Prenatal and Passive Smoke Exposure and Incidence of Asthma and Wheeze: Systematic Review and Meta-analysis

OBJECTIVES: Exposure to passive smoke is a common and avoidable risk factor for wheeze and asthma in children. Substantial growth in the prospective cohort study evidence base provides an opportunity to generate new and more detailed estimates of the magnitude of the effect. A systematic review and meta-analysis was conducted to provide estimates of the prospective effect of smoking by parents or household members on the risk of wheeze and asthma at different stages of childhood.

METHODS: We systematically searched Medline, Embase, and conference abstracts to identify cohort studies of the incidence of asthma or wheeze in relation to exposure to prenatal or postnatal maternal, paternal, or household smoking in subjects aged up to 18 years old. Pooled odds ratios (ORs) with 95% confidence intervals (CIs) were estimated by using random effects model.

RESULTS: We identified 79 prospective studies. Exposure to pre- or postnatal passive smoke exposure was associated with a 30% to 70% increased risk of incident wheezing (strongest effect from postnatal maternal smoking on wheeze in children aged ≤2 years, OR = 1.70, 95% CI = 1.24–2.35, 4 studies) and a 21% to 85% increase in incident asthma (strongest effect from prenatal maternal smoking on asthma in children aged ≤2 years, OR = 1.85, 95% CI = 1.35–2.53, 5 studies).

CONCLUSIONS: Building upon previous findings, exposure to passive smoking increases the incidence of wheeze and asthma in children and young people by at least 20%. Preventing parental smoking is crucially important to the prevention of asthma. Pediatrics 2012;129:735–744

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KEY WORDS asthma, wheeze, passive smoking exposure, meta-analysis

ABBREVIATIONS CI—confidence interval
HR—hazard ratio
NOS—Newcastle-Ottawa Quality Assessment Scale
OR—odds ratio

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It has been recognized for some years that asthma and wheezing are more common in children passively exposed to cigarette smoke, though much of the available evidence to date has been derived from cross-sectional rather than cohort studies. In a meta-analysis published in 1998 by Strachan and Cook,1 which informed the US Surgeon General’s report of 2006,2 the authors examined the effect of passive smoking on asthma and wheeze in children and found that if either parent smoked the risk of prevalent asthma in childhood was increased by around 40%, whereas postnatal maternal smoking increased the incidence of asthma or wheezing illness before the age of 6 by ~30%, and during school-age years by 13%. However, these estimates were each based on data from 4 cohort studies each. A more recent systematic review (2007) of exposure to household smoking revealed 33% increased risk of incident asthma in childhood; this estimate was based on data from only 8 studies.3

In recent years, several more prospective studies of this association have been reported, permitting a new meta-analysis to provide more reliable estimates and more detailed estimates of the effects of exposure to prenatal maternal and to postnatal maternal, paternal, or other household members smoking on the incidence of wheeze and asthma in childhood. We have, therefore, updated the original Strachan and Cook systematic review to provide summary estimates of the effects of passive smoking and to further expand these analyses by examining the effects in prenatal maternal and postnatal maternal, or paternal or household smoking from prospective studies of the risk of wheezing and asthma in childhood in various age groups. This work was initiated as part of a larger review on the effects of passive smoking in children for the Royal College of Physicians.4

**METHODS**

**Systematic Review Methods**

We used search strategies employed in the 1997 systematic review1 to identify all prospective epidemiologic studies assessing the association between passive smoke exposure and the incidence of wheeze or asthma in childhood published between 1997 and February 2011. This included searching the Medline, Embase, Cumulative Index to Nursing and Allied Health Literature, and AMED electronic databases, published reviews, reference lists from identified publications, and abstracts from major conference proceedings (European Respiratory Society and American Thoracic Society, 2006–2009).

**Search Strategy**

We search for articles in all languages, but in common with the previous search strategy, we included only articles published in English.1 Search terms were selected under guidance from the Centre for Reviews and Dissemination4 and the Airways Group Specialist5 search terms for wheeze and asthma. All references to passive smoking were selected by the MeSH heading Tobacco smoke pollution and/or text word combinations (passive, second hand, involuntary, parent, maternal, mother, paternal, father, or household) and (smoke, tobacco, cigarettes, or cotinine) in the title, keywords, or abstract. In addition, articles were then restricted to children or adolescents (aged 0–18 years). The results from searches were combined with studies identified in the previous review.1

**Inclusion Criteria**

We included all prospective epidemiologic studies assessing the association between passive smoke exposure and incidence of asthma or wheeze in children or young people up to the age of 18 years in which participants were free of disease (asthma or wheeze) at the start of the study and passive smoke exposure was documented at a time point before the incidence of disease was determined.

