Introduction

Our immunization programs for young children are one of the great public health success stories of the twentieth century. They have changed the face of childhood — literally saving the lives of thousands of children every year by minimizing or eliminating the risks of many serious infant and childhood illnesses.\(^1^,\,^2\) With the exception of safe water, no other modality, not even antibiotics, has had such a major impact on mortality reduction and so improved survival.\(^3\)

Subject

Recommended Immunization Schedule

The National Advisory Committee on Immunization (NACI) provides the federal government (ie, Health Canada) with ongoing and timely medical, scientific and public health advice relating to immunization.\(^1\) The current NACI-recommended immunization schedule for children is summarized in Table 1.\(^1^,\,^4\) Since health is a provincial not federal responsibility in Canada, each province and
territory individually decides which, when and whether specific vaccines will be included in the vaccine program funded by that province or territory.\textsuperscript{5} Unfortunately, this approach has led to an uneven patchwork of vaccine coverage for children across our land.\textsuperscript{5,6} For example, as of Dec 2003, the NACI-recommended newer vaccines (eg, varicella,\textsuperscript{7} conjugated pneumococcal,\textsuperscript{8} conjugated meningococcal,\textsuperscript{9} and adolescent acellular pertussis\textsuperscript{10} vaccines [See below.]) that are only routinely available in some provinces and territories.\textsuperscript{6} Even though there have been many calls for a National Immunization Strategy in Canada, as yet we do not have one, in contrast to the United States, Australia, the United Kingdom and others.\textsuperscript{5,6,11-14}

Benefits of Routine Immunization among Young Children: Lives Saved

Not so many years ago parents and health care workers alike saw first hand the potential consequences for infants and young children who became infected with the diseases now prevented by routine immunization programs. In the early 1900s, 5 out of every 1,000 children born in Canada and the United States died from pertussis (whooping cough) before they reached their 5th birthday\textsuperscript{15} and diphtheria (bacteria causing disease that hinders swallowing and breathing) was one of the most common causes of death in children from 1 to 5 years of age, killing thousands of children each year.\textsuperscript{1} Polio (an infectious viral disease affecting the central nervous system) was a much feared summer scourge that often killed or crippled.\textsuperscript{16} Table 2 presents a comparison of the prevalence of diseases before and after the introduction of routine vaccines.\textsuperscript{1,2,8-10,15-18}

To reap the benefit from these vaccines, children must be immunized and immunized on time. These diseases can still kill or maim, even when there is access to modern day intensive care and antibiotic therapy.\textsuperscript{19-21} In the mid 1990s, many families living in the Russian Federation were retaught the lesson of the dangers of diphtheria and the importance of immunization as diphtheria made a marked resurgence with more than 115,000 cases and 3,000 deaths reported.\textsuperscript{20} This outbreak occurred in a country where diphtheria had previously been well controlled. The break up of the former USSR led to profound social changes that included a dramatic fall off in immunization rates for infants and children and a failure to give booster doses to adults. Case control studies showed that those who were immunized were protected; those who were not were in trouble.\textsuperscript{22} This tragic epidemic was due not to vaccine failure, but to a failure to immunize.

Problems
Adverse Events Less Common with Vaccines than with Disease

Table 3 presents the effects of the disease and the known side effects of the vaccine for the routine vaccine-preventable diseases for infants and young children. In general, all of these diseases are serious and may be fatal, while the vaccine adverse events, if they occur, are usually minor such as local discomfort and/or inflammation at the site of the injection and/or mild fever or rash. Research has shown that the local pain of intramuscular infant immunization with DTaP/IPV/Hib can be diminished by the use of topical lidocaine-prilocaine without adversely affecting the development of the protective response from the multicomponent vaccine and the pain of multiple infant injections given during the same visit can be reduced by oral sucrose, oral tactile stimulation (a bottle or pacifier) and parental holding.

Serious vaccine adverse events occur with the routine immunizations but are a great deal rarer than serious events with the diseases. For example, aseptic meningitis (an infection of the membranes and fluid encasing the brain and spinal cord) occurs in 5% of those who get mumps (a viral disease which causes swelling of the salivary glands in the chin and face) and permanent deafness may occur in up to 0.5%. In contrast, aseptic meningitis following mumps vaccine with the Jeryl Lynn strain (the type of modified and weakened mumps viral vaccine strain used in Canada and several other countries) occurs after less than 1/800,000 doses and maybe as low as 1/3,000,000. Furthermore, the vaccine-associated aseptic meningitis is not followed by permanent problems, like deafness.

