Emergency department visits and hospitalisations for asthma, COPD and respiratory tract infections: what is the role of respiratory viruses, and return to school in September, January and March?

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ABSTRACT

Background: Asthma exacerbations increase in September coinciding with children returning to school. The aim of this study was to investigate whether this occurs 1) for COPD and respiratory tract infections (RTIs); 2) after school resumes in January and March; and 3) identify which viruses may be responsible.

Methods: Emergency department (ED) visits and admissions for asthma, COPD and RTIs and the prevalence of viruses in Ontario, Canada were analysed daily between 2003 and 2013. ED visits and admissions were provided by the Canadian Institute for Health Information. Viral prevalence was obtained from the Centre for Immunisation and Respiratory Infectious Diseases.

Results: ED visits and admissions rates demonstrated a biphasic pattern. Lowest rates occurred in July and August and the highest rates in September for asthma, and after December for COPD and RTI. The increase in rates for 30 days before and after school return in September was greatest for children with asthma <15 years (2.4–2.6×). Event rates fell after school return in January for all three conditions ranging from 10–25%, and no change followed March break for asthma and COPD. Human rhinovirus was prevalent in summer with a modest relationship to asthma rates in September. The prevalence of respiratory syncytial virus, influenza A and coronavirus was associated with sustained event rates for COPD and RTIs.

Conclusions: Asthma, COPD and RTIs increase in September but do not occur after return to school in January and March. Human rhinovirus is associated with ED visits and admissions only in September.

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Introduction

September is a worrying month for patients with asthma as it has been demonstrated in multiple jurisdictions that rates of asthma exacerbations increase after return to school after the summer vacation [1–6]. In Ontario, Canada a peak in asthma exacerbations requiring hospitalisation occurs among school-aged children 2–3 weeks after school resumes [7, 8]. This is followed approximately 1 week later by smaller increases in rates of asthma exacerbations in preschool-aged children and adults [8]. The hypothesised explanations for these observations were that exacerbations during this period are due to viral respiratory pathogens that first infect schoolchildren and are then transmitted by them to younger children and adults [8].

In the context of the current COVID-19 pandemic, children exposed to viral respiratory pathogens upon returning to school in September will be a serious public health concern. Children are less seriously affected by the SARS-CoV-2 virus than adults [9–11], but can spread to vulnerable populations such as the elderly, particularly given that asymptomatic carriers spread the disease is a major concern [12–15]. Patients with asthma have generally been under-represented in the current pandemic in large case series [16–18]. The predictable peak in September due to virally induced asthma raises concerns as to other respiratory conditions such as COPD and respiratory tract infections (RTIs). These may be more important as they affect the elderly more severely than the young with higher rates of hospital admission and increased mortality [19, 20].

Whether a September epidemic for COPD and RTI occurs is unknown. Viral spread would be equally expected on return to school in January after the Christmas holiday, and in March after the spring break holiday.

The objectives of this study were to investigate the daily rates of emergency department (ED) visits and admissions for asthma, COPD and RTI in the province of Ontario, Canada between 2003 and 2013: 1) in each month of the year; 2) during the return-to-school periods in September, January and March; and 3) to examine their relation to the community prevalence of respiratory viruses in those periods. We also wished to examine whether any sequences of exacerbation peaks between age groups occurred. Such insights into prior patterns of putative transmission of respiratory viruses and disease patterns that may be related to them should inform careful monitoring of the incidence of respiratory events in primary care following opening of schools this September and the development of contingency plans if increases in rates of respiratory morbidity are detected.

Methods

Data acquisition

The Centre for Immunisation and Respiratory Infectious Diseases collects data reported to the “Respiratory Virus Detection Surveillance System” by laboratories across all provinces in Canada. The number of nasopharyngeal swabs performed for respiratory virus detection and the number positive for each virus was obtained from data reported weekly by 6 hospital laboratories and 11 public health laboratories in the province of Ontario for the period 2003–2013. The swabs are taken at either family physician offices, urgent care centres or hospital EDs. The multiplex PCR panel included influenza A and B, respiratory syncytial virus (RSV), parainfluenza and adenovirus until 2010, after which the panel was expanded to include human metapneumovirus (hMPV), human rhinovirus (hRV), and coronavirus. The number of ED visits and admissions each day with asthma (ICD-10 J45), COPD (J40–44, J47), and RTI (J00–J06, J09–J18, J20–22) recorded as a primary diagnosis between 2003 and 2013 were acquired from the Canadian Institute for Health Information. Prevalence data for asthma were obtained from Statistics Canada and The Ontario Asthma Surveillance Information System. Data for the population size of Ontario were obtained from Statistics Canada.

