Impact of Asthma on the Severity of Serious Pneumococcal Disease

Ravneet K Dhillon, Barbara P Yawn, Kwang Ha Yoo, Thomas G. Boyce, Robert M Jacobson, Michaela E McGree, Amy L Weaver, and Young J Juhn

1Department of Pediatric and Adolescent Medicine, Mayo Clinic, Rochester, MN
2Department of Research, Olmsted Medical Center, Rochester, MN
3Department of Internal Medicine, Konkuk University College of Medicine, Seoul, Korea
4Department of Health Sciences Research, Mayo Clinic, Rochester, MN

Abstract

We recently reported an increased risk of serious pneumococcal disease (SPD) in asthmatics. Little is known about the impact of asthma status on the severity of SPD. We compared the severity of serious pneumococcal disease (SPD) between patients with asthma and those without asthma. The study subjects were Rochester, Minnesota residents who developed SPD between 1964 and 1983. SPD and asthma status were ascertained by using explicit predetermined criteria. Severity of SPD was assessed using intensive care unit (ICU) admission rate and total days of ICU stay and hospitalization associated with treatment for SPD. We found that there were no significant differences in severity outcomes between asthmatics (n=11) and non-asthmatics (n=163). Asthma status may increase the risk of SPD but not influence its severity. However, given a small sample size of our study, a larger study needs to be considered to clarify the relationship between asthma and severity of SPD.

Keywords

Asthma; Epidemiology; Pneumococcal disease; Atopy; Severity; Prognosis; Pneumonia; Invasive pneumococcal disease; Vaccine; Serious pneumococcal disease

Introduction

A previous study assessed underlying conditions among patients aged 2–64 years who developed invasive pneumococcal diseases (n=3,469) before the introduction of hepalvalent Pneumococcal Conjugate Vaccine (PCV-7) [1]. Only 50.6% (n=1,755) had at least one condition that was a known indication for either the pneumococcal polysaccharide or...
conjugate vaccine [1]. Thus, a significant proportion of patients who developed Invasive Pneumococcal Disease (IPD) did not have the high-risk conditions. Talbot et al. [2] assessed whether asthma is associated with the risk of IPD and they reported that having a diagnosis of asthma is associated with an increased risk of IPD, (odds ratio (OR): 2.4, 95% Confidence Interval (CI): 1.9–3.1) in their study population who received Medicaid insurance. Subsequently, Juhn et al. [3] reported that asthmatics had a higher risk of Serious Pneumococcal Disease (SPD) including IPD and/or pneumococcal pneumonia during the primarily pre-pneumococcal vaccine era, 1964–1983. As a result, the Advisory Committee on Immunization Practices (ACIP) now recommends a single dose of 23-valent Pneumococcal Polysaccharide Vaccine to adults (19–64 years of age) [4].

Given the increased risk of development of Serious Pneumococcal Disease (SPD) in individuals with asthma, whether asthma status influences severity of SPD may be an important question to be addressed because severity and risk of infection can have different risk factors and mechanisms. Addressing this question may be important in determining the prognosis and management of SPD when asthmatics acquire SPD. Currently, there is no population-based study that examined the relationship between asthma and the severity of SPD. To determine the association between asthma status and severity of SPD, we conducted a retrospective cohort study. In this study, we assessed the impact of asthma status on the severity of SPD in the same study population during the primarily pre-pneumococcal vaccine era, 1964–1983 who was studied for the association between asthma and the risk of SPD.

Methods

The study was approved by both the Institutional Review Boards at Mayo Clinic and Olmsted Medical Center. This is a population-based retrospective cohort study designed to assess whether asthma status is associated with the severity of SPD among the Rochester residents who developed SPD between 1964 and 1983, a primarily pre-pneumococcal vaccine era. SPD cases were identified through reviewing 3,941 medical records and asthma status was ascertained by predetermined criteria for asthma (Table 1).

