Invasive Bacterial Diseases in Northern Canada

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International Circumpolar Surveillance (ICS) is a population-based invasive bacterial disease surveillance network. Participating Canadian regions include Yukon, Northwest Territories, Nunavut, and northern regions of Québec and Labrador (total population 132,956, 59% aboriginal). Clinical and demographic information were collected by using standardized surveillance forms. Bacterial isolates were forwarded to reference laboratories for confirmation and serotyping. After pneumococcal conjugate vaccine introduction, crude annual incidence rates of invasive Streptococcus pneumoniae decreased from 34.0/100,000 population (1999–2002) to 23.6/100,000 population (2003–2005); substantial reductions were shown among aboriginals. However, incidence rates of S. pneumoniae, Haemophilus influenzae, and group A streptococci were higher in aboriginal populations than in non-aboriginal populations. H. influenzae type b was rare; 52% of all H. influenzae cases were caused by type a. Data collected by ICS contribute to the understanding of the epidemiology of invasive bacterial diseases among northern populations, which assists in formulation of prevention and control strategies, including immunization recommendations.

The circumpolar region of Canada is a sparsely populated area of 1.74 million square miles comprising 3 territories (Yukon, Northwest Territories, and Nunavut) and the northern regions of Québec and Labrador. The estimated population is 132,956, which represents 0.4% of the Canadian population (1). The circumpolar population is younger (Table 1) and has a larger proportion of aboriginal persons than the general Canadian population. Approximately 59% of the population in the region self-identify as Inuit, First Nations, or Métis, compared with 3.3% of the total Canadian population (2). Northern populations tend to have higher rates of invasive bacterial diseases, including those caused by Streptococcus pneumoniae and Haemophilus influenzae, with aboriginal persons at greatest risk for disease (3–5).

Canada has a universal healthcare system that includes access to both physician and hospital care. Publicly funded vaccination programs are a major component of disease control programs. Universal infant H. influenzae type b (Hib) vaccination programs were implemented in the Yukon, Northwest Territories, Nunavut, and northern regions of Québec and Labrador in the early 1990s. Pneumococcal polysaccharide vaccine, which protects against 23 serotypes of S. pneumoniae, has been available since 1983 and is recommended by the Canadian National Advisory Committee on Immunization (NACI) for all adults >65 years of age.

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age and children ≥2 years of age at high risk for infection. The 7-valent pneumococcal conjugate vaccine (PCV7) protects against 7 serotypes of S. pneumoniae and has been available in Canada since 2001; NACI recommends it for all children <2 years of age and children <5 years of age at high risk for disease. Meningococcal C conjugate vaccine has been recommended by NACI for all children <5 years of age, adolescents, and young adults since 2001 (6). Implementation of these NACI recommendations has occurred at various times throughout the region.

In Canada, communicable disease reporting is mandated at the provincial or territorial level; the list of reportable diseases varies by region. Reporting to national notifiable disease surveillance is not mandatory, and timely submission of case-by-case data with epidemiologic, clinical, and laboratory information is variable. Therefore, to increase the understanding of the epidemiology of invasive bacterial diseases in northern populations, Canada has participated in International Circumpolar Surveillance (ICS) since its inception in 1999. ICS is a population-based invasive bacterial disease surveillance network of circumpolar countries that includes the United States, Canada, Greenland, Iceland, Finland, Norway, and Sweden. We describe Canadian ICS data from 1999 through 2005, including the effect of universal PCV7 programs on invasive S. pneumoniae disease in children <2 years of age.

Methods

Case Reporting and Data Collection

Surveillance of invasive disease caused by S. pneumoniae began January 1, 1999. Surveillance for invasive H. influenzae, group A streptococci (GAS), group B streptococci (GBS), and Neisseria meningitidis commenced January 1, 2000. Cases reportable to ICS are defined as persons from whom an organism under surveillance is isolated from blood, cerebrospinal fluid, or other normally sterile site. Patients with clinical epiglottitis from whom H. influenzae is isolated from an epiglottis swab are also reportable to ICS. Cases are reported to public health officials by physicians or laboratories serving regions under surveillance; this includes patients managed outside of the region. Unconfirmed cases are not included. Standardized case report forms are completed in the region by trained communicable disease officers and include demographic, clinical, vaccination, and risk factor information. For the vaccine-preventable diseases (caused by S. pneumoniae, Hib, and N. meningitidis), details on the type of vaccine received are not currently available; however, information on the number of doses received is available. Reference laboratory representatives and communicable disease officers from each region participate in quarterly and annual data audits to ensure completeness of case finding and reporting.

