Since January 2020 Elsevier has created a COVID-19 resource centre with free information in English and Mandarin on the novel coronavirus COVID-19. The COVID-19 resource centre is hosted on Elsevier Connect, the company's public news and information website.

Elsevier hereby grants permission to make all its COVID-19-related research that is available on the COVID-19 resource centre - including this research content - immediately available in PubMed Central and other publicly funded repositories, such as the WHO COVID database with rights for unrestricted research re-use and analyses in any form or by any means with acknowledgement of the original source. These permissions are granted for free by Elsevier for as long as the COVID-19 resource centre remains active.
Viral infections in relation to age, atopy, and season of admission among children hospitalized for wheezing

Peter W. Heymann, MD, a Holliday T. Carper, BS,a Deborah D. Murphy, RN,a Thomas A. E. Platts-Mills, MD, PhD,b James Patrie, MS,c Anne P. McLoughlin, MD,a Elizabeth A. Erwin, MD,b Marcus S. Shaker, MD,a Martha Hellems, MD,a Jehanna Peerzada, MD,a Frederick G. Hayden, MD,a Tina K. Hatley, MD,a and Rachel Chamberlain, MDa Charlottesville, Va

Background: Viral respiratory tract infections and atopy are associated with attacks of wheezing during childhood. However, information about the relationship between viral infections and atopy among children whose attacks of wheezing lead to hospitalization is unclear.

Objective: To evaluate the prevalence of viral respiratory tract pathogens among infants and children hospitalized for wheezing and to analyze the results in relation to the patient’s age, atopic characteristics, and season of admission.

Methods: This was a case-control study of children (age 2 months to 18 years) admitted for wheezing to the University of Virginia Medical Center over a period of 12 months. Children without wheezing were enrolled as controls. Nasal secretions were evaluated for viral pathogens by using cultures, PCR tests, and antigen detection. Total IgE and specific IgE antibody to common aeroallergens was measured in serum.

Results: Seventy percent of children hospitalized for wheezing before age 3 years (n = 79) were admitted between December and March, whereas 46% of children age 3 to 18 years (n = 54) were hospitalized between September and November. Among children younger than 3 years, viral pathogens were detected in 84% (66/79) of wheezing children and 55% (42/77) of controls (P < .001). Respiratory syncytial virus was the dominant pathogen during the winter months, but rhinovirus was more common during other months. Total serum IgE levels were generally low, and values from wheezing and control subjects overlapped considerably. Among children 3 years and older, 61% (33/54) of subjects admitted for wheezing tested positive for virus (predominantly rhinovirus), compared with 21% (12/56) of controls (P < .001). The total serum IgE values among wheezing children (geometric mean, 386 IU/mL; 95% CI, 259-573) were substantially elevated compared with those of controls (geometric mean, 38 IU/mL; 95% CI, 26-56; P < .001). A significantly higher percentage of wheezing children compared with controls was sensitized to at least 1 of the inhaled allergens tested: 84% (36/43) compared with 33% (15/45; P < .001). The atopic characteristics of wheezing children who tested positive or negative for virus were similar.

Conclusions: Viral infections were the dominant risk factor for wheezing among children hospitalized before 3 years of age. By comparison, a large majority of the wheezing children age 3 to 18 years had striking atopic characteristics that may be critical as a risk factor for hospitalization and an adverse response to viral infections, especially infections caused by rhinovirus.

Key words: Wheezing, asthma, children, hospitalization, viral respiratory tract infections, rhinovirus, respiratory syncytial virus, influenza, total serum IgE, inhaled allergens

Hospital admissions for wheezing continue to be a significant health care problem for infants and children growing up in the United States and in developed countries.1,2 Attacks of wheezing treated in the hospital or emergency department are also associated with substantial health care costs.3,4 However, information regarding the major risk factors which contribute to these attacks, in particular the relationship between viral infections and allergic inflammation in the airways, remains unclear. With the development of more sensitive techniques to test for viral pathogens, especially PCR, it is now possible to examine the prevalence of viral respiratory tract infections among children treated for wheezing in greater detail.5,6 Combined with assessments for atopy, the results of these tests may provide information that can be used to improve methods for decreasing the frequency and severity of attacks that require hospital care.

