Serotype distribution and clinical characteristics associated with *Streptococcus pneumoniae* among Chinese children and adults with invasive pneumococcal disease: a multicenter observational study

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ABSTRACT

Few studies in China focused on serotypes of *Streptococcus pneumoniae* in patients with invasive pneumococcal disease (IPD). We aimed at investigating the serotype distribution for IPD-causing *S. pneumoniae* and vaccine coverage among Chinese children and adults. This was a multicenter, observational study to collect *S. pneumoniae* isolates from normal sterile sites and IPD-related clinical information among children and adults. Serotyping was performed by a Capsule-Quellung reaction test using type-specific antisera. The study collected a total of 300 eligible isolates (pediatric = 148, adult = 152) were serotyped in a central laboratory. The most prevalent serotypes were 19A (20.9%) and 23 F (20.3%) in the pediatric group; 3 (21.7%) and 19 F (11.8%) in the adult group. PCV10 had low-to-moderate serotype coverage rates for children (60.8%) and adults (34.2%). PCV13 and PPV23 had high coverage rates for children (89.9%, 93.2%) and adults (70.4%, 82.9%), respectively. Investigational PCVs including PCV15 and PCV20 had high estimated coverage rates in children (89.9%, 93.9%). The study identified 269 subjects with IPD reported as the primary diagnosis in the medical records. Sepsis (48/136, 35.3%) and pneumonia (48/133, 36.1%) had the highest occurrence in the pediatric and adult groups, respectively. Study findings showed that non-PCV7 *S. pneumoniae* 19A and 3 were the most prevalent serotypes in Chinese children and adults, respectively. High-valent vaccines had similar coverage rates and may have a greater potential in preventing IPD.

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Introduction

*Streptococcus pneumoniae* causes bacterial infections, namely pneumococcal disease, in children and the elderly worldwide. Pneumococcal disease includes serious infections such as meningitis and pneumonia, as well as common illnesses such as sinusitis and otitis media. Pneumococcal disease can be invasive (IPD), defined as an infection confirmed by the isolation of *S. pneumoniae* from a normally sterile body site. The WHO estimated that pneumococcal disease caused approximately 5% of total deaths among children <5 y in 2008. In the pre-vaccination era, incidence rates of IPD ranged from 15-49/100,000 in North America, 11-27/100,000 in Europe, and up to 216/100,000 in specific Asian regions. IPD is age-specific, in which 38%, 54% of all reported cases occurred in children <2 y and adults above 50 y of age, respectively. The incidence of IPD in children <2 y was 44.4/100,000 in Europe, 167/100,000 in the US, and 60-797/100 000 in African countries. In adults, mortality rates of IPD were 11-30% in the western world and 26–30% in Asia countries. Mortality is greatest in younger infants and elderly patients. The capsular polysaccharides (CPS) on the surface of *S. pneumoniae* are the primary bacterial virulence and the basis for serotyping, thereby serving the immunogenic targets as covered by *S. pneumoniae* vaccination. There are over 90 identified serotypes of *S. pneumoniae*, among which IPD attributes to a relatively small subset. Serotypes differ in...
exhibiting invasive potential to cause IPD and that the prevalence of different serotypes can vary by geographic region. Studies suggested that serotypes 1, 5 and 7 were more likely to cause invasive disease, and 6, 9, 14, 19 and 23 were predominantly in pediatric populations. In children <5 y, serotypes 1, 5, 6A, 6B, 14, 19 F, and 23 F are common causes of IPD globally whereas 18 C was common in some western countries. Therefore, inclusion of these serotypes in different vaccine preparations is critical.

Heptavalent pneumococcal conjugate vaccines (PCV) significantly reduced the burden of IPD in young children. PCV7 (covers serotypes 4, 6B, 9 V, 14, 18 C, 19 F and 23 F) also reduced the disease incidence among non-vaccinated children and adult groups due to herd immunity.

