Burden of pneumococcal disease in children in Cuba before the introduction of a novel pneumococcal conjugate vaccine

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Background The Cuban heptavalent conjugate pneumococcal vaccine will be introduced for children beginning in 2020. We estimated the burden of pneumococcal cases and deaths in children 1–59 months in 2015.

Methods Mortality and morbidity attributable to pneumococcus were estimated for each of the three primary syndromes commonly associated with pneumococcus: pneumonia, meningitis, and invasive non-pneumonia, non-meningitis (NPNM). Vaccine randomized clinical trial data were used to estimate the proportion of pneumonia deaths attributable to pneumococcus. Data from Cuba were obtained from two domestic sources: National Bacterial Meningitis Surveillance System and the laboratory register from the National Reference Laboratory. Syndrome-specific pneumococcal mortality proportions were applied to all-cause pneumonia and meningitis death estimates prepared by the World Health Organization Maternal Child Epidemiology Estimation (WHO/MCEE) collaboration. The proportion of pneumonia cases attributable to pneumococcus was applied to estimates of all-cause clinical and severe pneumonia also prepared by the WHO/MCEE collaboration. Pneumococcal NPNM morbidity and mortality estimates were prepared using the ratio pneumococcal meningitis to pneumococcal invasive disease. Estimates were adjusted for HIV prevalence and access to health care.

Results In 2015, pneumococcus was estimated to cause 970 severe cases (uncertainty range, UR=692–1209) and 39 deaths (UR=23–59) among children 1–59 months. The estimated incidence rate of severe pneumonia was 149 (UR=112–170) per 100,000 children 1–59 months. The estimated case fatality ratio (CFR) was 2% (UR=1–2%). Over a period of one year, 22 deaths were attributed to pneumococcal pneumonia (UR=14–19) to pneumococcal meningitis, and 8 (UR=3–17) to pneumococcal bacteremia. The CFR due pneumococcal meningitis was 18% (UR=8–37%) in 2015.

Conclusions Reduced morbidity and mortality due to pneumococcal disease in Cuba could be achieved with the accelerated introduction of a novel PCV product in children.

Keywords: Pneumococcal conjugate vaccine, burden of disease, Streptococcus pneumoniae, Cuba.
Streptococcus pneumoniae (pneumococcus) is a leading bacterial cause of pneumonia, meningitis, and other serious invasive diseases in children. In many settings, pneumococcal disease continues to represent an important cause of morbidity and mortality in children and the elderly. Researchers estimate that pneumococcus caused approximately 300,000 deaths in children globally in 2015 (1). The high burden of pneumococcal disease in many settings has been associated with substantial health care system costs to governments and society (2, 3).

Recognizing the substantial burden of pneumococcal disease occurring in children and the established safety and efficacy of current pneumococcal conjugate vaccine (PCV) products available for use in this age group, the World Health Organization (WHO) recommends that all countries use PCV in their national routine immunization programs (4). Evidence from Latin America has demonstrated the significant impact of PCV on hospitalizations in children due to radiograph-confirmed pneumonia, clinical pneumonia, meningitis, and invasive pneumococcal disease (5). However, concerns about the cost PCV has been a barrier to its introduction in some countries (6).

The Cuban government has a new heptavalent PCV product under advanced clinical development (7, 8). The candidate vaccine includes contains 2 μg of serotypes 1, 5, 14, 18C, 19F, 23F and 4 μg of 6B each conjugated to tetanus toxoid (TT). Pending positive clinical results, the Cuban government is planning to introduce the novel heptavalent PCV product in its national immunization program in 2019 using a 2+1 schedule (9). In the meantime, a randomized clinical trial enrolling infants 2–6 months of age is being conducted to assess the efficacy of this vaccine for use in this age group (RPCEC00000243). Estimates of pneumococcal disease burden are needed to inform the use of this PCV product and to assess the potential impact of the new vaccine. We use globally established methods to estimate pneumococcal morbidity and mortality in children younger than 5 years in Cuba for 2015.