Beyond these known but rare vaccine-associated serious adverse events (Table 3), there have been periodic allegations that infant vaccines may cause other serious problems such as SIDS (sudden infant death syndrome) and autism. However, research has shown that these claims are unsubstantiated. There is no causal relationship between infant immunization on one hand and SIDS or autism on the other hand. While an event may have been recognized as happening soon after the receipt of an infant vaccine (ie, establishing a possible temporal relationship), receipt of the vaccine is not the basis for the occurrence of an event.

In order to enhance the evaluation of reported serious vaccine events in Canada, the Advisory Committee on Causality Assessment (ACCA) was set up in 1994 by Health Canada. This expert committee is charged with the task of monitoring signals for vaccine safety. The committee regularly reviews all reports of serious and unusual vaccine associated adverse events to

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determine, through a systematic, standardized approach whether the association of the event to the receipt of the vaccine is likely causal, probably causal, possibly causal, unlikely causal, unrelated or unclassifiable. On the international level, the World Health Organization set up the Global Advisory Committee on Vaccine Safety in 1999 whose task is to respond promptly, efficiently and with scientific rigor to vaccine safety issues of potential global importance.

In 1991, to improve the detection of serious vaccine-associated adverse events, vaccine failures and selected infant and child infectious diseases that are now or are soon to be vaccine preventable, Health Canada, in collaboration with the Canadian Paediatric Society and others, piloted a cross-Canada paediatric hospital-based active surveillance network in 5 centres. The network was then expanded to 11 centres in 1995 and 12 centres in 1999 (IMPACT). Compiled network data has repeatedly shown that the routine vaccines for young children are very safe. In addition, IMPACT has proven valuable in detecting rare but unexpected serious events (eg, disseminated Bacille Calmette-Guerin (BCG) infections in aboriginal infants immunized with BCG) that have lead to policy reevaluation. IMPACT has also been able to show a sharp decline in disease following the introduction of a new vaccine and a decrease in side effects following a shift to a new, improved vaccine.

Research Context

Best Practices for Vaccine Programs

In 1995, NACI initiated a 2-year consultative process to develop guidelines for childhood immunization practices applicable to both the public and the private systems of vaccine delivery in Canada. Table 4 provides a brief overview of the guidelines. Research has shown that a number of factors can enhance vaccine uptake, including timely reminders, quality parent education materials, after-hours and weekend clinics, vaccine uptake monitoring, multiple vaccines given during one visit, standing orders for vaccines, multi-component provider education, and the elimination of financial barriers to immunization.

When it comes to giving consent for immunization, research has shown that what matters to parents is that they receive the information they need to make an informed decision, but the mode in which this information is given does not matter. Bearing this in mind, the 2002 edition of the NACI Canadian Immunization Guide was expanded to include a separate chapter on consent issues and parental concerns regarding immunization to help health care providers better counsel...
parents. Recognizing that the information on vaccines contained in the NACI Canadian Immunization Guide may be too technical for many parents, the Canadian Paediatric Society supported the development of a vaccine handbook specifically designed for parents entitled, “Your Child’s Best Shot,” which was first published in 1997, then updated to include the newer NACI-recommended vaccines in 2002.

As noted above, well-informed health care providers are an important factor in enhanced vaccine uptake, but research has shown that many are not well informed. Efforts to improve on this knowledge deficit include the revamping and upgrading of the NACI Canadian Immunization Guide, continuing vaccine education events for doctors and nurses, journal articles, further research, the formation of the Canadian Coalition for Immunization Awareness and Promotion, the formation of the Canadian Association for Immunization Research and Evaluation and the biannual National Immunization Conference.

Vaccine Programs for Young Children with Special Needs

While the routine NACI vaccine schedule (Table 1) is appropriate for the majority of Canadian children there are subgroups with special needs including:

1. infants and young children born outside of Canada who come as immigrants, refugees or foreign adoptees who may not have received all of the vaccines recommended in Canada, and/or may not have adequate vaccine documentation.
2. infants born prematurely
3. infants and children who are immunocompromised from birth or from disease
4. infants and children who have bleeding disorders or have a nonfunctional or absent spleen
5. infants and young children who travel to other countries. In each of these cases, the routine immunization requirements and schedule may need to be adapted.