Virus detection

Healthcare workers in Ontario performed the nasopharyngeal swabs on patients during routine clinical care. Specimens were processed at hospital or a public health laboratory and viruses were detected by multiplex PCR. The number of tests performed and the number positive for each virus is reported by the Centre for Immunisation and Respiratory Infectious Diseases on a weekly basis.

Statistical analysis

The daily rates for ED visits and admissions for asthma, COPD, and RTI were determined by expressing the number of events each day as a proportion of the population of Ontario at that time per 100 000 (105) and then taking the mean across each month of the year to obtain the average rate in five different age groups (<5, 5–15, 15–50, 50–70, and >70 years). The analysis for COPD was conducted only for 15–50, 50–70 and >70-year age groups. To assess the effect of children returning to school, the average event rate 30 days before and after the first day back to school in September, January and after March break holidays.
was assessed. The dates for the first day of school return after these breaks between 2003 and 2013 was obtained from the Ontario Ministry of Education.

The positive response rates for each respiratory virus was calculated by dividing the number of tests positive for each virus by the total number of tests performed and was expressed as the average response rate for each month of the year by averaging across all years (11 years of data for influenza A and B, RSV, parainfluenza, and adenovirus and 3 years of data for hMPV, hRV and coronavirus). The average positive response rate for each respiratory virus in the pre- and post-school vacation periods was determined by averaging the response rate over the 30 days before and after the day that school resumed. All daily rates are described in doubling logarithmic units per 10^5 to allow easier recognition of the relative rates across age, months and condition. All analyses and graphs were performed in TIBCO Statistica (Academic Package v.13.2).

**Results**

**Overall daily ED visits and admissions**

The population of Ontario increased from 12,251,405 in 2003 to 13,606,541 in 2013. There were 649,666 ED visits for asthma, of which 68,626 (10.6%) were hospitalised; 817,141 ED visits for COPD, of which 260,665 (31.9%) were hospitalised; and 4,365,578 ED visits for RTIs, of which 321,719 (7.4%) were hospitalised. The contribution of virus, age, and sex to ED visits and admissions has been described previously [21].

**Daily rates of ED visits and admissions throughout the whole year**

**Asthma**

The mean daily number of ED visits and admissions for asthma throughout the year demonstrated a biphasic pattern in the <5, 5–15, and, to a lesser extent, the 15–50-year age categories for both ED visits and admissions (figure 1a). The nadir for ED visits and admissions in children <5 and 5–15 years was in July (ED visits: 2.28 per 10^5 and 1.35 per 10^5, admissions: 0.44 per 10^5 and 0.11 per 10^5, respectively) with a 2.4-fold increase occurring in September in both age groups. The rates in all children <15 years steadily decreased after September until January and was similar from February to June. In all months, the daily rate of ED visits was always highest in the youngest age group and declined with age.

Rates of admissions followed a similar pattern with a nadir in July and August, and increased in September in the <5 (2.9×), 5–15 (3.5×) and near-doubling increase in the 15–50-year age group (1.8×). The proportion of ED visits that resulted in admission was consistently highest in the >70-years group throughout the year at 23%, with no further increase in September. In contrast, the proportion hospitalised increased only in the <5 (23% in September versus 18–20% for the rest of year) and 5–15-year age groups (13% in September versus 7–8% for the rest of year).

**COPD**

ED visits and admissions increased with age and demonstrated a biphasic response with a nadir in August in all age groups: 15–50, 50–70 and >70 years (0.49, 1.60, 2.59 per 10^5, respectively; figure 1b). This was followed by increases in ED visits in September in all three age groups; 15–50, 50–70, >70 years (1.6×/1.3×/1.6×) and October (1.4×/1.3×/1.4×), which was sustained until March (figure 1b).