Relatively newer PCVs (PCV10, PCV13) have been available to cover more IPD-specific serotypes. Evidence has suggested that the coverage of IPD serotypes by PCV7, PCV10/PCV13 was ≥49%, ≥70%, respectively, across geographic regions. Recently, 15- and 20-valent PCVs (PCV15, PCV20) with additional serotypes have been investigated in phase 2 and 3 clinical trials. In China, PCV7 was on the market since 2008 for healthy children under 5 y old. In addition, 23-valent pneumococcal polysaccharide vaccine (PPV23) is indicated for healthy children aged 2 y or older and the elderly. These vaccines were not included in the China's national immunization program (NIP) and the immunization coverages were generally low. There were no PCVs available since 2015 until PCV13 received approval in China in late 2016 and launched in the market in 2017.

Nevertheless, emergent non-vaccine serotypes increased as causes of IPD following the extensive use of PCVs. Serotype replacement by 19A was one of the primary causes of childhood IPD (>20.0%) in PCV7 accessed countries. Also, non-PCV13 serotypes such as 22 F, 12 F and 33 F contributed to 42.2% of childhood IPD cases in countries where higher valent vaccines were introduced. Therefore, examining epidemiological effects of conjugate vaccines by investigating serotype distribution associated with pneumococcal diseases is of priority before universal immunization in new settings. Although studies collecting serotype data were conducted globally for children with IPD, relatively fewer studies focused on adults. Limited data in China documented IPD epidemiology and serotype distribution. Our objectives were to study the serotype distribution of IPD-causing S. pneumoniae and potential coverage by conjugate and polysaccharide vaccines among Chinese children and adults.

Methods and materials

Study design

This is a multicenter, observational study to collect S. pneumoniae isolates and corresponding IPD clinical information from both pediatric and adult subjects treated in 27 tertiary hospitals in China. Study sites were screened and selected individually by balancing the geographic location (north vs. south) and the evaluation of the investigator's (clinical microbiologist) capability in collecting S. pneumoniae isolates, especially those from children, in the past three years at the time of the site initiation.

The site's routine clinical practice in diagnosing and treating IPD was also reviewed to meet the study needs. Between 2012 and 2015, the investigators obtained S. pneumoniae isolates once the subject had microbiologically confirmed IPD and clinical information was prospectively collected at the site. A retrospective study phase was conducted concurrently to collect cultured isolates from the microbiology laboratory and review the subject's medical records for IPD primary diagnosis within the last three years upon the initiation of each site. Merck Sharp & Dohme (MSD) designed and sponsored the study and analyzed the data. The study was conducted following the guidelines of the International Conference on Harmonization and local regulatory guidance and was approved by independent ethics committees of all study sites.

Patients and S. pneumoniae isolates

The patient was defined as a study case (subject) once IPD was microbiologically confirmed by S. pneumoniae cultured from specimens obtained through a normally sterile site, including blood and cerebrospinal fluid (CSF) or, less commonly, joint, pleural, and pericardial fluid. IPD was defined as isolation of S. pneumoniae from a normally sterile site in an individual with clinical signs and symptoms of invasive disease. Patients were not eligible as an IPD study case if they met the definition for non-IPD as proper S. pneumoniae cultures related to acute otitis media or pneumococcal pneumonia with an absence of bacteremia. Also excluded were patients whose S. pneumoniae cultures sampled from the nasopharynx or whose age information was missing. All patients were treated in the routine clinical practice where isolates were collected from both inpatient and outpatient departments at each site by the investigators with clinical microbiology specialty. Patients or their legal representatives gave written informed consent before any study procedure commenced. The ethics committee at each study site waived the informed consent process for IPD patients identified from the retrospective study phase in which acceptable conditions precluding the consent existed (e.g. death or discharge).