Laboratory Methods

A network of laboratories ascertains infection with any of the 5 organisms under surveillance within the region. Invasive isolates are submitted to 1 of 3 Canadian reference laboratories (National Centre for Streptococcus, National Microbiology Laboratory, and Laboratoire de Santé Public du Québec). The reference laboratory confirms the isolate’s identity, determines its serotype or serogroup, and tests for antimicrobial susceptibility. Laboratories also participate in an ongoing quality control program.

Isolates were confirmed as S. pneumoniae by using conventional methods of identification (7). Strains were classified by the capsular swelling technique (8,9) by using commercial antisera (Statens Serum Institut, Copenhagen, Denmark). Antimicrobial drug susceptibility testing was performed by using the broth microdilution method consistent with National Committee for Clinical Laboratory Standards guidelines current at the time of testing (10,11).

M typing was performed on all submitted GAS isolates according to standardized methods (12) by using M type-specific antisera prepared in-house. Antisera to 61 of 86 internationally recognized M types, representing the most common M types (13), were available; strains for which an M type could not be assigned were classified as M nontypeable. GBS serotyping was performed according to conventional serologic techniques (14) by using type-specific antisera prepared in-house. Antisera were available for all 9 internationally recognized serotypes (Ia, II, Ib, II, III, IV, V, VI, VII, and VIII). For both organisms, Lancefield hot-acid extracts were prepared from the clinical isolates and tested in Ouchterlony immunodiffusion agar slides with appropriate control strains.

H. influenzae was confirmed by standard biochemical tests (15), and biotypes were determined according to current nomenclature (16). Serotyping was conducted by using a slide agglutination assay with antisera from commercial sources (Difco, Oakville, Ontario, Canada, and Denka Seiken, Tokyo, Japan). N. meningitidis was identi-

<table>
<thead>
<tr>
<th>Region</th>
<th>Total</th>
<th>&lt;2 y</th>
<th>2–4 y</th>
<th>5–19 y</th>
<th>20–64 y</th>
<th>&gt;65 y</th>
</tr>
</thead>
<tbody>
<tr>
<td>Circumpolar*</td>
<td>132,956</td>
<td>4,849 (3.7)</td>
<td>7,414 (5.6)</td>
<td>37,431 (28.2)</td>
<td>77,823 (58.5)</td>
<td>5,439 (4.1)</td>
</tr>
<tr>
<td>Canada†</td>
<td>30,007,095</td>
<td>652,120 (2.2)</td>
<td>1,044,160 (3.5)</td>
<td>6,082,585 (20.3)</td>
<td>18,339,680 (61.1)</td>
<td>3,888,550 (13.0)</td>
</tr>
</tbody>
</table>

*Yukon, Northwest Territories, Nunavut, northern Quebec, and northern Labrador.
†Includes the circumpolar region.
fied by using standard biochemical tests (17). Serogrouping was conducted by using bacterial agglutination with rabbit antisera to the different serogroups. Serotyping and serosubtyping were conducted by using an indirect, whole-cell, ELISA with monoclonal antibodies (18).

Statistical Analysis
Statistical analysis was conducted by using SAS statistical package version 9.1 (SAS, Cary, NC, USA). The analyses were stratified by organism. Crude and age-specific annual incidence was calculated by using total and age-specific population estimates from the Demography Division of Statistics Canada for 2001 (J). Because of the small numbers of cases per year, 3-year period–based annual incidence rates were calculated for each organism to determine time trends. Rates were calculated for 2 periods (1999 for S. pneumoniae or 2000 for H. influenzae, GAS, and GBS to 2002 and 2003–2005). Regression analysis was conducted to detect trends in crude annual incidence rates over time. Crude incidence rates by ethnicity were calculated by using population data from the aboriginal population profile, which was developed from 2001 census data (2). Because of the lack of additional period estimates, no trend analyses were conducted on data by ethnicity.

