National Institute for Health and Care Excellence

FINAL

Indoor air quality at home

[2] Evidence review for exposure to pollutants and health outcomes

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Evidence review
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Final

These evidence reviews were developed by Public Health Internal Guideline Development team



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Exposure to pollutants and health outcomes

Review question

What signs and symptoms should prompt healthcare professionals to consider exposure to poor indoor air quality at home in people presenting to health services?

Introduction

People spend up to 90% of their lives indoors and 60% of that time at home. To minimize the health risks from pollutants occurring in homes, exposures to these pollutants should be controlled. The priority in this is to control the source of the pollutant and so reduce exposure. Often, especially in existing buildings, this may be difficult to achieve, in which case pollutant exposures should be controlled by providing enough ventilation air to dilute and remove the contaminants.

Recent reviews suggest that adequate ventilation results in more than 0.4 air changes per hour (Wargocki 2013) and that home ventilation ratios greater than 0.5 air changes per hour was associated with better health outcomes (Sundell 2010).

The aims of this review are to Identify clinical signs and symptoms that are associated with exposure to poor indoor air quality at home.

PICO table

Table 1 outlines the PICO elements of the review protocol which are available in Review protocol in Appendix A:

Table 1: PICO table for signs and symptoms

Eligibility criteria	Content
Population	People in all dwellings
Prognostic factors	 Clinical signs / symptoms associated with exposure to indoor air pollutants at home including:
	 Neurological symptoms for example: headache, drowsiness, fatigue, poor concentration, confusion
	 Respiratory symptoms for example: coughing, sneezing, wheezing, sinus congestion, phlegm, sore throat, nasal congestion, runny nose
	• Cardiovascular symptoms for example chest pain, shortness of breath
	Nausea
	Eye irritation
	 Signs and symptoms of immune response disorder for example asthma, allergic rhinitis, dermatological conditions for example atopic eczema, psoriasis

Eligibility criteria	Content
	 Pregnancy related for example low birth weight for gestational age, premature birth, infant mortality (but not sudden infant death (SID)), stillbirth
Outcomes	Risk ratios, odds ratios of exposure to indoor air pollutants at home

Methods and process

This evidence review was developed using the methods and process described in Developing NICE guidelines: the manual. Methods specific to this review question are described in the review protocol in Appendix A:.

As review questions 1 (individual and building factors associated with exposure to poor indoor air quality at home) and 2 (signs and symptoms should prompt healthcare professionals to consider exposure to poor indoor air quality at home) overlapped, both reviews were carried out using a single search. The results of this search were then parsed as follows

- Studies that examined the association between individual and building characteristics and health outcomes
- Studies that examined the association between sources of pollutants and health outcomes
- Studies that examined the association between exposure levels and health outcomes.

This review is concerned with the association between sources of pollutants or exposure and health outcomes. Please see Evidence review 1 for the association between individual or building characteristics and exposure levels.

Declarations of interest were recorded according to NICE's 2018 conflicts of interest policy.

Public health evidence

Included studies

33967 references were identified from literature searches outlined in Appendix B. 426 papers were ordered in full-text for questions 1 and 2. Of these 2878 were excluded from both reviews and 148 articles in total were included in the two reviews. 99 studies from 101 articles and 31 studies from 32 articles were included in the two parts of this review respectively and 15 studies from 16 papers were included in evidence review 1.

Excluded studies

The full list of excluded studies and reasons for exclusion are in Appendix I:

Quality assessment of studies included in the evidence review

For this review question, cohort studies were considered to be of highest quality and case control studies as next best evidence quality. Evidence quality started as 'high' for cohort studies and 'low' for case control studies.

See Appendix F: for the full GRADE tables.

Association between individual and building characteristics and health outcomes

Summary of public health studies included in the evidence review

A summary of the characteristics of the included studies are in the following table

Table 2: Characteristics of included studies

Study	(country)	Design	Population	Characteristic	Outcomes	Risk of bias
1.	Bajeux 2014 (France)	Prospective cohort	Pregnant women	Domestic products	Wheeze, Eczema, Food allergies	High
2.	Baker 2006 (US)	Prospective cohort	Children	Heating fuel	Lower respiratory illness	Low
3.	Bedard 2014 (France)	Nested case-control	Women with asthma	Domestic products	Asthma,	Moderate
4.	Belanger 2003 (US)	Prospective cohort	Infants at risk of asthma, Infants not at risk of asthma	Gas stove, House dust, Pets, Mould / mildew	Wheeze, Persistent cough	Moderate
5.	Belanger 2006 (US)	Prospective cohort	Children with asthma	Gas stove, Gas dryer	Wheeze, Persistent cough, Shortness of breath, Chest tightness	Moderate
6.	Bhinder 2014 (Canada)	Retrospective cohort	Adult lung transplant recipients	Proximity to traffic	Chronic lung allograft dysfunction	Moderate:
7.	Bornehag 2005 (Sweden)	Nested case-control	Children with respiratory symptoms	Ventilation rate	Asthma and allergic symptoms	Moderate
8.	Bowatte 2017 (Australia)	Prospective cohort	Adults	Proximity to traffic	Asthma, Wheeze	Low
9.	Brunekreef 1989 (US)	Prospective cohort	Children	Damp / mould	Wheeze, Cough, Bronchitis, Chest illness, Lower respiratory illness, Asthma, Hay fever, Non-chest illness	High
10.	. Cable 2014 (UK)	Prospective cohort	Adults	Damp	Cough, Phlegm	High
11.	Carlsten 2010 (Canada)	Prospective cohort	Children at risk of asthma	House dust, Pets	Asthma	Low
12.	Casas 2012 (Germany)	Prospective cohort	Children	Damp, Gas stove, Pets	Asthma, Persistent wheeze	High

FINAL Association between individual and building characteristics and health outcomes

Study (country)	Design	Population	Characteristic	Outcomes	Risk of bias
13. Casas 2013 (Spain)	Prospective cohort	Children	Domestic products	Wheezing	Moderate
14. Chang 2009 (US)	Retrospective cohort	Children with asthma	Proximity to traffic	Asthma exacerbations	Low
15. Clarke 2015 (US)	Prospective cohort	Women	Pets	Thyroid cancer	Moderate
16. de Bilderling 2005 (United Kingdom)	Prospective cohort	Children & adolescents	Gas stove, Gas oven, Gas heating	Wheezing	High
17. Diez 2002 (Germany)	Nested case control	Infants at risk of allergies	Restoration work, Pets	Pulmonary infection, Wheeze	Moderate
18. Diez 2003 (Germany)	Prospective cohort	Children at risk of asthma	Redecoration,	Obstructive bronchitis, Wheeze	High
19. Du Prel 2006 (Germany)	Prospective cohort	Children	Damp, Heating	Bronchitis, Respiratory symptoms, Eczema	Low
20. Emenius 2003 (Sweden)	Nested case control	Children	Building age	Recurrent wheezing	Moderate
21. Emenius 2004 (Sweden)	Nested case control	Children	Redecoration, Damp	Recurrent wheezing	Moderate
22. Engvall 2001 (Sweden)	Retrospective cohort	Adults	Damp, Humidity, History of water leaks	Eye irritation, Nasal irritation, Throat irritation, Cough, facial skin symptoms, headache, Tiredness	High
23. Engvall 2010 (Sweden)	Retrospective cohort	Adults	Tenancy status	Eye irritation, Nasal irritation, Throat irritation, Cough	Moderate
24. Farrow 2003 (UK)	Prospective cohort	Women and children	Domestic products	Diarrhoea, Vomiting, Earache	Moderate
25. Franck 2014 (Germany)	Cohort	Mother baby dyads	Redecoration	Recurrent wheeze, Obstructive bronchitis	Low
26. Gan 2010 (Canada)	Prospective cohort	Adults	Proximity to traffic	CHD mortality	Low

FINAL Association between individual and building characteristics and health outcomes

Study (country)	Design	Population	Characteristic	Outcomes	Risk of bias
27. Garshick 2003 (United States)	Prospective cohort	Adults	Proximity to traffic	Wheeze, Cough, Phlegm	Moderate
28. Habre 2014 (United States)	Prospective cohort	Children with asthma	Particulate matter	Wheeze, Cough	Moderate
29. Hagerhed- Engman 2009 (Sweden)	Nested case control	Children	Damp / mould	Asthma, Eczema, Rhinitis	Low
30. Hagmolen of Ten Have 2007 (The Netherlands)	Prospective cohort	Children with asthma	Damp, Pets	Airway hyper- responsiveness	High
31. Hart 2014 (US)	Prospective cohort	Women	Proximity to traffic	Sudden cardiac death	Low
32. Harville 2018 (UK)	Prospective cohort	Women who had given birth	Mould	Low birth weight, preterm birth, small for gestational age	Moderate
33. Heinrich 2013 (Germany)	Prospective cohort	Women	Proximity to traffic	Mortality	Low
34. Henderson 2008 (UK)	Prospective cohort	Pregnant women	Domestic products	Wheeze	High
35. Herr 2012 (France)	Prospective cohort	Infants	Pets, Redecoration, House dust, Cleaning prays	Wheeze	Low
36. Hjortebjerg 2012 (Denmark)	Prospective cohort	Pregnant women	Redecoration	Congenital malformations	Low
37. Hoffmann 2007 (Germany)	Prospective cohort	Adults	Proximity to traffic	Coronary artery calcification	Low
38. Hunt 2011 (US)	Prospective cohort	Infants at risk of asthma	Particulate matter	Wheeze	Low
39. Ibargoyen- Roteta 2007 (Spain)	Prospective cohort	Children	Glazing, Damp / mould	Allergic Rhino- conjunctivitis	High
40. Jaakkola 2005 (Finland)	Prospective cohort	Children	Damp, Mould	Asthma	High

FINAL Association between individual and building characteristics and health outcomes

Study (country)	Design	Population	Characteristic	Outcomes	Risk of bias
41. Jaakkola 2010 (Finland)	Prospective cohort	Children	Damp / mould	Allergic rhinitis	High
42. Jedrychowski 2010 (Poland)	Prospective cohort	Children with intrauterine exposure	PAHs, Particulate matter	Wheeze	Moderate
43. Jedrychowski 2011 (Poland)	Prospective cohort	Infants	Damp / mould, Particulate matter	Eczema	Moderate
44. Jedrychowski 2014 (Poland)	Prospective cohort	Children with intrauterine exposure	PAHs	Wheeze	Moderate
45. Jung 2012 (US)	Prospective cohort	Children	Particulate matter	Wheeze	Moderate
46. Karvonen 2015 (Finland)	Prospective cohort	Children	Damp / mould	Asthma, cough, wheeze	Low
47. Kingsley 2015 (US)	Prospective cohort	Women	Proximity to traffic	Incident hypertension	Low
48. Koloski 2015 (Australia)	Prospective cohort	Adults	Pets	Irritable bowel syndrome, Functional dyspepsia	Low
49. Korppi 2008 (Finland)	Cohort	Children at risk of asthma	Pets,	Asthma	Moderate
50. Larsson 2009	Cohort	Children	Flooring, Damp	Autistic spectrum disorders	Moderate
51. Larsson 2010 (Sweden)	Cohort	Children	Flooring	Asthma	Low
52. Le Moual 2012 (France)	Prospective cohort	Women with asthma	Domestic products	Asthma	High
53. Li 2006 (US)	Prospective cohort	Children	Gas stove, Heating, Housing area	Lower respiratory tract symptoms	High
54. Li 2016 (US)	Prospective cohort	Adults	Proximity to traffic	Obesity	Low
55. Lindgren 2013 (Sweden)	Prospective cohort	Children	Proximity to traffic	Asthma, Bronchiolitis, Obstructive bronchitis	Low
56. Litonjua 2002 (US)	Prospective cohort	Children at risk of atopy	Pets	Wheeze	Moderate

FINAL Association between individual and building characteristics and health outcomes

