence-based clinical preventive approach to complement existing public health approaches. In selecting topics, primary care practitioners considered not just the burden of illness but also health inequities and gaps in current knowledge. Public health concerns and predeparture migrant screening and treatment protocols were also consid-

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Table 1A: Classification of international migration to Canada (2007)*

<table>
<thead>
<tr>
<th>Immigration category</th>
<th>Annual migration (no.)†‡</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Permanent residents</strong>†</td>
<td></td>
</tr>
<tr>
<td>Economic class (business and economic migrants)</td>
<td>131 000</td>
</tr>
<tr>
<td>Family class (family reunification)</td>
<td>66 000</td>
</tr>
<tr>
<td>Humanitarian class (refugees resettled from abroad or selected in Canada from refugee claimants)</td>
<td>28 000</td>
</tr>
<tr>
<td>Others</td>
<td>11 000</td>
</tr>
<tr>
<td>Total</td>
<td>237 000</td>
</tr>
<tr>
<td><strong>Temporary residents</strong>‡</td>
<td></td>
</tr>
<tr>
<td>Migrant workers</td>
<td>165 000</td>
</tr>
<tr>
<td>International students</td>
<td>74 000</td>
</tr>
<tr>
<td>Refugee claimants (those arriving in Canada and claiming to be refugees)§</td>
<td>28 000</td>
</tr>
<tr>
<td>Other temporary residents§</td>
<td>89 000</td>
</tr>
<tr>
<td>Total§</td>
<td>357 000</td>
</tr>
<tr>
<td><strong>Other migrants</strong></td>
<td></td>
</tr>
<tr>
<td>Total irregular migrants,§ not annual migration</td>
<td>~ 200 000</td>
</tr>
<tr>
<td>Visitors§</td>
<td>~ 30 100 000</td>
</tr>
</tbody>
</table>

*Reproduced, with permission, from Gushulak et al.
†Numbers rounded to nearest 1000. Total in each category may not match sum of values reported because of rounding.
‡Unless otherwise indicated.
§No official migration status; this population includes those who have entered Canada as visitors or temporary residents and remained to live or work without official status. It also includes those who may have entered the country illegally and not registered with authorities or applied for residence.
tered, but these were not the driving force for the recommendations. We implemented evidence-based methods, which included searches for evidence on immigrant preferences and values, as well as incorporating the GRADE approach (Grading of Recommendations Assessment, Development and Evaluation), to formulate clinical preventive recommendations.20–22 Our evidence reviews synthesized data from around the world, and our recommendations focus on immigrants, refugees and refugee claimants, with special attention given to refugees, women and the challenges of integrating recommendations into primary care. Migrants living without official status are particularly vulnerable, but specific evidence for this population is limited.19 In these guidelines, the “health settlement period” refers to the first five years of residence in Canada for an immigrant or refugee, the time during which loss of the healthy immigrant effect begins to surface.

In recent years, there has been an increase in development of practice guidelines for international migrants.20 Notable publications have included Cultural Competency in Health;21 Immigrant Medicine22 and guidelines for refugees from the Australasian Society for Infectious Diseases.21 Many have been designed to address diseases and conditions of public health importance,22–23 and some highlight the importance of psychosocial problems and mental illness, issues of women’s health and chronic noninfectious diseases.21,26,27 Other practice guidelines include strategies to improve communication (e.g., interpreters), responsiveness to sociocultural background (e.g., cultural competence), empowerment (e.g., health literacy), monitoring (e.g., health and access disparities) and strategies for comprehensive care delivery.21

Our recommendations differ from other guidelines because of our insistence on finding evidence for clear benefits before recommending routine interventions. For example, in our guidelines for post-traumatic stress disorder, intimate partner violence and social isolation in pregnancy, we recommend not conducting routine screening, but rather remaining alert. With regard to screening for asymptomatic intestinal parasites, we recommend focusing on serologic testing for high burden of disease parasites, rather than traditional testing of stool for ova and parasites.

How were these guidelines developed?

We followed the internationally recognized Appraisal of Guidelines for Research and Evaluation (AGREE; www.agreetrust.org). We selected guideline topics using a literature review, stakeholder engagement and the Delphi process with equity-oriented criteria.20 In May 2007, we held a consensus meeting of experts in immigrant and refugee health to develop a systematic process for transparent, reproducible, evidence-based reviews. The guideline committee selected review leaders from across Canada on the basis of their clinical and evaluation expertise (see Appendix 1, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

The 14-step evidence review process (Box 1A)20 used validated tools to appraise the quality of existing systematic reviews, guidelines, randomized trials and other study designs. We searched MEDLINE, Embase, the Cochrane Library and other sources for admissible evidence, specifically reviews and related studies, from 1996 to 2010. We identified guidelines developed by other groups but based our recommendations on evidence from primary studies. We identified patient-important outcomes and used the GRADE approach to assess the magnitude of effect on benefits and harms and on quality of evidence. We included both direct evidence from immigrant and refugee populations and indirect evidence from other populations. We downgraded the quality of evidence for indirectness when there was concern that the evidence might not be applicable to immigrant and refugee populations (e.g., because of differences in baseline risk, morbidity and mortality, genetic and cultural factors, and compliance variations). We assessed whether benefits outweighed harms, the quality of evidence, and values and preferences to minimize the potentially negative effects of labelling on patients, families and communities (Table 1B).16–18

Each of the resulting evidence reviews for priority conditions of the Canadian Collaboration for Immigrant and Refugee Health provides detailed methods and results concerning the burden of illness for the immigrant populations relative to Canadian-born populations, along with information about effectiveness of screening and interventions, a discussion of clinical considerations, the basis for recommendations and gaps in research.

How should I begin to assess immigrants for clinical preventive care?

Determine each person’s age, sex, country of origin and migra-

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**Box 1A: Fourteen-step process for evidence reviews used by the Canadian Collaboration for Immigrant and Refugee Health**

1. Develop clinician summary table
2. Develop logic model and key questions
3. Set the stage for admissible evidence (using search strategy)
4. Assess eligibility of systematic reviews
5. Search for data specific to immigrant and refugee populations
6. Refocus on key clinical preventive actions and key questions
7. Assess quality of systematic reviews
8. Search for evidence to update selected systematic reviews
9. Assess eligibility of new studies
10. Integrate data from updated search
11. Synthesize final evidence bank and draft two key clinical actions
12. Develop table for summary of findings
13. Identify gaps in evidence and needs for future research
14. Develop clinical preventive recommendations using GRADE (Grading of Recommendations Assessment, Development and Evaluation)

*Adapted, with permission, from Tugwell and others.16
tion for socially disadvantaged populations, sequencing of care using checklists or algorithms can improve both the uptake and the delivery of preventive care and allows other members of the primary health care team to participate in the delivery of care. Working with interpreters, cultural brokers, patients’ families and community support networks can support culturally appropriate care. Most importantly, clinicians should recognize that the implementation of recommendations (vaccinations, for example) may take three or four visits, a process more akin to the delivery of well-baby care than to an annual examination. Our recommendations are aimed at primary care practitioners, but competencies related to immigrant and cross-cultural care will vary depending on training and experience, and expert support should be sought accordingly.

Which immigrant populations face the most significant health risks?

Refugees, who are by definition forcefully displaced, are at highest risk for past exposure to harmful living conditions, violence and trauma. Refugees undergo medical screening before admission to Canada but are protected by law from exclusion on the basis of noninfectious burden of illness (through the Immigration and Refugee Protection Act). The health risks of refugees and other migrants vary greatly depending on exposures (e.g., to vectors of disease such as mosquitos), trauma from war, living conditions (e.g., access to water and sanitation), neglect from long periods in refugee camps, susceptibilities (e.g., related to ethnicity and migration stress), social stratification (e.g., race, sex, income, education and occupation) and access to preventive services (e.g., pre-departure access to primary care, vaccinations and screening, access to Canadian services and access issues related to linguistic and cultural barriers).

Specifically, refugees are at risk for a rapid decline in self-reported health after arrival (OR 2.31, 95% CI 1.1–4.9), as are low-income immigrants (OR 1.5, 95% CI 1.3–1.7) and immigrants with limited English- or French-language proficiency. There is also an increased risk of reporting poor health among immigrants with limited English- or French-language proficiency (OR 2.0, 95% CI 1.5–2.7), those facing cost-related barriers to health care (OR 2.8, 95% CI 1.7–4.5), low-income immigrants and non-European immigrants (OR 2.3, 95% CI 1.6–3.3).

Clinical recommendations

Considering the burden of illness of immigrant populations, the quality of evidence for screening and interventions, and the feasibility of clinicians implementing the recommendations, we have organized our recommendations into four groups: infectious diseases, mental health and physical and emotional maltreatment, chronic and noncommunicable diseases, and women’s health.


Infectious diseases

Many immigrants are susceptible to vaccine-preventable diseases upon arrival in Canada. For example, 30%–50% of new immigrants are susceptible to tetanus, 32%–54% are susceptible to either measles, mumps or rubella, and immigrants from tropical countries are 5–10 times more susceptible to varicella, which has serious implications for adult immigrants.

A large proportion (20%–80%) of the immigrants who come from countries where chronic hepatitis B virus infection is prevalent are not immune. In addition, immigrants are more likely to be exposed to hepatitis B virus in their households and during travel to countries where hepatitis B is prevalent. Immigrants from countries where chronic hepatitis B virus infection is prevalent (affecting 2% or more of the population) can benefit from screening and treatment to prevent hepatitis and hepatocellular carcinoma.

Foreign-born people account for 65% of all active tuberculosis in Canada, and screening and treatment for latent tuberculosis remain priorities for immigrants from countries in sub-Saharan Africa, Asia, and Central and South America. To promote patients’ safety and adherence to therapy, patients must be informed of the risks and benefits of treatment in a culturally and linguistically appropriate manner. Refugees may already be aware of their HIV-positive status but may have limited knowledge of effective screening and treatment options. HIV-related stigma and discrimination put immigrants and refugees at risk for delayed diagnosis and unequal treatment rates for HIV infection. Immigrants are an unrecognized risk group for chronic hepatitis C virus infection and would benefit from early detection and appropriately timed treatment.

Subclinical strongyloidiasis and schistosomiasis can persist for decades after immigration and, if left untreated, can lead to serious morbidity or death through disseminated disease.

<table>
<thead>
<tr>
<th>Table 1B: Basis of recommendations*</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Issue</strong></td>
</tr>
<tr>
<td>Balance between desirable and undesirable effects</td>
</tr>
<tr>
<td>Quality of evidence</td>
</tr>
<tr>
<td>Values and preferences</td>
</tr>
</tbody>
</table>

*Reproduced, with permission, from Tugwell et al. Based on GRADE (Grading of Recommendations Assessment, Development and Evaluation).
Serologic tests for these intestinal parasites, rather than traditional stool testing, are recommended. Malaria is one of the leading causes of death worldwide, and delay in diagnosis and treatment of *Plasmodium falciparum* may lead to severe disease and even death. Migrant children are especially at risk for malaria and its complications.

Recommendations for infectious diseases are summarized in Box 1B.

**Mental health and physical and emotional maltreatment**

The mental health of immigrants has emerged as one of the most challenging areas for clinicians. Among refugees, depression commonly co-occurs with post-traumatic stress disorder and other anxiety disorders, which can complicate its detection and treatment. Conducting a systematic clinical assessment, or using a validated questionnaire in a language in which the patient is fluent, is recommended if the clinician practises in an integrated system that links patients with suspected depression to treatment programs with a stepped-care approach. Effective detection and treatment may also require the use of professional interpreters or trained culture brokers (not children or other family members) to identify patients’ concerns, explain illness beliefs, monitor progress, ensure adherence, and address the social causes and the consequences of depression. The majority of those who experience traumatic events will heal spontaneously after reaching safety. Empathy, reassurance and advocacy are key clinical elements of the recovery process. Pushing for disclosure of traumatic events could cause more harm than good.

The children of ethnic minorities, including some recently settled immigrants and/or refugees, are disproportionately over-screened (up to 8.75 times more likely) and over-reported as positive (up to four times more likely) for child maltreatment. False-positive reports could result in harm, leading to psychological distress, inappropriate family separation, impaired clinician–patient rapport and legal ramifications associated with the involvement of child protection services. Routine screening is not recommended; rather, clinicians should remain alert for maltreatment, either intimate partner violence or child maltreatment.

Recommendations related to mental health and maltreatment, both physical and emotional, are summarized in Box 1C.

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**Box 1B: Summary of evidence-based recommendations for infectious diseases**

<table>
<thead>
<tr>
<th>Measles, mumps and rubella</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaccinate all adult immigrants without immunization records using one dose of measles–mumps–rubella vaccine.</td>
</tr>
<tr>
<td>Vaccinate all immigrant children with missing or uncertain vaccination records using age-appropriate vaccination for measles, mumps and rubella.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Diphtheria, pertussis, tetanus and polio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaccinate all adult immigrants without immunization records using a primary series of tetanus, diphtheria and inactivated pertussis vaccine.</td>
</tr>
<tr>
<td>Vaccinate all immigrant children with missing or uncertain vaccination records using age-appropriate vaccination for diphtheria, pertussis, tetanus and polio.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Varicella</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaccinate all immigrant children &lt; 13 years of age with varicella vaccine without prior serologic testing.</td>
</tr>
<tr>
<td>Screen all immigrants and refugees from tropical countries ≥ 13 years of age for serum varicella antibodies, and vaccinate those found to be susceptible.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Hepatitis B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Screen adults and children from countries where the seroprevalence of chronic hepatitis B virus infection is moderate or high (i.e., ≥ 2% positive for hepatitis B surface antigen), such as Africa, Asia and Eastern Europe, for hepatitis B surface antigen, anti-hepatitis B core antibody and anti-hepatitis B surface antibody.</td>
</tr>
<tr>
<td>Refer to a specialist if positive for hepatitis B surface antigen (chronic infection).</td>
</tr>
<tr>
<td>Vaccinate those who are susceptible (negative for all three markers).</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Tuberculosis</th>
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</thead>
<tbody>
<tr>
<td>Screen children, adolescents &lt; 20 years of age and refugees between 20 and 50 years of age from countries with a high incidence of tuberculosis as soon as possible after their arrival in Canada with a tuberculin skin test.</td>
</tr>
<tr>
<td>If test results are positive, rule out active tuberculosis and then treat latent tuberculosis infection.</td>
</tr>
<tr>
<td>Carefully monitor for hepatotoxicity when isoniazid is used.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>HIV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Screen for HIV, with informed consent, all adolescents and adults from countries where HIV prevalence is greater than 1% (sub-Saharan Africa, parts of the Caribbean and Thailand).</td>
</tr>
<tr>
<td>Link HIV-positive individuals to HIV treatment programs and post-test counselling.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Hepatitis C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Screen for antibody to hepatitis C virus in all immigrants and refugees from regions with prevalence of disease ≥ 3% (this excludes South Asia, Western Europe, North America, Central America and South America).</td>
</tr>
<tr>
<td>Refer to a hepatologist if test result is positive.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Intestinal parasites</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strongyloides: Screen refugees newly arriving from Southeast Asia and Africa with serologic tests for <em>Strongyloides</em>, and treat, if positive, with ivermectin.</td>
</tr>
<tr>
<td>Schistosoma: Screen refugees newly arriving from Africa with serologic tests for <em>Schistosoma</em>, and treat, if positive, with praziquantel.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Malaria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do not conduct routine screening for malaria.</td>
</tr>
<tr>
<td>Be alert for symptomatic malaria in migrants who have lived or travelled in malaria-endemic regions within the previous three months (suspect malaria if fever is present or person migrated from sub-Saharan Africa). Perform rapid diagnostic testing and thick and thin malaria smears.</td>
</tr>
</tbody>
</table>

*Order of listing considers clinical feasibility and quality of evidence.*
Chronic and noncommunicable diseases
People of certain ethnic backgrounds (specifically Latin Americans, Africans and South Asians) face a twofold to four-fold higher prevalence of type 2 diabetes mellitus than white people, with earlier onset and poorer outcomes. People with hypertension have the most to gain from treatment of obesity, high cholesterol, hypertension and hyperglycemia. Culturally appropriate diabetes education and lifestyle interventions are effective in preventing the disease or improving disease management. Iron deficiency is the most common nutritional deficiency in the world, and immigrant women and children can benefit from screening and supplementation.

Dental disease is often challenging for medical practitioners, but screening and treating pain with nonsteroidal anti-inflammatory drugs can lead to better outcomes and more effective referrals for oral health care. In addition, there is value in recommending twice-daily tooth-brushing with fluoridated toothpaste, as some immigrants may not be familiar with this approach to oral health. Loss of vision is the final common pathway for all eye diseases, and all immigrants can benefit from having their visual acuity assessed soon after arrival in Canada.

Recommendations for chronic and noncommunicable diseases are summarized in Box 1D.

Women’s health
To prevent unintended pregnancy, screening for unmet contraceptive needs should begin soon after a woman’s arrival in Canada. Giving women their contraceptive method of choice (the intrauterine device being the most common contraceptive worldwide, although personal preferences vary), providing the contraceptive method on site and having a good interpersonal relationship all improve contraceptive-related outcomes.

School vaccination programs vary by province, and immigrant girls and women may miss school programs for human papillomavirus vaccination, depending on their age at the time of arrival. Subgroups of immigrants, most notably South Asian and Southeast Asian women, have substantially lower rates of cervical cytology screening than Canadian-born women. Women who have never undergone cervical screening and those who have not had cervical screening in the previous five years account for 60%–90% of invasive cervical cancers. Providing information to patients, building rapport...

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**Box 1C: Summary of evidence-based recommendations for mental health and physical and emotional maltreatment**

**Depression**
If an integrated treatment program is available, screen adults for depression using a systematic clinical inquiry or validated patient health questionnaire (PHQ-9 or equivalent).
Individuals with major depression may present with somatic symptoms (pain, fatigue or other nonspecific symptoms).
Link suspected cases of depression with an integrated treatment program and case management or mental health care.

**Post-traumatic stress disorder**
Do not conduct routine screening for exposure to traumatic events, because pushing for disclosure of traumatic events in well-functioning individuals may result in more harm than good.
Be alert for signs and symptoms of post-traumatic stress disorder (unexplained somatic symptoms, sleep disorders or mental health disorders such as depression or panic disorder).

**Child maltreatment**
Do not conduct routine screening for child maltreatment.
Be alert for signs and symptoms of child maltreatment during physical and mental examinations, and assess further when reasonable doubt exists or after patient disclosure.
A home visitation program encompassing the first two years of life should be offered to immigrant and refugee mothers living in high-risk conditions, including teenage motherhood, single parent status, social isolation, low socioeconomic status, or living with mental health or drug abuse problems.

**Intimate partner violence**
Do not conduct routine screening for intimate partner violence.
Be alert for potential signs and symptoms related to intimate partner violence, and assess further when reasonable doubt exists or after patient disclosure.

Note: PHQ-9 = nine-item Patient Health Questionnaire.
*Order of listing considers clinical feasibility and quality of evidence.

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**Box 1D: Summary of evidence-based recommendations for chronic and noncommunicable diseases**

**Type 2 diabetes mellitus**
Screen immigrants and refugees > 35 years of age from ethnic groups at high risk for type 2 diabetes (those from South Asia, Latin America and Africa) with fasting blood glucose.

**Iron-deficiency anemia**
*Women*
Screen immigrant and refugee women of reproductive age for iron-deficiency anemia (with hemoglobin).
If anemia is present, investigate and recommend iron supplementation if appropriate.

*Children*
Screen immigrant and refugee children aged one to four years for iron-deficiency anemia (with hemoglobin).
If anemia is present, investigate and recommend iron supplementation if appropriate.

**Dental disease**
Screen all immigrants for dental pain. Treat pain with nonsteroidal anti-inflammatory drugs and refer patients to a dentist.
Screen all immigrant children and adults for obvious dental caries and oral disease, and refer to a dentist or oral health specialist if necessary.

**Vision health**
Perform age-appropriate screening for visual impairment.
If presenting vision < 6/12 (with habitual correction in place), refer patients to an optometrist or ophthalmologist for comprehensive ophthalmic evaluation.

*Order of listing considers clinical feasibility and quality of evidence.*
and offering access to female practitioners can improve acceptance of Papanicolaou (Pap) testing.66

Finally, newly-arrived pregnant women are at increased risk for maternal morbidity.67 We identified social isolation, risks of unprotected or unregulated work environments, and sexual abuse (specifically in forced migrants) as priority areas for research.

Recommendations related to women’s health are summarized in Box 1E.

Knowledge translation

We developed a summary of our recommendations and have engaged multiple stakeholders as partners to share these recommendations with their constituencies, including the Public Health Agency of Canada, Citizenship and Immigration Canada, regional health and public health authorities, immigrant community groups and primary care practitioners. These recommendations and their related evidence reviews are available on the CMAJ website (see www.cmaj.ca/lookup/suppl /doi:10.1503/cmaj.090313/-/DC1). Forty primary care practitioners from across Canada with experience working with immigrants pilot-tested the recommendations, provided feedback on the presentation format and are helping to promote the guidelines through their networks. Finally, we sought feedback on our recommendations from our immigrant community partners (specifically, the Edmonton Multicultural Health Brokers Cooperative, which represents 16 ethnic communities) and continue to work with our community partners to improve access to health services.

Directions for future research

Immigrant populations are a heterogeneous group. Because of the selection processes that are in place, most immigrants arrive in good health, although some subgroups are at increased risk of chronic and infectious diseases and mental illness. More research is needed on strategies to address barriers to health services, most urgently for refugees, women and other immigrants with low income and language barriers. There is also a need to develop and study interventions for social isolation and intimate partner violence for pregnant immigrants and refugees. Data remain limited for immigrant children, refugee claimants and nonstatus persons and for many disease areas, including malaria morbidity, post-traumatic stress disorder and interventions for intimate partner violence.

More work must be done to improve immigrants’ access to health services. We hope this evidence-based initiative will provide a foundation for improved preventive health care for immigrant populations.


Podcasts for practitioners and additional information for patients can be found at www.ccirhken.ca.
2. Selection of potentially preventable and treatable conditions

Community-based primary health care practitioners see most of the immigrants and refugees who arrive in Canada. This is not only because Canada’s health care system centres on primary care practice, but also because people with lower socioeconomic status, language barriers and less familiarity with the system are much less likely to receive specialist care.64

Guideline development can be costly in terms of time, resources and expertise.69 Setting priorities is critical, particularly when dealing with complex situations and limited resources.70 There is no standard algorithm on who should determine top priorities for guidelines or how this should be done, although burden of illness, feasibility and economic considerations are all important.71 Stakeholder engagement, to ensure relevance and acceptability, and the use of an explicit procedure for developing recommendations are critical in guideline development.72–74 We chose primary care practitioners, particularly those who care for immigrants and refugees, to help the guideline committee in selecting conditions for clinical preventive guidelines for immigrants and refugees, with a focus on the first five years of settlement. A more detailed description of this Delphi process was published previously.15

Methods

We used a modified Delphi consensus process to select 20 high-priority conditions for guideline development.70,75,76 To begin, we identified key health conditions using an environmental scan, literature review and input from key informants from the Canadian Initiative to Optimize Preventive Care for Immigrants national network, a nascent network of immigrant health providers. This initial step identified 31 conditions. During the ranking process, survey participants were invited to list additional conditions. These conditions, if associated with potentially effective clinical preventive actions, were integrated into the pool of conditions for subsequent ranking.

We developed priority-setting criteria that emphasized inequities in health, building on a process developed for primary care guidelines affecting disabled adults.70,77 Importance or burden of illness is often used for setting priorities, usefulness or effectiveness is frequently used, and disparity is now a well-recognized component of many public health measures.78 We defined our criteria as importance, usefulness and disparity:

- Importance: Conditions that are the most prevalent health issues for newly arriving immigrants and refugees; conditions with a high burden of illness (e.g., morbidity and mortality).
- Usefulness: Conditions for which guidelines could be practically implemented and evaluated. Such guidelines refer to health problems that are easy to detect, for which the means of prevention and care are readily available and feasible, and for which health outcomes can be monitored.
- Disparity: Conditions that might not be currently addressed or that are poorly addressed by public health initiatives or illness-prevention measures that target the general population.

We (H.S., K.P., M.R., L.N.) purposively selected 45 primary care practitioners, including family physicians and nurse practitioners, recently or currently working in a setting serving recent immigrants and refugees. We sampled clinical settings from 14 urban centres across Canada to ensure in-depth experience with a variety of migrants. The settings also covered a range of health service funding models: community health centres (centres locaux de services communautaires in Quebec), refugee clinics, group and solo practices, and ethnic community practices. We aimed to select practitioners with substantial experience, academic expertise or local leadership roles who were willing to commit to offering future input into guideline development and dissemination.

Immigrant and refugee health is a new subdiscipline. The skills, knowledge and experience that define expertise have not yet been determined, and there are no examinations, certification or developed courses that can be used as a proxy for expertise. We believed that contextual knowledge, experience arising from engaged care of immigrants and refugees in Canada, and related work experience in international health were important factors in determining expertise. As a measure of expertise, we adapted a formula used by Médecins Sans Frontières. This criterion combines work with Médecins Sans Frontières in developed countries and in the field. Our criterion for experience was set at seven years or more and included all work in developing countries. It was calculated as number of years of experience with migrants in Canada + (2 × years of experience working in developing countries).

As prompts for decision-making, we asked our practitioner panel to make choices based on the defined criteria, imagining that the guidelines under development might be used at a clinic serving new immigrants or by physicians who do not often see immigrant and refugee patients. Just as clinical practice does, these criteria challenged practitioners to make choices based on competing demands.

This first round of the Delphi survey aimed to ensure that we had the appropriate health conditions under consideration and to begin developing some consensus as to priorities. Participants were asked to rank the 31 conditions identified initially and to propose conditions that were not on the initial list. We chose an a priori cut-off of 80% consensus for inclusion in the top 20. In the second round, we presented an unranked, modified version of this list, excluding all conditions that had already reached 80% consensus and adding newly proposed conditions. The remaining conditions to be included in the top 20 were determined by overall ranking in the second round. This list was reviewed by the codirectors of the Edmonton Multicultural Health Brokers Co-operative (www.mchb.org/OldWebsite2008/default.htm), a group representing over 16 ethnic communities that had initially requested preventive health guidance relevant for immigrant
communities. In addition, the panel of experts who would be developing the guidelines reviewed the list. Then, during the final round, we requested approval, through a simple agree/disagree vote, of the process and the resulting list of priorities, with one-on-one interviews to resolve concerns in the two months following the ranking process.

Consent to participate in the Delphi survey was determined by completion of a questionnaire. Demographic questions elicited personal, professional and practice characteristics of the study participants. With each round, we sent to participants (by email) an explanation of the process to date, the priority-setting criteria, instructions for filling out the survey and a link to the SurveyPro survey. Telephone follow-up was used to maximize response rate. We used Microsoft Excel for the analysis.

Results

Ninety per cent (40/45) of the selected practitioners agreed to participate. Four of the five participants who chose not to participate cited reasons of leave of absence or sabbatical leave, and the fifth cited workload. Ninety-five per cent of the consenting participants completed the first round of the survey, and 88% completed the second and third rounds (Figure 2A). The first two rounds of the Delphi consensus process took place between Mar. 5 and May 31, 2007.

The 40 participants consisted of 35 physicians and five nurse practitioners or nurses with expanded roles. Participants were predominantly women and had been in practice for an average of 14 years. They worked an average of 16 hours per week with immigrants and refugees. More than 80% spoke two or more languages (Table 2A).

The average length of experience working with refugees and immigrants in Canada was 7.5 years; 64% of participants had some experience working in developing countries, with a median overseas duration of 16 (range 1–120) months. Thirty-one per cent of primary care practitioners self-identified as being an immigrant or refugee; of the remainder, 38% self-identified as being the child of an immigrant or refugee (of the 35 practitioners who responded to this optional question).

Forty-five per cent of participants identified themselves as having had prior training in the field, which included accredited tropical medicine courses, designated rotations during residency, work exposures before becoming a health care practitioner, and conferences or self-directed studies in multicultural or cross-cultural medicine.

The refugees and immigrants with whom most practitioners interacted came from all parts of the world; using an average of straight ranking (1 to 6) of regions, south and central Africa was estimated as the most frequent source region of immigrants for these practitioners. Children formed, on average, 30% of clientele, and women, 41%. Seventy-one per cent of migrants were estimated to have been in Canada less than five years, and 73% were involuntary migrants. Involuntary migrants included refugee claimants, so-called Convention Refugees and internally displaced persons (although this is not really an issue for Canada).

Box 2A lists the top 20 conditions for which practitioners identified a current need for guidelines on the basis of our criteria. In the first round, 80% consensus was reached to include 11 conditions. Eighty per cent consensus was also reached to exclude three conditions from the process: Chagas disease, colon cancer and prostate cancer. Three well-defined and unique conditions were proposed for the second round of
ranking: osteoporosis, contraception and vision screening. The nine conditions selected in the second round were based on average ranking (Box 2A).

The list of top 20 conditions was reviewed and approved, with one modification, by the panel of key experts who would be developing the guidelines: routine vaccine-preventable diseases were considered a single priority, with tetanus, diphtheria and polio combined with measles, mumps and rubella for the purposes of guideline development. As a final step, we sent the 20 identified conditions to survey participants for approval and discussion; all 35 people who participated in this round approved (i.e., 88% of the 40 original participants).

**Discussion**

Refugees and many immigrants may have poor or deteriorating health, because of conditions experienced before, during or after arrival to Canada. A health care system that is poorly adapted to their needs compounds this situation, resulting in further marginalization. Our Delphi consensus process used practitioners’ years of field experience strategically to identify preventable and often unrecognized clinical care gaps that can result from such majority-system biases.

An overarching goal of our guideline development project was to supplement guidelines that exist for the general Canadian population by focusing on health inequities. We therefore selected a high proportion of practitioners who work with refugees, a particularly vulnerable subgroup of immigrants prone to disparities. Using practitioners to select conditions ensured both that the needs of the future guideline-users were given priority and that conditions presenting serious clinical challenges, but that might be under-represented in the literature, were included. In working with perceived needs of practitioners, we risked a reporting bias: overemphasizing popular stereotypes (e.g., the importance of infectious diseases), underemphasizing unrecognized or emerging conditions (e.g., vitamin D deficiency) and loss of precision in terms of specific populations (e.g., our list does not fully reflect the greatest needs of children).

Also, by deliberately selecting participants who work with refugees, we risked falsely stereotyping the health status of all immigrants by overemphasizing refugee-specific conditions and, conversely, underemphasizing common health risks, such as hypertension, that affect all immigrants.

The Delphi process generated 20 conditions for guideline development that reflected the needs and priorities of primary care practitioners working with immigrants and refugees. Although immigrant screening has historically focused on infectious diseases, the conditions selected by survey participants extended across a spectrum of diseases, including infectious disease, dentistry, nutrition, chronic disease, maternal

| Table 2A: Demographic characteristics of 40 participants in Delphi consensus process* |
|-------------------------------------|-------------------------------|
| Characteristic                      | No. (%) of participants†      |
| Sex, female (n = 40)                | 25 (62)                       |
| Age, yr, mean                       | 42.5                          |
| Length of practice, yr, mean        | 14.0                          |
| Province of practice (n = 40)       |                               |
| British Columbia                    | 7 (18)                        |
| Prairies (Alberta, Saskatchewan, Manitoba) | 4 (10)                      |
| Ontario                             | 17 (42)                       |
| Quebec                              | 8 (20)                        |
| Maritime (New Brunswick, Nova Scotia, Prince Edward Island, Newfoundland and Labrador) | 4 (10) |
| Type of practice (n = 39)           |                               |
| Solo                                | 2 (5)                         |
| Group (excluding those in a community health centre) | 19 (49)                     |
| Community health centre             | 18 (46)                       |
| Level of cross-cultural exposure and expertise | 7.5                           |
| Experience working with immigrants or refugees, mean, yr | 25 (64)                     |
| Medical experience in low- and middle-income countries (n = 39) | 26 (65)                     |
| ≥ 7 years’ experience (criteria adapted from Médecins Sans Frontières) (n = 40) | 33 (82)                     |
| Bilingual (n = 40)                  | 17 (42)                       |

*Adapted, with permission, from Swinkels and associates.† Except where indicated otherwise.

**Box 2A: High-priority conditions**

1. Abuse and domestic violence*
2. Anxiety and adjustment disorder*
3. Cancer of the cervix
4. Contraception
5. Dental caries, periodontal diseases*
6. Depression*
7. Diabetes mellitus*
8. Hepatitis B*
9. Hepatitis C
10. HIV/AIDS*
11. Intestinal parasites*
12. Iron-deficiency anemia*
13. Malaria
14. Measles, mumps, rubella, diphtheria, tetanus, pertussis, polio and Hib disease
15. Pregnancy screening
16. Syphilis
17. Torture and post-traumatic stress disorder*
18. Tuberculosis*
19. Varicella (chicken pox)
20. Vision screening

*Conditions identified by consensus in first round (the rest were selected in the second round).
and child health, and mental health. Mental health conditions were rated particularly high, and all four of the proposed mental health conditions reached 80% consensus in the first round of the Delphi survey. Four infectious diseases and three chronic diseases also reached 80% consensus. The inclusion of dental caries and periodontal disease in the top 11 conditions is notable, reflecting important cultural, as well as socio-economic, barriers that refugees and immigrants face in access to dental care.8 This range of conditions suggests that immigrant and refugee medicine covers the full spectrum of primary care. Although infectious disease continues to be an important area of concern, we are now seeing mental health and chronic diseases as key considerations for recently arriving immigrants and refugees.

**Take-home messages**

Preventable and treatable, but often-neglected, health conditions were selected for the development of guidelines for immigrant populations made vulnerable because of health system bias. Criteria that emphasized addressing inequities in health helped in identifying gaps in clinical care. This evidence-based guideline initiative marks the evolution of immigrant and refugee medicine from a focus on infectious diseases to a more inclusive consideration of such chronic diseases as mental illness, dental disease, diabetes mellitus and cancer. We hope that this practitioner engagement process will improve the practicality of the evidence-based guidelines, help practitioners who already to work in the area to target and streamline their efforts, and encourage new practitioners to enter this challenging and interesting discipline.

3. Evaluation of evidence-based literature and formulation of recommendations

A variety of methods are used for developing clinical guidelines and practice recommendations.44 We used the recently developed approach of moving away from recommendations classified by letters and numbers to the simplified classification system recommended by the Grading of Recommendations Assessment, Development and Evaluation (GRADE) Working Group45 and applied this to clinical preventive actions. Our guideline development process followed the Appraisal of Guidelines for Research and Evaluation (AGREE) instrument (www.agreetrust.org), which is recognized internationally as providing best-practice criteria for evidence-based guideline development.

We developed our recommendations on the basis of a pre-specified process overseen by the guideline committee of the Canadian Collaboration on Immigrant and Refugee Health. Defining a methods process ensured that each guideline was developed in a systematic, reproducible manner and was based on the best evidence available. This process was based on existing guidelines, including the Canadian Medical Association (CMA) handbook on developing clinical practice guidelines84 and the ADAPTE framework for adapting existing guidelines.68 Our process emphasized identifying immigrant- and refugee-specific evidence on efficacy and population characteristics from guidelines, systematic reviews and primary studies. When immigrant- and refugee-specific evidence was unavailable, we used specific criteria, adapted from the Cochrane Handbook,89 to judge how this evidence applied to our intended target population.

Conditions considered most important by practitioners caring for immigrants and refugees in Canada were assigned to groups of content experts, who were asked to develop evidence reviews with clinical conclusions for recent immigrants and refugees to Canada using a logic model and following a structured 14-step process. The guidelines focus on clinical care gaps44 during the “health settlement period,” which we define as the first five years of residence in a new country for an immigrant or refugee. This is the period during which health practitioners are likely to have initial contact with this population and the time during which stressors from a person’s country of origin and country of settlement are most likely to manifest. Immigrants and refugees are thus grouped together by this organizing period of resettlement; however, the heterogeneity, complexities and differences between and within these groups were recognized throughout the process.

In our process, we emphasized making clinically relevant recommendations and establishing an extension to existing guidelines rather than a replacement or revision.

Methods

We used the AGREE checklist to guide the overall development process: a panel of experts and a guideline committee set the scope and purpose of the guidelines, and stakeholders were engaged to select priority conditions and to merge recommendations. To ensure rigour and applicability, we developed 14 standardized steps (described below and summarized in Box 1A in section 1 of this article, above). The guideline committee and other guideline experts and practitioners provided feedback to improve clarity of presentation. We accepted funding only from university and government sources, to ensure editorial independence. Here we describe the steps in our standardized evidence review.

Step 1: Develop clinician summary table

A standardized clinician summary template was used in setting the framework for each selected condition. During subsequent steps, this clinician summary table was used to focus development of the preventive guidelines, on the basis of the condition’s prevalence in the population of interest, population-specific clinical considerations (e.g., stigma and awareness of screening and treatment options), clinical actions upon migration, screening tests, screening interval or timing, and treatment.

Step 2: Develop logic model and key questions

Our logic model, which illustrates a plausible causal pathway for each guideline, was adapted from the US Preventive Services Task Force,88 with the addition of a box to consider patient perspectives (for an example, see Figure 3A). The logic model outlines the population of interest (immigrants and refugees); the intervention (i.e., screening); the target condition; adverse effects of screening, diagnosis and treatment; treatment options and outcomes; and the link between treatment and reductions in morbidity and mortality. The model illustrates how identification of the condition can be expected to lead to treatment and reduced morbidity and mortality in the population of interest. This logic model identified the need to consider whether intermediate outcomes would be accepted as the basis for the recommendations, and if so, the strength of association between intermediate and clinical outcomes. For example, high-risk behaviour is an intermediate outcome in reducing morbidity and mortality from HIV.

Review group leaders were asked to use this logic model to define the PICO (population, intervention, comparison and outcome) format for each clinical action. These elements guided the search for evidence.