Study Design and Setting

The study setting and population were previously described in detail [3]. Briefly, Rochester, Minnesota is an excellent setting to conduct a retrospective study because medical care is virtually self-contained within the community and the Rochester Epidemiology Project provides information on all Rochester residents who had received medical care from one of two primary medical centers in Rochester. All diagnostic information has been indexed since 1935 using Berkson codes even before International Classification of Diseases (ICD) codes were available [5]. The incidence rate of asthma in Rochester was 238 per 100,000, which is comparable to those in other communities such as Tecumseh, Michigan (250/100,000) [6].
Study Subjects
We reported the details for identification of SPD cases previously [3]. Briefly, a total of 85 different medical index search codes (Berkson codes and ICD codes) were used to identify potential SPD cases and each potential case was then confirmed by medical records review. We reviewed medical records of all 3,941 persons and identified 174 SPD cases during the period 1964 to 1983. Case definition of SPD included IPD cases (isolation of *Streptococcus pneumoniae* from a normally sterile site such as blood or cerebrospinal fluid), and/or pneumococcal pneumonia requiring all three of the following criteria, 1) a physician diagnosis of pneumonia, 2) the isolation of pneumococcus from sputum gram-stain or culture, and 3) the documented pneumonia by chest radiograph. We defined the index date of onset of the SPD as the date of documented isolation of *S. pneumoniae*.

Ascertainment of asthma status
After all SPD cases were identified and confirmed, we used the previously collected data on asthma during the period 1964 to 1983 to ascertain the asthma status among the confirmed SPD cases (Table 1) [7]. In the original study, random samples of records were reviewed by different nurse abstractors and analyzed for inter-observer reliability and agreement rates between abstractors and a high degree of concordance was found [8]. A number of publications on asthma research used these criteria to define asthma [9–18].

The Severity of SPD
We used the number of days of inpatient hospitalization due to SPD as the primary surrogate measure for severity of SPD. A previous study showed that the number of days of inpatient hospitalization was well correlated with severity of IPD [19]. We defined the number of days of inpatient hospitalization for SPD as days spent on all inpatient floors (including intensive care units) from admission associated with subsequent diagnosis of SPD to discharge. In addition, the severity of SPD was measured using the intensive care unit (ICU) admission rate, days of ICU stay, and a total duration of outpatient and inpatient antibiotic therapy for SPD. Patients who were readmitted to the hospital had their length of stay calculated by adding the duration of the two hospitalizations together.

Data analysis
Socio-demographic and clinical characteristics of SPD cases with and without asthma were descriptively summarized (Table 2). The total days of ICU stay, hospitalization, and antibiotic therapy, respectively, were compared between the two groups (asthmatics vs. non-asthmatics) using the Wilcoxon rank sum test. Also, the percentage of patients admitted to the ICU was compared between the two groups (asthmatics vs. non-asthmatics) using the Fisher’s exact test. All calculated p-values were two-sided and p-values less than 0.05 were considered statistically significant.
Results

Study Cohort

A total of 3,941 records were reviewed and we identified 174 SPD cases. Of the 174 SPD cases, 11 cases (6.3%) were asthmatics. Socio-demographic and clinical characteristics of SPD cases with and without asthma are summarized in Table 2.

There were no significant differences in SPD severity between asthmatics and non-asthmatics (Table 3). The percentage of patients admitted to the ICU was 27.3% and 16.0% for asthmatics and non-asthmatics, respectively (p=0.40). The mean days of ICU stay were 0.9 ± 2.1 days in asthmatics (median=0) compared to 0.7 ± 2.5 in non-asthmatics (median=0, p=0.32). The mean days of hospitalization for asthmatics were 10.3 ± 8.9 days (median=7) whereas those for non-asthmatics were 9.7 ± 9.6 days (median=7, p=0.72).

Discussion

Our previous study showed an increased risk of SPD in asthmatics. However, the current study results suggest that there were no significant differences in all outcome measures for severity of SPD between asthmatics and non-asthmatics. The literature on IPD suggests that persons with more severe illness, such as meningitis, had longer duration of hospitalization compared with those who had less severe illness such as bacteremia or pneumonia [19]. Laurichesse et al. [19] reported that in adult patients with IPD, the mean duration of hospitalization was 10.6 days (range 1–56 days). They also reported that the duration of hospitalization ranged from 4.4 ± 4.7 days for patients with bacteremia without complication to 11.5 ± 9.5 days for those with sepsis with pneumonia and 18.5 ± 9.5 for those with meningitis. Therefore, our study results indicate that the duration of hospitalization for SPD in adult subjects (9.8 days with the range of 0–62 days, n=153) are similar to that (10.6 days) reported by Laurichesse et al. [19], although these results need be compared cautiously since we assessed SPD while the previous study included only IPD. At any rate, according to our data, asthma status has little or no influence on the severity of SPD, although no data is available to compare our study results on the hospital duration for SPD in asthmatics.