Study (country)	Design	Population	Characteristic	Outcomes	Risk of bias
57. Lynch 2014 (US)	Prospective cohort	Children at risk of atopy	Pets, House dust	Wheeze	Moderate
58. Mahalingaiah 2014 (US)	Prospective cohort	Women	Proximity to traffic	Uterine leiomyomata	Low
59. Mahalingaiah 2016 (US)	Prospective cohort	Women	Proximity to traffic	Infertility	Low
60. McConnell 2002 (US)	Prospective cohort	Children	Gas stove, Pets	Asthma, wheeze	Low
61. McConnell 2006 (US)	Prospective cohort	Children	Proximity to traffic	Asthma, wheeze	Low
62. Mommers 200 (Germany and the Netherlands)	Nested case control	Children	Gas stove, Heating, SES	Asthma symptoms, Cough	High
63. Morgernstern 2007 (Germany)	Prospective cohort	Children	Proximity to traffic	Wheeze, Cough, Bronchitis, Respiratory infections, Nasal symptoms	Moderate
64. Morgernstern 2008 (Germany)	Prospective cohort	Children	Proximity to traffic	Asthma, Hay fever, Eczema	Moderate
65. Nenna 2017 (Italy)	Case-control	Infants with bronchiolitis	Occupancy, Cooking oil	Bronchiolitis	High
66. Norback 2013 (Europe, Australia, US)	Prospective cohort	Adults	Damp / mould	Asthma, Bronchial hyper-responsiveness	High
67. Ostro 1993 (US)	Prospective cohort	Adults	Gas stove	Respiratory illness	Low
68. Pettigrew 2004 (US)	Prospective cohort	Infants at risk of asthma	Mould	Otitis media	Moderate
69. Pettigrew 2004 b (US)	Prospective cohort	Infants	Heating, Mould, Pets, Air conditioning	Otitis media	Low
70. Pindus 2016 (Estonia)	Prospective cohort	Not specified	Particulate matter	Cough, Wheeze, Asthma, Allergic	Moderate

FINAL Association between individual and building characteristics and health outcomes

Study (country)	Design	Population	Characteristic	Outcomes	Risk of bias
				rhinitis, Breathlessness, Chest tightness, Cardiac disease, Stroke, Hypertension, Heart infarction or angina pectoris	
71. Ponsonby 2001 (Australia)	Prospective cohort	Children	Gas appliances, House dust	Asthma	Moderate
72. Power 2015 (US)	Prospective cohort	Women	Proximity to traffic	Anxiety	Low
73. Puett 2014 (US)	Prospective cohort	Women	Proximity to traffic	Lung cancer	Low
74. Pujades- Rodriguez 2009 (UK)	Prospective cohort	Adults	Proximity to traffic	Wheeze, COPD, Bronchial hyper- responsiveness, Allergic sensitisation	Moderate
75. Reponen 2011 (US)	Prospective cohort	Children	Mould, Air conditioning, House dust	Asthma	Low
76. Rice 2015 (US)	Prospective cohort	Adults	Proximity to traffic	Asthma, Obstruction, Wheeze, Cough	Moderate
77. Roda 2011 (France)	Prospective cohort	Infants	Formaldehyde, Pets,	Lower respiratory infection, Lower respiratory infection with wheeze	Moderate
78. Samet 1993 (US	Prospective cohort	Infants	Gas stove	Respiratory illness, Cough, Wheeze	Low
79. Sbihi 2016 (Canada)	Prospective cohort	Children	Proximity to traffic	Asthma	Low
80. Sherriff 2005 (UK)	Prospective cohort	Pregnant women	Domestic products	Wheeze	High
81. Shmuel 2017 (US and Puerto Rico)	Prospective cohort	Adults	Proximity to traffic	Breast cancer	Moderate
82. Shu 2013 (Sweden)	Prospective cohort	Children	Flooring	Asthma	Moderate

FINAL Association between individual and building characteristics and health outcomes

Study (country)	Design	Population	Characteristic	Outcomes	Risk of bias
83. Sorensen 2010 (Denmark)	Prospective cohort	Pregnant women	Redecoration	Preterm birth, small for gestational age	Low
84. Stark 2005 (US)	Prospective cohort	Children at risk of asthma or allergy	Mould	Allergic rhinitis	Low
85. Thacher 2017 (Sweden)	Prospective cohort	Children	Damp, Mould	Asthma, Rhinitis	Low
86. Tiesler 2015 (Germany)	Prospective cohort	Children	Damp / mould	Sleep problems	High
87. Tin Tin 2016	Prospective cohort	Pregnant women,	Occupancy, Tenancy status, Heating, Mould	Acute respiratory infections	Moderate
88. Triche 2002 US)	Prospective cohort	Infants	Heating	Wheeze, Cough	Low
89. Triche 2005 (US)	Prospective cohort	Women with an infant child	Gas heating	Wheeze, Chest tightness	Moderate
90. Virtanen 2014 (Finland)	Prospective cohort	Children	Pets	Type 1 diabetes	Moderate
91. Weinmann 2017 (Germany)	Prospective cohort	Adults	Domestic products	Asthma, Wheeze	High
92. Weinmayr 2015 (Germany)	Prospective cohort	Adults	Proximity to traffic	Type 2 diabetes	Low
93. Wesselink 2017 (US)	Retrospective cohort	Pregnant women	Proximity to traffic	Pre-eclampsia, Placental abruption, Small for gestational age, Stillbirth	Moderate
94. White 2017 (US and Puerto Rico)	Prospective cohort	Women at risk of breast cancer	Heating, Cooking	Breast cancer	Low
95. Willers 2006 (the Netherlands)	Prospective cohort	Children at risk of asthma	Gas stove	Asthma, Wheeze, Nasal symptoms, Eczema	Moderate
96. Zhang 2016 (US)	Prospective cohort	Women	Proximity to traffic	Hypertension	Low

FINAL Association between individual and building characteristics and health outcomes

Study (country)	Design	Population	Characteristic	Outcomes	Risk of bias
97. Zhou 2013 (France)	Prospective cohort	Mother-child pairs,	Pets, Proximity to traffic, Heating, Damp	Asthma, Wheeze, Bronchiolitis	Moderate
98. Zock 2007 (10 European Countries)	Prospective cohort	Adults	Domestic products	Asthma, Wheeze. Nocturnal shortness of breath	High

See Appendix D:for full evidence tables.

Quality assessment of public health studies included in the evidence review

See appendix F for full GRADE tables.

Economic evidence

No economic evidence review was carried out for this review

Economic model

No economic modelling was carried out for this review

Evidence statements

Sources of NO₂

Gas heating (Grade F.1.1.1)

- This evidence review found low quality evidence from 1 study of 2,898 infants at risk of allergy that that gas heating was not associated with cough aOR 0.78 (95%CI 0.56 to 1.09) and infants not at risk of allergy aOR 1.01 (95%CI 0.69 to 1.46)
- This evidence review found low quality evidence from 1 study of 1,868 children showing that gas central heating was not associated with wheeze aOR 0.76 (95%CI 0.47 to 1.23)
- This evidence review found low quality evidence from 1 study of 1,868 children showing that use of a gas fire was not associated with wheeze aOR 0.97 (95%CI 0.67 to 1.39)
- This evidence review found high quality evidence for gas as heating fuel from 1 study with 50,884 women at risk of breast cancer showing that gas heating was associated with breast cancer aHR 1.15 (95%CI 1.00 to 1.32)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Gas space heater (Grade F.1.1.2)

- This evidence review found high quality evidence from 1 study with 890 infants showing that the use of a gas space heater was associated with wheeze aRR 1.25 (95%CI 1.05 to 1.50)
- This evidence review found moderate quality evidence from 1 study with 888 mothers of infants showing that use of a gas space heater was not associated with
 - o cough aRR 1.00 (95%CI 0.97 to 1.04),

- o wheeze aRR 1.03 (95%CI 0.94 to 1.13),
- o phlegm aRR 0.96 (95%CI 0.88 to 1.05),
- o runny / stuffy nose aRR 0.99 (95%CI 0.95 to 1.03),
- sore throat aRR 0.99 (95%CI 0.95 to 1.04)
- o laryngitis aRR 0.93 (95%CI 0.79 to 1.10).
- This evidence review found high quality evidence from 1 study with 890 infants showing that the use of a gas space heater was not associated with cough aRR 0.94 (95%CI 0.75 to 1.18)
- This evidence review found moderate quality evidence from 1 study with 3,535 children with no history of asthma showing that use of a gas space heater was not associated with asthma with wheeze aOR 1.20 (95%CI 0.70, 2.00)
- This evidence review found moderate quality evidence from 1 study of 888 mothers of infants showing that use of a gas space heater was not associated with chest tightness aRR 1.01 (95%CI 0.96 to 1.07)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - People with disabilities
 - Pregnant women
 - People with conditions associated with or exacerbated by indoor air pollution.

Gas for cooking (Grade F.1.1.3)

- This evidence review found high quality evidence from 1 study with 321 adults showing that the use of a gas stove was associated with lower respiratory tract infections aOR 1.23 1.03 to 1.47) but not with upper respiratory tract infections aOR 1.06 (95%CI 0.94 to 1.18)
- This evidence review found moderate quality evidence from 1 study of 728 children with asthma in multi-family housing showing that the use of a gas stove was associated with
 - o shortness of breath aOR 2.38 (95%CI 1.12 to 5.06) and
 - o chest tightness aOR 4.34 (95%CI 1.76 to 10.69)
- This evidence review found moderate quality evidence from 1 study with 3,148 children at risk of atopy showing that the use of a gas stove was associated with nasal symptoms aOR 1.34 (95%CI 1.06 to 1.71)
- This evidence review found low quality evidence from 1 study with 849 children showing
 that use of a gas stove was associated with cough aOR 1.52 (95%CI 1.06 to 2.18) but not
 with wheeze aOR 1.28 (95%CI 0.88 to 1.86) for those children whose mother did not have
 asthma or for wheeze or cough for children whose mother had asthma aOR 1.03 (95%CI
 0.59 to 1.79) and aOR 0.79 (95%CI 0.46 to 1.36) respectively
- This evidence review found high quality evidence from 1 study with 1,205 infants showing that use of a gas stove or gas for cooking was not associated with
 - respiratory illness aOR 0.98 (95%CI 0.90 to 1.07),
 - wheeze aOR 0.84 (95%CI 0.64 to 1.09)
 - o cough aOR 0.94 (95%CI 0.82 to 1.07)

- This evidence review found moderate quality evidence from 1 study with 5078 children showing that the use of gas for cooking was not associated with wheeze aOR 1.09 (95%CI 0.76 to 1.57)
- This evidence review found moderate quality evidence from 1 study with 1868 children and adolescents showing that any use of gas for cooking was not associated with wheeze aOR 1.02 (95%CI 0.77 to 1.36)
- This evidence review found low quality evidence from 1 study of 728 children with asthma
 in single-family housing showing that use of a gas stove or gas for cooking was not
 associated with
 - shortness of breath aOR 0.91 (95%CI 0.50 to 1.64)
 - chest tightness aOR 0.68 (95%CI 0.34 to 1.32)
- This evidence review found moderate quality evidence from 2 studies that use of a gas stove or gas for cooking was not associated with asthma aOR 1.3 (95%CI 0.80 to 2.00) for 1 study with 3535 children and aOR 1.33 (95%CI 0.88 to 2.00) for the second study with 5078 children.
- This evidence review found low quality evidence from 1 study with 3,148 children at risk of atopy showing that use of a gas stove or gas for cooking was not associated with eczema aOR 0.97 (95%CI 0.74 to 1.26)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - People with disabilities
 - Pregnant women

Other gas appliance (F.1.1.4)

- This evidence review found very low quality evidence from 1 study with 1,191 children showing that the use of an unvented gas geyser for water heating was not associated with cough aOR 1.74 (95%CI 0.74 to 4.12)
- This evidence review found very low quality evidence from 1 study with 1,191 children showing that the use of a vented gas geyser for water heating was not associated with wheeze aOR 1.28 (95%CI 0.85 to 1.94)
- This evidence review found low quality evidence from 1 study with 728 children in multifamily housing showing that the use of a gas dryer was not associated with
 - shortness of breath aOR 2.39 (95%CI 0.77 to 7.43)
 - o chest tightness aOR 1.09 (95%CI 0.31 to 3.90)
- This evidence review found low quality evidence from 1 study with 728 children in singlefamily housing showing that the use of a gas dryer was not associated with
 - shortness of breath aOR 0.91 (95%CI 0.50 to 1.64)
 - o chest tightness aOR 1.41 (95%CI 0.61 to 3.26)
- This evidence review found low quality evidence from 1 study with 456 children showing that the use of home gas appliances (appliances were not specified) was not associated with asthma aOR 1.30 (95%CI 0.74 to 2.29)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas

- o Older people
- o People with disabilities
- Pregnant women
- People with conditions associated with or exacerbated by indoor air pollution

Sources of particulate matter

Fireplace (Grade F.1.2.1)