Step 3: Set the stage for admissible evidence

We followed the process used by the US Preventive Services Task Force and the Canadian Task Force on Preventive Health Care to focus on evidence most critical to making a recommendation.44 We began with searches of specific guidelines and systematic reviews for the target population of immigrants and refugees, to document the current state of direct evidence. We extended these searches to capture evidence from the general population. The search strategy was modelled on that used by the Cochrane Collaboration89 and
was conducted by one of two clinical librarians. The following databases were searched: MEDLINE, PreMEDLINE, Cochrane Database of Systematic Reviews, Database of Abstracts of Reviews of Effectiveness, Embase, CINAHL, National Guideline Clearing House and the CMA Infobase. We also searched the databases and publications of the Canadian Task Force on Preventive Health Care, the US Preventive Services Task Force, the Centers for Disease Control and Prevention, and the World Health Organization. We asked authors to create flow charts of their searches, using the Preferred Reporting Items for Systematic Reviews and Meta-analyses (PRISMA) framework as a template.

**Step 4: Assess eligibility of systematic reviews**

Two members of the review group independently reviewed the search strategies, abstracts and relevant full-text articles on the basis of the inclusion criteria and specified outcomes of interest.

Data from each eligible systematic review were extracted and documented in a table with the following headings: author and year, objective, number and types of studies included, setting, participants, intervention and findings. If no eligible systematic review was found, then the review group team searched for the next best available study (randomized controlled trials, observational studies) that addressed the question.

**Step 5: Search for data specific to immigrant and refugee populations**

A tailored search process was used to gather information on population-specific considerations relevant to immigrants and refugees in the following areas:

- baseline risk (prevalence) versus the Canadian general population
- rate of clinically important beneficial and harmful outcomes (e.g., mortality, morbidity)
- genetic and cultural factors (e.g., knowledge, attitudes, practices, cultural preferences, dietary preferences)
- compliance variation (e.g., physicians’ and patients’ adherence to recommendations)

**Step 6: Refocus on key clinical preventive actions and key questions**

After reviewing the literature and available evidence, review group teams were asked to focus on the most relevant clinical action(s) and immigrant and refugee subpopulation(s) and to select three or fewer candidate recommendations with added value over and above existing guidelines.

**Step 7: Assess quality of systematic reviews**

For each recommendation, all relevant systematic reviews were compared to ensure consistency among findings. If the conclusions of the systematic reviews were consistent, the most recent review was selected. Any inconsistencies in reviews were explicitly addressed: reasons for inconsistencies, including the evidence base or the interpretation, were explored, and the most appropriate systematic review was selected, considering the purposes of these guidelines.

The most relevant systematic reviews were then assessed

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**Figure 3A:** Sample logic model for HIV (adapted from US Preventive Services Task Force). Open rectangles designate the potential screening population and patient factors to be considered; shaded rectangles designate interventions and related outcomes; and circles and numbers provide points in the evidence chain that were used to develop the search questions. Note: ARV = antiretroviral, EIA = enzyme immunoassay.
for quality to ensure they met the four criteria assessed in the National Institute for Health and Clinical Excellence (formerly the Health Development Agency) critical appraisal tool for evidence-based briefings or reviews of reviews: system

Step 8: Search for evidence to update selected systematic reviews
To find new primary studies published since the selected systematic review, a search was conducted using the same approach as in step 3.

Step 9: Assess eligibility of new studies
As in step 4, two reviewers independently screened for relevant studies and then assessed each study for eligibility. Each relevant study was summarized to describe study design, the clinical intervention, details about length of intervention and follow-up, outcomes, population characteristics and data analysis.

For studies evaluating the effectiveness or safety of treatment or screening, the Cochrane Effective Practice and Organisation of Care Review Group’s data collection checklist92 and the Newcastle–Ottawa Scale93 for assessing the quality of nonrandomized studies in meta-analyses were used to assess study limitations.

Step 10: Integrate data from updated search
Any new relevant and eligible studies that could modify or substantially strengthen the conclusions of the “reference” systematic review were assessed and added to the worksheet.

Step 11: Synthesize final evidence bank and draft two key clinical actions
The review group teams synthesized the evidence from the updated systematic reviews, explicitly incorporating clinical considerations and value judgments specific to immigrant and refugee populations to draft preferably no more than two key clinical actions, targeting (where necessary) specific populations or regions.


- High quality: Further research is very unlikely to change our confidence in the estimate of effect.
- Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.
- Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.
- Very low quality: We are very uncertain about the estimate.

Step 12: Develop table for summary of findings
Both desirable and undesirable effects of the intervention were summarized, in both absolute and relative terms, for each patient-important outcome using the summary-of-findings table format adopted by the Cochrane Collaboration. The quality for each outcome was assessed using the items specified by the GRADE Working Group (indirectness, consistency, precision, reporting bias and study limitations) (Box 3A). Observational studies that met these five criteria were upgraded if they also met one of three additional criteria (dose–response, influence of confounding variables, large effect). A separate table was developed for each clinical action or question. For dichotomous outcomes, relative risks or odds ratios were extracted from the reference systematic review (or next best available study). Number needed to treat for one person to benefit was calculated as 1/(control event rate × [1 – relative risk]). The control event rate was taken from the control group of the reference systematic review or best available study.

Step 13: Identify gaps in evidence and needs for future research
The review group teams identified gaps in the literature and outlined recommendations for future research on such topics as implementation, inequalities and vulnerable groups, cost-effectiveness and implications of applying the recommendations in health care settings.

Step 14: Develop clinical preventive recommendations
For each condition, the guideline committee reviewed the clinician summary table, the logic model and the summary-of-findings tables and met with the review group leader to clarify details. Then, for each key clinical action, the guideline committee discussed each of the issues in the GRADE system (see Table 1B in section 1 of this article, above): the balance between desirable and undesirable effects (the relative importance of burden, benefits and harms), quality of the available evidence, and values and preferences. We explicitly decided not to use cost and feasibility in judging the basis of the recommendation because we did not have sufficient confidence in the data. Rather than report the strength of the recommendation as weak or strong, the guideline committee chose to make the recommendation only in the event of net benefits and to report the basis for the recommendation, to provide clinicians with key information to consider when selecting or discussing the preventive recommendation with a patient. The guideline committee took votes if the agreement was not unanimous, and the majority prevailed.

Discussion

This 14-step process was useful for ensuring sufficient uniformity among the transdisciplinary teams for each condition. Specifically, this systematic approach enabled the review group teams to meet the requirements of the GRADE quality-assessment process and the steering group to apply the GRADE recommendation process. These steps were also designed to conform with AGREE, the current quality stan-
standard for guidelines. We worked with each review group leader and team to ensure we met the 23 AGREE criteria in six domains: scope and purpose, stakeholder involvement, rigour of development, clarity and presentation, applicability and editorial independence.16

**Take-home messages**

We combined the AGREE best-practice framework, the current quality standard for guidelines, with the recently developed GRADE approach to quality assessment to develop evidence-based clinical preventive guidelines for immigrants and refugees to Canada. Here, we have documented the systematic approach used to produce the evidence reviews and apply the GRADE approach. The 14-step approach included building on evidence from previous systematic reviews, searching for and comparing evidence between general and specific immigrant populations, and applying the GRADE criteria for making recommendations. The basis of each recommendation (balance of benefit and harm, quality of evidence, values) is stated explicitly to ensure transparency.

4. Measles, mumps, rubella, diphtheria, pertussis, tetanus and polio

Vaccination is one of the most beneficial and cost-effective measures for preventing disease.93-96 Before routine vaccination, the annual burden of measles, mumps, rubella, diphtheria, pertussis, tetanus and polio in the United States and Canada was considerable. The incidence of and mortality from these diseases have been reduced substantially: by more than 99% for measles, rubella, diphtheria and polio, by more than 95% for mumps, and by more than 92% for pertussis and tetanus relative to annual morbidity and mortality before introduction of the corresponding vaccines.97 Despite these successes, the recent outbreaks of pertussis in California, outbreaks of mumps in the United States and Canada in 2005–2006 and the ongoing transmission of polio in the past five years, with recent spread to Tajikistan, highlight the need to maintain high levels of herd immunity and to identify and vaccinate susceptible groups so that outbreaks can be prevented.98,99 Almost 20% of the Canadian population is foreign born,100 and in the past 30 years the majority of these people (more than 70%) have originated from countries where vaccination coverage may be suboptimal or where several of the childhood vaccines that are routine in Canada are not part of the national vaccination schedule.101 Immigrants are therefore likely to be an unrecognized group at risk for childhood vaccine-preventable diseases. We conducted an evidence review to guide primary care practitioners in the need to assess and update childhood vaccination in the immigrant population. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on updating vaccines are outlined in Box 4A.

Methods

We used the 14-step method developed by the Canadian Collaboration for Immigrant and Refugee Health102 (summarized in section 3 of this article, above). We considered the epidemiology of measles, mumps, rubella, diphtheria, pertussis, tetanus and polio in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE (Ovid), MEDLINE InProcess, Embase, CINAHL and the Cochrane Library from Jan. 1, 1950, to Jan. 14, 2010, for studies pertinent to immigrants and from Jan. 1, 1997, to Jan. 14, 2010, for studies pertinent to the general population. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for this topic (Appendix 3, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

Results

In the search for systematic reviews and guidelines for immigrants regarding measles, mumps, rubella, diphtheria, pertussis, tetanus and polio, we identified and screened 242 records, of which none met the eligibility criteria. In the search for systematic reviews and guidelines involving these diseases in the general population, we identified 6293 articles, of which 24 met the eligibility criteria. A search for articles reporting information about admission to hospital and mortality associated with measles, mumps, rubella, diphtheria, pertussis, tetanus and polio identified 3888 articles (after duplicates were removed), of which 59 were relevant, and one of these was critical for this review97 (Table 4A). In addition, a search for articles about immigrants and measles, mumps, rubella, diphtheria, pertussis, tetanus and polio identified 1177 articles (duplicates removed), of which 54 were relevant, addressing the following areas: epidemiology, prevaccination screening, knowledge and compliance, treatment and vaccination in the immigrant population.

What is the burden of vaccine-preventable diseases in immigrant populations?

A large proportion of immigrants and refugees, particularly adults, are likely to be susceptible to vaccine-preventable diseases because of underimmunization, waning immunity or both. Underimmunization likely plays an important role, given that vaccine coverage globally ranges from 50% to 90%, that routine childhood vaccination began only in the mid-1970s and that rubella and mumps vaccines are not administered routinely in most developing countries.101 Given the progress in global vaccination coverage, immigrant children and adolescents are more likely than their parents to have received vaccines that are part of the World Health Organization Extended Program on Immunization (measles, diphtheria, pertussis, tetanus, polio, bacille Calmette-Guérin), but many may not have received other vaccinations that are part of the routine childhood vaccination program in Canada (mumps, rubella, varicella, Hemophilus influenzae, Streptococcus pneumoniae). Many immigrants, especially adults, do not have vaccination records, and even when present, more than 50% of such records may not be current according to the host country’s vaccination schedules.98,102 Seroprevalence studies consistently have shown that a large proportion of adult immigrants are susceptible to rubella (about 80%–85% immune but as low as 75%) and tetanus (about 50%–60% immune among those 20–30 years of age, but decreasing with increasing age).97,98,103,104 A higher-than-expected proportion of immigrants are involved in rubella outbreaks, and most reported cases of congenital rubella syndrome and neonatal tetanus have occurred in children born to unimmunized foreign-born mothers.103,104 In adult immigrants, seroprevalence studies of measles (> 95% immune) and mumps (80%–92% immune but as low as 70%) have generally shown adequate antibody levels, with
some exceptions. However, immigrants have not been over-represented in recent measles and mumps outbreaks. Diphtheria seroprevalence in immigrants is low (range 35%–50%) and generally decreases with age. To maintain herd immunity in the population, certain threshold levels of antibodies need to be maintained: 91%–94% for measles, 90%–92% for mumps, 83%–85% for rubella, 80%–85% for diphtheria, 80%–85% for polio and 90%–94% for pertussis. Immigrants likely fall below this threshold for susceptible to tetanus and at risk for the morbidity and mortality associated with this disease. Any population in which a large proportion of individuals are susceptible to vaccine-preventable disease will be at risk for disease transmission. High-risk groups must therefore be identified so that targeted vaccination programs can be formulated and implemented.

### Does vaccination against specific vaccine-preventable diseases decrease associated morbidity and mortality?

#### Relative benefits and harms of vaccination

In the prevaccination era, diseases such as measles, mumps, rubella, diphtheria, pertussis, tetanus, smallpox and polio were very frequent and were a major cause of morbidity and mortality. These diseases also had enormous societal and economic costs, including time off school and work, physician visits and admissions to hospital. Childhood vaccination programs have decreased the morbidity from these diseases by more than 92%–99% and mortality by more than 99%. Childhood vaccination programs have repeatedly been found to be one of the most cost-effective medical interventions.

#### Measles, mumps and rubella

Measles–mumps–rubella vaccine is highly effective against measles and rubella. Almost 100% of individuals are protected against measles after two doses, and more than 95% are protected against rubella after a single dose, with antibodies persisting for at least 15 years. The effectiveness of mumps vaccine is lower and depends on the vaccine strain used, the time since vaccination and possibly the genotype of the wild type. In the recent US outbreaks, the effectiveness of mumps vaccine was estimated to be as low as 64% after one dose and 79% after two doses (Jeryl Lynn strain). Measle–mumps–rubella vaccine has been associated with fever (about 5%), febrile convulsions (0.3%), benign thrombocytopenia purpura (< 0.01%), parotitis (rarely) and arthritis (up to 25% in postpubertal women), usually within two weeks of vaccination. The frequency of adverse reactions in seronegative women, however, is higher among those who have never been vaccinated than among re-vaccinated seronegative women. In 1998 Wakefield and colleagues published a case series of 12 children with development disor-

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**Box 4A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: vaccine-preventable diseases**

**Measles, mumps and rubella**

- Vaccinate all adult immigrants without immunization records using one dose of measles–mumps–rubella vaccine.
- Vaccinate all immigrant children with missing or uncertain vaccination records using age-appropriate vaccination for measles, mumps and rubella.

**Basis of recommendations**

**Balance of benefits and harms**

Childhood vaccination programs have dramatically decreased the incidence of and associated mortality from measles, mumps, rubella and congenital rubella (absolute difference of 95.9%–99.9% in reduction of cases and 100% in reduction of deaths). Serious adverse events, including autism (relative risk 0.92, 95% confidence interval 0.68–1.24), are not significantly associated with measles–mumps–rubella vaccine. Mumps and rubella are not part of routine vaccination programs in most source countries of origin for the majority of new immigrants. A large proportion of adult immigrants may be susceptible to rubella (20%–30%) and at risk for having a child with congenital rubella syndrome.

**Quality of evidence**

High

**Values and preferences**

The committee attributed more value to preventing the risk of outbreaks and the individual burden due to these diseases and less value to the cost of vaccination.

**Diphtheria, pertussis, tetanus and polio**

- Vaccinate all adult immigrants without immunization records using a primary series of diphtheria, tetanus and inactivated polio vaccine (three doses), the first of which should include acellular pertussis vaccine to also protect against pertussis.
- Vaccinate all immigrant children with missing or uncertain vaccination records using age-appropriate vaccination for diphtheria, pertussis, tetanus and polio.

**Basis of recommendations**

**Balance of benefits and harms**

Childhood vaccination programs have dramatically decreased the incidence of and associated mortality from diphtheria, pertussis, tetanus and polio (absolute difference of 92.9%–99.9% in reduction of cases and 99.2%–100% in reduction of deaths) relative to the prevaccination period, without associated increases in serious adverse events. A large proportion of adult immigrants are susceptible to tetanus (40%–50%) and diphtheria (about 60%), and the proportion susceptible increases for both with increasing age. To prevent individual morbidity and mortality and to prevent outbreaks, susceptible individuals must be identified and vaccinated.

**Quality of evidence**

High

**Values and preferences**

The committee attributed more value to preventing the risk of outbreaks and the individual burden due to these diseases and less value to the cost of vaccination.
Diphtheria, acellular pertussis, tetanus and polio

Diphtheria and tetanus vaccines are highly effective, with protective levels of antibody developing in more than 95% of individuals after a primary series. Antibodies to diphtheria, which persist for up to 10 years, wane more rapidly than those for tetanus, which persist for up to 25 years. There is no immunologic correlate of protection for pertussis, but the clinical protective efficacy of the diphtheria, tetanus and acellular pertussis vaccine for children is about 85%. In over 99% of those vaccinated, protective levels of antibodies to all three serotypes of polio vaccine develop after three doses, and a one-time booster in adulthood is required to maintain immunity. For those who have received a primary series of tetanus and diphtheria vaccine, booster doses for tetanus–diphtheria vaccine (adult) are recommended every 10 years to maintain immunity. Individuals older than seven years of age who have not received a primary series should be given three doses of tetanus–diphtheria vaccine (adult), the first being tetanus–diphtheria–acellular pertussis, to provide protection against pertussis. In addition, during management of any wound other than clean or minor wounds, these individuals should receive a primary series of vaccine and also tetanus immune globulin. Given that infants constitute the group at highest risk for development of severe pertussis, that adults are the main reservoir for pertussis and that antibodies wane over time (5–10 years after a primary series), all adults should be given a one-time booster of tetanus–diphtheria–acellular pertussis.

Local pain, swelling and erythema are common after administration of the pediatric tetanus–diphtheria–acellular pertussis vaccine and are reported in up to 40% of those vaccinated; the corresponding rates are 75% after adult tetanus–diphtheria–acellular pertussis vaccine and 60%–70% after tetanus–diphtheria vaccine. Fever occurs in less than 5% during a primary series; however, the pediatric and adult tetanus–diphtheria–acellular pertussis vaccines have not been associated with an increased risk of serious adverse events.

Clinical considerations

All adults and children without written immunization records should restart a primary immunization schedule appropriate for their age. An alternative, though somewhat less practical, approach is to test for antibodies to the major vaccine antigens and administer any vaccines to which the person has no immunity. A limitation to this approach is that serologic testing for diphtheria and tetanus is not widely available in most settings. Similarly, before one or two doses of trivalent measles–mumps–rubella vaccine are given, three serologic

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**Table 4A: Summary of findings for vaccination to prevent rubella and tetanus**

**Outcome** | **Absolute effect** | **Risk for control group (annual average)** | **Difference with vaccination** | **Relative effect** | **No. of participants (studies)** | **GRADE quality of evidence**
---|---|---|---|---|---|---
**Rubella** | | | | | | |
Cases, rubella | | 47 745 (for 1966–1968) | 47 734 fewer cases (for 2006) | Absolute difference: 99.9% reduction | NA (1) | High**†
**Tetanus** | | | | | | |

Note: GRADE = Grading of Recommendations Assessment, Development and Evaluation; NA = not applicable.

*Only one study.
†Reduction in absolute numbers > 90%.
tests need to be done and the patient needs to return for the results before the vaccination series can be started. In one study, it was less costly and more effective to vaccinate all individuals if more than 80% had not completed the full diphtheria–tetanus vaccine series or if antibody seroprevalence to both diphtheria and tetanus was less than 51%.118

Children are more likely than adults to have written immunization records; however, the optimal approach for those with vaccination records is challenging. Interpreting written records can be difficult because of language barriers and the fact that immunization schedules and products used in other countries may differ from those used in Canada. Even if records can be translated, they may not necessarily predict immunity. In several studies of internationally adopted children with seemingly appropriate records, many (up to 50%) did not have serologic evidence of protective immunity to the specific antigen. This discordance has been ascribed to falsification of records, breaches in the cold chain in the countries of origin or host factors.119,120 Given this uncertainty, the most conservative approach would be to give a full vaccination series or, as mentioned above, perform prevaccination screening and vaccinate those without immunity.111

Barriers to vaccine uptake in children are well described, but a recent Canadian study showed that immigrant children were more likely to be vaccinated than children in the host population.121,122 In spite of this, many immigrant children may not have received all the vaccines that are part of the Canadian immunization schedule, and there are no structured catch-up vaccination programs for these children after arrival. Updating vaccination for adult immigrants is even more challenging than for children, in part because they are not a well-recognized group at risk, but also because of the lack of routine well-adolescent or well-adult visits to access this population, the lack of structured programs to update vaccines and the cost (many vaccines are often not covered). In addition, missed opportunities for catch-up vaccination in adult immigrants when they come into contact with the health care system are well documented.123,124 With this in mind, health care providers need to remain vigilant in assessing and updating vaccinations in newly arrived immigrant and refugee children and adults.

**Recommendations of other groups**

The Canadian National Advisory Committee on Immunization recommends that all persons without written vaccination records should receive an age-appropriate primary vaccination series. Adults with no records or an unclear history of prior vaccination should receive a primary series of tetanus–diphtheria vaccine (three doses, one of which should be tetanus–diphtheria–acellular pertussis vaccine, to provide protection against pertussis), as well as a primary series for polio, as inactivated poliovirus vaccine (three doses, at baseline, 4–8 weeks and 6–12 months). The National Advisory Committee on Immunization also recommends that a single dose of measles–mumps–rubella vaccine be given to adults born after 1970 who do not have a history of measles or who are seronegative for mumps or rubella.111 Our recommendations highlight the importance of making primary care providers aware of the gaps in vaccination in newly arrived immigrants and refugees of all ages and the need to update vaccination against measles, mumps, rubella, diphtheria, pertussis and tetanus in this population.

**Take-home messages**

- A large proportion of immigrants are susceptible to several childhood vaccine-preventable diseases (especially rubella, diphtheria and tetanus) and are at risk for associated morbidity and mortality.
- Individuals susceptible to vaccine-preventable diseases must be identified and vaccinated to maintain herd immunity and prevent outbreaks.
- Health care providers need to aware of these gaps and ensure that they take all opportunities to update vaccinations in newly arrived immigrants and refugee children and adults.

5. Varicella

Varicella occurs at older ages in tropical countries, and most of these countries do not have varicella vaccination programs, which means that a large proportion of adolescent and adult immigrants from tropical countries (about 30%) are susceptible to the disease.39,125-128 This is important because adults are more likely than children to contract severe varicella, with higher rates of pneumonia, encephalitis, admission to hospital and death.129 Varicella may also result in poor outcomes for pregnant women and their fetuses and infants. Several outbreaks of varicella have been documented in adult immigrants from tropical countries soon after arrival in temperate countries.130-134 In a recent US study, the mortality rate for varicella was higher in the foreign-born population than in the US-born population, likely because varicella occurs at older ages among immigrants.135 Canada has no programs to verify immune status to varicella, nor are there any systematic targeted or catch-up vaccination programs for the immigrant population. We conducted an evidence review to guide primary care practitioners in the need to assess varicella immune status in the immigrant population. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on varicella vaccination are outlined in Box 5A.

Methods

We used the 14-step method developed by the Canadian Collaboration for Immigrant and Refugee Health16 (summarized in section 3 of this article, above). We considered the epidemiology of varicella in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE (Ovid), MEDLINE InProcess, Embase, CINAHL and the Cochrane Database of Systematic Reviews from 1950 to Jan. 19, 2011, for relevant articles pertinent to immigrants and from Jan. 1, 1997, to Jan. 19, 2011, for articles pertinent to the general population. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for varicella vaccination (Appendix 4, available at www.cmaj.ca /lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

Results

In the search for systematic reviews and guidelines relevant to immigrants, we identified six records, but none of the studies followed a systematic review methodology. We identified 743 records for the general population, and of these, 11 were included: three recent guidelines from Canada and one guideline from the United States on preventing varicella, one review of varicella in pregnancy, two reviews on the efficacy of and adverse events due to varicella vaccine and the effectiveness of varicella vaccination programs, one review of the predictive value of a history of varicella infection, one systematic review on the cost-effectiveness of varicella vaccine, one review of barriers to adolescent vaccination and one systematic review on interventions to improve compliance with vaccination; we also identified a key article on the impact of varicella vaccination on health care utilization.136 In addition, the search for varicella AND immigrants or refugees identified 31 relevant articles that addressed epidemiology, cost-effectiveness of vaccination and screening, vaccine knowledge and compliance. A search for articles dealing with varicella-associated morbidity and mortality identified 3400 articles, of which 122 were relevant. These included articles on epidemiology, admission to hospital and mortality due to varicella, screening for varicella immunity, vaccination effi-

Box 5A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: varicella (chicken pox)

- Ensure that immigrants and refugees of all ages are immune to varicella.
- Vaccinate all immigrant children < 13 years of age with varicella vaccine without prior serologic testing.
- Screen all immigrants and refugees from tropical countries ≥ 13 years of age for serum varicella antibodies, and vaccinate those found to be susceptible.

Basis of recommendations

Balance of benefits and harms

Varicella vaccination programs have substantially decreased ambulatory care visits (number needed to vaccinate [NNV] 794, 95% confidence interval 688-990) and mortality (NNV 3 031 773) due to varicella in all age groups. The adverse effects of vaccination are minimal and include minor pain and redness at the injection site, rashes and fevers. A large proportion (> 30%) of adolescents and adult immigrants from tropical countries are susceptible to varicella because it occurs at an older age in tropical countries and because most of these countries do not have a varicella vaccination program. As a result, immigrants from tropical countries are at increased risk of developing severe varicella after arrival in Canada, as varicella develops at an older age and there are no systematic catch-up varicella vaccination programs for immigrants.

Quality of evidence

Moderate

Values and preferences

The committee attributed high value to reducing morbidity and mortality from varicella, which has a high burden of disease in adolescent and adult immigrant populations. For children < 13 years of age, it is cost-saving to vaccinate all without prior serologic testing. In a cost-effectiveness analysis of adult immigrants and refugees, it was most cost-effective to vaccinate without prior screening with a seroprevalence of < 84% (children, most adolescents and some adults) and to serotest before vaccination when the seroprevalence was between 85% and 95% (some adolescents and most adults). If serologic testing results in extra costs or presents a barrier to completion of the vaccination series, vaccination without prior serologic testing should be offered. The most effective strategy will be one that is tailored to the vaccination setting and that balances the cost and anticipated uptake of the vaccine and the availability of, compliance with and costs of serologic testing.
cacy and cost-effectiveness, and vaccine knowledge and compliance. The key article on varicella-associated mortality (used for Table 5A) was identified in this search.135

What is the burden of varicella in immigrant populations?

Larger proportions of adolescents and adults from tropical countries are susceptible to varicella (about 50% at age 15 years and about 10%–15% from age 30 to 35 years) than is the case for Canadians (about 10% at age 15 years and less than 2% from age 30 to 35 years). This is because the mean age at which varicella develops is older in tropical countries (10–15 years) than in temperate or cold countries (4–5 years). In certain tropical regions (the Caribbean, Sri Lanka, Singapore, Indonesia and rural areas of other countries such as Pakistan and India), a large proportion of adults older than 35 years may still be susceptible to varicella. Immigrants from tropical countries have been involved in several varicella outbreaks and have higher mortality from varicella.133,134,135

The severity of varicella increases with age. Older age of infection increases the risk of varicella in pregnancy, which may result in poor outcomes for both the mother and the baby. Pregnant women are more likely than other adults to develop varicella pneumonia, to be admitted to hospital and to die from varicella. Congenital varicella syndrome is associated with a fetal case fatality rate of up to 50%, and survivors may have congenital anomalies, including limb hypoplasia, microcephaly and dermatomal scarring.136 In addition, 20% of newborns born to women infected with varicella between five days before and three days after delivery will develop neonatal varicella, which carries a case fatality rate of 30%–50%140,141.

Does screening for immunity to varicella and vaccinating those who are susceptible decrease morbidity and mortality?

Diagnostic tests to screen for varicella antibodies

The most widely available tests to detect antibodies to varicella zoster virus are enzyme immunoassays. These tests are less sensitive than the fluorescent antibody to membrane antibody test, but they have good specificity (> 95%). Enzyme immunoassays are sufficiently sensitive to detect protective antibodies after natural infection with varicella zoster virus (60%–92%) but are inadequately sensitive to detect protective antibodies resulting from vaccination.142 Given that the majority of adult immigrants at risk for varicella in Canada originate from countries without routine childhood vaccination programs, enzyme immunoassay is an acceptable method of screening for immunity to varicella in this population.

History of prior varicella to determine varicella immune status

The majority of adults (> 95%) who report a history of varicella are immune to the virus.143 However, a positive history of varicella may less reliably predict protective varicella antibodies in immigrant populations where varicella seroprevalence is lower than expected for age.70,144 In adult immigrant populations, it may be more prudent to perform serologic testing on all individuals before vaccination or to vaccinate empirically (if serologic testing is not available), to avoid the potential poor outcomes associated with varicella.

Efficacy of varicella vaccination

A single dose of varicella vaccine is 80% to 85% effective in preventing disease of any severity and is more than 95% effective in preventing severe varicella.145 Vaccine efficacy for adults has been estimated at about 80%. Because of plateauing rates of varicella between 2003 and 2006 and ongoing outbreaks, despite a one-dose vaccination schedule, the United States recommended a universal two-dose childhood varicella vaccination program (first dose at 12–15 months, second dose at four to six years) in 2006.146 The National Advisory Committee on Immunization of Canada has also recently recommended that two doses of varicella vaccine be given to all children under 13 years of age.147 Adverse events associated with the vaccine include pain and redness at the injection site (22%–35%), rash (which is sometimes varicella-like; 1%–5%) and fever (4%–7%). Secondary transmission has been documented from five vaccine recipients.148 Varicella vaccine is a live attenuated vaccine that should be avoided in immunosuppressed individuals. It can be given to HIV-infected individuals with mild to moderate symptoms and a CD4 count above 200 × 10⁹/L or less than 15%.149

Effectiveness of varicella vaccination to decrease morbidity and mortality

Varicella was a common childhood illness before implementation of widespread universal childhood vaccination programs in Canada (2005) and the United States (1995). These programs have resulted in a significant decrease in the number of cases and in the number of varicella-associated hospital admissions and deaths (> 75%) in all age groups, including adults (Table 5A).133,134,135 A recent study of the effect of varicella vaccine in Ontario showed decreases in varicella-associated hospital admissions (by 53%, 95% confidence interval [CI] 48%–58%), emergency department visits (by 43%, 95% CI 41%–44%) and physician visits (by 45%, 95% CI 44%–45%) relative to the prevaccination period (1992–1998).150

Cost-effectiveness of varicella vaccination

Cost-effectiveness analyses of routine varicella vaccination of preschool-age children have demonstrated that the vaccine (one or two doses) is cost-saving from a societal point of view.151,152 Most cost-effectiveness studies support serologic testing before the vaccine is given to adolescents and adults. A recent cost-effectiveness analysis of strategies to prevent varicella in adult immigrants and refugees found that various vaccination strategies were cost-saving, relative to no intervention, at the following seroprevalence thresholds: vaccinate all without prior serologic testing (threshold 84%); would include immigrant children under 13 years of age, many ado-
lescents and some adults); provide serologic testing for all, and vaccinate those found to be susceptible (threshold 85%–92%; would include some adolescents and most adults); provide serologic testing only for those with a negative or unknown history of varicella (threshold 93%–95%) and vaccinate those found to be susceptible; and no intervention (threshold > 95%).134

Clinical considerations

What are the potential implementation issues?

We found no data specific to newly arrived immigrants or refugees. General factors associated with low uptake of varicella vaccine were similar to those for other vaccines (e.g., low socioeconomic status, low parental education, younger maternal age, lack of knowledge about the disease and vaccination, negative beliefs about and attitudes toward immunization, fear of adverse effects, lack of transport, inconvenient clinic hours), and cost was the most important barrier.122,143 Adolescents and adults face additional barriers to vaccination relative to children, including lack of awareness of the need for vaccination, lack of routine well-adolescent or well-adult visits and lack of coordinated vaccination programs for these populations.124,125 124 The two most important barriers to vaccination for adult immigrants are likely to be cost of the vaccine and lack of awareness among health care providers that adult immigrants are at increased risk for severe varicella. In certain settings, serologic testing may result in extra costs or may present a barrier to completion of the vaccination series, in which case vaccination without prior serologic testing should be offered.

The most effective interventions to improve uptake of vaccination in the general population were instituting reminder or recall systems, educating target populations and vaccine providers, and reducing out-of-pocket costs.155,156 The most effective interventions to improve vaccination coverage among adults are standing orders and expanding access to vaccination in nontraditional settings (e.g., schools, workplaces and social gathering places such as churches and sports clubs).154,156

<table>
<thead>
<tr>
<th>Table 5A: Summary of findings for vaccination to prevent varicella and related morbidity and mortality</th>
</tr>
</thead>
</table>
| **Patient or population:** People up to 49 years of age;126 national death records15
| **Setting:** MarketScan database with enrollees from over 100 health insurance plans of about 40 large US employers, from 1994 to 2002;126 *National Center for Health Statistics multiple cause-of-death mortality data for 1990 through 2001*15
| **Intervention:** Vaccination against varicella
| **Comparison:** Prevaccine era
<p>| <strong>Absolute effect</strong> |</p>
<table>
<thead>
<tr>
<th><strong>Outcome</strong></th>
<th><strong>Risk for control group (prevaccine era)</strong></th>
<th><strong>Difference with vaccination</strong></th>
<th><strong>Relative effect (95% CI)</strong></th>
<th><strong>No. of participants (studies)</strong></th>
<th><strong>GRADE quality of evidence</strong></th>
<th><strong>Comments (95% CI)</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Admission to hospital (primary diagnosis of varicella)</td>
<td>2.3 per 100 000</td>
<td>2.0 fewer per 100 000</td>
<td>RR 0.13 (0.04–0.41)</td>
<td>Not reported* (1)156</td>
<td>High†</td>
<td>NNT 49 975 (45 384 – 73 443)</td>
</tr>
<tr>
<td>Ambulatory visits (primary diagnosis of varicella)</td>
<td>215 per 100 000</td>
<td>126 fewer per 100 000</td>
<td>RR 0.41 (0.32–0.53)</td>
<td>Not reported* (1)156</td>
<td>Moderate‡</td>
<td>NNT 794 (688–990)</td>
</tr>
<tr>
<td>Death from varicella</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Overall</td>
<td>0.56 per 1 000 000‡</td>
<td>0.33 fewer per 1 000 000§</td>
<td>RR 0.41 (0.25–0.66)</td>
<td>Not reported* (1)155</td>
<td>Moderate‡</td>
<td>NNT 3 031 773 (2 393 719 – 5 314 626)</td>
</tr>
<tr>
<td>Foreign-born</td>
<td>0.35 per 1 000 000§</td>
<td>0.25 fewer per 1 000 000¶</td>
<td>RR 0.29 (0.14–0.58)</td>
<td>Not reported* (1)155</td>
<td>Moderate‡</td>
<td>NNT 4 024 145 (3 322 259 – 6 802 721)</td>
</tr>
<tr>
<td>US-born</td>
<td>0.19 per 1 000 000$</td>
<td>0.12 fewer per 1 000 000¶</td>
<td>RR 0.37 (0.15–0.88)</td>
<td>Not reported* (1)155</td>
<td>Moderate‡</td>
<td>NNT 8 354 219 (6 191 950 – 43 859 649)</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NNT = number needed to treat; RR = risk ratio.

*Because of the nature of the original studies, these studies did not report the total number of participants.

†Graded up two levels to reflect very strong evidence of association (RR < 0.2).
‡Graded up one level to reflect strong evidence of association (RR < 0.5).
§Age-adjusted rate from Nguyen et al.155
¶Age-adjusted rate from Nguyen et al.155

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Recommendations of other groups

The National Advisory Committee on Immunization in Canada recently recommended that all children between the ages of 12 months and 12 years should receive two doses of varicella vaccine as primary immunization. Persons 13 years of age and older should receive two doses of vaccine a minimum of six weeks apart. Certain groups at increased risk for varicella, such as immigrants from tropical countries, women of child-bearing age, household contacts of immunocompromised people, health care workers and adults who work in other occupations with increased exposure to varicella (teachers, daycare workers, etc.), should be considered for targeted varicella vaccination. Since 2007, the US Advisory Committee on Immunization Practice has recommended that both children and adults receive two doses of varicella vaccine. Our recommendations highlight the importance of varicella vaccination for susceptible immigrants.

Take-home messages

• Among immigrants from tropical countries, a large proportion of adolescents (up to 50%) and adults (up to 10%) are susceptible to varicella and are at increased risk of severe varicella.
• Pregnant immigrant women and their babies are at highest risk for complications of varicella.
• The mean age at which varicella develops varies for different world regions: it is acquired at an older mean age in tropical countries (15 years) than in temperate and cold countries (5 years).
• In certain tropical regions (the Caribbean, Sri Lanka, Singapore, Indonesia and rural areas of other countries such as Pakistan and India), the mean age of acquiring varicella may be even older, and a large proportion of adults older than 35 years may still be susceptible to the disease.
• Immigrants and refugees of all ages from tropical countries would benefit from having their varicella immune status verified and being offered varicella vaccine if found to be susceptible.

6. Hepatitis B

Hepatitis B virus infection is an important global health problem affecting 350 million people worldwide and leading to one million premature deaths from chronic liver disease and hepatocellular carcinoma annually. About 3% of immigrants to Canada but only 0.5% of Canadian-born people have chronic infection with hepatitis B virus. Most people with chronic infection are asymptomatic, and their disease goes undetected and untreated. This is likely why mortality from viral hepatitis and hepatocellular carcinoma among immigrants is two to four times higher than in the Canadian-born population.

Over the past 10 years, several medications have become available that decrease viral replication and morbidity from chronic liver disease. Screening in high-risk populations and targeted treatment have therefore become important strategies to decrease the burden of chronic infection with hepatitis B virus. An effective vaccine (available in Canada since 1982) is an important tool to control transmission of hepatitis B. Despite the disparity in populations and the availability of a safe and effective vaccine, no programs in Canada systematically screen immigrants for chronic hepatitis B virus infection, nor are there systematic targeted or catch-up hepatitis B vaccination programs outside of the childhood vaccination program. We conducted a review to quantify the burden of chronic hepatitis B among immigrants, to search for evidence of the effectiveness of screening and vaccination programs, and to identify barriers or challenges to implementing such programs. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on screening for and vaccinating against hepatitis B virus are outlined in Box 6A.