In previous studies, the major viral pathogen associated with wheezing during infancy was the respiratory syncytial virus (RSV).7,8 This was especially true during the winter months, when RSV was more prevalent in the northern hemisphere. In contrast, rhinovirus has been detected more frequently in community based studies of wheezing among school-age children.9 Other population surveys have shown that sensitization to inhaled allergens as well as elevations of total serum IgE were strongly associated with asthma in children and adults.10,11 When the prevalence of viral infections and atopy were considered together in a study of asthma in a pediatric emergency department, the strongest odds for wheezing...
METHODS

Study population

This was an observational, case-control investigation involving 133 children (age 2 months to 18 years) who were hospitalized for wheezing at the University of Virginia Medical Center from April 1, 2000, through March 31, 2001. These children represented 93% of all admissions for wheezing during the enrollment period. Children with chronic lung disease or congenital heart disease and children who were immunosuppressed were excluded. The control group included 133 children with no symptoms of wheezing at the time of admission. Exclusion criteria for the controls included immunosuppression but did not include the presence of other respiratory tract symptoms or a history of previous wheezing, asthma, or allergic disorders. Goals for matching among wheezing and control patients included age and sex. The majority of matched controls were enrolled within a month of each wheezing patient. Patient demographic information was obtained from hospital charts and from questionnaires administered to parents. The study was approved by the Human Investigation Committee at the University of Virginia. Informed consent was obtained from parents, and assent was obtained from children when they were old enough (usually ≥7 years old).

Virus detection

Nasal secretions were obtained from patients as previously described. Briefly, 1 mL PBS, pH 7.4, was instilled into each nostril. For infants and toddlers, the PBS and secretions were aspirated into a mucus trap (attached to wall suction) by inserting the tip of a flexible 18F Tri-Flow suction catheter (Allegiance Healthcare Corp) into the anterior nares. For children age ≥4 years, a Yankauer, Medi-Vac, suction catheter (Allegiance Healthcare Corp) was used. After obtaining the sample, 3 mL PBS was aspirated through the catheter to rinse residual secretions into the trap. A plastic transfer pipette was used to mix secretions vigorously with wash fluid. One milliliter of this mixture was added to viral transport medium (ViraTrans; Trinity Biotec Co, Wicklow, Ireland). Each sample was transported on ice to the Clinical Virology Laboratory at the University of Virginia for virus culture.

The remaining wash fluid and secretions were frozen (−80°C) and were used to test for common viral respiratory tract pathogens by using PCR techniques and assays for viral antigen. The primers and methods used for detecting rhinovirus, enterovirus, and coronavirus by RT-PCR have previously been described. Conserved RNA sequences for influenza A and B, RSV strains A and B, and parainfluenza types 1, 2, and 3 were detected by using the Hexaplex RT-PCR Enzyme Hybridization Assay (Prodess, Waukesha, Wis). To test for adenovirus, DNA was extracted from wash samples and amplified by PCR by using primers corresponding to the hexon-coding region. This region has a high degree of homology among adenovirus serotypes. Each nasal wash was also tested for RSV and influenza A and B antigens by using membrane enzyme immunoassays (Abbott Laboratories, Chicago, Ill).

Total serum IgE and allergen-specific IgE antibody

Blood was obtained by venipuncture (or heel stick in infants) and was analyzed for total serum IgE by using the Pharmacia CAP immunoassay (Uppsala, Sweden). Sera, available from 80% of children age 3 to 18 years from both wheezing and control groups, were analyzed for allergen-specific IgE antibody to dust mite (D farinae and D pteronyssinus), cat, dog, cockroach, Alternaria, grass, oak, and ragweed allergens by using Pharmacia CAP immunoassays. Sera with ≥0.35 IU/mL IgE antibody to any of the allergens tested were considered positive for allergen sensitization.