- This evidence review found high quality evidence from 1 study of 888 mothers of infants showing that the use of a fireplace for heating was associated with
 - o cough aOR 1.05 (95%CI 1.01 to 1.09)
 - o sore throat aOR 1.04 (95%CI 1.00 to 1.08)
- This evidence review found moderate quality evidence from 1 study of 890 infants showing that the use of a fireplace was not associated with
 - o cough aOR 0.99 (95%CI 0.81 to 1.21)
 - o wheeze aOR 0.25 (95%CI 0.04 to 1.43),
- This evidence review found low quality evidence from 1 study of 905 adults showing that the use of wood as heating fuel was not associated with
 - wheeze without cold aOR 1.14 (95%CI 0.75 to 1.73)
 - hay fever (reported as allergic rhinitis) aOR 0.63 (95%Cl 0.42 to 0.94)
 - o breathlessness aOR 0.97 (95%CI 0.64 to 1.48),
 - o chest tightness aOR 1.05 (95%CI 0.72 to 1.51),
 - cardiac disease aOR 0.92 (95%CI 0.60 to 1.39)
 - hypertension aOR 0.78 (95%CI 0.54 to 1.12),
 - o stroke aOR 0.85 (95%CI 0.27 to 2.71),
 - o heart infarction or angina pectoris aOR 0.67 (95%CI 0.28 to 1.56)
- This evidence review found high quality evidence from 1 study of 50884 women at risk of breast cancer showing that the use of a fireplace for heating was associated with breast cancer aHR 1.11 (95%CI 1.01 to 1.22)
- This evidence review found moderate quality evidence from 1 study of 50884 women at risk of breast cancer showing that the use of wood as heating fuel was not associated with breast cancer aHR 1.09 (95%CI 0.98 to 1.21)
- This evidence review found moderate quality evidence from 1 study of 50884 women at risk of breast cancer showing that the use of wood as the main source of heating fuel was not associated with breast cancer aHR 1.09 (95%CI 0.82 to 1.45)
- This evidence review found moderate quality evidence from 1 study with 813 infants showing that the use of a fireplace was not associated with
 - o any episodes of earache aOR 1.14 (95%Cl 0.90 to 1.45)
 - recurrent earache (four or more episodes separated by 21 days in 1 year) aOR 0.99 (95%CI 0.58 to 1.72)
- This evidence review found moderate quality evidence from 1 study of 888 mothers of infants showing that the use of a fireplace was not associated with
 - wheeze aOR 1.07 (95%CI 0.97 to 1.18),

- o laryngitis aOR 1.02 (95%CI 0.94 to 1.10),
- o phlegm aOR 1.04 (95%CI 0.99 to 1.09)
- runny / stuffy nose aOR 0.99 (95%CI 0.95 to 1.04)
- o chest tightness aOR 1.05 (95%CI 0.99 to 1.12)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - People with conditions associated with or exacerbated by indoor air pollution

Wood stove (Grade F.1.2.2)

- This evidence found moderate quality evidence from 1 study with 813 infants showing that the use of wood stoves was not associated with
 - o any episodes of earache aOR 1.22 (95%CI 0.66 to 2.23)
 - recurrent earache (four or more episodes separated by 21 days in 1 year) aOR 1.08 (95%CI 0.85 to 1.38)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - o People with disabilities
 - o Pregnant women
 - People with conditions associated with or exacerbated by indoor air pollution

Heating or cooking fuel (Grade F.1.2.3)

- This evidence review found high quality evidence from 1 study with 28,888 children showing that the use of coal, wood, gas or oil as heating fuel was associated with
 - bronchitis (ever diagnosed) aOR 1.15 (95%Cl 1.00 to 1.32) for homes in West Germany
 - more than 4 colds in past 12 months aOR 1.13 (95%Cl 1.03 to 1.23) for homes in East Germany
- This evidence review found moderate quality evidence from the same study found the use of coal, wood, gas or oil as heating fuel was not associated with
 - o frequent cough aOR 0.97 (95%CI 0.86 to 1.10) for homes in East Germany
 - o frequent cough aOR 0.88 (95%CI 0.68 to 1.15) in homes in West Germany,
 - sneeze attacks in past 12 months aOR 0.92 (95%Cl 0.80 to 1.06) for homes in East Germany and
 - sneeze attacks in past 12 months aOR 1.21 (95%Cl 0.88 to 1.66) for homes in West Germany,
 - o bronchitis (ever diagnosed) aOR 1.02 (95%CI 0.96 to 1.09) for homes in East Germany
 - more than 4 colds in past 12 months aOR 0.96 (95%Cl 0.79 to 1.18) for homes in West Germany.
 - Allergy (ever diagnosed) aOR 1.07 (95%CI 0.96 to 1.18) for homes in East Germany

- o Allergy (ever diagnosed) aOR 0.97 (95%CI 0.79 to 1.19) in West Germany
- Eczema (ever diagnosed) aOR 0.90 (95%CI 0.83 to 0.98) for homes in East Germany
- Eczema (ever diagnosed) aOR 1.07 (95%CI 0.87 to 1.32) in West Germany
- Overweight aOR 0.89 (95%Cl 0.78 to 1.01) for homes in East Germany
- o Overweight aOR 1.12 (95%CI 0.86 to 1.47) in West Germany
- This evidence review found moderate quality evidence for wood as heating fuel from 1 study with 50,884 women at risk of breast cancer showing that wood was not associated with breast cancer aHR 1.09 (95%CI 0.98 to 1.32)

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- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - People with conditions associated with or exacerbated by indoor air pollution

Coal heating (Grade F.1.2.4)

- This evidence review found high quality evidence from 1 study of 452 children showing that the use of coal for heating was associated with lower respiratory tract infections -aOR 1.45 (95%CI 1.07 to 1.97).
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Artificial logs (Grade F.1.2.5)

- This evidence review found moderate quality evidence from 1 study with 50,884 women at risk of breast cancer that the use of artificial logs for heating was not associated with breast cancer aHR 0.98 (95%CI 0.85 to 1.12)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - o Pregnant women
 - Children and young people
 - People with conditions associated with or exacerbated by indoor air pollution

Fuel oil (Grade F.1.2.6)

• This evidence review found moderate quality evidence from 1 study with 50,884 women at risk of breast cancer that the use of fuel oil for heating was not associated with breast cancer aHR 1.13 (95%CI 0.97 to 1.32)

- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - o Older people
 - o People with disabilities
 - Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Cooking oil (Grade F.1.2.7)

- This evidence review found very low quality evidence from 1 study with 416 infants showing that the use of seed oil for cooking was associated with bronchiolitis aOR 1.82 (95%CI 1.21 to 2.74)
- · No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - o Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Paraffin (Kerosene) heating (Grade F.1.2.8)

- This evidence review found high quality evidence from 1 study with 888 mothers of infants that the use of a paraffin heater was associated with wheeze aOR 1.06 (95%CI 1.01 to 1.11)
- This evidence review found moderate quality evidence from 1 study with 888 mothers of infants showing that the use of a paraffin heater was not associated with
 - o cough aOR 1.01 (95%CI 0.99 to 1.03),
 - o laryngitis aOR 1.01 (95%CI 0.97 to 1.04),
 - phlegm aOR 0.98 (95%CI 0.93 to 1.03),
 - o runny / stuffy nose aOR 1.01 (95%CI 0.99 to 1.03)
 - sore throat aOR 1.00 (95%CI 0.97 to 1.02)
 - o chest tightness aOR 1.02 (95%CI 0.99 to 1.05)
- This evidence review found moderate quality evidence from 1 study with 890 infants showing that the use of a paraffin heater was not associated with
 - o cough aOR 1.01 (95%CI.93 to 1.10),
 - wheeze aOR 0.90 (95%CI 0.64 to 1.25)
- This evidence review found very low quality evidence from 1 study with 1,137 children showing that the use of paraffin heaters was not associated with lower respiratory tract infections aOR 1.41 (95%CI 0.96 to 2.07)
- This evidence review found moderate quality evidence from 1 study with 813 infants showing that the use of a paraffin heater was not associated with
 - o any episodes of earache aOR 0.94 (95%CI 0.50 to 1.70)
 - recurrent earache (4 or more episodes separated by 21 days in 1 year) aOR 0.91 (95%CI 0.67 to 1.26)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas

- o Older people
- People with disabilities
- o Pregnant women
- People with conditions associated with or exacerbated by indoor air pollution

Sources of allergens

Pets (Grade F.1.3.1)

- This evidence review found low quality evidence from 1 study with 5,078 children showing that having pets at home was not associated with wheeze aOR 1.05 (95%CI 0.83 to 1.33)
- This evidence review found high quality evidence from 1 study with 1,879 infants showing that having a cat at home was protective against with wheeze aOR 0.65 (95%CI 0.47 to 0.89)
- This evidence review found moderate quality evidence from 1 study with 1,765 infants showing that prenatal and postnatal exposure to cats was not associated with
 - wheeze aOR 0.94 (95%CI 0.61 to 1.46)
 - o bronchiolitis aOR 0.69 (95%CI 0.47 to 1.03)
- This evidence review found moderate quality evidence from 1 study with 226 children at risk of atopy showing that dogs at home was protective against wheeze aOR 0.12 (95%CI 0.01 to 0.97)
- This evidence review found very low quality evidence from 1 study with 1,562 children showing exposure to pets was associated with cough
 - o aOR 1.58 (95%Cl 1.10 to 2.20) for previous short period of exposure to pets,
 - o aOR 1.64 (95%CI 1.09 to 2.46) for constant exposure to pets
- This evidence review found very low quality evidence from 1 study with 1,562 children showing previous long period of exposure to pets was not associated with cough aOR 1.10 (95%CI 0.72 to 1.68)
- This evidence review found very low quality evidence from 1 study with 526 children with asthma showing that pet ownership was not associated with airway hyper-responsiveness aOR 1.17 (95%CI 0.70 to 1.94)
- This evidence review found moderate quality evidence from 1 study with 3,535 children showing that for pet ownership in childhood was associated with asthma aOR 1.60 (95%CI 1.00 to 2.50)
- This evidence review found high quality evidence from 1 study with 1,765 infants showing that prenatal and postnatal exposure to cats was protective against asthma aOR 0.27 (95%CI 0.08 to 0.86)
- This evidence review found moderate quality evidence from 1 study with 5,078 children showing that pet ownership was protective against asthma aOR 0.69 (95%CI 0.52 to 0.91)
- This evidence review found low quality evidence from 1 study with 100 children with wheeze aOR showing that exposure to cats was not associated with asthma aOR 0.26 (95%CI 0.03 to 2.42) and exposure to dogs aOR 0.20 (95%CI 0.02 to 1.78)
- This evidence review found moderate quality evidence from 1 study with 3,535 children with wheeze showing exposure to pets was not associated with asthma aOR 1.10 (95%CI 0.60 to 2.00)

- This evidence review found high quality evidence from 1 study with 767 adults showing that having pets at home was associated with irritable bowel syndrome aOR 2.09 (95%CI 1.19 to 3.67) for exposure to an herbivore pet
- This evidence review found moderate quality evidence from 1 study with 767 adults showing that having pets at home was not associated with irritable bowel syndrome
 - o aOR 1.47 (95%CI 0.83 to 2.61) for any pet,
 - o aOR 1.58 (95%CI 0.90 to 2.76) for a carnivore pet,
 - o aOR 0.97 (95%Cl 0.26 to 3.59) for an omnivore pet
- This evidence review found high quality evidence from 1 study with 767 adults showing that having pets at home was associated with functional dyspepsia
 - o aOR 2.34 (95%CI 1.24 to 4.45) for exposure to an herbivore pet
 - o aOR 2.04 (95%CI 1.03 to 4.03) for exposure to an carnivore pet
- This evidence review found moderate quality evidence from 1 study with 767 adults showing that having pets at home was not associated with functional dyspepsia
 - o aOR 1.69 (95%CI 0.86 to 3.36) for any pet
 - o aOR 0.98 (95%Cl 0.21 to 4.50) for an omnivore pet
- This evidence review found low quality evidence from 1 study with 61,799 women that pet ownership in childhood was not associated with papillary thyroid cancer aRR 0.77 (95%CI 0.51 to 1.17)
- This evidence review found moderate quality evidence from 1 study with 813 infants showing that having pets at home was not associated with
 - o any episodes of otitis media aOR 0.76 (95%CI 0.47 to 1.26)
 - recurrent otitis media (four or more episodes separated by 21 days in 1 year) aOR 1.06 (95%CI 0.90 to 1.26)
- This evidence review found low quality evidence from 1 study with 3,143 children showing that having pets at home was not associated with type 1 diabetes (clinical or pre-clinical)
 - o aOR for a dog at home aOR 0.40 (95%CI 0.14 to 1.14)
 - o aOR for a cat at home aOR 1.34 (95%CI 0.58 to 3.10)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - People with disabilities
 - Pregnant women

