1	Can environment or allergy explain international variation in prevalence of wheeze in
2	childhood?
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34 Abstract

Asthma prevalence in children varies substantially around the world, but the contribution of knownrisk factors to this international variation is uncertain.

The International Study of Asthma and Allergies in Childhood (ISAAC) Phase Two studied 8-12 year old children in 30 centres worldwide with parent-completed symptom and risk factor questionnaires and aeroallergen skin prick testing. We used multilevel logistic regression modelling to investigate the effect of adjustment for individual and ecological risk factors on the between-centre variation in prevalence of recent wheeze.

Adjustment for single individual-level risk factors changed the centre-level variation from a reduction of up to 8% (and 9% for atopy) to an increase of up to 6%. Modelling the 11 most influential environmental factors among all children simultaneously, the centre-level variation changed little overall (2.4% increase). Modelling only factors that decreased the variance, the 6 most influential factors (synthetic and feather quilt, mother's smoking, heating stoves, dampness and foam pillows) in combination resulted in a 21% reduction in variance. Ecological (centre-level) risk factors generally explained higher proportions of the variation than did individual risk factors.

Single environmental factors and aeroallergen sensitisation measured at the individual (child) level
 did not explain much of the between-centre variation in wheeze prevalence.

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55 Introduction

56 Asthma poses an important health burden worldwide, but its aetiology is still not fully understood, particularly in low- and middle-income countries. For instance, allergic mechanisms which have been 57 58 widely studied in high-income countries appear to be less important in less affluent settings [1]. 59 There are substantial differences in childhood asthma prevalence worldwide [2] as described in 60 Phases One and Three of the International Study of Asthma and Allergy in Childhood (ISAAC), which 61 are also apparent in ISAAC Phase Two where allergic sensitization was assessed by aeroallergen skin 62 prick testing [3]. It is not known how much of this international variation in wheeze prevalence is 63 explained by differences in allergic sensitization or other individual-level risk factors. If the currently 64 well-established risk factors fail to explain a substantial part of the international variation this would 65 indicate that important risk factors are still undiscovered. In addition to these child-level risk factors (e.g. the child is vaccinated against measles), contextual factors at the population level (e.g. the 66 67 proportion of the population that is vaccinated against measles), may also be relevant in determining prevalence. Additionally, ecological (population-level) analyses may inform about risk factors that 68 69 vary little within a given population but vary markedly in prevalence between different populations, 70 e.g. factors related to a "Western" life style. Early attempts to exploit prevalence differences to 71 understand the role of individual level risk factors in allergic disease were undertaken in Germany 72 and China, by comparing one population with a highly Westernized lifestyle (e.g. West Germany, 73 Hong Kong) to a population of the same ethnic background that was much less Westernized (e.g. 74 East-Germany, mainland China) [4,5]. The Chinese study concluded that lifestyle and environmental 75 risk factors that varied between Hong Kong and mainland China could "explain away" the prevalence 76 difference between the two populations. However, such a comparison of only two centres is 77 inherently limited in terms of generalizability.

In this paper, we extended this approach to thirty diverse study centres, including the German and Chinese centres previously studied, to quantify the extent to which known and suspected individual and contextual (ecological) risk factors may explain the observed large international variation in the prevalence of wheeze in children using data from ISAAC Phase Two.

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84 METHODS

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86 Study population and fieldwork

The methods of ISAAC Phase Two have been described in detail elsewhere [3]. Briefly, random samples of at least 10 schools from defined geographical areas were chosen and children (n>l 000 89 per centre) attending classes with a majority of 9-11-year-olds were invited to participate.
90 Standardized parental questionnaires were used. In a few centres, skin prick tests and risk factor
91 questionnaires were carried out in stratified subsamples targeting 100 children with and without
92 wheeze in the past year, respectively (details see Online Resource).

93 Thirty centres in twenty two countries participated in the questionnaire survey and 29 centres from

94 21 countries performed the standardized skin prick test.

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97 Outcome:

The main symptom of asthma used in this analysis was "wheeze in the past year". Analyses were also carried out separately for wheeze among atopic and among non-atopic children. Children were defined as atopic if the skin prick test to any of the six aeroallergens (*Dermatophagoides pteronyssinus, D. farinae,* cat dander, *Alternaria tenuis,* mixed tree pollen and mixed grass pollen) or any other locally tested allergen was positive [1]. The standardized protocol can be found online [6].

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104 **Exposures**:

105 The detailed questionnaire for environmental risk factors is available online [6] and covers 106 environmental and life style risk factors in the domains of early day exposures, diseases and 107 immunizations, the child's home (indoor air, animals and other living conditions), exercise and food. 108 The guestionnaire enguired "Have you made any changes in your home because your child had 109 asthma or allergic problems?", with subsequent specifications which were each answered separately: removed pets, stopped or reduced smoking, changed pillows, changed bedding, changed floor 110 111 covering. This helps us to address concerns about reverse causality in this cross-sectional study. We compared child-level associations (within-centre) before and after exclusion of those reporting 112 113 changes to the relevant risk factor and present these results in supplementary table E2 on the Online 114 Resource

Furthermore, we retrieved potentially relevant ecological variables from publicly available data sources (for detailed description see Online Resource). Because many potentially relevant factors are not available from such sources we additionally derived, by aggregation, centre-level covariates from the questionnaires: from the individual data on risk factors, we constructed ecological variables giving the prevalence of the individual risk factor in the centre. - for details see Online Resource. We did this for all available risk factors acknowledging that some of the resulting variables may be indicators for other centre-level risk factors.