Statistical analysis

Patient demographic data and positive tests for virus or allergen sensitization were analyzed by nonparametric methods, as were the frequencies for positive tests for virus among wheezing and control patients who were enrolled in the same season. Because only 8 of the 266 patients were enrolled more than once, each of these patient enrollments was treated as an independent event. Tests of homogeneity with respect to the frequency data were formulated as a difference of 2 group proportions in which the group proportions were assumed equal under the null hypothesis. Nonparametric exact methods were used to have a test and CI with proper coverage probability, even when the contingency cell frequencies were small. Exact 2-sided 95% CIs for the differences of proportion were constructed as described by Agresti and Min. Multivariate analyses related to predicting wheezing as a function of the patients’ atopic status and evidence for viral infection were performed by multiple logistic regression. Tests of association were evaluated on the basis of the generalized Wald χ² statistic, and 95% CI construction for the adjusted odds ratio (OR) was based on the Wald approximation. Total serum IgE data were analyzed on the logarithmic scale by 1-way ANOVA. CI construction for the ratio of geometric means (GMs) was based on the Student r distribution. StatXact 5 (Cytel, Cambridge, Mass) was used to compute nonparametric exact tests and the nonparametric exact 95% CIs. The Proc Gennmod and the Proc Mixed procedures of SAS version 8.2 (SAS Institute, Cary, NC) were used to conduct the multiple logistic regression and the ANOVA, respectively.

RESULTS

Patient characteristics

Monthly admissions for wheezing showed seasonal peaks that differed among children <3 years old compared with children age 3 to 18 years (Fig 1, A). Seventy percent (55/79) of children <3 years old were hospitalized from December through March, and 46% (25/54) of children age 3 to 18 years were admitted from September through November. There were more male subjects admitted for wheezing, and the majority of children (65%; 86/133) were from low-income families (ie, families with a payment requirement of 0% to 5%). No significant differences in income status were observed.
between the wheezing and control groups (Table I). More African American children age 3 to 18 years were enrolled in the wheezing group, and 71% (20/28) were from low-income families. Altogether, 52% (137/264) of the children (wheezing and control subjects combined) were exposed to environmental tobacco smoke (ETS) at home. The proportion of children exposed to ETS was greater for children from low-income families (62%; 99/159) than for children from families of higher income (36%; 38/105; \( P < .001 \)). The frequency of ETS exposure was similar for children in the wheezing and control groups (Table I).

Although more wheezing than control children < 3 years old were exposed to ETS from their mothers (42% and 30%, respectively), the difference was not significant \( (P = .13) \).

**Virus identification**

*Children < 3 years old.* Eighty-four percent (66/79) of wheezing children tested positive for virus compared with 54% (42/77) of controls \( (P < .001) \). RSV was frequently detected through the age of 2 years (Fig 2) and was the dominant viral pathogen detected during the winter months when admissions for this age group peaked (Fig 3). Influenza was also significantly associated with wheezing during the winter; however, rhinovirus was detected more often among wheezing children who were hospitalized between April and November (58%; 14/24) compared with controls (26%; 5/19; \( P = .04 \); Fig 4, A). Overall, 29% (23/79) of the wheezing children compared with 16% (12/77) of controls tested positive for \( >1 \) virus \( (P = .04) \). In the wheezing group, this observation was most common among infants < 6 months old and declined with age (Fig 2). Among the controls, 22% (17/77) had diagnoses related to the respiratory tract, and 76% of them tested positive for virus, compared with 48% of controls without respiratory complaints \( (P < .05) \).

*Children age 3 to 18 years.* The percentage of positive tests for virus was 68% (23/34) among wheezing children age 3 to 9 years and 50% (10/20) among children age 10 to 18 years (Fig 2). The percentage of positive tests among controls in these age groups was significantly less (26%; 9/35; \( P < .001 \); and 14%; 3/21; \( P < .05 \), respectively). Rhinovirus accounted for 77% (48/62) of all positive tests for virus among the wheezing subjects and was the only virus significantly associated with wheezing (Fig 4, B). Between September and November, when the monthly admissions for wheezing in this age group increased, the percentage of wheezing patients with positive tests for rhinovirus was 48% (12/25). This percentage did not differ significantly from the percentage of wheezing patients who tested positive for rhinovirus and were admitted from April through August (60%; 9/15; \( P = .57 \)) or from December through March (36%; 5/14; \( P = .57 \)).