Carpet flooring (Grade F.1.3.2)

- This evidence review found high quality evidence from 1 study with 465 infants showing that having carpet flooring at home during pregnancy was associated with
 - wheeze aOR 5.39 (95%CI 1.75 to 16.54)
 - o obstructive bronchitis aOR 4.39 (95%CI 1.01 to 19.05)
- This evidence review found moderate quality evidence from 1 study with 465 infants showing that having carpet flooring at home in the first year of life was not associated with wheeze aOR 4.18 (95%CI 0.40 to 43.70)

- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - o People with conditions associated with or exacerbated by indoor air pollution

Second hand mattress (Grade F.1.3.3)

- This evidence review found moderate quality evidence from 1 study with 2,898 infants showing that having a used (second-hand) mattress was associated with cough aOR 1.47 (95%CI 1.00 to 2.17) in the infants at risk of allergies
- This evidence review found low quality evidence from 1 study with 2,898 infants showing having a used (second-hand) mattress was not associated with cough for infants not at risk of allergies aOR 1.22 (95%CI 0.80 to 1.88)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Sources of damp or mould

High air humidity in bathroom (Grade F.1.4.1)

- This evidence review found low quality evidence from 1 study with 9,808 adults showing that high bathroom air humidity was associated with both
 - o cough aOR 2.30 (95%CI 2.21 to 2.40)
 - o nasal symptoms aOR 1.94 (95%CI 1.88 to 2.01)
 - throat symptoms aOR 3.23 (95%CI 3.12 to 3.25)
 - o facial skin symptoms aOR 2.42 (95%Cl 2.33 to 2.51)
 - o headache aOR 3.07 (95%CI 2.96 to 3.17)
 - o tiredness aOR 2.16 (95%CI 2.11 to 2.22)
 - eye irritation aOR 2.94 (95%CI 2.83 to 3.05)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - Children and young people
 - o People with conditions associated with or exacerbated by indoor air pollution

Condensation on windows (Grade F.1.4.2)

- This evidence review found low quality evidence from 1 study with 9,808 adults showing that condensation on windows was associated with both
 - o Cough aOR 2.58 (95%CI 2.47 to 2.70)
 - Nasal symptoms aOR 2.72 (95%Cl 2.62 to 2.81)
 - Throat symptoms aOR 3.22 (95%CI 3.19 to 3.35)
 - o facial skin symptoms aOR 2.11 (95%CI 2.02 to 2.20)
 - headache aOR 3.30 (95%CI 3.19 to 3.43)
 - o tiredness aOR 2.19 (95%CI 2.12 to 2.25)
 - o eye irritation aOR 3.14 (95%CI 3.01 to 3.27)
- This evidence review found high quality evidence from 1 study with 4,779 children showing that more than 5 cm condensation on windows was associated with autism spectrum disorders
 - o aOR 2.05 (95%CI 1.03 to 4.10) for condensation in the child's room
 - o aOR 2.03 (95%CI 1.08 to 3.82) for condensation in the parent's room
- This evidence review found moderate quality evidence from 1 study with 4,779 children showing that between 1 cm and 5cm condensation on windows was not associated with autism spectrum disorders
 - o aOR 1.35 (95%CI 0.71 to 2.57) for condensation on windows in the child's room
 - o aOR 1.52 (95%CI 0.84 to 2.73) for condensation on windows in the parent's room
- This evidence review found low quality evidence from 1 study with 6,853 children showing that heavy condensation on windows was not associated with acute respiratory infection requiring hospitalisation
 - o aHR 1.01 (95%CI 0.86 to 1.17) for condensation on windows rarely
 - o aHR 1.05 (95%CI 0.88 to 1.27) for condensation on windows guite often
 - o aHR 1.00 (95%CI 0.77 to 1.31) for heavy condensation on windows quite often
- This evidence review found very low quality evidence from 1 study with 7,104 adults showing that condensation on windows was not associated with
 - o asthma aRR 1.07 (95%CI 0.75 to 1.53)
 - o asthma and airway hyper-responsiveness aRR 1.43 (95%CI 0.67 to 3.07)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - People with disabilities
 - Pregnant women
 - People with conditions associated with or exacerbated by indoor air pollution

Moisture on walls/surfaces (Grade F.1.4.3)

 This evidence review found very low quality evidence from 1 study with 1,916 children showing that moisture on walls was not associated with asthma aOR 0.92 (95%CI 0.54 to 1.54)

- This evidence review found low quality evidence from 1 study with 3,360 children showing that moisture on walls was associated with allergic rhino-conjunctivitis aOR 1.90 (95%CI 1.01 to 3.56)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - People with conditions associated with or exacerbated by indoor air pollution

History of water leakage (Grade F.1.4.4)

- This evidence review found low quality evidence from 1 study with 9,808 adults that a history of water leakage was associated with
 - o cough aOR 1.52 (95%CI 1.44 to 1.59)
 - nasal symptoms aOR 1.36 (95%CI 1.31 to 1.41)
 - throat symptoms aOR 2.18 (95%CI 2.09 to 2.28)
 - facial skin symptoms aOR 1.56 (95%CI 1.48 to 1.63)
 - o headache aOR 1.27 (95%CI 1.21 to 1.33)
 - tiredness aOR 2.19 (95%CI 2.12 to 2.25)
 - o irritation aOR 1.57 (95%CI 1.50 to 1.65)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - People with disabilities
 - Pregnant women
 - Children and young people
 - People with conditions associated with or exacerbated by indoor air pollution

Water damage (Grade F.1.4.5)

- This evidence review found low quality evidence from 1 study with 1,863 children showing that water damage was associated with hay fever (reported as allergic rhinitis) aOR 2.06 (95%CI 1.35 to 3.13)
- The evidence review found very low quality evidence from 1 study with 1916 children showing that water damage was not associated with asthma aOR 1.01 (95%CI 0.45 to 2.26)
- The evidence review found moderate quality evidence from 1 study with 499 infants showing that water damage was not associated with lower respiratory illness aOR 1.34 (95%CI 0.99 to 1.82) for infants at risk of asthma or allergy.
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas

- o Older people
- People with disabilities
- Pregnant women
- People with conditions associated with or exacerbated by indoor air pollution

Damp condition (Grade F.1.4.6)

- This evidence review found high quality evidence from 1 study with 26,888 children showing that damp conditions were associated with
 - o bronchitis (ever diagnosed) aOR 1.25 (95%Cl 1.13 to 1.37) for homes in East Germany
 - o bronchitis (ever diagnosed aOR 1.30 (95 % 1.03 to 1.65) for homes in West Germany,
 - frequent colds (more than 4 colds in last 12 months) aOR 1.41 (95%Cl 1.25 to 1.60) for homes in East Germany
 - frequent colds (more than 4 colds in last 12 months) aOR 1.62 95 % 1.21 to 2.17) for homes in West Germany,
 - o frequent cough aOR 1.66 (95%CI 1.42 to 1.95) for homes in East Germany
 - o frequent cough aOR 2.60 (95%CI 1.90 to 3.55) for homes in West Germany
 - sneeze attacks in the last 12 months aOR 1.52 (95%Cl 1.26 to 1.83) for homes in East Germany
 - sneeze attacks in the last 12 months aOR 2.25 (95%Cl 1.52 to 3.33) for homes in West Germany
 - o eczema aOR = 1.15 (95%Cl 1.01 to 1.31) for homes in East Germany
- This evidence review found moderate quality evidence from 1 study with 26888 children showing that damp conditions were not associated with
 - o allergies (ever diagnosed) aOR 1.09 (95%Cl 0.93 to 1.66) for homes in East Germany
 - o allergies (ever diagnosed) aOR 1.20 (95%Cl 0.87 to 1.66) for homes in West Germany
 - eczema (ever diagnosed) aOR 1.10 (95%CI 0.77 to 1.57) for homes in West Germany
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - People with disabilities
 - Pregnant women
 - People with conditions associated with or exacerbated by indoor air pollution

Sources of VOCs

Parquet flooring (Grade F.1.5.1)

- This evidence review found moderate quality evidence from 1 study with 465 infants showing that exposure to parquet flooring during pregnancy was not associated with wheeze aOR 5.78 (95%CI 0.30 to 111.08)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - People with disabilities
 - People with conditions associated with or exacerbated by indoor air pollution

Laminate flooring (Grade F.1.5.2)

- This evidence review found high quality evidence from 1 study with 465 infants showing that exposure to laminate flooring during pregnancy was associated with wheeze aOR 4.46 (95%CI 1.01 to 19.63)
- This evidence review found moderate quality evidence from 1 study with 465 infants showing that exposure to laminate flooring in in the first year of life was not associated with wheeze aOR 2.44 (95%CI 0.40 to 14.74)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - o People with disabilities
 - o People with conditions associated with or exacerbated by indoor air pollution

PVC flooring (Grade F.1.5.3)

- This evidence review found high quality evidence from 1 study with 465 infants showing that exposure to PVC flooring during pregnancy was associated with wheeze aOR 24.7 (95%CI 2.18 to 280.39)
- This evidence review found high quality evidence from 1 study with 465 infants showing that exposure to PVC flooring in the first year of life was associated with wheeze aOR 51.7 (95%CI 3.21 to 833.2)
- This evidence review found moderate quality evidence from 1 study with 4,779 children showing that exposure to PVC flooring in childhood was not associated with autism spectrum disorders at 6 to 8 years of age
 - o aOR 1.19 (95%CI 0.71 to 2.00) for PVC flooring in the child's bedroom
 - o aOR 1.59 (95%Cl 0.97 to 2.61) for PVC flooring in the parent's bedroom
- This evidence review found moderate quality evidence from 1 study with 3,228 children showing that exposure to PVC flooring in childhood was associated with asthma at 5 years of age
 - aOR 1.54 (95%CI 1.06 to 2.23) for PVC flooring in the child's bedroom when compared with wood flooring,
 - aOR 1.60 (95%CI 1.29 to 2.81) for PVC flooring in the parent's bedroom compared to wood flooring
 - aOR 1.71 (95%CI 1.05 to 2.80) for PVC flooring in the parent's bedroom when compared with other types of flooring
- This evidence review found moderate quality evidence from 1 study with 3,228 children showing that exposure to PVC flooring in childhood was associated with asthma at 10 years of age
 - aOR 1.54 (95%CI 1.06 to 2.23) for PVC flooring in the child's bedroom when compared with other types of flooring,
 - aOR 2.04 (95%CI 1.41 to 2.94) for PVC flooring in the parent's bedroom when compared with other types of flooring and
 - aOR 1.90 (95%CI 1.29 to 2.81) for PVC flooring in the parent's bedroom when compared with wood flooring

- This evidence review found low quality evidence from 1 study with 3,228 children showing that exposure to PVC flooring in child's room was not associated with
 - asthma at 5 years of age aOR 1.50 (95%Cl 0.91 to 2.47) when compared to other flooring
 - asthma at 10 years of age aOR 1.37 (95%Cl 0.92 to 2.04) when compared to wood flooring
- This evidence review found moderate quality evidence from 1 study with 2,779 children showing that
 - PVC flooring in the child's bedroom was not associated with asthma in the following 5 years aOR 1.52 (95%CI 0.99 to 2.35)
 - PVC flooring in the parent's bedroom was not associated with asthma in the following 5 years aOR 1.48 (95%CI 0.86 to 2.57)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - o People with disabilities
 - o People with conditions associated with or exacerbated by indoor air pollution

New furniture (Grade F.1.5.4)

- This evidence review found moderate quality evidence from 1 study with 465 infants showing that new furniture during pregnancy or in the first year of life was not associated with recurrent wheeze
 - o aOR 1.94 (95%CI 0.72 to 5.26) for new furniture during pregnancy
 - o aOR 2.26 95%Cl 0.83 to 6.17) for new furniture in the first year of life.
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - o People with disabilities
 - o People with conditions associated with or exacerbated by indoor air pollution

Home products - Air fresheners (Grade F.1.5.5)

- This evidence review found very low quality evidence from 1 study with 3,503 adults that the use of air fresheners was not associated with asthma
 - o aRR 1.29 (95%CI 0.74 to 2.26) for the use of any perfumed or scented product
 - o aRR 1.46 (95%Cl 0.78 to 2.70) for the use of air refreshing sprays
- This evidence review found moderate quality evidence from 1 study with 14,541 women showing that the use of air fresheners most days during pregnancy was associated with headache
 - o aOR 1.24 (95%CI 1.11 to 1.38) at 8 months after giving birth
 - o aOR 1.22 (95%CI 1.09 to 1.36) at between 9 to 21 months after giving birth
- This evidence review found moderate quality evidence from 1 study with 14,541 women showing that the use of air fresheners once a week during pregnancy was associated with headache