**Data Extraction**

Four authors (Drs Burke and McKeever, Mr Hashim, and Ms Pine-Abeta) independently reviewed the titles, abstracts, and then full text excluding irrelevant articles after each stage. Disagreements were resolved by discussion. Articles were then data extracted independently by 2 authors (Ms Burke and Dr McKeever; Mr Hashim and Dr Chen; or Ms Pine-Abeta and Dr McKeever) from all included text. Information on study design, methods, definitions of asthma and wheeze, passive smoke exposure (source and timing of exposure), location of study, age of study population (both when passive smoke exposure was attained and when outcome was collected) and results (adjusted estimates where possible) were obtained using a previously piloted data extraction form. Included articles were also independently scored by 2 authors for methodological quality by using the Newcastle-Ottawa Quality Assessment Scale (NOS).5 This scale awards points for representativeness of the cohort, ascertainment of exposure/outcome, adjustment for confounders in the analyses and follow-up (length and completeness), and has a maximum score of 9 points. Data were then entered into a standardized database and cross-checked. To ensure that no cohort was included more than once in the same meta-analysis, each article was given a study identification number relating to the birth cohort used. If different full texts presented similar outcomes and exposure at the similar age in the same cohort, then the criteria for choosing which 1 to use in the analyses included the more robust definition of asthma/wheeze (for example, persistent wheeze was used over wheeze ever),
the younger age of outcome, and multivariate results were chosen over univariate results. If a dose-response relation was reported, then the highest level of smoking was used.

Analysis: Exposure Comparisons

For each study, for a given outcome, we sought to obtain an odds ratio (OR) and its 95% confidence interval (CI) for any of the 4 exposures: prenatal maternal smoking, maternal smoking, paternal smoking, and household smoke exposure (or equivalently either parent smokes). Because the phenotype of asthma is potentially different according to age of diagnosis, we also divided the outcomes up into the following age groups: incidence of asthma/wheeze in children aged 0 to 2 years, incidence of asthma/wheeze in children aged 3 to 4 years, and incidence of asthma/wheeze in children aged over 5 years.

Meta-analyses were performed to calculate weighted effect measures and 95% CIs across studies by using random effects models using the DerSimonian and Laird method to calculate weights. Heterogeneity was expected from the analyses due to differences in study populations and exposures. The amount of heterogeneity between the studies was assessed by using $I^2$ and if substantial levels of heterogeneity were detected ($I^2 > 50$%), subgroup analyses were conducted to investigate potential reasons based on methodological quality score (using forest plots $>6$ vs $\leq 6$) and examining differences in size of effect and heterogeneity between studies. Studies presenting data from using hazards ratios (HRs) were analyzed separately. Publication bias was examined visually by using funnel plots, where there were more than 10 studies available. The presentation of the meta-analyses adhered to the Meta-analysis of Observational Studies in Epidemiology guidelines. All analyses were performed by using Stata version 10.

RESULTS

From an initial 5074 articles identified from the literature searches, 583 abstracts were reviewed and 180 full texts initially examined (Fig 1). We excluded those which did not report data on the relation between smoking and respiratory diseases, were published in non-English language, were review articles, were not prospective cohort studies, or ascertained smoking status at the same time as disease outcome. A further potential 28 articles were identified after reference list review and 9 identified as potentially eligible articles from the previous meta-analyses.