Recent Research Results

Newer NACI-recommended Vaccines for Young Children

There are 3 vaccines recently recommended by NACI for infants and young children which are not yet covered by funded vaccine programs in all of the provinces and territories. These include the varicella vaccine for prevention of chicken pox, conjugated pneumococcal vaccine for the
prevention of blood infection, pneumonia and meningitis due to pneumococcal bacteria, and conjugated meningococcal vaccine for the prevention of meningitis and blood infection, again, due to this organism. The risks of these diseases, vaccine benefits and side effects are summarized in Tables 2 and 3. In all three cases these vaccines have been shown to be safe and effective in preventing serious diseases in infants and young children, but each is also relatively expensive compared to the cost of the “regular” infant immunizations. The prohibitive costs of these vaccines has led to delays and disparities in having these vaccines added to the “routine” list covered by each province and territory. A similar problem exits for the acellular pertussis vaccine for adolescents and adults. While acellular pertussis vaccine is available across Canada for infants and young children, despite NACI recommendations, the acellular vaccine for adolescents and adults is not yet routinely available across the country. Widespread use of this vaccine in adolescents and adults has the potential to decrease pertussis in families and thus decrease exposure of infants who are too young to be immunized (ie, less than 6 weeks of age), the group at highest risk for fatal disease.

Conclusions

The NACI-recommended vaccines for young children are a safe and effective means of eliminating or minimizing the risks of many serious infant and childhood illnesses. Infants and children who are not immunized continue to be at risk. The NACI Canadian Guide to Immunization is the best detailed source of information on all aspects of immunization for health care providers and Your Child’s Best Shot provides quality information for parents.

Implications

A National Immunization Program is needed to improve equity of access across this country to all of the NACI-recommended vaccines for infants and young children in order to be able to protect all of our children from the potential damage incurred by a vaccine-preventable disease. Not ensuring equity of access means many infants and children remain at risk for problems such as acquired deafness from meningitis due to pneumococcal infection, along with its profound developmental implications. Policy makers at the federal, provincial, and territorial levels must work together to ensure a National Immunization Program for Canadian infants and children so that all have access to NACI-recommended vaccines.
### NACI-recommended Immunization Schedule for Infants and Children (from references 1, 8-10, 15-18)

<table>
<thead>
<tr>
<th>Age at Vaccination</th>
<th>DTaP</th>
<th>IPV</th>
<th>Hib</th>
<th>MMR</th>
<th>dTap or Td</th>
<th>HepB (3 doses)</th>
<th>V</th>
<th>PC</th>
<th>MC</th>
<th>Influenza</th>
</tr>
</thead>
<tbody>
<tr>
<td>Birth</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td></td>
<td></td>
<td>Infancy or pre-adolescence</td>
<td></td>
<td></td>
<td></td>
<td>Before influenza season in those over 6 months, esp. in high risk categories</td>
</tr>
<tr>
<td>2 months</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td></td>
<td></td>
<td></td>
<td>x</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 months</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td></td>
<td></td>
<td></td>
<td>x</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6 months</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>x</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>12 months</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td></td>
<td></td>
<td>X</td>
<td></td>
<td>x</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>18 months</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td></td>
<td></td>
<td>x or</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4–6 years</td>
<td>x</td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>14–16 years</td>
<td>x</td>
<td></td>
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<td></td>
<td></td>
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</tr>
</tbody>
</table>

**DTaP** Diphtheria, tetanus, pertussis (acellular), infant and young-child-type vaccine  
**IPV** Inactivated polio vaccine  
**Hib** *Haemophilus influenzae* type-b conjugate vaccine  
**MMR** Measles, mumps, rubella vaccine  
**dTap** Tetanus and diphtheria toxoid, (acellular) pertussis, adolescent/adult-type vaccine  
**Td** Tetanus and diphtheria toxoid, adult-type vaccine  
**HepB** Hepatitis B vaccine  
**V** Varicella vaccine  
**PC** Pneumococcal conjugate vaccine  
**MC** Meningococcal conjugate vaccine  
**Influenza** Influenza virus vaccine

* Conjugated pneumococcal vaccine: Doses at 2, 4 and 6 months, followed by one dose at 12–15 months of age.  
** Conjugated meningococcal vaccine: If started at 2 months — 3 doses; if started at 4 to 11 months — 2 doses; if started at ≥ 12 months — 1 dose.