- o aOR 1.29 (95%CI 1.14 to 1.47) at between 9 to 21 months after giving birth
- This evidence review found that the use of air fresheners once a week during pregnancy was not associated with headache in mothers 8 months after birth - low quality evidence from 1 study with 14,541 women aOR 1.06 (95%CI 0.94 to 1.19)
- This evidence review found that the use of air fresheners most days during pregnancy was associated with depression at 8 months after giving birth - moderate quality evidence from 1 study with 14,541 women aOR 1.19 (95%CI 1.05 to 1.36)
- This evidence review found that the use of air fresheners once a week during pregnancy was not associated with depression in mothers 8 months after the birth low quality evidence from 1 study with 14,541 women aOR 1.11 (95%CI 0.96 to 1.29)
- This evidence review found low quality evidence from 1 study with 14,541 women showing that the use of air fresheners during pregnancy was not associated with
 - cough or cold 9 to 21 months after the birth aOR 1.03 (95%Cl 0.87 to 1.20) for use of air fresheners once a week during pregnancy
- This evidence review found low quality evidence from 1 study with 14,541 women showing that the use of air fresheners during pregnancy was associated with a reduction in cough or cold 9 to 21 months after the birth aOR 0.82 (95%Cl 0.72 to 0.93) for air freshener use most days during pregnancy
- This evidence review found low quality evidence from 1 study with 2,292 mothers of infants showing that the use of air fresheners was not associated with
 - o Wheeze aOR 1.09 (95%CI 0.87 to 1.37) for air freshener use,
 - Wheeze aOR 1.39 (95%Cl 0.85 to 2.29) for the use of air fresheners during pregnancy only
 - Wheeze aOR 1.23 (95%Cl 0.79 to 1.93) for the use of air fresheners during and after pregnancy
 - Lower respiratory tract infections aOR 1.31 (95%CI 0.77 to 2.21) for the use of air fresheners during pregnancy only
- This evidence review found moderate quality evidence from 1 study with 2,292 children showing that the use of air fresheners was associated with
 - wheeze aOR 1.75 (95%Cl 1.01 to 3.04) for the use of air fresheners after pregnancy only
 - Lower respiratory tract infections aOR 1.29 (95%CI 1.03 to 1.63)
 - Lower respiratory tract infections aOR 1.85 (95%Cl 1.04 to 3.30) for the use of air fresheners after pregnancy only
 - Lower respiratory tract infections aOR 1.59 (95%Cl 1.00 to 2.55) for the use of air fresheners during and after pregnancy
- This evidence review found very low quality evidence from 1 study with 3,503 adults showing that the was perfumed or scented products were not associated with wheeze
 - o aRR 1.11 (95%CI 0.83 to 1.49) for the use of any perfumed or scented products
 - o aRR 1.36 (95%CI 0.98 to 1.88) for the use of air refreshing sprays
- This evidence review found moderate quality evidence from 1 study with 13,971 infants showing that the use of air fresheners during pregnancy was associated with earache in infants
 - aOR 1.24 (95%CI 1.02 to 1.50) for the use of air freshener once a week during pregnancy

- aOR 1.30 (95%Cl 1.09 to 1.54) for the use of air fresheners most days during pregnancy
- This evidence review found moderate quality evidence from 1 study with 14,541 mothers
 of infants showing that the use of air fresheners was associated with diarrhoea 9 to 21
 months after birth
 - aOR 1.14 (95%CI 1.00 to 1.31) for the use of air freshener once a week during pregnancy
 - o aOR 1.14 (95%CI 1.01 to 1.28) for the use of air freshener most days during pregnancy
- This evidence review found low quality evidence from 1 study with 13,971 infants showing that the use of air fresheners was not associated with
 - Diarrhoea aOR 1.10 (95%Cl 0.99 to 1.23) for the use of air freshener most days during pregnancy,
 - vomiting aOR 1.06 (95%CI 0.93 to 1.20) for the use of air fresheners once a week during pregnancy
 - vomiting aOR 1.09 (95%CI 0.97 to 1.22) for the use of air freshener most days during pregnancy
- This evidence review found moderate quality evidence from 1 study with 13,971 infants showing that the use of air fresheners once a week during pregnancy was associated with diarrhoea aOR 1.20 (95%CI 1.06 to 1.35).
- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - Older people
 - People with disabilities
 - o People with conditions associated with or exacerbated by indoor air pollution

Home products - Cleaning sprays (Grade F.1.5.6)

- This evidence review found very low quality evidence from 1 study with 633 adults showing that the use of cleaning sprays was not associated with
 - o wheeze aOR 1.53 (95%CI 0.88 to 2.65) for low use,
 - o aOR 1.34 (95%CI 0.75 to 2.39) for medium use,
 - o aOR 1.71 (95%CI 0.80 to 3.67) for high use.
- This evidence review found low quality evidence from 1 study with 1,157 infants showing that the use of cleaning sprays after pregnancy was not associated with wheeze aOR 1.34 (95%CI 0.80 to 2.24).
- This evidence review found moderate quality evidence from 1 study with 1,879 infants showing that that daily use of cleaning sprays was not associated with wheeze aOR 1.50 (95%CI 0.97 to 2.32).
- This evidence review found moderate quality evidence from 1 study with 2,292 infants that the use of cleaning sprays was associated with wheeze
 - o aOR 1.37 (95%CI 1.10 to 1.69) for the use of sprays,
 - o aOR 1.62 (95%CI 1.11 to 2.36) for the use of sprays during pregnancy only
 - o aOR 1.61 (95%CI 1.08 to 2.41) for the use of sprays during and after pregnancy
- This evidence review found low quality evidence from 1 study with 2,292 infants that the
 use of cleaning sprays after pregnancy was not associated with wheeze aOR 1.37 (95%CI
 1.10 to 1.69)

- This evidence review found low quality evidence from 1 study with 683 women showing that the use of 2 or more types of cleaning sprays more than 1 day per week was associated with asthma aOR 1.67 (95%CI 1.08 to 2.56)
- This evidence review found very low quality evidence from 1 study with 683 women showing that the use of cleaning sprays was not associated with asthma aOR 0.68 (95%Cl 0.44 to 1.04) for the use of 1 type of cleaning spray more than 1 day per week
- This evidence review found very low quality evidence from 1 study with 1,895 adults showing that the use of household sprays was not associated with asthma
 - o aOR 0.70 (95%CI 0.23 to 2.06) for low use of household spray,
 - o aOR 0.78 (95%CI 0.26 to 2.36) for medium use of household spray
 - o aOR 2.79 (95%Cl 0.84 to 9.20) for high use of household spray
- This evidence review found low quality evidence from 1 study with 570 women with asthma showing that the spray use more than 1 day per week was not associated with asthma aOR 1.45 (95%CI 0.94 to 2.24)
- This evidence review found low quality evidence from 1 study with 3,503 adults showing that the use of cleaning sprays was associated with asthma
 - o aOR 2.11 (95%CI 1.15 to 3.89) for the use of sprays 4 to 7 days per week,
 - aOR 2.96 (95%Cl 1.33 to 6.56) for the use of three or more sprays more than 1 day per week,
 - o aOR 2.46 (95%CI 1.26 to 4.80) for the use of furniture sprays,
 - o aOR 1.49 (95%CI 1.12 to 1.99) for the use of any spray
- This evidence review found very low quality evidence from 1 study with 3,503 adults the
 use of household sprays was not associated with asthma
 - o aOR 1.28 (95%CI 0.78 to 2.09) for the use of any spray,
 - o aOR 0.93 (95%CI 0.51 to 1.67) for the use of sprays 1 to 3 days per week,
 - o aOR 0.97 (95%CI 0.53 to 1.77) for use of 1 type of spray more than 1 day per week,
 - o aOR 1.47 (95%Cl 0.70 to 3.06) for use of 2 types of spray more than 1 day per week,
 - o aOR 1.43 (95%Cl 0.84 to 2.44) for the use of glass-cleaning sprays,
 - o aOR 0.80 (95%CI 0.11 to 5.93) for the use of sprays for carpets, rugs and curtains,
 - o aOR 0.93 (95%Cl 0.30 to 2.85) for the use of sprays for mopping floors,
 - aOR 0.63 (95%CI 0.09 to 4.64) for the use of oven sprays
 - o aOR 1.51 (95%CI 0.46 to 4.96) for the use of ironing sprays
- This evidence review found low quality evidence from 1 study with 3,503 adults showing that the use of cleaning sprays was associated with asthma attacks and / or nocturnal shortness of breath
 - o aOR 1.75 (95%Cl 1.21 to 2.54) for use of sprays 4 to 7 days per week
 - aOR 2.40 (95%Cl 1.47 to 3.91) for the use of 3 or more types of spray more than 1 day per week
- This evidence review found very low quality evidence from 1 study with 3,503 adults showing that the use of sprays was not associated with asthma attacks and / or nocturnal shortness of breath
 - o aOR 1.36 (95%CI 0.99 to 1.89) for the use of sprays 1 to 3 days per week,
 - aOR 1.37 (95%Cl 0.99 to 1.90) for the use of 1 type of spray more than 1 day per week,

- aOR 1.45 (95%Cl 0.92 to 2.27) for the use of 2 types of spray more than 1 day per week
- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - o Older people
 - o People with disabilities
 - Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Home products - Solvents (Grade F.1.5.7)

- This evidence review found very low quality evidence from 1 study with 3,503 adults that exposure to solvents was not associated with asthma aOR 0.48 (95%Cl 0.12 to 1.97) for use of solvents/strain removers
- This evidence review found low quality evidence from 1 study with 3,503 adults showing that exposure to solvents/strain removers was associated with wheeze aRR 2.00 (95%CI 1.30 to 3.07).
- This evidence review found moderate quality evidence from 1 study with 2,292 infants showing that exposure to solvents was associated with
 - wheeze aOR 1.30 (95%CI 1.03 to 1.62)
 - o lower respiratory tract infections aOR 1.54 (95%CI 1.11 to 2.14).
- This evidence review found moderate quality evidence from 1 study with 2,292 infants showing that exposure to solvents was not associated with lower respiratory tract infections aOR 1.19 (95%CI 0.95 to1.48)
- This evidence review found very low quality evidence from 1 study with 1,157 infants showing that exposure to solvents was not associated with wheeze
 - Wheeze aOR 1.04 (95%CI 0.71 to 1.51) for prenatal but not postnatal exposure,
 - Wheeze aOR 0.87 (95%Cl 0.55 to1.37) for postnatal but not prenatal exposure
 - o Wheeze aOR 1.81 (95%CI 0.98 to 3.37) for both prenatal and postnatal exposure.
- This evidence review found low quality evidence from 1 study with 1,505 children up to 2
 years of age showing that exposure to solvents was associated with wheeze
 - o aOR 1.66 (95%CI 1.11 to 2.47) for postnatal but not prenatal exposure
 - o aOR 2.50 (95%CI 1.45 to 4.33) for both prenatal and postnatal exposure
- This evidence review found very low quality evidence from 1 study with 1,505 children up to 2 years of age showing that exposure to solvents was not associated with wheeze
 - o aOR 0.89 (95%CI 0.34 to 2.31) for prenatal but not postnatal exposure
- This evidence review found very low quality evidence from 1 study with 1,505 children up to 2 years of age showing that exposure to solvents was not associated with
 - Eczema aOR 0.72 (95%Cl 0.35 to 1.50) for prenatal but not postnatal exposure to solvents,
 - Eczema aOR 1.03 (95%CI 0.79 to 1.36) for postnatal but not prenatal exposure to solvents
 - Eczema aOR 1.23 (95%Cl 0.84 to 1.82) for both prenatal and postnatal exposure to solvents
 - Food allergies aOR 1.25 (95%Cl 0.41 to 3.80) for prenatal but not postnatal exposure to solvents,

- Food allergies aOR 1.28 (95%CI 0.80 to 2.03) for postnatal but not prenatal exposure to solvents
- Food allergies aOR 1.32 (95%Cl 0.71 to 2.46) for prenatal and postnatal exposure to solvents

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- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - o People with disabilities
 - o People with conditions associated with or exacerbated by indoor air pollution

Home products - Aerosols (Grade F.1.5.8)