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123 Statistical approach

124 In this analysis, we were interested in the variation of wheeze prevalence in this international 125 multicentre study that is due to true underlying variation between centres and not to the play of 126 chance (sampling error). We also sought to estimate how much of the non-sampling variation could 127 be explained by between-centre differences in the prevalence of individual or ecological factors associated with wheeze. Within the framework of a multilevel logistic regression analysis with 128 129 individuals as the first level and centres as the second level, the "true" wheeze prevalence for each 130 centre is reflected (on a logodds scale) by the random intercept for that centre, and the between-131 centre variation can be summarised by the variance of the distribution of the random intercepts 132 (tau²).

When introducing explanatory variables the variance tau² changes: introducing ecological (centrelevel) variables will always lead to a reduction because only the between-centre variance is affected. For individual-level variables, where both the individual-level and centre-level variation are affected, a change in tau² can occur in either direction [7].

137 In order to reduce the between-centre variation, an individual-level variable must be either a risk 138 factor (increasing child's risk of wheezing) and more common in centres with higher prevalence of 139 disease, or a protective factor at the individual level and inversely correlated with the prevalence of 140 disease at the centre level. There are also cases where a risk factor may be inversely correlated with 141 wheeze prevalence at the centre level, or a protective factor may be more common in centres with 142 higher prevalence of disease. In those instances, adjustment for the child-level associations in the 143 multi-level model will increase (not decrease) the between-centre variation (tau²). Thus, adjustment 144 for individual-level variables can either decrease ("explain away") or increase ("accentuate") 145 between-centre differences in disease prevalence.

146 In contrast to continuous outcomes and linear models, the variance at individual level in the logistic 147 model is determined by the binomial distribution of the dichotomous outcome and therefore, 148 models that differ in explanatory variables cannot be compared directly regarding their coefficients 149 and their tau². To allow a direct comparison, we used a scaling method [8], as described in detail 150 elsewhere [7]. Hence we compared the rescaled tau² of risk factor models to the rescaled tau² of the 151 null model without any explanatory factors.

For some risk factors in some centres, the case-control design gives artificially high intercepts because of the stratified subsample is enriched for wheezy children. This was corrected in our multilevel model by using the appropriate sampling weights for wheezy and non-wheezy children in these stratified subsamples [9] (for details see Online Resource).

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158 Construction of models with explanatory variables and determination of the change in tau²

A detailed description can be found in the Online Resource. In brief, we first tested all individual level and ecological variables in single risk factor models i.e. only one explanatory variable was introduced. The tau² of these models was compared to the tau² of the null model: the relative change (in percent) in tau² with regard to the tau² of the null model was calculated.

163 The individual risk factors to be introduced in multivariate models were chosen from the risk factors 164 that engendered the greatest change in tau².From previous work with the ISAAC data we know that 165 so far adjustment with potential confounders had very little influence on effect estimates in this 166 multicentre international context (see e.g. [10,11])

To avoid important losses in the number of children analysed in the multivariate models, we adopted a simple approach to substitute the missing values with mean values (for details see the Online Resource). We also performed analyses stratified by atopy because the relevant environmental risk factors may differ between atopic and non-atopic children reflecting atopic and non-atopic wheeze [1,12].

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All analyses were carried out using Stata releases 10 and 14 using gllamm (http://www.gllamm.org).
The rescaling of coefficients and tau² was carried out according to a do-file developed by the authors
and published elsewhere [7].

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178 RESULTS

Up to 53748 children (depending on availability of risk factor information) from 30 centres in 22 countries were included. The prevalence of wheeze in the past year ("recent wheeze") across the 30 centres ranged from 0.8% in Pichincha Province, Ecuador to 25.6% in Uruguaiana, Brazil [1]. Only 2% of the corresponding between centre variation in prevalence could be attributed to binomial sampling error (heterogeneity l^2 = 98%). When the analysis was stratified by skin prick test positivity, the prevalence of recent wheeze among atopic children ranged from 1.1% to 40.5% (l^2 = 92%) and among non-atopic children from 0.5% to 24.1% (l^2 = 98%).

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188 Single risk factor models:

189 Individual level environmental variables

Table 1 shows the 30 variables ascertained at the individual (child) level that lead to the greatest changes in tau² when included, each in turn, in the two-level model. The maximum decrease of tau² that was related to adjustment for a single individual-level variable was 8.4% for use of a synthetic quilt at present (Table 1, left side). This risk factor had a wide range of prevalence among centres from 1.9% to 87.9%, and was associated with recent wheeze within centres, with an individual-level odds ratio (OR) of 1.33 (95% confidence interval (CI): 1.09 to 1.61).

Adjustment for 14 other factors (singly) reduced the scaled tau² by more than 2% each. These pertained to bedding, smoking habits of the mother, heating of the home and dampness/mould in the living area.

199 Introducing explanatory individual-level factors into the multi-level model sometimes increased the 200 between-centre variation. The variable resulting in the strongest increase in tau² (by 6.8%) was 201 having carpets or rugs in the child's bedroom. This factor had a within-centre OR of 0.78 (95% CI: 202 0.65 to 0.94) and ranged in prevalence among centres from 6.2% to 98%. Adjustment for worm 203 infection, whooping cough infection, tuberculosis infection, no pillow use and cooking with 204 wood/coal at present, each resulted in an increase of more than 2%.

