Parental occupational exposure pre- and post-conception and development of asthma in offspring

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Abstract

Background: While direct effects of occupational exposures on an individual’s respiratory health are evident, a new paradigm is emerging on the possible effects of pre-conception occupational exposure on respiratory health in offspring. We aimed to study the association between parental occupational exposure starting before conception and asthma in their offspring (at 0–15 years of age).

Methods: We studied 3985 offspring participating in the Respiratory Health in Northern Europe, Spain and Australia (RHINESSA) generation study. Their mothers or fathers (n = 2931) previously participated in the European Community Respiratory Health Survey (ECRHS). Information was obtained from questionnaires on parental job history pre- and post-conception which was linked to an asthma-specific job-exposure matrix (JEM). We assessed the association between parental occupational exposure and offspring asthma, applying logistic regression models, clustered by family and adjusted for study centre, offspring sex, parental characteristics (age, asthma onset, place of upbringing, smoking) and grandparents’ level of education.

Results: Parental occupational exposure to microorganisms, pesticides, allergens or reactive chemicals pre-conception or both pre- and post-conception was not related to offspring asthma; in general, subgroup analyses confirmed this result. However, maternal exposure both pre- and post-conception to allergens and reactive chemicals was associated with increased odds for early-onset asthma in offspring (0–3 years of age); odds ratio 1.70 (95% CI: 1.02–2.84) and 1.65 (95% CI: 0.98–2.77), respectively.

Conclusions: This study did not find evidence that parental occupational exposure, defined by an asthma JEM before conception only or during pre- and post-conception vs non-exposed, was associated with offspring asthma.

Key words: Epidemiology, asthma, generation study, occupational exposure, air pollutants, occupation, job-exposure matrices

Introduction

In the early 1990s, associations were found between low birth weight or preterm birth and increased risk of ischaemic heart disease in adulthood.1,2 These findings led to the theory that exposures in utero could affect offspring health in adulthood,3–5 the Barker Hypothesis,1,2 and further developed into the Developmental Origins of Health and Diseases, including the role of early life environmental exposures such as chemicals in the aetiology of adult disease.5

During the last 30 years, the incidence of asthma has increased considerably,6,7 and asthma is now one of the most common global chronic diseases in children and adolescents.6–8 Occupational exposures to e.g. chemical and biological agents are known to affect respiratory health in adult workers,9–12 and a substantial number of agents have been linked to occupation-related asthma.13 It is estimated that ~15% of adult asthma cases could be prevented if occupational exposures were avoided.14 A few studies15,16...
have also suggested that maternal occupational exposures, for example to low molecular weight agents (from e.g. cleaning agents, hairdressing and dentistry chemicals, wood dust) during pregnancy can pose a risk for asthma in offspring. Furthermore, a recent study using data from the Respiratory Health in Northern Europe (RHINE) study\(^{20,21}\) found that paternal pre-conception exposure to occupational welding may influence the risk of asthma among their children born years after this exposure. These results suggest that exposures after the age of 15 years and before conception may constitute a ‘window of susceptibility’ among males to having future offspring with asthma.\(^{17}\) Evidence from mainly animal studies suggests that pre-conception environmental exposures in males may induce epigenetic alterations or damage sperm DNA, which may later be transferred to the offspring through the male germline.\(^{18,19}\)

In this study, we aimed to investigate whether a range of parental occupational exposures influenced the risk of asthma in offspring, relative to the timing of exposure (pre-conception only vs both pre- and post-conception) and parental sex. We hypothesized that parental sex modifies the development of offspring asthma through pre-conception exposures in fathers and around pregnancy in mothers.

### Methods

#### Study population

The study population was adult offspring \(\geq 18\) years of age from 10 centres who provided self-reported information collected in the RHINESSA generation study \((n = 8204)\)—Respiratory Health in Northern Europe, Spain and Australia (www.rhinessa.net). Either the father or the mother \((n = 6026)\) of each offspring in the RHINESSA study had previously participated in the longitudinal European Community Respiratory Health Survey (ECRHS) initiated in 1989–1992 (www.ecrhs.org). The ECRHS was initially conducted to estimate variations in the prevalence of asthma and allergy in young adults in Europe and other parts of the world in a randomized sample of the general population aged 20–44 years. The ECRHS was performed in three stages, and we used the second stage (ECRHS II) which provides information on parents’ previous total job history. A detailed description of the RHINESSA generation study can be found elsewhere.\(^{20,21}\)

In this two-generation, population-based cohort study, we used information from the 3985 eligible adult offspring \((18–52\) years—collected in RHINESSA, 2013–2016) with information on one of their parents, \(n = 2931\) \((26–54\) years—collected in ECRHS II, 1999–2000). Information about the parents was obtained from parent’s self-reported data and the offspring’s information was obtained from the offspring’s self-reported data. We also obtained information about the educational level of the parents’ parents.

#### Data collection

For the seven Northern European ECRHS centres [Reykjavik (Iceland), Bergen (Norway), Umeå, Uppsala and Göteborg (Sweden), Aarhus (Denmark) and Tartu (Estonia)], parental information was available from a detailed self-administered questionnaire survey, RHINE II (the RHINE study, www.rhine.nu). Melbourne (Australia) and Huelva and Albacete (Spain) used interviewer-administered questionnaires from ECRHS II. Importantly, offspring information was obtained directly from the offspring themselves and available from the RHINESSA questionnaire. All questionnaires are available online from the study webpages listed above.