- This evidence review found moderate quality evidence from 1 study with 14,541 mothers
 of infants showing that aerosol use once a week during pregnancy was associated with
 headache
 - o aOR 1.16 (95%CI 1.00 to 1.35) at 8 months after the birth
 - o aOR 1.35 (95%Cl 1.15 to 1.59) at 9 to 21 months after the birth
- This evidence review found moderate quality evidence from 1 study with 14,541 mothers
 of infants showing that aerosol use most days during pregnancy was associated with
 headache in mothers
 - o aOR 1.25 (95%Cl 1.13 to 1.39) at 8 months after the birth
 - o aOR 1.21 (95%Cl 1.10 to 1.34) at between 9 to 21 months after the birth
- This evidence review found that the use of aerosols most days during pregnancy was not associated with depression at 8 months after giving birth - low quality evidence from 1 study with 14,541 women aOR 1.03 (95%CI 0,91 to 1.17)
- This evidence review found that the use of aerosols once a week during pregnancy was not associated with depression in mothers 8 months after the birth - low quality evidence from 1 study with 14,541 women aOR 1.06 (95%CI 0.88 to 1.27)
- This evidence review found low and moderate quality evidence from 1 study with 14,541 mothers of infants showing that exposure to aerosols was not associated with influenza in mothers 9 to 21 months after the birth
 - aOR 1.03 (95%CI 0.85 to 1.24) for aerosol use once a week during pregnancy
 - aOR 0.87 (95%CI 0.77 to 0.99) for aerosol use daily or most days during pregnancy
- This evidence review found low quality evidence from 1 study with 13,971 infants showing that that exposure to aerosols during pregnancy was not associated with earache in infants
 - o aOR 1.00 (95%CI 0.78 to 1.29) for aerosol use once a week during pregnancy
 - o aOR 1.05 (95%CI 0.84 to 1.25) for aerosol use daily or most days during pregnancy
- This evidence review found moderate quality evidence from 1 study with 13,971 infants showing that exposure to aerosols was associated with diarrhoea aOR 1.22 (95%Cl 1.09 to 1.36) for aerosol use daily or most days during pregnancy
- This evidence review found low quality evidence from 1 study with 13,971 infants showing that exposure to aerosols was not associated with diarrhoea aOR 1.09 (95%CI 0.93 to 1.28) for aerosol use once a week during pregnancy

- This evidence review found moderate quality evidence from 1 study with 13,971 infants showing that exposure to aerosols was associated with vomiting
 - o aOR 1.17 (95%CI 1.00 to 1.37) for aerosol use once a week during pregnancy
 - o aOR 1.14 (95%Cl 1.02 to 1.27) for aerosol use daily or most days during pregnancy
- This evidence review found low quality evidence from 1 study with 14,541 mothers of infants that exposure to aerosols was associated with urinary tract infections in mothers 9 to 21 months after the birth aOR 1.23 (95%Cl 1.04 to 1.45) for aerosol use daily or most days during pregnancy
- This evidence review found moderate quality evidence from 1 study with 14,541 mothers
 of infants that exposure to aerosols was not associated with urinary tract infections in
 mothers 9 to 21 months after the birth aOR 1.16 (95%CI 0.89 to 1.52) for aerosol use
 once a week during pregnancy
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - People with disabilities
 - o People with conditions associated with or exacerbated by indoor air pollution

Paint (Grade F.1.5.9)

- This evidence review found moderate quality evidence from 1 study with 465 infants showing that exposure to paint fumes was not associated with recurrent wheeze
 - o aOR 2.35 (95%CI 0.89 to 6.20) for exposure during pregnancy
 - o aOR 2.53 (95%CI 0.85 to 7.49) for exposure during 1st year of life
- This evidence review found high quality evidence from 1 study with 465 infants that
 exposure to paint fumes during pregnancy was associated with obstructive bronchitis in
 first of life aOR 5.46 (95%CI 1.09 to 27.20)
- This evidence review found high quality evidence from 1 study with 20,103 women that exposure to paint fumes in first trimester of pregnancy was associated with congenital renal anomalies aOR 2.16 (95%CI 1.02 to 4.58) for exposure
- This evidence review found moderate quality evidence from 1 study with 20,103 women showing that exposure to paint fumes in first trimester of pregnancy was not associated with
 - o congenital anomalies (all) aOR 0.95 (95%Cl 0.74 to 1.21),
 - o aOR 2.19 (95%CI 0.76 to 6.32) for congenital anomalies to the nervous system),
 - o aOR 1.79 (95%CI 0.70 to 4.57) for congenital anomalies to the eyes,
 - o aOR 2.15 (95%Cl 0.84 to 5.55) for congenital anomalies to the ear, face and neck),
 - o aOR 0.76 (95%CI 0.39 to 1.49) for heat defects,
 - o aOR 1.13 (95%Cl 0.27 to 4.79) for congenital anomalies to the respiratory system,
 - o aOR 1.06 (95%CI 0.33 to 3.46) for cleft lip and cleft palate,
 - o aOR 0.61 (95%CI 0.15 to 2.50) for congenital anomalies to the digestive system,
 - o aOR 0.83 (95%CI 0.48 to 1.43) for congenital anomalies to the genitals,
 - o aOR 0.82 (95%Cl 0.54 to 1.24) for limb defects,
 - o aOR 1.77 (95%CI 0.75 to 4.16) for congenital anomalies to the muscular and skeletal,

- o aOR 1.24 (95%Cl 0.62 to 2.46) for other congenital anomalies.
- This evidence review found moderate quality evidence from 1 study with 19,000 women that exposure to paint fumes during pregnancy was not associated with
 - o small for gestational age aOR 0.89 (95%CI 0.81 to 0.98)
 - pre-term birth aOR 0.95 (95%CI 0.82 to 1.11)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - o People with conditions associated with or exacerbated by indoor air pollution

Any type of redecoration (Grade F.1.5.10)

- This evidence review found moderate quality evidence from 1 study with 465 infants showing that redecoration during pregnancy was associated with recurrent wheeze aOR 2.04 (95%CI 0.78 to 5.28).
- This evidence review found moderate quality evidence from 2 studies with 2344 infants showing that redecoration in the first year of life was associated with recurrent wheeze aOR 1.22 (95%Cl 0.96 to 1.54) in the first study with 1879 infants and aOR 1.89 (95%Cl 0.71 to 5.06) in the second study with 465 infants.
- This evidence review found low quality evidence from 1 study with 475 premature infants at risk of allergies that exposure to redecoration during pregnancy was associated with pulmonary infections aOR 5.6 (95%CI 1.3 to 24.0)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - People with disabilities
 - People with conditions associated with or exacerbated by indoor air pollution

Building characteristics and health outcomes

Building age (Grade F.1.6.1)

- This evidence review found moderate quality evidence from 1 study with 540 children that building age was associated with recurrent wheeze
 - o aOR 1.69 (95%Cl 1.01 to 2.89) for buildings built between 1940 and 1975 in Sweden
 - o aOR 1.86 (95%CI 1.05 to 3.27) for buildings built after 1975 in Sweden compared to houses built before 1940.
- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - Older people
 - People with disabilities

- Pregnant women
- People with conditions associated with or exacerbated by indoor air pollution

Dwelling size (Grade F.1.6.2)

 This evidence review found very low quality evidence from 1 study with 1,137 children showing that dwelling size was not associated with lower respiratory tract infections a OR 0.99 (95%CI 0.92 to 1.06) per room increase in household

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- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - People with conditions associated with or exacerbated by indoor air pollution

Central air conditioning (Grade F.1.6.3)

- This evidence review found high quality evidence from 1 study with 176 children showing that central air conditioning was protective against asthma aOR 0.3 (95%CI 0.14 to 0.83)
- This evidence review found moderate quality evidence from 1 study with 813 infants showing that central air conditioning was not associated with
 - o any episodes of earache aOR 0.52 (95%Cl 0.27 to 1.03)
 - recurrent earache (four or more episodes separated by 21 days in 1 year) aOR 0.93 (95%CI 0.77 to 1.11)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - People with disabilities
 - Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Ventilation rate (Grade F.1.6.4)

- This evidence review found very low quality evidence from 1 study with 400 children showing that ventilation rate was not associated with asthma and allergic symptoms
 - o aOR 1.17 (95%CI 0.57 to 2.42) for ventilation rate (third quartile versus fourth quartile),
 - aOR 1.35 (95%Cl 0.66 to 2.74) for ventilation rate (second quartile versus fourth quartile)
 - o aOR 1.95 (95%Cl 0.94 to 4.04) for ventilation rate (first quartile versus fourth quartile)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas

- o Older people
- People with disabilities
- Pregnant women
- People with conditions associated with or exacerbated by indoor air pollution

Proximity to traffic – Traffic intensity (Grade F.1.6.5)

- This evidence review found high and moderate quality evidence from 1 study with 7,898 children that for 8,640 or more cars per day within 100m of the birth address was not associated with obstructive bronchiolitis aHR 1.0 (95%CI 0.9 to1.2)
- This evidence review found high and moderate quality evidence from 1 study with 7,898 children that for 8,640 or more cars per day within 100m of the birth address was protective against bronchiolitis aHR 0.7 (95%CI 0.6 to 0.9)
- This evidence review found high and moderate quality evidence from 1 study with 7,898 children that for 8,640 or more cars per day with 100m of the birth address and had never moved was not associated with
 - o bronchiolitis aHR 0.7 (95%CI 0.6 to 0.9)
 - obstructive bronchiolitis aHR 1.0 (95%CI 0.8 to 1.2)
- This evidence review found high quality evidence from 1 study with 7,898 children showing that proximity to traffic was not associated with asthma
 - aHR 0.7 (95%Cl 0.6 to 0.9) for 8,640 or more cars per day with 100m of the birth address
 - aHR 0.7 (95%Cl 0.6 to 0.9) for 8,640 or more cars per day with 100m of the birth address and never moved
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - People with conditions associated with or exacerbated by indoor air pollution

Located within 50 m of major traffic (Grade F.1.6.6)

- This evidence review found moderate quality evidence from 1 study with 2,628 adults showing that the location of the dwelling within 50 metres of a major road was associated with wheeze aOR 1.31 (95%CI 1.00 to 1.71)
- This evidence review found moderate quality evidence from 1 study with 71,271 women showing that the location of the dwelling within 50 metres of a major road was not associated with anxiety symptoms aOR 1.01 (95%CI 0.95 to 1.08)
- This evidence review found low quality evidence from 1 study with 2,628 adults showing that the location of the dwelling within 50 metres of a major road was not associated with
 - o chronic cough aOR 1.24 (95%CI 0.92 to 1.68)
 - chronic phlegm aOR 1.18 (95%CI 0.88 to 1.56)

- This evidence review found low quality evidence from 1 study with 3,577 children showing that the location of the dwelling within 50 metres of a major road was not associated with
 - o cough without infection aOR 0.74 (95%CI 0.55 to 1.00),
 - dry cough at night aOR 0.84 (95%CI 0.61 to 1.16),
 - o wheeze aOR 1.14 (95%CI 0.92 to 1.42),
 - o sneezing, runny, stuffy nose aOR 1.10 (95%Cl 0.87 to 1.39)
 - o respiratory infections aOR 1.03 (95%CI 0.86 to 1.23)
 - o asthmatic / spastic / obstructive bronchitis aOR 1.12 (95%CI 0.88 to 1.44)
- This evidence review found moderate quality evidence from 1 study with 5,921 children showing that the location of the dwelling within 50 metres of a major road was associated with asthma aOR 1.66 (95%CI 1.01 to 2.59)
- This evidence review found moderate quality evidence from 1 study with 68,195 children of pre-school age showing that the location of the dwelling within 50 metres of a major road was associated with asthma aOR 1.25 (95%CI 1.04 to 1.49)
- This evidence review found moderate quality evidence from 1 study with 3,297 children showing that the location of the dwelling within 50 metres of a major road was not associated with asthma exacerbations requiring hospitalisations aHR 1.11 (95%CI 0.92 to 1.33)
- This evidence review found low quality evidence from 1 study with 5,921 children showing that the location of the dwelling within 50 metres of a major road was not associated with
 - hay fever aOR 1.16 (95%CI 0.67 to, 2.00)
 - o eczema aOR 0.96 (95%CI 0.72 to 1.11)
- This evidence review found that the location of the dwelling within 50 metres of a major road was associated with
 - Coronary Heart Disease (CHD) Mortality/sudden cardiac death high quality evidence from 1 study with 450,283 adults aRR 1.29 (95%CI 1.18 to 1.41)
 - Coronary Heart Disease (CHD) Mortality/sudden cardiac death high quality evidence from 1 study with 107,130 women aHR 1.38 (95%CI 1.04 to 1.82)
- This evidence review found high quality evidence from 1 study with 103,650 women showing that the location of the dwelling within 50 metres of a major road was associated with lung cancer incidence aHR 2.01 (95%CI 1.06 to 3.80)
- This evidence review found moderate quality evidence from 1 study with 2,372 adults showing that the location of the dwelling within 50 metres of a major road was not associated with obesity aOR 1.10 (95%CI 0.97 to 1.25)
- This evidence review found moderate quality evidence from 1 study with 85,251 women showing that the location of the dwelling within 50 metres of a major road was not associated with uterine leiomyomata aOR 1.01 (95%CI 0.93 to 1.09)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - o People with disabilities
 - Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Located within 75 m of major traffic (Grade F.1.6.7)