205 Changes made because of the asthma or allergy of the child partly influenced the results, depending 206 on the risk factor. Table E2 in the Online Resource shows the results for the centres that have asked 207 these questions which, depending on the risk factor, encompasses more than half up to most of the 208 affluent centres where one would expect changes to occur more often because of the frequency of 209 allergies and the higher education regarding allergic disease. The change is most marked for carpets 210 and rugs where the OR changes to one. For the other factors small to moderate changes were seen 211 which, given the precision of the estimates, are within the limits of chance. In line with this is the fact 212 that these small changes occurred in both directions when excluding children with changes e.g. an 213 increase in the OR for ETS and a decrease for the mother smoking at present.

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215 Analyses of individual level variables stratified by allergic sensitization

Among participants assessed for allergic sensitization (N=31301), a positive skin prick test was associated at the individual level with recent wheeze (OR 3.3, 95%CI: 2.8 to 4.0). Adjustment for this measure of atopy, which ranged in prevalence across centres from 1.7% to 45.3%, resulted in a 8.5% decrease of between-centre variation in wheeze prevalence.

220 When restricting the population to atopic children and non-atopic children, respectively, the pattern 221 (as shown in Table 2 which contains the same variables as Table 1) only partly corresponded to the 222 one for wheeze overall (Table 1). In general, greater changes in tau² were seen among atopic children. While adjustment for synthetic bedding resulted in a higher variance change in atopic 223 224 children compared to non-atopic children, results for feather (quilt and pillow) were inconsistent. Restricting to children where no changes in bedding occurred, yielded an increase in the OR related 225 226 to synthetic bedding for atopic children and decrease for non-atopic children. However, the changes 227 of the ORs were well within the limits of precision (i.e. the 95%-CI). Maternal smoking, especially in

228 pregnancy and at present, seemed to be more influential in non-atopic than in atopic children, in 229 terms of the effect of adjustment on between-centre variation.

Regarding infections, the changes in variance observed in all children when adjusting for worm infection seemed to be mainly driven by non-atopic children. In contrast, adjustment for whooping cough infection and tuberculosis vaccination had a stronger effect among atopic children, with increases of variance of 7.5% and 15.8%, respectively. In both subgroups many OR were imprecisely estimated so the above observations should be interpreted with caution.

235 The variables inducing the highest changes in tau² also differed between atopic and non-atopic 236 children (Table E3 in Online Resource which contains the 30 variables that lead to the highest 237 changes in atopic and non-atopic children, respectively). In atopic children, the numerically most 238 important changes were the increase in variation of 15.8% related to tuberculosis vaccination and 239 the decrease in variation of 15.6% related to synthetic bedding at present. Variables that do not 240 appear in Table 1 but were of importance among atopic children are the number of all siblings (OR 241 1.08, 95%-CI: 1.00 to 1.16; 7.8% increase in tau²), the number of older siblings (OR 1.08, 95%-CI: 1.01 242 to 1.15; 4.6% increase in tau²) and measles infection (OR 1.40, 95%-CI: 1.18 to 1.65; 4.4% increase in 243 tau²) (Table E3 in Online Resource). Among non-atopic children most of the variables with the 244 strongest changes in tau² related to indoor air quality (smoking, heating and dampness), but also 245 included bedding and whooping cough.

Overall, adjustment for the most influential risk factors tended to lead predominantly to a decrease in variance of prevalence among non-atopic children.; in atopic children, however, adjustment for the most influential risk factors more often resulted in an increase in variance.

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250 <u>Centre/country level variables</u>

251 Ecological variables introduced into the model generally resulted in markedly higher decreases in 252 tau² than those seen for individual level variables (Table 1 and Online Resource Tables E4 and E5). 253 The highest decrease of almost 50% was caused by the centre-level prevalence of contact with a dog 254 in the first year of life (Online Resource Table E4). Of the factors in Table 1, the highest decrease was 255 related to centre-level prevalences of maternal smoking, heating with wood, use of synthetic quilt 256 and tuberculosis vaccination. Ecological factors that appeared to influence notably the wheeze prevalence variation, but which were not strongly associated with wheeze at the individual level, 257 258 were contact to animals, bedroom sharing and cooked green vegetables (Online Resource Table E4). 259 In comparison with individual level variables, the effect estimates for the centre-level average 260 exposures were imprecise due to the limited number of centres (contrasting with the large number 261 of children for estimation of within-centre associations with individual risk factors). Given this 262 limitation, which also applies to the variables from open access data sources, we chose to put our

263 emphasis on an in depth analysis of the individual-level variables and to not pursue the analysis of264 the ecological variables with multivariate model.

Among the ecological variables obtained from open access data sources, the strongest reduction in variation of wheeze prevalence was linked to the country-level variables: the proportion of the population living in urban areas (32% reduction) and other indicators of affluence such as migration, and annual urban population growth (Table E4 in Online Resource). The most important centre-level variables were related to temperature variability (inverse association with wheeze) and coastal location (positive association with wheeze).

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272 Multivariate models

273 Given the uncertainty regarding estimation of the effect of centre-level variables, we only introduced 274 individual-level variables into the multivariate model. In the model incorporating only variables that resulted in a decrease in between centre variation by 1.5% and more, we obtained a 21% reduction 275 276 of the between centre prevalence variance tau² (Table 3). When risk factors that caused an increase 277 in tau² in the univariate models were also introduced, these factors counteracted the influence of 278 factors decreasing tau² and the resulting overall change was an increase by 2.4%. The resulting 279 changes in predicted prevalence (converting centre-specific random intercepts from logodds to 280 prevalence) are shown in Figure 1 (for details of the calculation of the predicted prevalence, see 281 Online Resource).

In the corresponding models for atopic children, we obtained a decrease of tau² by 16.8% and an
 increase of 11.6%, respectively. In non-atopic children, both models yielded a decrease in tau², by
 27% and 16%, respectively.