#### Eligibility and exclusion criteria

The eligibility criteria for inclusion were: offspring \(\geq 18\) years of age when they participated in the RHINESSA study, with a parent who had earlier participated in the ECRHS II/RHINE II. A flowchart of the study population is provided in Figure 1. We excluded 2439 parents and their 3288 offspring due to (i) missing information of job history, (ii) exposures to occupational exposures other than those of interest, (iii) no exposure in the relevant exposure windows, or (iv) missing information about the timing of exposures. Furthermore, 344 offspring and their 168 parents were excluded due to missing information about offspring asthma or the late onset of offspring asthma \((> 15\) years of age) to avoid contamination of data by occupational exposures of offspring. Information on parental educational level was not available for the centres in Melbourne, Huelva and Albacete. To keep these centres in our analysis, we applied two different adjustment models of co-variables, performing analysis both with and without adjustment for educational level. Due to missing co-variables (except parental educational level), we excluded 597 offspring and 488 parents; and due to missing information on parental educational level, we excluded 246 offspring and 165 parents in our fully adjusted model. A comparison of those included in the study vs those excluded is available in Supplementary Table S1, available as Supplementary data at IJE online. Those excluded had a lower educational level and a higher percentage were smokers compared with those included in the study.
Figure 1 Flowchart of the exclusion process of the study population. Model 1 + 2—full information of study centre, offspring’s sex, parent’s characteristics (sex, age, asthma onset, place of upbringing and smoking) and grandmother’s and grandfather’s educational level. Model 3—full information of Model 1 + 2 and parent’s educational level. *Occupational exposures of interest: 1, microorganisms (moulds, endotoxin); 2, pesticides (herbicides, insecticides, fungicides); 3, allergens (animals, flour, house dust mites, storage mites, plant mites, enzymes, latex, fish/shellfish); 4, reactive chemicals (high level chemical disinfectants, isocyanates, acrylates, epoxy resins, persulfates/henna, aliphatic amines, bleach).
Parental occupational exposures

The question about parental job history was formulated as:

1. RHINE II—‘List your jobs including branch, your work tasks and the period of employment. Periods shorter than six months need not to be specified. Employment also includes work done by people with their own company.’

2. ECRHSII—‘If employed or self-employed or a full time house-person go to Q28: “Q 28. If you had more than one job in the same company, or if you were doing more than one job at the same time, we would like to talk about them separately. Please start with your current or last job.’

Parental job history was linked to an occupational asthma-specific job-exposure matrix (OAsJEM) to yield information about exposure to 30 specific agents. For each of these agents, experts have evaluated exposure for each job code in the International Standard Classification of Occupations, 1988 (ISCO-88). We focused on the 20 most specific agents grouped into four main exposure groups.

1. Microorganisms (molds, endotoxin).
2. Pesticides (herbicides, insecticides, fungicides).
3. Allergens (animals, flour, house dust mites, storage mites, plant mites, enzymes, latex, fish/shellfish).
4. Reactive chemicals (high-level chemical disinfectants, isocyanates, acrylates, epoxy resins, persulfates/henna, aliphatic amines, bleach).

Each of the four groups of agents was dichotomized with respect to exposure (yes/no) for each of the exposure windows investigated: (i) pre-conception only; (ii) both pre- and post-conception; (iii) post-birth only; and (iv) unexposed (the reference group). However, due to the small sample size of exposed post-birth only (n = 145) even before stratification by each exposure category and parent’s sex, post-birth only was not included. In contrast, the pre-conception window included 330 exposed and 51 asthma cases and during both the pre- and post-conception windows combined, 994 were exposed including a total of 147 asthma cases. Occupational exposure pre-conception only was chosen if the employment ended ≥2 years before the birth of a parent’s offspring. Occupational exposures during both pre- and post-conception periods were chosen if the parent was exposed both before and after the offspring’s year of birth (Figure 2).

Asthma phenotypes in offspring

Adult offspring themselves provided information in the RHINESSA study on their ever asthma status through the following questions: ‘Do you have or have you ever had asthma?, and if yes, information about age at asthma onset: ‘How old were you when you first experienced asthma symptoms?’ We included offspring asthma with onset between 0 and 15 years of age as the main outcome, which was further divided into asthma onset between 0 and 3 years (early-onset asthma) and between 4 and 15 years (late-onset asthma). An affirmative answer to the questions: ‘Do you have any nasal allergies including hay fever?’ or ‘Have you ever had eczema or any kind of skin allergy?’ was defined as offspring atopy.

Statistical procedures

Multivariable logistic regression models were applied to estimate the odds ratio (OR) of the association between parental occupational exposure and offspring asthma (onset at 0–15 years of age) using the PROC GENMOD function.
in the software package SAS. Each of the four different agent exposure groups was analysed separately. The reference group consisted of working parents not exposed to any of the 30 agents from the OAsJEM.

A priori, it was decided to separate all analyses by sex of the parent, adjust for study centre and include clustering by family to account for multiple offspring (siblings) from the same parent. We identified potential co-variables based on the literature and discussed direction and relationships of co-variables using directed acyclic graphs (DAGs) (Supplementary Figure S1, available as Supplementary data at IJE online). To be identified as a confounder in the DAGs, the covariate had to be a risk factor for both exposure and outcome. The following covariates were included as confounders from the DAGs: parent’s characteristics (age, early/late asthma onset, place of upbringing, educational level) and educational level of grandparents. Smoking and offspring sex were not considered as confounders according to the DAGs; however, smoking and offspring sex were included as co-variables in accordance with earlier studies.17,25–27 We applied the following three models.

- Model 1: Clustered by family and adjusted for centre.
- Model 2: Model 1 and further adjustment for offspring sex, the parent's characteristics (age, early/late asthma onset, place of upbringing and smoking) and educational level of grandparents.
- Model 3: Model 2 and further adjustment for parent’s educational level.

In the description of results, we generally referred to Model 2. The significance level was set at a P-value of <0.05 (two-sided) and 95% confidence intervals (CIs) were calculated.

A priori, subgroup analyses were included to examine if offspring age at asthma onset, offspring atopy or offspring sex modified the association between parental occupational exposure and risk of offspring asthma. The impact of offspring age on asthma onset was investigated by multinomial logistic regression models (0–3 and 4–15 years of age) using the PROC GEE function in SAS (SAS Institute Inc., Cary, NC, USA). Due to statistical power issues, subgroup analyses were only performed for allergens and chemicals. All statistical analyses were performed using SAS version 14.1.

**Results**

The baseline characteristics of 3985 offspring, 2931 parents and grandparents are presented in Table 1. Offspring mean age was 30 years [standard deviation (SD) 7.5] when they responded to the questionnaire. A slightly higher proportion of offspring of exposed parents developed early-onset asthma (4.5%) compared with the offspring of non-exposed parents (3.6%). Parental mean age was 43 years (SD 6.5) when they responded to the questionnaire. More women than men participated, both among offspring (57.0% women) and parents (59.5% women). Exposed parents were more likely to have asthma, have lived on a farm and to have a lower educational level compared with unexposed parents.