Study procedure and assessments

All isolates were transported to a central microbiology laboratory for S. pneumoniae identification and in vitro susceptibility tests (Department of Laboratory Medicine, Peking Union Medical College Hospital). In the study, one enrolled patient contributed one isolate to the central laboratory. An additional isolate was not required unless sample processing was not appropriate during the transportation or in the laboratory. The subject's demographic and clinical data were collected via a prospective fashion or a review of medical records at the site. All isolates were then transported to a disease control central laboratory (National Institute for Communicable Disease Control and Prevention, Chinese Center for Disease Control and Prevention) for serogrouping and serotyping after all microbiology testings have been completed.

Sero-typing

A simplified chessboard system processed by Pneumotest-kit (Demark, Serum Institute) was applied to determine the
serotype grouping of all isolates. While S. pneumoniae serotyping was done by a Capsule-Quellung reaction test using type-specific antisera (Statens Serum Institut, Copenhagen, Denmark) against all 90+ serotypes. The isolates identified as S. pneumoniae that reacted negatively were classified as ‘nontypable’.

Antimicrobial in vitro susceptibility
The Agar dilution method and E-test methods were used to determine the antibiotic susceptibility of isolates to 17 commonly prescribed antibiotics. The CLSI 2016 criteria for minimum inhibitory concentrations (MICs) were applied to classify the susceptible, intermediate, and resistant isolates. Separate interpretive breakpoints for nonmeningeal versus meningeal isolates were used to define penicillin and parenteral cephalosporin resistance.

Demographics and clinical information
The demographic characteristics including age, gender, rural/urban, and geographic region were collected for both retrospective and prospective study phases. Clinical information including IPD primary and secondary diagnoses, site of culture specimen and antimicrobial regimen were also collected.

Study objectives
The study objectives were to determine the serotype distribution for IPD-causing S. pneumoniae and potential serotype coverage by conjugate vaccine (PCV10, PCV13, and PCV15) and polysaccharide vaccine (PPV23) in Chinese pediatric and adult populations. Post-hoc analysis was made to investigate serotype coverage by PCV20 which is under clinical investigation.

Statistical analysis
Sample size consideration
The study was descriptive and no priori hypothesis was made for study objectives. Sample size was dependent upon the precision of estimation (two-sided 95% confidence interval) on the proportions of combined serotypes 22 F/33 F which the investigational PCV15 serotype may potentially cover. A sample size of 140 from a studied population (pediatric or adult) yielded a two-sided precision of 8.0% for the estimation, assuming that the combined serotypes 22 F/33 F accounted for 5.0% of serotypes identified for all isolates. Meanwhile, this sample size was statistically sufficient if an estimated proportion of PCV15 coverage was 80% with a two-sided precision of 14%. The Clopper-Pearson Exact method was applied as the formula for the calculation. This sample size was also statistically sufficient if an estimated proportion of PCV15 coverage was 80% with a two-sided precision of 14%. Considering a loss by an approximately 5-10% of all isolates during study execution, approximately 150 isolates was to be collected from in pediatric and adult populations, respectively. Therefore, a total sample size of 300 isolates was collected for the study. Due to operational difficulties, randomized sampling was not conducted for the study.

Analysis population
Subjects with proven S. pneumoniae isolates and evaluable clinical information were included for the analyses. The analyses were performed in pediatric (<18 y of age) versus adult study populations.

Analysis of study objectives and other variables
For serotype distribution, number and percentage of total isolates were provided by serotype. Vaccine coverage was summarized by number and percentage and corresponding 95%CI was given. Resistant isolates to antimicrobials were interpreted by MICs and were summarized by number and percentage. For demographics and clinical data, descriptive statistics were made to display the results. Chi-square or t-test was used to test the significance of difference in categorical or continuous variables wherever appropriate. Subgroup analysis by age intervals was performed for serotype distribution and vaccine coverage. There was no imputation for missing IPD primary diagnosis in the study. Statistical analyses were performed using SAS 9.1 and SAS JMP 14.3 (SAS Institute, Cary, NC, USA), and a p value of 0.05 was considered statistically significant whenever applicable.

