**Ambient air pollution and the prevalence of rhinoconjunctivitis in adolescents: A worldwide ecological analysis**

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**Abstract**

Whether exposure to outdoor air pollution increases the prevalence of rhinoconjunctivitis in children is unclear. Using data from Phase Three of the International Study of Asthma and Allergies in childhood (ISAAC) we investigated associations of rhinoconjunctivitis prevalence in adolescents with model-based estimates of ozone, and satellite-based estimates of fine (diameter <2.5µm) particulate matter (PM2.5) and nitrogen dioxide (NO2). Information on rhinoconjunctivitis (defined as self-reported nose symptoms without a cold or flu accompanied by itchy watery eyes in the past 12 months) was available on 505,400 children aged 13-14years,in 183 centres in 83 countries. Centre-level prevalence estimates were calculated and linked geographically with estimates of long-term average concentrations of NO2, ozone and PM2.5. Multi-level models were fitted adjusting for population density, climate, sex and Gross National Income. Information on parental smoking, truck traffic and cooking fuel was available for a restricted set of centres (77 in 36 countries). Between-centres within-countries, the estimated change in rhinoconjunctivitis prevalence per 100 children, was 0.171 (95% Confidence Interval; -0.013, 0.354) per 10% increase in PM2.5, 0.096 (-0.003, 0.195) per 10% increase in NO2 and -0.186 (-0.390, 0.018) per 1 ppbV increase in ozone. Between-countries, rhinoconjunctivitis prevalence was significantly negatively associated with both ozone and PM2.5. In the restricted dataset, the latter association became less negative following adjustment for parental smoking and open fires for cooking. In conclusion, there were no significant within-country associations of rhinoconjunctivitis prevalence with study pollutants. Negative between-country associations with PM2.5 and ozone require further investigation.

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**Introduction**

The International Study of Asthma and Allergies (ISAAC) is a programme of cross-sectional surveys of allergic disease in children conducted in centres across the world and based on standardised protocols (Ellwood et al. 2005, 2010). From comparisons within ISAAC study centres over a seven year period there is evidence that the prevalence and hence the health care burden associated with rhinoconjunctivitis is increasing (Asher et al. 2006). It is therefore important to try and establish what factors contribute to variations in disease prevalence at the population level. The notion of a link between allergic diseases and air pollution is well established although findings from studies are inconsistent (Wyler et al. 2000; Hajat et al. 2001; Janssen et al. 2003; Lee et al. 2003; Gehring et al. 2010; Pénard-Morand et al. 2010; Carlsten and Melén 2012; Fuertes et al. 2013; Gehring et al. 2015; Burte et al. 2018). In previous analyses of ISAAC Phase Three, positive individual-level within-centre associations of rhinoconjunctivitis were reported with markers of exposure to combustion products i.e. frequency of truck traffic (Brunekreef et al. 2009), and parental smoking (Mitchell et al. 2012). And in a previous meta-analysis of ISAAC Phase I data (restricted to 24 countries with more than one centre and using city-specific estimates of PM10 from the World Bank Global Model of ambient particles), an overall positive within country association of PM10 with centre-level rhinoconjunctivitis prevalence was observed in children aged 13-14 years (Anderson et al, 2010). The aim of our ecological study was therefore to investigate whether ambient concentrations of nitrogen dioxide (NO2), fine particulate matter of aerodynamic diameter <2.5 µm (PM2.5) and ozone, might explain the wide variation in rhinoconjunctivitis symptom prevalence observed in adolescents in the ISAAC Phase Three study centres.

**Methods**

Phase Three of ISAAC was mainly conducted between 2000 and 2003 and included surveys of children aged 6-7 and 13-14 years (Aït-Khaled et al. 2009). A priori we confined our current analyses to the surveys of children aged 13-14 years as for this age group information on the variables of interest and combinations thereof were available from a larger number of centres. The core self-completed questionnaire contained questions relating to allergic disease and was completed by 798,685 children in 233 centres in 97 countries. Based on these data, a child was considered to have rhinoconjunctivitis if they reported “a problem with sneezing or a runny or blocked nose when [they] did not have a cold or flu” in the past 12 months which was “accompanied by itchy-watery eyes”. An additional optional self-completed questionnaire, the environmental questionnaire, was used in a subset of ISAAC Phase Three centres and was completed by 358,982 children in 121 centres in 54 countries. The information obtained included markers of indoor and outdoor personal exposure to products of combustion (i.e. frequency of truck traffic in street of residence ( “Never”, “Seldom”, “Frequently throughout the day” and “Almost the whole day”); mother smokes (“Yes”, “No”); father smokes (“Yes”, “No”); usually use gas for cooking (“Yes”, ”No”); and usually cook on open fires (“Yes”, ”No”).