**METHODS**

We used the same methods previously published to estimate pneumococcal disease burden in Cuba (1). We updated a systematic review of pneumococcal invasive disease from 1980–2005 (10) with published and unpublished data through 2014 for the meningitis and NPNM models. We followed the same methods and quality assessment criteria described previously. We searched six global databases (ie, PubMed, EMBASE, Biosis, Cochrane, Global Health, and Pascal) and five regional databases (ie, IMEMR, IMSEAR, LILACS, WHOLIS, WPRIM). Additional pneumococcal data were obtained from two domestic sources: National Bacterial Meningitis Surveillance System and the laboratory register from the National Reference Laboratory at Tropical Medicine Institute “Pedro Kouri” (11). The mortality and morbidity attributable to pneumococcus in Cuba were estimated separately for pneumonia, meningitis, and NPNM. All analyses were conducted using Stata 14 of StataCorp (Stata Corp, College Station, TX, USA).

To estimate pneumococcal pneumonia mortality and morbidity, we applied etiologic fractions to estimates of all-cause pneumonia deaths and cases in Cuba in 2015 prepared by WHO and the Maternal and Child Epidemiology Estimation (MCEE) collaboration (12, 13) For Cuba, all-cause pneumonia deaths are based on hospital-based vital registration data. Table 1 describes the ICD10 codes used to assign deaths to pneumonia and meningitis causes of death (14). All-cause pneumonia and severe pneumonia cases were based on the prevalence of pneumonia risk factors from Multiple Indicator Cluster Surveys conducted in Cuba (15–21). To determine the proportion of pneumonia mortality and morbidity attributable to pneumococcus, we used the vaccine probe approach we previously described in detail (1, 22) Briefly, efficacy values

<table>
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<tr>
<th>CAUSE OF DEATH</th>
<th>ICD-10 CODES</th>
<th>ICD-9 CODES</th>
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<tbody>
<tr>
<td>Acute respiratory infections</td>
<td>H65-H66, J00-J22, J85, P23, U04</td>
<td>460-466, 480-487, 381-382, 513, 770.0</td>
</tr>
<tr>
<td>Meningitis/encephalitis</td>
<td>A20.3, A32.1, A39.1, G00-G09</td>
<td>036, 320, 322-326</td>
</tr>
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from PCV randomized control trials were adjusted to account for pneumococcal serotype distribution, incomplete vaccine efficacy, and the proportion of pneumonia attributable to *Haemophilus influenzae* type b (Hib), since all PCV trials were conducted following the introduction of Hib vaccine. Following these adjustments, efficacy values were combined into summary estimates. We standardized WHO pneumonia case definitions (15, 16, 23). PCV trials did not assess vaccine efficacy against pneumonia mortality. Therefore, we used efficacy against radiography-confirmed, primary end-point pneumonia as a proxy for this value.

We estimated pneumococcal meningitis deaths by applying an estimate of the proportion of meningitis deaths in Cuba attributable to pneumococcus to all-cause meningitis deaths in Cuba prepared by the WHO/MCEE collaboration. We divided the pneumococcal meningitis death estimate by an estimate of pneumococcal meningitis CFR for Cuba derived from the literature and adjusted for access to care. The proportion of meningitis deaths attributable to pneumococcus in Cuba was determined by meta-analyses of observational studies from the literature that provide data for the most common causes of bacterial meningitis (ie, pneumococcus, Hib, and *Neisseria meningitidis*). We estimated pneumococcal meningitis case fatality using data from the literature from Cuba and epidemiologically relevant settings. Reported case fatality values reflect mortality in children with access to care. Therefore, we adjusted estimates of pneumococcal meningitis case fatality to account for the higher case fatality assumed for those without access to care. We used the proportion of children seeking care for pneumonia symptoms from Multiple Indicator Cluster Surveys in Cuba as a proxy for this value. We estimated pneumococcal morbidity due to other invasive disease using studies reporting on the ratio of pneumococcal NPNM cases to pneumococcal meningitis cases. Estimates from settings that are epidemiologically to Cuba in 2015 were then combined into summary estimates. We stratified pneumococcal NPNM cases by severe and non-severe cases. We used a summary estimate of case fatality for pneumococcal NPNM from the literature to estimate NPNM in children.