Results
S. pneumoniae isolate collection
Figure 1a displays the study enrollment and Figure 1b shows the site geographic distribution. The investigators identified a total of 312 S. pneumoniae isolates cultured between 2010 and 2015, among which 300 (pediatric = 148, adult = 152) were included for serotyping and study analysis. The most frequent reason for the exclusion from serotyping was storage issue in the central microbiology laboratory (pediatric = 3, adult = 4). In the pediatric group, more isolates (102/148, 68.9%) were collected through a retrospective phase; whereas in the adult group, more isolates (93/152, 61.2%) were obtained in the prospective study phase.

In the study, there were 39 distinct S. pneumoniae serotypes identified in the central laboratory. Fewer serotypes (21) were obtained from the pediatric group as compared with those (37) from the adult group.

Demographics and clinical characteristics
The subject’s demographic and clinical information are shown in Table 1. The subject’s mean age was 2.6 and 54.3 y in the pediatric and adult groups, respectively. Gender and rural residence did not differ significantly between the two groups; however, the North China region had more isolates for the adult group as compared to that for the pediatric group (62.8%) versus 55 (37.2%). Most isolates were cultured from the blood (72.3%) and the cerebrospinal fluid (19.3%), among which there was no significant difference in the source of specimens between the two groups. Cephalosporins (58.8%) and carbepenems (29.6%) were more frequently prescribed in the pediatric and the adult groups, respectively.
Serotype distribution and vaccine coverage

The most prevalent *S. pneumoniae* serotypes identified were 19A (15.7%), 19F (15.7%), 23F (13.7%), 3 (12.7%) and 14 (7.0%) among all subjects. Figure 2 exhibits numbers of *S. pneumoniae* isolates and cumulative percentages of serotype distribution and corresponding vaccine coverages by PCVs and PPV23 in the pediatric and adult groups. Full results of coverage rates and corresponding 95% CIs are shown in Supplemental Table 1. The most prevalent serotypes were 19A (20.9%), 23F (20.3%), 19F (19.6%), 14 (8.1%) and 6B
There was one isolate serotyped 22 F (0.7%, 95% CI: 0.0%, 3.6%) in the adult group. Rates of serotype coverage were generally higher in the pediatric group as compared with those in the adult group. PCV10 had low-to-moderate serotype coverage rates for the adult group (34.2%, 95% CI: 26.7, 42.3) and the pediatric group (60.8%, 95% CI: 52.5, 68.7). PCV13, PPV23 and the investigational PCV15 and PCV20 had higher serotype coverage rates in the pediatric group.

| Variable                        | Children n = 148 | Adults n = 152 | Total n = 300 |
|---------------------------------|------------------|----------------|---------------|
| Age (years)                     | 2.6 (3.3)        | 54.3 (17.4)    | 28.9 (28.8)   |
| Gender                          |                  |                |               |
| Male                            | 95 (64.2)        | 102 (67.1)     | 197 (65.7)    |
| Female                          | 52 (35.1)        | 50 (32.9)      | 102 (34.0)    |
| Rural residence†                | 58 (39.2)        | 66 (43.4)      | 124 (41.3)    |
| Geographic region&              |                  |                |               |
| North                           | 55 (37.2)        | 94 (62.8)      | 149 (49.7)    |
| South                           | 93 (62.8)        | 58 (38.2)      | 151 (50.3)    |
| Duration of antibiotic treatment (Day) | 5.3 (3.8)        | 5.3 (3.9)      | 5.3 (3.8)     |
| Blood                           | 114 (77.0)       | 103 (67.8)     | 217 (72.3)    |
| Cerebrospinal fluid             | 26 (17.6)        | 32 (21.1)      | 58 (19.3)     |
| Joint                           | 0 (0.0)          | 2 (1.3)        | 2 (0.7)       |
| Pleural                         | 7 (4.7)          | 4 (2.6)        | 11 (3.7)      |
| Others                          | 1 (0.7)          | 11 (7.2)       | 12 (4.0)      |
| Antibiotic used                 |                  |                |               |
| Cephalosporins                  | 87 (58.8)        | 67 (44.1)      | 154 (51.3)    |
| Carbapenems                     | 29 (19.6)        | 45 (29.6)      | 77 (25.7)     |
| Quinolones                      | 7 (4.7)          | 8 (5.3)        | 15 (5.0)      |

*Data are expressed as mean (SD) or count (%); †Based on patient’s self-reporting and/or medical charts review. &Judged by patient’s information or the site location if patient information missing.
Figure 3 shows vaccine coverage rates by age subgroups.