Among the 30703 children in the multivariate model from 29 centres on whom skin prick tests were performed, adjustment for individual environmental factors that decreased tau^2 reduced tau^2 by 23%. Adjusting further for atopy (as measured by skin prick test positivity) increased the reduction in tau² to 31%. The corresponding results when all environmental factors (that increased or decreased tau²) were included were a 9.3% increase before adjustment for atopy and a 0.4% reduction after this adjustment. These tau^2 differentials are broadly consistent with the effect of adjustment for atopy as a single risk factor (8.5% reduction, see above).

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294 DISCUSSION

To our knowledge this is the first analysis investigating the influence of risk factors on the international variation of disease symptoms. Single environmental factors and aeroallergen sensitisation (atopy) measured at the individual (child) level each explained less than 10% of the between-centre variation in wheeze prevalence", and adjustment for some environmental factors accentuated the variation in prevalence. When all the most influential child-level variables were modelled together, the variation in prevalence was little changed (2.4% increase without including atopy, 0.4% decrease if atopy was included).

302 So far attempts to unravel the influence of risk factors on prevalence differences have been limited 303 to the comparison of two locations [4,5,13]. Those studies investigated smaller prevalence 304 differences of 3.7 vs 1.2% [13], 5.8 vs 3.4% [5] and 27 vs 17% [4]. In the Ethiopian study [13], 305 adjusting for housing and mattress material did reduce the magnitude of the OR between rural and 306 urban location for wheeze and sensitization to house dust mite. In the Chinese study [5], the factors 307 that reduced the difference between mainland China and Hong Kong most were foam pillow, cooking with gas, damp housing and raw vegetables. The generalizability of such two centre comparisons is 308 309 uncertain but in our study we could improve this by analysing 30 diverse study centres.

310 In such a multi-centre study, the variation in disease prevalence between the centres reflects three 311 components: sampling variation (the play of chance when recruiting individual children); true (non-312 sampling) variation (between children and between centres) which can be explained by measured 313 risk factors or protective factors which themselves vary in prevalence among the study centres; and 314 true variation between centres which is not (yet) explained. We investigated the changes in this third 315 component (unexplained variation between centres) as different combinations of risk factors or 316 protective factors were included in a multi-level logistic regression model. In such a model, the 317 centre-specific prevalences are reflected by a set of intercepts (log-odds) and the parameter tau² 318 measures the variance of these centre-specific intercepts.

The individual level environmental and life style factors that caused the highest and most consistent changes in tau² were factors related to bedding material, indoor air quality, mostly smoking, and infectious diseases. However, while centre-level variables always result in a decrease of the variation our results illustrate that adjustment for individual risk factors can actually lead to changes in tau² in both directions. Overall, individual risk factors explained only a small to moderate amount of the prevalence variation.

325 Several of the most influential child-level variables leading to changes in tau² were potentially prone 326 to reverse causality, if changes had been made to the home environment following (and due to) the 327 onset of asthma or allergy in the child. The bias thereby introduced could be in either direction. For 328 instance, avoidance of pets by allergic families would tend to attenuate a harmful association of pets 329 with wheeze in the child. In contrast, avoidance of feather bedding following the child's asthma 330 diagnosis would accentuate risks associated with synthetic pillows and bedding. Reverse causality is 331 less of a concern for exposures in the first year of life, although selective avoidance by allergic 332 families could still introduce reverse causality biases. Our analyses restricted to children whose

parents reported making no such changes are generally reassuring. Except for carpets, the associations with the environmental factors were not affected substantially and making this allowance for possible reverse causality had less effect than the centre selection in this complementary analysis.

Therefore reverse causality does not seem to influence much our broad conclusions regarding the amount of centre-level variation that could be explained by the investigated environmental factors Nevertheless, these measured factors may actually reflect some other underlying unmeasured risk/protective factor. If the "true" determinant is measured imperfectly, the change in tau² will be only partial.

With the cross-sectional design we cannot safely infer causality even for the influential risk factors regarding the reduction of tau². These factors may actually reflect some other underlying unmeasured risk/protective factor. Because they therefore measure imperfectly the unmeasured factor the reduction in tau² will be only partial.

346 A positive skin prick test resulted in the same variance reduction as the most prominent 347 environmental risk factor. This occurred despite the fact that non-atopic asthma is important 348 worldwide [1] but seems plausible given the strong association of atopy with wheeze within centres 349 and the wide range of atopy prevalence across our study centres. In our previous work, we have 350 found an attributable fraction of atopy on asthma of 40.7% among the affluent centres and 20.3% 351 among the non-affluent centres that already highlighted the importance of atopy on the population 352 level especially for the affluent world [1]. Therefore, risk factors influencing strongly the 353 development of atopy can also be expected to account for some of the international variation in 354 wheeze prevalence, in addition to factors influencing asthma through mechanisms independent of 355 allergy.

356 In our dataset, ecological risk factors had a considerably greater explanatory potential than individual 357 risk factors, consistent with findings from social sciences regarding the importance of so-called 358 contextual factors. For an epidemiological example, it has been found that the wealth of a 359 neighbourhood has an effect on adult asthma prevalence independently of the individual's 360 socioeconomic status [14] reducing the between centre variation by 37%. For several risk factors in 361 the child's environment one could imagine a similar scenario as children move not only within their 362 homes but are in contact with their friends' and extended families' homes and public locations. For 363 example, it has been shown that community prevalence of cat keeping is a statistically significant 364 determinant of mattress cat allergen levels for non-cat owners [15].