![FIGURE 1 Mean daily rates of emergency department (ED) visits and admissions per 10^5 persons by month of the year from 2003 to 2013 for a) asthma, b) COPD, and c) respiratory tract infections (RTIs). The y-axis is expressed on a log_2 scale and is eight-fold greater for RTIs. Error bars represent 95% confidence intervals.](https://doi.org/10.1183/23120541.00593-2020)
Relative to ED visits, the proportion admitted was highest in those >70 years throughout the year (55–99%) compared to 15–50 (3–5%) and 50–70 years (26–29%).

Respiratory tract infections
ED visits and admissions followed a similar monthly pattern with a nadir in August over all the age categories but with far higher rates than asthma or COPD (figure 1c). ED visits with RTIs were lowest in August in all age groups: <5, 5–15, 15–50, 50–70, >70 years (18.6, 6.3, 4.0, 2.7, 5.2 per 10^5, respectively) and increased monthly from September till December and was sustained until peaking in February. The greatest single monthly increase was in October for all five age categories (1.5×/1.2×/1.4×/1.3×, respectively).

Admissions were highest in the <5 and >70 years but with a pronounced bifascial monthly pattern in the <5 but not seen in the >70-year age group. Admissions were relatively infrequent in patients aged 5–70 years (0.1–0.5 per 10^5). Relative to ED visits, the proportion admitted was highest in the >70-year group throughout the year (33–36%), and lowest in 5–15 (2%), 15–50 (2%), and 50–70-year groups (10%) with minor monthly variations. The proportion of ED visits that led to admission in the <5-years group ranged from 10% in January and February and halved to 5% between June and October.

Daily rates for ED visits and admissions before and after school return
Asthma
The highest proportionate increase in ED visits and admissions for asthma occurred within 30 days after return to school in September in children <5 and 5–15 years (ED visits: 2.4×/2.6×; admissions: 2.8×/3.8×, respectively; figure 2). The proportionate increase in ED visits and admissions in September decreased as age group increased; 15–50, 50–70, >70 years (ED visits: 1.5×/1.3×/1.1×, admissions: 1.9×/1.1×/1.1×). After returning to school in January, ED visits declined with increasing age groups by 15–25% and admissions were either the same or declined. There were no changes in ED visits or admissions after the March break.

COPD
The highest proportionate increase in ED visits and admissions for COPD occurred within 30 days after children had returned to school in September in age groups 15–50, 50–70, >70 years (ED visits: 1.7×/1.3×/1.7×; admissions: 1.3×/1.3×/1.1×, respectively, figure 3). After children returned to school in January, ED visits and admissions declined proportionately in all three age groups by 20% and 10% respectively. There were no changes in ED visits or admissions after the March break.

Respiratory tract infections
The greatest proportionate increase in ED visits and admissions for RTIs occurred within 30 days after children returned to school in September and decreased as age group increased; <5, 5–15, 15–50, 50–70, >70 years (ED visits: 1.5×/1.4×/1.2×/1.2×/1.1×, respectively; admissions: 1.8×/1.9×/1.3×/1.2×/1.1×, respectively; figure 4). After children returned to school in January, ED visits declined in all age groups by 20% and admissions were either the same (<5) or declined (5–50: 20%, >70: 10%). After the March break,
ED visits and admissions declined in age groups <5, 5–15, 15–50 (ED visits: 30%, 20%, 10%, admissions: 40%, 20%, 10%) and was the same in the >50-year group.

**Peaks in ED visits and admission in September**

The rates of ED visits and admissions 30 days after children’s school return were assessed for all age group and conditions to evaluate the relationship between peak rates (figure 5). The peak in children <15 with

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**FIGURE 3** COPD emergency department (ED) visit and admission rates per $10^5$ persons averaged over the 30 days prior to (pre) and after (post) return to school after the summer (Sept), winter (Jan), and spring (March) vacation periods. a) Age 15–50 years; b) age 50–70 years; c) age >70 years. The y-axis is expressed on a log$_2$ scale. Error bars represent 95% confidence intervals.

**FIGURE 4** Respiratory tract infection emergency department (ED) visit and admission rates per $10^5$ persons averaged over the 30 days prior to (pre) and after (post) return to school after the summer (Sept), winter (Jan), and spring (March) vacation periods. a) Age <5 years; b) age 5–15 years; c) age 15–50 years; d) age 50–70 years; e) age >70 years. Error bars represent 95% confidence intervals.
Asthma was on day 13 (figure 5a), with a slow gradual increase in daily rates for all other age groups, with no obvious peak. The pattern was the same for COPD (figure 5b) and RTI (figure 5c); however, rates continued to increase beyond September. Of interest, there was a progressive increase in all event rates for asthma and RTI in the younger group peaking on a Sunday.