λ While all recommended by NACI, acellular pertussis for adolescents, varicella, conjugated pneumococcal, and conjugated meningococcal vaccines are not currently available in all Canadian provincial and territorial infant and childhood immunization programs.

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**TABLE 2**

**Serious Illnesses in Infants, Children and Youth in Canada in the Pre- and Post-Vaccine Eras (from references 1, 2, 8-10,15-18)**
<table>
<thead>
<tr>
<th>Disease/Organism</th>
<th>Incidence Before Vaccination</th>
<th>Incidence After Vaccination</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polio</td>
<td>2.5 to 28.3/100,000</td>
<td>Disease eradicated from Canada and from most countries in the world.</td>
</tr>
<tr>
<td>3 types of polio virus</td>
<td>Epidemics: up to 20,000 cases of paralysis.</td>
<td></td>
</tr>
<tr>
<td>Diphtheria</td>
<td>In 1924, 9,000 cases reported. Major cause of death in 1 to 5 year olds.</td>
<td>No cases reported since 1996, and prior to this only 2–5 per year.</td>
</tr>
<tr>
<td>Tetanus (lock jaw)</td>
<td>60 to 75 cases per year, with 40 to 50 deaths</td>
<td>Less than 2 cases per year in past 15 years.</td>
</tr>
<tr>
<td>Pertussis (whooping cough)</td>
<td>Over 150/100,000 cases per year with 50 to 100 deaths.</td>
<td>10/100,000 cases per year with 1 to 3 deaths in very young infants.</td>
</tr>
<tr>
<td><em>Haemophilus influenzae</em> type b</td>
<td>Overall 2,000 cases per year with 1500 in those &lt;5 years of age. Leading cause of bacterial meningitis in infancy.</td>
<td>Less than 50 cases per year. Rare cause of bacterial meningitis in infants.</td>
</tr>
<tr>
<td>Measles</td>
<td>Cyclic epidemics every 2-3 years. 300,000–400,000 cases per year.</td>
<td>Now fewer than 400 cases per year.</td>
</tr>
<tr>
<td>Mumps</td>
<td>About 30,000 reported cases per year but many more not reported</td>
<td>Less than 500 cases per year.</td>
</tr>
<tr>
<td>Rubella</td>
<td>About 250,000 cases per year, with over 200 congenital rubella syndrome (CRS)/year</td>
<td>Less than 100 cases reported per year, 1 to 2 congenital rubella syndrome (CRS) /year.</td>
</tr>
<tr>
<td>Disease/Organism</td>
<td>Incidence Before Vaccination</td>
<td>Incidence After Vaccination</td>
</tr>
<tr>
<td>------------------</td>
<td>-----------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>Influenza</td>
<td>Yearly epidemics with up to 20% of the population being infected. Wide variation in annual incidence. Last major outbreak: 1968 with 50 million cases, 33,000 deaths.</td>
<td>Since influenza virus changes on a yearly basis, yearly vaccination is required. Current program is aimed at high risk with limited uptake by others. With well-matched vaccine-influenza in community, 70–90% of illness is prevented, if immunocompetent. If ill-matched, it is only 30–60% protective. Less effective if not immunocompetent.</td>
</tr>
<tr>
<td>Hepatitis B</td>
<td>20,000 new infections per year, 1 in 200 people in population is a chronic carrier. BC had a rate of 33.7/100,000 in 1992. Risk of transmission from an infected mother to her newborn infant is 90%.</td>
<td>From 1992 to 2002 in BC after adopting a Grade 6 vaccine program, overall rate acute infection fell from 7 to 2/100,000 and in 12 to 21 year olds from 1.7 to 0/100,000. Immunization of newborn infants prevents transmission from mother in &gt; 90% of cases.</td>
</tr>
<tr>
<td>Varicella (Chicken pox)</td>
<td>Infection in 50% of children by age 5 and 90% by age 12.</td>
<td>Varicella mortality in the US has decreased by 76% with national program and coverage rates of 80% in less than 3 year olds.</td>
</tr>
<tr>
<td>Disease / Organism</td>
<td>Incidence Before Vaccination</td>
<td>Incidence After Vaccination</td>
</tr>
<tr>
<td>-------------------</td>
<td>------------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td><em>Streptococcus pneumoniae</em></td>
<td>About 500,000 cases of pneumococcal diseases per year with over 200,000 in children under 5 years of age. Rate of invasive disease &lt;5 years 35-64/100,000; &lt; 2 years 59-112/100,000.</td>
<td>Conjugated heptavalent vaccine for infants only licensed in Canada in 2001. In the US, clinical trials in infants showed vaccine efficacy of 94% for invasive diseases due to strains in vaccine and 89% for invasive disease due to any pneumococcal strain.</td>
</tr>
<tr>
<td><em>Neisseria meningitidis</em></td>
<td>Endemic in Canada with epidemics every 10 to 15 years. 200 to 350 endemic cases per year. Rates in 3 highest age groups: &lt;1 years 11.3/100,000; 1-4yr 2.4/100,000; 15 to19 years 1.5/100,000. Serogroups in 2001: A — rare; B — 28%; C — 59%; W 135 — 3%; Y — 10%.</td>
<td>Conjugated group C vaccine for infants only licensed in Canada in 2001. In UK, routine infant immunization started in 1999 with follow up campaign for children and adolescents has decreased disease by &gt;90%.</td>
</tr>
</tbody>
</table>