Information on Gross National Income (GNI) per capita for 2001 (Central Intelligence Agency 2003; World Bank 2009), population density for 2005 (Centre for International Earth Science Information Network 2005), vapour pressure, daily mean temperature, monthly precipitation (averaged over 1991-2000) (Mitchell 2004; Mitchell and Jones 2005), satellite based estimates of annual mean ground-level PM2.5 (averaged over 2001-2006) (van Donkelaar et al. 2010) and NO2 (average for 2005) (Lamsal et al. 2008), and chemical transport model-based estimates of seasonal ground-level daily 1-hour maximum ozone (maximal 3 monthly running mean of daily max hourly ozone for 2005) were obtained as described in detail elsewhere (Anderson et al. 2012). Data on pollutants and population density were available at a spatial resolution of 0.1o latitude by 0.1o longitude while information on climate was available at a resolution of 0.5o latitude by 0.5o longitude. Information from these external sources was linked geographically to each ISAAC centre via a previously identified location grid of dimensions 0.1o latitude by 0.1o longitude (Anderson et al. 2012). The identification of location grids is described in detail elsewhere (Anderson et al. 2012), but in brief the aim was to obtain grids that encapsulated the centre of population of each study area.

With respect to pollutants, estimates of ground-level PM2.5 were based on measures of aerosol optical depth from spectroradiometers on the satellite Terra (van Donkelaar et al. 2010) while estimates of ground level NO2 were based on tropospheric NO2 columns derived from the Ozone Monitoring Instrument on the satellite Aura (Lamsal et al. 2008). In both cases information on vertical pollutant profiles was provided by the GEOS-Chem chemical transport model and applied to the column value retrieved from the satellite instruments (Lamsal et al. 2008; van Donkelaar et al. 2010). Three-month running averages of daily 1 hour maximum ozone were derived from the TM5 chemical transport model and extrapolated to a finer spatial resolution using linear interpolation (Krol et al. 2005).

**Sample attrition**

As our focus was on air pollution, which may vary substantially over relatively small areas, centres that sampled children from widely dispersed schools (centre not broadly contained within 1,000 km2), were deliberately excluded from our analyses (Anderson et al. 2012). As a result, our principal analyses were based on 183 centres in 83 countries, all of which had complete centre-level information on rhinoconjunctivitis prevalence, sex, pollutants, climate, population density and GNI per capita.

Individual level data on sex, truck traffic, parental smoking and cooking fuel from the environmental questionnaire were available on 215,552 subjects in 82 (of the 183) centres in 38 countries. However for five centres, complete information on these variables was provided by less than 65% of participants. Analyses involving environmental factors were therefore based on a restricted dataset of 210,665 subjects in 77 centres in 36 countries.

**Statistical methods**

The calculations of all centre-level aggregates were based on individual-level data in the dataset under analysis. The associations between centre-level rhinoconjunctivitis prevalence and centre-level pollution concentrations are illustrated graphically in Figure 1. For consistency with our previous publication (Anderson et al. 2012) we log transformed both PM2.5 and NO2 prior to analysis Centre-level associations between prevalence and potential confounding factors were investigated using Spearman’s correlation (see Online Resource 2 Table S1). Country-level pollution, climate and population density variables were obtained by taking an unweighted average of the corresponding centre-level variables of constituent centres.

**Two-level models (183 centres in 83 countries)**

We first investigated the association between rhinoconjunctivitis prevalence and pollution at both centre (i.e. between-centres within-country) and country-levels (i.e. between-country). We used 2-level mixed-effects linear regression models (XTMIXED) in STATA (StataCorp 2007) with adjustment for centre-level and country-level sex, climate, population density, and country-level GNI per capita. Country was modelled as a random effect and the improvement in model fit from allowing the centre-level association with pollutant to vary between countries was investigated (Begg and Parides 2003; Steele 2009). Meta-analysis regression (METAREG) in STATA was used to investigate the effect of adjusting for differences in sample size between centres (Pattenden et al. 2000).