Prevalence of respiratory viruses each month of the year and relative to school return
Overall, there were 462 823 multiplex PCR tests performed and collected in the Ontario Canadian Viral Surveillance Program between 2003 and 2013 for five respiratory viruses (influenza A/B, RSV, parainfluenza, adenovirus). In addition, there were 79 511 tests performed and reported between 2010 and 2013 for three additional viruses (hRV, hMPV and coronavirus).

hRV was the most prevalent virus for 7 months of the year and started to increase in March and peaked in September/October (figure 6). RSV, influenza A were the dominant virus for 4 months from December to March, and started to increase in October and peaking in February. Parainfluenza viruses followed a similar pattern to hRV. Adenovirus did not vary throughout the year. Coronavirus, hMPV and influenza B mirrored influenza A but at lower rates.

Relative to 30 days before and after school return in September, only hRV increased, whilst the other viruses either declined to remained the same (figure 7). In January, the prevalence of RSV, influenza A, coronavirus, hMPV, influenza B increased in the community. hRV, adenovirus and parainfluenza all decreased. After March break, RSV, influenza A, coronavirus all decreased, whilst hMPV, hRV, parainfluenza increased modestly in the community.

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FIGURE 5 Mean daily rates of emergency department visits and admissions per 10^5 persons within 30 days of return to school for a) asthma, b) COPD, and c) respiratory tract infections. The y-axis is expressed on a log_2 scale and is eight-fold greater for RTIs. Error bars represent 95% confidence intervals. Dashed line indicates peak for <5 and 5–15-year age group.

FIGURE 6 Proportion of nasopharyngeal swabs positive for each virus by month. The y-axis is expressed on a log_2 scale. Error bars represent 95% confidence intervals. #: Viruses that were available only between 2010 and 2013.
Discussion

This report is of a retrospective analysis over an 11-year period of ED visits and hospital admissions occurring in the province of Ontario, Canada and their relation to the community detection of respiratory viruses. These data highlight some important observations: 1) the increase in ED visits and admissions in September is preceded by the lowest yearly rates in July and August for asthma, COPD and RTIs; 2) the increase is most pronounced in children with asthma <15 years and occurs only after return to school in September within 14 days, but event rates fall or are similar following school return in January and March for all ages and condition; 3) rates for RTIs and COPD are orders of magnitude higher than asthma throughout the year; and 4) hRV is associated with an increase in rates only in September, whilst influenza A, RSV and coronavirus is associated with sustained high rates for COPD and RTI between October and March.

Observations that seasonal and monthly variations in asthma, COPD, RTIs or detections of respiratory viruses have been previously described [6, 22–26], however, the data from this study show that September is unusual because it is the month in which the highest increases for RTI and COPD also occur, not just asthma. Furthermore, analysis across all months shows that although September represents the peak in children under 15 years old with asthma, rates continue to increase for COPD and RTIs, and remain high until March. The reasons for the rise in September are unclear. Return to school is one possible explanation; however, these data challenge this supposition, as we found no increase in rates of ED visits or admissions for any condition after return to school in January or March. This is particularly interesting because: 1) rates of community viruses such as influenza A, RSV and coronavirus are on the rise in December and January; 2) the Christmas period is associated with mixing between friends and family of multiple generations, increasing the likelihood of viral transmission; and 3) rates increase for children <15 years within the first 14 days on return to school and then begin to fall, yet rates in all other age groups and condition continue to rise well beyond September. Alternative explanations that have been highlighted include reduced adherence to medication in August [27], changes in weather [28], colder temperatures [24, 29], humidity [30] and pollution [31] and returning from holiday travel. However, this is unlikely in Ontario, as previous evidence suggests the September epidemic is independent of the degree of urbanisation and geographical location [8, 32].