Data are expressed as percent (%) and are stratified by age subgroups (n): <2 (76), 2–4 (47), 5–17 (25), 18–64 (113), and ≥65 (39).

Invasive pneumococcal disease

The results of IPD overall and by serotype distribution are summarized in Table 2. Individual serotypes were defined if numbers of collected isolates >10. A total of 269 patients with a primary diagnosis of IPD were confirmed by the investigators (children = 136, adult = 133). Only the primary diagnosis for each IPD patient was identified at the time of isolate collection. Sepsis had the highest occurrence [35.3% (48/136)] in the pediatric group and pneumonia was the most prevalent [36.1% (48/133)] in the adult group. Meningitis accounted for 21.3% (29/136), 17.3% (23/133) in the pediatric and the adult groups, respectively.

In each IPD primary diagnosis, serotype 19A (37.8%, 14/37) and 3 (25.0%, 12/48) had the highest proportions in children and adults who had pneumonia, respectively. In children with sepsis, 19 F accounted for 22.9% (11/48) of all serotypes identified. Serotype 23 F infected 31.0% (9/29) of all IPD children diagnosed with meningitis.

Non-PCV7/PCV13 serotypes

Non-PCV7 and PCV13 serotypes and associated IPD and in vitro susceptibility results are presented in Table 3. Serotype 19A was the prevalent among all non-PCV7 isolates in the pediatric group [49.2% (31/63)]. Among 29 IPD children contracted 19A, 14 (48.3%) had pneumonia and 8 (27.6%) developed sepsis. Serotype 3 was most commonly seen in non-PCV7 isolates in the adult group [30.8% (33/107)]. Among 28 IPD adults contracted serotype 3, 12 (42.9%) had pneumonia and 6 (21.4%) had bacteremia. Serotype 19A had exhibited a relatively high non-susceptibility to commonly prescribed antibiotics against IPD including ceftriaxone and meropenem in both children [ceftriaxone: 48.4% (15/31)] and adults [meropenem: 87.5% (14/16)].

The number of individual non-PCV13 isolates (n = 15) was relatively fewer in the pediatric and the adult groups. Serotype 15 C had 3 (20.0%) isolates in the pediatric group and serotype 20 had 6 (13.3%) in the adult group, respectively.

Discussion

Findings from this multicenter, observational study suggested that the most prevalent S. pneumoniae serotypes were 19A, 19 F, 23 F, and 3 across different regions in China. Non-PCV7 serotype 19A and 3 was most prevalent in children and in adults, respectively, and 22 F/33 F were rare. New conjugate vaccines provided higher rates of serotype coverage in the pediatric and adult groups.

Relatively fewer epidemiological studies conducted in China34 and in Asia35 to study S. pneumoniae serotypes in IPD. Studies on IPD children reported data from the 1980s36 or focused serotypes from invasive pneumonia or non IPDs although more recently conducted.37,38 The ANSORP Study collected 642 isolates from China but only obtained 33 IPD strains.35 Other Asian studies were either geographically different39 or had no associated clinical information40 or limited to specimens from the respiratory tract.41 This study, on the other hand, had obtained 300 serotyped isolates from normal sterile sites in both Chinese children and adults, and is
Table 2. Invasive pneumococcal disease overall and by serotype.