Why does asthma status have a different impact on the incidence and severity of SPD? We can only postulate it. The literature suggests asthmatics have Th2-predominant biological milieu [20] and have suboptimal or impaired innate and adaptive immunity against microbial agents [21–29] including pneumococci specifically [30]. Th2-cytokines (e.g., IL-4 or IL-5) have a reciprocal counter-regulatory relationship with Th1-cytokines (e.g., IFN-gamma) [31]. Thus, suboptimal immunity in asthmatics may be associated with the increased risk of SPD but Th2 cytokines may be beneficial in down-regulating Th1-cytokines or their related pro-inflammatory mediators, which may result in excessive inflammatory response leading to tissue damages or complications such as acute respiratory distress syndrome (ARDS) or severe inflammatory response syndrome (SIRS) due to immune dysregulation during SPD [32,33]. Thus, asthma status as a Th2-predominant immune condition may not necessarily result in more severe SPD. Whether this stipulation is true needs to be further studied.
The strengths of the study include population-based study design during the primarily pre-pneumococcal vaccine era, self-contained health care system with unified medical record system for research, and independent ascertainment of SPD cases and asthma status. Also, there are inherent limitations in the study due to its retrospective design. We did not measure any laboratory or clinical parameters for severity of SPD. Although the hospital duration is a commonly used indicator for severity, future study needs to include laboratory or clinical measures for severity of SPD. Our study had a small sample size so that the study findings may be subject to type-2 error. Our study subjects were predominantly Caucasian population and it is cautious to generalize our study findings to other study settings.

In conclusion, although asthma status has been reported to be associated with the increased risk of SPD, it does not influence the severity of SPD. Given a small sample size of our study, a study with a larger sample size needs to be considered to clarify the relationship between asthma and severity of SPD.

Acknowledgments

We thank Kathy Inabnit and Chun Shan as well as other staff of the Pediatric Asthma Epidemiology Unit who made this study possible. We thank John Yunginger, M.D. for his comments and support to use the database for the Rochester Asthma Epidemiology Project. This work was supported by an NIH grant (R01 AI 56133) from the National Institute of Allergy and Infectious Diseases and made possible by the Rochester Epidemiology Project (R01-AR30582) from the National Institute of Arthritis and Musculoskeletal and Skin Diseases.

Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| IPD          | Invasive Pneumococcal Disease |
| SPD          | Serious Pneumococcal Disease |
| 95% CI       | 95% Confidence Interval |
| ACIP         | Advisory Committee on Immunization Practices |
| PCV-7        | heptavalent Pneumococcal Conjugate Vaccine |
| PPV-23       | 23-valent Pneumococcal Polysaccharide Vaccine |
| ICD          | International Classification of Diseases |
| RR           | Risk Ratio |

References

1. Robinson KA, Baughman W, Rothrock G, Barrett NL, Pass M, et al. Epidemiology of invasive Streptococcus pneumoniae infections in the United States, 1995–1998: Opportunities for prevention in the conjugate vaccine era. JAMA. 2001; 285:1729–1735. [PubMed: 11277827]
2. Talbot TR, Hartert TV, Mitchel E, Halasa NB, Arbogast PG, et al. Asthma as a risk factor for invasive pneumococcal disease. N Engl J Med. 2005; 352:2082–2090. [PubMed: 15901861]
3. Juhn YJ, Kita H, Yawn BP, Boyce TG, Yoo KH, et al. Increased risk of serious pneumococcal disease in patients with asthma. J Allergy Clin Immunol. 2008; 122:719–723. [PubMed: 18790525]
4. http://www.cdc.gov/vaccines/recs/provisional/downloads/pneumo-Oct-2008-508.pdf
5. Kurland LT, Molgaard CA. The patient record in epidemiology. Sci Am. 1981; 245:54–63. [PubMed: 7027437]
6. Broder I, Higgins MW, Mathews KP, Keller JB. Epidemiology of asthma and allergic rhinitis in a total community, Tecumseh, Michigan. IV. Natural history. J Allergy Clin Immunol. 1974; 54:100–110. [PubMed: 4851886]