Allergen exposure is commonly considered as a risk factor for asthma exacerbations [33]. The ragweed season in Ontario, Canada starts in mid-August and allergic asthma is the more common in school age children. Hence it is plausible that the combination of allergen exposure and hRV cause the increase rates in September, rather than return to school. Evidence supporting this hypothesis comes from studies showing exposure to rhinovirus after allergen exposure shows increases risk of exacerbations [34, 35] and a greater degree of bronchoconstriction [36]. Opposing this view is the fact that no increase in asthma ED visits and admissions occur between April and August, which corresponds with tree and grass pollen seasons even though the percentage positive tests for hRV is increasing. We speculate that although hRV is being detected between April and August, the virus may not be as pathogenic resulting milder disease which might result in an ED visit. This explains why the strength of the relationship with hRV in the community is not as strong as might be expected [21]. Secondly, this does not explain why rates also increase for COPD and RTIs, which are generally not mediated by allergen exposure.
The effect of changes in emotion and stress on perceptions of symptoms and visits to ED for asthma, COPD and RTI is unclear. The increase in September for all conditions is preceded by falling rates in July and August. The increase was greatest in school-aged children under 15 years, raising the possibility that the start of the summer holidays lasting at least 8 weeks is associated with a reduction in stress and anxiety, whilst the start of school in September the opposite. We would also speculate that children under 5 years may be in a daycare or nursery facility and hence show similar pattern to school-aged children. Children at home may have siblings at school with who they are interacting. Our data show that Sunday had the highest rates of for asthma and RTI for children under 15 years, suggesting that the availability and threshold of parents to take children to hospitals for assessment may also be an important contributor.

Psychological triggers have been shown to be associated with a decrease in asthma control, greater medical treatment, and ED visits [37–39]. Mood changes, attention, and perception of symptoms have all been shown to impact symptom [40–44] and poor symptom perception is associated with near-fatal asthma [45]. Experimentally evoked stress has been associated with greater bronchoconstriction [46], and increased stress during school examinations also enhances airway inflammation induced by antigen challenge [47]. This may to some extent explain the fall and rise of ED visits and admission for asthma and RTI after school ends in July and begins in September. However, why this occurs for COPD in the elderly over 50 and 70 years in this study needs further attention.

The pandemic has drastically changed human behaviour. Social distancing, reduced travel and widespread public use of personal protective equipment have become common, so it is unclear whether the same patterns observed over the previous years will be repeated. Despite these caveat, this study has some important clinical implications: 1) a potential second wave of infections in September and beyond is likely to involve other non-SARS-CoV-2 viruses and hence testing for symptomatic individuals of all ages should be performed using multiplex PCR panels that include traditional viruses; 2) the turnaround time for these multiplex PCR panels should be minimised despite the expected increase in respiratory virus testing in September to avoid strictly isolating those with confirmed non-SARS-CoV-2 viruses; and 3) provisions need to be made for the care of elderly patients with COPD and RTIs who are likely to have increased mortality from SARS-CoV-2 and already have high rates of hospital admission from usual respiratory viruses in the autumn and winter. This includes assessment of lung function and sputum cultures, which is currently extremely limited due to fear of aerosol generation.

Our study has several important limitations. First, the database recording daily rates of ED visits and admissions is not directly linked to the respiratory virus surveillance program or to individual level data, so it is difficult to assess comorbidities, medication and coinfection with multiple viral or bacterial pathogens. Furthermore, at best these are associations and not an inference of causation. Second, the number and cause of ED visits and admissions due to each condition were determined using the ICD-10 codes applied by physicians and administrators to each visit, so there is potential for misclassification, particularly of asthma in children under 5 years. The diagnosis of asthma under 5 years is very difficult to differentiate from viral-induced wheezing. Third, it is unlikely that we have not captured all infections as many milder cases will not attend ED. Fourth, we do not have a more detailed breakdown of the 15–50-year age group, as this would have been useful to differentiate those most likely to have COPD. Fifth, we do not have information about whether children under 5 years are in daycare or nursery. Six, laboratories validate their own multiplex PCR assays used for virus testing, though each laboratory undergoes audits for quality assurance that are conducted by The Institute for Quality Management in Healthcare.

Conclusions
There is an increase in ED visits and admissions in September every year in Ontario, Canada for asthma, COPD and RTI, which occurs within 30 days after return to school in September but not in January or March. The increase in September is associated with an increase in prevalence of hRV only, whilst for other months of the year, the dominant viruses are influenza A, RSV and coronavirus.