Multivariable logistic regression analyses (Table 2) showed no clear associations between parent’s occupational exposure and offspring asthma at 0–15 years of age, neither for pre-conception exposures only nor for exposures both pre- and post-conception. These findings were consistent for both fathers and mothers in all four groups of exposure agents. Of note, no results are presented for maternal exposure to microorganisms and pesticides pre-conception only, due to a very low number of exposed individuals.

**Subgroup analyses**

Subgroup analyses of the age of asthma onset (Table 3) supported the main results (Table 2) with two exceptions. Maternal exposure to allergens during both pre- and post-conception periods was associated with increased odds for early-onset asthma, OR_{Model2} 1.70 (1.02–2.84). Similarly, maternal exposure to chemicals both pre- and post-conception increased the odds for early-onset asthma [OR_{Model2} 1.65 (0.98–2.77)], also seen in Models 1 [(OR 1.87 [1.12–3.13]) and 3 (OR 1.82 (1.06–3.12)]. However, no association was found for maternal allergen or chemical exposures and late-onset asthma, OR_{Model2} 1.03 (0.73–1.45) and OR_{Model2} 1.03 (0.73–1.45), respectively.

The subgroup analyses stratified by offspring allergic status (Table 4) and by offspring sex (Table 5) showed similar results for allergic and non-allergic asthma as well as offspring sex. No results are available for parental exposure to allergens pre-conception only and early-onset asthma due to very few exposed individuals. Similarly, no results are presented for maternal exposure to chemicals pre- and post-conception and early-onset asthma.

**Discussion**

In this study, parental occupational exposure to microorganisms, pesticides, allergens or reactive chemicals pre-conception only as well as both pre- and post-conception did not show clear associations with offspring development of childhood asthma. In general, subgroup analyses by age of asthma onset, offspring atopic status and offspring sex confirmed the negative finding except for an increased
Table 1 Characteristics of offspring, parents and grandparents by parents exposed to either microorganisms, pesticides, allergens and/or chemicals or non-exposed

|                      | Exposed | Non-exposed | Total  |
|----------------------|---------|-------------|--------|
| **Offspring, n**     | 1324    | 2661        | 3985   |
| Female, n (%)        | (56.1)  | (57.4)      | (57.0) |
| Age at data collection mean, SD | 28.5 (6.7) | 30.3 (7.7) | 29.7 (7.5) |
| Asthma 0–15 years, n (%) | 198 (15.0) | 377 (14.2) | 575 (14.4) |
| Asthma onset 0–3 years, n (%) | 59 (4.5) | 97 (3.6) | 156 (3.9) |
| Asthma onset 4–15 years, n (%) | 139 (10.5) | 280 (10.5) | 419 (10.5) |
| Allergy, n (%)       | 748 (56.5) | 1535 (57.7) | 2283 (57.3) |
| Centre               |         |             |        |
| Aarhus, n (%)        | 80 (6.0) | 239 (9.0)   | 319 (8.0) |
| Albacete, n (%)      | <5 (<0.4) | <25 (<0.9) | <30 (<0.8) |
| Huelva, n (%)        | <5 (<0.4) | <15 (<0.6) | <20 (<0.3) |
| Reykjavik, n (%)     | 134 (10.1) | 399 (15.0) | 533 (13.4) |
| Bergen, n (%)        | 206 (15.6) | 581 (21.8) | 787 (19.7) |
| Gothenburg, n (%)    | 83 (6.3) | 204 (7.7) | 287 (7.2) |
| Umea, n (%)          | 422 (31.9) | 397 (14.9) | 819 (20.6) |
| Uppsala, n (%)       | 327 (24.7) | 529 (19.9) | 856 (21.5) |
| Tartu, n (%)         | 57 (4.3) | 155 (5.8) | 212 (5.3) |
| Parents, n           | 995     | 1936        | 2931   |
| Female               | 572 (57.5) | 1065 (55.0) | 1637 (55.9) |
| Age at data collection mean, SD | 42.4 (6.6) | 43.8 (6.4) | 43.3 (6.5) |
| Asthma <10 years, n (%) | 40 (4.0) | 68 (3.5) | 108 (3.7) |
| Asthma >10 years, n (%) | 81 (8.1) | 149 (7.7) | 230 (7.8) |
| Place of upbringing, n (%) |         |             |        |
| Farm                 | 240 (24.1) | 315 (16.3) | 555 (18.9) |
| Village, small town, suburb of city | 636 (65.9) | 1317 (68.0) | 1973 (67.3) |
| Inner city           | 99 (9.9) | 304 (15.7) | 403 (13.7) |
| Educational levelb   |         |             |        |
| Primary, n (%)       | 112 (11.6) | 173 (9.6) | 285 (10.3) |
| Secondary, n (%)     | 411 (42.5) | 637 (35.4) | 1048 (37.9) |
| University/College, n (%) | 445 (46.0) | 988 (54.9) | 1433 (51.8) |
| Smokers, n (%)       | 510 (51.3) | 1015 (52.4) | 1525 (52.0) |
| Centre               |         |             |        |
| Aarhus, n (%)        | 66 (6.6) | 192 (9.9) | 258 (8.8) |
| Albacete, n (%)      | <5 (<0.5) | <20 (<1.0) | <25 (<0.9) |
| Huelva, n (%)        | <5 (<0.5) | <10 (<0.5) | <15 (<0.5) |
| Reykjavik, n (%)     | 112 (11.3) | 306 (15.8) | 418 (14.3) |
| Bergen, n (%)        | 149 (15.0) | 405 (20.9) | 554 (18.9) |
| Gothenburg, n (%)    | 66 (6.6) | 153 (7.9) | 219 (7.5) |
| Umea, n (%)          | 304 (30.6) | 276 (14.3) | 580 (19.8) |
| Uppsala, n (%)       | 236 (23.7) | 380 (19.6) | 616 (21.0) |
| Tartu, n (%)         | 49 (4.9) | 131 (6.8) | 180 (6.1) |
| Grandmothers, n      | 995     | 1936        | 2931   |
| Educational level    |         |             |        |
| Primary, n (%)       | 692 (69.5) | 1287 (66.5) | 1979 (67.5) |
| Secondary, n (%)     | 216 (21.7) | 470 (24.3) | 686 (23.4) |
| University/College, n (%) | 87 (8.7) | 179 (9.2) | 266 (9.1) |
| Grandfathers, n      | 995     | 1936        | 2931   |
| Educational level    |         |             |        |
| Primary, n (%)       | 584 (58.7) | 1019 (52.6) | 1603 (54.7) |
| Secondary, n (%)     | 271 (27.2) | 579 (29.9) | 850 (29.0) |
| University/College, n (%) | 140 (14.1) | 338 (17.5) | 478 (16.3) |

aDue to data security, i.e. the European Data Protection Regulation, we are not allowed to show results based on microdata (<5 persons); data are available on request for authorized research units.