**TABLE 3**

**Comparison of Effects of Serious Infant and Childhood Diseases and Adverse Effects of Vaccines (from references 1, 2, 8-10, 15-17)**
<table>
<thead>
<tr>
<th>Disease</th>
<th>Outcomes</th>
<th>Vaccine Reactions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polio</td>
<td>4–8% have minor illness, 1% get severe disease- paralytic polio, 1 in 20 hospitalized patients die and 50% remain paralyzed.</td>
<td>Local discomfort or redness at the site of injection in 5%. Killed vaccine so no risk of vaccine-associated polio.</td>
</tr>
<tr>
<td>Diphtheria</td>
<td>5–10% of cases die even with ICU care, antitoxin and antibiotics. The toxin may lead to neurological and cardiac complications.</td>
<td>DTaP vaccine: Local discomfort, swelling and/or redness at the site of injection in 20%, fever in 5%. A transient nodule may occur at the injection site, lasting for a few weeks. Up to 70% develop redness and swelling at the 4-6yr booster.</td>
</tr>
<tr>
<td>Tetanus (lock jaw)</td>
<td>10% of cases die, even with ICU care, antitoxin and antibiotics. Risk is greatest for the very young and the very old.</td>
<td>See above for DTaP. Local reactions increase with age, esp. in adults with Td boosters. Peripheral nerve damage has rarely been reported (&lt;1/1,000,000).</td>
</tr>
<tr>
<td>Pertussis (whooping cough)</td>
<td>1/400 infants with pertussis die, 1/400 sustain permanent brain damage. If under 6 months, 1% of cases die from pneumonia or fatal oxygen deprivation of the brain.</td>
<td>As above for DTaP. Far fewer side effects with the acellular pertussis (aP) vaccine than the previous whole-cell pertussis vaccine (P).</td>
</tr>
<tr>
<td>Haemophilus influenzae type b</td>
<td>5% of cases of meningitis die, 10–15% have permanent brain damage and 10–20% have deafness.</td>
<td>Usually in combination, as with DTaP/IPV/Hib. See above for side effects (same as for DTaP).</td>
</tr>
</tbody>
</table>
Measles
10% have complications such as pneumonia, ear infections. 1/1,000 have encephalitis (infection of the brain) with 10% dying and 25% being left with permanent brain damage, 1/25,000 have SSPE (a delayed fatal degenerative brain disease). Usually in combination, as with MMR. 5-10% have discomfort or local swelling and fever, with or without a rash. 1/24,000 have low platelets < 1/1,000,000 have encephalitis.

Mumps
1/20 develop aseptic meningitis (viral infection of tissues and fluids around the brain). 1/200 develop encephalitis. 1/200,000 are left deaf. Inflammation of testicles in 20-30% of males; inflammation of ovaries in 5% of post-pubertal females. Local discomfort, swelling and redness or fever in 5-10%.

Rubella
50% have rash, swollen glands, fever; 50% of adolescents and adults have arthritis and arthralgias; 1/6,000 have encephalitis. In the first 10 weeks of pregnancy, 85% risk of congenital rubella syndrome causes death of fetus, deafness, blindness and/or heart disease. 10% have local discomfort and fever, 5% have swollen glands, arthralgias (esp. adults), stiff neck. 1% develop noninfectious rash.
Influenza

Highest mortality rate in those over 65 years and in infants aged <12 months. Complications: pneumonia, febrile seizures, encephalitis, myocarditis, and myositis, Reye’s syndrome.