| Population | Children (by serotype) n = 136 | Adults (by serotype) n = 133 |
|------------|--------------------------------|-----------------------------|
|            | Total N = 269                   | Adults                     |
|            | 19A 23 F 19 F 3 14 6B Other     | 19A 23 F 19 F 3 14 6B Other|
| Pneumonia  | 37(27.2) 85(31.6) 14(37.8) 8(21.6) 5(13.5) 0(0.0) 4(10.8) 2(5.4) 4(10.8) | 5(10.4) 1(2.1) 5(10.4) 12(25.0) 2(4.2) 2(4.2) 21(43.8) |
| Sepsis     | 48(35.3) 70(26.0) 8(16.7) 6(12.5) 11(22.9) 3(6.3) 3(6.3) 14(29.2) | 0(0.0) 3(13.6) 3(13.6) 4(18.2) 3(13.6) 1(4.5) 8(36.4) |
| Meningitis | 29(21.3) 52(19.3) 4(13.8) 9(31.0) 5(17.2) 13(4.0) 2(6.9) 1(3.4) 7(24.1) | 3(13.0) 1(4.3) 4(17.4) 3(13.0) 0(0.0) 0(0.0) 12(52.2) |
| Bacteremia | 5(3.7) 20(15.0) 25(9.3) 1(20.0) 1(20.0) 0(0.0) 1(20.0) 1(20.0) 0(0.0) | 2(10.0) 2(10.0) 2(10.0) 6(30.0) 1(5.0) 0(0.0) 7(35.0) |
| Other      | 17(12.5) 20(15.0) 37(13.8) 2(11.8) 5(29.4) 4(23.5) 0(0.0) 2(11.8) 0(0.0) 4(23.5) | 3(15.0) 3(15.0) 3(15.0) 3(15.0) 1(5.0) 6(30.0) |

Data are expressed as n(%); 269 isolates out of 300 corresponding to 269 patients’ IPD information were collected; IPD was deemed as the primary diagnosis of the invasive disease caused by S. pneumoniae infections; Column percentage was calculated for IPDs in children and adults by population and total; Row percentage was calculated for IPDs by serotype in children and adults; Most clinically significant IPDs are listed; Individual serotypes accounted for >5.0% in one IPD in children or adults are displayed; IPD: invasive pneumococcal disease.
| Non-PCV7 Serotype | Susceptibility n = 63 | Children | Non-PCV13 Serotype | Susceptibility n = 45 | Adults | IPDlogan 153 |
|-------------------|-----------------------|----------|-------------------|-----------------------|--------|--------------|
|                   | TX        | MP        | IPD (n = 58)       | TX        | MP        | IPD (n = 94)       |
| 19A               | 15/31(48.4) | 26/31(83.9) | 14/29(48.3)       | 8/29(27.6) | 4/29(13.8) | 1/29(3.4)       | 2/29(6.9) | 3/16(18.8) | 14/16(87.5) | 5/13(38.5) | 3/13(23.1) | 2/13(15.4) | 0/13(0.0) | 3/13(23.1) |
| 6A                | 1/7(14.3)  | 6/7(85.7)  | 0/6(0.0)          | 2/6(33.3) | 2/6(33.3) | 0/6(0.0)          | 2/6(33.3) | 0/6(0.0) | 1/6(16.7)  | 2/6(33.3) | 1/3(33.3) | 0/3(0.0) | 0/3(0.0) |
| 3                 | 0/5(0.0)   | 1/5(20.0)  | 0/4(0.0)          | 3/4(75.0) | 1/4(25.0) | 0/4(0.0)          | 0/4(0.0) | 1/3(33.3) | 3/3(100.0) | 12/28(42.9) | 3/28(10.7) | 6/28(21.4) | 4/28(14.3) | 3/28(10.7) |
| 20                | 0/1(0.0)   | 0/1(0.0)   | 1/1(100.0)        | 0/1(0.0)  | 0/1(0.0)  | 0/1(0.0)          | 0/1(0.0) | 0/6(0.0) | 0/6(0.0)   | 3/6(50.0) | 2/6(33.3) | 0/6(0.0) | 0/6(0.0) |
| Other             | 4/19(21.1) | 7/19(36.8) | 3/18(16.7)        | 11/18(61.1)| 2/18(11.1)| 0/18(0.0)        | 2/18(11.1)| 2/46(4.3)| 6/46(13.0) | 16/44(36.4) | 8/44(18.2) | 7/44(15.9) | 8/44(18.2) | 5/44(11.4) |
|                   | TX        | MP        | IPD (n = 15)       | TX        | MP        | IPD (n = 43)       |
| 20                | 0/1(0.0)   | 0/1(0.0)   | 1/1(100.0)        | 0/1(0.0)  | 0/1(0.0)  | 0/1(0.0)          | 0/1(0.0) | 0/6(0.0) | 0/6(0.0)   | 3/6(50.0) | 0/6(0.0) | 2/6(33.3) | 0/6(0.0) |
| 15 C              | 0/3(0.0)   | 2/3(66.7)  | 1/3(33.3)         | 2/3(66.7) | 0/3(0.0)  | 0/3(0.0)          | 0/3(0.0) | 0/2(0.0) | 1/2(50.0)  | 1/2(50.0) | 0/2(0.0) | 1/2(50.0) | 0/2(0.0) |
| 23A               | 0/2(0.0)   | 0/2(0.0)   | 1/2(50.0)         | 0/2(0.0)  | 0/2(0.0)  | 0/2(0.0)          | 1/2(50.0) | 0/1(0.0) | 1/2(50.0)  | 1/2(50.0) | 0/1(0.0) | 1/2(50.0) | 0/1(0.0) |
| Other             | 2/9(22.2)  | 3/9(33.3)  | 1/9(11.1)         | 5/9(55.6) | 2/9(22.2) | 0/9(0.0)          | 1/9(11.1) | 1/35(2.2) | 5/45(9.1)  | 13/33(39.4) | 7/33(21.2) | 3/33(9.1) | 5/33(15.2) |