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References
1 Scheuereman O, Meyerovitch J, Marcus N, et al. The September epidemic of asthma in Israel. J Asthma 2009; 46: 652–655.
2 Sears MR, Johnston NW. Understanding the September asthma epidemic. J Allergy Clin Immunol 2007; 120: 526–529.
3 Johnston NW, Sears MR. Asthma exacerbations. 1: epidemiology. Thorax 2006; 61: 722–728.
4 Lincoln D, Morgan G, Sheppeard V, et al. Childhood asthma and return to school in Sydney, Australia. Public Health 2006; 120: 854–862.
5 Monteil MA, Juman S, Hassanally R, et al. Descriptive epidemiology of asthma in Trinidad, West Indies. J Asthma 2006; 37: 677–684.
6 Harju T, Keistinen T, Tuuponen T, et al. Seasonal variation in childhood asthma hospitalizations in Finland, 1972–1992. Eur J Pediatr 1997; 156: 436–439.
7 Johnston NW, Johnstone SL, Duncan JM, et al. The September epidemic of asthma exacerbations in children: a search for etiology. J Allergy Clin Immunol 2005; 115: 132–138.
8 Johnston NW, Johnstone SL, Norman GR, et al. The September epidemic of asthma hospitalization: school children as disease vectors. J Allergy Clin Immunol 2006; 117: 557–562.
9 Huang C, Wang Y, Li X, et al. Clinical features of patients infected with 2019 novel coronavirus in Wuhan, China. Lancet 2020; 395: 497–506.
10 Wang D, Hu B, Hu C, et al. Clinical characteristics of 138 hospitalized patients with 2019 novel coronavirus–infected pneumonia in Wuhan, China. JAMA 2020; 323: 1061–1069.
11 Li Q, Guan X, Wu P, et al. Early transmission dynamics in Wuhan, China, of novel coronavirus–infected pneumonia. N Engl J Med 2020; 382: 1199–1207.
12 CDC Covid-19 Response Team. Severe outcomes among patients with coronavirus disease 2019 (COVID-19)—United States, February 12–March 16, 2020. MMWR Morb Mortal Wkly Rep 2020; 69: 343–346.
13 Yu X, Yang R. COVID-19 transmission through asymptomatic carriers is a challenge to containment. Influenza Other Respir Viruses 2020; 14: 474–475.
14 Hu Z, Song C, Xu C, et al. Clinical characteristics of 24 asymptomatic infections with COVID-19 screened among close contacts in Nanjing, China. Sci China Life Sci 2020; 63: 706–711.
15 Bai Y, Yao L, Wei T, et al. Presumed asymptomatic carrier transmission of COVID-19. JAMA 2020; 323: 1406–1407.
16 Carli G, Cecchi L, Stebbing J, et al. Is asthma protective against COVID-19? Allergy 2020; in press [https://doi.org/10.1111/all.14426].
17 Dong X, Cao YY, Lu XX, et al. Eleven faces of coronavirus disease 2019. Allergy 2020; 75: 1699–1709.
18 Li X, Xu S, Yu M, et al. Risk factors for severity and mortality in adult COVID-19 inpatients in Wuhan. J Allergy Clin Immunol 2020; 146: 110–118.
19 Aljbotani JS, Oyelade T, Aldhabir AM, et al. Prevalence, severity and mortality associated with COPD and smoking in patients with COVID-19: a rapid systematic review and meta-analysis. PLoS ONE 2020; 15: e0233147.
20 Liu K, Chen Y, Lin R, et al. Clinical features of COVID-19 in elderly patients: A comparison with young and middle-aged patients. J Infect 2020; 80: e14–e18.
21 Satia I, Cusack R, Greene JM, et al. Prevalence and contribution of respiratory viruses in the community to rates of emergency department visits and hospitalizations with respiratory tract infections, chronic obstructive pulmonary disease and asthma. PLoS ONE 2020; 15: e0228544.
22 Weiss KB. Seasonal trends in US asthma hospitalizations and mortality. JAMA 1990; 263: 2323–2328.
23 Wise RA, Calverley PM, Carter K, et al. Seasonal variations in exacerbations and deaths in patients with COPD during the TIOSPIR trial. Int J Chron Obstruct Pulmon Dis 2018; 13: 605–616.
24 Donaldson GC, Wedzicha JA. The causes and consequences of seasonal variation in COPD exacerbations. Int J Chron Obstruct Pulmon Dis 2014; 9: 1101–1110.