bEducational level, missing: exposed n = 27, non-exposed n = 138, total n = 165.
odds for early-onset offspring asthma in the case of maternal pre- and post-conception exposure to allergens and chemicals.

Comparison with other studies

Maternal occupational exposure and offspring asthma
We did not identify any other studies investigating pre-conception occupational exposure in mothers, but maternal exposure to occupational factors during pregnancy has been reported to be related to offspring asthma in previous studies. Christensen et al. found that combined intrauterine and postnatal occupational exposure to low molecular weight agents, but not high molecular weight agents, might be related to asthma in 7-year-old children. We did find associations between mothers exposed to allergens both pre- and post-conception, though only for early-onset asthma (0–3 years). Christensen et al. did not

### Table 2
Multivariable logistic regression analyses clustering by family of paternal or maternal occupational exposure in relation to asthma in offspring (0–15 years)

|          | n   | Asthma, % | ORModel1 \(^a\) | 95 % CI | ORModel2 \(^b\) | 95 % CI | ORModel3 \(^c\) | 95 % CI |
|----------|-----|-----------|------------------|--------|------------------|--------|------------------|--------|
| **Microorganisms** |     |           |                  |        |                  |        |                  |        |
| **Fathers** |     |           |                  |        |                  |        |                  |        |
| Unexposed | 1216 | 15.5 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 |
| Exposed pre-conception only | 40 | 20.0 | 1.22 | (0.52; 2.86) | 1.19 | (0.49; 2.90) | 1.26 | (0.53; 3.01) |
| Exposed both pre- and post-conception | 109 | 11.0 | 0.57 | (0.28; 1.13) | 0.53 | (0.26; 1.06) | 0.61 | (0.30; 1.24) |
| **Mothers** |     |           |                  |        |                  |        |                  |        |
| Unexposed | 1445 | 13.0 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 |
| Exposed pre-conception only | 18 | 0.0 | – | – | – | – | – | – |
| Exposed both pre- and post-conception | 31 | 12.9 | 0.96 | (0.36; 2.60) | 0.66 | (0.26; 1.64) | 0.56 | (0.21; 1.4) |
| **Pesticides** |     |           |                  |        |                  |        |                  |        |
| **Fathers** |     |           |                  |        |                  |        |                  |        |
| Unexposed | 1216 | 15.5 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 |
| Exposed pre-conception only | 51 | 17.6 | 1.12 | (0.49; 2.55) | 1.09 | (0.46; 2.54) | 1.13 | (0.48; 2.65) |
| Exposed both pre- and post-conception | 126 | 13.5 | 0.72 | (0.39; 1.30) | 0.67 | (0.37; 1.22) | 0.74 | (0.40; 1.37) |
| **Mothers** |     |           |                  |        |                  |        |                  |        |
| Unexposed | 1445 | 13.0 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 |
| Exposed pre-conception only | 16 | 0.0 | – | – | – | – | – | – |
| Exposed both pre- and post-conception | 32 | 9.4 | 0.77 | (0.24; 2.42) | 0.56 | (0.19; 1.69) | 0.46 | (0.14; 1.51) |
| **Allergens** |     |           |                  |        |                  |        |                  |        |
| **Fathers** |     |           |                  |        |                  |        |                  |        |
| Unexposed | 1216 | 15.5 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 |
| Exposed pre-conception only | 76 | 10.5 | 0.70 | (0.31; 1.59) | 0.66 | (0.29; 1.51) | 0.67 | (0.29; 1.59) |
| Exposed both pre- and post-conception | 235 | 12.8 | 0.78 | (0.50; 1.23) | 0.77 | (0.48; 1.21) | 0.83 | (0.52; 1.31) |
| **Mothers** |     |           |                  |        |                  |        |                  |        |
| Unexposed | 1445 | 13.0 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 |
| Exposed pre-conception only | 118 | 17.8 | 1.36 | (0.79; 2.31) | 1.04 | (0.59; 1.83) | 0.96 | (0.54; 1.70) |
| Exposed both pre- and post-conception | 523 | 16.3 | 1.26 | (0.95; 1.69) | 1.15 | (0.85; 1.54) | 1.16 | (0.85; 1.57) |
| **Chemicals** |     |           |                  |        |                  |        |                  |        |
| **Fathers** |     |           |                  |        |                  |        |                  |        |
| Unexposed | 1216 | 15.5 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 |
| Exposed pre-conception only | 116 | 15.5 | 1.09 | (0.65; 1.85) | 0.98 | (0.58; 1.66) | 0.98 | (0.55; 1.66) |
| Exposed both pre- and post-conception | 300 | 14.0 | 0.88 | (0.60; 1.31) | 0.85 | (0.57; 1.26) | 0.86 | (0.57; 1.29) |
| **Mothers** |     |           |                  |        |                  |        |                  |        |
| Unexposed | 1445 | 13.0 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 |
| Exposed pre-conception only | 112 | 18.8 | 1.49 | (0.88; 2.54) | 1.20 | (0.69; 2.11) | 1.11 | (0.63; 1.95) |
| Exposed both pre- and post-conception | 520 | 16.3 | 1.26 | (0.94; 1.68) | 1.17 | (0.87; 1.57) | 1.18 | (0.87; 1.61) |