Information about PCV7 program implementation was collected from the regions (Table 2). The prevaccination period was defined as 1999–2002, and the program implementation period was defined as 2003–2005. Because Labrador implemented its vaccination program during the second period, data from this region on the effect of the vaccine were excluded from the analysis. Sensitivity analyses that included data from Labrador in both arms was conducted to ensure this did not alter the results. Annual incidence of S. pneumoniae for all ages and the number of cases in children <2 years of age were compared for the 2 periods. Bivariate analysis was conducted by using $\chi^2$ and Fisher exact tests.

Results
There were 251 confirmed cases of invasive disease caused by S. pneumoniae in northern Canada from 1999 through 2005. During 2000–2005, 62 cases of invasive disease caused by H. influenzae, 45 caused by GAS, 17 caused by GBS, and 6 caused by N. meningitidis were reported. Because of the small number of N. meningitidis cases reported, no further disease-specific analyses were conducted for this organism.

Cases and Incidence Rates
In the ICS region, the crude annual incidence rate for S. pneumoniae was highest in 2001 (38.4/100,000 population) and lowest in 2005 (17.3/100,000 population), but this downward trend was not statistically significant ($p = 0.119$ by F test for slope). The age-specific incidence rate in children <2 years of age decreased during 2000–2004 but increased in 2005. However, these incidence rates are based on a small number of cases and changes in rates over time should be interpreted with caution. The incidence rates in the population ≥65 years of age, who were eligible for the 23-valent polysaccharide vaccination, did not show any trend (Table 3). During 1999–2002 and 2003–2005, the crude annual incidence rates were 34.0 and 23.6/100,000 population/year, respectively. Although this finding suggests a decreasing incidence over the 2 periods, data from additional periods are necessary to determine if this is reflective of a trend.

Among the 240 (95.6%) of 251 S. pneumoniae cases with serotype information, the most common serotypes were type 1 (30.4%), type 8 (8.8%), type 14 (7.9%), type 4 (6.3%), and type 6B (5.8%). A total of 47 (60%) of 76 cases in children <2 years of age were caused by PCV7 serotypes. Among persons ≥65 years of age, 23 (88.5%) of 26 cases were caused by serotypes in the polysaccharide pneumococcal vaccine.

There were no trends in overall crude annual incidence rates of H. influenzae or GBS (Table 4). The crude annual incidence rate of H. influenzae was lowest in 2003 (4.5/100,000 population) and highest in 2001 (13.5/100,000 population). Among 59 cases with serotype information, 31 (59%) were H. influenzae type a (Hia); 73.3% of these cases were in children <2 years of age. Eight cases (13.6%) of Hib were reported during the surveillance period: 6 in infants <5 months of age, 1 in a child 18 months of age, and 1 in an adult. Two infants had no vaccine information. The adult and 1 infant had not been vaccinated; the remaining 4 children had received only 1 Hib dose. Thus, none of these cases were considered vaccine failures. GAS incidence in...
creased significantly during 2001–2005 ($F = 229.371, p = 0.01$). The largest number of cases ($n = 14$) was reported in 2005 with a crude incidence rate of 10.5/100,000 population. The increase in GAS cases was not clustered by region, period, or serotype. A total of 1–4 cases of GBS were reported in the region annually, for a crude annual incidence range of 0.8–3.0/100,000 population.

**Demographic Characteristics**

Infections with *S. pneumoniae*, *H. influenzae*, and GAS were more common in male patients (59.8%, 58.3%, and 62.2%, respectively). Seventy-one percent of all cases of GBS were among female patients; 17.6% (3/17) of cases of GBS were among newborns <1 month of age. All of the newborn cases occurred in the early neonatal period. These 4 organisms disproportionately affect children <2 years of age and persons >65 years of age. Although <4% of the surveillance population was <2 years of age, 21%–67% of the infections occurred within this age group. Similarly, adults >65 years of age had a higher proportion of cases than the surveillance population they represent (Figure).

Data on patient ethnicity was missing for 42 (11.0%) of 381 cases of invasive bacterial disease. Aboriginal persons represented 59% of the surveillance population and 84% of cases of *S. pneumoniae*, 92% of *H. influenzae*, 93% of GAS, and 53% of GBS. To assess changing incidence over time by ethnicity, the surveillance period was divided into 2 periods (1999–2002 and 2003–2005 for all other organisms). For all but GBS, the crude annualized incidence rates were higher in the aboriginal population than in the non-aboriginal population (Table 5). For *S. pneumoniae*, the disparity between aboriginal persons and non-aboriginal persons decreased from 4.6-fold to 2.5-fold between the 2 periods. Among aboriginal persons, the GAS rate in the second period was nearly double that seen in the first period. Six of 8 case-patients with Hib and all 27 case-patients with Hia for whom ethnicity data were available were among aboriginal persons.