The alternative explanation is that these ecological factors are indicators of different life styles between regions of the world, which would be the underlying overall cause. Indeed, Pearce and Douwes, in their review, propose that there is a Western "package" of environmental and social

368 factors that influence asthma prevalence while there is no known risk factor that would be able to 369 explain on its own either prevalence differences between populations or changes observed within 370 populations over time [16].

371 In our analysis, the variance explained by average centre-level exposure was generally not 372 diminished when incorporating the corresponding individual-level variable (Online Resource Table 373 E6). We therefore interpret these centre-level correlations as an indirect indication of the potential 374 role of contextual factors, and/or a surrogate for undiscovered individual-level or population-level 375 determinants. A strength of the present analysis is that it is the first multicentre comparison, made 376 possible by adapting new methodology (i.e. Bauer's scaling method [8]) and therefore being able to 377 compare multilevel logistic regression models that contain different individual risk factors. The study involved a large number of children, therefore the power for investigating individual risk factors is 378 379 high.

380 A limitation of our study is that even 30 centres worldwide represent a relatively small number of 381 potential centres especially when investigating centre-level variables and consequently uncertainty 382 around the estimates of between-centre variation is high. When looking at the estimates of tau² and 383 its standard error (of the null model), our estimate of 0.32 (SE 0.15) shows similar imprecision to 384 some other studies, e.g. 0.052 (SE 0.026) for the variance in psychiatric health care utilization [17] 385 (235 neighbourhoods) but higher than in other studies, e.g. the study on asthma prevalence in the 386 287 Chicago neighbourhoods (0.14 (SE 0.02) [18]). This calls for caution when gauging the 387 quantitative importance of the risk factors and part of the changes observed may well lie within the 388 range of uncertainty. To our knowledge, no paper has so far tackled this issue but generally just the 389 percentage of change was reported ([14,17,19,20]).

390 Our approach to handling missing observations was a fairly crude one. Unfortunately, almost none of 391 the currently available statistical software offers missing imputation for dichotomous variables in a 392 multilevel framework. However, in sensitivity analyses treating several centres with risk factor 393 information the same as a centre having no information for risk factors, our method to replace the 394 values for all children in the centre with the mean international prevalence proved quite robust i.e. 395 comparable to the original results. Overall, substituting the missing values with mean values is a 396 conservative approach which is expected to lead to an underestimation of the change in variance.

In conclusion, we found several risk factors, both at the individual level and the centre (population) level, that explained part of the large worldwide variation in prevalence of wheeze among children. Overall, individual risk factors explained a moderate amount of the variation in this international study, the most important remediable exposures being bedding material and maternal smoking. Atopy, measured by aeroallergen skin prick tests, also explained a proportion of the worldwide variation in wheeze prevalence. Our multi-centre study design overcomes the limitations of two-

- 403 centre comparisons and the multi-level modelling approach permits adjustment for the effect of
- 404 individual-level risk factors, which are excluded in most conventional ecological (centre-level)
- 405 analyses.

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447 References:

448 1. Weinmayr G, Weiland SK, Bjoïksteń B, Brunekreef B, Buchele G, Cookson WOC, et al. Atopic
449 sensitization and the international variation of asthma symptom prevalence in children. Am J Respir
450 Crit Care Med. 2007;176:565-74.

451 2. ISAAC Steering Comittee 1, Beasley R, Keil U, Von Mutius E, Pearce N. Worldwide variation in
452 prevalence of symptoms of asthma, allergic rhinoconjunctivitis, and atopic eczema: ISAAC. Lancet.
453 1998;351:1225-32.

3. Weiland SK, Björksteín B, Brunekreef B, Cookson WOC, von Mutius E, Strachan DP. Phase II of the
International Study of Asthma and Allergies in Childhood (ISAAC II): rationale and methods. Eur
Respir J. 2004;24:406-12.

457 4. von Mutius E, Martinez FD, Fritzsch C, Nicolai T, Roell G, Thiemann HH. Prevalence of asthma and 458 atopy in two areas of West and East Germany. Am J Respir Crit Care Med. 1994;149:358-64.

459 5. Wong GWK, Ko FWS, Hui DSC, Fok TF, Carr D, von Mutius E, et al. Factors associated with

difference in prevalence of asthma in children from three cities in China: multicentre epidemiological
survey. BMJ. 2004;329:486.

- 462 6. ISAAC Steering Comittee. ISAAC Phase Two.
- 463 <u>http://isaac.auckland.ac.nz/phases/phasetwo/phasetwo.html</u>. Date last updated: 5 Apr 2017. Date
 464 last accessed: 11 Oct 2017.
- 465 7. Weinmayr G, Dreyhaupt J, Jaensch A, Forastiere F, Strachan DP. Multilevel regression modelling to
 466 investigate variation in disease prevalence across locations. Int J Epidemiol. 2017;46:336-47.
- 467 8. Bauer DJ. A Note on Comparing the Estimates of Models for Cluster-Correlated or Longitudinal
 468 Data with Binary or Ordinal Outcomes. Psychometrika. 2008;74:97-105.
- 469 9. Richardson DB, Rzehak P, Klenk J, Weiland SK. Analyses of case-control data for additional
 470 outcomes. Epidemiology. 2007;18:441-5.
- 471 10. Weinmayr G, Gehring U, Genuneit J, Buchele G, Kleiner A, Siebers R, et al. Dampness and moulds
- in relation to respiratory and allergic symptoms in children: results from Phase Two of the
- 473 International Study of Asthma and Allergies in Childhood (ISAAC Phase Two). Clin Exp Allergy.
- 474 2013;43:762-74.
- 475 11. Weinmayr G, Forastiere F, Buchele G, Jaensch A, Strachan DP, Nagel G. Overweight / Obesity and
- 476 Respiratory and Allergic Disease in Children : International Study of Asthma and Allergies in
- 477 Childhood (ISAAC) Phase Two. PLoS One. 2014;9:ell3996.
- 478 12. García-Marcos L, Castro-Rodriguez JA, Suarez-Varela MM, Garrido JB, Hernandez GG, Gimeno
- AM, et al. A different pattern of risk factors for atopic and non-atopic wheezing in 9-12-year-old
 children. Pediatr Allergy Immunol. 2005;16:471-7.
- 481 13. Yemaneberhan H, Bekele Z, Venn A, Lewis S, Parry E, Britton J. Prevalence of wheeze and asthma
 482 and relation to atopy in urban and rural Ethiopia. Lancet. 1997;350:85-90.
- 483 14. Basagaña X, Sunyer J, Kogevinas M, Zock J-P, Duran-Tauleria E, Jarvis D, et al. Socioeconomic
 484 status and asthma prevalence in young adults: the European Community Respiratory Health Survey.
- 485 Am J Epidemiol. 2004;160:178-88.