Methods

We used the 14-step approach developed by the Canadian Collaboration for Immigrant and Refugee Health (summarized in section 3 of this article, above). We considered the epidemiology of disease in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE, Embase, CINAHL, the Cochrane Library and other sources from Jan. 1, 1996, to Jan. 1, 2010. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for hepatitis B (see Appendix 5, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

Results

In the search for systematic reviews and guidelines for immi-
grants, we identified and screened 54 records. Of the six that met our eligibility criteria, all were narrative reviews on screening immigrants for hepatitis B and were excluded because none followed a systematic review method. A total of 2565 records were identified and screened from the search for systematic reviews and guidelines for the general population and the Web-based search. Fourteen records met the eligibility criteria: two guidelines from the United States on screening for chronic hepatitis B, three guidelines addressing treatment of chronic hepatitis B virus infection, one systematic review and one guideline on screening for hepatocellular carcinoma, one systematic review and one guideline on preventing hepatitis B virus infection among neonates, one systematic review on the adverse effects of hepatitis B vaccination, one systematic review on improving immunization rates and three guidelines on vaccinating against hepatitis B. In the search for articles relevant to immigrants and hepatitis B, we identified 148 articles addressing the following areas: epidemiology, screening, knowledge and compliance, treatment, and vaccination in the immigrant population.

**How does hepatitis B virus affect immigrant populations?**

Canada has low rates of hepatitis B virus infection and an overall seroprevalence of chronic infection less than 0.5%. Over the past 40 years, most immigrants who have arrived in Canada (> 70% of 250 000 per year) have originated from countries with intermediate or high rates of endemic hepatitis B (Table 6A).

These immigrants have an overall seroprevalence of chronic infection with hepatitis B virus of about 4%. People with chronic hepatitis B virus infection have a 15%–25% lifetime risk of dying from cirrhosis and hepatocellular carcinoma. They are typically asymptomatic until they present with end-stage liver disease or hepatocellular carcinoma several decades after infection. Hepatocellular carcinoma is one of the most fatal types of cancer, and the five-year survival rate is less than 11%, because symptoms usually appear only at an advanced stage, when the disease is not treatable. The growing pool of asymptomatic, undetected and untreated hepatitis B virus infection in the immigrant population (8000 to 26 000 new imported infections per year) is likely why mortality is higher among immigrants than in the Canadian-born population: 1.8–3.8 times higher from viral hepatitis and 2.2–4.9 times higher from hepatocellular carcinoma. The burden of undetected chronic infection with hepatitis B virus in immigrants is likely also in part responsible for the 8.4-fold higher mortality from chronic infection with hepatitis B virus and the 2.2-fold higher incidence of hepatocellular carcinoma over the past 30 years in Canada (between 1969 and 1997).

Young children living in a household that includes someone with chronic hepatitis B virus infection have rates of acquiring new infection of 1%–2% per year during the first decade of life. Immigrant children, therefore, are at risk of acquiring hepatitis B in Canada, as they are more likely to live in a household that includes someone with silent hepatitis B virus infection. Many immigrant children have not been vaccinated in the universal childhood immunization program because they are too young or arrived after the age of vaccination. This risk was highlighted in a study in which Quebec’s school-based vaccination program (for children in grade 4) was evaluated 10 years after initiation. In that study, rates of acute infection with hepatitis B virus decreased in all age groups except those 10 years of age and younger, and 53% of these cases occurred in foreign-born children. The study underscored both the need to provide protection against

<table>
<thead>
<tr>
<th>Disease characteristic</th>
<th>High prevalence</th>
<th>Intermediate prevalence</th>
<th>Low prevalence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prevalence of chronic infection with HBV, %</td>
<td>≥ 8</td>
<td>2–7</td>
<td>&lt; 2</td>
</tr>
<tr>
<td>Geographic distribution</td>
<td>Southeast Asia, China, Pacific Islands, sub-Saharan Africa, Alaska, Peru, northwestern Brazil</td>
<td>Mediterranean basin, eastern Europe, central Asia, Japan, Amazon basin, Middle East</td>
<td>United States and Canada, western and northern Europe, Australia, New Zealand</td>
</tr>
<tr>
<td>Percentage of global population, %</td>
<td>45</td>
<td>43</td>
<td>12</td>
</tr>
<tr>
<td>Predominant mode of infection</td>
<td>Maternal-infant, percutaneous or mucosal</td>
<td>Maternal-infant, percutaneous or mucosal, sexual</td>
<td>Sexual, percutaneous or mucosal</td>
</tr>
<tr>
<td>Predominant age at acquisition of infection</td>
<td>Perinatal (vertical) and early childhood (horizontal)</td>
<td>All age groups</td>
<td>Adult</td>
</tr>
<tr>
<td>Likelihood of chronic infection after acute infection with HBV, %*</td>
<td>80–90</td>
<td>30–60</td>
<td>&lt; 5</td>
</tr>
<tr>
<td>Mean age at which hepatocellular carcinoma develops, yr*</td>
<td>57</td>
<td>57</td>
<td>75</td>
</tr>
<tr>
<td>Lifetime probability of infection (immunity), %</td>
<td>&gt; 60</td>
<td>20–60</td>
<td>&lt; 20</td>
</tr>
</tbody>
</table>

*Likelihood that chronic infection with HBV will develop after acute infection with HBV is inversely proportional to age at acquisition. The earlier the infection is acquired, the earlier hepatocellular carcinoma will develop.
hepatitis B as early as possible for children (possibly changing to routine infant rather than childhood vaccination programs) and the need to provide catch-up vaccination for immigrants.

**Does screening for chronic infection with hepatitis B virus decrease morbidity and mortality?**

**Screening tests**
Serologic tests to detect hepatitis B virus are inexpensive, and commercially available tests are sensitive and specific.\(^{108}\) Antigens and antibodies associated with hepatitis B virus infection include hepatitis B surface antigen and its corresponding antibody, antibody to hepatitis B core antigen, and hepatitis B e antigen and its corresponding antibody. The most frequently used serologic markers to differentiate acute, resolving and chronic infection are hepatitis B surface antigen, the antibody to hepatitis B core antigen (immunoglobulin M for acute, immunoglobulin G for chronic or resolved) and the antibody to hepatitis B surface antigen; they should be used for initial screening for hepatitis B virus infection.\(^{109}\) The antibody to hepatitis B core antigen is a useful marker in the diagnosis of hepatitis B virus infection, as it appears during the course of acute infection and usually persists for life in the presence of either chronic infection (in which case the person is also positive for hepatitis B surface antigen) or resolved infection, with or without the presence of concomitant antibody to hepatitis B surface antigen.

**Relative benefits and harms of treatment**
Several antiviral agents that suppress chronic hepatitis B virus infection have become available over the past 10 years. Although follow-up with the newer antiviral agents has not been long enough to show improved clinical outcomes, these agents have uniformly been shown to decrease surrogate markers of chronic liver disease (normalization of alanine aminotransferase, decrease in hepatitis B DNA, loss of hepatitis B e antigen and even loss of hepatitis B surface antigen).\(^{108-110}\) However, in a landmark randomized controlled trial of lamivudine versus placebo in patients with advanced liver disease and high-level viral replication, the risk of progressive liver failure (hazard ratio 0.45, \(p = 0.02\)) and of hepatocellular carcinoma (hazard ratio 0.49, \(p = 0.047\)) were both lower in the lamivudine arm than the placebo arm after a mean of 32 months of treatment (Table 6B).\(^{109}\) Lamivudine is well tolerated, but with prolonged use as a single agent, resistance develops progressively. Since publication of that trial, several other antiviral agents (adefovir, entecavir, telbivudine and tenofovir) have become available, all of which are relatively well tolerated and more potent than lamivudine.\(^{109,111-113}\) Treatment of chronic hepatitis B virus infection is rapidly evolving and complex; the decision of who should be treated, when to initiate therapy and which medication to use should be made by professionals with expertise in this area.\(^{111}\)

**Effectiveness of surveillance for hepatocellular carcinoma**
Certain people with chronic hepatitis B virus infection (i.e.,

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**Table 6B: Summary of findings for efficacy of lamivudine to decrease mortality in patients with chronic HBV infection**

<table>
<thead>
<tr>
<th>Patient or population:</th>
<th>“Adults &gt; 16 yr, positive for HBsAg for at least 6 mo, and either positive or negative for HBeAg, HBV DNA ≥ 10⁷ IU/mL at screening, who had a liver biopsy showing an Ishak fibrosis score of at least 4 (where 0 indicates no fibrosis and 6 indicates cirrhosis) at screening or during the previous 2 yr.”(^{111})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Setting:</td>
<td>Multicentre trial with patients from Taiwan, Hong Kong, China, Singapore, Thailand and Australia</td>
</tr>
<tr>
<td>Intervention:</td>
<td>Lamivudine treatment</td>
</tr>
<tr>
<td>Comparison:</td>
<td>Placebo</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Absolute effect</th>
<th>Outcome (period of follow-up)</th>
<th>Risk for control group</th>
<th>Difference with lamivudine (95% CI)</th>
<th>Relative effect (95% CI)</th>
<th>No. of participants (studies)</th>
<th>GRADE quality of evidence</th>
<th>Comments (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Increase in Child–Pugh score (median 32.4 mo)</td>
<td>88 per 1000</td>
<td>54 fewer per 1000 (70 fewer to 23 fewer per 1000)</td>
<td>RR 0.39 (0.20–0.74)</td>
<td>651 (1)</td>
<td>Moderate*</td>
<td>NNT 19 (15–44)</td>
<td></td>
</tr>
<tr>
<td>Hepatocellular carcinoma incidence (median 32.4 mo)</td>
<td>74 per 1000</td>
<td>35 fewer per 1000 (54 fewer to 1 more per 1000)</td>
<td>RR 0.52 (0.27–1.01)</td>
<td>651 (1)</td>
<td>Moderate*</td>
<td>NNT not statistically significant</td>
<td></td>
</tr>
<tr>
<td>Serious adverse event(^{1}) (5 yr)</td>
<td>177 per 1000</td>
<td>53 fewer per 1000 (92 fewer to 5 more per 1000)</td>
<td>RR 0.70 (0.48–1.03)</td>
<td>651 (1)</td>
<td>Moderate*</td>
<td>NNT not statistically significant</td>
<td></td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; HBeAg = hepatitis B e antigen; HBsAg = hepatitis B surface antigen; HBV = hepatitis B virus; NNT = number needed to treat; RR = relative risk.

*Data were analyzed by GlaxoSmithKline; article written by committee including employees.

1Adverse events were considered to be serious if the investigator determined that they jeopardized the patient’s health, were life-threatening or would result in admission to hospital, disability or death.
those with cirrhosis, Asian men > 40 years of age, Asian women > 50 years of age, Africans > 20 years of age and people with a family history of hepatocellular carcinoma) should be screened for hepatocellular carcinoma with ultrasonography and α-fetoprotein serologic testing every six months.172–174 This screening will detect cancer at an earlier stage, when it may be amenable to therapeutic intervention resulting in improved survival (five-year disease-free survival of about 50% v. 0%–10%).162,175 In a randomized controlled trial of screening every six months with ultrasonography and α-fetoprotein testing versus no screening in more than 18 000 Chinese people with positive results for hepatitis B markers, those in the screening arm had a 37% reduction in hepatocellular carcinoma–related mortality after a mean follow-up of five years. This was despite suboptimal adherence rates (< 60%) by the end of the study and represents the minimum benefit that can be expected from surveillance.175

**Effectiveness of screening programs for chronic infection with hepatitis B virus (pregnant women)**

Screening for chronic infection with hepatitis B is routinely recommended for pregnant women in Canada and the United States.111,176,177 There is good evidence that screening and giving immunoprophylaxis to infants born to mothers with chronic hepatitis B virus infection markedly decreases transmission to the newborn. In a recent meta-analysis, the risk of transmission of hepatitis B virus was 92% lower (95% confidence interval [CI] 83%–97%) among infants born to mothers with chronic hepatitis B virus infection who received hepatitis B vaccine and hepatitis B immunoglobulin within 12 hours of birth relative to those whose mothers received placebo.178

**Does vaccination decrease morbidity and mortality from hepatitis B?**

Hepatitis B vaccine (which is about 88% effective in preventing transmission) has been available in Canada since 1982, and a universal childhood hepatitis vaccination program has been recommended and operating in most provinces since 1991.111,177,179 It is an effective vaccine that substantially decreases the risk of acute or chronic hepatitis B virus infection and hepatocellular carcinoma. It is well tolerated, with only mild and transient adverse events. Additionally, several studies have demonstrated no link between hepatitis B vaccine and multiple sclerosis or other neurologic or rheumatologic disorders.109,111,177,179

**Effectiveness of hepatitis B vaccination to decrease morbidity and mortality**

**Children**

Several cohort studies in settings where the incidence of hepatitis B is high have shown the effectiveness of universal infant and childhood vaccination programs in decreasing the incidence of acute hepatitis B, of chronic infection with hepatitis B virus and of hepatocellular carcinoma. The incidence of acute hepatitis B virus infection decreased by 93.5%, to 1/15th the initial rate (i.e., from 215 to 14 per 100 000 population) over a four-year period in an Alaskan Native population after 90% vaccine coverage rate of the whole population.180 The relative risk of chronic infection with hepatitis B virus among vaccinated preschoolers relative to historical school-aged controls ranged from 0.1 to 0.34 in four South Pacific islands.181 Finally, after 15 years of follow-up, a universal infant and childhood vaccination program in Taiwan showed a dramatic decrease in seropositivity for hepatitis B surface antigen (from 9.8% to 0.7%) and a 49% decrease in incidence of hepatocellular carcinoma, to just over half the initial rate (from 0.7 to 0.36 per 100 000 population) (Table 6C).182–184

**Adults**

Hepatitis B vaccination in adults is also effective in decreasing acquisition of acute hepatitis B. A recent meta-analysis of the effectiveness of hepatitis B vaccination among health care workers (in countries where the incidence of hepatitis B virus infection is low) showed that vaccination decreased the acquisition of acute infection by 68% (95% CI 35%–84%) relative to those who were unvaccinated.185 Because in adults it is uncommon for chronic infection with hepatitis B virus to develop following acute infection (< 5%), we were unable to identify any studies that measured the effectiveness of vaccine to decrease morbidity or mortality associated with chronic infection with hepatitis B virus in adults.

**Prevaccination screening for prior immunity**

From 50% to 80% of adults from countries where hepatitis B is highly endemic (prevalence of hepatitis B surface antigen ≥ 8%) and 20% to 30% of adults from countries where hepatitis B is moderately endemic (prevalence of hepatitis B surface antigen 2%–7%) will have serologic evidence of prior infection with and thus immunity to hepatitis B virus (Table 6B). Cost-effectiveness studies have shown that the “breakpoint seroprevalence” of prior hepatitis B infection, above which prevaccination screening for prior immunity is worth doing, ranges from 17% to 35%.186,187 Therefore, it would likely be cost-effective to do prevaccination screening for prior infection among adult immigrants originating from countries that are moderately to highly endemic for hepatitis B.188 In two studies of refugee children with mean ages of 7 and 10 years, respectively, from several different world regions (Africa, former Soviet Union and Yugoslavia, Asia), the seroprevalence of hepatitis B surface antigen was 6.5% and 4%, respectively, and the prevalence of prior infection was 30% and 21%, respectively.189,190 Although these data might not be representative of all immigrant children, they suggest that it would also be cost-effective to do serologic testing for prior infection before vaccinating children.

**Clinical considerations**

**What are the potential implementation issues?**

Studies in immigrant populations have shown that they have relatively little knowledge (40%–60%) of the importance of hepatitis B virus infection and its long-term consequences.
Table 6C: Summary of findings for efficacy of universal childhood vaccination to decrease chronic HBV infection

**Patient or population:** Taiwanese children < 15 years old, born after HBV vaccination program was implemented, and adolescents and young adults > 15–20 years old, born before HBV vaccination program began

**Setting:** Chronic infection with HBV measured before and after implementation of vaccination program in Taiwan

**Intervention:** Perinatal and childhood HBV vaccination

**Comparison:** No HBV vaccination


<table>
<thead>
<tr>
<th>Outcome</th>
<th>Risk in control group, 1984</th>
<th>Difference with vaccination, 1999 (95% CI)</th>
<th>Relative effect (95% CI)</th>
<th>No. of participants (studies)</th>
<th>GRADE quality of evidence</th>
<th>Comments (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HBSAg carriage, &lt; 15-year-olds</td>
<td>98 per 1000</td>
<td>91 fewer per 1000 (94 to 85 fewer per 1000)</td>
<td>RR 0.07 (0.04–0.13)†</td>
<td>2557 (1)</td>
<td>High†</td>
<td>NNT 11 (11–12)</td>
</tr>
<tr>
<td>Anti-HBc, &lt; 15-year-olds</td>
<td>262 per 1000</td>
<td>233 fewer per 1000 (241 to 220 fewer per 1000)</td>
<td>RR 0.11 (0.08–0.16)†</td>
<td>2557 (1)</td>
<td>High†</td>
<td>NNT 5 (5–5)</td>
</tr>
</tbody>
</table>

Note: anti-HBc = hepatitis B core antibody; CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; HBSAg = hepatitis B surface antigen; HBV = hepatitis B virus; NNT = number needed to treat; RR = relative risk.

‡Taiwan’s mass-vaccination program against HBV was launched in July 1984.‡‡‡ For first 2 years, program covered only neonates born to mothers who were HBSAg carriers, but it was extended to all neonates in July 1986, to preschool children in July 1987, to primary-school children in 1988, to middle-school children in 1989, and to adults in 1990.†

†Because of uncertainty regarding the sampling method, it is unclear to which populations these results can be generalized.

‡Upgraded two levels for large effect sizes.

Low proportions (< 50%) of at-risk immigrants have been screened for hepatitis B.191,192 Similarly, low proportions of immigrants are vaccinated (24%–76%) against hepatitis B, and vaccination rates are lower among adults than among children.191-193 Vaccination rates can be enhanced by reducing language and cultural barriers, educating immigrant and refugee populations and improving social supports for using reminder and recall interventions (these being most effective but also most costly).155,193,194 These barriers and limitations need to be considered to improve uptake of screening and vaccination programs among immigrants. In general, when provided with the proper education and access, immigrants and refugees seem to accept vaccines with very little “anti-vaccination” sentiment.195-199 No data on how acceptance and compliance rates differ between cultures are available for hepatitis B treatment. Cultural and language barriers might need to be overcome to optimize the clinical impact of antiviral therapy and to minimize drug-resistant hepatitis B mutants.

**Recommendations of other groups**

With regard to screening, all pregnant women are screened for chronic hepatitis B virus infection in Canada and the United States to prevent transmission to their neonates.171,179 The US Centers for Disease Control and Prevention have also recently recommended screening all immigrants originating from countries where hepatitis B is endemic (prevalence of hepatitis B surface antigen ≥ 2%).196

With regard to vaccination, hepatitis B vaccination is routinely given to all children (age differs from province to province) as part of the national immunization program in Canada. There is no routine catch-up vaccine for immigrants; however, it is recommended that children younger than seven years whose families have immigrated to Canada from areas where prevalence of hepatitis B is high be targeted for vaccination.199,111

Our recommendations highlight the need to screen for chronic hepatitis B virus infection and to vaccinate susceptible high-risk immigrant groups.

**Take-home messages**

- Among newly arrived immigrants and refugees, the prevalence of chronic hepatitis B virus infection is about 3%, as compared with 0.5% in the Canadian-born population.
- The mortality rate from chronic viral hepatitis and hepatocellular carcinoma is higher among immigrants than in the Canadian-born population, likely primarily because of greater prevalence of undetected and untreated chronic hepatitis B virus infection.
- Immigrants would benefit from screening for chronic hepatitis B virus infection and vaccination against hepatitis B.

7. Tuberculosis

Tuberculosis is an airborne transmissible disease that causes a substantial burden to patients, their contacts and society. Although tuberculosis is relatively uncommon in Canada (1621 cases, or about 5 per 100,000 population, reported in 2006), it is costly ($58 million in direct costs in Canada in 2004), treatment is lengthy, many patients require admission to hospital, and the mortality rate among patients with tuberculosis is still high (11%).

The foreign-born population bears a disproportionate burden of tuberculosis in Canada: 65% of all cases of active tuberculosis occur in foreign-born patients, although they make up only 20% of the population. This is because most recent immigrants and refugees originate from countries with a high incidence of tuberculosis, and up to half of them harbour latent tuberculosis infection and are at risk for development of active tuberculosis. Successful control of tuberculosis in Canada will depend on decreasing the rates of tuberculosis in the foreign-born population. We conducted a review to quantify the burden of tuberculosis in the migrant population, to identify those at highest risk for development of active tuberculosis, to describe the effectiveness of screening and treatment programs for latent tuberculosis, to identify barriers or challenges to implementation of such programs and to highlight possible interventions to improve these programs. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on preventing tuberculosis are outlined in Box 7A.

**Box 7A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: tuberculosis**

*Children:* Screen children and adolescents < 20 years of age from countries with a high incidence of tuberculosis (smear-positive pulmonary tuberculosis > 15 per 100,000 population) as soon as possible after their arrival in Canada with a tuberculin skin test, and recommend treatment for latent tuberculosis infection if results are positive, after ruling out active tuberculosis.

*Adults:* Screen all refugees between 20 and 50 years of age from countries with a high incidence of tuberculosis as soon as possible after their arrival in Canada with a tuberculin skin test. Screen all other adult immigrants who have risk factors that increase the risk of active tuberculosis by means of a tuberculin skin test, and recommend treatment for latent tuberculosis infection if results are positive, after ruling out active tuberculosis.

**Basis of recommendations**

**Balance of benefits and harms***

The decision about whom to screen and offer treatment for latent tuberculosis is based on the balance between the potential benefit of treatment (decreasing the lifetime risk of active tuberculosis, which is influenced by age, presence of underlying medical conditions and immigration category) versus the potential harm of hepatotoxicity (which increases with age) and the poor effectiveness of isoniazid in many settings because of suboptimal uptake of screening and treatment. For several groups, screening for latent tuberculosis should be routinely performed, and those with positive results should be offered treatment. These groups are children from countries with a high incidence of tuberculosis (number needed to treat [NNT] 20–26, number needed to harm [NNH] 134–268), adults with risk factors for active tuberculosis (NNT 3–20, NNH variable) and refugees < 50 years of age (NNT 15–26, NNH 49). Screening for latent tuberculosis and offering treatment could also be considered for adult refugees 50–65 years of age (NNT 20–51, NNH 9–18) and other adults without underlying medical conditions < 65 years of age if adherence to treatment can be assured and hepatotoxicity carefully monitored to minimize harms. A decision to screen is a decision to offer treatment and to ensure adherence to treatment with appropriate counselling and monitoring.

**Quality of evidence**

High

**Values and preferences**

The committee attributed more value to screening and treating latent tuberculosis infection to prevent active disease in patients and to prevent transmission of active disease and less value to the practitioner burden of screening and counselling.

*Estimated number needed to treat (NNT) and number needed to harm (NNH) are based on the following assumptions: seven years after arrival, the annual risk of active tuberculosis is 0.1%; the relative risk of active tuberculosis is highest upon arrival and decreases with time (relative risk 5.1, compared with 1.4 seven years after arrival); the patient will live to age 80 years; the efficacy of isoniazid is 90%; and adherence is 70%.

**Methods**

We used the 14-step approach developed by the Canadian Collaboration for Immigrant and Refugee Health (summarized in section 3 of this article, above). We considered the epidemiology of the disease in immigrant populations, as well as potential key clinical actions. We searched MEDLINE, Embase, CINAHL, the Cochrane Library and other sources from Jan. 1, 1996, to Jan. 1, 2010. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for tuberculosis, as presented by Greenaway and associates.

**Results**

In the search for systematic reviews and guidelines for immigrants, we found nine documents that met the eligibility criteria: two guidelines from the United States that addressed screening for latent tuberculosis infection in immigrants, three articles that addressed tuberculosis screening issues in foreign-born patients, one review of postlanding surveillance in Canada and three narrative reviews that recommended screening for latent tuberculosis infection in immigrants and refugees. However, none had used a systematic review method.

In the search for systematic reviews and guidelines for research involving tuberculosis in the general population, we
What is the burden of tuberculosis in immigrant populations?

More than one billion people are infected with latent tuberculosis, which results in 9.2 million new active cases and 1.5 million deaths per year (> 95% of these occurring in low- to middle-income countries).199 Canada is a low-incidence country, with an overall rate of active tuberculosis of five cases per 100,000 population.41,200,201 Most of these cases (> 65%) occur in foreign-born patients, among whom the incidence of tuberculosis is 20 times that in the non-Aboriginal Canadian-born population (16 v. 0.8 cases per 100,000 population), but with rates as high as 500 times greater in certain subgroups of immigrants.202-204 In the past 40 years, most new immigrants have originated from high-incidence countries (i.e., > 15 cases of smear-positive pulmonary tuberculosis per 100,000 population), 30%–50% of whom are infected with latent tuberculosis, which has resulted in a reservoir of about 1.5 million people in Canada with latent tuberculosis infection who are at risk for development of active tuberculosis.196

Among people with positive results for tuberculin skin tests who live in a low-incidence country and have no risk factors, the estimated annual probability of development of active tuberculosis is only 0.1% per year. This means that active disease will develop in only 5%–10% of those with latent tuberculosis infection.205 Recent transmission of tuberculosis confers an increased risk of active tuberculosis. The highest risk of active tuberculosis occurs in the first year after exposure and decreases to the baseline risk (0.1% per year) 5 to 10 years after exposure.198,206

The strongest predictors for development of active tuberculosis in immigrant populations are global region of origin, immigration category (e.g., refugee), the presence of underlying medical comorbidity and the time since arrival.

Region of origin

Rates of tuberculosis are highest in immigrant populations that originate from world regions with the highest rates of tuberculosis, such as sub-Saharan Africa and Asia. Immigrants from these regions (which have rates of smear-positive pulmonary tuberculosis of 200 to 300 per 100,000 population) are more likely to be heavily exposed to tuberculosis, to have positive results on tuberculin skin testing and to have been recently exposed to tuberculosis.199,202-204

Immigration category

The risk of active tuberculosis in refugee populations is about double that in other immigrant populations.207-210 This difference is probably due to both a higher prevalence of latent tuberculosis infection and to having lived in crowded conditions that increase the likelihood of recent exposure to tuberculosis.211

Presence of underlying comorbidity

Underlying medical illnesses, especially any conditions that decrease local or systemic immunity, increase the rate of active tuberculosis to varying degrees (Table 7A), with HIV being the strongest risk factor.212 The issue of HIV screening for new immigrants and refugees is discussed in section 8 of this article, below.

Time since arrival

Rates of tuberculosis in immigrant and refugee populations, from all world regions, are highest within the first five years.

<table>
<thead>
<tr>
<th>Variable</th>
<th>RR†</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>High risk (RR &gt; 6‡)</strong></td>
<td></td>
</tr>
<tr>
<td>AIDS</td>
<td>110–170</td>
</tr>
<tr>
<td>HIV infection</td>
<td>10–110</td>
</tr>
<tr>
<td>Transplantation (related to immunosuppressant therapy)</td>
<td>20–74</td>
</tr>
<tr>
<td>Leukemia, lymphoma</td>
<td>1.0–35</td>
</tr>
<tr>
<td>Silicosis</td>
<td>1.5–33</td>
</tr>
<tr>
<td>Chronic renal failure requiring hemodialysis</td>
<td>1.6–25</td>
</tr>
<tr>
<td>Carcinoma of head and neck</td>
<td>16</td>
</tr>
<tr>
<td>Recent tuberculosis infection (≤ 2 yr)</td>
<td>15</td>
</tr>
<tr>
<td>Abnormal results of chest radiography: fibronodular disease</td>
<td>6–19</td>
</tr>
<tr>
<td>Tumour necrosis factor α inhibitors</td>
<td>1.7–9</td>
</tr>
<tr>
<td><strong>Intermediate risk (RR = 3–6‡)</strong></td>
<td></td>
</tr>
<tr>
<td>Treatment with glucocorticoids</td>
<td>4.9</td>
</tr>
<tr>
<td>Diabetes mellitus (all types)</td>
<td>2.0–4.1</td>
</tr>
<tr>
<td>Young when infected (0–4 yr)</td>
<td>2.2–5</td>
</tr>
<tr>
<td><strong>Low-intermediate risk (RR = 1.3–3‡)</strong></td>
<td></td>
</tr>
<tr>
<td>Underweight (&lt; 90% ideal body weight; for most people, body mass index ≤ 20)</td>
<td>1.6–3</td>
</tr>
<tr>
<td>Cigarette smoker (1 pack/day)</td>
<td>2–3</td>
</tr>
<tr>
<td>Abnormal results of chest radiography: granuloma</td>
<td>2</td>
</tr>
<tr>
<td>Refugee</td>
<td>2</td>
</tr>
<tr>
<td><strong>Low risk (RR = 1)§</strong></td>
<td></td>
</tr>
<tr>
<td>Infected person, no known risk factor, normal results of chest radiography (“low-risk reactor”)</td>
<td>1</td>
</tr>
</tbody>
</table>

‡Adapted, with permission, from Greenaway and associates.42
§See Greenaway and associates for sources for these RR values.
†Mean RR for each variable falls in this range.
§Incidence of development of active tuberculosis 0.1%/yr.
after arrival in a low-incidence region but decrease dramatically after the first year after arrival. Rates of active tuberculosis in the immigrant population, relative to five years after arrival, are 5 to 10 times greater in the first year and twofold greater one to four years after arrival.202–204 These higher rates are most likely caused by the effect of recent exposure to tuberculosis before arrival. A practical Web-based tool can be used to help calculate the lifetime risk of active tuberculosis on the basis of these factors (www.tstin3d.com/index.html).21

Do screening and treatment for latent tuberculosis decrease morbidity from active tuberculosis?

Screening tests
The tuberculin skin test and interferon gamma release assays are available for diagnosing latent tuberculosis infection. The sensitivity of these tests is estimated to be 70%–90%, and the specificity for all tests is above 95%, except for the tuberculin skin test in patients vaccinated with bacille Calmette-Guérin (specificity 60%, because of cross-reactivity).213 With tuberculin skin tests, the likelihood of a false-positive result caused by bacille Calmette-Guérin decreases with time since vaccination, but it also depends on the age when the person was vaccinated. In the first 10 years after vaccination, up to 42% of patients vaccinated after two years of age will have positive results on tuberculin skin testing (the rate being lower among those vaccinated as neonates), but data on the rate of decline are conflicting.214–216 Among those receiving a tuberculin skin test more than 10 years after vaccination as a neonate, only 1%–2% of results will be positive, compared with 21% for those vaccinated after two years of age. Interpreting the results of a tuberculin skin test is therefore particularly difficult for

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**Table 7B: Summary of findings for isoniazid to prevent active tuberculosis**

<table>
<thead>
<tr>
<th>Outcome; risk category</th>
<th>Risk for control group</th>
<th>Difference with isoniazid (95% CI)</th>
<th>Relative effect (95% CI)</th>
<th>No. of participants (studies)</th>
<th>GRADE quality of evidence</th>
<th>Comments (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active tuberculosis</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intermediate risk*</td>
<td>17 per 1000</td>
<td>10 fewer per 1000 (12 fewer to 8 fewer per 1000)</td>
<td>RR 0.40 (0.31–0.52)</td>
<td>73 375 (11)</td>
<td>Moderate†§</td>
<td>NNT 99 (86–123)</td>
</tr>
<tr>
<td>Highly compliant (&gt; 80% of doses taken)*</td>
<td>10 per 10 000</td>
<td>7 fewer per 10 000 (8 fewer to 5 fewer per 10 000)</td>
<td>RR 0.20 (0.13–0.31)</td>
<td>15 696 (1)</td>
<td>High</td>
<td>NNT 85 (78–98)</td>
</tr>
<tr>
<td>High risk†</td>
<td>53 per 1000</td>
<td>3 fewer per 1000 (40 fewer to 19 fewer per 1000)</td>
<td>RR 0.40 (0.24–0.65)</td>
<td>1 875 (5)</td>
<td>Moderate‡</td>
<td>NNT 32 (25–54)</td>
</tr>
<tr>
<td>Hepatitis (follow-up 5 yr)**</td>
<td>1 per 1000</td>
<td>5 more per 1000 (2 more to 11 more per 1000)</td>
<td>RR 5.54 (2.56–12)</td>
<td>20 874 (1)</td>
<td>Moderate</td>
<td>NNH 220 (91–642)</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NNH = number needed to harm; NNT = number needed to treat; RR = risk ratio.

*Numbers taken from Smieja et al.202
†In 1999 systematic review of isoniazid for tuberculosis in HIV-positive patients, this was the risk among HIV-positive, tuberculin skin test–positive patients.211
‡Test for heterogeneity, p = 0.02.
§Only one study examined 6 months vs. 12 months of isoniazid therapy; risk of hepatitis and active tuberculosis not significantly different between groups.
¶Downgraded for directness, as the data were from developing countries.
**As reported by the International Union Against Tuberculosis Committee on Prophylaxis.222
children under 10 years of age. All HIV-positive patients should be screened for latent tuberculosis infection with a tuberculin skin test.198

Because there is no gold standard test for the diagnosis of latent tuberculosis infection, assessing and comparing the performance of these tests is challenging, especially when there are discrepancies. The major advantage of the tuberculin skin test is that the risk of active disease for different sizes of induration is well described, whereas very few prospective data exist for interferon gamma release assays. The most recent Canadian guidelines recommend using the tuberculin skin test as the primary screening tool for both adults and children and using interferon gamma release assays sequentially, after tuberculin skin testing, in people with a high likelihood of a false-positive result on the tuberculin skin test (i.e., low risk of tuberculosis infection).209 This recommendation is supported by a recent cost-effectiveness analysis.220 The major limitation of these tests is their inability to distinguish the 10% of people with latent tuberculosis infection in whom active tuberculosis will develop from the 90% in whom the disease will not develop.

All patients for whom the results of a tuberculin skin test are positive should undergo chest radiography to rule out active tuberculosis; they should also be questioned for symptoms of active tuberculosis (chronic cough, weight loss, fever, night sweats). If there is any suspicion of active tuberculosis, three samples of sputum or specimens from other sites (e.g., lymph node, cerebrospinal fluid) should be gathered for smear and culture before treatment for latent tuberculosis infection is started.

Relative benefits and harms of treatment

The efficacy of isoniazid relative to that of placebo in decreasing the likelihood that active tuberculosis will develop in people with latent tuberculosis infection has been established in a Cochrane review of 11 randomized controlled trials (relative risk 0.40, 95% confidence interval 0.31–0.52) (Tables 7B and 7C).213–216 The overall efficacy of isoniazid is 62% after 12 months of treatment, but efficacy increases to 93% among those who adhere to treatment (i.e., take > 80% of their doses).221 Although the efficacy of treatment for 9 versus 12 months has not been directly compared, a recent reanalysis showed that the maximal benefit of isoniazid was achieved at 9 months (Tables 7D).222 A study on the effect of resistance to isoniazid on the efficacy of isoniazid chemoprophylaxis showed that, at a mean prevalence of 7%–10% isoniazid resistance (the level in the immigrant population), isoniazid was the drug of choice and that only at very high rates of isoniazid resistance (> 15%–20%) were other regimens preferred.228,229

Hepatotoxicity is a limitation of isoniazid therapy. It most commonly manifests as a transient, asymptomatic increase in liver function (10%–20%), rarely causes clinical hepatitis (0.5%) (which resolves when treatment with isoniazid is stopped230,231) and very rarely causes fulminant hepatitis and liver failure leading to death or liver transplantation (< 0.01%).230,231 Initial higher overall rates of hepatotoxicity (1%) were reported among adults in the 1970s, but these data were likely confounded by unrecognized underlying cirrhosis. Hepatotoxicity among patients taking isoniazid is greater among those with pre-existing liver disease, alcoholism, concomitant use of hepatotoxic drugs and older age. Although clinical and fulminant hepatitis are rare, they can occur at any age. This possibility underscores the importance of monthly monitoring for all patients and of teaching them to recognize the symptoms of hepatitis (nausea, vomiting, abdominal pain, jaundice) and to stop medication as soon as worrisome symptoms occur.230,231 Adequate time must be taken, through interpreters if necessary, to ensure that all patients are appropri-

|---|---|---|---|---|

<table>
<thead>
<tr>
<th>Absolute effect</th>
<th>Relative effect (95% CI)</th>
<th>No. of participants (studies)</th>
<th>GRADE quality of evidence</th>
<th>Comments (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Risk for control group</td>
<td>Difference with older age (95% CI)</td>
<td>Age 35–49</td>
<td>Age ≥ 50</td>
<td>Age 35–49</td>
</tr>
<tr>
<td><strong>Toxicity:</strong></td>
<td>RR</td>
<td>1982</td>
<td>Very low*</td>
<td>NNH not statistically significant</td>
</tr>
<tr>
<td>elevation of transaminases by &gt; 5 times upper limit of normal</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 per 1000</td>
<td>4 more per 1000 (2 fewer to 21 more per 1000)</td>
<td>15 more per 1000 (2 more to 58 more per 1000)</td>
<td>RR (0.59–6.34)</td>
<td>RR (1.40–15.49)</td>
</tr>
</tbody>
</table>
| Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NNH = number needed to harm; RR = risk ratio. *Fewer than 300 events.
ately informed and understand the risks and benefits of isoniazid, and they must be given a clear description of what to do if symptoms arise.

Clinical considerations

What should be considered in screening and treatment?

Children

Children less than 11 years of age do not undergo prearrival radiographic screening. For this and several other reasons, they could benefit greatly from screening for latent tuberculosis infection. Children face many years of life in which active tuberculosis could develop, and they have a relatively low potential for hepatotoxicity. When active tuberculosis develops in children, it is often difficult to diagnose because it is more often paucibacillary or extrapulmonary, and young children (especially those younger than five years of age) are more likely susceptible to severe or rapidly progressive disease.

Refugees

Refugee populations have consistently had about a two-fold greater risk of active tuberculosis than the immigrant population, at least within the first year after arrival. A higher prevalence of latent tuberculosis infection in the refugee population and having lived in crowded conditions, which increase the likelihood of recent exposure to tuberculosis, are contributing factors.

What are the potential implementation issues?