Data are expressed as n(%); non-PCV7 isolates: 170; non-PCV13 isolates: 60; MIC<sub>90</sub> (range) for non-PCV7 isolates (mg/L) – TX: 2 (0.002, 32) in children and 1 (0.002, 2) in adults; MP: 0.75 (0.003, 6) in children and 2 (0.002, 4) in adults; MIC<sub>90</sub> (range) for non-PCV13 isolates (mg/L) – TX: 2 (0.002, 3) in children and 1 (0.002, 2) in adults; MP: 1.9 (0.003, 4) in children and 3.4 (0.003, 4) in adults.

Most clinically significant IPDs were listed; Individual serotypes accounted for >1.0% or n ≥ 4 for non-PCV7 or non-PCV13 in the pediatric or adult group were displayed; IPD was deemed as the primary diagnosis of the invasive disease caused by S. pneumoniae infections.

IPD: invasive pneumococcal disease; TX: ceftriaxone parenteral; MP: meropenem.
the largest in China to study serotype distribution and associated IPD spectrum. The study design allowed a concurrent retrospective collection of isolates. Study data can thus cover serotype distribution in a 6-y interval between 2010 and 2015. In addition, this study emphasized the subject’s exact age information on isolate collection. An analysis on all age subgroups was performed, which provided descriptive results on a specific age group, including those who were younger than 2 or older than 65 y of age.