We estimated pneumococcal pneumonia and meningitis deaths in HIV-infected children using estimates of HIV prevalence in children less than 59 months provided by UNAIDS (personal communication) and estimates of the relative risk for invasive pneumococcal disease in HIV-infected children from published meta-analyses (22) Cuba has not yet introduced PCV and so we did not adjust for vaccine use for 2015. These estimates are reported with uncertainty ranges (UR) using the same methods described elsewhere (1).

### RESULTS

The estimates of pneumococcal disease burden for 2015 are presented in Table 2. We estimated that pneumococcus is responsible for 39 deaths in children 1–59 months in Cuba in 2015. 22 deaths were attributed to pneumococcal pneumonia (UR=16–23), 9 deaths to meningitis (UR=14–19) and 8 to bacteremia (UR=3–17). Despite the child mortality rate in children less than five years is low (5.5 per 1,000 live births) compared with other Latin America and the Caribbean countries, pneumococcal pneumonia deaths represent 14% of all deaths in children 1–59 months and pneumococcal meningitis deaths represent 5% of all deaths.

In addition to the 22 deaths (UR=16–23) due to pneumococcal pneumonia in 2015, we estimated 1330 (UR=1149–1581) pneumococcal pneumonia cases, of which 862 (UR=64–983) were estimated to be severe pneumococcal pneumonia cases. Pneumococcal pneumonia deaths in Cuba correspond to a mortality rate of 4 (UR=3–4) deaths per 100,000 children 1–59 months.

<table>
<thead>
<tr>
<th>Indicator and source</th>
<th>Value</th>
</tr>
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<tbody>
<tr>
<td>Population less than 5 years in 2015 (27)</td>
<td>587755</td>
</tr>
<tr>
<td>Child mortality rate in 2015 (28)</td>
<td>5.5 per 1000 live births</td>
</tr>
<tr>
<td>Access to care in 2015 (29, 30)</td>
<td>93%</td>
</tr>
<tr>
<td>% of deaths in children 1–59 months due to meningitis in 2015 (15)</td>
<td>5%</td>
</tr>
<tr>
<td>Pneumococcal conjugate vaccine coverage in 2015 (16)</td>
<td>0%</td>
</tr>
<tr>
<td>% of deaths in children 1–59 months due to pneumonia in 2015 (26)</td>
<td>14%</td>
</tr>
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</table>

Table 2. Indicators used for modeling pneumococcal disease burden for Cuba in 2015
The pneumococcal pneumonia incidence rate in 2015 was 230 (UR=198–273) cases per 100,000 children and the severe pneumococcal pneumonia incidence rate was 149 (UR=112–170) cases per 100,000 children (Table 3).

Pneumococcal meningitis was estimated to cause 51 (UR=22–105) cases in 2015 with a corresponding incidence rate of 9 (UR=4–18) cases per 100,000 children 1–59 months. The pneumococcal meningitis case fatality in 2015 was estimated to be 18% (UR= 8–37%) in 2015 (Table 4). Similarly, there were 152 (UR=66–315) cases of non-severe pneumococcal NPNM in 2015 in Cuba and 57 (UR=25–119) cases of severe pneumococcal NPNM in 2015. The case fatality for pneumococcal NPNM was also high, at approximately 14% (UR=6–29%) in 2015 (Table 5).

In summary in 2015 in Cuba, there were an estimated 970 (UR=692–1209) severe cases of pneumococcal disease and 39 deaths (UR=23–59) in children 1–59 months.

**DISCUSSION**

This paper documents the burden of pneumococcal disease in Cuba using international standardized international methodology (1). The scope includes a description of invasive and non-invasive disease in children less than five years of age. This constitutes the first comprehensive country-level estimation of pneumococcal disease burden in Cuba that will allow policy maker to develop estimates of the potential impact of childhood disease intervention and determine the cost-effectiveness of vaccination.

Previous studies estimating the burden of pneumococcal meningitis in Cuban children have been conducted (24–26) and were included as part of evidence revised to generate the WHO document reporting the Burden of Pneumococcal Disease and Cost-Effectiveness of a Pneumococcal Vaccine in Latin America and the Caribbean (27).