Barriers to the uptake of screening and completion of treatment for latent tuberculosis infection include a combination of patient, provider and institutional factors. Patient-related barriers include the stigma of tuberculosis and its association with HIV, linguistic barriers and difficulties getting to appointments because of inconvenient clinic locations or limited clinic hours. Provider-related barriers to offering screening to migrants are related to inadequate knowledge of which migrants should be screened and how they should be followed. Low adherence to treatment for latent tuberculosis infection is associated with barriers similar to those for screening for latent tuberculosis. These barriers include linguistic barriers, cultural taboos and stigmatization, low education level, perceived low risk of progression from latent tuberculosis infection to active disease, belief that positive results from tuberculin skin tests are due to bacille Calmette-Guérin, not wanting to undergo venipuncture and economic factors (costs of travel, lack of insurance, delays in obtaining insurance, missed days at work).

Increased adherence to tuberculin skin test screening has been achieved with reminders to patients (e.g., letters, phone calls), education of patients and physicians, and novel strat-

<table>
<thead>
<tr>
<th>Age, yr</th>
<th>Time lived in Canada, yr</th>
<th>Cumulative lifetime risk of active tuberculosis, % †‡</th>
<th>Course of isoniazid completed§; NNT</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>100%</td>
</tr>
<tr>
<td>10</td>
<td>0</td>
<td>8.1</td>
<td>14</td>
</tr>
<tr>
<td>2</td>
<td>0.5</td>
<td>7.5</td>
<td>15</td>
</tr>
<tr>
<td>5</td>
<td></td>
<td>7.1</td>
<td>16</td>
</tr>
<tr>
<td>20</td>
<td>2.5</td>
<td>7.1</td>
<td>16</td>
</tr>
<tr>
<td>5</td>
<td></td>
<td>6.5</td>
<td>18</td>
</tr>
<tr>
<td>35</td>
<td>0</td>
<td>5.6</td>
<td>18</td>
</tr>
<tr>
<td>2</td>
<td></td>
<td>5.0</td>
<td>20</td>
</tr>
<tr>
<td>5</td>
<td></td>
<td>4.6</td>
<td>23</td>
</tr>
<tr>
<td>50</td>
<td>0</td>
<td>4.1</td>
<td>24</td>
</tr>
<tr>
<td>2</td>
<td></td>
<td>3.5</td>
<td>28</td>
</tr>
<tr>
<td>5</td>
<td></td>
<td>3.1</td>
<td>32</td>
</tr>
<tr>
<td>65</td>
<td>0</td>
<td>2.6</td>
<td>36</td>
</tr>
<tr>
<td>2</td>
<td></td>
<td>2.0</td>
<td>44</td>
</tr>
<tr>
<td>5</td>
<td></td>
<td>1.6</td>
<td>57</td>
</tr>
</tbody>
</table>

Note: NNT = number needed to treat.

*Adapted, with permission, from Greenaway and associates.
†Assume 0.1% annual risk of infection, with a relative risk of development of active tuberculosis for each year since arrival of 5.08 for < 1 year, 2.96 for 1.1–2 years, 2.35 for 2.1–3 years, 2.06 for 3.1–4 years, 1.87 for 4.1–5 years, 1.89 for 5.1–6 years, and 1.36 for 6.1–7 years (all v. > 7 years).
‡Assume individuals live to age 80 years.
§For 100% completion, assume 90% efficacy of isoniazid.
egies, such as drive-by tuberculin skin test readings for taxi drivers. In one study, educating primary care providers about how and whom to screen for tuberculosis not only increased screening and identification of people with latent tuberculosis, but also increased identification of those with active tuberculosis. In this randomized clinical trial, screening rates were 0.4% in the nonintervention group and 57% in the intervention groups, and identification of both those with latent tuberculosis infection (9% to 19%) and those with active tuberculosis (34% v. 47%) was higher in the intervention group. Strategies that have increased adherence to treatment for latent tuberculosis infection in immigrant and refugee populations include patient reminders (calendar stickers for self-monitoring, phone calls, letters and directly observed therapy), adherence coaches who speak the same language as the patient, ongoing education of patients and providers, and cultural case management. Goldberg and colleagues found that when case managers were matched to the ethnic and linguistic background of patients and provided treatment for latent tuberculosis and monitoring during monthly home visits, adherence with treatment improved substantially over standard clinic-based management before the intervention (82% v. 37%).

Recommendations of other groups

The Canadian Tuberculosis Committee, the Canadian Thoracic Society and the Canadian Paediatric Society recommend using a tuberculin skin test to screen the following groups from countries with a high incidence of tuberculosis: children younger than 15 years living in Canada for less than two years and people 15 years of age or older with factors increasing the risk of active tuberculosis or within two years after arrival if they have had known contact with tuberculosis (Table 7A). Our recommendations highlight the importance of screening for latent tuberculosis with Mantoux tests in high-risk immigrant groups.

Take-home messages

- Foreign-born people account for 65% of all those with active tuberculosis in Canada, and subgroups have up to a 500-fold greater risk of active tuberculosis relative to the non-Aboriginal Canadian-born population.
- The Canadian Collaboration for Immigrant and Refugee Health recommends screening certain groups as soon as possible after arrival in Canada, with a tuberculin skin test, and initiating treatment for latent tuberculosis infection in those with a positive result, after ruling out active tuberculosis.
- Although isoniazid is highly efficacious in decreasing the development of active tuberculosis in those with latent tuberculosis infection, monitoring for hepatotoxicity is required for patients of all ages.
- Close monitoring is required for those over 50 years of age and those with pre-existing liver disease, alcoholism or concomitant use of hepatotoxic drugs.
- Adherence to screening and treatment for latent tuberculosis infection can be increased if delivered in a culturally sensitive manner.

8. HIV

An estimated 33.2 million women, men and children are infected with HIV worldwide.241 In Canada, populations at high risk for HIV include women and men from countries where HIV is endemic,242 who are at risk of heterosexual transmission and mother-to-child transmission.243 Although HIV testing is now part of the immigration medical examination, persons with HIV are not necessarily excluded from immigrating. Delayed disclosure of positive results by patients to their partners and practitioners, as well as the ongoing risk of HIV transmission attributable to a variety of factors (such as travel to the country of origin, where HIV could be endemic244), remains a concern. We conducted an evidence review to guide primary care practitioners in the early detection, prevention and treatment of HIV for newly arriving immigrants. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on HIV screening and treatment are outlined in Box 8A.

Methods

We used the 14-step method developed by the Canadian Collaboration for Immigrant and Refugee Health16 (summarized in section 3 of this article, above). We considered the epidemiology of HIV in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE, Embase, CINAHL, the Cochrane Library and other sources from Jan. 1, 1995, to Jan. 1, 2010. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for HIV (Appendix 6, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

Box 8A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: HIV

Screen for HIV, with informed consent, all adolescents and adults from countries where HIV is prevalent (> 1%).

Basis of recommendations

Balance of benefits and harms

The decision to screen men and women for HIV is based on a dramatic reduction in mortality with treatment, e.g., with a combination of three versus two antiretrovirals (number needed to treat [NNT] 132, 95% confidence interval [CI] 91–357) and reduction of high-risk behaviour (NNT 5, 95% CI 4–7). Prevalence of HIV infection is higher among immigrants from countries where HIV is prevalent (> 1%) than among other Canadians (< 0.18%). Harms included adverse drug reactions requiring change in regimen. Data on harms related to anxiety and possible discrimination related to HIV status are unavailable.

Quality of evidence

Moderate

Values and preferences

The committee attributed more value to identifying HIV-positive women and men for appropriate treatment, support and prevention and less value to uncertain risk of couple discord and risk of discrimination and less concern for burden of testing with informed consent.

Results

We initially identified eight papers, including a systematic review of HIV prevalence studies among sub-Saharan African refugees245 and a systematic review of retention of African patients in antiretroviral treatment programs.246 Although these papers detailed the complexities of HIV issues among immigrants, highlighting the need for sensitivity to sociocultural context in the interventions, none provided evidence of the benefits and harms of screening. Without restricting the search to papers specific to immigrants, we identified 13 systematic reviews and guideline articles providing evidence on screening for HIV.247−259 We identified and appraised longitudinal studies of antiretroviral treatment,260,261 a meta-analysis on behaviour change related to HIV screening tests,262 a meta-analysis on antiretroviral therapy263 and a Cochrane systematic review on antiretroviral treatment.264 One hundred and four titles addressed the burden of HIV and barriers to care for immigrant populations.

What is the burden of HIV in immigrant populations?

The HIV infection rate is about 12.6 times higher among immigrants and refugees from countries where HIV is endemic than it is in the Canadian-born population,265 accounting for 7% of HIV cases in large urban centres.10 In subgroup mortality analyses of the Canadian Mortality Database, both male and female immigrants from the Caribbean had higher mortality from HIV infection (standardized mortality rates 4.2 for males and 27.4 for females). In a study based on screening of immigration applicants to Canada, 70% of those who tested positive were refugees or refugee claimants from regions where HIV is endemic (Table 8A), and the seropositivity among applicants from sub-Saharan Africa was above 3%.266

Table 8A: Prevalence of HIV by region in 2007*

<table>
<thead>
<tr>
<th>Region</th>
<th>Prevalence of HIV, % (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sub-Saharan Africa</td>
<td>5.0 (4.6–5.5)</td>
</tr>
<tr>
<td>Caribbean</td>
<td>1.0 (0.9–1.2)</td>
</tr>
<tr>
<td>Eastern Europe and central Asia</td>
<td>0.9 (0.7–1.2)</td>
</tr>
<tr>
<td>North America</td>
<td>0.6 (0.5–0.9)</td>
</tr>
<tr>
<td>Latin America</td>
<td>0.5 (0.4–0.6)</td>
</tr>
<tr>
<td>Oceania</td>
<td>0.4 (0.3–0.7)</td>
</tr>
<tr>
<td>Middle East and north Africa</td>
<td>0.3 (0.2–0.4)</td>
</tr>
<tr>
<td>Southeast Asia</td>
<td>0.3 (0.2–0.4)</td>
</tr>
<tr>
<td>Western and central Europe</td>
<td>0.3 (0.2–0.4)</td>
</tr>
<tr>
<td>East Asia</td>
<td>0.1 (&lt; 0.2)</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval.

*Adapted, with permission, from Joint United Nations Programme on HIV/AIDS (UNAIDS).241
Women 20–39 years of age accounted for 66% of positive HIV test reports among adult immigrant women in Canada in 2006. Most of these women had been exposed to HIV in areas where infection was endemic. Sexual transmission of HIV from a man to a woman is two to eight times more likely than transmission from a woman to a man. A woman’s susceptibility to HIV infection is further increased if she or her partner has a sexually transmitted infection, if she has experienced genital trauma or if her partner is HIV-positive with a high viral load. Determinants of vulnerability to HIV include increased mobility (e.g., 1.5–1.8 times higher for mobile populations in South Africa), experience of war-related violence (e.g., 10%–12% of women experiencing sexual violence in the Democratic Republic of Congo contract HIV), certain sexual practices (studies suggest a twofold risk of HIV with vaginal douching), limited knowledge about HIV and AIDS, and limited language proficiency.

Does screening for HIV decrease related morbidity and mortality?

Screening tests
Most laboratories in Canada use a two-step testing strategy for HIV. Enzyme-linked immunosorbent assays are sensitive (≥99%) and specific (≥99%) for both HIV-1 and HIV-2 after about three weeks of infection. Risk-reduction counselling (e.g., regarding condom use) has been shown to be most effective when targeted to HIV-positive or high-risk women and men.

Relative benefits and harms of treatment
In a meta-analysis of 27 studies on behaviour change in HIV-positive women and men, Weinhardt and associates found decreases in high-risk sexual behaviour (including unprotected intercourse) with provision of counselling and screening (Table 8B). We were unable to find quantitative data on anxiety or depression, changes in relationships with sexual partners or discrimination associated with false-positive results.

A meta-analysis by Enanoria and colleagues comparing three-drug and two-drug antiretroviral treatment showed a significant decrease in mortality (risk ratio 0.62, 95% confidence interval 0.45–0.86) (Table 8C). In another meta-analysis, Siegfried and coauthors reported a relative risk of 1.3 for adverse events associated with antiretroviral treatment. However, most adverse events, including metabolic disturbances associated with cardiovascular events, can be ameliorated by changes in regimen or appropriate treatment. Evidence for a greater than 80% reduction in HIV transmission among HIV-discordant couples receiving antiretroviral treatment has been obtained from Spain, Thailand and Uganda. Combination antiretroviral therapy is associated with dramatic decreases in mortality, regardless of sex, race, age and risk factors for transmission. Recent studies have reported that voluntary counselling and testing for HIV are cost-effective in populations with HIV prevalence above 0.1%.

Clinical considerations
What are the potential implementation issues?
Refugees and refugee claimants may delay testing and/or

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**Table 8B:** Summary of findings on how pretest counselling and screening affect high-risk behaviours

<table>
<thead>
<tr>
<th>Patient or population:</th>
<th>HIV-positive patients, mostly men</th>
</tr>
</thead>
<tbody>
<tr>
<td>Setting:</td>
<td>United States</td>
</tr>
<tr>
<td>Intervention:</td>
<td>Counselling and screening tests</td>
</tr>
<tr>
<td>Comparison:</td>
<td>No screening procedures</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Risk for control group</th>
<th>Difference with counselling</th>
<th>Relative effect* (95% CI)</th>
<th>No. of participants (studies)</th>
<th>GRADE quality of evidence</th>
<th>Comments (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unprotected intercourse</td>
<td></td>
<td>NA</td>
<td>d+ 0.47 (0.32–0.61)</td>
<td>402 (5)</td>
<td>Very low</td>
<td>NNT 5 (4–7)†</td>
</tr>
<tr>
<td>HIV-positive</td>
<td>No baseline data available</td>
<td>NA</td>
<td>d+ 0.19 (0.08–0.31)</td>
<td>599 (7)</td>
<td>Very low</td>
<td>NA</td>
</tr>
<tr>
<td>HIV-negative</td>
<td>No baseline data available</td>
<td>NA</td>
<td>d+ 0.75 (0.59–0.92)</td>
<td>293 (2)</td>
<td>Very low</td>
<td>NA</td>
</tr>
<tr>
<td>Discordant couples</td>
<td>No baseline data available</td>
<td>NA</td>
<td>d+ 0.65 (0.42–0.87)</td>
<td>160 (4)</td>
<td>Very low</td>
<td>NA</td>
</tr>
<tr>
<td>Condom use, HIV-positive</td>
<td>No baseline data available</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Adverse effects</td>
<td>No numeric data available‡</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td></td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; d+ = standardized mean difference index; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NA = not applicable; NNT = number needed to treat.

*The value for d+, the standardized mean difference index, is computed by behaviour before and after HIV counselling and testing. An effect size of 0.20 is considered small and 0.5 is considered medium. Positive effect sizes indicate a reduction in high-risk sexual behaviour.

†We are very uncertain about the estimate.

‡Qualitative studies have highlighted various issues but have also suggested that immigrants favour increased access to testing.
treatment because they fear failing to obtain legal immigration status if HIV-positive.\textsuperscript{44} Immigrants suffering from post-traumatic stress or depression could require additional social support and reassurance on many fronts before finding the treatment for HIV infection acceptable. Many HIV-positive immigrants adopt codes of silence, which precludes providing sex education to their children.\textsuperscript{41,42} This highlights a crucial role for primary care practitioners in providing education about sex and HIV to youth.

Limited education, lower health literacy, linguistic barriers, traditional and religious beliefs,\textsuperscript{45} psychological issues that stem from HIV-related stigma, practitioners’ reluctance to offer screening tests and perceptions of low risk among women and men suffering from HIV have all been identified as factors impeding HIV testing.\textsuperscript{27} HIV-related stigma is magnified by structural inequities, sex roles, negative attitudes and discrimination by health care providers, and social attitudes in general.\textsuperscript{27} Strategies to improve access to care include sensitivity to emotional and trauma issues, provision of comprehensive and holistic care, presence of HIV-positive peer educators and integration of HIV information within settlement services.\textsuperscript{27} Qualitative studies\textsuperscript{23} have shown that most immigrant women support HIV screening procedures as part of routine medical care.

Recommendations of other groups

The US Preventive Services Task Force\textsuperscript{44} recommends that clinicians screen all pregnant women, as well as all adolescents and adults at increased risk for HIV (e.g., from countries with HIV prevalence > 1%). The Centers for Disease Control and Prevention\textsuperscript{29} recommend screening tests for all sexually active adolescents and adults (15–64 years), citing substantial individual and population benefits of early detection. The UK guidelines\textsuperscript{35} recommend HIV testing for all who present to genitourinary clinics. Our recommendations highlight the importance of routine HIV screening for high-risk immigrant groups.

Take-home messages

- Immigrants and refugees from countries where HIV is prevalent (> 1%) are vulnerable because of the high HIV prevalence in their home countries and because of high levels of HIV-related stigma.
- Stigma and discrimination related to HIV are associated with avoidance of or delays in seeking HIV testing, delays in disclosure of seropositive status to partners and practitioners, and postponement or rejection of treatment.
- The effects of HIV-related stigma are magnified among socially vulnerable minority groups.
- Providing information about HIV testing options and effectiveness of treatment can improve the likelihood of testing and acceptance of care.

9. Hepatitis C

Chronic infection with hepatitis C virus affects an estimated 170 million people worldwide. The chronic form of infection develops in about 70% of those who have had acute infection. Cirrhosis develops in 20% of individuals with chronic infection and hepatocellular carcinoma in 1%–5% during the two decades following the initial infection. Canada has a low burden of hepatitis C, with an estimated seroprevalence of about 0.8%. The majority of chronic hepatitis C virus infections in Canada occur in individuals who have previously or are currently injecting drugs. However, an estimated 20% of cases occur in immigrants, and immigrant populations have increased mortality due to viral hepatitis. There is no effective vaccination to prevent acquisition of hepatitis C virus infection, but the virus can be detected, and treatment regimens are moderately successful in eradicating chronic infections. The treatment regimen averages 24–48 weeks and is often difficult to tolerate.

Emerging data on combination therapies with protease inhibitors have shown substantially improved efficacy with shorter duration of treatments. We conducted an evidence review to determine the benefits and harms of screening for hepatitis C virus infection in immigrant populations. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on screening for hepatitis C are outlined in Box 9A.

Methods

We used the 14-step method developed by the Canadian Collaboration for Immigrant and Refugee Health (summarized in section 3 of this article, above). We considered the epidemiology of hepatitis C in immigrant populations and defined clinical preventive actions (intervention), outcomes and key clinical questions. We searched MEDLINE (Ovid), MEDLINE InProcess, Embase, CINAHL and the Cochrane Library from 1950 to Jan. 28, 2010, for studies pertinent to immigrants and from Jan. 1, 1997, to Jan. 28, 2010, for studies pertinent to the general population. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for hepatitis C (Appendix 7, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

Results

In the search for systematic reviews and guidelines for hepatitis C in immigrants, we identified 31 records, but none met our eligibility criteria. In the search for systematic reviews and guidelines for hepatitis C in the general population, we identified 4714 articles, of which 24 met the eligibility criteria. In addition, a search for articles relevant to hepatitis C and immigrants yielded 250 articles, of which 50 were relevant and addressed the following areas: epidemiology and knowledge of and compliance with screening and treatment in the immigrant population.

What is the burden of hepatitis C in immigrant populations?

Canada is a low-prevalence country for chronic infection with hepatitis C virus. In Canada, deaths from non-A, non-B hepatitis (the majority of which is presumed to be due to chronic hepatitis C virus infection) have increased 3.4-fold (from 0.12 to 0.41 per 100 000) over the past 30 years, and the incidence of infection has increased 2.2-fold during the same period. These increases are thought to be due to the uncontrolled epidemic among injection drug users and the importation of virus by immigrants from countries where chronic hepatitis C virus infection is endemic. The seroprevalence of chronic infection in the immigrant population is estimated to be about 3% (range 0.1%–18% for various global regions of origin) and likely reflects rates in the countries of origin (Figure 9A). According to a recent Canadian study, mortality from viral hepatitis was 1.8- to 3.8-fold greater and mortality from hepatocellular carcinomas was 2.2- to 4.9-fold greater among immigrants than in the Canadian-born population.

It is unclear what proportion of deaths due to viral hepatitis and hepatocellular carcinomas...
are attributable to chronic infection with hepatitis C virus in the immigrant population, but it may be as high as 30%. This figure is extrapolated from the fact that hepatitis B and hepatitis C account for 80%–90% of all hepatocellular carcinomas worldwide and that 30% of all cases are attributable to chronic hepatitis C virus infection.276

The primary mode of transmission of hepatitis C virus is percutaneous; sexual or perinatal transmission is much less frequent. As a result, injection drug users represent the group at greatest risk for hepatitis C in Canada. Immigrants are an important unrecognized risk group for chronic hepatitis C infection in Canada, but they are likely to have acquired their infection through unsafe health care–related injections or through other medical equipment, unscreened blood products or surgical procedures in their countries of origin. It is estimated that up to 40% of chronic infections globally are acquired through unsafe injections.278 The proportion of immigrants ineligible for treatment because of comorbidities such as psychiatric illness, substance abuse or medical conditions may therefore not be as high as for other at-risk populations such as injection drug users. More than 70% of the chronic infections in North America are due to genotype 1, whereas in certain countries elsewhere in the world, other genotypes may predominate (e.g., genotype 4 in Egypt and genotype 3 in Pakistan). Immigrants are more likely to have concurrent chronic hepatitis B virus infection or HIV infection, and co-infection will increase the risk and rate of liver fibrosis associated with chronic hepatitis C virus infection.276

Does screening for hepatitis C virus decrease morbidity?

Screening tests

Widely available third-generation serologic enzyme immunoassays to detect anti–hepatitis C virus antibodies are highly sensitive (97%) and specific (99%). False-positive results occur in populations where the prevalence of hepatitis C is low, and false-negative results may occur in the setting of severe immunosuppression, such as in patients with HIV, hypogammaglobulinemia or agammaglobulinemia; those who have undergone solid organ transplantation; or those receiving hemodialysis.278 If the result is positive, a nucleic acid test to detect hepatitis C RNA (qualitative or quantitative) should be performed to confirm the presence of circulating virus.

Relative benefits and harms of treatment

The current standard for treatment is combination therapy with pegylated interferon and ribavirin, which achieves an overall sustained virologic response of about 50% in all patients278,279–280 (Table 9A).290,291 Sustained virologic response has been associated with improved clinical outcomes in patients with cirrhosis due to chronic hepatitis C virus infection. In this population, those who did not achieve a sustained virologic response had higher rates of hepatocellular carcinoma (hazard ratio [HR] 2.59, 95% confidence interval [CI] 1.13–5.97) and liver-related mortality (HR 6.97, CI 1.71–28.42) than those who achieved sustained virologic response.292–294 The response rate, dosage of the medications and duration of treatment (ranging from 24 to 48 months) are determined by the viral genotype. For infection with genotype 1 virus, treatment response ranges from 42% to 46% and requires 48 weeks of treatment. The response is better for those with genotype 2 (74%) and genotype 3 (69%), and therapy for these genotypes is of shorter duration (usually 24 weeks).278,284–286 For other genotypes (4, 5, 6, 7, 8), the results are less well defined: they appear to be better than for genotype 1 but not as good as for genotypes 2 and 3.295,296 The treatment is associated with numerous adverse effects, and 10% to 14% of patients discontinue therapy because of an

Figure 9A: Estimated worldwide prevalence of hepatitis C, 1999. Based on data from the World Health Organization.284
Guidelines

adverse event, most commonly psychiatric symptoms or severe anemia.\textsuperscript{278,299} Persons with advanced cirrhosis are less likely to have a response to treatment, which highlights the importance of initiating treatment before advanced liver disease develops.\textsuperscript{300,301} Recent results for combination regimens with protease inhibitors and standard therapy have shown substantially improved efficacy (70% v. 50%) for those with genotype 1, as well as shorter treatment durations.\textsuperscript{270,302} These new treatment options will likely change the standard of therapy in the near future, making screening and appropriately timed treatment an important strategy for controlling the burden of chronic hepatitis C virus infection. Management during therapy often requires a multidisciplinary approach, and all patients found to be positive should be referred to a health care professional with expertise in managing hepatitis C virus infection.

In most studies, screening has been found to be cost-effective only when the prevalence of hepatitis C is high (> 10%) because of the relatively poor efficacy of treatment (overall 50%), the high adverse-effect profile and the high proportion of persons ineligible for treatment because of underlying comorbidities such as substance abuse, psychiatric illness or medical diseases (37%).\textsuperscript{303} Plunkett and Grobman\textsuperscript{304} found that screening pregnant women was not cost-effective. They assumed a 1% seroprevalence of hepatitis C virus and 48 weeks of treatment with pegylated interferon and ribavirin, but they calculated only direct costs. Singer and Younossi\textsuperscript{305} also found that screening for hepatitis C virus in the general US population was not cost-effective. They assumed that the seroprevalence was 3%, that only 20% of individuals found to be positive would be given treatment (because of underlying comorbidities), that the response rate (to interferon and ribavirin) in patients with genotype 1 would be 37% and that 72% of all patients would have genotype 1. In a sensitivity analysis, however, they found that if 50% of individuals positive for hepatitis C virus started treatment, then screening would be cost-effective at a seroprevalence of 3%.\textsuperscript{304,305} Certain ethnic groups, such as Southeast Asians, have a better response to therapy than other ethnic groups.\textsuperscript{306}

In summary, we recommend routine screening for immigrant populations in whom the estimated prevalence of chronic hepatitis C virus infection is 3% or higher, given their increased risk of death from viral hepatitis and hepatocellular carcinoma, the more favourable response to treatment before development of cirrhosis (which favours early detection), the likelihood that immigrants are more likely to be eligible for treatment and the likelihood that certain immigrant groups can be expected to respond more favourably to treatment than the general Canadian population.

Clinical considerations

What are the potential implementation issues?
Few studies have evaluated immigrant populations’ knowledge of the importance of hepatitis C, its consequences and the risk factors for transmission. The data suggest that knowledge of hepatitis C is generally low and that it is significantly lower for immigrants than for nonimmigrants.\textsuperscript{306,307} Predictors of better knowledge are higher levels of education, employment and being highly acculturated.\textsuperscript{307} We found no data on the

Table 9A: Summary of findings for comparison of pegylated interferon plus ribavirin compared with interferon plus ribavirin for treatment of chronic hepatitis C virus infection

<table>
<thead>
<tr>
<th>Patient or population: Patients with chronic hepatitis C virus infection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Settings: Multiple countries (including Italy, Egypt, Japan, Taiwan, Germany, Saudi Arabia, Belgium)</td>
</tr>
<tr>
<td>Intervention: Pegylated interferon plus ribavirin</td>
</tr>
<tr>
<td>Comparison: Interferon plus ribavirin</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Absolute effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Outcome</td>
</tr>
<tr>
<td>Risk with interferon plus ribavirin</td>
</tr>
<tr>
<td>Difference with pegylated interferon plus ribavirin (95% CI)</td>
</tr>
<tr>
<td>Relative effect (95% CI)</td>
</tr>
<tr>
<td>No. of participants (studies)</td>
</tr>
<tr>
<td>GRADE quality of evidence</td>
</tr>
<tr>
<td>Comments (95% CI)</td>
</tr>
</tbody>
</table>

| Sustained virologic response\textsuperscript{301} | 489 per 1000 | 147 fewer per 1000 (181 to 108 fewer per 1000) | RR 0.70 (0.63–0.78) | 1878 (2) | High | NNT 7 (6–10) |
| Sustained virologic response\textsuperscript{300} | 617 per 1000 | 123 fewer per 1000 (160 to 74 fewer per 1000) | RR 0.80 (0.74–0.88) | 4659 (16) | Low*† | NNT 9 (7–14) |
| Dose reductions\textsuperscript{310} | 290 per 1000 | 128 more per 1000 (41 to 238 more per 1000) | RR 1.44 (1.14–1.82) | Unknown (8)† | Moderate* | NNH 8 (5–25) |

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NNH = number needed to harm; NNT = number needed to treat; RR = risk ratio.

* Lack of blinding, inadequate allocation concealment
† Significant heterogeneity ($I^2 = 56\%$, $p = 0.003$).
\textsuperscript{1} Actual number of participants not reported; only percentages and RR values provided.
proportion of immigrants who accept screening for hepatitis C virus or their compliance with treatment of infection. In addition to potential patient barriers to the uptake of screening, there are likely important barriers that prevent primary care practitioners from recommending screening. More than half of primary care physicians have little experience in treating patients who test positive for infection, have a poor understanding of the natural history of the disease and incorrectly identify risk factors for infection. A large proportion (60%) of primary care physicians do not routinely ask patients about risk factors for hepatitis C on an initial visit. Of those who did identify higher-risk patients, only 50% to 75% ordered the appropriate diagnostic tests and correctly interpreted the results.

**Recommendations of other groups**

The US Preventive Services Task Force does not recommend screening for hepatitis C in the general US population (seroprevalence of about 1.8%) but rather recommends screening high-risk groups such as injection drug users. Neither the recent Canadian or US guidelines for managing hepatitis C identify immigrants as an at-risk group that should be targeted for screening. Our guidelines highlight the potential benefits of routine hepatitis C screening for at-risk immigrant groups.

**Take-home messages**

- About 3% of immigrants are infected with chronic hepatitis C virus (up to 18% in certain populations), and a large proportion of these will likely be eligible for treatment.
- The majority of immigrants acquire infection through unsafe injections or other medical procedures in their countries of origin, rather than through injection drug use.
- Immigrants are more likely to have concurrent infection (e.g., with hepatitis B or HIV), which increases the risk of progression of chronic hepatitis C virus–associated disease.
- Screening individuals for chronic hepatitis C virus infection and offering treatment before development of cirrhosis is important, because sustained virologic response is higher and treatment is better tolerated in the absence of cirrhosis.

10. Intestinal parasites: *Strongyloides* and *Schistosoma*

About one-third of the world’s population is infected with intestinal parasites, and most of these infections are sustained through cycles of repeated exposure from the environment. When populations emigrate from parts of the world where intestinal parasites are endemic and resettle in countries where they do not exist, most infections will clear without treatment within a few years after immigration. Two intestinal parasites — *Strongyloides* and *Schistosoma* — are notable exceptions in that they may persist for decades as subclinical infections or as low-grade disease with nonspecific clinical manifestations. In the presence of immunosuppression, strongyloidiasis can evolve rapidly into life-threatening disseminated disease, whereas chronic schistosomiasis can result in complications causing future morbidity and death. We conducted an evidence review to guide primary care practitioners in the early detection and treatment of strongyloidiasis and schistosomiasis for newly arriving refugees. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on screening for and treatment of these parasites are outlined in Box 10A.

**Methods**

We used the 14-step method developed by the Canadian Collaboration for Immigrant and Refugee Health (summarized in section 3 of this article, above). We considered the epidemiology of intestinal parasites in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We search MEDLINE, Embase, the Database of Abstracts of Reviews of Effectiveness, the HTA Database, the Cochrane Library and other sources from Jan. 1, 1980, to Jan. 1, 2010. Detailed methods, search terms, case studies, clinical considerations and research recommendations can be found in the complete evidence review for intestinal parasites (Appendix 8, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

**Results**

Our electronic search strategy for systematic reviews yielded 5039 articles. We identified an additional 150 articles through an open-ended search. We reviewed the titles of all 5189 articles, and evaluated 445 abstracts for potential relevance, eliminating 437 because they did not fulfill our inclusion criteria. Of the remaining eight manuscripts, five pertained to the diagnosis of *Strongyloides* or *Schistosoma*, and the remaining three pertained to treatment of these conditions. Although our search did not identify any systematic reviews on the diagnosis of either parasite or the treatment of strongyloidiasis, we found a systematic review pertaining to the treatment of schistosomiasis. Furthermore, we found two major guidelines pertaining to the evaluation and management of parasitic infections in newly arriving refugees.

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**Box 10A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: intestinal parasites**

**Strongyloides**

Screen refugees newly arriving from Southeast Asia and Africa with serologic tests for *Strongyloides*, and treat, if positive, with ivermectin (first-line therapy) or albendazole (if there are contraindications to ivermectin).

**Basis of recommendation**

**Balance of benefits and harms**

*Strongyloides* is estimated to infect 100 million people worldwide. Among immigrant populations, refugees from Southeast Asia and Africa appear to have the highest risk of infection. Subclinical infections or low-grade disease can persist for decades after immigration and in the presence of immunosuppression may transform into life-threatening disseminated disease. Treatment with ivermectin is of short duration, is highly effective (number needed to treat [NNT] 2, 95% confidence interval [CI] –1 to 3) and has a favourable adverse-effect profile.

**Quality of evidence**

Moderate

**Values and preferences**

The committee attributed more value to the availability of a highly sensitive and specific serologic test and effective treatment options to prevent potentially life-threatening disseminated disease than to the potential limitations of serologic testing in distinguishing current from remote infection in high-risk newly arriving refugees.

**Schistosoma**

Screen refugees newly arriving from Africa with serologic tests for *Schistosoma*, and treat, if positive, with praziquantel.

**Basis of recommendation**

**Balance of benefits and harms**

*Schistosoma* is estimated to infect 200 million people worldwide, of whom approximately 85% live in Africa. Among immigrant populations, refugees from Africa have the highest risk of infection. Subclinical infections or low-grade disease can persist for decades after immigration and may cause future morbidity or death. Serologic testing is the most sensitive diagnostic modality currently available. Treatment with praziquantel is of short duration, is highly effective (NNT 4, 95% CI –1 to 124) and has a favourable adverse-effect profile.

**Quality of evidence**

Moderate

**Values and preferences**

The committee attributed more value to the availability of a highly sensitive and specific serologic test and effective treatment to prevent future morbidity or death than to the limitations of serologic testing in distinguishing current from remote infection in high-risk newly arriving refugees.
What is the burden of strongyloidiasis and schistosomiasis in immigrant populations?

Quantifying the burden of parasitic infection is challenging because existing studies have not involved systematic or random testing of immigrant populations. Most estimates of burden have been derived from small observational studies and included primarily refugees from selected countries. Furthermore, a significant number of these studies used stool microscopy, a diagnostic test that is known to have limited sensitivity in the detection of each of these parasites.

In the case of strongyloidiasis, existing data have been derived primarily from refugee populations originating from Southeast Asia and Africa (Figure 10A). Studies using stool microscopy have reported prevalence rates between 0.8% and 4.3%, the highest burden being identified in refugees from Southeast Asia. Studies using serologic enzyme immunoassays have reported significantly higher prevalences of infection (between 9% and 77%), the highest burden being identified in refugees from Southeast Asia and Africa.

About 85% of the global burden of schistosomiasis is believed to occur in Africa. Studies using stool microscopy to detect *Schistosoma* in African refugee populations have reported prevalences from 0.4% to 7%. In contrast, studies using serologic enzyme immunoassays have reported significantly higher prevalences, ranging from 2.2% in East African pediatric populations to 64% in Sudanese refugees and 73% in Somalia refugees.

Does screening for strongyloidiasis and schistosomiasis decrease morbidity?

The scope and analytic horizon of existing studies prevent establishment of a direct link between screening for strongyloidiasis or schistosomiasis and an improvement in health outcomes. The association can be derived indirectly, however, since highly sensitive and specific diagnostic tests are currently available to detect each parasite, and effective treatment is known to mitigate the risk of future morbidity or death.

Screening tests

Stool microscopy for ova and parasites is the only definitive way to confirm the presence of intestinal infection with either Strongyloides or Schistosoma, but this diagnostic modality has suboptimal sensitivity. Although overall sensitivity can be improved by increasing the number of stool specimens examined, the costs associated with this approach can be substantial, and many patients are reluctant to provide multiple specimens. The sensitivity of a single stool examination to detect Strongyloides is estimated at just 30%, but this increases to over 90% when seven specimens are examined.

By contrast, serologic testing is the most sensitive diagnostic modality to detect Strongyloides and Schistosoma, making such tests ideal screening tools. Although these tests are also quite specific, serologic positivity cannot definitively distinguish current from remote infection. The National Reference Centre for Parasitology in Montréal, which performs serologic testing for both parasites in Canada, estimates that its enzyme immunoassays have 100% sensitivity and 88% specificity for Strongyloides stercoralis and 96% sensitivity and 82% specificity for Schistosoma mansoni. Several studies have reported levels of antibodies to Strongyloides declining after treatment, which suggests that serologic positivity is indicative of current infection; however, this has not been a universal finding. Levels of antibodies to Schistosoma do not appear to decline after treatment. Nonetheless, given the potential for these infections to persist for decades and to cause potentially life-threatening disease, it is generally presumed that a positive result on serologic testing for either parasite in high-risk newly arriving refugees (without a history of recent effective treatment) represents current infection.

Relative benefits and harms of treatment

The detection and subsequent treatment of strongyloidiasis and schistosomiasis can prevent future morbidity and potentially life-threatening complications. A two-day course of ivermectin (200 µg/kg orally once daily) is the preferred treatment for strongyloidiasis (Table 10A); however, among refugees from areas of the world where *Loa loa* is endemic, a seven-day course of albendazole (400 mg orally twice daily) should be used. This is because cases of encephalopathy have been reported with use of ivermectin during large-scale treatment campaigns in West and Central Africa, where *Loa loa* is endemic (see the guidelines of the US Centers for Disease Control and Prevention for a table of countries where *Loa loa* is endemic). This rare but potentially serious event may occur in persons who have a high load of *Loa loa* microfilaria, which are rapidly killed by ivermectin. Alternatively, because ivermectin is the most effective treatment option available for strongyloidiasis, practitioners may screen refugees at risk for *Loa loa* infection with a daytime-collected, thin blood smear for microfilaria and treat with ivermectin if high-level microfilaremia is not identified. For *Schistosoma* species found in Africa, a one-day course of praziquantel (40 mg/kg divided in

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**Figure 10A:** Number of immigrants and refugees screened for strongyloidiasis by birth country, as identified in our review of the medical literature between 1988 and 2010.
two doses) is the preferred treatment strategy (Table 10B).