\(^a\)ORModel1 cluster by family, adjusted for study center.
\(^b\)ORModel2 cluster by family, adjusted for study center, offspring’s sex and parent’s characteristic (age, asthma before the age of 10 years, asthma after the age of 10 years, place of upbringing, and smoking) as well as grandmother and grandfathers educational level.
\(^c\)ORModel3 cluster by family, same adjustment as ORModel2 plus parent’s educational level.
\(^d\)The category was included in the model, but due to very few exposed individuals, the result is not presented. Due to data security, i.e. the European Data Protection Regulation, we are not allowed to show results based on microdata (<5 persons). Data are available on request for authorized research units.
Table 3: Multivariable logistic regression analyses clustering by family of paternal or maternal occupational exposure in relation to asthma in offspring; sensitivity analyses of age at asthma onset (0–3 years and 4–15 years)

| Allergens—fathers | Asthma 0–3 years | Asthma 4–15 years |
|-------------------|------------------|------------------|
|                   | n                | Asthma, %        | OR Model1 | 95 % CI | OR Model2 | 95 % CI | OR Model3 | 95 % CI |
| Unexposed         | 1216             | 4                | 1.00      | 1.00    | 1.00      | 1.00    |
| Exposed pre-conception only | 76 | <7 | - | - | - | - |
| Exposed both pre- and post-conception | 235 | 3.8 | 0.93 (0.43; 2.01) | 0.77 (0.34; 1.78) | 0.88 (0.39; 1.99) |
| Exposed both pre- and post-conception | 1216 | 11.1 | 1.00 | 1.00 | 1.00 | 1.00 |
| Exposed pre-conception only | 76 | 6.6 | 0.60 (0.21; 1.78) | 0.59 (0.20; 1.76) | 0.61 (0.21; 1.81) |
| Exposed both pre- and post-conception | 235 | 8.9 | 0.73 (0.43; 1.24) | 0.74 (0.44; 1.26) | 0.79 (0.46; 1.36) |

| Allergens—mothers | Asthma 0–3 years | Asthma 4–15 years |
|-------------------|------------------|------------------|
|                   | n                | Asthma, %        | OR Model1 | 95 % CI | OR Model2 | 95 % CI | OR Model3 | 95 % CI |
| Unexposed         | 1445             | 3.0              | 1.00      | 1.00    | 1.00      | 1.00    |
| Exposed pre-conception only | 118 | <5 | - | - | - | - |
| Exposed both pre- and post-conception | 523 | 5.4 | 1.99 (1.20; 3.30) | 1.70 (1.02; 2.84) | 1.85 (1.08; 3.15) |
| Exposed pre-conception only | 1445 | 10.0 | 1.00 | 1.00 | 1.00 | 1.00 |
| Exposed both pre- and post-conception | 118 | 16.1 | 1.55 (0.89; 2.69) | 1.21 (0.69; 2.14) | 1.07 (0.60; 1.90) |
| Exposed both pre- and post-conception | 523 | 10.9 | 1.07 (0.76; 1.49) | 0.99 (0.70; 1.40) | 0.98 (0.69; 1.40) |

| Chemicals—fathers | Asthma 0–3 years | Asthma 4–15 years |
|-------------------|------------------|------------------|
|                   | n                | Asthma, %        | OR Model1 | 95 % CI | OR Model2 | 95 % CI | OR Model3 | 95 % CI |
| Unexposed         | 1216             | 4.4              | 1.00      | 1.00    | 1.00      | 1.00    |
| Exposed pre-conception only | 116 | 6.9 | 1.90 (0.85; 4.26) | 1.37 (0.57; 3.28) | 1.35 (0.54; 3.36) |
| Exposed both pre- and post-conception | 300 | 3.7 | 0.94 (0.46; 1.89) | 0.77 (0.35; 1.68) | 0.65 (0.29; 1.47) |
| Exposed pre-conception only | 1216 | 11.1 | 1.00 | 1.00 | 1.00 | 1.00 |
| Exposed both pre- and post-conception | 116 | 8.6 | 0.81 (0.41; 1.61) | 0.80 (0.41; 1.55) | 0.75 (0.37; 1.52) |
| Exposed both pre- and post-conception | 300 | 10.3 | 0.87 (0.55; 1.37) | 0.86 (0.55; 1.37) | 0.92 (0.58; 1.47) |

| Chemicals—mothers | Asthma 0–3 years | Asthma 4–15 years |
|-------------------|------------------|------------------|
|                   | n                | Asthma, %        | OR Model1 | 95 % CI | OR Model2 | 95 % CI | OR Model3 | 95 % CI |
| Unexposed         | 1445             | 3.0              | 1.00      | 1.00    | 1.00      | 1.00    |
| Exposed pre-conception only | 112 | <5 | - | - | - | - |
| Exposed both pre- and post-conception | 520 | 5.2 | 1.87 (1.12; 3.13) | 1.65 (0.98; 2.77) | 1.82 (1.06; 3.12) |
| Exposed pre-conception only | 1445 | 10.0 | 1.00 | 1.00 | 1.00 | 1.00 |
| Exposed both pre- and post-conception | 112 | 16.1 | 1.62 (0.93; 2.84) | 1.34 (0.76; 2.36) | 1.18 (0.67; 2.09) |
| Exposed both pre- and post-conception | 520 | 11.2 | 1.09 (0.78; 1.52) | 1.03 (0.73; 1.45) | 1.03 (0.72; 1.46) |

*OR Model1: cluster by family, adjusted for study center.
*OR Model2: cluster by family, adjusted for study center, and parent’s characteristic (age, asthma before the age of 10 years, asthma after the age of 10 years, place of upbringing, and smoking) as well as grandmother and grandfathers educational level.
*OR Model3: cluster by family, same adjustment as OR Model2 + parent’s educational level.
*The category was included in the model, but due to very few exposed individuals, the result is not presented. Due to data security, i.e. the European Data Protection Regulation, we are not allowed to show results based on microdata (<5 persons). Data are available on request for authorized research units.

distinguish between age at asthma onset and included only children up to the age of 7 years.15