**Two-level models (210,665 individuals in 77 centres)**

Using the restricted dataset of 210,665 subjects in 77 centres, 2-level (centre, individual) mixed-effects logistic regression models were fitted to the individual-level data. All models included GNI per capita (country-level); temperature, water vapour pressure, precipitation and population density at centre-level ; and maternal smoking, paternal smoking, gas for cooking, open fires for cooking, frequent truck-traffic and sex at the individual level. Cross-level interactions between centre-level pollutants and individual-level exposures to combustion products were introduced one at a time and any improvement in model fit assessed using likelihood ratio tests. Odds ratios and 95% confidence intervals are presented with the potential effect modifier set equal to its 25th and 75th percentiles.

**Results**

The following describes the results from fitting single pollutant models having adjusted for centre- and country-level sex, population density and climate and country-level GNI per capita (model 3, Table 1). For loge(PM2.5) there was little evidence of a between-centre within-country association with the prevalence of rhinoconjunctivitis, although there was evidence of a negative association between-countries. The estimated difference in country-level prevalence per 10% higher country-level PM2.5 concentration was -0.379 (95% CI -0.600 to -0.159) per 100 children. For loge(NO2) there was some suggestion of a small but not statistically significant (p=0.057) positive centre-level (i.e. between-centres within-country) association with prevalence but no evidence of an association at country-level (i.e. between-countries). For ozone there was no evidence of an association at centre-level but a negative association at country-level. The estimated difference in country-level prevalence per 1 ppbV higher country-level ozone was -0.173 (95% CI -0.305 to -0.041) per 100 children.

When all three pollutants were included in the same model (model 5, Table 1), the negative country-level association with loge(PM2.5) persisted and increased in magnitude while the negative country-level association with ozone was reduced in magnitude and no longer statistically significant. The three pollutant model also resulted in a significant positive country-level association between rhinoconjunctivitis and loge(NO2) and a significant negative centre-level association with ozone.

In modelling the relationship between prevalence and pollution using 2-level mixed effects linear regression rather than 2-level mixed effects logistic regression we avoided the problems of over-dispersion associated with the latter but failed to adjust for differences in sample size between centres. For the full data set (i.e. 183 centres) the median sample size was 3007 (interquartile range: 2341 to 3181). However the smallest sample size was 66 and the largest 6378. We therefore re-estimated the between-centre within-country associations in Table 1, model 3, adjusting for sample size using METAREG in STATA. As a result estimates (estimated difference in prevalence per 100 children per incremental increase in pollutant) changed almost imperceptibly from 0.171 (-0.013 to 0.354) per 10% PM2.5 to 0.174 (-0.017 to 0.364), from 0.096 (-0.003 to 0.195) per 10% NO2 to 0.097 (-0.007 to 0.200) and from -0.186 (-0.390 to 0.018) per 1ppbV ozone to -0.186 (-0.397 to 0.024).

Information on markers of exposure to combustion products was available for a sub-set of centres (i.e. this restricted analysis was based on 77 centres in 36 countries rather than 183 centres in 83 countries). At the country-level rhinoconjunctivitis prevalence was strongly negatively correlated with paternal smoking (Spearman’s r = -0.35; p<0.05) and open fires for cooking (r = -0.48; p<0.01). Correlations with maternal smoking (r = 0.11), frequent truck traffic (r = 0.19) and gas for cooking (r = 0.30) were positive but non-significant.

When we additionally adjusted associations between rhinoconjunctivitis prevalence and pollution for centre and country-level maternal smoking, paternal smoking, frequent truck traffic, gas cooking and open fires for cooking (Table 2), the estimated difference in country-level rhinoconjunctivitis prevalence per 100 children per 10% higher country-level PM2.5 was reduced in absolute magnitude from -0.208 (-0.567 to 0.151) to 0.024 (-0.330 to 0.378).