Treatment with ivermectin or praziquantel in patients with underlying neurocysticercosis may lead to an acute inflammatory reaction and could precipitate seizure activity. Therefore, these drugs should not be used in persons with known neurocysticercosis or an unexplained seizure disorder. Otherwise, ivermectin, albendazole and praziquantel each have a generally favourable side-effect profile.34

Clinical considerations

On the basis of this review, we propose routine serologic screening for all newly arriving refugees at high risk for strongyloidiasis and schistosomiasis. With limited data on the burden of these two parasites in nonrefugee immigrant populations, there is currently insufficient evidence to justify routine screening of all nonrefugee populations, even though some high-risk groups may be missed.44 Further research is needed to clarify the burden of illness in other immigrant populations.

Practitioners should consider testing foreign-born persons for strongyloidiasis and/or schistosomiasis if they have lived in areas of the world where these parasites are endemic and (i) they have compatible signs and/or symptoms of infection (independent of the time elapsed since their arrival into Canada) and/or (ii) they have evidence of peripheral blood eosinophilia. Clinicians should also be aware that persons infected with the retrovirus human T-lymphotropic virus 1 (HTLV-1) have a modified immune response that complicates the treatment of strongyloidiasis.44 Some areas of the world are

<table>
<thead>
<tr>
<th>Patient or population</th>
<th>Patients with strongyloidiasis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Setting:</td>
<td>Okinawa, Japan</td>
</tr>
<tr>
<td>Intervention:</td>
<td>Ivermectin</td>
</tr>
<tr>
<td>Comparison:</td>
<td>Pyrvinium pamoate</td>
</tr>
</tbody>
</table>

Table 10A: Summary of findings for comparison of ivermectin and pyrvinium pamoate for strongyloidiasis

<table>
<thead>
<tr>
<th>Absolute effect</th>
<th>Risk with pyrvinium pamoate</th>
<th>Difference with ivermectin (95% CI)</th>
<th>Relative effect (95% CI)</th>
<th>No. of participants (studies)</th>
<th>GRADE quality of evidence</th>
<th>Comments (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cure rate*</td>
<td>233 per 1000</td>
<td>969 per 1000 (610 to 1000 per 1000)</td>
<td>RR 4.16 (2.62–6.59)</td>
<td>127 (1)</td>
<td>Moderate†</td>
<td>NNT 2 (-1 to 3)</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NNT = number need to treat; RR = risk ratio.

*Common adverse effects with treatment are pruritus, fever and tenderness of the lymph nodes. Serious or fatal encephalopathy has been reported (rarely) during treatment of patients with loiasis. Therefore, pretreatment assessment for Loa loa infection is recommended for any patient emigrating from areas where such infection is endemic (West and Central Africa).

†Not randomized, no blinding.

Table 10B: Summary of findings for comparison of praziquantel and placebo for treatment of schistosomiasis

<table>
<thead>
<tr>
<th>Patient or population</th>
<th>Patients with schistosomiasis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Setting:</td>
<td>Sudan, Zambia, Burundi, Democratic Republic of Congo, Kenya, Tanzania, Philippines, China, Republic of Congo, Niger</td>
</tr>
<tr>
<td>Intervention:</td>
<td>Praziquantel</td>
</tr>
<tr>
<td>Comparison:</td>
<td>Placebo</td>
</tr>
</tbody>
</table>

Table 10B: Summary of findings for comparison of praziquantel and placebo for treatment of schistosomiasis

<table>
<thead>
<tr>
<th>Absolute effect</th>
<th>Risk for control group</th>
<th>Difference with praziquantel (95% CI)</th>
<th>Relative effect (95% CI)</th>
<th>No. of participants (studies)</th>
<th>GRADE quality of evidence</th>
<th>Comments (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parasitologic cure</td>
<td>NA</td>
<td>NA</td>
<td>RR 28.29 (1.81–441.61)</td>
<td>69 (1)</td>
<td>Moderate†</td>
<td>NNT 4 (–1 to 124)</td>
</tr>
<tr>
<td>Adverse effects†</td>
<td>60 per 1000</td>
<td>361 per 1000 (173 to 750 per 1000)</td>
<td>RR 6.01 (2.89–12.5)</td>
<td>436 (1)</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NA = not applicable; NNT = number need to treat; RR = risk ratio.

†Dizziness, headache, malaise, abdominal pain.

*No details on randomization or blinding.
endemic for both *Strongyloides* and HTLV-1, and clinicians should therefore consider screening for the retrovirus if a patient (i) tests positive for *Strongyloides* and originates from an area with high prevalence of HTLV-1 (i.e., South America, the Caribbean, Japan or Africa) and/or (ii) has persistent strongyloidiasis that responds poorly to antiparasitic treatment.

Finally, the studies in this review primarily involved adult refugees, but children are well known to be at risk of infection with intestinal parasites. Consequently, screening newly arriving refugees of all ages from areas of the world endemic for strongyloidiasis and schistosomiasis is suggested. However, treatment with albendazole is not recommended in children under one year of age. Similarly, treatment with ivermectin is not recommended for children weighing less than 15 kg or less than 90 cm in length, and praziquantel is not recommended for children under four years of age. Infected children in these circumstances should be referred to a practitioner experienced in the management of intestinal parasites in pediatric populations.

Serologic samples to be tested for *Strongyloides* and *Schistosoma* are processed at the National Reference Laboratory for Parasitology in Montréal, so primary care providers should anticipate delays from the time a specimen is collected until test results are received. Furthermore, if patients require treatment for strongyloidiasis with ivermectin or albendazole, these drugs must be obtained through Health Canada’s Special Access Programme. Instructions on how to obtain these medications can be found at Health Canada’s website. By contrast, praziquantel can be obtained locally through a physician’s prescription.

Serologic testing after treatment for schistosomiasis is not recommended, as the antibodies tend to persist over time. By comparison, several studies have reported declining *Strongyloides* antibody titres 6 to 12 months after successful treatment, and the use of serologic testing has been advocated as a marker for clearance of this parasite. Although there is a body of evidence demonstrating this post-treatment effect, this finding has not been universally observed. A practice of post-treatment serologic testing may be considered by practitioners, but at a minimum, all treated individuals should be followed prospectively for clinical signs or symptoms of persistent infection and to ensure that eosinophil counts remain within or return to normal limits within six months of receiving effective treatment. Should patients have persistent symptoms and/or eosinophilia after six months, further investigations — including the option of repeat serologic testing for *Strongyloides* — should be pursued.

**Recommendations of other groups**

Two sets of national guidelines offer recommendations on the diagnosis and management of intestinal parasites in refugee populations. Both the US Centers for Disease Control and Prevention and the Australasian Society for Infectious Diseases support the use of serologic testing as part of screening for strongyloidiasis and schistosomiasis. Our recommendations focus routine serologic screening on migrant populations known to have high prevalence rates for these two parasitic infections.

**Take-home messages**

- Strongyloidiasis and schistosomiasis are parasitic infections that can persist for years to decades and consequently can cause serious morbidity or death long after an immigrant resettles in a new country.
- The burden of strongyloidiasis appears greatest in refugee populations originating from Southeast Asia and Africa, whereas the burden of schistosomiasis is greatest in refugee populations from Africa.
- Detection of strongyloidiasis or schistosomiasis is limited by subclinical infection or low-grade disease and by the suboptimal sensitivity of stool microscopy.
- Serologic testing substantially enhances diagnostic sensitivity.
11. Malaria

In 2006, worldwide, there were 250 million cases of malaria, leading to one million deaths.49 A large proportion of Canada’s new immigrants and refugees come from malaria-endemic countries (Figure 11A).100-348 The symptoms of malaria are nonspecific; they include fever and a constellation of other findings ranging from mild illness to fulminant organ failure and death. Plasmodium falciparum is the most virulent causative species, with an overall 1% case fatality rate.350 This rate increases to 10%–20% or higher for those with severe disease, defined by high parasitemia or end-organ damage.348,350 Most imported cases of malaria, particularly those due to *P. falciparum*, present within three months of leaving an endemic area.351-353 Canadian health care providers may have difficulties in diagnosing and treating malaria because of the rarity of the disease and unfamiliarity with laboratory diagnosis and clinical management.348,350 As well, drugs for treatment of malaria may not be readily accessible in pharmacies across the country. These delays, which may be compounded by migrants’ unfamiliarity with the Canadian health care system, can lead to severe disease and even death.351-357 In addition, malarial illness may interfere with the migrant’s successful integration into the host community because of issues such as physical incapacity and added financial stress.

We undertook this review to determine whether Canadian primary care physicians should routinely screen for malaria in asymptomatic adult and child migrants from low- to middle-income countries. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on malaria are outlined in Box 11A.

### Methods

We used the 14-step method developed by the Canadian Collaboration for Immigrant and Refugee Health46 (summarized in section 3 of this article, above). We considered the epidemiology of malaria in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE, Embase, CINAHL, HealthSTAR, the Cochrane Library and other sources from Jan. 1, 1996, to Dec. 31, 2010. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for malaria (Appendix 9, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

### Results

The initial searches yielded 1421 articles, of which 101 were selected for detailed appraisal. Initial searches for documents about screening for malaria in immigrants and refugees generated many retrospective and prospective reviews of screening in asymptomatic migrants, several of which related to refugees.351-361 Some of these articles provided recommendations on screening for malaria in migrants from areas where the disease is endemic; however, none reported use of a systematic review methodology. We found no randomized controlled trials investigating the effect of routine screening of asymptomatic individuals on morbidity and mortality related to malaria. The search for articles about treatment of malaria yielded 247 records, including two systematic reviews.352,380

<table>
<thead>
<tr>
<th>Box 11A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: malaria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do not conduct routine screening for malaria.</td>
</tr>
<tr>
<td>Be alert for symptomatic malaria in migrants who have lived or travelled in malaria-endemic regions within the previous three months, particularly in the context of fever or migration from sub-Saharan Africa, and perform timely diagnostic inquiry and testing (rapid diagnostic testing and thick and thin malaria smears).</td>
</tr>
</tbody>
</table>

### Basis of recommendation

**Balance of benefits and harms**

Individuals from malaria-endemic regions, particularly migrants from sub-Saharan Africa, remain vulnerable to acute *Plasmodium falciparum* malaria for the first three months after arrival. Clinical trials demonstrating the value of routine screening for asymptomatic malaria are lacking, prevalence data remain poor, the performance characteristics of malaria screening tests in asymptomatic individuals are uncertain, and local transmission of malaria is nonexistent in Canada. Thus, the focus of recommendations is on timely diagnosis and treatment of symptomatic malaria, where medications are effective and harms from adverse effects are minimal.

**Quality of evidence**

Low

### Values and preferences

The committee attributed more value to avoiding burden and cost from routine screening in the absence of clear evidence of prevalence of *P. falciparum* and uncertainty of performance of screening tests in asymptomatic individuals and determined that malaria was best addressed by primary care practitioners remaining alert for signs and symptoms of the disease and performing timely clinical diagnostic inquiry and treatment of symptomatic individuals.
groups are not screened before departure. Published reports of screening protocols for new migrants have indicated variable prevalence of malaria. Specifically, prevalence ranges from 6.8% to 64% among African migrants, most of whom are refugees. The prevalence among people from other malaria-endemic areas is much lower. A limited number of follow-up studies suggest that the risk of development of symptomatic malaria after screening is 20% to 40%.

The burden of symptomatic malaria among Canadian migrants is difficult to ascertain. The Public Health Agency of Canada receives reports of 350 to 1000 imported malaria cases per year, but the reason for travel in these cases is not reported. Such information is available, through the Canadian Malaria Network, only for cases of malaria requiring parenteral therapy, with 150 cases being reported from June 2001 through January 2010. Of these, 31 cases (20.7%) occurred in new migrants. Sixty-five percent of these (20/31) were children, whereas the overall percentage of pediatric cases was much lower (26.7% [40/150]). All but one migrant case originated from Africa, the exception occurring in a recent Karen refugee (from Burma). In Alberta, new migrants accounted for 20.7% (79/382) of malaria cases over 10 years (unpublished information, Alberta Health and Wellness). Data compiled from three Canadian tropical medicine clinics (GeoSentinel Surveillance Network Sites in Montréal, Ottawa and Toronto), reporting mostly illness in adults, for January 2006 through October 2010, found that migrants accounted for 15.1% (22/146) of malaria cases, with children accounting for 6.8% (10/146) of all cases and 27.3% (6/22) of cases involving migrants; overall, 60% of childhood cases occurred in migrants (D. Freedman, GeoSentinel Surveillance Network, Birmingham, Ala.: personal communication, 2010). In the United States, malaria in migrants accounted for 7.9% (362/4597) of all malaria cases from 2005 to 2007. Between 2007 and 2008, children were disproportionately represented among US migrant cases, accounting for 66.3% of malaria cases in that population, compared with about 20% of all malaria cases (P. Arguin, Surveillance Reports to Domestic Malaria Branch, Centers for Disease Control and Prevention. Atlanta, Ga.: personal communication, 2010). Most nonsurveillance studies report on the movements of refugees or case series in specialty clinics. In one pediatric case series from British Columbia, 28.6% of cases (over 17 years) occurred in immigrants. These data confirm the occurrence of symptomatic malaria, including severe malaria, in new migrants, including those to Canada, and highlight the disproportionate burden of the disease in migrant children.

![Figure 11A: Map of migrant source countries with overlay of malaria endemicity. Based on data from the 2006 Census of Canada and the World Health Organization.](image-url)
Does screening for malaria decrease morbidity and mortality?

Screening tests
Microscopic diagnosis with Giemsa-stained thick and thin blood smears has traditionally been the gold standard for diagnosing malaria and determining the level of parasitemia. These tests are inexpensive and ubiquitous, but time-consuming, and they rely on substantial expertise that is not readily available in laboratories outside of malaria-endemic areas. Rapid diagnostic tests detect antigens from lysed parasite-infected red blood cells, providing results within 5–20 minutes. These tests have the advantage of portability and ease of use. The usual lower limit of detection is 100 parasites/µL, with sensitivity declining at lower levels of parasitemia. A nonsystematic review concluded that the sensitivity of rapid diagnostic tests for P. falciparum was 95.3% (95% confidence interval [CI] 93%–97%); this dropped to 89.2% (95% CI 75%–97%) for 100–500 parasites/µL. The specificity for P. falciparum was 94.2% (95% CI 93%–95%). However, the ability to detect P. vivax malaria was not as robust, with a sensitivity of 68.9% (95% CI 66%–72%) and specificity 99.8% (95% CI 99%–100%). There is some uncertainty about the performance of rapid diagnostic tests in asymptomatic individuals. Polymerase chain reaction–based assays are also used to diagnose malaria and have lower detection limits. However, this significant improvement in detection levels is offset by the cost, institutional support, accessibility and time required for test results.

In Canada, there are many regions where laboratories are unaccustomed to diagnosing malaria by microscopy. The technology, convenience and affordability make rapid diagnostic tests a suitable alternative for Canadian health laboratories.

Relative benefits and harms of treatment
Data from predeparture treatment protocols for asymptomatic African refugees to Australia and the United States indicate that there has been a reduction in malaria cases among migrants. Of concern, however, are reports of treatment failures, often thought to be due to problems with completion of therapy or delays in departure following therapy. The efficacy of treating symptomatic malaria is well documented. The World Health Organization recommends artemisinin combination therapy as first-line treatment for P. falciparum malaria. However, these drugs are not licensed or available in Canada, where atovaquone and proguanil or quinine plus doxycycline (clindamycin in pregnancy or under the age of eight years) are recommended. Diagnosis and management for symptomatic individuals should follow the Canadian malaria guidelines.

In summary, although detecting and treating asymptomatic parasitemia may decrease morbidity and mortality from malaria, there is currently a paucity of clinical trials demonstrating the benefits and harms of routine screening and insufficient data to establish the true risk in migrants to Canada.

Clinical considerations
In 2009, Citizenship and Immigration Canada began screening for malaria, by means of rapid diagnostic testing, all refugees destined to Canada from East Africa; screening is performed 48–72 hours before departure. When the result of such testing is positive, treatment is provided with a full course of artemether and lumefantrine (Coartem). For lactating women and for children under 5 kg, a combination of artemunate and amodiaquine is used. For pregnant women, quinine is used. Australia follows the same protocol as Canada, whereas the United States provides presumptive treatment with artemether and lumefantrine to all persons except those who are pregnant, lactating or under 5 kg. For the latter groups, rapid diagnostic testing is provided, and only those with a positive result are treated.

It is important to remember that the majority of individuals migrating from areas of malaria risk, particularly sub-Saharan Africa, are not refugees from East Africa and will not have received predeparture malaria screening. As well, malaria may develop even in those who have undergone screening, including those who have received therapy, especially in the first three months after migration. Many new migrants to Canada are unfamiliar with the health care system, and those from malaria-endemic countries may be accustomed to having access to malaria therapy without visiting a physician or obtaining a prescription, and the perceived cost of medical care may be a barrier to receiving care. As well, migrants may be interacting with health care providers who are unfamiliar with diagnosis and management of the disease. Depending on the area of relocation, effective drugs for treating malaria may not be readily available. These factors may delay detection and treatment, thereby increasing the risk of severe malaria disease.

Numerous studies have documented the increased risk of malaria, including severe malaria, in migrants and their children who return to visit friends and family in malaria-endemic countries. Practitioners can take the opportunity to introduce the concept of future malaria risk and the need to seek pretravel advice for recommendations on preventing and managing malaria.

Recommendations of other groups
Canada has guidelines for prevention and treatment of malaria, but they do not address the topic of screening immigrants and refugees. In the United States, the Centers for Disease Control and Prevention have technical instructions addressing predeparture treatment of refugees, which recommend postarrival presumptive therapy for those who missed predeparture therapy. In Australia, there are guidelines for screening refugees from Africa who have resided in or travelled through a malaria-endemic region. The Australian guidelines advocate screening (with both thick and thin smear and rapid diagnostic testing) and treatment instituted by or in consultation with a specialist infectious disease service. Our guidelines highlight the risk of malaria but also the limitations of current evidence to support routine screening of asymptomatic migrants.
Take-home messages

• Routine screening is not recommended, but clinicians should be vigilant for symptoms of malaria.
• Migrants who have lived or travelled in malaria-endemic areas are vulnerable to acute malaria, particularly within the first three months after arrival.
• The symptoms of this disease (malaise, myalgia, headache and fever) are nonspecific, and primary care practitioners may not readily recognize them as symptoms of malaria.
• Delays in the diagnosis and treatment of *P. falciparum* infection may lead to severe disease and even death.
• Improved surveillance for malaria is needed in Canada, as well as more research related to the utility of screening immigrants and refugees for this disease.

12. Depression

Depression is a common and costly health care problem. Nearly all people with major depression are seen only in primary care, but up to 60% of cases go undetected and untreated. The level of underdiagnosis and inadequate treatment for depression is higher among migrants, who face cultural, linguistic and other barriers to accessing mental health care. Although migration in itself does not lead to an increase in depression, specific stressors and challenges can contribute to the onset of depression or influence its course, particularly among refugees. In general, immigrants to Canada have lower rates of depression than the general Canadian population, whereas refugees have comparable rates of depression but higher rates of post-traumatic stress disorder. Over time, the rate of depression in immigrant groups increases to match that of the general population. We undertook this review to determine whether existing approaches to screening for depression are appropriate for immigrants and refugees and to identify strategies that could improve the quality of care. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on screening for depression are outlined in Box 12A.

Box 12A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health:

**depression**

If an integrated treatment program is available, screen adults for depression using a systematic clinical inquiry or validated patient health questionnaire (PHQ-9 or equivalent).

Link suspected cases of depression with an integrated treatment program and case management or mental health care.

**Basis of recommendations**

**Balance of benefits and harms**

The number needed to treat to prevent one case of persistent depression was 18 (95% confidence interval 10–91) in studies of 1–12 months’ duration. Treatment in enhanced depression-care models accounts for an additional 1%–2% reduction in depressive symptoms relative to usual care. The prevalence of depression is similar among Canadians and among immigrants and refugees (10.7%), but access to care may be limited for migrants. No data on harms were reported, which would include patients’ out-of-pocket costs and adverse effects of medication.

**Quality of evidence**

Moderate

**Values and preferences**

The committee attributed more value to screening and treating depression to improve quality of life and less value to concerns about impairing rapport in therapeutic relationships, cultural acceptability and potential stigma of diagnostic labels, the cost and inconvenience of additional follow-up assessments, and the possible adverse effects or costs associated with treating patients with an incorrect diagnosis.

Note: PHQ-9 = nine-item Patient Health Questionnaire.

**Methods**

We used the 14-step approach developed by the Canadian Collaboration for Immigrant and Refugee Health (summarized in section 3 of this article, above). We considered the epidemiology of depression in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE, Embase, CINAHL, PsychLIT, the Cochrane Library and other sources from Jan. 1, 1998, to Jan. 1, 2010. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for depression (Appendix 10, available at www.cma.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

**Results**

Recommendations from the Canadian Task Force on Preventive Health Care and the US Preventive Services Task Force make scant mention of immigrants and refugees. In its guidelines for the treatment of depression, the American Psychiatric Association notes that language and other cultural variables may hamper accurate diagnostic assessment and treatment; it also mentions ethnic differences in the response to pharmacotherapy. The guidelines of the UK National Institute for Health and Clinical Evidence include statements on ethnic variations in prevalence and on the importance of social and cultural factors in choice of treatment. More recent studies, discussed in the complete evidence review (Appendix 10, available at www.cma.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1), provide evidence that can inform the implementation of screening for depression and integrated care for immigrants and refugees in primary care.

**What is the burden of depression in immigrant populations?**

The Canadian Community Health Survey (version 1.2) revealed a lifetime prevalence of depression of 10.8% in the general population. Immigrants who had arrived in Canada in the previous four years had the lowest rates of depression (3.3%–3.5%). Among those who had arrived 10–14 years ago (rate 8.5%) or more than 20 years ago (rate 6.8%–7.2%), rates were similar to those of the Canadian-born population. Proficiency in English or French and employment status did not affect these rates. A meta-analysis of studies on serious mental disorders among refugees found rates of depression similar to those in the general population but much higher levels of post-traumatic stress disorder, often in association with depression.

Pregnancy and the postpartum period have been associated with symptoms of depression in immigrant women. Risk factors may include stressful life events, lack of social support.
or isolation, physical health problems, inability to speak the language of the host country, the demands of multiple roles and separation from children who have remained in the country of origin.396,397

**Does screening for depression decrease morbidity and mortality?**

**Screening tools**

Many screening instruments for depression have been validated in primary care settings, and little evidence suggests that any particular instrument performs better than other instruments, although brief tools tend to be less specific.398 Both brief screening tools (two or three items) and longer ones tend to have relatively high false-positive rates (60%–70%) when the prevalence of depression is 10%.399 Therefore, positive results on screening must be confirmed by a full diagnostic interview. Most screening instruments have not been validated for many of the immigrant groups commonly seen in primary care in Canada, although the patient health questionnaire has been validated with Chinese, South Asian and other populations.

**Relative benefits and harms of treatment**

In a systematic review of screening for depression conducted in 2002, the US Preventive Services Task Force found that clinical trials of integrated programs have demonstrated modest improvements in patient outcomes, but benefits have not been observed when screening results are simply reported to physicians without coordinated treatment and follow-up.400 Subsequent reviews have confirmed this finding.401 Adverse effects among immigrants have not been systematically studied, but they may include impaired rapport and less use of general medical services if patients believe they are being labelled and stigmatized or are being treated improperly, the cost and inconvenience of additional follow-up assessments, and possible adverse effects or costs associated with treating patients with an incorrect diagnosis.

In a recent meta-analysis, Gilbody and associates402 found no benefit for screening alone, although there was some benefit in high-risk populations. However, a cumulative meta-analysis showed modest benefit when an integrated system of collaborative care was in place for follow-up (Table 12A).402 In a low-quality longitudinal study conducted in the United States, Wells and colleagues403 examined the effect of screening for depression within an integrated system of care, with follow-up by nurses and with other quality-associated improvements. The greatest improvement was seen for minority groups, specifically African Americans and Latinos.404

**Clinical considerations**

**Screening**

Screening should be conducted in a language in which the patient is fluent, either with translated instruments or through a trained interpreter. Cultural variations in presentation of symptoms, ways of coping and the stigma attached to mental health problems may complicate detection and treatment.405 The presence of prominent somatic symptoms and patients' tendency to attribute their depressed mood to somatic distress can also reduce primary care physicians' recognition of depression.406 Among refugee patients with depression, more than half also have post-traumatic stress disorder, and this comorbidity can complicate the recognition of depression.407 Many cultures strongly stigmatize mental health problems, which may limit disclosure of behavioural or emotional difficulties.408 Depression can be distinguished from other forms of mental health problems and can be explained as a state of “energy depletion” and demoralization, which may provide a rationale for psychosocial assessment and treatment.

**Child-bearing women**

Guidelines from obstetrical groups have proposed that women be screened for depressive symptoms in each trimester of pregnancy, at 1–2 weeks postpartum, and possibly at 2, 4, and 6

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<tr>
<td><strong>Table 12A: Summary of findings for effects of collaborative care for depression</strong></td>
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<td><strong>Absolute effect</strong></td>
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<tr>
<td>Outcome</td>
<td>Risk for control group</td>
<td>Difference with collaborative care</td>
<td>Relative effect (95% CI)</td>
<td>No. of participants (studies)</td>
</tr>
<tr>
<td>Depression at 6 mo</td>
<td>See comment</td>
<td>0.25 (0.18–0.32)</td>
<td>NA</td>
<td>12 344 (35)</td>
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Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NA = not applicable; NNT = number needed to treat.

*Standardized depression scales.
†Directness uncertain because the studies were conducted in the US health system setting, and it is unclear whether their results would apply to immigrants and refugees in the Canadian health care system.
months postpartum. The Edinburgh Postnatal Depression Scale or nine-item patient health questionnaire have been used with immigrant women. Immigrant women’s multiple roles in the home and the workplace may impede access to health services. Availability of child care facilities, transportation and support from family members and spouses can facilitate their seeking help. Group meetings can be an effective way to provide social support and health-promotion information.

Adolescents and children
The US Preventive Services Task Force has recommended screening adolescents (age 12–18 years) when integrated systems of treatment are available, including assessment, psychotherapy and follow-up. It is unclear which of the more than 30 available depression scales is best for screening and diagnosing depression among immigrant and refugee youth.

Elderly people
Migrant elderly people have not been well studied but may have a high risk of depression because of social isolation, loss of familiar surroundings and the changing nature of the family as members adapt to the new social context.

What are the potential implementation issues?
Linguistic and cultural differences may constitute substantial barriers to recognition of depression and subsequent treatment negotiation and delivery. Medical interpreters, “culture brokers,” bilingual and bicultural mental health practitioners, clinician training in cultural competence and cultural consultation may mitigate these potential barriers. Screening for depression produces benefits only when it is linked to an integrated system of care. An integrated system involves the following elements: systematic patient education, availability of allied health professionals to support continuity of care, frequent follow-up, a caseload registry to track patients, caseload supervision by a psychiatrist if indicated, stepped care and a plan for preventing relapse. Stepped care involves a progression of levels from patient education and self-management to medication or psychotherapy and, for complex cases, referral to a mental health practitioner.

The clinical relationship is central to detection and treatment of mental health problems in primary care. Screening with structured questionnaires cannot replace clinical sensitivity, systematic inquiry and relationship-building. Given the great diversity of immigrant and refugee patients, no single approach is likely to be sufficient for optimal recognition and appropriate treatment of depression.

Recommendations of other groups
The Canadian Task Force on Preventive Health Care recommends screening adults for depression in primary care when integrated systems that include diagnostic, treatment and follow-up components are in place. The US Preventive Services Task Force recommends screening adolescents (age 12–18 years) when integrated systems of treatment, including assessment, psychotherapy and follow-up, are in place; however, it concludes that evidence is insufficient to make any recommendation for children 7–11 years of age. Our recommendations highlight the value of screening for depression in the context of integrated treatment programs.

Take-home messages

- Rates of depression are lower among new immigrants to Canada, but over time these rates generally rise to match the rate in the general population.
- The prevalence of depression among refugees is comparable to that in the general population.
- Existing guidelines for depression suggest that all patients should be screened for depression when integrated systems are in place to provide follow-up treatment.
- For immigrants, use information about depression in relevant languages, translated screening questions and trained interpreters, as well as systematic inquiries about losses, stressors and symptoms.
- Moderate to severe depression should be treated with a stepped-care model, beginning with psychoeducation and antidepressant medication, close follow-up and culturally appropriate counselling.


More detailed information and resources for screening, assessment and treatment of depression can be found at: www.mnhrc.ca.
13. Post-traumatic stress disorder

A large proportion of new immigrants to Canada come from countries experiencing social turmoil, and some are directly affected by protracted conflicts or war. Refugees and others who face significant trauma and loss are at risk for mental health consequences, including post-traumatic stress disorder. For three main reasons, primary care practitioners play a key role in the recognition and management of post-traumatic stress disorder in immigrants and refugees. First, immigrants and refugees underutilize formal mental health services. Second, an integrated treatment approach is often needed for extreme traumas, common in refugees, such as torture and rape, which have severe and long-lasting consequences for both physical and mental health. Third, a family perspective is essential because trauma stemming from organized violence tends to affect the whole family, particularly children, who may not display dramatic or easily recognizable symptoms. We conducted an evidence review to determine the burden of post-traumatic stress disorder within immigrant and refugee populations, to evaluate the effectiveness of screening and treatment, and to identify barriers for primary care. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on post-traumatic stress disorder are outlined in Box 13A.

**Methods**

We used the 14-step method developed by the Canadian Collaboration for Immigrant and Refugee Health (summarized in section 3 of this article, above). We considered the epidemiology of post-traumatic stress disorder in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE, Embase, CINAHL, PsychLIT, the Cochrane Library and other sources from Jan. 1, 2002, to Dec. 31, 2010. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for post-traumatic stress disorder (Appendix 11, available at www.cma.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

**Results**

We identified 16 systematic reviews relevant to immigrants and refugees and five guidelines. We selected the 2005 guidelines commissioned by the National Institute for Clinical Excellence for the management of post-traumatic stress disorder in primary care, but none of the selected intervention studies in those guidelines provided evidence for immigrants or refugees. We also selected four Cochrane reviews on treatment of post-traumatic stress disorder, the practice guidelines from the International Society for Traumatic Stress Studies and a systematic review on treatment of this condition in refugees and asylum seekers.

**What is the burden of illness of post-traumatic stress disorder in immigrant populations?**

Most persons who experience traumatic events have a favourable mental health prognosis. When symptoms of post-traumatic stress disorder or acute stress disorder develop, there is, in most cases, substantial natural recovery (estimated at about 80%). However, those in whom post-traumatic stress develops may remain symptomatic for years and are at risk of secondary problems, such as substance abuse. A meta-analysis of studies involving adult refugees resettled in developed countries reported a 9% prevalence of post-traumatic stress disorder, and 5% had major depression. Among refugees with major depression, 71% also had post-traumatic stress disorder. Conversely, 44% of refugees with post-traumatic stress disorder also had major depression. Studies of child refugees report 11% prevalence of post-traumatic stress disorder. Symptoms may be reactivated when faced with new traumas, particularly if reminiscent of earlier traumatic experiences. Torture and cumulative trauma are the strongest predictors of post-traumatic stress disorder and are associated with chronic physical and mental health problems.

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**Box 13A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: post-traumatic stress disorder**

- **Do not conduct routine screening for exposure to traumatic events, because pushing for disclosure of traumatic events in well-functioning individuals may result in more harm than good.**
- **Be alert for signs and symptoms of post-traumatic stress disorder, especially in the context of unexplained somatic symptoms, sleep disorders or mental health disorders such as depression or panic disorder, and perform clinical assessment as needed to address functional impairment.**

**Basis of recommendation**

**Balance of benefits and harms**

Many persons who have been exposed to trauma do fine once they find safety and social supports. Brief screening instruments overestimate the rate of disease because they focus on symptoms and do not measure functional impairment. Detailed inquiry and pushing for disclosure without indications of distress or disorder could be harmful. There are no clinical trials demonstrating the benefits of routine screening for post-traumatic stress disorder.

**Quality of evidence**

Low (evidence available for refugee populations)

**Values and preferences**

The committee attributed more value to preventing potential harms from routine screening in the absence of clear evidence of benefits and determined that post-traumatic stress disorder was best dealt with through primary care practitioners remaining alert for signs and symptoms of this condition and performing clinical assessment to address functional impairment.
Longitudinal studies from Canada indicate that most adults and children with refugee status adapt well, in spite of a high level of exposure to premigratory trauma. A population-based health survey from Quebec similarly found that non-refugee immigrants also experienced high levels of premigratory trauma, but that most immigrants were in good mental health.

**Does screening for post-traumatic stress disorder decrease morbidity and mortality?**

**Screening**

Several short screening instruments practical for primary care settings have been developed. The four-item primary care post-traumatic stress disorder screening scale and the Breslau seven-item screening scale (available at http://ajp.psychiatryonline.org/cgi/content/full/156/6/908#T2) are two simple means of identifying symptoms in primary care patients. In both cases, their cultural validity is unknown. Very few screening instruments have been tested for diagnostic accuracy among immigrants, refugees and asylum seekers. However, it may be reasonable to use questionnaires to assist in identifying symptoms, as part of a clinical assessment when addressing functional impairment.

**Relative benefits and harms of psychological treatment (adults and children)**

The systematic review and meta-analysis commissioned by the National Institute for Clinical Excellence provided evidence that psychological treatments, including trauma-focused cognitive–behavioural therapy and eye movement desensitization and processing, reduce the symptoms of post-traumatic stress disorder. We rated the quality of this evidence as low because of study limitations and inconsistency of results. Two Cochrane reviews provided similar evidence of effectiveness. A recent systematic review showed that psychological treatments (cognitive–behavioural therapy and narrative exposure therapy) can reduce symptoms of post-traumatic stress disorder among refugees, but we rated this evidence as very low quality. Other authors have reported that patients may experience adverse effects with therapy, such as re-experiencing traumatic events, and rates of withdrawal from active therapy may approach 30%.

**Clinical considerations**

**What are the potential implementation issues?**

Primary care practitioners need to be aware that immigrants and refugees may have been exposed to traumatic events. If a patient discloses a traumatic experience, it may be helpful to acknowledge the pain and suffering associated with the experience, to explain that a reaction is common for anyone who has undergone trauma and to offer empathetic reassurance that the situation is likely to get better. Several Canadian cities have centres and experts available to help care for survivors of trauma and torture.

Exploration of trauma and its consequences is not recommended in the first meeting with a patient unless it is the patient’s primary complaint. However, certain symptom presentations should alert clinicians to the need for assessment for post-traumatic stress disorder, including unexplained physical complaints, sleep disorders, depression, panic disorder and somatoform disorder. Other presentations, such as severe dissociation mimicking brief reactive psychosis, dissociative disorders (amnesia and conversion) and psychotic depression, although less frequent, may also be related to post-traumatic stress disorder. Key elements of the assessment include level of psychological distress, the impairment associated with the symptoms in the patient and his or her family, substance abuse and suicidality.

Familiarity with the cultural background of the patient is recommended, and assessment should involve a professional interpreter if the patient’s language ability is inadequate to express psychological distress and narrate the experience. Disclosing traumatic experience through relatives, family members or, particularly, children can be traumatic.

Although not supported by clinical trials, the National Institute for Clinical Excellence recommends a phased intervention model, reflecting a pragmatic approach for refugees and asylum seekers who face the possibility of being returned to a traumatic environment. Phase I is defined as the period in which safety has not yet been established and during which intervention should focus on practical, family and social support. Phases II and III should focus on the patient’s priorities, which may include social integration and/or treatment of symptoms. Unemployment, isolation and discrimination may overshadow the efficacy of mental health treatment in many patients, which suggests that multifaceted interventions that include primary care, community organizations and other social institutions may be effective.

**Recommendations of other groups**

The UK National Institute for Clinical Excellence recommends against routine systematic provision of brief, single-session interventions. It recommends that consideration be given to the use of a brief screening instrument to detect post-traumatic stress disorder among refugees and asylum seekers, but does not suggest any specific instrument for screening or provide evidence of effectiveness of treatment in refugees. It also recommends that children and youth with post-traumatic stress disorder be offered a course of trauma-focused cognitive behaviour therapy. For sleep disorders, the National Institute for Clinical Excellence recommends the short-term use of hypnotic medication for adults or, if longer-term treatment is required, the use of suitable antidepressants to reduce the risk of dependence. For significant comorbid depression or severe hyperarousal, the National Institute for Clinical Excellence recommends paroxetine and mirtazapine. The US Centers for Disease Control and Prevention state that, in general, the majority of people who experience reactions to stress after disasters and emergencies show resilience and do not go on to experience long-term psychopathology. Our recommendations highlight the paucity of evidence for routine screening and the potential for harms.
Take-home messages

- Forty percent of Canadian immigrants and refugees from countries involved in war or with significant social unrest have been exposed to traumatic events before migration.
- Most (estimated at 80%) individuals who experience traumatic events heal spontaneously after reaching safety.
- Empathy, reassurance and advocacy are key clinical elements of the recovery process.
- Pushing for disclosure of traumatic events by well-functioning individuals may result in more harm than good.
14. Child maltreatment

Child maltreatment is an important public health problem worldwide. The 2003 Canadian incidence study of reported child abuse and neglect estimated an incidence rate of 22 per thousand for child maltreatment. Of reported cases, 15% involved emotional maltreatment, 28% involved exposure to domestic violence, 24% involved physical abuse, 30% involved neglect and 3% involved sexual abuse. Surveys conducted with nonrepresentative ethnic minority samples (which have likely included immigrants and refugees) have yielded higher rates of maltreatment than appear in official reports. This review was undertaken to clarify reports of child maltreatment in ethnic communities, to determine whether existing tools to screen for child maltreatment are appropriate for immigrant and refugee children, and to recommend strategies to improve the quality of care for these populations. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health related to child maltreatment are outlined in Box 14A.

Methods

We used the 14-step approach developed by the Canadian Collaboration for Immigrant and Refugee Health (summarized in section 3 of this article, above). We considered the epidemiology of child maltreatment in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE, Embase CINAHL, PsychLIT, the Cochrane Library and other sources from Jan. 1, 1995, to Dec. 31, 2010. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for child maltreatment (Appendix 12, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

Results

We found no systematic reviews or guidelines on screening, prevention or treatment for child maltreatment in recently settled immigrants or refugees. The general literature search identified 180 titles with reference to child maltreatment. Seventeen citations were selected, and five key reviews retained as evidence. Studies conducted with general population and ethnic minority samples provided additional evidence that informed our recommendations related to child maltreatment among immigrants and refugees.