After excluding studies that presented similar outcomes from the same cohorts, there were 70 articles (representing 71 studies) eligible for inclusion in the systematic review and meta-analyses (Fig 1, Supplemental Tables 3 [included studies] and 4[excluded studies]). Thirty-two of the 71 studies assessed asthma as an outcome (46%), 31 assessed wheeze (44%), and 8 assessed both asthma and wheeze (11%). Thirty-seven studies assessed exposure to prenatal maternal smoking (52%), 26 assessed postnatal maternal smoking (37%), 7 assessed postnatal paternal smoking (10%), and 28 assessed household smoking (39%). Twenty-three studies presented the incidence of respiratory disease in children aged $\leq 2$ years, 18 in children aged 2 to 4 years, and 32 studies in children aged 5 to 18 years. Only 1 analysis had more than 10 studies, and there was no evidence of publication bias when examining the funnel plot for prenatal maternal passive smoke and wheeze in children aged $\leq 2$ years (funnel plot not presented). The NOS scores for the 71 included studies ranged from 5 to 7 with a median of 6. Thirty-one of the 71 studies (44%) were scored as being of moderate or high level ($>6$) methodological quality. The main reasons for lower NOS scores were lack of an objective measure of smoking since the vast majority relied upon self-reported smoking (only 1 study provided objective data in the form of cotinine levels), lack of adjustment for confounding factors, and lack of study sample representativeness due to restrictions to children whose parents had an allergic disease.

Meta-analysis Findings

Wheeze as the Outcome

All except for 2 of the pooled analyses revealed that exposure to passive smoking was associated with a significant increased risk of onset of wheeze in the child (Table 1, Supplemental Figs 3–12).

Prenatal Maternal Smoking

Exposure to prenatal maternal smoking was associated with 40% increase in risk of wheeze in children aged $\leq 2$ years (OR = 1.41, 95% CI = 1.20–1.67, $I^2 = 82.5\%$, 14 studies). The high levels of heterogeneity between the studies was not attributable to quality (higher quality: OR = 1.35, 95% CI = 1.13–1.61, $I^2 = 85.3\%$, 9 studies; poorer quality: OR = 1.66, 95% CI = 0.93–2.96, $I^2 = 87.7\%$, 5 studies). A similar magnitude of effect was observed for the relation between prenatal maternal smoking and incidence of wheeze between ages 3 and 4 (OR = 1.28, 95% CI = 1.14–1.44, $I^2 = 65.5\%$, 8 studies). High levels of heterogeneity in this analysis was partly due to the study quality (higher quality studies: OR = 1.44, 95% CI = 0.92–2.25, $I^2 = 59.5\%$, 3 studies; poorer quality studies: OR = 1.35, 95% CI = 1.13–1.62, $I^2 = 72.6\%$, 5 studies). Prenatal passive smoke exposure was associated with a 52% increased risk of wheeze in children aged 5 to 18 years (OR = 1.52, 95% CI = 1.23–1.87, $I^2 = 21.1\%$, 5 studies). We could not include 1 article in the meta-analyses because it provided no data other than $P$ values; however, it
revealed that maternal smoking in pregnancy was associated significant increased risk of wheeze in the last 12 months in boys age 11 ($P = .003$) but no significant association in girls.\textsuperscript{18}

**Maternal Smoking**

Exposure to postnatal maternal smoking was associated with the strongest effects on the incidence of wheeze, effects on incidence of wheeze in children aged $\leq 2$ years ($OR = 1.70$, 95% CI = $1.24$–$2.35$, $I^2 = 0\%$, 4 studies), on incidence of wheeze in children aged 3 to 4 years ($OR = 1.65$, 95% CI = $1.20$–$2.68$, $I^2 = 48.5\%$, 4 studies), and on the incidence of wheeze in children aged 5 to 18 years ($OR = 1.18$, 95% CI = $0.99$–$1.40$, $I^2 = 1.4\%$, 3 studies).

**Paternal Smoking**

Data on exposure to paternal smoking were more limited; the only available estimate was for incidence of wheeze in children aged 3 to 4 years ($OR = 1.06$, 95% CI = $0.88$–$1.27$, $I^2 = 54.5\%$, 4 studies). The moderate levels

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**TABLE 1** Passive Smoke Exposure and Incidence of Wheeze