- 486 15. Heinrich J, Bedada GB, Zock J-P, Chinn S, Norbäck D, Olivieri M, et al. Cat allergen level: its
 487 determinants and relationship to specific IgE to cat across European centers. J Allergy Clin Immunol.
 488 2006;118:674-81.
- 489 16. DouwesJ, Pearce N. Asthma and the westernization "package". IntJ Epidemiol. 2002;31:1098490 102.
- 491 17. Ivert AK, Torstensson Levander M, Merlo J. Adolescents' utilisation of psychiatric care,
- 492 neighbourhoods and neighbourhood socioeconomic deprivation: A multilevel analysis. PLoS One.493 2013;8.
- 494 18. Gupta RS, Zhang X, Sharp LK, Shannon JJ, Weiss KB. The protective effect of community factors on
 495 childhood asthma. J Allergy Clin Immunol. 2009;123:1297-1304.e2.
- 496 19. Kogan MD, Singh GK, Dee DL, Belanoff C, Grummer-Strawn LM. Multivariate analysis of state
 497 variation in breastfeeding rates in the United States. Am J Public Health. 2008;98:1872-80.
- 498 20. Merlo J, Ostergren PO, Broms K, Bjorck-Linne A, Liedholm H. Survival after initial hospitalisation
- for heart failure: a multilevel analysis of patients in Swedish acute care hospitals. J EpidemiolCommunity Health. 2001;55:323-9.
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Individual level variables Centre level variables⁰ (30 centres; 45297 - 51081 children¹³) (30 centres; 53748 children⁰¹) OR per 10% relative centre-level correlation relative increase in risk OR prevalence Risk factor change in change in factor between wheeze prevalence (LCL-UCL) range tau²(%) tau²(%) and risk factor prevalence prevalence (LCI-UCI) 1.33 115 Bedding: synthetic quilt (present) 1.9-87.9 -8.4 -23.6 0.5661 (1.09-1.61) (1.04 - 1.26)0.78 1.05 0.2553 Floor covering: fitted or loose carpets (present) 6.2-98.1 6.8 -6.6 (0.65-0.94) (0.98-1.13) 1 28 1 15 Bed Bed Mot Mot Mot Pillo Dise

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Table 1: Wheeze prevalence - change in the between-centre variance tau² by individual level variables and centre level variables

Bedding: synthetic quilt (fy)	2.2-75.2	-6.7	(1.08-1.53)	-21.8	(1.04-1.28)	0.5546
Bedding: feather quilt (present)	1.2-61.7	-6.7	0.56 (0.47-0.65)	-8.2	0.88 (0.76-1.04)	-0.2701
Mother smoked during the child's fy of life	0.1-43.8	-5.2	1.23 (1.12-1.36)	-32.7	1.27 (1.11-1.45)	0.6252
Mother smoked during pregnancy	0.2-33.9	-4.8	1.28 (1.13-1.44)	-34.4	1.38 (1.16-1.64)	0.6580
Mother smokes at present	0.1-48.3	-4.1	1.19 (1.07-1.32)	-24.1	1.20 (1.06-1.37)	0.5677
Pillow: feather (present)	0.4-91.9	-3.9	0.60 (0.42-0.85)	-3.3	0.95 (0.86-1.05)	-0.2822
Disease: worm infection	2.4-99.9	3.4	1.31 (1.03-1.65)	-5.0	0.89 (0.78-1.02)	0.1963
Disease: whooping cough	0.6-48.9	3.1	1.62 (1.46-1.80)	-0.8	0.96 (0.81-1.14)	-0.1100
Bedding: feather quilt (fy)	1.2-86.2	-3.0	0.66 (0.58-0.75)	-3.2	0.94 (0.82-1.07)	-0.2075
Heating inside home (fy)	0.8-98.5	-2.9	1.18 (1.04-1.35)	-5.8	1.05 (0.97-1.15)	0.2913
Heating: wood (fy)	0.1-79.6	-2.9	1.22 (0.96-1.55)	-5.7	1.06 (0.96-1.18)	0.4170
Damp or mould (fy)	6.5-36.7	-2.8	1.65 (1.43-1.89)	-5.9	1.24 (0.89-1.73)	0.4396
Heating inside home (present)	0.8-99.9	-2.6	1.11 (1.02-1.21)	-9.0	1.06 (0.99-1.14)	0.4308
Heating: wood (present)	0-77.2	-2.6	1.10	-24.4	1.16	0.6639