25 Wilkinson TM, Aris E, Bourne S, et al. A prospective, observational cohort study of the seasonal dynamics of airway pathogens in the aetiology of exacerbations in COPD. Thorax 2017; 72: 919–927.
26 Brittain-Long R, Andersson L-M, Olofsson S, et al. Seasonal variations of 15 respiratory agents illustrated by the application of a multiplex polymerase chain reaction assay. Scand J Infect Dis 2012; 44: 9–17.
27 Julious S, Campbell M, Bianchi S, et al. Seasonality of medical contacts in school-aged children with asthma: association with school holidays. Public Health 2011; 125: 769–776.
28 Newson R, Strachan D, Archibald E, et al. Acute asthma epidemics, weather and pollen in England, 1987–1994. Eur Respir J 1998; 11: 694–701.
29 Goldstein IF. Weather patterns and asthma epidemics in New York City and New Orleans, USA. Int J Biometeorol 1980; 24: 329–339.
30 Nastos PT, Palaiotassos AG, Papadopoulos M, et al. The effect of weather variability on pediatric asthma admissions in Athens, Greece. J Asthma 2008; 45: 59–65.
31 Villeneuve PJ, Chen L, Rowe BH, et al. Outdoor air pollution and emergency department visits for asthma among children and adults: a case-crossover study in northern Alberta, Canada. Environ Health 2007; 6: 40.
32 Larsen K, Zhu J, Feldman LY, et al. The annual September peak in asthma exacerbation rates. Still a reality? Ann Am Thorac Soc 2016; 13: 231–239.
33 Rosas I, McCartney H, Payne R, et al. Analysis of the relationships between environmental factors (aeroallergens, air pollution, and weather) and asthma emergency admissions to a hospital in Mexico City. Allergy 1998; 53: 394–401.
34 Murray CS, Poletti G, Kebadze T, et al. Study of modifiable risk factors for asthma exacerbations: virus infection and allergen exposure increase the risk of asthma hospital admissions in children. Thorax 2006; 61: 376–382.
35 Murray CS, Simpson A, Custovic A. Allergens, viruses, and asthma exacerbations. Proc Am Thorac Soc 2004; 1: 99–104.
36 Lemanske R, Dick EC, Swenson CA, et al. Rhinovirus upper respiratory infection increases airway hyperreactivity and late asthmatic reactions. J Clin Invest 1989; 83: 1–10.
37 Ritz T, Steptoe A, Bobb C, et al. The asthma trigger inventory: validation of a questionnaire for perceived triggers of asthma. Psychosom Med 2006; 68: 956–965.
38 Göksel Ö, Çelik GE, Erkekol FO, et al. Triggers in adult asthma: are patients aware of triggers and doing right? Allergologia et immunopathologia 2009; 37: 122–128.
39 Peterson MG, Gaeta TJ, Birkhahn RH, et al. History of symptom triggers in patients presenting to the emergency department for asthma. J Asthma 2012; 49: 629–636.
40 Affleck G, Apter A, Tennen H, et al. Mood states associated with transitory changes in asthma symptoms and peak expiratory flow. Psychosom Med 2000; 62: 61–68.
41 Meek PM. Influence of attention and judgment on perception of breathlessness in healthy individuals and patients with chronic obstructive pulmonary disease. Nurs Res 2000; 49: 11–19.
42 Main J, Moss-Morris R, Booth R, et al. The use of reliever medication in asthma: the role of negative mood and symptom reports. J Asthma 2003; 40: 357–365.
43 von Leupoldt A, Ehnes F, Dahme B. Emotions and respiratory function in asthma: a comparison of findings in everyday life and laboratory. Br J Health Psychol 2006; 11: 185–198.
44 Peuter SD, Put C, Lemaigre V, et al. Context-evoked overperception in asthma. Psychol Health 2007; 22: 737–748.
45 Davenport PW, Cruz M, Stecenko AA, et al. Respiratory-related evoked potentials in children with life-threatening asthma. Am J Respir Crit Care Med 2000; 161: 1830–1835.
46 Ritz T, Kullowatz A, Goldman MD, et al. Airway response to emotional stimuli in asthma: the role of the cholinergic pathway. J Appl Physiol 2010; 108: 1542–1549.
47 Liu LY, Coe CL, Swenson CA, et al. School examinations enhance airway inflammation to antigen challenge. Am J Respir Crit Care Med 2002; 165: 1062–1067.