<table>
<thead>
<tr>
<th>Smoking Exposure</th>
<th>Age the Outcome Was Collected</th>
<th>No. of Studies</th>
<th>Pooled OR</th>
<th>95% CIs</th>
<th>$I^2$, %</th>
<th>Ref. Nos.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prenatal maternal</td>
<td>$\leq 2$</td>
<td>14</td>
<td>1.41</td>
<td>1.19–1.67</td>
<td>87.9</td>
<td>17;30;38–48</td>
</tr>
<tr>
<td>Maternal</td>
<td>$\leq 2$</td>
<td>4</td>
<td>1.70</td>
<td>1.24–2.35</td>
<td>0.0</td>
<td>17;38;46;49</td>
</tr>
<tr>
<td>Paternal</td>
<td>$\leq 2$</td>
<td>0</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Household</td>
<td>$\leq 2$</td>
<td>10</td>
<td>1.35</td>
<td>1.10–1.64</td>
<td>64.5</td>
<td>39;40;42;43;46;50–53</td>
</tr>
<tr>
<td>Prenatal maternal</td>
<td>3–4</td>
<td>8</td>
<td>1.28</td>
<td>1.14–1.44</td>
<td>65.6</td>
<td>17;28;38;42;54–57</td>
</tr>
<tr>
<td>Maternal</td>
<td>3–4</td>
<td>4</td>
<td>1.65</td>
<td>1.20–2.28</td>
<td>48.5</td>
<td>17;38;54;58</td>
</tr>
<tr>
<td>Paternal</td>
<td>3–4</td>
<td>0</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Household</td>
<td>3–4</td>
<td>4</td>
<td>1.06</td>
<td>0.88–1.27</td>
<td>54.5</td>
<td>42;55;56;59</td>
</tr>
<tr>
<td>Prenatal maternal</td>
<td>5–18</td>
<td>5</td>
<td>1.52</td>
<td>1.23–1.87</td>
<td>21.1</td>
<td>29;57;58;60;61</td>
</tr>
<tr>
<td>Maternal</td>
<td>5–18</td>
<td>3</td>
<td>1.18</td>
<td>0.99–1.40</td>
<td>1.40</td>
<td>62–64</td>
</tr>
<tr>
<td>Paternal</td>
<td>5–18</td>
<td>2</td>
<td>1.39</td>
<td>1.05–1.85</td>
<td>0.0</td>
<td>60;63</td>
</tr>
<tr>
<td>Household</td>
<td>5–18</td>
<td>5</td>
<td>1.32</td>
<td>1.12–1.56</td>
<td>1.7</td>
<td>57;63;65–67</td>
</tr>
</tbody>
</table>

**FIGURE 1**

Flowchart for identifying studies.

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of heterogeneity were partly attributable to high levels of heterogeneity and in this case were attributable to study quality (higher quality: OR = 1.20, 95% CI = 1.02–1.41, I² = 0%, 3 studies; poorer quality: OR = 0.95, 95% CI = 0.87–1.02, 1 study). Exposure to household smoking also increased the risk of incidence of wheeze in children aged 5 to 18 years (OR = 1.32, 95% CI = 1.12–1.55, I² = 0%, 5 studies).

In addition, we were unable to include 1 article in the meta-analyses because the exposure was measured as per hour exposed to passive smoke, and this study revealed that increased exposure to passive smoking was associated with an increased risk of persistent wheeze in children aged 2 years (OR per hour per week exposed = 1.14, 95% CI = 1.02–1.27). There was also 1 study that used an objective measure of passive smoke exposure and revealed that children with detectable cord blood levels were not at significantly increased risk of wheeze at age 1 or 3 years or of wheeze at age 3 years in relation to blood cotinine levels at age 1.

### Asthma as the Outcome

The effects of passive smoke exposure on the incidence of asthma tended to be weaker than those on wheeze, with exposures resulting in ~20% increased risk of asthma and most of which were statistically significant (Table 2, Supplemental Figs 13–21).

#### Prenatal Maternal Smoking

The strongest significant effect was for prenatal maternal smoking and incidence of asthma in children aged ≤2 years (OR = 1.85, 95% CI = 1.35–2.53, I² = 41.9%, 5 studies). The effect of prenatal maternal smoking became progressively weaker in relation to asthma incidence with increasing age but remained significantly associated with asthma onset between the ages of 5 and 18 years (OR = 1.23, 95% CI = 1.12–1.36, I² = 50%, 11 studies). In addition, the authors of 1 study examined the effect of prenatal smoking and incidence of asthma by using HRs, and therefore data could not be included in the meta-analyses, but the study revealed that prenatal maternal smoking was associated with a significant increased risk of incidence of asthma (HR = 1.5, 95% CI = 1.1–2.1).