			(0.90-1.34)		(1.05-1.29)	
Vaccination: tuberculosis	16.5-99.7	2.4	1.07 (0.94-1.21)	-21.2	0.90 (0.83-0.97)	-0.3518
Pillow: no pillow (fy)	2.7-56.1	2.3	0.82 (0.69-0.99)	-8.9	1.14 (0.97-1.34)	0.0979
Pillow: feather (fy)	1.3-75.8	-2.2	0.82 (0.71-0.94)	-4.4	0.94 (0.83-1.05)	-0.3412
Cooking: coal/wood (present)	0-99.3	2.1	1.21 (1.11-1.31)	-2.6	0.94 (0.84-1.06)	0.1318
Damp or mould (present)	2.2-47.1	-2.1	1.51 (1.29-1.76)	-2.6	1.09 (0.88-1.35)	0.3975
Pillow: synthetic fibre (fy)	1.1-85.4	2.0	1.23 (1.06-1.42)	-2.4	0.95 (0.85-1.07)	0.0054
Pillow: foam (fy)	0-79.9	-1.7	1.10 (0.97-1.24)	-16.3	1.17 (1.02-1.33)	0.6186
Cooking: coal/wood (fy)	0-99.2	1.7	1.13 (0.93-1.37)	-5.0	0.93 (0.83-1.05)	-0.1531
Cooking: electricity (fy)	0.3-99.2	1.5	0.86 (0.72-1.03)	-0.6	1.01 (0.94-1.09)	0.1574
Heating inside home / Cooking: gas, oil, coal, coke, wood (present)	0.8-100	-1.4	1.08 (0.95-1.22)	-6.0	1.06 (0.97-1.15)	0.2283
ETS: 10 or more cigarettes	4.5-34.4	-1.3	1.17 (1.03-1.32)	-12.8	1.33 (1.00-1.77)	0.5051
Air conditioning: present	1.2-95.4	-1.3	0.93 (0.85-1.01)	-9.3	0.91 (0.82-1.02)	-0.4027
Breastfeeding 6 months or more	13.5-99.4	-1.2	0.95 (0.86-1.06)	-4.4	0.94 (0.85-1.03)	0.0776
Cooking: gas (fy)	0.5-98.9	-1.1	1.10 (0.95-1.29)	-1.8	1.03 (0.96-1.10)	0.0215

a: This table contains all variables that lead to a change of >1% in tau² when investigated as individual level variables, b: differences in the number of children relate to different numbers of missing values for the respective questions: this is because, in the case of stratified subsamples, we did not impute missing values (details see Online Resource); c: each child has the value of the mean exposure for children in its centre, i.e. all children in the same centre have the same contextual exposure; d.: all children with information on wheeze got assigned a value; fy = first year of life of the child; ETS = Environmental tobacco smoke;

Table 2: Wheeze prevalence among atopic and non-atopic children - change in the between-centre variance tau² by individual level variables³.

	Wheeze among atopies			Wheeze among non-atopics		
	(29 centres; 6390 - 7302 children ⁶)(29 cer			ntres; 20335 - 2		
Risk factor	prevalence range	relative change in tau ² (%)	OR (LCL-UCL)	prevalence range	relative change in tau ² (%)	OR (LCL-UCL)
Bedding: synthetic quilt (present)	4.0-89.8	-15.6	1.54 (1.25-1.90)	3.8-88.3	-4.4	1.29 (1.02-1.62)
Floor covering: fitted or loose carpets (present)	3.9-98.5	8.4	0.68 (0.56-0.83)	6.2-98.2	0.6	0.98 (0.77-1.24)
Bedding: synthetic quilt (fy)	0-85.0	-10.8	1.36 (0.99-1.86)	1.5-74.0	-4.1	1.33 (0.99-1.78)
Bedding: feather quilt (present)	0-59.1	-1.1	0.62 (0.48-0.80)	0.9-61.7	-9.4	0.62 (0.49-0.77)
Mother smoked during the child's fy of life	0-45.7	-3.0	1.20 (1.03-1.39)	0.1-45.4	-7.2	1.50 (1.33-1.69)
Mother smoked during pregnancy	0-32.2	-3.8	1.27 (1.01-1.61)	0.2-35.6	-4.4	1.40 (1.18-1.66)
Mother smokes at present	0-49.5	-1.9	1.16 (1.03-1.32)	0.1-50.4	-4.9	1.37 (1.23-1.52)
Pillow: feather (present)	0-84.8	-2.3	0.66 (0.39-1.13)	0.4-93.8	-2.7	0.81 (0.62-1.07)
Disease: worm infection	2.3-99.4	-0.7	0.96 (0.78-1.18)	2.8-100.0	1.3	1.58 (1.26-1.98)
Disease: whooping cough	0-44.1	7.5	1.86 (1.50-2.31)	0.7-45.6	3.6	1.55 (1.26-1.90)
Bedding: feather quilt (fy)	0-85.9	-2.9	0.63 (0.44-0.90)	1.0-86.7	-3.7	0.68 (0.43-1.08)
Heating inside home (fy)	1.0-98.2	-1.6	1.36 (1.09-1.69)	0.5-96.3	-3.1	1.24 (0.97-1.57)
Heating: wood (fy)	0-80.0	-2.5	1.48 (1.26-1.74)	0.1-79.2	-2.5	1.29 (0.92-1.82)
Damp or mould (fy)	5.6-45.8	-2.7	1.80 (1.50-2.16)	5.6-38.0	-3.1	1.62 (1.29-2.05)
Heating inside home (present)	1.0-96.6	-1.9	1.17 (0.98-1.40)	0.6-98.4	-5.0	1.22 (0.99-1.50)
Heating: wood (present)	0-75.0	-5.9	1.30	0-78.1	-4.0	1.24