Local mild reactions at injection site and/or low fever for 1 to 2 days in up to 60%. Occasional mild oculorespiratory syndrome. Rare: Guillian-Barre syndrome 1/1,000,000.

Hepatitis B

Variable: asymptomatic to overwhelming liver disease.
Neonate asymptomatic, 5-15% of 1 to 5 year olds have symptoms, 33-50% older children egg nausea, jaundice, fever, vomiting, big liver, spleen.
< 1% fulminating fatal liver failure.
Chronic disease 90% infants, 25-50% of 1 to 5 years, and 6 to 10% older children. Risk liver cancer, liver failure with chronic disease.

15% experience local discomfort and occasionally experience low-grade fever.

Varicella (chicken pox)

Death rate 1–3 /100,000 cases in children. Complications in 5–10% of previously healthy children: pneumonia, encephalitis (1/5,000), cerebellar ataxia (1/4,000), osteomyelitis, hepatitis, septic arthritis. In 50% of children who get flesh-eating disease (necrotizing fascitis), chicken pox precedes it. Shingles in adults. Congenital varicella syndrome.

15–20% experience mild swelling, discomfort at injection site and/or fever.
1–5% develop mild rash.
Anaphylaxis, a potentially life-threatening allergic reaction, occurs rarely (0.11 to 0.31 reports per 100,000 doses of vaccine distributed). It is rarer in infants and young children and occurs within 30 minutes of receipt of vaccine. Can be treated with an injection of epinephrine.

**Streptococcus pneumoniae** Leading cause of invasive bacterial disease in young children. Annual cases: 65 meningitis (hearing loss 20–30%, brain damage 15–20%), 700 cases bacteremia, 2,200 cases hospitalized with pneumonia, 9,000 cases non-hospitalized pneumonia. Case fatality rate <6 months 4.3%, 12 years 2%. 15 deaths/year in <5 years. Sickle cell disease, HIV more at risk bad disease.

**Neisseria meningitidis** Meningitis 30-50% (MR 5%), meningitis + bacteremia 40%, bacteremia alone 7-10% (MR 20-40%). Other complications: arthritis, pneumonia, peritonitis.

Case fatality rate 10% despite ICU/antibiotics. Highest mortality rate (MR): <1 year 1/100,000

* Anaphylaxis, a potentially life-threatening allergic reaction, occurs rarely (0.11 to 0.31 reports per 100,000 doses of vaccine distributed). It is rarer in infants and young children and occurs within 30 minutes of receipt of vaccine. Can be treated with an injection of epinephrine.

**TABLE 4**

National Guidelines for Childhood Immunization Practices: Summary*

1 Immunization services should be readily available.

2 There should be no barriers or unnecessary prerequisites to the receipt of vaccines.
Providers should use all clinical encounters to screen for needed vaccines and, when indicated, vaccinate children.

Providers should educate parents in general terms about immunization.

Providers should inform parents in specific terms about the risks and benefits of the vaccines their child is to receive.

Providers should recommend deferral or withholding of vaccines for true contraindications only.

Providers should administer all vaccine doses for which a child is eligible at the time of each visit.

Providers should ensure that all vaccinations are accurately and completely recorded.

Providers should maintain easily retrievable summaries of the vaccination records to facilitate age-appropriate vaccination.

Providers should report clinically significant adverse events following vaccination promptly, accurately, and completely.

Providers should report all cases of vaccine-preventable diseases as required under provincial and territorial legislation.

Providers should adhere to appropriate procedures for vaccine management.

Providers should maintain up-to-date, easily retrievable protocols at all locations where vaccines are administered.

Providers should be properly trained and maintain ongoing education regarding current immunization recommendations.

Providers should operate a tracking system.
Audits should be conducted in all immunization clinics to assess the quality of immunization records and assess immunization coverage levels.

*Adapted from the Canadian Immunization Guide, 6th edition.¹

References


42. Le Saux N, Barrowman NJ, Moore DL, Whiting S, Scheifele D, Halperin S, for Members of the Canadian Paediatric Society/Health Canada Immunization Monitoring Program–Active (IMPACT). Decrease in hospital admissions for febrile seizures and


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