Finally Table 3 investigates whether individual-level associations between rhinoconjunctivitis and exposure to combustion products are modified by centre-level pollution. For both loge(PM2.5) and ozone, odds ratios for exposure to frequent truck-traffic and paternal smoking were marginally but significantly higher in centres with high rather than low background concentrations, although for the full truck-traffic variable (see Online Resource 2 Table S2) the pattern of any effect modification was not consistent across categories. A significant cross-level interaction was also observed between gas cooking and loge(NO2), although in the absence of any association at the individual-level (Wong 2013). This interaction is difficult to interpret and may be spurious.

**Discussion**

**Main findings**

In our ecological analysis of the association between the prevalence of rhinoconjunctivitis and yearly pollution concentration adjusted for centre and country-level sex, climate and population density and country-level GNI, we found evidence of differences in centre-level and country-level associations. At centre-level, associations with both loge(PM2.5) and loge(NO2) though positive were small and not statistically significant, while at country-level there were significant negative associations of rhinoconjunctivitis prevalence with both loge(PM2.5) and ozone.

**Centre-level associations**

Evidence of a link between air pollution, particularly diesel exhaust particles, and allergic sensitisation comes from experimental, (Diaz-Sanchez D et al. 2000; Carlsten and Melén 2012) and epidemiological studies (Wyler et al. 2000; Janssen et al. 2003; Pénard-Morand et al. 2010). It is thought that air pollutant exposure may induce oxidative stress leading to inflammation and facilitating the enhanced presentation of allergens to mast cells, which in turn results in an increase in histamine release and the severity of allergy-related symptoms. (Diaz-Sanchez et al. 2000; Saxon and Diaz-Sanchez 2005; Carlsten and Melén 2012) This sort of mechanism suggests that short-term (e.g. day to day) as well as long-term (e.g. annual average) pollutant exposures may have a role to play in the pathogenesis of rhinoconjunctivitis.

Outside of the ISAAC programme, however, there are relatively few epidemiological studies of air pollution or air pollution markers considering hay fever or rhinoconjunctivitis as outcomes and the findings are inconsistent (Wyler et al. 2000; Hajat et al. 2001; Janssen et al. 2003; Lee et al. 2003; -Gehring et al. 2010; Pénard-Morand et al. 2010; Fuertes et al. 2013; Gehring 2015; Burte et al. 2018). Nevertheless, in Taiwan a large study of 312,873 middle-school children within 55 communities reported a weak negative community-level association of allergic rhinitis prevalence with annual average ozone which the authors suggested might be due to scavenging by traffic exhaust emissions and a positive community-level association with annual average NOx (oxides of nitrogen) (Lee et al. 2003). Similarly a study of over 2,000 children aged 7-12 years in 24 Dutch schools situated close to motorways, reported a positive association of hay fever ever with school-level annual average PM2.5 and positive associations of current conjunctivitis (i.e. in the past 12 months) with both school-level annual average NO2 and PM2.5 (Janssen et al. 2003). Though consistent in direction with our own centre-level findings, in our study, there were small positive but not statistically significant associations with loge(NO2) and loge(PM2.5) and a small negative and not significant association with ozone. Further between-community associations (e.g. centre-level, school-level etc.) may be very different to those at the individual or within-community level. In a large time-series study of London children a strong positive (rather than negative) association was observed between ozone (averaged over 0-3 days prior) and general practice consultations for rhinoconjunctivitis; although the focus here was on the exacerbation of symptoms rather than prevalence (Hajat et al. 2001).

**Country-level associations**

From our two and three pollutant models in Table 1 it would appear that at country-level the negative association with ozone may be explained by the negative association with loge(PM2.5). This is not surprising given the strong country-level correlation (Spearman r= 0.56; p=0.0004) between the two pollutants. However, the dominance of one association over the other could possibly result from the different uncertainties associated with each pollutant. Negative country-level associations with these two pollutants have previously been observed in ISAAC Phase Three with severe asthma prevalence (Anderson et al. 2012). Such associations may therefore be driven by some factor or factors common to allergic disease rather than specific to rhinoconjunctivitis. They are nevertheless at odds with positive individual-level associations observed in ISAAC Phase Three: between both rhinoconjunctivitis and severe asthma and exposure to paternal smoking, maternal smoking (Mitchell et al. 2012), and frequent truck traffic (Brunekreef et al. 2009); and between severe asthma and the use of open fires for cooking (Wong et al. 2013). When we adjusted our analyses for these exposures (i.e. their averages at centre and country-level), we found some evidence that the negative country-level association of loge(PM2.5) with rhinoconjunctivitis (Table 2) was explained by country-level parental smoking (particularly paternal smoking) and country-level open fires for cooking.