No wheezing subject and only 3 controls in this age group tested positive for \( >1 \) virus. Twenty percent of the controls (11/56) were hospitalized with respiratory tract diagnoses; not all were thought to be linked to infection (eg, aspiration pneumonia). Thirty-six percent of these controls tested positive for virus, compared with 18% of controls without respiratory complaints \( (P = .12) \). Thirteen (24%) of the wheezing subjects had received the influenza vaccine in the year before enrollment. Only 3
TABLE I. Demographics and other patient characteristics

|                    | <3 y old; n = 79 | Control n = 77 | 3-18 y old; n = 54 | Control n = 56 |
|--------------------|------------------|----------------|-------------------|----------------|
| Mean age, y (range)| 1.0 (0.20-2.8)   | 0.9 (0.20-2.8) | 8.2 (3.0-18.9)    | 8.3 (3.0-18.0) |
| Gender (% male)    | 57%              | 56%            | 63%               | 59%            |
| Race (% white, % African American)§ | 53%, 42% | 66%, 30%    | 43%, 52%          | 75%, 23%       |
| Payment requirements|                  |                |                   |                |
| 100%               | 24% (19)         | 32% (25)       | 31% (17)          | 41% (23)       |
| 10% to 75%         | 9% (7)           | 8% (6)         | 7% (4)            | 9% (5)         |
| 0% to 5%           | 67% (53)         | 60% (46)       | 61% (33)          | 50% (28)       |
| ETS exposure*      | 57% (79)         | 50% (76)       | 50% (54)          | 49% (55)       |
| ETS exposure from mother† | 42% (78) | 30% (73)     | 35% (54)          | 41% (54)       |
| Family history of allergy | 73% (77) | 71% (73)    | 83% (54)*         | 56% (50)       |
| History of eczema  | 18% (79)         | 8% (75)        | 32% (53)*         | 7% (56)        |
| History of previous wheezing | 66% (79)†      | 29% (73)      | 94% (54)*         | 25% (51)       |

Significant differences between wheezing and controls groups:
*P < .01
†P < .001.
‡Two controls who were matched for age with wheezing subjects <3 years old fell into the older age group.
§Four percent of the patients enrolled (n = 12) were from other racial groups.

Payment requirements show the percentage of the hospital bill charged to the patient’s family based on their annual income. The percentage of families who were charged 100%, from 10% to 75%, or 0% to 5% of their child’s bill is shown for each patient group. In parentheses are the number of patients enrolled within each income group.

The percentage of patients exposed to ≥1 smokers living in the house who smoked ≥5 cigarettes per day. Shown in parentheses is the number of families responding to each question on the questionnaire. One wheezing (age, 18 years) and one control patient (age, 16 years) reported smoking.

#Percentage of patients whose mothers smoked ≥5 cigarettes per day.

wheezing subjects tested positive for influenza (none of them had received the influenza vaccine), and only 1 tested positive for RSV. The prevalence of positive tests for virus did not differ significantly among wheezing patients on the basis of sex, race, or income status.

Assessments of IgE and IgE antibody

Children <3 years old. Total IgE levels were low, and there was significant overlap in the intraquartile ranges for total IgE among the wheezing and control groups (Fig 5). There was also overlap among the values from children who had total IgE levels >75th percentile of the distribution (range, 35-440 IU/mL for wheezing patients and 19-304 IU/mL for controls).

Children age 3 to 18 years. Total serum IgE levels were significantly elevated among wheezing children compared with controls for children age 3 to 9 years and 10 to 18 years (Fig 5). For wheezing and control patients, respectively, the GM values and 95% CIs for total IgE within each age group were 12.4 IU/mL (9.0-17.1) and 8.1 IU/mL (5.8-11.2; P = .07) for children <3 years old; 377 IU/mL (229-619) and 43 IU/mL (26-68; P < .001) for children age 3 to 9 years; and 400 IU/mL (211-759) and 33 IU/mL (18-61; P < .001) for children age 10 to 18 years. In addition, among children who had a complete set of tests for allergen specific IgE antibody, the percentage of patients with at least 1 positive test was significantly increased for wheezing subjects (84%; 36/43) compared with controls (33%; 15/45; P < .001). The most frequent response among wheezing patients was to dust mite (D farinae and/or D pteronyssinus): 58% compared with 16% among controls (P < .001). The prevalence of positive tests to the other allergens among wheezing and control subjects, respectively, was as follows: cat, 49% and 22%, P < .01; grass, 42% and 9%, P < .001; dog, 40% and 11%, P < .01; ragweed, 30% and 4%, P < .01; oak, 30% and 11%, P < .03; cockroach, 26% and 7%, P < .02; and Alternaria, 26% and 2%, P < .01. When adjusted for sensitization to the other allergens, dust mite was the only allergen that remained significantly associated with wheezing (adjusted OR, 4.7; 95% CI, 1.3-16.1). The prevalence of allergic sensitization (a positive test for IgE antibody to at least 1 allergen) and total IgE levels were not significantly different among wheezing children on the basis of sex, race, income status, or previous hospitalizations.