Tagiyeva et al. found that maternal antenatal occupational exposure to latex and biocides/fungicides was associated with an increased likelihood of asthma in 7-year-old children.28 However, no effect was seen after adjustment for postnatal exposures, indicating that the main effect could be due to the ‘carry-home’ effect. We wanted to include an exposure window for those only exposed after the birth of the offspring to compare with the window of those exposed both pre- and post-conception to examine possible carry-home effects. Unfortunately, we did not have sufficient statistical power to do this in the post-birth exposure window as we a priori used a cut-off point at age 3 years of the offspring to minimize the risk of offspring developing asthma before the parent was occupationally exposed. We had too few cases to produce firm results for the exposure window pre-conception only for pesticides and...
microorganisms; the results for allergens and chemicals did not reveal any clear results for offspring asthma. Due to data security, i.e. the European Data Protection Regulation, we are not allowed to show results based on microdata (<5 persons). Data are available on request for authorized research units. Magnusson et al. found that some maternal jobs (e.g. engineering technicians, social workers and librarians) during pregnancy were associated with an elevated risk of asthma in their children (14–18 years old).16

In agreement with other studies,29,30 Svanes et al. reported that among parents of 24 168 offspring from the RHINE study,17 there was an increased early onset of offspring asthma if mothers smoked around pregnancy. This is in line with our findings for maternal occupational exposure to reactive chemicals during both pre- and post-conception, with a significant association observed.

Table 4 Multivariable logistic regression analyses clustering by family of paternal or maternal occupational exposure in relation to asthma in their offspring (0–15 years); subgroup analysis by allergic and non-allergic offspring

|                   | n   | Asthma, % | OR Model1a | 95 % CI | OR Model2b | 95 % CI | OR Model3c | 95 % CI |
|-------------------|-----|-----------|------------|---------|------------|---------|------------|---------|
| **Allergens—fathers** |     |           |            |         |            |         |            |         |
| Asthma 0–15 years atopic |     |           |            |         |            |         |            |         |
| Unexposed          | 684 | 20.5      | 1.00       | 1.00    | 1.00       | 1.00    |
| Exposed pre-conception only | 51  | 13.7      | 0.72 (0.30; 1.74) | 0.72 (0.29; 1.78) | 0.74 (0.30; 1.80) |
| Exposed both pre- and post-conception | 129 | 16.3      | 0.70 (0.40; 1.22) | 0.74 (0.42; 1.29) | 0.83 (0.48; 1.45) |
| Asthma 0–15 years non-atopic |     |           |            |         |            |         |            |         |
| Unexposed          | 532 | 9.2       | 1.00       | 1.00    | 1.00       | 1.00    |
| Exposed pre-conception only | -  | 4.0       | -          | -       | -          | -       |
| Exposed both pre- and post-conception | 106 | 8.5    | 0.89 (0.39; 1.99) | 0.79 (0.34; 1.81) | 0.81 (0.36; 1.82) |
| **Allergens—mothers** |     |           |            |         |            |         |            |         |
| Asthma 0–15 years atopic |     |           |            |         |            |         |            |         |
| Unexposed          | 851 | 16.8      | 1.00       | 1.00    | 1.00       | 1.00    |
| Exposed pre-conception only | 66  | 24.2      | 1.48 (0.79; 2.77) | 1.28 (0.69; 2.38) | 1.15 (0.62; 2.15) |
| Exposed both pre- and post-conception | 292 | 21.6      | 1.32 (0.93; 1.87) | 1.18 (0.82; 1.71) | 1.19 (0.82; 1.75) |
| Asthma 0–15 years non-atopic |     |           |            |         |            |         |            |         |
| Unexposed          | 594 | 7.6       | 1.00       | 1.00    | 1.00       | 1.00    |
| Exposed pre-conception only | 52  | 9.6       | 1.29 (0.48; 3.45) | 0.79 (0.27; 2.32) | 0.82 (0.28; 2.40) |
| Exposed both pre- and post-conception | 231 | 9.5    | 1.34 (0.77; 2.35) | 1.21 (0.69; 2.13) | 1.25 (0.70; 2.20) |
| **Chemicals—fathers** |     |           |            |         |            |         |            |         |
| Asthma 0–15 years atopic |     |           |            |         |            |         |            |         |
| Unexposed          | 684 | 20.5      | 1.00       | 1.00    | 1.00       | 1.00    |
| Exposed pre-conception only | 77  | 18.2      | 1.04 (0.58; 1.87) | 1.01 (0.55; 1.85) | 0.98 (0.52; 1.81) |
| Exposed both pre- and post-conception | 175 | 17.7      | 0.82 (0.51; 1.30) | 0.81 (0.50; 1.29) | 0.82 (0.51; 1.33) |
| Asthma 0–15 years non-atopic |     |           |            |         |            |         |            |         |
| Unexposed          | 532 | 9.2       | 1.00       | 1.00    | 1.00       | 1.00    |
| Exposed pre-conception only | -  | 10.3     | -          | -       | -          | -       |
| Exposed both pre- and post-conception | 125 | 8.8    | 0.91 (0.44; 1.92) | 0.80 (0.37; 1.71) | 0.79 (0.36; 1.75) |
| **Chemicals—mothers** |     |           |            |         |            |         |            |         |
| Asthma 0–15 years atopic |     |           |            |         |            |         |            |         |
| Unexposed          | 851 | 16.8      | 1.00       | 1.00    | 1.00       | 1.00    |
| Exposed pre-conception only | 63  | 23.4      | 1.58 (0.85; 2.93) | 1.45 (0.79; 2.67) | 1.31 (0.71; 2.42) |
| Exposed both pre- and post-conception | 291 | 22.0      | 1.35 (0.93; 1.92) | 1.24 (0.86; 1.79) | 1.25 (0.85; 1.83) |
| Asthma 0–15 years non-atopic |     |           |            |         |            |         |            |         |
| Unexposed          | 549 | 8.2       | 1.00       | 1.00    | 1.00       | 1.00    |
| Exposed pre-conception only | 44  | 11.4      | 1.45 (0.54; 3.86) | 0.95 (0.33; 2.79) | 0.94 (0.32; 2.74) |
| Exposed both pre- and post-conception | 208 | 10.1    | 1.24 (0.71; 2.19) | 1.14 (0.65; 2.01) | 1.18 (0.66; 2.09) |

aOR Model1 clust by family, adjusted for study centre.
bOR Model2 clust by family, adjusted for study center, and parent’s characteristic (age, asthma before the age of 10 years, asthma after the age of 10 years, place of upbringing and smoking) as well as grandmother and grandfathers educational level.
cOR Model3 clust by family, same adjustment as OR Model2 + parent’s educational level.
dThe category was included in the model, but due to very few exposed individuals, the result is not presented. Due to data security, i.e. the European Data Protection Regulation, we are not allowed to show results based on microdata (<5 persons); data are available on request for authorized research units.
post-conception periods related to early-onset asthma in offspring. The parents in our study constitute a subpopulation of the larger study population used by Svanes et al. However, we obtained information directly from the offspring of the parents themselves in the RHINESSA study. Similar to Svanes et al.,17 we did not find a clear association for mothers only exposed pre-conception in the main analysis. These findings should be interpreted with caution as we had too few cases to produce firm results for the exposure window pre-conception only in relation to early-onset asthma.