Our study showed a high prevalence of non-PCV7 serotype 19A in children, especially in those who were less than 5 y of age. This finding was consistent with data previously reported from studies in China between 2005 and 2011.12 Emergent 19A was an issue in western countries and in some Asian countries after the introduction of PCV7. But studies in Singapore,43 South Korea14 and China42 showed an increase of serotype 19A among IPD children before PCV7 became available. PCV7 was marketed in China since 2008 for healthy children <5 y under private health care Results from this study, along with previous evidence, indicated that prevalence of 19A may be due to other factors instead of serotype replacement caused by PCV7 in China. In the adult group, a high proportion of non-PCV7 serotype 3 was observed. A recent study reported 14.0% of serotype 3 in Portuguese adults.45 Similar results were seen in Japan where serotype 3 accounted for 14.7% among serotypes cultured from respiratory tract.41 But a significantly lower distribution of serotype 3 was reported from recently cultured IPD isolates in China and India.53,55,51,54 Studies from 46 USD and Germany47 also reported a low prevalence of serotype 3 (7-8%) in the adult. A few studies demonstrated the lower efficacy of the serotype 3 compared with other serotypes in PCV13 for children,48-50 but with conflicting results in different regions.51,52 A study in Hong Kong monitored PCV 13 immunization program on children between 2015 and 2017. An increase by 59% in the IPD caused by serotype 3 was seen.53 This finding supported the lower efficacy of PCV13 against serotype 3 in an Asian pediatric population. However, studies identifying serotypes in adults were very limited in China, and PCV7 and PCV13 were not indicated for the adult. Results from our study on the prevalence of serotype 3 may warrant further investigations.

In the study, coverage rates of PCV10 in young children were relatively lower compared with that in western countries.1 This can be partly explained by findings that non-PCV7 serotypes were more frequent in China and PCV10-specific serotypes were few. Nevertheless, rates of serotype coverage by higher valent vaccines (PCV13, PCV15 and PCV20) increased greatly as compared with PCV10 among children and adults. Considering high non-susceptibility to antibiotics and associated major IPDs caused by non-PCV7 strains, newer vaccination programs including PCV13, 15 and 20 are in need to substantially prevent IPD episodes. In addition, this study identified 21 and 37 serotypes in children and adults, respectively. Because more serotypes were identified from adults, rates of vaccine coverage are generally lower in the group. Due to the difference in serotype numbers and distribution between these two populations, observed herd effects toward adults from children among western countries22,23 may not benefit to the similar extent in China when new vaccines are firstly indicated in children. In addition, an expected difference between PCV13 and PCV15 did not occur. Data from literature review of pediatric isolates and studies on IPD isolates have indicated 22 F/33 F serotypes were scarce (<1.0%) in China34,37,38,42 as opposed to western countries, for example, the Netherlands.54

Clinically, findings presented from 269 study subjects when analyzing serotype distribution. Pneumonia had the highest occurrence in the adult group. Although the study was not able to distinguish if it was community or hospital acquired, the results were consistent with previously reported.55 Sepsis, however, had been the most common in the pediatric group, followed by pneumonia and meningitis. A majority of isolates in the study were from the blood. Since clinical practice for blood culture may differ in study sites, sampling procedure might not be performed until primary infections clinically progress in pediatric patients. Several serotypes including 19A, 3, 23 F, and 19 F accounted for high proportions in individual IPDs in the pediatric and the adult group. Unlike some studies,55,56 serotypes categorized as invasive did not appear frequently. Because the adult group had a wide serotype spectrum, the role of conjugate and polysaccharide vaccines for individual IPDs must be monitored carefully if vaccinated.

This study has several limitations. Firstly, this study was not designed as a national surveillance program. Comparison in serotype distribution and monitoring particular serotypes over time was not feasible. In addition, the study was not conducted as a cross-sectional study to balance enrollment numbers across site and geographic region by age group. The study team did not perform additional tests including multilocus sequence typing (MLST) to determine specific strains from a certain serotype which caused multiple IPDs. Retrospective data collection was based on review of medical records. Incomplete data caused ascertainment bias in confirming a primary IPD diagnosis. Moreover, the sample size was not sufficient for PCV-targeted serotypes in a particular age subgroup. Last, the study was conducted in major tertiary hospitals in China and was not population-based. The results may not be generalizable to the entire population, especially that in the community and remote areas.

Conclusion
Study findings suggested that non-PCV7 covered S. pneumoniae 19A and 3 were the most prevalent serotypes in Chinese children and adults, respectively. Higher valent conjugate and polysaccharide vaccines had favorable rates of serotype coverage and may have a greater potential in preventing IPD caused by S. pneumoniae.

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B.P., T.W.M and W.J. are employees of MSD China, Shanghai, China. Other authors had none to declare.
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