Overall, the adjusted OR for wheezing among the children age 3 to 18 years was 7.6 (95% CI, 2.5-22.8) for children who tested positive for virus, 13.8 (95% CI, 4.4-43.5) for children with allergic sensitization, and 104.1 (95% CI, 16.3-663.7) for children who tested positive for both virus and IgE antibody to allergen. Total IgE levels from wheezing patients who tested positive for virus (GM, 443 IU/mL; 95% CI, 265-740) were similar to the values from wheezing patients with negative tests (GM, 354 IU/mL; 95% CI, 238-1529; P = .50). The percentages of wheezing children with sensitization to at least 1 allergen among children with positive or negative tests for virus were also similar (87% and 83%, respectively). In addition, similarities were apparent when the pattern of monthly admissions for wheezing was compared among the virus-positive and virus-negative groups: 45% (15/33) of wheezing children with positive tests and 48% (10/21)
with negative tests were hospitalized between September and November \( (P = .79; \text{ Fig 1, B}) \).

**DISCUSSION**

Previous studies of wheezing admissions during childhood have focused on either viral respiratory tract infections or atopy as risk factors for acute exacerbations. However, controversies persist regarding the relationship between viral infections and the atopic characteristics of the children who have these attacks. In our current study, viral infections were frequently detected among wheezing patients, and their prevalence declined with age. From age 3 years on, children admitted for wheezing had striking
atopic characteristics, and 40% had no evidence of virus infection, despite the comprehensive assessment for viral pathogens that included PCR tests, cultures, and antigen detection for RSV and influenza. Moreover, the wheezing children >3 years old with positive tests for virus had atopic characteristics similar to those of children who had negative tests, and both groups had similar monthly admission patterns, including an increased risk for hospitalization during the fall. Taken together, the data suggest that atopy among the children age 3 to 18 years may be critical as a risk factor for hospital admission and an adverse response to infection, especially infections caused by rhinovirus.

Among children <3 years old, the results confirmed that RSV was the major virus associated with wheezing during the winter months, when most of these children were hospitalized. Sixty percent (33/55) of patients enrolled during the months of December through March tested positive for RSV. A higher risk for recurrent wheezing during childhood has been reported for children who were hospitalized for wheezing during infancy. Although wheezing induced by RSV has been linked to more frequent episodes of wheezing as children grow older, prospective studies have not been able to confirm this association.

Asthma diagnosis and treatment
not been able to show that wheezing with RSV during infancy is associated with the development of atopy, or with positive skin tests to inhaled allergens.\textsuperscript{25,26} Consistent with observations in our emergency department,\textsuperscript{14} total serum IgE levels in the current study were generally low among both wheezing and control children who were <3 years old. Prospective studies have shown, however, that increased levels of IgE detected during infancy track with age, and we speculate that the children with values in the upper quartile range may have a higher risk for developing allergic respiratory tract symptoms as they grow older.\textsuperscript{27,28}

Exposure to ETS was common among children enrolled in this study, and the frequency of exposure was similar among the wheezing and control groups. This was also observed in our previous pediatric emergency department studies.\textsuperscript{29} Among wheezing children, however, especially children whose mothers smoke, there is substantial evidence that passive smoke exposure aggravates airway hyperresponsiveness and the frequency and persistence of symptoms.\textsuperscript{30,31} Nevertheless, it seems doubtful that fluctuations in ETS exposure at home would coincide with or significantly influence the seasonal peaks in wheezing admissions observed in the current study. Consistent with previous reports of racial and ethnic differences in asthma, more African American children age 3 to 18 years were enrolled in the wheezing group.\textsuperscript{32} Most of the African American children admitted for wheezing were also from low-income families; however, the frequency of positive tests for allergen specific IgE antibody and for rhinovirus were not significantly different among these children on the basis of race or family income status.