**Paternal occupational exposure and offspring asthma**

We did not find any associations between paternal occupational exposure and offspring asthma. In comparison, Svanes et al.17 suggested that paternal occupational exposure to welding as well as smoking increased the risk of asthma in offspring if exposed to welding pre-conception.

| Table 5 Multivariable logistic regression analyses clustering by family of paternal or maternal occupational exposure in relation to asthma in their offspring (0–15 years); subgroup analysis by offspring sex |
|--------------------------------------------------|--------|---------|---------|---------|---------|---------|
| Allergens—fathers                               | n     | Asthma, % | ORModel1 | 95 % CI | ORModel2 | 95 % CI | ORModel3 |
| Asthma 0–15 years—boys                          |       |          |         |        |         |        |         |
| Unexposed                                        | 519   | 15.4     | 1.00    | 1.00   | 1.00    |        |
| Exposed pre-conception only                      | 39    | 12.8     | 0.90    | (0.34; 2.37) | 0.74 | (0.28; 1.96) | 0.80 | (0.31; 2.04) |
| Exposed both pre- and post-conception            | 102   | 14.7     | 0.94    | (0.48; 1.82) | 0.82 | (0.42; 1.59) | 0.95 | (0.49; 1.84) |
| Asthma 0–15 years—girls                          |       |          |         |        |         |        |         |
| Unexposed                                        | 697   | 15.6     | 1.00    | 1.00   | 1.00    |        |
| Exposed pre-conception only                      | -     | -        | -       | -      | -       | -      |
| Exposed both pre- and post-conception            | 133   | 11.3     | 0.66    | (0.35; 1.23) | 0.67 | (0.35; 1.27) | 0.71 | (0.37; 1.35) |
| Allergens—mothers                                |       |          |         |        |         |        |         |
| Asthma 0–15 years—boys                          |       |          |         |        |         |        |         |
| Unexposed                                        | 614   | 14.0     | 1.00    | 1.00   | 1.00    |        |
| Exposed pre-conception only                      | 51    | 17.6     | 1.36    | (0.63; 2.96) | 1.05 | (0.46; 2.37) | 0.98 | (0.43; 2.22) |
| Exposed both pre- and post-conception            | 224   | 15.2     | 1.15    | (0.75; 1.77) | 1.03 | (0.66; 1.61) | 0.98 | (0.62; 1.56) |
| Asthma 0–15 years—girls                          |       |          |         |        |         |        |         |
| Unexposed                                        | 831   | 12.3     | 1.00    | 1.00   | 1.00    |        |
| Exposed pre-conception only                      | 67    | 17.9     | 1.37    | (0.64; 2.94) | 1.05 | (0.48; 2.31) | 0.94 | (0.42; 2.11) |
| Exposed both pre- and post-conception            | 299   | 17.1     | 1.38    | (0.92; 2.06) | 1.23 | (0.81; 1.86) | 1.29 | (0.84; 1.97) |
| Chemicals—fathers                                |       |          |         |        |         |        |         |
| Asthma 0–15 years—boys                          |       |          |         |        |         |        |         |
| Unexposed                                        | 519   | 15.4     | 1.00    | 1.00   | 1.00    |        |
| Exposed pre-conception only                      | 57    | 14.0     | 0.97    | (0.44; 2.11) | 0.81 | (0.36; 1.78) | 0.83 | (0.37; 1.87) |
| Exposed both pre- and post-conception            | 135   | 13.3     | 0.82    | (0.46; 1.48) | 0.74 | (0.41; 1.33) | 0.76 | (0.42; 1.40) |
| Asthma 0–15 years—girls                          |       |          |         |        |         |        |         |
| Unexposed                                        | 697   | 15.6     | 1.00    | 1.00   | 1.00    |        |
| Exposed pre-conception only                      | 59    | 16.9     | 1.17    | (0.56; 2.43) | 1.11 | (0.54; 2.28) | 1.05 | (0.48; 2.27) |
| Exposed both pre- and post-conception            | 165   | 14.5     | 0.96    | (0.57; 1.61) | 0.94 | (0.55; 1.59) | 0.94 | (0.55; 1.62) |
| Chemicals—mothers                                |       |          |         |        |         |        |         |
| Asthma 0–15 years—boys                          |       |          |         |        |         |        |         |
| Unexposed                                        | 614   | 14.0     | 1.00    | 1.00   | 1.00    |        |
| Exposed pre-conception only                      | 50    | 20.0     | 1.61    | (0.77; 3.36) | 1.27 | (0.59; 2.73) | 1.24 | (0.58; 2.68) |
| Exposed both pre- and post-conception            | 231   | 16.5     | 1.25    | (0.82; 1.90) | 1.15 | (0.74; 1.79) | 1.12 | (0.71; 1.76) |
| Asthma 0–15 years—girls                          |       |          |         |        |         |        |         |
| Unexposed                                        | 831   | 12.3     | 1.00    | 1.00   | 1.00    |        |
| Exposed pre-conception only                      | 62    | 17.7     | 1.38    | (0.63; 3.06) | 1.11 | (0.49; 2.54) | 0.97 | (0.42; 2.22) |
| Exposed both pre- and post-conception            | 289   | 16.3     | 1.28    | (0.86; 1.93) | 1.16 | (0.76; 1.76) | 1.20 | (0.78; 1.87) |

aORModel1 cluster by family, adjusted for study center.
bORModel2 cluster by family, adjusted for study center, and parent’s characteristic (age, asthma before the age of 10 years, asthma after the age of 10 years, place of upbringing and smoking) as well as grandmother and grandfather's educational level.
cORModel3 cluster by family, same adjustment as ORModel2 plus parent’s educational level.
dThe category was included in the model, but due to very few exposed individuals, the result is not presented. Due to data security, i.e. the European Data Protection Regulation, we are not allowed to show results based on microdata (<5 persons). Data are available on request for authorized research units.
The association was found for only non-allergic asthma with an onset before the age of 10 years, suggesting effects on children’s lung function rather than allergy-related outcomes.\textsuperscript{17} Similarly, Accordini \textit{et al}. in a study of 1964 fathers participating in the ECRHS study reported that paternal smoking during early puberty was associated with a higher risk of non-allergic offspring asthma.\textsuperscript{25}