			(1.04-1.64)			(0.93-1.65)
Vaccination: tuberculosis	14.3-100.0	15.8	1.47 (1.10-1.98)	16.2-99.7	-2.0	0.92 (0.65-1.29)
Pillow: no pillow (fy)	0-57.1	2.0	0.89 (0.78-1.02)	2.6-56.0	1.6	0.77 (0.55-1.07)
Pillow: feather (fy)	1.9-82.1	2.3	1.21 (0.58-2.52)	1.1-77.1	-1.7	0.75 (0.42-1.32)
Cooking: coal/wood (present)	0-100.0	1.2	1.22 (0.86-1.74)	0-99.5	0.8	1.39 (1.18-1.64)
Damp or mould (present)	0-51.9	-2.6	1.61 (1.36-1.90)	1.1-48.1	-2.9	1.41 (1.09-1.83)
Pillow: synthetic fibre (fy)	0.5-97.0	-0.1	1.02 (0.65-1.60)	0.3-85.5	4.6	1.30 (0.85-1.99)
Pillow: foam (fy)	0-77.2	-1.8	1.11 (0.85-1.44)	0-82.2	-2.0	1.13 (0.95-1.35)
Cooking: coal/wood (fy)	0-94.4	0.6	1.13 (0.87-1.45)	0-99.2	2.0	1.25 (0.99-1.57)
Cooking: electricity (fy)	1.0-99.3	3.4	0.74 (0.55-0.99)	0.2-99.2	0.2	0.85 (0.69-1.06)
Heating inside home / Cooking: gas, oil, coal, coke, wood (present)	0-100.0	-3.1	1.25 (0.99-1.57)	0.9-100.0	-0.2	1.01 (0.89-1.14)
ETS: 10 or more cigarettes	4.5-38.3	-0.1	1.10 (0.79-1.53)	4.8-34.5	-2.4	1.28 (1.11-1.48)
Air conditioning: present	0-97.0	-0.4	0.83 (0.70-0.98)	0.7-96.9	-2.0	0.91 (0.77-1.06)
Breastfeeding 6 months or more	8.4-100.0	-0.2	0.99 (0.81-1.21)	12.6-99.4	-0.3	0.91 (0.82-1.01)
Cooking: gas (fy)	0-99.0	-0.7	1.39 (1.10-1.76)	0.6-98.9	0.7	0.95 (0.82-1.10)

a: This table contains the same variables as Table 1 listing the variables that resulted in the strongest changes of tau² in the whole population (as opposed to non-atopic or atopic children only, which is shown in Table E2 in the Online Resource)b: differences in the number of children relate to different numbers of missing values for the respective questions;; fy = first year of life of the child; ETS = Environmental tobacco smoke

Table 3: Wheeze prevalence - change in the between-centre variance tau² by individual level variables in multivariate models

	All children (N centres=30, N children=50,852ª)	Atopic children (N centres=29, N children=7,285 ^{a\$})	Non-atopic children (N centres=29, N children=23,418 ^a)
	Relative change in tau² (%)	Relative change in tau²(%)	Relative change in tau² (%)
factors that result in increase or decrease in tau ²		<u>, k</u>	<u> </u>
Bedding: synthetic quilt at present add Floor covering: fitted or loose	-8.4	-16.4	-4.3
carpet at present add Bedding: feather quilt at present add Mother smoked during first year	-1.0 -3.7	-5.2 -3.3	-6.3 -14.1
of life	-8.2	-5.0	-20.0
add Worm infection	-4.4	-6.5	-19.0
add Whooping cough	-3.3	-4.7	-16.9
add Heating inside home fy	-5.7	-4.9	-19.7
add Damp or mould fy	-7.9	-5.7	-22.8
add Vaccination: tuberculosis	-3.2	4.1	-20.0
add Pillow: no pillow fy	-0.9	5.6	-18.2
add Cooking: coal/wood at present	2.3	7.3	-15.3
add Pillow: foam fy	2.2	6.7	-15.7
add Cooking: electricity fy	2.4	11.6	-15.9
only factors that result in a decrease in tau ² in all children			
Bedding: synthetic quilt at present	-8.4	-16.4	-4.3
add Bedding: feather quilt at present add Mother smoked during first year	-10.5	-13.4	-12.0
of life	-15.0	-15.1	-17.6
add Heating inside home fy	-17.2	-15.0	-21.3
add Damp or mould fy	-19.0	-15.2	-24.9
add Pillow: foam fy	-20.5	-16.8	-26.8

fy: first year of life of the child; for choice of variables for multivariate model see Methods section in the Online Resource, a: in the stratified subsamples only children included with risk factor information were included resulting in 50852 children (in contrast to the 53748 children who had information on wheeze).

Figure legend:

Fig.I : Predicted prevalence in the centres in the null model and after incorporating the risk factors for the latter a reference population with risk factor prevalences equal to the arithmetic mean of all centres was used. Model with decrease and increase tau²: risk factors are included irrespective of the direction of change in the between-centre variance; Model with decrease tau²: only risk factors that lead to a reduction in the between-centre variance are included for illustrative purpose, (for detailed methods see Online Resource).



Figure 1: Predicted prevalence in the centres in the null model and after incorporating the risk factors. Model with decrease and increase tau: risk factors are included irrespective of the direction of change in the between-centre variance; Model with decrease tau: only risk factors that lead to a reduction in the between-centre variance are included for illustrative purpose.