Among children 3 to 18 years old, positive tests for virus, especially rhinovirus, were common among the wheezing patients. Rhinovirus was detected in close to half of the wheezing subjects and was the only virus significantly associated with wheezing. This observation is consistent with a recent report showing that infections with rhinovirus, but not metapneumovirus, were significantly associated with wheezing among children hospitalized for asthma.\textsuperscript{33} A trend toward lower rates of virus detection was observed among the older children (10-18 years old) in our study. Even lower rates of infection were reported in a study of asthma exacerbations in adults, suggesting that protective immunity may develop despite the existence of approximately 100 rhinovirus serotypes.\textsuperscript{34,35} The evaluation of subjects for infections with mycoplasma and \textit{Chlamydia} remains a challenge.\textsuperscript{36} Nonetheless, recent studies of children and adults who were hospitalized for asthma detected mycoplasma or \textit{Chlamydia} in less than 5% of patients.\textsuperscript{35,37}

The atopic characteristics of the children admitted for wheezing from age 3 years on did not appear to diminish with age. IgE levels were also not significantly different among children with previous hospitalizations for wheezing compared with children who were admitted for the first time. Other surveys of children and adults have shown that asthma and airway hyperresponsiveness are closely linked to the serum IgE level, and that elevations of total IgE may be associated with asthma independent of allergen specific IgE responses.\textsuperscript{11-13,38} The contribution of IgE to wheezing episodes among children who require hospitalization compared with asthmatic children who do not need hospitalization is difficult to judge from the current study. However, increased levels of IgE have been observed in adults hospitalized for asthma whose values were significantly higher than values from adults with stable asthma.\textsuperscript{35} In addition, the odds for wheezing associated with sensitization to aeroallergens was recently shown to be greater among school-age children requiring hospital or emergency department care for their asthma (OR, 16.95) compared with children with stable asthma (OR, 2.09).\textsuperscript{39}

Collectively, these observations suggest that sensitization and exposure to aeroallergens may be a predisposing

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure5}
\caption{Total serum IgE levels. Median and intraquartile ranges are indicated by solid lines within box plots. GM values are shown next to the boxes and are also indicated by dotted lines within the boxes.}
\end{figure}
risk factor for an augmented response to acute infections with rhinovirus. In keeping with this, young adults with mild asthma and elevated levels of total serum IgE developed increased lower respiratory tract symptoms along with evidence for increased airway inflammation in response to an experimental rhinovirus challenge. In addition, the asthmatics with higher IgE levels had evidence for increased airway inflammation and reduced lung function before viral inoculation. Moreover, in our pediatric emergency department, the strongest odds for wheezing were observed among asthmatic children who were atopic and who tested positive for virus. This observation was even more striking in the current study among wheezing children who were 3 to 18 years old, and it provides further evidence that the interaction between allergen-induced and virus-induced inflammation may be amplified and synergistic rather than additive.

Almost half of the older children in this study were hospitalized during the months of September through November. This has also been observed annually in our pediatric emergency department. Rhinovirus infections, acquired when children return to school, have been thought to contribute to this peak. However, we did not detect a higher prevalence of rhinovirus during the fall months in this study. Moreover, the monthly pattern of admissions, especially peak admissions in the fall, was very similar among wheezing children who tested positive or negative for virus. During the fall, exposure to ragweed allergen and Alternaria increases, and significant elevations in dust mite allergen have been detected in homes of asthmatic subjects living in central Virginia. In the current study, sensitization to dust mite was significantly associated with wheezing even after adjusting for sensitization to other aeroallergens. In conclusion, the children hospitalized for wheezing from the age of 3 years on had striking atopic characteristics, including levels of total serum IgE that could potentially be treated with anti-IgE antibody. Combined with observations from previous studies, the results also suggest that efforts to reduce allergic airway inflammation might help children and young adults with asthma tolerate their infections with rhinovirus better.

We thank Sandra Turner-Purvis for her help in preparing this article. We also thank Dr John Steinke for his assistance in the development of primers used in the PCR analysis for adenovirus and the nursing staff working on the pediatric inpatient units at the University of Virginia for their assistance in enrolling patients.

REFERENCES

1. Centers for Disease Control and Prevention. Measuring childhood asthma prevalence before and after the 1997 redesign of the National Health Interview Survey—United States. MMWR Morb Mortal Wkly Rep 2000;49:908-11.