Mitchell et al (2012), have already shown that whereas at the individual level there is evidence in ISAAC Phase Three that paternal smoking is positively associated with symptoms of asthma and rhinoconjunctivitis in adolescents, at an ecological level in ISAAC Phase One (i.e. across centres, adjusted for country-level GNP) the relationship is negative (Mitchell and Stewart 2001). In our current study we observed strong negative correlations of rhinoconjunctivitis prevalence with both paternal smoking and open fires for cooking at country-level.

As we observed significant positive associations at the individual-level and negative relationships at the country-level, this suggests an important role for other factors, besides pollution, on country-level prevalence. Centre-level disease prevalence depends, not only on the risk associated with a given exposure at the individual-level or the proportion of the centre population exposed but also the baseline risk, that is the risk of disease due to other exposures and genetic predisposition.

While there may be further scope in linking ISAAC data with other global databases in order to try and explain differences in country-level disease prevalence, these differences may themselves suggest novel risk factors or risk factor interactions to be investigated in future epidemiological studies.

**Study limitations**

Ambient air pollution is a universal exposure, i.e. all in the population are exposed to some extent. However, the level of that exposure will vary depending on factors such as time spent indoors, type of building, proximity to roads, distance of home from school etc. Our pollution data, whether model based or satellite based, estimate outdoor background ground-level exposure at centre-level and may therefore tell us little about individual-level exposure, in particular to components which occur in the indoor environment. Nevertheless we found some evidence that in centres with higher levels of PM2.5 and ozone, previously reported positive individual-level associations between rhinoconjunctivitis and both frequent truck traffic (Brunekreef et al. 2009), and paternal smoking (Mitchell et al. 2012),were slightly more marked (Table 3).

In interpreting between-country associations we need to be careful as our measures of country-level prevalence are not based on a representative sample of 13-14 year old children from that country. However our country-level associations are of interest only as signposts to any real causes by which they may be “confounded”. Our focus is therefore on associations between centres within countries as given our method of analysis these are adjusted for country-level differences both measured and unmeasured. These centre-level associations are effectively based on data from 128 centres in 28 countries (i.e. countries with at least two centres), 16 of which provided data from at least 3 centres (i.e. New Zealand (3 centres), Spain (9), Italy(9), Portugal (4), Serbia and Montenegro(5), Iran (4), Chile(4), Brazil(19), Argentina(4), Mexico(9), China(5), India(16), Thailand(4), Kyrgyzstan(3), Lithuania(3), Syria(3)). Our findings may therefore be disproportionately influenced by associations within Brazil and India but also to a lesser extent by associations in Spain, Italy and Mexico.

**Conclusion**

In our global ecological analysis we found no evidence of centre-level associations between rhinoconjuntivitis and pollutants. The observed negative country-level associations likely reflect complex relationships involving genetic and multiple social, demographic and environmental factors rather than exposure to air pollution.

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**Compliance with Ethical Standards**

**Funding:** The ISAAC Phase III Studies were supported by many funding bodies worldwide. The analyses presented in this paper were unfunded.

**Potential Conflicts of Interest:** BKB owns shares in Royal Dutch Shell and Scottish and Southern Energy and her spouse has a deferred Shell pension. All other authors declare they have no potential conflict of interest.

**Informed consent:** Approval for the ISAAC Phase III study was obtained from local ethics committees or other appropriate bodies. An information letter was sent home via pupils to their parents/guardians in advance of the study taking place in school. In most study centres consent was passive i.e. parents or guardians were asked to contact researchers if they or the study subject did not want to take part and no such contact was taken as consent (Ellwood et al, 2010).