#### Maternal Smoking

Exposure to postnatal maternal passive smoking was not significantly associated with incidence of asthma in children aged ≤2 years or 3 to 4 years but demonstrated a borderline significant association with incidence of asthma in children aged 5 to 18 years (OR = 1.20, 95% CI = 0.98–1.46, P = .08, I² = 65.3%, 8 studies). Subgroup analysis by quality score accounted for only a little of the heterogeneity, which appeared to be located in the higher quality studies (higher quality: OR = 1.06, 95% CI = 0.67–1.68, I² = 76.5%, 4 studies; poorer quality: OR = 1.25, 95% CI = 1.00–1.57, I² = 55.5%, 4 studies). However, studies that presented the data by using HRs revealed that exposure to maternal smoking was associated with a 21% increased risk of incidence of asthma (HR = 1.21, 95% CI = 1.01–1.45, I² = 74.7%, 5 studies, Fig 2).

Again, a high amount of heterogeneity was revealed that existed in the lower quality studies (poorer quality: HR = 1.17, 95% CI = 0.99–1.37, I² = 72.9%, 4 studies; higher quality: HR = 3.17, 95% CI = 1.42–7.04, 1 study). The age of diagnosis of incidence of asthma in these 4 studies ranged considerably from birth age 5 (birth until age 11, or age 10 or older).

#### Paternal Smoking

There were more limited data on the effect of exposure to paternal smoking with no studies with data for children aged ≤2 years and only 1 study for children aged 3 to 4 years that revealed a significant effect (OR = 1.34, 95% CI = 1.23–1.46). Paternal exposure was not associated with incidence of asthma in children aged 5 to 18 years (OR = 0.98, 95% CI = 0.71–1.36, I² = 0%, 4 studies). In 1 study, the authors reported HRs and found that having a father as a current smoker resulted in a significantly increased risk of incidence of asthma from birth up to age 11 (HR = 1.34, 95% CI = 1.24–1.46).

### Table 2 Passive Smoke Exposure and Incidence of Asthma

<table>
<thead>
<tr>
<th>Smoking Exposure</th>
<th>Age the Outcome Was Collected</th>
<th>No. of Studies</th>
<th>Pooled OR</th>
<th>95% CI</th>
<th>I², %</th>
<th>Ref. No(s.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prenatal maternal</td>
<td>≤2</td>
<td>5</td>
<td>1.85</td>
<td>1.35–2.53</td>
<td>41.9</td>
<td>31,43,46,56,68</td>
</tr>
<tr>
<td>Maternal</td>
<td>≤2</td>
<td>2</td>
<td>2.47</td>
<td>0.85–9.39</td>
<td>3.7</td>
<td>46,68</td>
</tr>
<tr>
<td>Paternal</td>
<td>≤2</td>
<td>0</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Household</td>
<td>≤2</td>
<td>3</td>
<td>1.14</td>
<td>0.94–1.38</td>
<td>1.7</td>
<td>43,46,56</td>
</tr>
<tr>
<td>Prenatal maternal</td>
<td>3–4</td>
<td>1</td>
<td>1.50</td>
<td>0.88–1.92</td>
<td>0</td>
<td>70</td>
</tr>
<tr>
<td>Maternal</td>
<td>3–4</td>
<td>4</td>
<td>1.05</td>
<td>0.88–1.25</td>
<td>0.0</td>
<td>70–73</td>
</tr>
<tr>
<td>Paternal</td>
<td>3–4</td>
<td>1</td>
<td>1.34</td>
<td>1.23–1.46</td>
<td>100</td>
<td>72</td>
</tr>
<tr>
<td>Household</td>
<td>3–4</td>
<td>5</td>
<td>1.21</td>
<td>1.00–1.47</td>
<td>72.7</td>
<td>55,74–77</td>
</tr>
<tr>
<td>Prenatal maternal</td>
<td>5–18</td>
<td>8</td>
<td>1.23</td>
<td>1.12–1.56</td>
<td>50.0</td>
<td>32–34,36,48,68,78–82</td>
</tr>
<tr>
<td>Maternal</td>
<td>5–18</td>
<td>8</td>
<td>1.20</td>
<td>0.98–1.44</td>
<td>65.3</td>
<td>34,35,62,78,81,83–85</td>
</tr>
<tr>
<td>Paternal</td>
<td>5–18</td>
<td>3</td>
<td>0.98</td>
<td>0.71–1.36</td>
<td>0.0</td>
<td>35,84,85</td>
</tr>
<tr>
<td>Household</td>
<td>5–18</td>
<td>6</td>
<td>1.30</td>
<td>1.04–1.62</td>
<td>37.7</td>
<td>36,66,67,81,86,87</td>
</tr>
</tbody>
</table>
Household Smoking