7. Yunginger JW, Reed CE, O’Connell EJ, Melton LJ 3rd, O’Fallon WM, et al. A community-based study of the epidemiology of asthma. Incidence rates, 1964–1983. Am Rev Respir Dis. 1992; 146:888–894. [PubMed: 1416415]

8. Beard CM, Yunginger JW, Reed CE, O’Connell EJ, Silverstein MD. Interobserver variability in medical record review: an epidemiological study of asthma. J Clin Epidemiol. 1992; 45:1013–1020. [PubMed: 1432015]

9. Silverstein MD, Reed CE, O’Connell EJ, Melton LJ 3rd, O’Fallon WM, et al. Long-term survival of a cohort of community residents with asthma. N Engl J Med. 1994; 331:1537–1541. [PubMed: 7969322]

10. Hunt LW Jr, Silverstein MD, Reed CE, O’Fallon WM, et al. Accuracy of the death certificate in a population-based study of asthmatic patients. JAMA. 1993; 269:1947–1952. [PubMed: 8464126]

11. Yunginger JW, Reed CE, O’Connell EJ, Melton LJ 3rd, O’Fallon WM, et al. A community-based study of the epidemiology of asthma. Incidence rates, 1964–1983. Am Rev Respir Dis. 1992; 146:888–894. [PubMed: 1416415]

12. Bauer BA, Reed CE, Yunginger JW, Wollan PC, Silverstein MD. Incidence and outcomes of asthma in the elderly. A population-based study in Rochester, Minnesota. Chest. 1997; 111:303–310. [PubMed: 9041973]

13. Silverstein MD, Yunginger JW, Reed CE, Petterson T, Zimmerman D, et al. Attained adult height after childhood asthma: effect of glucocorticoid therapy. J Allergy Clin Immunol. 1997; 99:466–474. [PubMed: 9111490]

14. John YJ, Sauver JS, Katusic S, Vargas D, Weaver A, et al. The influence of neighborhood environment on the incidence of childhood asthma: a multilevel approach. Soc Sci Med. 2005; 60:2453–2464. [PubMed: 15814171]

15. John YJ, Weaver A, Katusic S, Yunginger J. Mode of delivery at birth and development of asthma: a population-based cohort study. J Allergy Clin Immunol. 2005; 116:510–516. [PubMed: 16159617]

16. John YI, Kita H, Lee LA, Smith RW, Bagniewski SM, et al. Childhood asthma and human leukocyte antigen type. Tissue Antigens. 2007; 69:38–46. [PubMed: 17212706]

17. Yawn BP, Yunginger JW, Wollan PC, Reed CE, Silverstein MD, et al. Allergic rhinitis in Rochester, Minnesota residents with asthma: frequency and impact on health care charges. J Allergy Clin Immunol. 1999; 103:54–59. [PubMed: 9893185]

18. John YJ, Kita H, Lee LA, Swanson RJ, Smith R, et al. Childhood asthma and measles vaccine response. Ann Allergy Asthma Immunol. 2006; 97:469–476. [PubMed: 17069101]

19. Laurichesse H, Romaszkio JP, Nguyen LT, Souweine B, Poirier V, et al. Clinical characteristics and outcome of patients with invasive pneumococcal disease, Puy-de-Dôme, France, 1994–1998. Eur J Clin Microbiol Infect Dis. 2001; 20:299–308. [PubMed: 11453589]

20. Smart JM, Kemp AS. Increased Th1 and Th2 allergen-induced cytokine responses in children with atopic disease. Clin Exp Allergy. 2002; 32:796–802. [PubMed: 11994108]

21. Fahy JV, Corry DB, Boushey HA. Airway inflammation and remodeling in asthma. Curr Opin Pulm Med. 2000; 6:15–20. [PubMed: 10608420]