**Clinical Findings and Outcomes**

Information on clinical findings was available for 380 of 381 cases (Table 6). The most common primary clinical finding for invasive *S. pneumoniae* was pneumonia (64.5%), followed by bacteremia/septicemia (21.5%). Among invasive GAS cases, the most common primary clinical finding was cellulitis (31.1%); necrotizing fasciitis accounted for 11.1%. For cases with *H. influenzae* or GBS, the most common primary clinical findings were bacteremia/septicemia (33.9% and 47.1%, respectively). There were no reported cases of epiglottitis caused by *H. influenzae*.

Cases with GAS had the highest case-fatality rate (18.2%, 8 of 44 cases with outcome data). Two (40%) of 5 cases with GAS and necrotizing fasciitis resulted in death; however, this difference was not statistically significant ($p = 0.065$, by Fisher exact test). The case-fatality rates for infections with the other 3 organisms were 4.8% (11/230) for *S. pneumoniae*, 6.1% (3/49) for *H. influenzae*, and 7.1% (1/14) for GBS. The relative risk for death did not vary by ethnicity ($p = 0.550$, by Fisher exact test).

**Effect of PCV7 Immunization Programs**

Fifty-two cases of *S. pneumoniae* occurred in children <2 years of age. Eight of these case-patients had received ≥1 dose of pneumococcal vaccination; 1 case-patient with PCV7-preventable *S. pneumoniae* had received only 1 dose of vaccine (serotype 6b), 6 case-patients had serotypes that were not preventable with PCV7 (serotypes 19A, 20, 13, 15a, 22, and 22F), and 1 case-patient had no information on serotype. These findings suggest that there were no known cases of vaccine failure.

Numbers of cases of disease caused by *S. pneumoniae* in children <2 years of age were compared during the pre-vaccination and program implementation periods (Table 7). In regions where universal PCV7 infant programs were implemented in 2002 ( Nunavut and northern Québec), 19 cases with PCV7 serotypes were reported during the pre-vaccination period and no cases were reported during the program implementation period. In the other Canadian ICS regions where universal PCV7 infant programs were implemented after 2002 (excluding Labrador), 6 cases of PCV7-
preventable S. pneumoniae disease occurred in the prevaccination period and 3 cases in the program implementation period. A χ² analysis showed that the number of cases of PCV7-preventable illness by vaccination region was statistically significant (p = 0.019, by Fisher exact test). These results are conservatively biased because PCV7 was available in all regions in the program implementation period and a universal vaccination program was started during the later half of 2005 in the Yukon, which may have reduced the number of cases seen in the comparison area. Our findings suggest that early implementation of universal PCV7 programs was associated with a reduction in PCV7-preventable illness in children <2 years of age. A sensitivity analysis including Labrador in both arms (program or no program) did not change the statistical significance of the difference because 1 case in Labrador was not vaccine preventable.

### Discussion

To our knowledge, this study is the first comprehensive surveillance report on invasive bacterial diseases in the Canadian Arctic. Disease caused by S. pneumoniae continues to be a serious problem in northern Canada. The annual rate for 2003–2005 was 23.6/100,000 population/year, which is more than twice the reported rate of invasive pneumococcal disease in the overall Canadian population (9.1/100,000 population in 2004) (19). Although this rate is lower than that seen in the earlier period (1999–2002, 34.0/100,000 population), additional data will be needed to determine if the decreasing trend is sustained. The decrease in the disease incidence may be partly attributed to PCV7 programs, as well as the mass pneumococcal polysaccharide vaccination campaigns launched in 2001 and 2002 in response to outbreaks of serotype 1 disease in parts of the region, which reduced the occurrence of this predominant serotype in subsequent years (20,21). Reduction in the number of cases among children <2 years of age in regions where universal infant PCV7 programs were implemented in 2002 (northern Québec and Nunavut) is an early indicator of the effect of the vaccination program. This finding is likely a conservative assessment of the effect of the program, given the staggered implementation of universal vaccine programs in the circumpolar region.