**Fig.1** Scatterplots illustrating the association of rhinoconjunctivitis prevalence at ages 13-14 with PM2.5 (µg/m3), NO2 (ppbV) and ozone (ppbV) based on 183 centres in 83 countries

Table 1: The association of rhinoconjunctivitis prevalence ages 13-14 years with PM2.5, NO2 and ozone

|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
| Model No. | Adjustment | Estimated change in rhinoconjunctivitis prevalence (95% CI) per 100 children per 10% increase in: | | | | Estimated change in rhinoconjunctivitis prevalence (95% CI) per 100 children per 1ppbV increase in: | |
| PM2.5 | | NO2 | | Ozone | |
| Between-centre within-country | Between-country | Between-centre within-country | Between-country | Between-centre  within-country | Between-country |
| Using data from 183 centres in 83 countries¶ | | | | | | | |
| 1 | Unadjusted | 0.136  (-0.032 to 0.305) | -0.300\*\*  (-0.505 to -0.096) | 0.018  (-0.067 to 0.104) | -0.031  (-0.133 to 0.071) | -0.242\*  (-0.441 to -0.044) | -0.156\*  (-0.278 to -0.034) |
| 2 | Sex, climate, GNI | 0.051  (-0.120 to 0.223) | -0.319\*\*  (-0.524 to -0.114) | 0.004  (-0.079 to 0.086) | 0.030  (-0.100 to 0.160) | -0.238\*  (-0.436 to -0.040) | -0.171\*  (-0.302 to -0.040) |
| 3 | Sex, climate, GNI, population density | 0.171  (-0.013 to 0.354) | -0.379\*\*\*  (-0.600 to -0.159) | 0.096  (-0.003 to 0.195) | 0.036  (-0.107 to 0.179) | -0.186  (-0.390 to 0.018) | -0.173\*  (-0.305 to -0.041) |
| 4 | Sex, climate, GNI, population density + log(NO2) [but log(PM2.5) if log(NO2) already in the model] | 0.111  (-0.092 to 0.315) | -0.556\*\*\*  (-0.809 to -0.304) | 0.067  (-0.044 to 0.178) | 0.208\*\*  (0.054 to 0.362) | -0.199  (-0.399 to 0.000) | -0.185\*\*  (-0.320 to -0.051) |
| 5 | Sex, climate, GNI, population density + the 2 other pollutants | 0.114  (-0.086 to 0.314) | -0.521\*\*\*  (-0.830 to -0.213) | 0.073  (-0.036 to 0.182) | 0.204\*\*  (0.049 to 0.359) | -0.200\*  (-0.396 to -0.005) | -0.032  (-0.189 to 0.125) |
| ¶Based on the 128 centres of the 28 countries with >=2 centres and having adjusted for sex, population density and GNI per capita as in model 3, the test for a random slope in loge(PM2.5) was non-significant (χ2=4.31 (degrees of freedom=2) p>0.05) as was the test for a random slope in loge(NO2) (χ2= 0.13 (degrees of freedom=2) p>0.05) and the test for a random slope in ozone(χ2= 0.004 (degrees of freedom=2) p>0.05). All analyses are therefore based on 183 centres in 83 countries (although only countries with >=2 centres provide any information on between-centre within-country associations)  \*p<0.05; \*\* p<0.01; \*\*\*p<0.001 | | | | | | | |

Table 2: The association of rhinoconjunctivitis prevalence and pollution in children ages 13-14 years: adjusting for exposure to combustion products.