What is the burden of child maltreatment in immigrant populations?

The prevalence and incidence of child maltreatment among immigrant and/or refugee children in Canada are unknown. The evidence on maltreatment among ethnic minority children in the United States and Canada suggests that some ethnic minority children are disproportionately over- and under-represented in child protection services. These children are more likely to be screened for child maltreatment and also more likely to be reported to child protection services. Higher rates of screening result in a higher rate of potential risks associated with increased reporting to child protection services.

<table>
<thead>
<tr>
<th>Box 14A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: child maltreatment</th>
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<tbody>
<tr>
<td><strong>Screening</strong></td>
</tr>
<tr>
<td>Do not conduct routine screening for child maltreatment.</td>
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<tr>
<td>Be alert for signs and symptoms of child maltreatment during physical and mental examinations, and assess further when reasonable doubt exists or after patient disclosure.</td>
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<tr>
<td><strong>Basis of recommendations</strong></td>
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<tr>
<td><strong>Balance of benefits and harms</strong></td>
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<tr>
<td>The committee recommends against routine screening because of poor performance of screening instruments and the potential harms caused by the very high false-positive rates. Sensitivity ranged between 25% and 100%, specificity between 16.5% and 94.3%, and positive predictive value (when available) between 1.7% and 28.2%.</td>
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<tr>
<td><strong>Quality of evidence</strong></td>
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<tr>
<td>Low</td>
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<tr>
<td><strong>Values and preferences</strong></td>
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<tr>
<td>The committee attributed more value to evidence for the negative effects of screening in relation to the high potential for harms. Harms could result from false positives leading to inappropriate labelling, psychological distress, inappropriate family separation, impaired clinician–patient rapport, potential reduction in use of general medical services and legal ramifications associated with involvement of child protection services.</td>
</tr>
<tr>
<td><strong>Prevention of child maltreatment and associated outcomes</strong></td>
</tr>
<tr>
<td>A home visitation program encompassing the first two years of life should be offered to immigrant and refugee mothers living in high-risk conditions, including teenage motherhood, single parent status, social isolation, low socioeconomic status, or living with mental health or drug abuse problems.</td>
</tr>
<tr>
<td><strong>Basis of recommendation</strong></td>
</tr>
<tr>
<td><strong>Balance of benefits and harms</strong></td>
</tr>
<tr>
<td>Home visitation programs for high-risk mothers, provided by nurses, reduced days in hospital for children (p &lt; 0.001). Harms from surveillance and reporting to child protection services were not clearly demonstrated.</td>
</tr>
<tr>
<td><strong>Quality of evidence</strong></td>
</tr>
<tr>
<td>Moderate</td>
</tr>
<tr>
<td><strong>Values and preferences</strong></td>
</tr>
<tr>
<td>The committee attributed more value to supporting high-risk mothers with an offer of a home visitation program to provide practical support for families and the program’s potential to improve health outcomes for children than to the potential risks associated with increased reporting to child protection services.</td>
</tr>
</tbody>
</table>
inappropriate referral to child protection services. Ethnic minority children who received medical examinations were twice as likely (p < 0.001) to be reported to child protection services.51

The Canadian incidence study of reported child abuse and neglect found44 that ethnic minority children had a 1.8 times greater likelihood to be over-represented, whereas white and Arab children were under-represented. The higher rates were found among Aboriginals, Blacks, Latinos and Asians (the latter group for only physical abuse). This racial bias may be one explanation why ethnic minority children are disproportionately represented at all levels of the child protection process, despite the fact that they do not seem to be at higher risk of maltreatment. Another explanation may be professionals’ divergent views as to what should be considered grounds for clinical suspicion of child maltreatment, which is associated with recency of training in child abuse, prejudices about the perpetrator and the professionals’ beliefs in the positive or negative consequences of reporting a given family to child protection services.54

### Does screening for child maltreatment reduce harm and premature death or disability?

#### Screening tools

Most screening methods consist of self-administered questionnaires generally completed by the mother, interviews or checklists completed by the professional who collects information directly from the child or clinical judgments by nurse or professional teams. All screening methods attempt to predict child maltreatment on the basis of either parents’ potential for maltreatment or the presence or level of risk factors associated with maltreatment, rather than on the occurrence of actual maltreatment. Three systematic reviews have reported that these instruments tend to have high sensitivity but poor specificity and false-positive rates too high for use in clinical settings. Sensitivity ranged between 25% and 100%, specificity between 16.5% and 94.3%, and positive predictive value (when available) between 1.7% and 28.2%.

#### Table 14A: Summary of findings for home visitation by nurses to prevent child maltreatment

<table>
<thead>
<tr>
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</thead>
<tbody>
<tr>
<td><strong>Outcome</strong></td>
<td><strong>Risk for control group</strong></td>
<td><strong>Difference with home visitation by nurses (95% CI)</strong></td>
<td><strong>Relative effect (95% CI)</strong></td>
<td><strong>No. of participants (studies)</strong></td>
</tr>
<tr>
<td>Out-of-home placements (follow-up: 16 mo)</td>
<td>226 per 1000</td>
<td>31 more per 1000 (70 fewer to 201 more per 1000)</td>
<td>RR 1.14 (0.69–1.89)**†</td>
<td>197 (1)</td>
</tr>
<tr>
<td>Mean no. of substantiated reports of child abuse and neglect over 15 yr</td>
<td>0.54†‡</td>
<td>0.25 fewer‡</td>
<td>0.77 (0.34–1.19)**</td>
<td>245 (1)‡‡</td>
</tr>
<tr>
<td>Mean no. of days in hospital for injuries and ingestions over 2 yr</td>
<td>0.16</td>
<td>0.13 fewer</td>
<td>NA</td>
<td>697 (1)‡‡‡</td>
</tr>
<tr>
<td>Mean no. of health care encounters for injuries and ingestions over 2 yr</td>
<td>0.55</td>
<td>0.12 fewer</td>
<td>NA</td>
<td>697 (1)‡‡‡</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NA = not applicable; NNT = number needed to treat; RR = risk ratio.

†Because RR crosses 0 (i.e., not statistically significant), the NNT could not be estimated.
‡Pregnant women with “specified psychosocial risk factors”: substance abuse, homelessness, domestic violence, psychiatric illness, incarceration, HIV infection or lack of social support.
§“When the recommendation is in favour of an intervention and the 95% confidence interval (or alternative estimate of precision) around the pooled or best estimate of effect includes no effect and the upper confidence limit includes an effect that, if it were real, would represent a benefit that would outweigh the downsides” (GRADE Pro software).
‡‡Adjusted for socioeconomic status, marital status, maternal age, education, locus of control, support from husband or boyfriend, working status, and husband or boyfriend use of public assistance at registration.
‡‡‡“Comparison log incidence” – (intervention log incidence).
Relative benefits and harms from screening

False-positive ratings, which are the most common result in low-risk populations, can lead to a number of negative consequences, such as inappropriate labelling and punitive attitudes, psychological distress, inappropriate separation of children from family support systems, destruction of family supports, loss of resources and loss of autonomy for those falsely accused. This may leave parents wary of any subsequent assistance that may be offered, thus reducing their access to care. A systematic review of the performance of screening tests concluded that adding a screening protocol to the clinical encounter yielded additional false-positives that exceeded additional abused children detected.

Compared with the general population, immigrant and refugee families may be more likely to suffer from the direct and indirect harms related to screening. Screening instruments have not been culturally validated and are less likely to be accurate because of factors such as language barriers, different cultural norms of behaviours and different attitudes toward institutional authority. Given the limited state of knowledge in immigrant populations, potential harms from routine screening for child maltreatment outweigh benefits, which have not yet been clearly established.

Relative benefits and harms of preventing child maltreatment

Home visitation programs by nurses aim to prevent child maltreatment by assessing and supporting families. To date, the 15-year longitudinal study by Olds and associates has provided the best evidence for the effectiveness of a nurse–family partnership program in reducing actual child maltreatment. The effectiveness of this program is particularly evident for first-time mothers who are younger than 19 years of age, single or economically disadvantaged (Table 14A). Another prevention program (the Early Start Program) has also shown efficacy in reducing hospital admissions for child injuries at 36 months (17.5% vs. 26.3% for control group).

Relative benefits and harms of treatment for child maltreatment

Several specific forms of intervention have been devised to reduce the consequences of child maltreatment. Trauma-focused cognitive–behavioural therapy reduces sexually abused children’s symptoms of anxiety, depression and sexual behaviour problems in both general population and ethnic minority children. Table 14B presents the outcomes of cognitive behavioural interventions. Parent–child interaction therapy showed a reduction in repeated reports of physical abuse in treatment relative to control groups (standard psychoeducational program) (19% vs. 49%). In most other studies, the outcomes were not statistically significant but there was a consistent tendency in favour of treatment programs. The lack of evidence of efficacy for immigrant or
refugee children precludes extrapolation of the findings to these groups.

**Clinical considerations**

**What are the potential implementation issues?**

Some forms of child discipline may be unusual or outside Canadian social norms but are not pathological or dangerous for the child. Immigrant or refugee families may resort to other disciplinary behaviours (e.g., hitting a child with an object) that are condoned in their cultural context but that contravene child protection laws in Canada. Some cultural practices (e.g., scarification as part of life cycle rituals among some African children or cupping, a common traditional healing method in some Asian cultures that leaves circular ecchymoses) may be misinterpreted as signs of child abuse. Other culture-specific practices (e.g., female genital cutting) contravene child protection and civil laws in Canada. In situations where child maltreatment is suspected, observed or disclosed, the practitioner must take action in accordance with the child protection law in his or her region.

Language barriers, fear of separation from the child, fear of punitive institutional power and fear of deportation may constitute major barriers to disclosure of child maltreatment. Failure to investigate family dynamics and inter-generational conflicts, after disclosure of maltreatment by an immigrant child, may further disempower the parents and attribute greater power to the child, consequently aggravating his or her problem. Immigrant and refugee children placed in foster care may suffer from loss of connection with language of origin and religious, familial and cultural traditions. As a preventive strategy, clinicians may want to provide families with sources of information about their province’s child protection law, their legal rights and their obligations regarding children, in addition to addressing other risk factors for child maltreatment. Recent research is showing that the SEEK (Safe Environment for Every Kid) model is promising.467

**Recommendations of other groups**

The US Preventive Services Task Force concluded that there is insufficient evidence for or against routine screening of child abuse.456 The Canadian Task Force on Preventive Health Care concluded that there is fair evidence to exclude screening for child maltreatment.79 The American Academy of Paediatrics and the American Medical Association do not support universal screening, but recommend that physicians be alert for signs and symptoms of child maltreatment during routine physical examination. The Task Force on Community Preventive Services of the US Centers for Disease Control and Prevention recommends early childhood home visitation for the prevention of child maltreatment in high-risk families and families with low-birth-weight infants.471 Our recommendations highlight the importance of prevention and the potential harms of routine screening in the context of cultural and linguistic diversity.

**Take-home messages**

- Children from ethnic minorities, including recently settled immigrants and refugees, are eight times more likely to be subjected to screening for child maltreatment than children in the general population.
- Immigrant and refugee families may be particularly vulnerable to the harms that can occur because of legal and institutional interventions consequent to false-positive screening results, such as over-reporting for child maltreatment and unnecessary separation of the child from his or her family.


More detailed information and resources on cultural aspects of child maltreatment can be found at: www.mmbrc.ca.
15. Intimate partner violence

Intimate partner violence, defined as physical, emotional, financial and/or sexual abuse perpetrated against the victim by his or her intimate partner, is a significant public health problem worldwide. In Canada, a 1999 study of a nationally representative sample of 26,000 participants reported 8% intimate partner violence against a female and 7% against a male by a previous or current partner in the past five years. Women, however, are more likely than men to be the victims of serious violent acts such as sexual abuse, beatings (25% v. 10%), being choked (20% v. 4%) or being threatened or having a weapon used against them (13% v. 7%). They are also more likely than men to be injured during the violent act (40% v. 13%) and to be fearful for their lives (40% v. <10%). In this review we aimed to determine whether existing screening tools and approaches for intimate partner violence are appropriate for immigrant and refugee women and to identify care barriers for these populations. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health related to intimate partner violence are outlined in Box 15A.

Methods

We used the 14-step approach developed by the Canadian Collaboration for Immigrant and Refugee Health (summarized in section 3 of this article, above). We considered the epidemiology of intimate partner violence in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE, Embase, CINAHL, PsychLIT, the Cochrane Library and other sources from Jan. 1, 1995, to Dec. 31, 2010. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for intimate partner violence (Appendix 13, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

Results

We found no systematic reviews or evidence-based guidelines on screening, prevention or treatment for intimate partner violence in immigrants or refugees. The general literature search identified 409 titles on intimate partner violence, and after appraisals, we retained two key reviews as evidence. After the search update, we selected two additional key reviews and one randomized controlled trial. Studies conducted with general population and ethnic minority samples informed our clinical recommendations.

What is the burden of intimate partner violence in immigrant populations?

Three studies provided secondary analyses of the 1999 Statistics Canada General Social Survey. Women born in developing countries reported the highest prevalence rates of intimate partner violence, followed by Canadian-born women and immigrant women from developed countries. However, when all other variables in the model were controlled for, the analysis showed that recently settled immigrant women (i.e., in Canada for less than 10 years) had significantly lower odds of intimate partner violence victimization than longer-term immigrants and Canadian-born women. Single, divorced, separated or widowed immigrant women were 10 times more likely to report intimate partner violence than immigrant women married or in a common-law relationship. Immigrant women reported higher rates of emotional abuse than Canadian-born women (14.7% v. 8.7%), with the strongest risk factor being their partner’s low educational level.

Regional surveys on intimate partner violence have yielded higher rates. MacMillan and colleagues reported rates that ranged from 4.1% to 17.7% for Canadian-born women and 12.6% for foreign-born women. Ahmad and coauthors reported a 22% rate of intimate partner violence following computer screening. Prevalence rates also vary in relation to the health care setting (highest prevalence in emergency departments). Finally, women in war zones, disaster zones, during flight or displaced in refugee camps in countries of asylum may be at higher risk for intimate partner violence.

Does screening for intimate partner violence reduce morbidity or mortality?

Screening tools

Screening for intimate partner violence differs from tradi-

Box 15A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: intimate partner violence

Do not conduct routine screening for intimate partner violence.

Be alert for potential signs and symptoms related to intimate partner violence, and assess further when reasonable doubt exists or after patient disclosure.

Basis of recommendation

Balance of benefits and harms

Current evidence does not demonstrate clear benefits from screening women for intimate partner violence, and harms have resulted from screening. Compared with the general population, there may be greater risk among immigrant and refugee women for harm directly related to screening (e.g., risk of loss of migration status and sponsorship agreements). Harm may occur indirectly through impaired patient–physician rapport and subsequent reduction in use of medical and mental health services.

Quality of evidence

Moderate

Values and preferences

The committee attributed more value to evidence of harms and lack of evidence of benefits and less value to recommending uncertain interventions, even in the face of significant concerns.
tional screening for medical disorders because the target of clinical concern is a behavioural event, which women usually recognize as a problem but which they may not view as appropriate for medical attention. Four short self-report questionnaires have received the most study. The “Hurt, Insulted, Threatened, or Screamed at” questionnaire (four items) yields sensitivity ranging from 30% to 100% and specificity from 86% to 99%. The Partner Violence Screening Tool (three items) provides sensitivity from 35% to 71% and specificity from 80% to 94%. The Women Abuse Screening Tool (five items) yields sensitivity ranging from 32% to 94% and specificity from 55% to 99%.

A Canadian randomized controlled trial found women preferred self-completed approaches. However, other studies comparing administration methods of screening instruments (e.g., face-to-face interviews, computer screening, written screening) have shown inconsistent results. Furthermore, it is unknown whether these results apply to immigrant and refugee women.

Relative benefits and harms of screening
A Canadian trial on the effect of screening found no statistically significant differences between women screened or not screened at 6, 12 or 18 months follow-up for recurrence of intimate partner violence (Table 15A). More than half of the women who disclosed being victims of intimate partner violence on screening did not discuss the violence with their practitioner during the health care visit. An important study limitation was that no specific intervention was provided to women who disclosed or screened positive.

Other studies have found screening benefits such as decreasing isolation, increasing support, relief, breaking the silence and validating women’s feelings. However, these same studies identified several harms, including feeling that the practitioner is too busy or not interested, feeling judged and being disappointed by the practitioner’s response, increased anxiety, concerns about privacy, breaches of confidentiality and legal repercussions, fear of being reported to child protective services, and concern about or actual increased risk of retaliation or further harm from the partner.

Relative benefits and harms of treatment
The strongest evidence for treatment has come from studies of the Experimental Social Innovation and Dissemination program, which reported decreased physical and emotional abuse at 12–24 months follow-up and improvement of women’s quality of life at 12 months follow-up. Ramsay and coworkers reported that, while promising, the results were inconclusive. In Table 15B, we report the efficacy of the Experimental Social Innovation and Dissemination advocacy and counselling intervention program in decreasing the incidence of intimate partner violence in an ethnically

| Table 15A: Summary of findings on screening for intimate partner violence to reduce morbidity due to such violence |

| Patient or population: English-speaking female patients |
| Settings: Health care settings in Ontario |
| Intervention: Screening for intimate partner violence |
| Comparison: No screening |

<table>
<thead>
<tr>
<th>Outcome (18-mo follow-up)</th>
<th>Absolute effect</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Risk for control group</td>
</tr>
<tr>
<td>Intimate partner violence, by Composite Abuse Scale</td>
<td>530 per 1000</td>
</tr>
<tr>
<td>Post-traumatic stress disorder screening, by SPAN (startle, physically upset by reminders, anger, numbness)</td>
<td>601 per 1000</td>
</tr>
<tr>
<td>Quality of life, by WHO Brief</td>
<td>Mean score 52.7</td>
</tr>
<tr>
<td>Depression</td>
<td>Mean score 24.4</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NA = not applicable; NNT = number needed to treat; RR = relative risk; WHO = World Health Organization.

*Calculated using Review Manager on the basis of observed counts.
†Only one study.
‡Dichotomous outcome: total number of events was less than 300.
§Continuous outcome: total population size was less than 400.
diverse sample of women who had spent at least one night in a shelter.

Clinical considerations

What are potential implementation issues?
Signs and symptoms of intimate partner violence differ significantly among women. They may be absent in some women or be of a psychological (depression, anxiety, suicidal ideation, alcohol or drug abuse), social (social isolation) and/or physical (injuries, bruises and aches) nature in other women. Patient–physician rapport thus remains a key element in the detection of intimate partner violence.

Recently settled immigrant women in Canada are more likely to report intimate partner violence to the police than women in the general population but are less likely to use social services.\(^{64}\) Barriers to help-seeking included fear of deportation or not accessing Canadian citizenship, lack of knowledge of services or language-specific services, experiences of racism or discrimination.\(^{65}\) Culturally specific perceptions of spousal relationships, gender roles, negative experiences with authorities, aggression and abuse may affect reporting and disclosure.\(^{66}\) Involvement with police or criminal proceedings may put immigrant women at risk of losing their sponsorship agreements.\(^{67,68}\)

Intimate partner violence is now considered a form of child maltreatment. Women may delay disclosure of violence because of fear of losing custody of their children (child protection services often cite the mother’s failure to protect her children).\(^{45,69}\) In addition, some women feel coerced into staying in a shelter to keep custody of their children. Although this may protect them from further intimate partner violence, it may also isolate them from extended family and community networks that might otherwise be integrated effectively into the intervention plan.\(^{69}\)

Services that can defuse conflict situations and reduce family stress include social welfare, reliable childcare, safe housing, language classes, and other educational and vocational training opportunities. Community grassroots organizations can provide information and support groups in appropriate languages and in a culturally competent manner.\(^{46–48}\) Research is beginning to show benefits when screening and interventions target women with specific conditions, for example pregnancy, mental illness and substance abuse, but this work has yet to consider the immigrant context.

Recommendations of other groups

National clinical preventive screening committees, the Canadian Task Force on Preventive Health Care, the UK National Screening Committee and the US Preventive Services Task Force have not found sufficient evidence to recommend for or against screening all women for intimate partner violence.\(^{67,68}\) The UK National Screening Committee concluded that “screening for domestic violence should not be introduced” in periodic health examinations. The American Medical Association, the American Academy of Family Physicians and the American College of Obstetricians and Gynecologists have recommended routinely screening all women for intimate partner violence.\(^{70}\) However, these organizations have not based their recommendations on systematic reviews of effectiveness. Our guidelines

| Table 15B: Summary of findings for advocacy programs to prevent further intimate partner violence |
|-----------------|-----------------|-----------------|-----------------|
| **Patient or population:** Women in a Midwest shelter program for women with abusive partners who had (i) spent at least one night in the shelter and (ii) planned on staying in the general vicinity for the first three months after leaving the shelter | **Setting:** Community setting | **Intervention:** Advocacy programs | **Comparison:** No advocacy program |
| **Sources:** Wathen CN, Macmillan HL. Interventions for violence against women: scientific review. *JAMA* 2003;289:589-600.\(^{47}\) Sullivan CM, Bybee DI. Reducing violence using community-based advocacy for women with abusive partners. *J Consult Clin Psychol* 1999;67:43-53.\(^{68}\) | **Absolute effect, mean score** | **Risk for control group** | **Difference with advocacy programs (95% CI)** | **No. of participants (studies)** | **GRADE quality of evidence** |
| **Outcome** | **Self-reported severity or frequency of abuse (scale 0–3; follow-up 24 mo)** | 0.85 | 0.15 higher | 265 (1)\(^{68}\) | Low*†† |
| | **Effectiveness in obtaining community resources (scale 1–4; follow-up 10 wk)** | 2.7 | 0.50 higher (0.34 higher to 0.66 higher) | 265 (1) | Low*†† |
| | **Quality of life (scale 1–7; follow-up 24 mo)** | 4.94§ | 0.25 higher (0.02 lower to 0.52 higher) | 265 (1) | Low*†† |
| | **Depression (scale 0–3; follow-up 24 mo)** | 2.00 | 0.08 lower (0.24 lower to 0.08 higher) | 265 (1) | Low*†† |

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation.

*Only one study.
†Concerns about directness and applicability only to women seen in primary care who have been in a shelter.
‡Fewer than 300 events.
§Postintervention scores.
highlight the paucity of data on the effectiveness of screening programs and the concern for potential harms from routine screening.

**Take-home messages**

- The rate of reporting of intimate partner violence is lower among recently settled immigrant women than among longer-term immigrants and Canadian-born women.
- Linguistic barriers, financial dependencies, fear of losing custody of children and limited knowledge of laws and health services constitute significant barriers to both disclosure and adherence to interventions among immigrant and refugee women.

- To decrease the rate of abuse, practitioners should refer women who report spending at least one night in a shelter to a structured program of patient-centred (advocacy) support services.


More detailed information and resources on cultural aspects of intimate partner violence can be found at: www.mnhrc.ca.
16. Type 2 diabetes mellitus

More than two million Canadians have diabetes mellitus and, by the end of this decade, this number is expected to rise to three million.49 People of South Asian, Latin American and African ethnicity have a two to four times greater risk for development of type 2 diabetes than the white population.57,580 Up to 50% of Canada’s recent immigrants come from South Asia, Latin America and the Caribbean.39 The efficiency of diabetes screening improves when targeted to undiagnosed and high-risk populations.501 We conducted an evidence review to estimate the diabetes burden for immigrant populations, to evaluate the effectiveness of screening for prevention and treatment, and to identify barriers and facilitators to prevention and treatment of diabetes in primary care. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on screening and treating prediabetes and diabetes are outlined in Box 16A.

Methods

We used the 14-step approach developed by the Canadian Collaboration for Immigrant and Refugee Health16 (summarized in section 3 of this article, above). We considered the epidemiology of diabetes in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE, Embase, CINAHL, the Cochrane Library and other sources from Jan. 1, 1995, to July 14, 2010. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for type 2 diabetes (Appendix 14, available at www .cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

Results

We found no randomized controlled trials evaluating the effect of diabetes screening programs on morbidity and mortality. We identified three reviews focusing on the genetic risk factors predisposing high-risk ethnic groups to diabetes,502-504 Additionally, we identified a systematic review505 and a meta-analysis506 on the disparities in type 2 diabetes treatment between high-risk ethnic groups and the general white population and two systematic reviews on culturally appropriate treatment of type 2 diabetes in high-risk populations.507,508 An updating search, building on the 2008 guidelines of the Canadian Diabetes Association509 and the systematic review of the US Preventive Services Task Force510 on type 2 diabetes, yielded an additional 21 articles, of which two were selected.48,511 We also identified a meta-analysis of randomized controlled trials512 that studied the impact of tight glycemic controls on morbidity and mortality.

What is the burden of type 2 diabetes in immigrant populations?

The prevalence of diabetes is increasing worldwide, most notably in South Asia, sub-Saharan Africa and Latin America, regions that are major contributors to international migration. Contributory factors cited for increased prevalence of type 2 diabetes include genetic susceptibility,503,504 increasing sedentary lifestyle, changing food habits512 and increasing levels of adult and childhood obesity.

In Canada, up to 50% of new immigrants face a twofold to fourfold higher prevalence of type 2 diabetes than whites513 and experience a younger age of onset.514 A prevalence study in Ontario509 reported higher prevalences for South Asians (odds ratio [OR] 4.01 for men, 3.22 for women), Latin Americans (OR 2.18 for men, 2.40 for women) and sub-Saharan Africans (OR 2.31 for men, 1.83 for women). The average age at diagnosis of diabetes is decreasing, for example, from 52.0 to 46.0 years in the United States (p < 0.05)514 and, most dramatically, in black, Asian and Hispanic populations. In India, for example, there is a significantly higher prevalence of type 2 diabetes in people 35 to 44 years (8.5%–11.5%, p = 0.05).515 Central adiposity (overweight or obesity) results in a higher degree of insulin resistance and hence confers a higher risk for diabetes and premature coronary artery disease.516 A study conducted in a high-risk population from Jamaica found that the sensitivity of waist circumference in predicting type 2 diabetes was 71% in men and 65% in women and that the specificity of this measure was 79% in men and 60% in women.517 which

Box 16A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: type 2 diabetes mellitus

Screen immigrants and refugees > 35 years of age from ethnic groups at high risk for type 2 diabetes (South Asian, Latin American and African) with fasting blood glucose.

Basis of recommendation

Balance of benefits and harms

Detecting impaired fasting blood glucose and treating with diet and exercise can delay the onset of diabetes (number needed to treat [NNT] 5, 95% confidence interval [CI] 4–6). Treating patients with diabetes with intensive blood pressure interventions can decrease mortality (NNT 38, 95% CI 23–203), and tight glucose control can decrease myocardial infarctions (NNT 131, CI 87–298). People of South Asian, Latin American or African ethnicity face a twofold to fourfold higher prevalence of type 2 diabetes with earlier onset compared with white people. Minimal harms are reported for lifestyle interventions and adverse effects of antihyperglycemic agents.

Quality of evidence

Moderate

Values and preferences

The committee attributed more value to delaying the onset of diabetes than to the current uncertainty of impact on mortality for lifestyle interventions. The committee also attributed greater value to the potential to decrease morbidity and mortality with treatment of hypertension and hyperglycemia in high-risk ethnic populations than to concern about harms due to treatments.
suggests that waist circumference can be a useful alternative to body mass index. The International Diabetes Federation has recommended the use of ethnoscopic waist circumference as a cut-off parameter for screening these high-risk populations (Table 16A).51,519

**Does screening decrease the incidence of diabetes and/or related cardiovascular complications?**

**Screening tests**

Fasting blood glucose is considered the most appropriate screening test for diabetes because of its feasibility and specificity.510 A fasting blood glucose level of 7.0 mmol/L has a moderate sensitivity (40%–87%), but good specificity (96%–99%), to predict a blood glucose level of 11.1 mmol/L in a two-hour oral glucose tolerance test. A recent systematic review also concluded that haemoglobin A1c and fasting blood glucose are equally effective screening tools, but haemoglobin A1c is more costly than fasting blood glucose.520

**Relative benefits and harms of treatment**

The 2008 meta-analysis of the US Preventive Services Task Force on lifestyle interventions for prediabetes560 showed a significant decrease in onset of diabetes with minimum harms. We rated the quality of this evidence as high but noted the need to study the long-term effects of delaying diabetes onset (Table 16B).521 Additionally, we identified a Cochrane systematic review58 showing that culturally appropriate health education was effective in reducing blood glucose and increasing diabetes knowledge in ethnic minority groups. None of these trials reported significant adverse effects from lifestyle interventions. The evidence was insufficient to comment on the negative theoretical impact of labelling.

Blood pressure control in diabetes reduces cardiovascular morbidity and mortality.510,521 Trials reported minor adverse drug reactions such as cough with some antihypertensives and no harms related to the socioeconomic implications of diagnosis and treatment (Table 16C).521 A recent meta-analysis515 of five prospective randomized controlled trials evaluating the effect of intensive glucose control showed a small but significant decrease in myocardial infarctions in the treatment group, which had a haemoglobin A1c level 0.9% lower than the control group, but no effect on events of stroke or mortality. Very intensive glucose control (haemoglobin A1c < 6.5%) has not been shown to decrease mortality and has been associated with hypoglycaemia. Populations and individuals who would benefit the most from screening have undiagnosed, longstanding disease, and intensive lifestyle and/or hypertension interventions would have the greatest absolute effect.

**Clinical considerations**

**What are the potential implementation issues?**

A Cochrane review521 found that diet and lifestyle interventions reduced weight and the incidence of diabetes, but these measures were more effective in younger populations. Others have found that diabetes education is less effective in situations of low socioeconomic status and cultural factors. Language barriers, migration stress and cultural diversity (diets, lifestyles, perceptions of weight, fasting at Ramadan) may complicate diabetes care and education.

Ethnic groups prefer culturally appropriate dietary and exercise advice.58 A Cochrane review58 on culturally appropriate health education showed improved glycemic control and diabetic knowledge scores compared with standard diabetic education approaches at 12 months. Other outcomes, such as lipid levels and blood pressure, showed no significant difference from control groups. Another review,524 looking at diabetic care in socially disadvantaged populations, identified interventions that may improve outcomes, including culturally tailored care, a focus on individuals in assessment and feedback, and incorporation of treatment algorithms.

**Recommendations of other groups**

The US Preventive Services Task Force recommends screening in all asymptomatic hypertensive populations523 and suggests population-targeted screening for those with hyperten-

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**Table 16A:** Country- and ethnic-specific values for waist circumference51,519

<table>
<thead>
<tr>
<th>Country or ethnic group</th>
<th>Waist circumference, cm* (as measure of central obesity)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Europid†</strong></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>≥ 94</td>
</tr>
<tr>
<td>Female</td>
<td>≥ 80</td>
</tr>
<tr>
<td><strong>South Asian‡</strong></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>≥ 90</td>
</tr>
<tr>
<td>Female</td>
<td>≥ 80</td>
</tr>
<tr>
<td><strong>Chinese</strong></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>≥ 90</td>
</tr>
<tr>
<td>Female</td>
<td>≥ 80</td>
</tr>
<tr>
<td><strong>Japanese§</strong></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>≥ 85</td>
</tr>
<tr>
<td>Female</td>
<td>≥ 90</td>
</tr>
<tr>
<td><strong>Ethnic South and Central American</strong></td>
<td>Use South Asian recommendations until more specific data are available</td>
</tr>
<tr>
<td><strong>Sub-Saharan African</strong></td>
<td>Use European data until more specific data are available</td>
</tr>
<tr>
<td><strong>Eastern Mediterranean and Middle East</strong></td>
<td>Use European data until more specific data are available for (Arab) populations</td>
</tr>
</tbody>
</table>

*These are pragmatic cut points, and better data are required to link them to risk. Ethnicity rather than country of residence should be the basis for classification. In future epidemiologic studies of populations of Europid origin, preference should be given to using both European and North American cut points to allow better comparisons.
†In the United States, the Adult Treatment Panel III values (102 cm for males, 88 cm for females) are likely to continue to be used for clinical purposes.
‡Based on a Chinese, Malay and Asian-Indian population.
§Subsequent analyses suggested that Asian values (90 cm for males, 80 cm for females) should be used for the Japanese population until more data are available.
sion, advanced age and obesity. The Canadian Diabetes Association recommends diabetes screening for all adults over 40 years of age. Earlier testing should be considered in people with additional risk factors, and ethnic-specific waist circumference cut-off points should be used for risk stratification. The American Diabetes Association recommends screening

### Table 16B: Summary of findings for lifestyle modifications to prevent type 2 diabetes mellitus

**Patient or population:** Patients at high risk for type 2 diabetes, with impaired fasting glucose  
**Settings:** Outpatient or community health clinics  
**Intervention:** Lifestyle modifications for patients with impaired fasting glucose  
**Comparison:** No intervention  

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Risk in control group</th>
<th>Risk with lifestyle modifications (95% CI)</th>
<th>Relative effect (95% CI)</th>
<th>No. of participants (studies)</th>
<th>GRADE quality of evidence</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Decreased incidence of type 2 diabetes: FPG, OGTT (mean follow-up 3.8 yr)</td>
<td>471 per 1000</td>
<td>245 fewer per 1000 (169 to 254 fewer per 1000)</td>
<td>RR 0.48 (0.4-0.58)</td>
<td>5275 (S)</td>
<td>High+†‡</td>
<td>NNT 5 (4-6)</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; FPG = fasting plasma glucose; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NNT = number needed to treat; OGTT = oral glucose tolerance test; RR = risk ratio.  
*Test for heterogeneity: Q = 6.104, p = 0.192.  
†These studies applied to our population group (i.e., age > 35 yr, Canadian, ethnic groups).  
‡There were sufficient participants, and most events recorded were minimal in the intervention group compared with the control group.

### Table 16C: Summary of findings for intensive antihypertensive treatment in patients with type 2 diabetes mellitus to prevent cardiovascular disease

**Patient or population:** Patients with type 2 diabetes and hypertension  
**Settings:** Hospital outpatient or community health clinics  
**Intervention:** Intensive blood pressure control  
**Comparison:** Moderate blood pressure control  

<table>
<thead>
<tr>
<th>Outcome* (mean follow-up 166.9 yr)</th>
<th>Risk with moderate blood pressure control</th>
<th>Risk with intensive blood pressure control (95% CI)</th>
<th>Relative effect (95% CI)</th>
<th>No. of participants (studies)</th>
<th>GRADE quality of evidence</th>
<th>Comments (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Decrease in incidence of stroke</td>
<td>46 per 1000</td>
<td>17 fewer per 1000 (9 to 29 fewer per 1000)</td>
<td>RR 0.64 (0.46-0.89)</td>
<td>3599 (27)</td>
<td>Moderate†‡</td>
<td>NNT 61 (41-198)</td>
</tr>
<tr>
<td>Decrease in cardiovascular events</td>
<td>140 per 1000</td>
<td>35 fewer per 1000 (15 to 62 fewer per 1000)</td>
<td>RR 0.75 (0.61-0.94)</td>
<td>3599 (27)</td>
<td>Moderate†‡</td>
<td>NNT 29 (19-120)</td>
</tr>
<tr>
<td>Decrease in cardiovascular death</td>
<td>64 per 1000</td>
<td>21 fewer per 1000 (4 to 52 fewer per 1000)</td>
<td>RR 0.67 (0.40-1.12)</td>
<td>3599 (27)</td>
<td>Low††‡</td>
<td>NNT not statistically significant</td>
</tr>
<tr>
<td>Decrease in total mortality</td>
<td>99 per 1000</td>
<td>27 fewer per 1000 (9 to 48 fewer per 1000)</td>
<td>RR 0.73 (0.56-0.95)</td>
<td>3599 (27)</td>
<td>Moderate†‡</td>
<td>NNT 38 (23 to 203)</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NNT = number needed to treat; RR = risk ratio.  
*Definitions from the International Classification of Disease, Ninth Revision.  
†Heterogeneity was not indicated for the studies involving diabetic patients; reported heterogeneity value is for all studies, including patients with and without diabetes.  
‡The population of interest would consist of people at least 35 years of age, whereas the mean age of patients in these studies was 59.6 to 60 years of age. Additionally, most study participants with diabetes might have been diagnosed many years ago and be asymptomatic, whereas our clinical action would involve screening patients who are asymptomatic.  
§Wide confidence interval including null effect.
with fasting glucose or hemoglobin $A_{1c}$ in adults who are overweight or obese (body mass index $\geq 25$ kg/m$^2$) and who have one additional risk factor (e.g., physical inactivity, first-degree relative with diabetes, members of high-risk ethnic populations, hypertension). The International Diabetes Federation guidelines$^4$ do not recommend universal screening but suggest that detection programs should target high-risk people identified by assessment of risk factors. Our recommendations focus on improving the precision of screening for high risk ethnic populations.

**Take-home messages**

- Persons of South Asian, Latin American and African origin develop hyperglycemia at a younger age and face a twofold to fourfold higher prevalence of type 2 diabetes than white people.
- Persons with hypertension and hypercholesterolemia are at high risk for complications from diabetes and have the most to benefit from treatment of obesity, high cholesterol, hypertension and hyperglycemia.
- Culturally appropriate diabetes education and lifestyle interventions are more effective at controlling levels of hemoglobin $A_{1c}$ than standard approaches.

17. Iron-deficiency anemia

Iron-deficiency is the most common cause of anemia, and iron-deficiency anemia is the most common nutritional disorder in the world. Other causes of anemia (such as malaria and hemoglobinopathies) may coexist, depending on patients’ diets, living conditions and genetic predispositions. We focus here on iron-deficiency anemia, which can lead to poor pregnancy outcomes, impaired physical and cognitive development in children and reduced work productivity in women. No routine iron-deficiency screening or supplementation program is offered in Canada for immigrants, either before or after their arrival. We conducted an evidence review on the effectiveness of screening and iron-supplement interventions for immigrant populations. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on screening for and treatment of iron-deficiency anemia are outlined in Box 17A.