2. Anderson HR. Increase in hospital admissions for childhood asthma: trends in referral, severity, and readmissions from 1970 to 1985 in a health region of the United Kingdom. Thorax 1989;44:614-9.

3. Loyano P, Sullivan SD, Smith DH, Weiss KB. The economic burden of asthma in US children: estimates from the National Medical Expenditure Survey. J Allergy Clin Immunol 1999;104:957-63.

4. Gergen PJ. Understanding the economic burden of asthma. J Allergy Clin Immunol 2001;107:445-85.

5. Johnston SL, Sanderson G, Pattermore PK, Smith S, Bardin PG, Bruce CB, et al. Use of polymerase chain reaction for the diagnosis of picornavirus infection in subjects with and without respiratory symptoms. J Clin Microbiol 1993;31:111-7.

6. Arruda E, Hayden FG. Detection of human rhinovirus RNA in nasal washings by PCR. Mol Cell Probes 1993;7:373-9.

7. La Via WV, Mark MI, Stuntman HR. Respiratory syncytial virus puzzle: clinical features, pathophysiology, treatment and prevention. J Pediatr 1992;121:503-10.

8. Henderson FW, Clyde WA, Collier AM, Denny FW, Senior RJ, Sheaffer CI, et al. The etologic and epidemiologic spectrum of bronchiolitis in pediatric practice. J Pediatr 1979;95:183-90.

9. Johnston SL, Pattermore PK, Sanderson G, Smith S, Lampe F, Josephs L, et al. Community study of role viral infections in exacerbations of asthma in 9-11 year old children. BMJ 1995;310:1225-8.

10. Sears MR, Herbison GP, Holdaway MD, Hewitt CJ, Flannery EM, Silva PA. The risks of sensitivity to grass pollen, house dust mite, and cat dander in the development of childhood asthma. Clin Exp Allergy 1989; 19:419-24.

11. Poillart SM, Chapman MD, Fiocco GP, Rose G, Platts-Mills TAE. Epidemiology of acute asthma: IgE antibodies to common inhalant allergens as a risk factor for emergency room visits. J Allergy Clin Immunol 1989;83:875-82.

12. Sears MR, Burrows B, Flannery EM, Herbison GP, Hewitt CJ, Holdaway MD. Relation between airway responsiveness and serum IgE in children with asthma and in apparently normal children. N Engl J Med 1991;325:1067-71.

13. Burrows B, Martinez FD, Halonen M, Barbee RA, Cline MG. Association of asthma with serum IgE levels and skin-test reactivity to allergens. N Engl J Med 1989;320:271-7.

14. Rakes GP, Arruda E, Ingram JM, Hoover GE, Zambrano JC, Hayden FG, et al. Rhinovirus and respiratory syncytial virus in wheezing children requiring emergency care. Am J Respir Crit Care Med 1999;159:785-90.

15. Pitkaranta A, Virolainen A, Jers J, Arruda E, Hayden FE. Detection of rhinovirus, respiratory syncytial virus, and coronavirus infections in acute otitis media by reverse transcriptase polymerase chain reaction. Pediatrics 1998;102:291-5.

16. Hierholzer JC, Halonen PE, Duhlen PO, Bingham PG, McDonough MM. Detection of adenovirus in clinical specimens by polymerase chain reaction and liquid-phase hybridization quantitated by time-resolved fluorometry. J Clin Microbiol 1993;31:1886-91.

17. Wu TC, Kanayama MD, Hsuan RB, Gaskin FB, Hutchins GM. Virus-associated RNA’s (VA-1 and VA-2): an efficient target for the detection of adenovirus infections by in situ hybridization. Am J Pathol 1992;140:991-8.

18. StatsXact 5 Statistical Software for Exact Nonparametric Inference, user manual. Vol 2. Cambridge (MA): Cytel Software Corp; 2001.

19. Agresti A, Min Y. On small sample confidence intervals for parameters in discrete distributions. Biometrics 2000;57:963-71.

20. Agresti A. Categorical data analysis. New York: Wiley & Sons; 1990.

21. McCullagh P, Nelder JA. Generalized linear models. 2nd ed. New York: Chapman & Hall; 1989.

22. Neuzil KM, Mellon BG, Wright PF, Mitchel EF, Griffin MR. The effect of influenza on hospitalizations, outpatient visits, and courses of antibiotics in children. N Engl J Med 2000;342:225-31.