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| Adjustment | Estimated change in rhinoconjunctivitis prevalence (95% CI) per 100 children per 10% increase in: | | | | Estimated change in rhinoconjunctivitis prevalence (95% CI) per 100 children per 1ppbV increase in: | |
| PM2.5 | | NO2 | | Ozone | |
| Between-centre within-country | Between-country | Between-centre within-country | Between-country | Between-centre  within-country | Between-country |
| Using data from 183 centres in 83 countries | | | | | | |
| Sex, climate, population density, GNI | 0.171  (-0.013 to 0.354) | -0.379\*\*\*  (-0.600 to -0.159**)** | 0.096  (-0.003 to 0.195) | 0.036  (-0.107 to 0.179) | -0.186  (-0.390 to 0.018) | -0.173\*  (-0.305 to -0.041) |
| Using data from 77 centres in 36 countries | | | | | | |
| Sex, climate, population density, GNI | 0.169  (-0.183 to 0.520) | -0.208  (-0.567 to 0.151) | 0.108  (-0.053 to 0.269) | 0.127  (-0.171 to 0.425) | -0.301\*  (-0.580 to -0.022) | -0.231\*  (-0.408 to -0.054) |
| Sex, climate, population density, GNI, paternal smoking, open fires for cooking. | 0.141  (-0.232 to 0.514) | -0.085  (-0.456 to 0.287) | 0.095  (-0.074 to 0.264) | 0.218  (-0.085 to 0.521) | -0.278  (-0.566 to 0.010) | -0.210\*  (-0.390 to -0.029) |
| Sex, climate, population density, GNI, paternal smoking, open fires for cooking, maternal smoking | 0.175  (-0.223 to 0.573) | -0.023  (-0.404 to 0.358) | 0.120  (-0.063 to 0.304) | 0.208  (-0.090 to 0.506) | -0.283  (-0.576,0.010) | -0.194\*  (-0.386 to -0.003) |
| Sex, climate, population density, GNI, paternal smoking, open fires for cooking, maternal smoking, frequent truck traffic, gas for cooking | 0.139  (-0.273 to 0.550) | 0.024  (-0.330 to 0.378) | 0.117  (-0.070 to 0.304) | 0.194  (-0.083 to 0.471) | -0.245  (-0.561 to 0.072) | -0.154  (-0.346 to 0.039) |
| \*p<0.05; \*\*p<0.01; \*\*\*p<0.001 | | | | | | |

Table 3: Investigating the effect of centre-level pollution variables on the individual-level associations between rhinoconjunctivitis and exposure to combustion products¶

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Exposure  (Yes vs No) | Potential effect modifier | | | | | | | | |
| loge(PM2.5) set equal to its: | | Test for effect modification by loge(PM2.5) | loge(NO2) set equal to its: | | Test for effect modification by loge(NO2) | Ozone set equal to its: | | Test for effect modification  by ozone |
| 25th percentile  OR (95% CI) | 75th percentile  OR (95% CI) | 25th percentile  OR (95% CI) | 75th percentile  OR (95% CI) | 25th percentile  OR (95% CI) | 75th percentile  OR (95% CI) |
| Using data from 210,665 individuals in 77 centres | | | | | | | | | |
| Mother smokes | 1.15 (1.11 to 1.20) | 1.20 (1.14 to 1.25) | P=0.093 | 1.18 (1.13 to 1.25) | 1.16 (1.11 to 1.21) | P=0.410 | 1.15  (1.10 to 1.20) | 1.20  (1.14 to 1.26) | P=0.177 |
| Father smokes | 1.09 (1.05 to 1.12) | 1.14 (1.10 to 1.18) | P=0.002 | 1.13 (1.09 to 1.17) | 1.10 (1.06 to 1.14) | P=0.224 | 1.08  (1.05 to 1.12) | 1.14  (1.10 to 1.19) | P=0.025 |
| Frequent truck traffic | 1.24 (1.20 to 1.28) | 1.30 (1.26 to 1.34) | P=0.003 | 1.25 (1.21 to 1.29) | 1.28 (1.24 to 1.32) | P=0.255 | 1.23  (1.19 to 1.27) | 1.31  (1.26 to 1.36) | P=0.005 |
| Gas for cooking | 0.98  (0.93 to 1.02) | 1.01 (0.97 to 1.06) | P=0.140 | 0.95 (0.91 to 1.00) | 1.02 (0.98 to 1.07) | P=0.018 | 0.98  (0.93 to 1.03) | 1.01  (0.96 to 1.06) | P=0.374 |
| Open fires for cooking | 1.22 (1.12 to 1.32) | 1.27 (1.17 to 1.37) | P=0.184 | 1.27 (1.17 to 1.38) | 1.21 (1.09 to 1.34) | P=0.343 | 1.25  (1.13 to 1.39) | 1.25  (1.15 to 1.37) | P=0.970 |

¶All models include GNI per capita at country-level; temperature, water vapour pressure, precipitation and population density at centre-level; maternal smoking, paternal smoking, gas for cooking, open fires for cooking, frequent truck traffic and sex at the individual-level.

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