- This evidence review found high quality evidence from 1 study with 5,341 children showing that the location of the dwelling within 75 metres of a major road was associated with
 - o asthma (lifetime) aOR 1.29 (95%CI 1.01 to 1.66)
 - o wheeze aOR 1.40 (95%CI 1.09 to 1.78)
- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - Older people
 - People with disabilities
 - Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Located within 100 m of major traffic (Grade F.1.6.8)

- This evidence review found moderate quality evidence from 1 study with 397 adults who
 had received a lung transplant that the location of the dwelling within 100 metres of a
 major road was associated with chronic lung allograft dysfunction aHR 4.72 (95%CI 2.13
 to 10.47)
- This evidence review found low quality evidence from 1 study with 6,339 adults showing that the location of the dwelling within 100 metres of a major road was not associated with
 - o chronic cough aOR 1.22 (95%CI 0.89 to 1.66)
 - wheeze aOR 1.02 (95%CI 0.84 to 1.25)
 - asthma aOR 1.18 (95%CI 0.95 to 1.46)
- This evidence review found low quality evidence from 1 study with 3,309 pregnant women that the location of the dwelling within 100 metres of a major road was not associated with
 - o pre-eclampsia aRR 0.46 (95%CI 0.16 to 1.29),
 - o placental abruption aRR 1.75 (95%CI 0.82 to 3.76)
 - o small for gestational age aRR 0.91 (95%Cl 0.63 to 1.31)
 - stillbirth aRR 1.71 (95%CI 0.56 to 5.23)
- This evidence review found high quality evidence from 1 study with 4,494 adults showing that the location of the dwelling within 100 metres of a major road was associated with coronary artery calcification aOR 1.45 (95%CI 1.15 to 1.82)
- This evidence review found high quality evidence from 1 study with 3,607 adults showing that the location of the dwelling within 100 metres of a major road was associated with diabetes incidence aRR 1.37 (95%CI 1.04 to 1.81)
- This evidence review found moderate quality evidence from 1 study with 121,700 women showing that the location of the dwelling within 100 metres of a major road was not associated with Incident hypertension aHR 1.01 (95%CI 0.88 to 1.15)
- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - Older people
 - People with disabilities
 - People with conditions associated with or exacerbated by indoor air pollution

Located within 150 m of major traffic (Grade F.1.6.9)

- This evidence review found that low quality evidence from 1 study with 2,644 adults showing that the location of the dwelling within 150 metres of a major road was not associated with
 - o wheezing in the last year aOR 0.86 (95%CI 0.68 to 1.08),
 - o COPD aOR 0.97 (95%CI 0.68 to 1.37)
 - o bronchial hyper-responsiveness aOR 0.92 (95%Cl 0.68 to 1.24)
 - o allergic sensitization aOR 0.87 (95%CI 0.70 to 1.07)
- This evidence review found moderate quality evidence from 1 study with 68,195 children showing that the location of the dwelling within 150 metres of a major road was not associated with asthma
 - o aOR 1.03 (95%CI 0.98 to 1.09) for pre-school age children
 - o aOR 1.04 (95%CI 0.92 to 1.16) for school age children.
- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - o Older people
 - o People with disabilities
 - Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Located within 200 m of major traffic (Grade F.1.6.10)

- This evidence review found high quality evidence from 1 study with 36,294 women showing that the location of the dwelling within 200 metres of a major road was associated with infertility aHR 1.11 (95%CI 1.02 to 1.20)
- This evidence review found high quality evidence from 1 study with 1,405 adults showing that the location of the dwelling within 200 metres of a major road was associated with wheeze aOR1.38 (95%CI 1.06 to 1.80)
- This evidence review found moderate quality evidence from 1 study with 1,405 adults showing that the location of the dwelling within 200 metres of a major road was not associated with asthma aOR1.21 (95%CI 0.91 to 1.59)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - Children and young people
 - o People with conditions associated with or exacerbated by indoor air pollution

Individual characteristics and health outcomes

Tenancy status (Grade F.1.7.1)

- This evidence review found moderate quality evidence from 1 study with 7,640 adults showing that tenancy status was associated with
 - o eye irritation aOR 2.07 (95%CI 1.19 to 3.58) for rented (versus owner) status
 - o nasal irritation aOR 2.07 (95%Cl 1.33 to 3.20) for rented (versus owner) status
- This evidence review found that low quality evidence from 1 study with 7,640 adults showing that tenancy status was not associated with
 - o cough aOR 1.85 (95%CI 0.94 to 3.65) for rented (versus owner) status
 - o throat irritation aOR 1.98 (95%CI 0.98 to 3.97) for rented (versus owner) status
- · No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - Children and young people
 - o People with conditions associated with or exacerbated by indoor air pollution

Household occupant density (Grade F.1.7.2)

- This evidence review found low quality evidence from 1 study with 416 infants that higher occupancy (4 or more people per household) was associated with bronchiolitis requiring hospitalisation aOR1.75 (95%Cl 1.36 to 2.13)
- This evidence review found moderate quality evidence from 1 study with 2,779 children showing that occupancy was not associated with newly diagnosed asthma aOR 1.48 (95%CI 0.86 to 2.57) for multi-family household
- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - Older people
 - People with disabilities
 - Pregnant women
 - People with conditions associated with or exacerbated by indoor air pollution

Socio-economic status (SES) (Grade F.1.7.3)

- This evidence review found very low quality evidence from 1 study with 1,191 children that socio-economic status (SES) was associated with cough
 - o aOR 1.53 (95%Cl 1.12 to 2.10) for middle SES (compared to high SES)
 - o aOR 3.37 (95%Cl 2.01 to 5.71) for low SES (compared to high SES)
- This evidence review found very low quality evidence from 1 study with 1,191 children that SES was associated with asthmatic symptoms
 - o aOR 1.43 (95%CI 1.00 to 2.04) for middle SES compared to high SES households

- o aOR 3.32 (95%CI 1.88 to 5.93) for low SES compared to high SES household
- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - o Older people
 - o People with disabilities
 - o Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Association between exposure levels and health outcomes

Public health evidence

Summary of public health studies included in the evidence review

A summary of the characteristics of the included studies are in the following table

Table 3: Characteristics of included studies

Study (country)	Design	Population	Exposure	Outcomes	Risk of bias
1. Belanger 2003 (US)	Prospective cohort	Infants	NO ₂	Cough, Wheeze	Moderate
2. Belanger 2013 (US)	Prospective cohort	Children	NO ₂	Asthma, Wheeze	Low
3. Bertelsen 2010	Prospective cohort	Children	Cat allergen	Asthma, bronchial hyperresponsiveness	Low
4. Brussee 2005 (The Netherlands)	Prospective cohort	Children	House dust	Asthma, Wheeze	Low
5. Casas 2015 (4 EU countries)	Prospective cohort	Children	HDM allergen	Asthma, Persistent wheeze	Low
6. Cho 2006 (US)	Prospective cohort	Infants	House dust	Wheeze	Moderate
7. Cole Johnson 2004 (US)	Prospective cohort	Children	Allergen	Asthma	Low
8. Cullinan 2004 (UK)	Prospective cohort	Children	Allergen	Wheeze	Low
9. Dales 1991	Retrospective cohort	Adults	Damp / mould	Respiratory symptoms, chronic respiratory disease, asthma, eye irritation	High
10.Diez 2002 (Germany)	Nested case control	Children at risk of allergies	VOCs	Pulmonary infections	Moderate
11.Emenius 2004 (Sweden)	Nested case control	Children	NO ₂	Recurrent wheezing	Moderate
12.Gent 2009 (US)	Prospective cohort	Children with asthma	Allergens	Wheeze, Asthma exacerbations	Low
13.Gent 2012 (US)	Prospective cohort	Children with asthma	Allergens	Wheeze, Persistent cough, Asthma exacerbations	Moderate
14.Habre 2014 (US)	Prospective cohort	Children with asthma	PM _{2.5}	Cough, Wheeze	Moderate
15.Hansel 2008 (US)	Prospective cohort	Children with asthma	NO ₂	Asthma symptoms	Low
16.Harris 2007 (UK)	Prospective cohort	Children	Allergens	Eczema	Low
17.Hunt 2011 (US)	Cohort	Infants at risk of asthma	PM _{2.5}	Wheeze	Low
18.lossifova 2009 (US)	Prospective cohort	Children	Mould	Wheeze	Low

Study (country)	Design	Population	Exposure	Outcomes	Risk of bias
19.Jedrychowski 2011 (Poland)	Prospective cohort	Infants	PM _{2.5}	Eczema	Moderate
20.Jung 2012 (US)	Prospective cohort	Children	PAHs	Asthma, Wheeze	Moderate
21.Jung 2012 b (US)	Prospective cohort	Children	PM _{2.5}	Wheeze	Moderate
22.Jung 2014 (US)	Prospective cohort	Children	VOCs	Asthma	Moderate
23.Lau 2000 (Germany)	Prospective cohort	Children at risk of asthma	Allergens	Asthma, Wheeze	Low
24.Litonjua 2002 (US)	Prospective cohort	Children at risk of atopy	Allergens	Wheeze	Moderate
25.Lynch 2014 (US)	Prospective cohort	Children at risk of atopy	Allergens	Wheeze	Low
26.McCormack 2009 (US)	Prospective cohort	Children with asthma	Particulate matter	Asthma symptoms	Moderate
27.0'Connor 2017 US)	Prospective cohort	Children at risk of asthma	Allergens, NO ₂	Asthma	Low
28.Raaschou-Nielsen 2010 (Denmark)	Prospective cohort	Infants at risk of asthma	Particulate matter, NoO2, Formaldehyde	Wheezing	Moderate
29.Roda 2011 (France)	Prospective cohort	Infants	PM _{2.5}	Cough, Wheeze without cold, asthma, Allergic rhinitis, Breathlessness, Chest tightness, Cardiac disease, Hypertension, Stroke, Heart infarction or angina pectoris	Moderate
30.Stark 2003 (US)	Prospective cohort	Infants	Fungal spores	Lower respiratory illness	Low
31.Torrent 2007	Prospective cohort	Infants	Allergens, Nitrogen dioxide	Asthma, wheeze	Low

See appendix D for full evidence tables.

Quality assessment of clinical studies included in the evidence review

See appendix F for full GRADE tables.