22. Wark PA, Johnston SL, Bucchieri F, Powell R, Puddicombe S, et al. Asthmatic bronchial epithelial cells have a deficient innate immune response to infection with rhinovirus. J Exp Med. 2005; 201:937–947. [PubMed: 15781584]

23. Contoli M, Message SD, Laza-Stanca V, Edwards MR, Wark PA, et al. Role of deficient type III interferon-lambda production in asthma exacerbations. Nat Med. 2006; 12:1023–1026. [PubMed: 16906156]

24. Grove DI, Burston TO, Wellby ML, Ford RM, Forbes JJ. Humoral and cellular immunity in asthma. J Allergy Clin Immunol. 1975; 55:152–163. [PubMed: 1089697]
25. Lee HJ, Kang JH, Henrichsen J, Konradsen HB, Jang SH, et al. Immunogenicity and safety of a 23-valent pneumococcal polysaccharide vaccine in healthy children and in children at increased risk of pneumococcal infection. Vaccine. 1995; 13:1533–1538. [PubMed: 8578838]

26. Lahood N, Emerson SS, Kumar P, Sorensen RU. Antibody levels and response to pneumococcal vaccine in steroid-dependent asthma. Ann Allergy. 1993; 70:289–294. [PubMed: 8466093]

27. Fischer JE, Johnson JE, Kuli-Zade RK, Johnson TR, Aung S, et al. Overexpression of interleukin-4 delays virus clearance in mice infected with respiratory syncytial virus. J Virol. 1997; 71:8672–8677. [PubMed: 9343225]

28. Kincy-Cain T, Clements JD, Bost KL. Endogenous and exogenous interleukin-12 augment the protective immune response in mice orally challenged with Salmonella dublin. Infect Immun. 1996; 64:1437–1440. [PubMed: 8606114]

29. Chaplin DD, Zindl CL, Duffy LB, Atkinson TP, Lai J. Clearance of Mycoplasma pneumoniae is impaired in mice with established allergic airway inflammation. J Allergy Clin Immunol. 2007; 119:S132.

30. Jung JA, Kita H, Dhillon R, Jacobson RM, Nahm MH, et al. Influence of asthma status on serotype-specific pneumococcal antibody levels. Postgrad Med. 2010; 122:116–124. [PubMed: 20861595]

31. Khan AQ, Shen Y, Wu ZQ, Wynn TA, Snapper CM. Endogenous pro-and anti-inflammatory cytokines differentially regulate an in vivo humoral response to Streptococcus pneumoniae. Infect Immun. 2002; 70:749–761. [PubMed: 11796608]

32. Iwasaka H, Noguchi T. Th1/Th2 balance in systemic inflammatory response syndrome (SIRS). Nihon Rinsho. 2004; 62:2237–2243. [PubMed: 15597970]

33. Muret J, Marie C, Fitting C, Payen D, Cavaillon JM. Ex vivo T-lymphocyte derived cytokine production in SIRS patients is influenced by experimental procedures. Shock. 2000; 13:169–174. [PubMed: 10718372]
Table 1

Definition of Asthma.