23. Chiou SS, Lan L, Chan KH, Wong WHS, Malik Pevus JS. Influenza-related hospitalizations among children in Hong Kong. N Engl J Med 2002;347:2097-103.

24. Morgan WJ, Martinez FD. Risk factors for developing wheezing and asthma in childhood. Pediatr Clin North Am 1992;36:1185-203.

25. Welliver RC, Duffy L. The relationship of RSV-specific immunoglobulin E antibody responses in infancy, recurrent wheezing, and pulmonary function at age 7 to 8 years. Pediatr Pulmonol 2001;33:111-7.
28. Sherrill DL, Stein R, Halonen M, Holberg CJ, Wright A, Martinez FD. Total serum IgE and its association with asthma symptoms and allergic sensitization among children. J Allergy Clin Immunol 1999;104:28-36.
29. Chang MY, Hogan AD, Rakes GP, Ingram JM, Hoover GE, Platts-Mills TAE, et al. Salivary cotinine levels in children presenting with wheezing to an emergency department. Pediatr Pulmonol 2000;29:257-63.
30. Young S, Le Souef PN, Geelhoed GC, Stick SM, Chir B, Turner KJ, et al. The influence of a family history of asthma and parental smoking on airway responsiveness in infancy. N Engl J Med 1991;324:1168-73.
31. Martinez FD, Cline M, Burrows B. Increased incidence of asthma in children of smoking mothers. Pediatrics 1992;89:21-6.
32. Lester LA, Rich SS, Blumenthal MN, Togias A, Murphy S, Malveaux F, et al. Ethnic differences in asthma and associated phenotypes: collaborative study on the genetics of asthma. J Allergy Clin Immunol 2001;108:357-62.
33. Rawlinson WD, Waliuzzaman Z, Carter IW, Belessis YC, Gilbert KM, Morton JR. Asthma exacerbations in children associated with rhinovirus but not human metapneumovirus infection. J Infect Dis 2003;187:1314-8.
34. Nicholson KG, Kant J, Ireland DC. Respiratory viruses and exacerbations of asthma in adults. BMJ 1993;307:982-6.
35. Green RM, Custovic A, Sanderson G, Hunt J, Johnston SL, Woodcock A. Synergism between allergens and viruses and risk of hospital admission with asthma: case-control study. BMJ 2002;324:763-6.
36. Isaacs D, Joshi P. Respiratory infections and asthma. Med J Aust 2002;177(suppl):S30-1.
37. Thumereille C, Deschidde A, Bouquillon C, Santos C, Sardet A, Scalbert M, et al. Role of viruses and atypical bacteria in exacerbations of asthma in hospitalized children: a prospective study in the Nord-Pas de Calais Region (France). Pediatr Pulmonol 2003;35:75-82.
38. Sumyer J, Anto JM, Castellsague J, Soriano JB, Rocca J, the Spanish Group of the European Study of Asthma. Total serum IgE is associated with asthma independently of specific IgE levels. Eur Respir J 1996;9:1880-4.
39. Ponsonby A-L, Gatenby P, Glasgow N, Mullins R, McDonald T, Hurwitz M. Which clinical subgroups within the spectrum of child asthma are attributable to atopy? Chest 2002;121:135-42.
40. Zambrano JC, Carper HT, Rakes GP, Patrie J, Murphy DD, Platts-Mills TAE, et al. Experimental rhinovirus challenges in adults with mild asthma: the response to infection in relation to IgE. J Allergy Clin Immunol 2003;111:1008-16.
41. Duff AL, Pomeranz ES, Gelber LE, Price GW, Farris H, Hayden FG, et al. Risk factors for acute wheezing in infants and children: viruses, passive smoke, and IgE antibodies to inhalant allergens. Pediatrics 1993;92:535-40.
42. Heymann PW, Zambrano JC, Rakes GP. Virus-induced wheezing in children: respiratory syncytial virus (RSV) and rhinovirus. Immunol Allergy Clin North Am 1998;18:35-47.
43. Platts-Mills TAE, Hayden ML, Chapman MD, Wilkins SR. Seasonal variation in dust mite and grass pollen allergens in dust from the houses of patients with asthma. J Allergy Clin Immunol 1987;79:781-91.