Males are suggested to have three possible vulnerability windows in sperm development: \textit{in utero}, before completion of puberty and post-puberty (reproductive cycle), where the pre-puberty window is suggested to be the most vulnerable.\textsuperscript{17-19} Even though we included parent job history throughout their working life, it is most likely that they entered the labour market after their pre-puberty years. In contrast, the germ cells in women are believed to be more vulnerable during the period around pregnancy.\textsuperscript{17} This could explain why we find an effect of pre- and post-conceptional exposure and early-onset offspring asthma in mothers but not for fathers. Epigenetic research on exposure effects across generations is novel and sparse. Environmental exposures could potentially affect the development of germ cells in a fetus, which could directly affect one or two generations, referred to as intergenerational inheritance.\textsuperscript{31} The intergenerational inheritance of epigenetic information is, though, far from understood.

\textbf{Strengths and limitations}

To our knowledge, this is the first study to investigate the association between multiple parental occupational exposures pre-conception and offspring risk of childhood asthma. An important strength of this study is that information about parental occupational exposure was collected independently and before the offsprings’ report of their asthma status.

Several limitations also need to be considered. Parental job information of the parent who did not participate in the study is lacking. Even if there was a tendency towards couples being more likely to have the same type of job, the associations found in mothers are unlikely to be explained by their partner’s exposure, as we did not observe associations for paternal exposure; however, the true contrast between the sexes of the parents could be larger than observed.

Even though the study design was prospective, questionnaire data were collected retrospectively and recall bias could have been introduced. Parents’ total employment history could thus be incomplete but it is unlikely that this would be different between exposed and unexposed parents. The use of the OAsJEM will leave out the risk of recall bias concerning exposure to specific agents at their jobs, as these are based on expert evaluations. However, the OAsJEM introduces non-differential misclassifications, as the job titles only constitute a proxy for the exposure to specific agent exposures. It is, therefore, uncertain whether the individual parents were exposed to the agents for which they were chosen. Hence, parents could have been exposed to agents other than those included in the OAsJEM. Even though we excluded all individuals exposed to any of the 30 other occupational asthma-specific agents from the reference group, other (non-occupational) exposures could have constituted a risk factor for the development of asthma, which may have contaminated the reference group data. However, it is unlikely that the findings are due to exposure misclassification of the JEMs. JEM-based exposure assessment is mainly affected by the Berkson error, which results in nearly unbiased effect estimates at the expense of loss of statistical power.\textsuperscript{32} We did not include measurements of air pollution, which is suspected to be a risk factor for childhood asthma\textsuperscript{33} and this might have influenced the associations. However, adjustment for various possible confounders including place of upbringing, parental smoking as well as parental and grandparents’ highest educational level did not substantially alter the associations. However, we cannot rule out residual confounding. We used complete case analysis to deal with missing data, which potentially could have under- or over-estimated the effect.\textsuperscript{34}

Asthma is a heterogenous disorder with multiple phenotypes where age at asthma onset may define some of the heterogeneity. We included offspring with asthma onset at 0–15 years of age, which was further divided into onset between 0–3 years (early-onset asthma) and 4–15 years (late-onset asthma) to account for possible different underlying mechanisms for different phenotypes.\textsuperscript{35} As asthma and age at asthma onset were assessed based on adult offsprings’ self-report, recall bias could have led to misclassification. Svanes \textit{et al}.\textsuperscript{17} found smoking and welding pre-conception to be associated with an increased risk of only non-allergic early-onset asthma. Due to the few available cases, it was not feasible to stratify the early-onset groups of asthma into atopic status or sex of the offspring. We cannot rule out that having allergic offspring in the analysis would have biased the association towards the null. However, no clear associations were seen in the main analyses of childhood asthma between 0 and 15 years of age, and similar results were seen in the main analysis among atopic and non-atopic children. As an isolated finding, maternal occupational exposure to allergens and reactive chemicals during the pre- and post-conception period was associated with increased odds of early-onset offspring asthma. Although this finding is not supported by the other analyses and could be a result of multiple comparisons, it holds
some biological plausibility and could be considered as hypothesis-generating to be further investigated in animal or observational studies. Current results should be interpreted and generalized with caution until further studies are conducted.

**Conclusion**

In conclusion, we did not find evidence to support the impact of only pre-conception or both pre-and post-conception parental occupational exposure measured by JEMs on childhood asthma in offspring in this study.

**Ethics approval and consent to participate**

The study was approved by regional committees of medical research ethics in each study centre according to national legislation. Furthermore, informed consent was obtained from each participant in all study centres. For a list of ethics committees and numbers for each centre, please see www.rhinessa.net.

**Availability of data and material**

The dataset is held and managed by the RHINESSA coordinating centre at the Department of Occupational Medicine, Haukeland University Hospital, Bergen, Norway. Data cannot be made freely available as they are subject to Norwegian Data Protection regulations, but de-identified data can be made available to researchers upon request. Requests for data can be sent to the principal investigator of RHINESSA: C.S., Cecilie.Svanes@med.uib.no.

**Supplementary Data**

Supplementary data are available at IJE online

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**Author Contributions**

K.P.: principal author, conception and design of the work, statistical analysis, interpretation of data, drafting the manuscript. All other authors: conception and design of the work, acquisition and interpretation of data and revision of the manuscript for important intellectual content. All authors approved the final version.

**Conflict of Interest**

None declared.

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