Household passive smoke exposure was not significantly associated with incidence of asthma in children aged ≤2 years (OR = 1.14, 95% CI = 0.94–1.38, I² = 0.1%, 3 studies) but was associated with incidence of asthma in children aged 3 to 4 years (OR = 1.21, 95% CI = 1.00–1.47, I² = 72.7%, 5 studies) and aged 5 to 18 years (OR = 1.30, 95% CI = 1.04–1.62, I² = 37.7%, 5 studies). The relation between household passive smoke exposure and incident asthma in children aged 3 to 4 years revealed a high level of heterogeneity that could be entirely explained by subgroup analyses based on quality score (higher quality: OR = 1.18, 95% CI = 0.92–1.53, I² = 0%, 2 studies; poorer quality: OR = 1.21, 95% CI = 0.92–1.59, I² = 85.8%, 3 studies). Two further studies presented data by using HRs, and a pooled analysis of these revealed household exposure to passive smoke was not significantly related to an increased risk of incidence of asthma (HR = 1.33, 95% CI = 0.72–2.44, I² = 74.2%, 2 studies; Supplemental Fig 22).

Finally, 1 study that could not be combined due to its classification of exposure revealed no significant association for doctor diagnosed asthma or reactive airways disease in relation per house exposed to passive smoke (OR per hour exposed = 1.04, 95% CI = 0.94–1.15)19.

There was insufficient data to combine the results from studies of dose-response associations; 3 studies for wheeze revealed a significant dose-response effect,28–30 and 3 studies for asthma31–34 revealed a significant dose-response effect; the significance could not be determined for 1 study, although the results did demonstrate increasing ORs.32 However, 2 of the above studies also demonstrate a nonsignificant trend.26,34

DISCUSSION

This updated systematic review and meta-analysis, which has drawn data from over 70 studies and is thus by far the largest review of the topic reported to date, revealed that exposure to passive smoking, in particular prenatal or postnatal maternal smoking, is associated with significantly increased risks of onset of wheeze and asthma in children. The much greater evidence base available now than at the times of previous systematic reviews of this area has permitted us to estimate effects of exposure to smoking both pre- and postnatally by the mother and by the father or any household member, on asthma or wheeze during 3 different age ranges in childhood. Our estimates update those published in the previous definitive review of these associations, which was conducted over 10 years ago and included both cross-sectional and cohort studies and are based exclusively on prospective studies in which smoking status was documented before the development of disease, thereby minimizing reporting bias and recall biases.

Our findings indicate that the effects of passive smoking on the incidence of wheeze and asthma are substantially higher than previously estimated, particularly for the effect of maternal postnatal smoking exposure. Previous results revealed that maternal postnatal smoking exposure increased the

FIGURE 2
Exposure to postnatal maternal smoking and incidence of asthma.
risk of asthma or wheezing illness in the first 5 to 7 years after birth by 31% (4 studies), and the effect was smaller when examining the incidence during schools (13% increased risk [4 studies]). The most comparable data from our study revealed that maternal smoking was associated with a 52% increased risk of wheeze and 20% increased risk of asthma at age 5 to 18 years. The other available recent systematic review revealed that maternal passive smoke exposure was associated with a 24% increased risk of incidence of asthma, and household passive smoke exposure was associated with a nonsignificant 13% increased risk of incidence of asthma in children aged 0 to 18 years. Both of these estimates are at the lower ends of our estimates, as we found that passive smoke exposure was associated with a 20% to 85% increased risk of incidence of asthma. Our estimates for the effect of passive smoke on wheeze demonstrate stronger effects ranging from 28% to 70% increased risk of incidence of wheeze. This more recent meta-analyses had more restrictive inclusion criteria, including postnatal exposure to smoking, asthma, and studies that allowed for the confounding effect of atopy/allergic studies. Only allowing studies that restrict the study population or controlled for atopy may be the reason their estimates are lower than the estimates that we attained. This current meta-analyses, 38 of 71 studies allowed for atopy/allergic disease in the analyses.