Establishing the burden of disease and creating demand for the pneumococcal conjugate vaccine is one of the critical first steps required when considering the introduction of a new vaccine (28, 29). Without recognition of the burden of pneumococcal disease, health decision makers will underestimate the value of pneumococcal vaccination. However, estimating the total burden of pneumococcal disease is challenging because standard diagnostics have a low sensitivity and therefore, observational and surveillance studies systematically underestimate pneumococcal disease burden. In Cuba, the epidemiological surveillance system is currently being strengthened and will include collection of more epidemiological data including incidence data from sentinel sites.

The proportion of deaths in Cuban children 1–59 months due to pneumonia in 2015 (14%) was lower than global estimation for the same year, (1) but is 2-fold greater than that reported for America region where several countries have already introduced pneumococcal vaccine as part of routine immunization programs (30) According our current estimations, the incidence rate of pneumonia in Cuban children, mortality rate and case fatality are very
similar to global estimations for 2015. To establish population-based surveillance for X-ray confirmed pneumonia and data from some of these surveillance systems has recently been implemented in Cuba data including incidence data at three sentinel sites and a working group was created in each sentinel site to standardize the categorization of radiological pneumonia.

The median percentage of confirmed cases of bacterial meningitis due to pneumococcus reported in Cuba in 2002 was 27% (UR=26–28). For 2015, we reported 51 cases (UR=22–105) with higher incidence rate than reported for American region, but lower than the global estimation for the same year (13; UR=5–26) (25). However, in comparison, the mortality is very similar than reported by Europe and American region. In addition, the case fatality is lower (18%) compared with global estimations (44%) and American countries (27%).

The main limitation in our data about pneumococcal meningitis is associated with pneumococcal isolation in cerebral fluid and in general, the incidence of pneumococcus may have been underestimated due to lack of collection of blood cultures, poor yield of culture techniques, variation in laboratory practice (including hours of operation), prior antibiotic use and non-uniformly applied case definitions. In this way, the decision to implement a sentinel surveillance system four years prior to PCV introduction (ie, since 2014) will substantially improve the data quality to conduct realistic estimation of burden disease that will permit international comparisons and to measure the impact of pneumococcal vaccine introduction.

CONCLUSIONS

Despite limitation in epidemiological surveillance and microbiological diagnostic that typically underestimate the true burden of pneumococcal disease, we can conclude that pneumococcal disease is relatively common in Cuba and represents an important public health problem. If we take into account that many of these infections are serious and may lead to hospitalizations, permanent disability, and death and the large burden of pneumococcal acute otitis media is also substantial contributor to healthcare system costs in Cuba, the introduction of a novel heptavalent PCV should be a high priority for the country as it will likely reduce the morbidity and mortality due to pneumococcal infections in Cuban children.

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Ethics approval: This study was submitted to the Tropical Medicine Institute “Pedro Kourí” Institutional Ethics Committee (IPK-IEC) and was approved under the agreement IPK IEC # 17-14/37-16. In addition, the Scientific Council of the Finlay Vaccine Institute approved the study.
Availability of data and materials: The data are available from the corresponding author (Nivaldo Linares-Pérez, nlinares@finlay.edu.cu) upon reasonable request, subject to obtaining clearance from the Pneumococcal Project Coordination Working Group of the Finlay Vaccine Institute of Cuba. For information on how to obtain clearance to access the data, contact the Pneumococcal Project Coordinator at +53 72086086 or dsantana@finaly.edu.cu.

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Contributors of authors: NLP, BW and MET drafted the manuscript. NLP, BW and MET participated in the design. BW worked on the methodology, the literature review, the mathematic modelling, and the interpretation of the data. MR and GT were instrumental in the data collection for the study. All authors contributed to manuscript revisions and read and approved the final manuscript.

Competing interests: The authors completed the Unified Competing Interest form at www.icmje.org/coi_disclosure.pdf (available upon request from the corresponding author), and declare no conflicts of interest.

REFERENCES