Methods

We used the 14-step method developed by the Canadian Collaboration for Immigrant and Refugee Health (summarized in section 3 of this article, above). We considered the epidemiology of iron-deficiency anemia in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE, Embase, CINAHL, the Cochrane Library and other sources from Jan. 1, 1996, to Jan. 1, 2010. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for iron-deficiency anemia (Appendix 15, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

Results

We found no systematic reviews or guidelines on screening for iron-deficiency anemia among immigrants and refugees. We based our search update on the US Preventive Services Task Force’s systematic review on iron-deficiency anemia. We identified relevant clinical trials that would help assess the effectiveness of screening for and treating iron-deficiency anemia and that added new and primary evidence to the published systematic reviews. We identified 76 articles as relevant to screening and treatment of iron-deficiency anemia among immigrants and refugees.

What is the burden of iron-deficiency anemia in immigrant populations?

Newly arriving immigrant and refugee children and women have a higher prevalence of anemia (15%–28%) than the Canadian-born population (2%–10%), excluding First Nations populations. The World Health Organization has estimated that the prevalence of iron-deficiency anemia among preschool children ranges from 21% to 68%. The prevalence of iron-deficiency anemia for women of reproductive age ranges from 18% to 48% (Table 17A).

The main risk factors for iron-deficiency anemia are low iron intake, poor absorption of iron related to diets high in phytate or phenolic compounds, and periods of life with high iron demand. Studies on newly arriving refugee children have identified exclusive breastfeeding after six months of age without the use of iron supplements or iron-rich complemen-

Box 17A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: iron-deficiency anemia

Women
Screen immigrant and refugee women of reproductive age for iron-deficiency anemia (with hemoglobin).

Basis of recommendation

Balance of benefits and harms
Treating iron-deficiency anemia provides an average net change in hemoglobin concentration of 15 g/L (number needed to treat [NNT] 2, 95% confidence interval [CI] 2–3) and an increase in function and provides a net change in the productivity ratio (NNT 4, 95% CI 3–8). The prevalence of iron deficiency is higher among immigrant women than among Canadian-born women (> 15% v. < 15%). Harms are minimal and include diarrhea and the personal costs of iron supplements.

Quality of evidence
Moderate

Values and preferences
The committee attributed more value to improving health among women of child-bearing age and less value to uncertainty about whether asymptomatic immigrant and refugee women value the treatment outcomes.

Children
Screen immigrant and refugee children aged one to four years for iron-deficiency anemia (with hemoglobin).

Basis of recommendation

Balance of benefits and harms
Treating children with iron-deficiency anemia improves cognitive development, with a standardized mean difference of 0.30, equivalent to a modest effect of 1.5–2 intelligence quotient points (NNT 7, 95% CI 5–14). Immigrant and refugee children have a higher prevalence of iron-deficiency anemia than Canadian-born children (> 20% v. < 20%). Adverse effects from iron treatment are minimal. The NNT for immigrant and refugee children is expected to be similar because many of the studies were conducted in developing countries.

Quality of evidence
Moderate

Values and preferences
The committee attributed more value to ensuring optimal opportunities for immigrant children and potential reduction of disparities in education, literacy and wages between immigrant and Canadian-born populations and less value to the discomfort of testing and treatment risk of diarrhea.
tary foods, use of cow’s milk or nonfortified infant formula as weaning food, early and frequent consumption of tea and relatively infrequent consumption of meat. Other important considerations in children are the presence of other illnesses and obesity.

Migrants from regions in the developing world with a high prevalence of hookworm and malaria, high parity and genetic predisposition for red blood cell disorders (hemoglobinopathies) are at increased risk for anemia. Alpha- and beta-thalassemias are most common in Africa, the Mediterranean,

Table 17A: Anemia prevalence and number of people affected among preschool-age children and nonpregnant women in each World Health Organization (WHO) region*

<table>
<thead>
<tr>
<th>WHO region</th>
<th>Preschool-age children†</th>
<th>Nonpregnant women†</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Prevalence, % (95% CI)</td>
<td>No. affected, millions (95% CI)</td>
</tr>
<tr>
<td>Africa</td>
<td>67.6 (64.3–71.0)</td>
<td>83.5 (79.4–87.6)</td>
</tr>
<tr>
<td>Americas</td>
<td>29.3 (26.8–31.9)</td>
<td>23.1 (21.2–25.1)</td>
</tr>
<tr>
<td>Southeast Asia</td>
<td>65.5 (61.0–70.0)</td>
<td>115.3 (107.3–123.2)</td>
</tr>
<tr>
<td>Europe</td>
<td>21.7 (15.4–28.0)</td>
<td>11.1 (7.9–14.4)</td>
</tr>
<tr>
<td>Eastern Mediterranean</td>
<td>46.7 (42.2–51.2)</td>
<td>0.8 (0.4–1.1)</td>
</tr>
<tr>
<td>Western Pacific</td>
<td>23.1 (21.9–24.4)</td>
<td>27.4 (25.9–28.9)</td>
</tr>
<tr>
<td>Global</td>
<td>47.4 (45.7–49.1)</td>
<td>293.1 (282.8–303.5)</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval.
*Adapted, with permission, from the World Health Organization.
†Preschool-age children: 0.00–4.99 yr.
‡Nonpregnant women: 15.00–49.99 yr.

Table 17B: Summary of findings for iron supplements to treat iron deficiency in children

Patient or population: Children with iron deficiency
Setting: International locations (developed and developing countries), participants identified within communities or schools
Intervention: Iron supplements
Comparison: Children without iron deficiency

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Absolute effect</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Risk in control group</td>
</tr>
<tr>
<td>Mental development (scale 0–100; median follow-up 3 mo)</td>
<td>No data</td>
</tr>
<tr>
<td>Weight for height (follow-up 3–6 mo)</td>
<td>No data</td>
</tr>
<tr>
<td>Infectious illnesses</td>
<td>1430 per 1000</td>
</tr>
<tr>
<td>Diarrhea events per child-years</td>
<td>1160 per 1000</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NA = not applicable; NNT = number needed to treat; SMD = standard mean difference.
*This is a modest effect, equivalent to 1.5–2.0 intelligence quotient points. In younger children (age ≤ 27 months), no effect of iron supplementation on mental development was detected.
†Consistency in infectious illnesses: heterogeneity Q = 78.29, df = 28, p < 0.0001.
‡Intervention and control group numbers are the number of events per child-years (observation/exposure time).
India and Southeast Asia, whereas sickle cell anemia is most often found in people of African descent. Glucose-6-phosphate deficiency (worldwide equatorial distribution) can lead to hemolysis in relation to oxidative injury from certain medications (e.g., nitrofurantoin, sulfamethoxazole) and thus can contribute to iron deficiency.56

Does screening for iron-deficiency anemia decrease morbidity?

Screening
The positive predictive value of hemoglobin concentration below 110 g/L, alone for iron deficiency in children (12–35 months) is 29% (95% confidence interval [CI] 20%–38%), and the sensitivity is 30% (95% confidence interval [CI] 20%–40%).531 Diagnostic measures used to confirm iron-deficiency anemia include serum ferritin and hemoglobin or hematocrit response to administration of iron.

Relative benefits and harms of treatment
Iron deficiency can be addressed through primary prevention by ensuring adequate iron intake27,533–535 or through secondary prevention by detecting the condition and treating it with iron supplements or diet education.90

Cognitive development in children
Sachdev and colleagues538 evaluated changes in weight-for-height between placebo and iron-treatment groups in children and found no statistically significant difference. Sachdev and colleagues539 also conducted a meta-analysis combining 17 trials (10 of which were conducted in developing countries) of iron supplementation in infants or in children up to 12 years of age and found increased mental development scores in the treatment group (Table 17B). In the subgroup analyses, improvement in mental development scores was attributed mainly to five trials in children seven years of age or older, and the effect was only intermediate for children between two and five years of age. Common adverse effects associated with iron supplements include dose-related reversible gastrointestinal symptoms534,535 and unintentional overdose.

Work productivity in women of reproductive age
We identified three randomized controlled trials studying the effect of iron supplements on work productivity.90,541,542 These trials, conducted in China, Indonesia and Sri Lanka, all reported increases in work productivity with iron supplements. Table 17C presents the study from China, which we chose as the best-quality study for women of child-bearing age who are not pregnant.540 Because of the complexity of measuring work productivity and the significance for only one of the two primary measures, we rated the quality of this evidence as low.

Clinical considerations
Poverty, food insecurity, culture, customs and limited education attainment can affect the choice and availability of iron-rich foods.543 Many refugees and some immigrants have ingested iron-insufficient diets for an extended period and do not recognize symptoms of iron deficiency. Culturally appropriate nutritional assessment and counselling by a registered dietician, when available, can identify specific nutrition problems and support appropriate change.531

What are the potential implementation issues?
Accessibility issues include language barriers, lower levels of education, low levels of ongoing support and follow-up, and

Table 17C: Summary of findings for iron supplements compared with placebo for women of reproductive age

<table>
<thead>
<tr>
<th>Patient or population:</th>
<th>Women of reproductive age (19–44 yr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Setting:</td>
<td>Factory in China (workers in cotton mill)</td>
</tr>
<tr>
<td>Intervention:</td>
<td>Oral iron supplementation</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Absolute effect</th>
<th>Placebo group</th>
<th>Difference with supplementation (95% CI)</th>
<th>Relative effect (95% CI)</th>
<th>No. of participants (studies)</th>
<th>GRADE quality of evidence</th>
<th>Comments (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemoglobin, g/L</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>p &lt; 0.001 NNT 2 (2–3)</td>
</tr>
<tr>
<td></td>
<td>115 (baseline)</td>
<td>15 higher (10.53–19.47 higher)</td>
<td>NA</td>
<td>80 (1)</td>
<td>Moderate</td>
<td></td>
</tr>
<tr>
<td>Energy expenditure at work, kJ/d (mean follow-up 12 wk)</td>
<td>4162 (baseline)</td>
<td>538 lower (862.34–213.66 lower)</td>
<td>NA</td>
<td>80 (1)</td>
<td>Moderate</td>
<td>p &lt; 0.001</td>
</tr>
<tr>
<td>Productivity efficiency, yuan/MJ (mean follow-up 12 wk)</td>
<td>1.86 (baseline)</td>
<td>0.33 higher (0.18–0.46 higher)</td>
<td>NA</td>
<td>80 (1)</td>
<td>Low</td>
<td>p &lt; 0.001 NNT 4 (3–8)</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NA = not applicable; NNT = number needed to treat.
access to supplements. Gastrointestinal side effects, changes in stool colour and three times daily dosing have been shown to reduce adherence rates.

Recommendations of other groups

The US Preventive Services Task Force recommends screening for iron-deficiency anemia in children 6–12 months of age and pregnant women who are at risk for iron deficiency. The US Centers for Disease Control and Prevention recommend routine screening for all children and women of reproductive age, citing the linked risks of iron-deficiency anemia in pregnancy affecting both women and infants. The World Health Organization recommends screening for at-risk women and children. Our recommendations highlight the importance of screening for iron-deficiency anemia in immigrant children and women of child-bearing age.

Take-home messages

- Immigrants and refugees coming from regions with limited access to iron-rich foods, higher rates of infectious disease and higher parity are at risk for iron deficiency.
- To improve their cognitive development, growing children aged one to four years should be screened for iron deficiency by means of hemoglobin measurement.
- To improve hemoglobin levels and work productivity, immigrant and refugee women of reproductive age should also be screened by means of hemoglobin measurement.

18. Dental disease

Dental caries affect virtually 100% of adults and 60%–90% of children worldwide, and periodontal disease is found in 5%–20% of most adult populations.546 Both are preventable chronic infectious diseases influenced by sociobehavioural, economic and environmental risk factors. In addition to oral pain, infection, tooth loss and associated dysfunction, chronic oral conditions can have a profound effect on general health and quality of life. Pain and disability associated with poor oral health can compromise a person’s ability to eat properly, which affects nutrition status and body weight of both children and older adults.546 We conducted an evidence review to guide primary care practitioners in the early detection, prevention and treatment of common oral conditions for newly arriving immigrants. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on screening for and treatment of dental and periodontal disease are found in Box 18A.

Methods

We used the 14-step method developed by the Canadian Collaboration for Immigrant and Refugee Health55 (summarized in section 3 of this article, above). We considered the epidemiology of oral disease in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE, Embase, CINAHL, HealthSTAR, the Cochrane Library and other sources from Jan. 1, 1996, to Jan. 1, 2010. Detailed methods, search terms, case studies, clinical considerations and research needs can be found in the complete evidence review for dental disease (Appendix 16, available at www.cmaj.ca/lookup/suppl /doi:10.1503/cmaj.090313/-/DC1).

Results

We found no systematic reviews or evidence-based guidelines specific to immigrants and refugees. The next search produced 35 relevant manuscripts, and five high-quality systematic reviews met the inclusion criteria.547–549 The focused guideline searches resulted in five relevant evidence-based, consensus-driven guidelines. The search for preventive and restorative interventions for dental referral yielded 10 high-quality systematic reviews.547–549 Fifty-six articles addressing prevalence, burden of disease, disease risk and access to care in immigrant populations were selected for detailed review.

What is the burden of oral disease in immigrant populations?

Immigrant adolescent children were five times more likely to have dental caries than children born in Canada, and 22.9% required restorative dental care for carries, compared with only 3.5% of those born in Canada.540 Although levels of oral disease in immigrants decreased relative to length of time in Canada, immigrant adolescents continued to be at a disadvantage for dental caries, gingivitis and level of oral hygiene when compared with their Canadian counterparts.

Epidemiologic information from the World Health Organization shows that development of caries is on the rise in developing countries in Africa and Asia. The increased consumption of refined sugar and inadequate exposure to the topical fluorides available in toothpastes and professionally applied fluoride products available in developed nations contribute to high rates of disease.546

Does screening decrease morbidity from common oral diseases?

Screening

The mouth should be examined using a tongue depressor to determine swelling, bleeding gums, loose teeth or holes in teeth, odour, mouth ulcers and sores. A systematic review and randomized controlled trial provide evidence that...
physicians can screen preschool children for dental caries with a high degree of accuracy.\textsuperscript{62} The evidence supports the effectiveness of prevention through application of sealants in children (Table 18A).\textsuperscript{657}

Relative benefits and harms of treatment
The management of underlying causes of dental pain by dentists is effective for prevention and management of dental disease.\textsuperscript{646} Potential harms include the cost to patients to access dental care and adverse reactions to treatment. Nonsteroidal anti-inflammatory drugs (NSAIDs) manage oral pain effectively (Table 18B).\textsuperscript{658} Antibiotics should be prescribed only in the presence of concomitant systemic symptoms, such as lymphadenopathy, fever, and associated cellulitis.\textsuperscript{661} Single-dose studies showed no difference in short-term adverse effects between oral ibuprofen and placebo.\textsuperscript{656} In patients given NSAIDs over longer periods, adverse reactions included abdominal pain, diarrhea, edema, dry mouth, rash, dizziness, headache and fatigue. These are generally considered to be mild to moderately severe.\textsuperscript{653}

Clinical considerations
What are potential implementation issues?
Lower income and immigrant status are both associated with fewer visits for preventive dental care.\textsuperscript{663} Language barriers reduce access to services and quality of care.\textsuperscript{665} Experiential influences (fear of dentists, history of inadequate care, embarrassment about oral condition) are likely hindrances for individuals who require professional dental care. Financial barriers decrease access to dental care. When nondentists refer on the basis of an oral health screening, people are twice as likely to go to the dentist.\textsuperscript{62} However, active referral (e.g., specific clinics identified, a patient information notice) is warranted.\textsuperscript{666}

Financial coverage of dental care
During the settlement period, Convention Refugees, refugee claimants and other protected people are eligible to apply for Interim Federal Health Program coverage for dental care. Details of services covered, the application process and a hand-

<table>
<thead>
<tr>
<th>Table 18A: Summary of findings for use of sealant to prevent carious lesions</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Patient or population:</strong> Children and adolescents (aged 6–19 yr) with carious lesions on permanent teeth</td>
</tr>
<tr>
<td><strong>Setting:</strong> Dental clinics in Brazil, United States, Canada, Zimbabwe</td>
</tr>
<tr>
<td><strong>Intervention:</strong> Sealants</td>
</tr>
<tr>
<td><strong>Comparison:</strong> No sealant</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Absolute effect</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Outcome</strong></td>
</tr>
<tr>
<td>Progressing lesions</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NNT = number needed to treat; RR = risk ratio.

<table>
<thead>
<tr>
<th>Table 18B: Summary of findings for use of NSAIDs as treatment for pain and swelling of dental origin</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Patient or population:</strong> Adults with pain (without localized swelling) of dental origin and no systemic symptoms (such as fever or lymphadenopathy)</td>
</tr>
<tr>
<td><strong>Setting:</strong> Dental offices and hospitals in United States, United Kingdom</td>
</tr>
<tr>
<td><strong>Intervention:</strong> NSAIDs</td>
</tr>
<tr>
<td><strong>Comparison:</strong> Placebo</td>
</tr>
<tr>
<td><strong>Source:</strong> Sutherland SE, Matthews DC. Emergency management of acute apical periodontitis in the permanent dentition: a systematic review of the literature. J Can Dent Assoc 2003; 69:160.\textsuperscript{666}</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Absolute effect</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Outcome</strong></td>
</tr>
<tr>
<td>Mean pain relief (100-mm visual analogue scale) at 24 h</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NA = not applicable; NSAID = nonsteroidal anti-inflammatory drug.

*Data in source article were insufficient to allow estimation of 95% CI.
book for health care providers are available through the Citizenship and Immigration Canada website (www.cic.gc.ca). In addition to the provincial public dental programs, there are low-cost or no-cost clinics in several of the larger cities. Information on many of these programs and clinics is available from the website of the Federal, Provincial and Territorial Dental Working Group (www.fptdwg.ca) and the Canadian Association of Public Health Dentistry (www.caphd-acsdp.org). In the absence of comprehensive dental services for marginalized populations, physicians’ involvement in oral health care is increasing, and the application of fluoride varnish by primary care practitioners shows great promise.

**Recommendations of other groups**

The Canadian Collaboration for Clinical Practice Guidelines in Dentistry, in a 2003 guideline, suggested that the most efficacious method to reduce tooth pain with or without localized swelling (when dental therapy cannot be started immediately) is with NSAIDs. Other guideline development groups have reported moderate- to high-quality evidence to support recommendations for the application of fluoride varnish to teeth of children at high risk for caries and the recommendation that teeth be brushed twice daily with toothpaste containing 1000 ppm fluoride. Our recommendations focus on screening all immigrants for dental pain and highlight the benefit of practitioner referrals for those with dental pathology.

**Take-home messages**

- Dental pain can be reduced if physicians ask whether patients have problems with their mouth, teeth or dentures. NSAIDs can be used effectively to treat dental pain.
- Migrants arriving from countries with limited dental care and where diets are high in sugar are at the highest risk for disease.
- Screening and referral for dental disease can facilitate treatment and prevention of dental disease.
- Patients are twice as likely to go for dental treatment when they are actively examined and referred by a physician.
- Tooth-brushing twice daily with fluoridated toothpaste is effective in reducing the risk of dental decay.

19. Vision health

Together, blinding disorders are the seventh leading cause of burden of disease, ahead of diabetes and cancer.\textsuperscript{571,572} Vision loss can limit the ability to work, drive and complete other activities of daily living. Even modest visual impairment (visual acuity < 6/12) is associated with substantial morbidity\textsuperscript{573–575} Uncorrected refractive error is the leading cause of visual impairment worldwide.\textsuperscript{576} Vision loss from most of these conditions is largely preventable with timely diagnosis and treatment. Even age-related macular degeneration is yielding somewhat to interventions.\textsuperscript{577,578} However, access to even basic vision care is often limited in developing countries.\textsuperscript{579} In light of evidence that source countries for new immigrants and refugees to Canada have higher burdens of vision loss,\textsuperscript{61} we reviewed evidence on prevalence, screening of asymptomatic populations, treatment effectiveness, population-specific concerns and implementation. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on vision health are outlined in Box 19A.

Methods

We used the 14-step method developed by the Canadian Collaboration for Immigrant and Refugee Health\textsuperscript{16} (summarized in section 3 of this article, above). We defined clinical preventive actions (interventions) and relevant clinical outcomes, concentrating on the body of evidence for screening tests for vision loss. We searched MEDLINE, Embase, CINAHL, the Cochrane Library and other sources from Jan. 1, 1996, to Jan. 1, 2010. Detailed methods, search terms and references, case studies and clinical considerations can be found in the complete evidence review for vision health (Appendix 17, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

Results

Our search did not yield any systematic reviews or guidelines related to vision screening tests in immigrants or refugees. However, we identified 23 systematic reviews and guidelines related to screening tests for vision loss in the general population. Our review identified four clinical trials of screening tests for vision loss in community-dwelling elderly people.\textsuperscript{680} These studies demonstrated no benefit, but none of them could be generalized to newly arrived immigrants and refugees. Eight guidelines for adults and one article focusing on amblyopia in children were identified; all recommended vision screening tests in varying forms and frequency.

What is the burden of vision loss in immigrant populations?

Visual acuity is the single most important indicator of ocular health. Decreased presenting vision is most frequently due to uncorrected refractive error but can signify underlying sight-threatening eye disease. International data\textsuperscript{61} suggest that vision loss and undiagnosed sight-threatening eye disease should be more common in new immigrants and refugees to Canada who originate from developing countries, but we could not find any Canadian data to confirm this assertion. Much of the high prevalence of blindness in developing countries can be attributed to disparities in access to care.\textsuperscript{579} The World Health Organization has estimated that 80% of blindness in developing countries could be avoided through cost-effective prevention and treatment. Glaucoma, the leading cause of irreversible blindness worldwide,\textsuperscript{61} reflects the global disparity in access to care, with only 10% of cases of open-angle glaucoma diagnosed in developing countries compared with 50% in developed countries.\textsuperscript{581} Similar global disparities in access to care are anticipated for other common causes of vision loss.\textsuperscript{579}

In the United States, minority populations have higher levels of vision loss and undiagnosed eye disease\textsuperscript{582–585} than does the general population. Numerous factors contribute to this disparity, including socioeconomic status, access to care, societal conventions, and physiologic and anatomic differences.\textsuperscript{584} These factors, in turn, influence the prevalence of vision loss from cataract, diabetic retinopathy, glaucoma and

\begin{boxedtext}
Box 19A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: vision health

Perform age-appropriate screening for visual impairment.*
If presenting vision < 6/12 (with habitual correction in place), refer patients to an optometrist or ophthalmologist for comprehensive ophthalmic evaluation.

Basis of recommendation

Balance of benefits and harms
Uncorrected refractive error, the most common cause of visual impairment, is amenable to correction with eyeglasses (number needed to screen to find one person with vision worse than 6/15 or 20/50 due to uncorrected refractive error = 19). Prevalence of uncorrected refractive error in immigrant populations is higher than in the general population; however, economic and cultural barriers could reduce rates of referral and use of corrective eyeglasses. Harms are minimal and can include out-of-pocket costs.

Quality of evidence

Very low

Values and preferences

The committee attributed more value to ensuring that visual acuity is adequate for daily functioning and employment and to detecting serious underlying ocular disease. The committee attributed less value to the burden of screening and the cost of eyeglasses.

*Visual acuity should be measured with distance glasses or contact lenses in place if worn habitually. Age-appropriate measurement in children is required at 0–3 months (infant should react to light), at 6–12 months (baby’s eyes should fix and follow light) and at 3–5 years (child should use visual acuity chart where possible). Additional screening manoeuvres are useful for children: at each screening interval, assess for red reflex and inspect external ocular structures. For patients 6 months and older, also assess for strabismus.
\end{boxedtext}
uncorrected refractive error. Recent immigration has been identified as a risk factor for vision loss, the most common cause of which is uncorrected refractive error. The prevalence of vision loss (< 6/15) due to uncorrected refractive error is estimated at 5.4% for those 12 and older in the general population and substantially higher in minority populations.

Although trachoma, onchocerciasis (river blindness) and vitamin A deficiency figure prominently as causes of blindness in some low-income countries, these conditions have not been reported among immigrants and refugees in Canada. Anecdotal evidence suggests that the rare cases in immigrants and refugees to Canada most frequently take the form of subconjunctival scarring from cicatricial trachoma (noninfectious) evident on eversion of the upper eyelids. Typically, the threat to vision posed by these conditions in asymptomatic patients dissipates harmlessly on arrival to Canada through a vitamin A–sufficient diet or by breaking the cycle of re-infection that underlies the decades-long descent into blindness from trachoma or onchocerciasis.

Do screening tests and treatment for eye disease decrease morbidity?

Screening
Measuring visual acuity is a simple manoeuvre that can be carried out accurately (sensitivity of 94% and specificity of 89%) in a primary care setting. When measuring vision in immigrants and refugees, a standardized chart with the tumbling E or the Landholt C optotypes could be helpful, because familiarity with an alphabet or numbers is not required. In some cases, a cut-out “E” or “C” for the reader to orient to the chart will simplify communication. The ETDRS-type of visual acuity chart is the preferred chart layout and has several advantages, including five optotypes on each line. However, an inexpensive Snellen chart, which uses ambient room light, is adequate for screening purposes. A pinhole viewing device can be added as a simple adjunct to determine whether the visual impairment is due to refractive error (sensitivity 79%, specificity 98%).

Relative benefits and harms of treatment
We did not find direct evidence that routinely screening and treating immigrant children in primary care for visual impairment results in improved visual acuity. However, a randomized clinical trial has shown that intensive screening procedures, compared with usual vision surveillance, leads to improved visual acuity. Amblyopia and its leading causes (strabismus, astigmatism and anisometropia) are reversible if diagnosed and treated early after onset within the first seven or eight years of life, after which reversibility is minimal.

For adults, we focused our effectiveness synthesis on the prevalence of vision loss from uncorrected refractive error, which is amenable to simple correction with eyeglasses. We were unable to identify any evidence to estimate the effect of adherence to recommendations for an eye examination on whether subsequent prescriptions for eyeglasses are filled. Using the GRADE rating system, we rated the quality of this evidence as very low. Our evidence review found no data on harm to the patient for measuring vision (Table 19A).

Clinical considerations

What are the potential implementation issues?
Comprehensive ophthalmic examination is not covered for those aged 18 to 64 in any Canadian province. However, specific nonrefractive indications for performing an evaluation — such as being at increased risk for glaucoma, screening for diabetic retinopathy or excluding clinically evident cataract as a cause of vision loss — are typically covered by provincial health plans. Convention refugees, covered under the Interim Federal Health Program, receive coverage for cor-

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Table 19A: Summary of findings for vision screening and correction to reduce visual impairment

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Absolute effect</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Outcome</strong></td>
<td><strong>Presenting vision, % (95% CI)</strong></td>
<td><strong>Best corrected vision, % (95% CI)</strong></td>
<td><strong>Relative effect (95% CI)</strong></td>
<td><strong>No. of participants (studies)</strong></td>
</tr>
<tr>
<td>Visual impairment</td>
<td>6.4 (6.0–6.8)</td>
<td>1.1 (0.7–1.5)</td>
<td>Difference 5.3 (4.9–5.7)</td>
<td>14 203 (1)</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NNS = number need to screen to find one person with vision worse than 6/15 or 20/50 because of uncorrected refractive error.

*Presenting visual impairment is based on vision with habitual correction, if worn, in place. Best-corrected visual impairment is based on vision with best possible refractive correction in place.

†Acceptance of correction through eyeglasses can only be inferred. The study was a large population-based examination survey, not a treatment study. Proportion of participants who would benefit from correction using eyeglasses was estimated by the change in prevalence of visual impairment based on refraction compared with presenting vision. In practice, the NNS to improve vision would be greater because not everyone whose vision would improve to better than 20/50 would accept the cost and inconvenience of eyeglasses.
rection with eyeglasses and other urgent care for up to one year after arrival.

Beyond issues of financial access, age and sex can be influential. Age can influence the decision to seek care or to treat, a decision that is sometimes beyond the control of elderly patients with vision loss. Female patients appear to face greater barriers to assessment and treatment, and sex discrimination is more influential in low- and middle-income countries than in high-income countries.59 Finally, stigma associated with wearing glasses can also influence eye care for refugees and immigrants to Canada. The desire to conform with societal norms and beauty standards is often stronger than the desire for treatment.596

**Recommendations of other groups**

The recommendations of other groups regarding vision health are presented below. Our recommendations highlight the importance of screening all immigrants for visual impairment.

**Screening tests for visual impairment in children**
The Canadian Task Force on the Periodic Health Examination597,598 (now the Canadian Task Force on Preventive Health Care) and the US Preventive Services Task Force recommend screening tests for visual impairment and strabismus for children younger than five years. Additional manoeuvres recommended for children include assessing for a red reflex and inspecting the external eye beginning with newborns.599

**Screening tests for visual impairment in adults**
For adults, both the Canadian Task Force on Preventive Health Care600 and the US Preventive Services Task Force601 recommend screening tests for visual impairment by primary care practitioners but suggest limiting these tests to people older than 65 years of age. Since these influential guidelines were published more than a decade ago, experts have recognized that the epidemiology of “best corrected” visual acuity on which they are based does not address uncorrected refractive error,602 now recognized as the leading cause of vision loss worldwide.598

**Other conditions**
In the case of glaucoma and diabetic retinopathy, patients do not typically present with vision impairment until damage has become irreversible. The Canadian Task Force on Preventive Health Care and the US Preventive Services Task Force have presented compelling evidence supporting recommendations for referral of asymptomatic adults at high risk (those older than 65 years, blacks older than 40 years, those with a first-degree relative who has glaucoma and myopia exceeding –6 diopters) for glaucoma testing.600,603 Similarly, the Canadian Diabetes Association clinical practice guidelines make a compelling case for periodic screening tests for diabetic retinopathy in people with diabetes.604

**Take-home messages**
- Loss of vision and undiagnosed sight-threatening eye diseases are more common among new immigrants and refugees than in the general population.
- New immigrants should be screened for vision loss within their first year in Canada and should be referred to an optometrist or ophthalmologist if their presenting vision (with habitual correction in place) is less than 6/12 (i.e., less than 20/40).
- Referral for assessment is also warranted in the presence of other risk factors for blinding eye disease, including diabetes, age older than 65 years (or age older than 40 years for those of black ethnicity), glaucoma in a first-degree relative and myopia exceeding –6 diopters.
- Regionally prominent “tropical” eye diseases, such as onchocerciasis (river blindness), active trachoma and xerophthalmia, have not been reported in immigrants or refugees to Canada.
- Asymptomatic forms of these diseases should resolve or stabilize away from endemic conditions after arrival in Canada.

20. Contraception

Reproductive and sexual health care is central to improving the health of women and is a basic human right. Couples and individuals have the right to “decide … the number, spacing and time of their children and to have the information and the means to do so.” Worldwide, about 120 million women have an unmet need for contraception. As a consequence, many experience unintended pregnancies, which can be associated with such negative outcomes as abortion, failure to adopt healthy pregnancy recommendations and limitation of women’s ability to achieve educational, employment and economic goals. We reviewed the evidence for the effectiveness of screening and counselling for unmet contraceptive needs for immigrant populations. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health related to contraception are outlined in Box 20A.

Methods

We used the 14-step method developed by the Canadian Collaboration on Immigrant and Refugee Health (summarized in section 3 of this article, above). We considered the epidemiology of unmet needs for contraception in immigrant populations and defined clinical considerations and potential key clinical actions. We searched MEDLINE, Embase, CINAHL, POPLINE, the Cochrane Library and other sources from Jan. 1, 1996, to Jan. 1, 2010. Detailed methods, search terms, case studies and clinical considerations, as well as the complete reference list, can be found in the evidence review for contraception (Appendix 18, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313//DC1).

Results

We found no systematic reviews or guidelines for immigrant screening related to contraception. Our search on contraceptive counselling in the general population yielded 789 titles. We retained four reviews, two guidelines and six primary studies as the basis for our evidence. In our search to update the reference systematic reviews, we found 203 new studies. Our evidence comes from studies in both general and high-risk populations of sexually active women from developed, low-income and middle-income countries.

What unmet contraceptive needs affect immigrant and refugee women?

Most immigrants to Canada come from developing countries, where unmet needs for contraception range from 5% to 40%. Rates are highest in sub-Saharan Africa, among young women and women who have had more than three births. Rural, uneducated and poor women are also at higher risk. Worldwide, more than one-third of pregnancies are unintended, and 50% of these end in abortion.

Studies from Europe suggest that immigrant and refugee women have higher rates of unintended pregnancy and abortion than native-born women, and more than half of immigrants who seek abortion are not using any form of contraception. In a recent US study, foreign-born and native-born women had similar abortion rates, although rates were higher for visible minority and poorer women, characteristics that are associated with immigrant status in Canada. Immigrant and refugee women are also less likely than the general population to seek counselling for family planning.

Box 20A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: contraception

| Screen immigrant women of reproductive age for unmet contraceptive needs. |
| Provide culturally sensitive, patient-centred contraceptive counselling to decrease unintended pregnancy and promote patient satisfaction. |

Basis of recommendation

| Balance of benefits and harms |
| Contraceptive counselling led to improved patient satisfaction (number needed to treat [NNT] 3, 95% confidence interval [CI] 2–5) and improved continuation rates (NNT 4, 95% CI 3–7). Evidence that in-depth counselling reduces unintended pregnancy rates shows some uncertainty (relative risk 0.47, 95% CI 0.16–1.34); however, the committee judged that contraceptive continuation rates are an acceptable surrogate for unintended pregnancy rates. There is a high prevalence of unmet need for contraception among immigrant and refugee women (5%–40%). Harms were minimal. No data were available on couple or family discord. |

Quality of evidence

| Moderate |

Values and preferences

| The committee attributed more value to supporting informed choice to meet future family needs and the woman’s personal needs (empowerment) and less value to concern about causing couple and family discord. |
than those who received regular counselling (Table 20B).64 In another randomized trial, which provided moderate-quality evidence for individualized counselling and follow-up, women attending a sexually transmitted disease clinic who had individualized contraceptive counselling and follow-up had higher rates of effective contraceptive use at four and eight months than those receiving regular information on options, although the effect diminished over time and by 12 months was not statistically significant.61 The cohort study by Nawar and colleagues61 of 590 Egyptian women provided moderate-quality evidence that women receiving client-centred care are more satisfied than those receiving usual care (RR 2.17, 95% CI 1.76–2.68). We found no data on harm of screening or contraceptive counselling, other than personnel and costs.

Evidence from systematic reviews, observational studies and guidelines suggests that a client-centred approach, giving women their method of choice, providing the contraceptive method on site and having a good personal relationship, improve patient satisfaction and contraception continuation rates.60,61,62 Provider pressure to adopt a method has been shown to be associated with method discontinuation. A randomized trial demonstrated better knowledge improvement when a simpler rather than more detailed chart of contraceptive effectiveness was used.64 High-quality contraceptive care respects each woman’s human and reproductive rights and enables her to make an informed contraceptive choice consistent with her personal values, needs and beliefs.62,63 (Tables 20A and 20B).

### Clinical considerations

#### Which women might need special consideration for contraceptive counselling?

With migration to a more stable environment, fertility rates of refugee women sometimes increase as they choose to rebuild families. Refugee women who have been in their country of resettlement for less than three months appear to have the highest levels of fertility of all resettled populations, although whether these pregnancies are intended or unintended is unknown.67 Alternatively, a desire to prevent pregnancy could increase if women perceive uncertainty and instability with migration.68,69 Contraceptive needs can fluctuate because of family reunification and an ability to visit partners in their home countries. Pregnancy intention, contraceptive options and emergency contraception should therefore be discussed early in resettlement and should be reassessed as circumstances change.

Unmarried women could be vulnerable to unintended pregnancy if cultural proscriptions on premarital sex prevent them from identifying and seeking support for their contraceptive needs. The most common reason for not using contraception among sexually active, unmarried women surveyed across all developing regions was a low perceived risk of pregnancy because of infrequent sexual activity.66

Adolescent risk-taking and experimentation with sexuality put teenagers at risk for unintended pregnancy. Most young people become sexually active between 15 and 19 years of age. In Canada, teenagers who are recent immigrants have lower rates of sexual activity and pregnancy than Canadian-

### Table 20A: Summary of findings for contraceptive screening and counselling of women of reproductive age with unmet contraceptive needs

<table>
<thead>
<tr>
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<tbody>
<tr>
<td></td>
<td>Absolute effect</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Risk for control group</td>
<td>Difference with screening and counselling (95% CI)</td>
<td>Relative effect (95% CI)</td>
</tr>
<tr>
<td>Informed choice** (received assistance in selecting method of contraception)</td>
<td>40 per 1000†</td>
<td>90 more per 1000 (22 to 233 more per 1000)</td>
<td>RR 3.25 (1.55–6.82)†</td>
<td>480 (1)</td>
</tr>
<tr>
<td>Services per visit for family planning**</td>
<td>21 per 1000</td>
<td>104 more per 1000 (66 to 160 more per 1000)</td>
<td>RR 5.96 (4.12–8.64)</td>
<td>2678 (1)</td>
</tr>
<tr>
<td>Harms</td>
<td>No data</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NNT = number needed to treat; RR = risk ratio.

*The committee judged that the results of these studies of populations in Guatemala and Bolivia would apply to immigrant and refugee populations.
†Moderate risk for control group taken from study and considered reasonable for refugee and immigrant women in Canada.
‡Large effect shown for assisting with decisions and referral for a method (RR 2).
§Pre–post test design: intervention-trained providers were to use screening tool to determine unmet contraceptive needs for women visiting clinic. Study occurred in a conservative environment with reduced access to family planning services; therefore, there were some concerns about generalizability.
**Pre–post test design: use of screening to promote provision of family planning services.
born teenagers. However, other countries of resettlement show the opposite pattern. Like North American–born teenagers, those from the developing world are interested in discussing sexual health concerns with health care providers, although needing to raise the topic, confidentiality concerns and parental presence often discourage this discussion. Adolescent newcomers sometimes experience conflict between their families’ attitudes toward teenagers’ sexuality and attitudes in their country of resettlement.

What social and cultural factors influence contraceptive counselling?