Our study was able to draw data from nearly 9 times more articles than the previous work and is thus more likely to reflect the true estimates of the effect of passive smoking on wheeze and asthma. We chose to separate the time of diagnoses into 3 different age categories (≤2 years, 3 to 4 years, and 5 to 18 years) to examine whether the effect of passive smoke was only an effect on disease diagnosed early in life or whether these effects were still present when the onset of wheeze or asthma presented later in childhood. In addition, this is the first time estimates for the effect of prenatal smoking on respiratory have been determined. We found estimates tended to be higher at a younger age of diagnosis, though we still found that exposure to household passive smoke still significantly increased the risk of new diagnosis of asthma after the age of 5 years.

One limitation to our systematic review is that in many of the studies we identified, the effect of smoking on asthma or wheeze was not the primary objective. It is therefore possible that we have missed other studies in which relevant data were collected but not evident in the title or abstract. Restriction of analysis to articles published in English is another potential limitation, though in practice this excluded only 8 studies. We also recognize that exposure to prenatal and postnatal smoking and smoking by the mother, father, or other household member are correlated and their effects are therefore difficult to disentangle. There was substantial heterogeneity in some of our analyses, which was not always attributable to study quality. Reasons for high levels of heterogeneity include wide range of cohort sizes, with some large cohorts providing some very tight range of effect size, some studies not allowing for confounding effect, also differences in the defined study population such as parents having allergic disease to be included into the study versus a random sample of population. Twelve of the cohort studies were selected on the basis of parents with allergic disease and of these 2 of them had ORs and 95% CIs that were outside the pooled estimate; however, 1 was an overestimate of the effect size and 1 was an underestimate, therefore no systematic bias was revealed within these study populations. In addition, each of these 2 studies also only contributed 3% of the overall weight to the meta-analyses and therefore overall conclusions remain the same when the studies were excluded from the meta-analyses. Another factor that must be considered is the child's own smoking status, and there were only 7 studies in which the incidence of disease was after age 12 and of these only 1 of them controlled for the child's smoking behavior. Finally, we must recognize the difficulty in diagnosing asthma in young children, and therefore we chose to present the results for both wheeze and asthma; however, we must recognize that early wheeze in children will be a combination of early transient wheezers and children with early signs of asthma.

Exposure to passive smoking is an important risk factor for the incidence of wheeze and asthma throughout childhood. In the United Kingdom over 7000 or 8% of new cases of wheeze in children under the age of 2 and over 15 000 or 14% of new cases of asthma in children over the age of 3 are due to passive smoke exposure, and internationally it is estimated that 651 000 disability adjusted life-years are results of exposure to passive smoking and asthma. These new cases of disease pose a significant public health burden that is potentially avoidable. Therefore, it is important to limit children's exposure to passive smoke both during gestation and throughout the child's life.

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WISH UPON A TWINKLING PLANET? Everyone knows there are billions and billions of stars. On a clear night, one can see evidence of their existence scattered across the night sky. What is a little less clear is how many planets exist. Until recently, most scientists thought planets relatively uncommon. As reported in The New York Times (Science: January 11, 2012), however, several new studies have shown that there may actually be more planets than stars. Scientists used a few different ways to detect the presence of planets around stars outside our solar system. NASA scientists used a space-based telescope to detect planets. This telescope can detect planets as small as Earth and planets quite close to their sun. Others examine the brightness of stars as seen by several land-based telescopes to infer the presence of planets. This method is good for detecting larger planets usually a bit farther from their sun. The surprising find is that many stars in the Milky Way have planets. Evidently, most do and very conservative estimates suggest that on average stars in the Milky Way have at least 1.6 planets. According to the article, the existence of more than 700 exoplanets, planets outside our solar system, has already been confirmed. Thousands are waiting independent confirmation. Planets have even been detected around stars thought very unlikely to have a planet. For example, planets have been found around solar systems with two stars. Somehow, the planets are not destroyed by the competing gravitational fields. Given that the Milky Way has more than 100 billion stars, the likelihood of finding a planet with similar characteristics as Earth seems much more likely. If so, scientists may have to brush up on their Greek mythology, as there will be plenty of naming opportunities.

Noted by WVR, MD
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