Although aboriginal persons represented 84% of invasive S. pneumoniae cases, a substantial reduction in disease incidence was demonstrated in the program implementation period. Progress toward elimination of this health disparity has also been reported for indigenous populations in Alaska (22) and Australia (23), where PCV7 has been available since 2001 to all indigenous children <2 years of age. The incidence of S. pneumoniae may be expected to decrease further among young children throughout northern Canada, particularly in aboriginal children, as universal PCV7 programs become fully implemented with sustained high coverage rates.

In contrast, there continues to be a health disparity for invasive H. influenzae disease. Annual period incidence rates for H. influenzae during 2003–2005 were >4-fold higher among aboriginal persons than among non-aboriginal persons. Although Hib disease is rare because of universal Hib vaccination, the greatest number of cases occurred among aboriginal persons, a group known to be at increased risk for Hib disease (24–26). Studies in Alaskan
aboriginal populations suggest that continued low-level nasopharyngeal colonization facilitates transmission to susceptible children (4). Environmental and housing conditions, including overcrowding, are also potential contributing factors to these health disparities (27–29). The data also indicate an apparent emergence of type a disease, with all Hia cases occurring among aboriginal persons. Hia disease has also been reported in aboriginal populations in the United States and Australia (30–32). A possible shift in disease epidemiology to non-b serotypes has been suggested from findings in an adjacent Canadian region (33), whereas a sustained increase in non-b serotypes has not been detected in Alaska (4).

The incidence of GAS in the ICS region has been increasing since 2001. Although changes in rates over time should be interpreted with caution because the number of cases is small, this apparent increase in GAS disease is being monitored. The rate in northern regions is greater than that in the overall Canadian population. In 2004, the incidence of GAS in the ICS population was 7.5 compared with the Canadian rate of 2.7/100,000 population (Public Health Agency of Canada, unpub. data). Aboriginal persons represented the greatest proportion of GAS cases; this has also been observed among indigenous populations in the United States (34) and Australia (35). As with S. pneumoniae and H. influenzae, this may be partially attributed to poverty and crowded living conditions in these populations (36).

A major limitation of the data is that the number of reported cases is too small to permit analysis of smaller areas and subpopulations within the region. In addition, changes in rates over time should be interpreted with caution due to small numbers of cases. It is also expected that the number of reported cases of invasive bacterial diseases is an underestimate. However, the enhanced nature of the surveillance system and regular data audits represent improvements over routine passive surveillance. Laboratory specimens may not have been taken before initiation of empiric treatment, and collection and transportation of clinical specimens are difficult in remote areas experiencing extreme temperatures (20,21). Unfortunately, because vaccine registries have not yet been fully implemented in Canada, immunization coverage rates are not available. This situation limits our ability to evaluate the effect of vaccination programs.

Despite these limitations, data collected by ICS contribute to understanding the epidemiology of invasive bacterial diseases among northern populations in Canada and throughout the world. These data assist in formulation of prevention and control strategies, including immunization recommendations (6). ICS data have also been instrumental in identifying potentially emerging pathogens such as Hia. Continued collection of data will be used to assess the effect of vaccination in this population and monitor serotype replacement, antimicrobial drug resistance, and reductions in disparities in Northern populations.

Dr Degani works at the Child Health Evaluative Sciences Unit at the Hospital for Sick Children in Toronto, Canada. Her research interests include infectious diseases and their prevention in marginalized populations.

### Table 7. Effect of universal PCV7 programs for children <2 y of age in the Canadian circumpolar region*

<table>
<thead>
<tr>
<th>Location, period</th>
<th>No. cases with PCV7 serotypes</th>
<th>No. cases without PCV7 serotypes</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Northern Québec and Nunavut</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pre Vaccination (1999–2002)</td>
<td>19</td>
<td>5</td>
<td>24</td>
</tr>
<tr>
<td>Program implementation (2003–2005)</td>
<td>0</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>Total cases</td>
<td>19</td>
<td>13</td>
<td>32</td>
</tr>
<tr>
<td>Northwest Territories and Yukon</td>
<td></td>
<td></td>
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<tr>
<td>Pre Vaccination (1999–2002)</td>
<td>6</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>Program implementation (2003–2005)</td>
<td>3</td>
<td>3</td>
<td>6</td>
</tr>
<tr>
<td>Total cases</td>
<td>9</td>
<td>5</td>
<td>14</td>
</tr>
</tbody>
</table>

*PCV7, 7-valent pneumococcal conjugate vaccine.

References