| Condition                                                                 |
|---------------------------------------------------------------------------|
| Patients were considered to have definite asthma if a physician had made a diagnosis of asthma and/or if each of the following three conditions were present, and they were considered to have probable asthma if only the first two conditions were present: |
| 1  History of cough, dyspnea, and/or wheezing, OR history of cough and/or dyspnea plus wheezing on examination, |
| 2  Substantial variability in symptoms from time to time or periods of weeks or more when symptoms were absent, and |
| 3  Two or more of the following:                                          |
|   • Sleep disturbance by nocturnal cough and wheeze                       |
|   • Nonsmoker (14 years or older)                                        |
|   • Nasal polyps                                                          |
|   • Blood eosinophilia higher than 300/uL                                 |
|   • Positive wheal and flare skin tests OR elevated serum IgE             |
|   • History of hay fever or infantile eczema OR cough, dyspnea, and wheezing regularly on exposure to an antigen |
|   • Pulmonary function tests showing one FEV1 or FVC less than 70% predicted and another with at least 20% improvement to an FEV1 of higher 70% predicted OR methacholine challenge test showing 20% or greater decrease in FEV1 |
|   • Favorable clinical response to bronchodilator                        |
| 4  Patients were excluded from the study if any of these conditions were present: |
|   • Pulmonary function tests that showed FEV1 to be consistently below 50% predicted or diminished diffusion capacity |
|   • Tracheobronchial foreign body at or about the incidence date          |
|   • Hypogammaglobulinemia (IgG less than 2.0 mg/mL) or other immunodeficiency disorder |
|   • Wheezing occurring only in response to anesthesia or medications      |
| 5  The following diseases excluded the patient from study if they occurred before the incidence date: |
|   • Bullous emphysema or pulmonary fibrosis on chest radiograph          |
|   • PiZZ alpha1-antitrypsin                                               |
|   • Cystic fibrosis                                                      |
| 6  Other major chest disease such as juvenile kyphoscoliosis or bronchiectasis FVC forced vital capacity; FEV1, forced expiratory volume in 1 sec. |
Table 2
Socio-demographic and Clinical Characteristics of SPD with and without asthma.

|                                | No asthma n=163 | Asthma n=11 |
|--------------------------------|-----------------|-------------|
| **Age at index date**          |                 |             |
| Mean (SD)                      | 56.9 (26.7)     | 57.6 (24.3) |
| **Gender**                     |                 |             |
| Male                           | 82 (50.3%)      | 6 (54.5%)   |
| Female                         | 81 (49.7%)      | 5 (45.5%)   |
| **Ethnicity**                  |                 |             |
| Hispanic/Latino                | 1 (0.6%)        | 0 (0%)      |
| Asian                          | 2 (1.2%)        | 0 (0%)      |
| White                          | 153 (93.9%)     | 11 (100%)   |
| Unknown                        | 7 (4.3%)        | 0 (0%)      |
| **Education**                  |                 |             |
| <High school                   | 34 (20.9%)      | 1 (9.1%)    |
| High school graduate           | 30 (18.4%)      | 2 (18.2%)   |
| Some college                   | 11 (6.7%)       | 0 (0%)      |
| College graduate               | 18 (11.0%)      | 2 (18.2%)   |
| Unknown                        | 70 (42.9%)      | 6 (54.5%)   |
| **Tobacco smoke exposure at index date** | | |
| No                             | 50 (30.7%)      | 4 (36.4%)   |
| Active                         | 51 (31.3%)      | 3 (27.3%)   |
| Passive                        | 13 (8.0%)       | 2 (18.2%)   |
| Unknown                        | 49 (30.1%)      | 2 (18.2%)   |
| **High-risk conditions* for SPD prior to the index date** | | |
| No                             | 114 (69.9%)     | 9 (81.8%)   |
| Yes                            | 49 (30.1%)      | 2 (18.2%)   |

*High-risk conditions were based on the ACIP recommended pneumococcal vaccine-eligible conditions.

Epidemiology (Sunnyvale). Author manuscript; available in PMC 2014 September 17.
### Table 3
Comparison between the Variables for SPD Severity in Asthmatics and Non-asthmatics.

| Variable                  | Asthmatics (n=11) | Non-asthmatics (n=163) | p value |
|---------------------------|-------------------|------------------------|---------|
| ICU admission, n (%)      | 3 (27.3%)         | 26 (16.0%)             | 0.40    |
| Days of ICU stay          |                   |                        |         |
| Mean ± SD                 | 0.9 ± 2.1         | 0.7 ± 2.5              | 0.32    |
| Median (IQR)              | 0 (0, 1)          | 0 (0, 0)               |         |
| Days of hospitalization   |                   |                        |         |
| Mean ± SD                 | 10.3 ± 8.9        | 9.7 ± 9.6              | 0.72    |
| Median (IQR)              | 7 (2, 18)         | 7 (3, 13)              |         |
| Days of antibiotic therapy|                   |                        |         |
| Mean ± SD                 | 13.2 ± 10.3       | 15.7 ± 13.5            | 0.61    |
| Median (IQR)              | 14 (5, 16)        | 13 (9, 17)             |         |

IQR, Interquantile range, 25th and 75th percentiles.