Women arriving from developing countries might have insufficient knowledge about reproduction and contraception to make an informed decision about family planning. Increased education and knowledge about reproduction correlate with more positive attitudes and increased use of and adherence to contraception. Women of all cultures use social networks for much of their sexual health information, which might be inaccurate or incomplete.

Cultural attitudes toward pregnancy and family planning vary. Providential (“children are God’s will”) or pronatalist cultures discourage pregnancy prevention. In some cultures, women who bear many children are highly esteemed. Religious beliefs about the acceptability of contraceptive practices also influence some women. Contraception used to space births is acceptable in most religions. Among women of any particular faith, attitudes toward contraception vary widely. Health care providers should avoid assumptions and should assess each woman or couple individually. Longer residence in the host country, educational and professional attainment, and youth favour positive attitudes toward modern contraception.

A woman might not perceive herself to be the decisionmaker for contraception, but she could be strongly influenced by her spouse, mother-in-law, sex role and religious beliefs. Worldwide, 11%–12% of married women do not use contra-

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**Table 20B: Summary of findings for contraceptive counselling of women of reproductive age with unmet contraceptive needs**

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Absolute effect</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Risk for control group</td>
<td>Difference with contraceptive counselling (95% CI)</td>
<td>Relative effect (95% CI)</td>
<td>No. of participants (studies)</td>
<td>GRADE quality of evidence</td>
<td>Comments (95% CI)</td>
</tr>
<tr>
<td>Continuation* (mean follow-up 12 mo)</td>
<td>Medium-risk population*</td>
<td>560 per 1000</td>
<td>258 more per 1000 (151 to 392 more per 1000)</td>
<td>RR 1.46 (1.27–1.7)</td>
<td>350 (1)</td>
<td>High</td>
</tr>
<tr>
<td>Effective contraceptive use** (self-reported use of effective method for &gt; 75% of coitus or sexual abstinence; mean follow-up 12 mo)</td>
<td>High-risk population</td>
<td>260 per 1000</td>
<td>57 more per 1000 (10 fewer to 143 more per 1000)</td>
<td>RR 1.22 (0.96–1.55)</td>
<td>632 (1)</td>
<td>Moderate†</td>
</tr>
<tr>
<td>Satisfaction† (no. of people endorsing 10–13 of 13 items related to satisfaction with services; mean follow-up 7 mo)</td>
<td>Medium-risk population</td>
<td>270 per 1000</td>
<td>316 more per 1000 (205 to 454 more per 1000)</td>
<td>RR 2.17 (1.76–2.68)</td>
<td>590 (1)</td>
<td>Low</td>
</tr>
<tr>
<td>Unintended pregnancy*** (self-reported; mean follow-up 12 mo)</td>
<td>Low-risk population</td>
<td>30 per 1000</td>
<td>5 fewer per 1000 (11 fewer to 3 more per 1000)</td>
<td>RR 0.83 (0.63–1.09)</td>
<td>632 (1)</td>
<td>Low‡</td>
</tr>
<tr>
<td></td>
<td>Medium-risk population</td>
<td>104 per 1000</td>
<td>18 fewer per 1000 (38 fewer to 9 more per 1000)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NNT = number needed to treat; RR = risk ratio.

*Control-group risk of 56% continuation for medroxyprogesterone acetate from US national survey of family growth.

†Serious limitations because 30% lost to follow-up, self-reported unintended pregnancy rate, only 45% of women invited agreed to participate, women who declined were older, and method of randomization and blinding was not described.

‡Unintended pregnancy data were rated as imprecise because RR was 0.47 (95% CI 0.00–1.34).
Contraception because of opposition from one or more influential parties. In sub-Saharan Africa, this figure is 23%,628 Recognition of the partner’s influence and his involvement, where appropriate, are important in counselling and supporting women’s choices. Some men consider contraception to be their spouse’s responsibility, but often this responsibility is shared, and involvement in counselling may be welcomed. In a population-based study of six African countries,629 women with supportive male partners were more likely to use modern contraceptives. In some traditional cultures, fathering many children is a sign of masculinity; however, perceived economic advantages of smaller families and better future opportunities for children can encourage men’s support for contraception.

**How acceptable are specific contraceptive methods?**

Contraceptive use is increasing worldwide, but the mix and acceptability of contraceptive methods vary (Table 20C).626 The average rate of contraceptive use by married women from low- and middle-income countries is 60%, with highest rates in Latin America and the Caribbean and much lower rates in sub-Saharan Africa.626 Effectiveness and freedom from adverse effects are the most important characteristics influencing contraceptive choice. Resistance to use modern methods (e.g., oral contraceptives) can be influenced by culture-specific fear of adverse effects. For example, although many North American women choose to eliminate menstrual bleeding, those from African cultures often prefer monthly bleeding. Spotting and bleeding associated with some methods are problems for women who have religious and cultural restrictions on intercourse or other activities related to bleeding.640

In many African and Latin American populations, condom use has connotations of infidelity, promiscuity, extramarital relationships or sexually transmitted infection. Although used by only 1%–2% of North American women aged 15–44 years, the intrauterine device is the most commonly used and most effective reversible contraceptive method worldwide.626,630,631

In sub-Saharan Africa, about 25% of women practising contraception use injectables. Breastfeeding is commonly used in low- and middle-income countries to control fertility.

Worldwide, many couples depend on tubal ligation for contraception. Some religions prohibit sterilization, and in many countries, tubal ligation is illegal or spousal consent is required. Most women from developing countries are unaware of emergency contraception. Availability of legal abortion varies greatly around the world, as does its cultural acceptance. Abortion rates are highest in Eastern Europe and central Asia, where contraceptive options have been limited and abortions freely available.632

**What are the medical considerations?**

Condom use should be encouraged for women at risk for sexually transmitted infections, irrespective of additional contraceptive use. More common in immigrant and refugee women, HIV infection and sickle cell anemia bear special consideration. Guidelines suggest that hormonal methods can be safely used by HIV-positive women. Although serum levels of contraceptive hormones can be reduced by some antiretroviral medications, the clinical implications are unknown.625 Intrauterine devices are considered safe for women with HIV who are immunocompetent.629 Pregnancy is risky for women with sickle cell anemia. Although combination hormonal contraceptives are considered reasonably safe for women with sickle cell anemia,625 progestin-only contraceptives such as medroxyprogesterone have the added benefit of reducing sickle cell crises.633,634

**What are the potential implementation issues?**

Language barriers and lack of familiarity with the Canadian

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**Table 20C: Regional prevalence of use of main contraceptive methods, from most effective (sterilization) to least effective (periodic abstinence)**626

<table>
<thead>
<tr>
<th>Region</th>
<th>Any method</th>
<th>Sterilization</th>
<th>Intrauterine device</th>
<th>Implant or injection</th>
<th>Oral contraceptives</th>
<th>Condom</th>
<th>Withdrawal</th>
<th>Periodic abstinence</th>
</tr>
</thead>
<tbody>
<tr>
<td>More developed regions</td>
<td>67.4</td>
<td>13.1</td>
<td>9.4</td>
<td>1.0</td>
<td>16.5</td>
<td>13.9</td>
<td>6.8</td>
<td>4.3</td>
</tr>
<tr>
<td>Less developed regions</td>
<td>62.4</td>
<td>24.0</td>
<td>16.5</td>
<td>3.7</td>
<td>7.2</td>
<td>4.4</td>
<td>2.3</td>
<td>3.4</td>
</tr>
<tr>
<td>Sub-Saharan Africa</td>
<td>21.5</td>
<td>1.5</td>
<td>0.5</td>
<td>6.2</td>
<td>4.2</td>
<td>1.8</td>
<td>1.2</td>
<td>3.8</td>
</tr>
<tr>
<td>Asia overall</td>
<td>67.9</td>
<td>27.0</td>
<td>19.6</td>
<td>3.2</td>
<td>6.1</td>
<td>5.3</td>
<td>2.5</td>
<td>3.4</td>
</tr>
<tr>
<td>East Asia</td>
<td>87.6</td>
<td>34.3</td>
<td>40.4</td>
<td>0.3</td>
<td>3.5</td>
<td>6.9</td>
<td>0.0</td>
<td>1.2</td>
</tr>
<tr>
<td>South Asia</td>
<td>54.2</td>
<td>29.4</td>
<td>3.5</td>
<td>1.9</td>
<td>6.0</td>
<td>5.0</td>
<td>3.1</td>
<td>4.9</td>
</tr>
<tr>
<td>West Asia/Middle East</td>
<td>54.5</td>
<td>3.3</td>
<td>15.4</td>
<td>0.7</td>
<td>8.9</td>
<td>5.3</td>
<td>14.9</td>
<td>2.6</td>
</tr>
<tr>
<td>Latin America and Caribbean</td>
<td>71.4</td>
<td>29.8</td>
<td>7.4</td>
<td>4.1</td>
<td>15.8</td>
<td>6.8</td>
<td>2.7</td>
<td>3.9</td>
</tr>
<tr>
<td>Eastern Europe</td>
<td>63.7</td>
<td>2.3</td>
<td>21.1</td>
<td>0.0</td>
<td>6.5</td>
<td>11.0</td>
<td>12.5</td>
<td>9.4</td>
</tr>
<tr>
<td>North America</td>
<td>73.0</td>
<td>32.5</td>
<td>1.9</td>
<td>3.6</td>
<td>17.9</td>
<td>11.9</td>
<td>2.9</td>
<td>1.4</td>
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</table>
health care system can limit immigrant and refugee women’s access to contraceptive care. In some communities of origin, the husband’s accompaniment or written consent is required to obtain contraception. In others, hormonal contraception is available in pharmacies without a prescription.

The caregiver’s sex is important for women in many cultures, particularly for refugee women, many of whom have a history of sexual assault and abuse. The advantages of ethnic and sex matching must be weighed against research suggesting a preference for “Canadian” professionals by immigrant and refugee women. Unnecessary medical barriers, such as examinations, blood tests and Papanicolaou smears, and a lack of culturally appropriate teaching aids are additional obstacles to contraceptive use.

The Interim Federal Health Program covers the cost of contraceptives for Convention refugees, refugee claimants and protected people. Newcomers without health insurance can be guided to publicly funded sexual health clinics that provide services and low-cost contraceptives, regardless of health insurance status. The Society of Obstetricians and Gynaecologists of Canada also has a Compassionate Contraceptive Assistance Program that assists women in financial need (www.sogc.org/compassionate/pdf/compassionate_form_e.pdf).

Recommendations of other groups

In 1996, the US Preventive Services Task Force recommended periodic counselling to prevent unintended pregnancy in teenagers and women of reproductive age, based on information taken from a sexual history. However, subsequent guidelines from this task force do not include this recommendation. The Institute for Clinical Systems Improvement gave prevention counselling for unintended pregnancy a level III recommendation (incomplete evidence and action left to judgment of group or clinician). Our recommendations highlight the importance of early screening of immigrant women for unmet contraceptive needs.

Take-home messages

- Screening for unmet contraceptive needs among immigrant women should begin soon after their arrival in Canada.
- Women from developing countries are often unaware of emergency contraception.
- Acceptability of contraception and preferences for particular methods vary across world regions (e.g., use of intrauterine devices is predominant in Asia and Latin America).
- In some communities, condoms have connotations of infidelity, promiscuity or sexually transmitted infection or are used only with nonmarital partners.
- Giving women their method of choice, providing the contraceptive method on site and having a good personal relationship improve outcomes.

21. Cervical cancer

Cervical cancer is one of the most preventable forms of cancer, yet deaths from cervical cancer persist among socially disadvantaged groups. The introduction of cervical cancer screening programs is associated with dramatic decreases in morbidity and mortality from cervical cancer. However, Canadian studies have documented significantly lower rates of screening among immigrants and refugees. Recent data from the Canadian Human Mortality Database (2000–2002) suggest a high mortality rate from cervical cancer among foreign-born women. We reviewed the evidence related to cervical cancer to guide practitioners in the prevention, early detection and treatment of cervical cancer for new immigrants. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on prevention of and screening for cervical cancer are outlined in Box 21A.

Methods

We used the 14-step method developed by the Canadian Collaboration for Immigrant and Refugee Health (summarized in section 3 of this article, above). We considered the epidemiology of cervical cancer in immigrant populations and defined clinical preventive actions (interventions), outcomes and key clinical questions. We searched MEDLINE, Embase, CINAHL, the Cochrane Library and other sources from Jan. 1, 1996, to Jan. 1, 2010. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for cervical cancer (Appendix 19, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.090313/-/DC1).

Results

We found no systematic reviews or guidelines specifically focused on immigrants. We identified 13 (out of 934) general population titles as relevant systematic reviews and, after critical appraisal, retained eight. Our search for new and pertinent studies identified four cohort studies relating to cervical cytology screening. A related study provided historical time trend mortality data related to cervical cancer screening programs. Finally, we identified one additional study reporting on the adverse events from human papillomavirus (HPV) vaccination in Australia and one meta-analysis reporting adverse pregnancy outcomes associated with treatment of cervical dysplasia. We retrieved 104 articles that addressed epidemiology, knowledge and compliance, and vaccination in immigrant populations.

What is the burden of cervical cancer in immigrant populations?

Data from the Canadian Human Mortality Database (non–age-standardized) showed that mortality rates from cervical cancer were 1.4 times higher among foreign-born women than among Canadian-born women (2000–2002). In the United States, the incidence of cervical cancer among Vietnamese-American women has been estimated at five times the incidence among white American women (incidence rate 43 per 100 000 v. 8.7 per 100 000). The Public Health Agency of Canada linked a sample of immigrants arriving to Canada (1980–1990) with cancer incidence data for the period (1980–1998) and found that overall in this cohort, foreign-born women had lower incidence rates of cervical cancer than Canadian-born women. However, rates among refugee women and Canadian-born women were similar, and older refugee women had higher rates of cervical cancer than Canadian-born women. Women who have never had cervical screening, or have not had cervical screening in the previous five years, account for 60%–90% of invasive cervical cancers overall. Also, several cross-sectional Canadian studies have...
documented lower rates of screening among immigrant populations.\(^6\) Foreign-born women aged 25–64 years, especially those born in Asia, are at higher risk of having never had a Papanicolaou (Pap) test (odds ratio 10.8).\(^6\)

Infection with HPV is strongly associated with cervical cancer. Prevalence estimates for HPV are particularly high for Africa (22.1%, 95% confidence interval [CI] 20.9%–23.4%) and Central America (20.4%, 95% CI 19.3%–21.4%).\(^6\) Women with HIV and women who have been victims of sexual trauma are at higher risk for HPV infection and cervical cancer.\(^6\) Refugee women in particular are disproportionately victims of sexual and sex-based violence, which can include rape, domestic violence and female genital mutilation.

**Does vaccination against HPV decrease morbidity and mortality?**

**Relative benefits and harms of vaccination**

A systematic review of vaccination against HPV showed reduction of high-grade cervical cancer lesions with no serious adverse events\(^6\) (Table 21A). We downgraded the quality of this evidence to moderate because of indirectness, since high-grade cervical lesions are considered surrogate outcomes for cervical cancer mortality. There were fewer than 15 anaphylactic events in a longitudinal study of more than one million doses of HPV vaccination among women in Australia.\(^6\) This adverse reaction occurred within 15 minutes of vaccination and was amenable to treatment.

**Does cervical cancer cytology screening and treatment decrease morbidity and mortality?**

**Screening tests**

Cervical cytology testing (liquid-based or conventional) is 60%–80% sensitive for high-grade lesions and 98% specific.\(^6\) Identifying cervical cancer using testing for HPV DNA is more sensitive but less specific than cervical cytology.\(^6\) Ninety-two per cent of women will survive five years when cervical cancer is localized, but only 13% will survive distant disease.\(^6\)

**Relative benefits and harms of screening and treatment programs**

For cervical cancer screening programs, we found two large-scale observational studies\(^6\) and two systematic reviews.\(^6\) The screening programs used invitation and reminder letters and practitioner incentives to increase screening rates from 61% to 83% of the population. Data from Rieck and coworkers\(^6\) showed a reduction in cervical cancer (relative risk 0.48, 95% CI 0.23–0.98) (Table 21B). Adverse effects resulting from referral for colposcopy include anxiety about pain and discomfort.\(^6\) However, no data quantifying these adverse effects were found. Perinatal

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### Table 21A: Summary of findings for prophylactic HPV vaccination against cervical cancer in women

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<tr>
<td><strong>Outcome</strong></td>
</tr>
<tr>
<td>High-grade cervical lesion</td>
</tr>
<tr>
<td>Persistent HPV infection, 12 mo</td>
</tr>
<tr>
<td>≥ 1 serious adverse event*</td>
</tr>
<tr>
<td>Death from adverse event</td>
</tr>
<tr>
<td>Death from cervical cancer</td>
</tr>
</tbody>
</table>

Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; HPV = human papillomavirus; NA = not applicable; NNT = number needed to treat; RR = relative risk.

*Serious adverse events: bronchospasm, gastroenteritis, headache, hypertension, pain at injection site or impaired joint movement in injected limb.
mortality and adverse pregnancy outcomes have also emerged as rare though important potential harms in treatment of cervical intraepithelial neoplasia relevant for young women; however, the quality of this evidence remains very low\(^{651}\) (Table 21B).

Introduction of screening programs to populations naive to screening reduces cervical cancer rates by 60%–90% within three years of implementation.\(^{653}\) Key factors that improve the effectiveness of programs include high participation rate, quality control in smear interpretation, reliable follow-up for abnormal results and facilities for adequate treatment.\(^{654}\)

**Clinical considerations**

Initiating a pelvic examination with simple nonmedical language can build rapport, increase comfort and empower patients.\(^ {655}\) Cervical cancer screening will be a new concept for many immigrant women. Factors that can reduce rates of screening among immigrant women included limited English- or French-language proficiency, employment and educational demands, transportation and childcare difficulties, as well as male physicians trying to respect patient’s modesty.\(^ {656}\) Before screening for cervical cancer in women who have been victims of sexual violence, practitioners should develop rapport, a process that can take several visits. Abuse may continue or increase after arrival in Canada because of additional stresses, so it is important to take time to listen to patients.

Women who perceive their risk of HPV infection as high are more accepting of HPV vaccination.\(^ {657}\) Other factors influencing acceptance of HPV vaccine include perceived efficacy of the vaccine and physicians’ recommendation of vaccination. The most common barrier to vaccine acceptance is cost. Evidence demonstrating the acceptability of vaccination to those at highest risk for cervical cancer (including ethnic minorities) is limited.\(^ {657}\) In most provinces and territories in Canada, the HPV vaccine is publicly funded only for girls through a school-based immunization program, with no catch-up vaccination provision for newly arriving older immigrant girls.

**What are the potential implementation issues?**

Immigrant women often have little knowledge and many misconceptions about the benefits of screening for cervical cancer. Many cultures are very private regarding sexuality, leaving women reluctant to ask for cervical screening, and some women feel uncomfortable undressing in front of a stranger.\(^ {658}\) Patients, especially female Muslim patients, may prefer female practitioners;\(^ {658}\) and some also prefer a caregiver from the same culture. Immigrant community health workers and other community interventions that provide information and offer transport, female physicians and interpreters in informal clinic settings may improve uptake of screening.\(^ {658}\)

**Recommendations of other groups**

The Canadian Immunization Committee recommends vaccination of Canadian female patients against HPV types 16 and 18.\(^ {659}\) The Society of Obstetricians and Gynaecologists of Canada\(^ {660}\) and the Canadian Task Force on the Periodic Health Examination (now the Canadian Task Force on Preventive

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**Table 21B:** Summary of findings for organized screening program compared with opportunistic screening to prevent cervical cancer

<table>
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<tr>
<td><strong>Absolute effect in medium-risk population</strong></td>
</tr>
<tr>
<td>Outcome</td>
</tr>
<tr>
<td>Cervical cancer rate(^ {654})</td>
</tr>
<tr>
<td>Invasive cervical cancer(^ {653})</td>
</tr>
<tr>
<td>Death from cervical cancer (surveillance)(^ {654})</td>
</tr>
</tbody>
</table>

*Note: CI = confidence interval; GRADE = Grading of Recommendations Assessment, Development and Evaluation; NNT = number needed to treat; RR = relative risk*
Health Care\textsuperscript{671} recommend cervical cytology screening programs for sexually active adolescents and women. The International Agency for Research on Cancer recommends cervical screening for women 25–65 years of age and suggests a three-year screening interval be considered in countries with adequate resources.\textsuperscript{638} Our recommendations highlight the need to improve the delivery of cervical cancer screening for immigrant women and vaccination against HPV.

**Take-home messages**

- Vaccination against HPV is recommended for 9- to 26-year-old female patients to reduce invasive changes related to cervical cancer.
- All sexually active women should be screened for cervical abnormalities (with Pap smear) to detect and treat invasive changes.
- Providing clear information about cervical screening, building rapport and offering access to a female practitioner improve acceptance of Pap tests.
- Immigrant girls may miss out on school vaccination programs, depending on their age upon arrival in Canada.

22. Pregnancy

Every migrant woman, regardless of her status, has the fundamental right to receive complete prenatal, birth and postnatal care with dignity. However, evidence shows that access to pregnancy care is inadequate. From 1996 to 2001, 22% of all births in Canada were to foreign-born women. Some births to migrant women are the result of sexual abuse in the case of civil unrest, sexual abuse during their flight from civil unrest, or sexual persuasion or favours granted to those organizing their transit to receiving countries. Higher rates of cesarean section among newly arrived women (33.0%–35.8%) than among Canadian-born women (26%) have been reported. Maternal mortality rates in the United Kingdom for black African women are 5.6 times higher than for white women; for black Caribbean women they are 3.7 times higher, and for Middle Eastern women they are 2.9 times higher. Upon investigation, these higher mortality rates were found to be related to poor overall health, including unrecognized medical conditions, practitioners’ ignorance of female genital mutilation, cultural practices, attitudes of male partners, inadequate interpreter services and suboptimal care. In this section we review and compile existing evidence for pregnancy care of women who have arrived in Canada within the past five years. The recommendations of the Canadian Collaboration for Immigrant and Refugee Health on pregnancy care are outlined in Box 22A.

Methods

We used the 14-step approach developed by the Canadian Collaboration for Immigrant and Refugee Health (summarized in section 3 of this article, above). We searched MEDLINE, Embase, CINAHL, HealthSTAR, the Cochrane Library and other sources from Jan. 1, 1996, to Jan. 1, 2010. We limited the literature search to the most salient issues for newly arrived pregnant women, and searched as systematically as possible, by focusing on issues with evidence of disease burden affecting this population more than other pregnant women, as well as on suggestions that clinical responses for this population should differ from responses for other pregnant women and on issues for which a concurrent evidence review was not already being conducted. Detailed methods, search terms, case studies and clinical considerations can be found in the complete evidence review for pregnancy (see Appendix 20, available at www.cmaj.ca/cgilookup/doi/10.1503/cmaj.090313/-/DC1).

Results

The search for guidelines and systematic reviews on pregnancy specific to immigrants yielded 20 titles, of which none were systematic reviews. After we applied our exclusion criteria to the search for articles on pregnancy for the general population, psychosocial concerns were restricted to social isolation, violence was restricted to sexual abuse, and prenatal screening for blood disorders was restricted to hemoglobinopathy. Work exposure and female genital mutilation remained as factors to consider in their entirety. Only four of 251 articles fulfilled our inclusion criteria. A further three were subsequently excluded because they lacked specificity for the screening or treatment intervention under consideration. The Committee of Obstetrics of the American College of Obstetricians and Gynecologists published clinical guidelines in 2007 and the Society of Obstetricians and Gynaecologists of Canada published clinical guidelines in 2008, both focused on hemoglobinopathy.

How do pregnancy-related issues affect immigrant populations?

Social isolation

One Canadian report showed that, after giving birth, 14.7% of asylum seekers and 7.7% of refugee women live alone, whereas 2.7% of Canadian-born women live alone. Perceived lack of social support was reported among 15.4% of immigrants versus 7.2% of Canadian-born women. Lack of psychosocial resources for nonmigrants has been found to lead to small-for-gestational-age infants.

Sexual abuse

Sexual abuse is common in armed conflict and internal strife. Abuse is sometimes used to dominate, sexual torture can be used as a method of interrogation, and refugee camp guards and male refugees sometimes regard unaccompanied women and girls as common sexual property. Violence and rape

Box 22A: Recommendations from the Canadian Collaboration for Immigrant and Refugee Health: pregnancy

Develop and study interventions to reduce social isolation, given the risk for maternal morbidity and small-for-gestational-age infants.

Basis of recommendation

Balance of benefits and harms

Pregnant immigrant and refugee women face an elevated risk of social isolation (15% v. 7.5% for Canadian-born women), which is associated with maternal morbidity and small-for-gestational-age infants. However, in the absence of evidence showing that social interventions work, such interventions could cause harm. Therefore, the committee recommends development and study of interventions for pregnant immigrant and refugee women who are socially isolated.

Quality of evidence

Very low, with no intervention evidence available

Values and preferences

The committee attributed more value to preventing uncertain harms than to providing uncertain benefits through unstudied social interventions.
among refugee women in camps ranges from 24.4% to 40.0%. Many refugee women (23%–50%) also report violence at the hands of their husbands. Pregnancy-related effects include unwanted pregnancies, sexually transmitted infections, chronic pelvic infection, reproductive tract trauma, psychological trauma, social rejection, and mistreatment or abandonment of resulting infants.

Hemoglobinopathy
Hemoglobinopathy (predominantly thalassemia and sickle-cell anemia) is more common among newly arrived women from certain regions of the world. Hemoglobinopathy can lead to serious maternal complications, severe anemia in infants and painful vaso-occlusive crises.

Exposure to hazards in the workplace
Newly arrived women are often exposed to unprotected and unregulated work environments, where the risk of adverse pregnancy outcomes is great. Their lack of knowledge of employee rights exaggerates these exposures.

Female genital mutilation
The prevalence of female genital mutilation ranges from 5% to 97% among women in countries where it is practised. There are four types of female genital mutilation: type I, excision of the prepuce, with or without excision of all or part of the clitoris; type II, excision of the clitoris with partial or total removal of the labia minora; type III, excision of part or all of the external genitalia and stitching or narrowing of the vaginal opening, also called “infibulation” and type IV, “unclassified,” which include pricking, piercing, incising, stretching, cauterizing, scraping, cutting, introducing corrosive substances or other methods. Type III causes a mechanical barrier to delivery; however, types I, II and IV can produce severe vulval and vaginal scarring that may also obstruct delivery. Types II and III are associated with greater risks of cesarean section, postpartum hemorrhage, extended hospital stay, infant resuscitation, and stillbirth or early neonatal death. An estimated 15% of all circumcised women have undergone type III mutilation. Most cases of female genital mutilation in Djibouti, Somalia and Sudan (80%–90%) are type III. Practitioners attending Somali births in Canada have been found to lack knowledge of female genital mutilation and to manifest unprofessional attitudes toward these women.

Does screening for social isolation or exposure to unregulated work environments decrease pregnancy-related morbidity and mortality?
Currently, no screening tools or interventions, apart from provision of information (www.workrights.ca), focus specifically on social isolation or exposure to unprotected work environments among pregnant women. Although these topics are recognized as issues in the literature, no relevant guidelines or systematic reviews were specific to pregnant women.

Clinical considerations
Language barriers and cultural considerations, as well as eligibility for health services, play important roles in access to care. Patients may fear that requests for additional services will reduce their chances of successfully remaining in Canada. Practitioners should consider the patient’s interest in screening for and treating these conditions, and interacting in culturally sensitive ways about these issues. Treatment preferences and expectations are often based on what was done in the country of origin, which could differ from Canadian care (e.g., expectation of a greater number of ultrasound examinations). The procedures themselves could be unacceptable (e.g., amniocentesis) or the notion of statistical probability might not be well understood. Further, if the testing must be paid for and families have limited resources, they could be unable to have the tests done. Closely spaced pregnancies (resulting in physical health problems and difficult social integration) are another culturally driven consideration.

Primary care practitioners in the author group noted that the woman’s health and pregnancy history might be incorrect or incomplete for several reasons. She could fear jeopardizing her asylum or claimant application (e.g., if the application states she has no other children, she might not want to report now that she has other children), her records from the previous country might be unavailable, or she could have delayed the start of prenatal care. These issues could be particularly important in determining the appropriateness of a cesarean birth.

In Canada, immigrants face a three-month waiting period for provincial health coverage in certain provinces, limited numbers of clinicians accept the Interim Federal Health Program plan, and migrants often lack awareness of services. Delayed prenatal care in Ontario has been found to be as high as 60% in migrant pregnant women.

Recommendations of other groups
The recommendations of other groups in three specific areas are outlined below. Our recommendations highlight the research needed to develop interventions and to reduce social isolation for pregnant immigrant women.

Female genital mutilation
Recommendations from the World Health Organization are available, as is a review of clinical practice. Guidelines for care in cases of female genital mutilation were published by Health Canada, although no evidence of the effectiveness of suggested interventions was provided. An article entitled “What Somali women say about giving birth in Canada” provides suggestions for providing more respectful and less interventionist care, as well as greater sensitivity for cross-cultural practices.

Sexual abuse
Health Canada’s guidelines on family-centred maternity and newborn care (published in 2000) recommend using the ALPHA form to assess current and past abuse (including
sexual abuse), although sensitivity and specificity (and other measures) are not provided for this tool. No recommendations are available specific to the screening and care of sexual abuse victims during pregnancy, a time during which bodily changes and vaginal examinations can elicit untoward memories of sexual abuses.

**Hemoglobinopathy**

The guidelines of the Society of Obstetricians and Gynaecologists of Canada recommend that those who originate from countries outside northern Europe should be considered at high risk for hemoglobinopathy. As well “the definitive test for carriers of hemoglobin S, C, or D is hemoglobin electrophoresis, which should be offered to all couples of African or Caribbean descent.” The guidelines of the American College of Obstetricians and Gynecologists’ (based on level A scientific evidence, i.e., good and consistent) note that “Individuals of African, Southeast Asian, and Mediterranean descent are at increased risk for being carriers of hemoglobinopathies and should be offered carrier screening and, if both parents are determined to be carriers, genetic counselling.”

**Take-home messages**

- Women newly arrived to Canada are responsible for an important proportion of total births.
- In the hopes of reducing maternal morbidity and reducing the number of infants who are small for their gestational age, further development and study is required of interventions to address the social isolation of immigrants and refugees who are pregnant.
- However, we lack data to support actions to be taken by primary care practitioners related to these disparities.

evidence in southern Italy. Vaccine 2003;21:399-400.


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Marc Deschenes has attended meetings of the Gilead Sciences Advisory Board. Sheila Dunn was on the advisory board of Trimedic in 2008, received speaker fees from Palabin Labs in 2008, received research grants for clinical trials of contraceptives from Organon (Scherwing-Plough) and provided information (without compensation) to the National Drug Safety and Advisory Committee, in collaboration with Palabin Labs, to support deregulation of emergency contraception in 2007. John Feightner has advised and consulted with the Canadian Task Force on Preventive Health Care. Elizabeth Harvey has served as a national advisory board member for Sanofi Aventis and as a consultant for Lifsescan; and her institution receives program funding from the Ontario Diabetes Strategy. In addition, Pfizer, Becson, Dickinson, Novo-Nordisk, Eli Lilly, Sanofi Aventis, Bayer, Merck, Astra Zeneca and Bristol Myers Squibb have provided grants to her institution and payments for lectures related to the development and presentation of educational programs about diabetes for primary care providers. Jenny Heathcote has served as a consultant to Axcan Pharma, Gilead Sciences, Hoffmann-La Roche, Merck, Schering-Plough and Tibotec; has received unrestricted grants from Axcan Pharma, Boehringer Ingelheim, Gilead Sciences, Hoffmann-La Roche, Schering-Plough and Vertex; has received unrestricted grants from Axcan Pharma, Boehringer Ingelheim, Gilead Sciences, Hoffmann-La Roche, Schering-Plough and Vertex; and has received speaker’s fees from Axcan Pharma, Boehringer Ingelheim, Hoffmann-La Roche, Schering-Plough and Tibotec. Charles Hui has received grant support through his institution, as well as payment for lectures and travel expenses, from Abbott International; has also served on a scientific advisory board for Novartis. Jay Keystone has received speaker fees and payments for development of educational presentations from GlaxoSmithKline, Sanofi Pasteur and Merck. Stan Kutcher occasionally serves as an expert witness. Noni MacDonald, Section Editor, Population and Public Health, CMAJ, is a coauthor of this article; she was not involved in the vetting of this article before publication. Anne McCarthy has served as an ad hoc consultant on travel medicine to Shorelands Inc. Lavanya Narasiah received speaker fees from GlaxoSmithKline for presentations on health. Pierre Blaise is a consultant to Roche, GlaxoSmithKline for presentations on travel health and tropical medicine. Peter Tugwell, chair of the CMA Journal Oversight Committee, is a coauthor of this article; he was not involved in the vetting of this article before publication. David Wong has received educational grants related to hosting the University of Toronto Sheila Sherlock Liver Research Day and the University of Toronto Hepatology Update; he also has received honoraria for educational sessions from Roche, Schering, Gilead, Bristol-Myers Squibb and Novartis (in all instances, the slides used for sessions have been his own). No competing interests declared by Deborah Assayag, Beverly Brocket, Ralf Buhrmann, Giovani Burgos, Glenn Campbell, Andrea Chambers, Angie Chan, Marianne Cheetham, Walter Delpero, Shafik Dharmoo, Arumoozhi Dominic, Ann Doggan, Nancy Durand, Allison Eyer, Anita Gagnon, Jennifer Grant, Christina Greenaway, Doug Gruner, Sinclair Harris, Stewart Harris, Ghayda Hassan, Christine Heidebrecht, William Hodge, Danielle Hone, Susan Hun, Praseedha Janakiram, Khausriv Jivani, Tomas Jurcik, Kamran Khan, Laurence Kirkmayr, Ian Kita, Srinivasan Krishnamurthy, Susan Kuhn, Robert LaRoche, Carmen Logie, Michelle Martin, Dominique Ellen Massenet, Debora Matthews, Barry Maze, Mary McIlrath, Dick Menzies, Marie Rundell, Félicité Murangira, Amy Nolen, Kevin Pottie, Meb Rashid, Cécile Rousseau, Hélène Rousseau, Andrew Ryder, Amelia Sandoe, Kevin Schwartzman, Jennifer Sears, William Stauffer, Helena Swinkels, Brett Thoms, Patricia Topp, Andrew Toren, Sara Torres, Erin Ueffing, Ahsan Ullah, Sunil Varghese, Bilkis Vissandjee, Vivian Welch, Michel Welt, Wendy Wobeser, Phyllis Zelkowitz, Jianwei Zhong, Stanley Zlotkin.

Contributors: Section 1 (Review): Kevin Pottie, Christina Greenaway, John Feightner, Vivian Welch, Helena Swinkels, Meb Rashid, Lavanya Narasiah, Laurence J. Kirkmayr, Erin Ueffing, Noni E. MacDonald, Ghayda Hassan, Mary McNally, Kamran Kahn, Ralf Buhrmann, Sheila Dunn, Arunmozhi Dominic, Anne E. McCarthy, Anita J. Gagnon, Cécile Rousseau, and Peter Tugwell contributed as members of the steering committee and/or lead authors on reviews and played an active role in formulating the recommendations. Other coauthors of the Canadian Collaboration for Immigrant and Refugee Health (see complete list on the title page) provided substantive contributions for the individual systematic reviews as listed below.

Section 2 (Selection of potentially preventable and treatable conditions): Helena Swinkels, Kevin Pottie, Peter Tugwell, Meb Rashid and Lavanya Narasiah contributed to the conception and refinement of the study design and to the analysis and interpretation of the data. Helena Swinkels drafted the initial manuscript, and all authors provided critical revisions and approved the final manuscript submitted for publication.

Section 3 (Evaluation of evidence-based literature): Peter Tugwell, Kevin Pottie, Vivian Welch, Erin Ueffing, Andrea Chambers and John Feightner contributed to the conception, writing and critical revision of the manuscript and approved the final version submitted for publication.

Section 4 (Measles, mumps, rubella, diphtheria, tetanus, pertussis and polio): Christina Greenaway, Marie Munoz, Elizabeth Barnett, Amelia Sandoe, Erin Ueffing, Kevin Pottie, Susan Kuhn and Jay Keystone contributed to the conception and refinement of the study design and the analysis and interpretation of the data. Christina Greenaway drafted the initial draft and all authors provided critical revisions and approved the final manuscript submitted for publication.

Section 6 (Hepatitis B): Christina Greenaway, Lavanya Narasiah, Pierre Plourde, Erin Ueffing, Kevin Pottie, Marc Deschenes, David K.H. Wong, Susan Kuhn and Jenny Heathcoote contributed to the conception and refinement of the study design and the analysis and interpretation of the data. Christina Greenaway drafted the initial manuscript, and all authors provided critical revisions and approved the final manuscript submitted for publication.

Section 7 (Tuberculosis): Christina Greenaway, Amelia Sandoe, Bilkis Vissandjee, Ian Kitai, Doug Gruner, Kevin Pottie, Erin Ueffing, Wendy Wobeser, Dick Menezes and Kevin Schwartzman contributed to the conception and refinement of the study design and to the analysis and interpretation of the data. Christina Greenaway wrote the initial draft, and all authors provided critical revisions and approved the final manuscript submitted for publication.

Section 9 (Hepatitis C): Christina Greenaway, David K.H. Wong, Deborah Assayag, Marc Deschenes, Chuck Hui, Erin Ueffing, Kevin Pottie, Amelia Sandoe, Meb Rashid and Jenny Heathcoote contributed to the conception and refinement of the study design and to the analysis and interpretation of the data. Christina Greenaway wrote the initial draft, and all authors provided critical revisions and approved the final manuscript submitted for publication.

Section 10 (Intestinal parasites): Kamran Khan, Christine Heidebrecht, Jennifer Sears, Angie Chan, Meb Rashid, Christina Greenaway, William Stauffer, Lavanya Narasiah and Kevin Pottie contributed to the conception and refinement of the study design and the analysis and interpretation of the data. Kamran Khan drafted the initial manuscript, and all authors provided critical revisions and approved the final manuscript submitted for publication.

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