Economic evidence

No economic evidence review was carried out for this review

Economic model

No economic evidence modelling was carried out for this review

Evidence statements

Damp

Damp (Grade F.2.1.1)

- This evidence review found low quality evidence from 1 study with 7,320 adults showing that exposure to marked dampness was associated with
 - o cough and phlegm aRR 2.73 (95%CI 1.88 to 3.99)
 - o phlegm aOR 2.05 (95%CI 1.07 to 3.91)
- This evidence review found very low quality evidence from 1 study of 7,320 adults showing that exposure to marked dampness was not associated with cough aRR 0.85 (95%CI 0.67 to 1.09)
- This evidence review found very low quality evidence from 1 study of 7,320 adults showing that exposure to slight to moderate dampness was not associated with
 - o Cough aRR 1.26 (95%CI 0.80 to 1.99)
 - o cough and phlegm aRR.1.24 (95%CI 0.99 to 1.56)
 - o phlegm aOR 0.82 (95%CI 0.54 to 1.27)
- This evidence review found moderate quality evidence from 1 study with 369 children showing that exposure to damp or mould was associated with wheeze IRR^a 1.67 (95%CI 1.39 to 2.01)
- This evidence review found moderate quality evidence from 1 study with 322 infants showing that exposure to damp or mould was associated with wheeze aHR 1.22 (95%CI 1.07 to 1.40)
- This evidence review found moderate quality evidence from 1 study with 7,104 adults (showing that exposure to damp was associated with asthma aRR 1.49 (95%CI 1.00 to 2.22)
- This evidence review found moderate quality evidence from 1 study with 398 children showing that exposure to major moisture damage or any damage with mould in the child's main living area was not associated with cough aOR 1.27 (95%CI 0.77 to 2,09)
- This evidence review found low quality evidence from 1 study with 14,799 adults showing that exposure to dampness and mould was associated with
 - o chronic respiratory disease aOR 1.45 (95%CI 1.29 to 1.64)
 - lower respiratory symptoms aOR 1.62 (95%CI 1.48 to 1.78)

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^a IRR: incidence rate ratio

- o upper respiratory symptoms aOR 1.50 (95%CI 1.38 to 1.61)
- o asthma aOR 1.56 (95%CI 1.25 to 1.95)
- This evidence review found high quality evidence from 1 study with 1,765 infants showing that exposure to damp was associated with
 - wheeze aOR 2.12 (95%CI 1.30 to 3.46)
 - o asthma aOR 2.19 (95%CI 1.06 to 4.53)
- This evidence review found moderate quality evidence from 1 study with 1,765 adults showing that exposure to damp was not associated with bronchiolitis aOR 1.32 (95%CI 0.80 to 2.18)
- This evidence review found low quality evidence from 1 study with 4,625 children showing that exposure to damp was associated with
 - o cough aOR 2.16 (95%CI 1.64 to 2.84)
 - o asthma aOR 1.42 (95%CI 1.04 to 1.94)
 - o bronchitis aOR 1.32 (95%CI 1.05 to 1.67)
 - o chest illness aOR 1.52 (95%CI 1.20 to 1.93)
 - o non-chest illness aOR 1.55 (95%CI 1.25 to 1.93)
 - o wheeze aOR 1.23 (95%CI 1.10 to 1.39)
 - o lower respiratory illness aOR 1.68 (95%CI 1.41 to 2.01)
 - o upper respiratory illness aOR 1.57 (95%CI 1.31 to 1.74)
- This evidence review found low quality evidence from 1 study with 400 children showing that exposure to damp was not associated with rhinitis
 - o aOR 1.39 (95%CI 0.73 to 2.67) for mild damp stains,
 - o aOR 0.37 (95%CI 0.04 to 3.43) for severe damp stains,
 - o aOR 1.16 (95%CI 0.36 to 3.76) for mild floor damp,
 - aOR 1.58 (95%Cl 0.10 to 26.14) for severe floor damp
- This evidence review found moderate quality evidence from 1 study with 5,078 children showing that exposure to damp was not associated with
 - wheeze aOR 1.11 (95%CI 0.87 to 1.43)
 - o asthma aOR 1.16 (95%CI 0.87 to 1.53)
- This evidence review found low quality evidence from 1 study with 6,853 children showing that exposure to damp was not associated with acute respiratory infection requiring hospitalisation
 - o aHR 0.95 (95%CI 0.82 to 1.11) for infrequent dampness in the house,
 - o aHR 1.08 (95%CI 0.91 to 1.29) for frequent dampness in the house,
 - o aHR 1.07 (95%CI 0.84 to 1.37) for always dampness in the house
- This evidence review found moderate quality evidence from 1 study with 6,853 children showing that exposure to mould was protective against acute respiratory infection requiring hospitalisation
 - aHR 0.81 (95%CI 0.67 to 0.99) for mould or mildew in the walls or ceilings of the room where the child sleeps at night in the past 2 weeks
- This evidence review found moderate quality evidence from 1 study with 6,853 children showing that exposure to mould was not associated with acute respiratory infection requiring hospitalisation
 - aHR 0.81 (95%CI 0.67 to 0.99) for mould or mildew in the walls or ceilings of the room where the child sleeps at night in the past 2 weeks,
 - o aHR 1.08 (95%CI 0.91 to 1.29) for frequent dampness in the house,
 - o aHR 1.07 (95%Cl 0.84 to 1.37) for always dampness in the house

- This evidence review found low quality evidence from 1 study of 528 children with asthma showing that exposure to damp was associated with airway hyper-responsiveness aOR 3.95 (95%CI 1.82 to 8.57)
- This evidence review found low quality evidence from 1 study with 593 infants at risk of asthma showing that exposure to mould or mildew was not associated with wheeze aOR 1.22 (95%CI 0.80 to 1.88)
- This evidence review found moderate quality evidence from 1 study with 499 infants at risk of asthma or allergy showing that exposure to water damage, mould or mildew was not associated with lower respiratory illness aOR 1.34 (95%CI 0.99 to 1.82)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - People with conditions associated with or exacerbated by indoor air pollution

Mould (Grade F.2.1.2)

- This evidence review found high quality evidence from 1 study with 593 infants not at risk of asthma showing that exposure to mould or mildew was associated with cough aOR 1.53 (95%CI 1.01 to 2.30)
- This evidence review found moderate quality evidence from 1 study with 256 infants at risk of allergy showing that exposure to mould or mildew was associated with
 - Wheeze aOR 2.51 (95%CI 1.37 to 4.62)
 - o Cough aOR 1.91 (95%CI 1.07 to 3.42)
- This evidence review found low quality evidence from 1 study with 593 infants not at risk of asthma showing that exposure to mould or mildew was not associated with wheeze aOR 1.22 (95%CI 0.80 to 1.88)
- This evidence review found moderate quality evidence from 1 study with 7,104 adults (showing that exposure to mould was not associated with asthma aRR 1.15 (95%CI 0.71 to 1.85)
- The evidence review found low quality evidence from 1 study with 1916 children showing that exposure to mouldy odour was associated with asthma aOR 2.44 [95%CI 1.07 to 5.60)
- This evidence review found low quality evidence from 1 study with 4,625 children showing that exposure to mould was associated with
 - o non-chest illness aOR 1.40 (95%CI 1.13 to 1.74)
 - o chest illness aOR 1.40 (95%CI 1.11 to 1.78)
 - bronchitis aOR 1.48 (95%CI 1.17 to 1.87)
 - o cough aOR 2.12 (95%CI 1.64 to 2.73)
 - hay fever aOR 1.57 (95%CI 1.31 to 1.74)
 - o lower respiratory illness aOR 1.57 (95%CI 1.31 to 1.87)
 - o wheeze aOR 1.79 (95%CI 1.44 to 2.32)
- This evidence review found moderate quality evidence from 1 study with 4,625 children showing that exposure to mould was not associated with asthma aOR 1.27 (95%CI 0.93 to 1.74)
- This evidence review found low quality evidence from 1 study with 400 children showing that exposure to mild mould was not associated with eczema aOR 1.39 (95%CI 0.73 to 2.67)
- This evidence review found low quality evidence from 1 study with 400 children showing that exposure to severe mould was not associated with eczema aOR 0.37 (95%CI 0.04 to 3.43)

- This evidence review found high quality evidence from 1 study with 483 children with atopy showing that exposure to high levels of visible mould was associated with wheeze aOR 6.16 (95%CI 1.38 to 27.44)
- This evidence review found moderate quality evidence from 1 study with 483 children with atopy showing that exposure to low levels of visible mould was not associated with wheeze aOR 1.86 (95%CI 0.86 to 4.00)
- This evidence review found low quality evidence from 1 study with 1,916 children showing that exposure to mould odour was associated with asthma aOR 2.44 (95%Cl 1.07 to 5.60)
- This evidence review found very low quality evidence from 1 study with 1,916 children showing that exposure to visible mould was associated with asthma aOR 0.65 (95%Cl 0.24 to 1.72)
- This evidence review found moderate quality evidence from 1 study with 3,798 children showing that exposure to mould was not associated with rhinitis
 - Asthma aOR 1.16 (95%CI 0.93 to 1.44) for single indicator of damp (mould odour or visible mould or dampness damage)
 - o aOR 1.03 (95%Cl 0.87 to 1.22) for a single indicator of damp (mould odour or visible mould or dampness damage),
 - aOR 1.18 (95%CI 0.92 to 1.52) for two indicators of damp (mould odour or visible mould or dampness damage and
 - aOR 1.23 (95%CI 0.82 to 1.85) for three indicators of damp (mould odour or visible mould or dampness damage)
- This evidence review found moderate quality evidence from 1 study with 3,798 children showing that exposure to mould was associated with asthma
 - aOR 1.37 (95%CI 1.01 to 1.86) for two indicators of damp (mould odour or visible mould or dampness damage)
 - aOR 1.73 (95%CI 1.10 to 2.74) for three indicators of damp (mould odour or visible mould or dampness damage)
- This evidence review found moderate quality evidence from 1 study with 398 children showing that exposure to exposure to minor moisture damage with or without mould spots in child's main living area was not associated with asthma aOR 1.31 (95%CI 0.72 to 2.36)
- This evidence review found moderate quality evidence from 1 study with 398 children showing that exposure to major moisture damage or any moisture damage with visible mould in child's main living area was not associated with asthma aOR 1.33 (95%CI 0.60 to 2.98)
- This evidence review found moderate quality evidence from 1 study with 3,535 children with no history of asthma showing that exposure to mould was not associated with asthma aRR 1.10 (95%CI 0.80 to 1.60)
- This evidence review found very low quality evidence from 1 study with 1916 children showing that exposure to visible mould was not associated with asthma aOR 0.65 (95%CI 0.24 to 1.72)
- This evidence review found low quality evidence from 1 study with 1,863 children showing that exposure to mould on walls or visible mould was associated with hay fever (reported as allergic rhinitis) aOR 1.73 (95%Cl 1.27 to 2.38) and aOR 1.98 (95%Cl 1.32 to 2.99) respectively
- This evidence review found moderate quality evidence from 1 study with 405 children showing that exposure to water damage or mould/mildew in past year was not associated with allergic rhinitis aOR 1.66 (95%CI 0.88 to 3.15)
- This evidence review found very low quality evidence from 1 study with 3,360 children showing that exposure to mould on walls was not associated with allergic rhinoconjunctivitis aOR 1.34 (95%CI 0.64 to 2.79)
- This evidence review found high quality evidence from 1 study with 3,535 children with wheeze
 at baseline showing that exposure to mould was protective against asthma aRR 0.60 (95%CI
 0.40 to 0.90)

- This evidence review found low quality evidence from 1 study with 14,799 adults showing that exposure to damp or mould was associated with eye irritation aOR 1.63 (95%CI 1.46 to 1.82)
- This evidence review found low quality evidence from 1 study with 9,808 adults showing that mouldy odour was associated with
 - o eye irritation aOR 3.75 (95%CI 3.60 to 3.92)
 - o tiredness aOR 2.38 (95%CI 2.31 to 2.46)
 - headache aOR 3.37 (95%CI 3.24 to 3.51)
 - o facial skin symptoms aOR 2.93 (95%Cl 2.80 to 3.06)
 - o cough aOR 3.30 (95%CI 3.16 to 3.46)
 - o nasal symptoms aOR 2.83 (2.73 to 2.93)
 - o throat symptoms aOR 3.48 (3.33 to 3.62)
 - o tiredness aOR 2.58 (95%CI 2.31 to 2.46)
- This evidence review found low quality evidence from 1 study with 1,719 children showing that exposure to dampness or visible mould was associated with
 - o any sleep problems aOR 1.80 (95%Cl 1.22 to 2.66)
 - o problems sleeping throughout the night aOR 2.36 (95%Cl 1.15 to 4.84)
 - o sleep less than 9 hours aOR 1.60 (95%CI 1.02 to 2.51)
- This evidence review found low quality evidence from 1 study with 1,719 children showing that exposure to dampness or visible mould was not associated with problems falling asleep aOR 1.50 (95%CI 0.98 to 2.30)
- This evidence review found low quality evidence from 1 study with 1,719 children showing that exposure to visible mould was associated with
 - o Any sleep problems aOR 1.70 (95%CI 1.13 to 2.54)
 - Sleep less than 9 hours aOR 1.67 (85%CI 1.06 to 2.65)
- This evidence review found low quality evidence from 1 study with 1,719 children showing that exposure to visible mould was not associated with problems falling asleep aOR 1.50 (95%CI 0.97 to 2.33)
- This evidence review found very low quality evidence from 1 study with 1,719 children showing that exposure to mould was not associated with
 - o problems falling asleep aOR 1.50 (95%Cl 0.97 to 2.33)
 - o problems sleeping throughout the night aOR 1.91 (95%Cl 0.89 to 4.13)
- This evidence review found low quality evidence from 1 study with 398 children showing that exposure to mould or mildew was not associated with wheeze aOR 1.34 (95%Cl 0.90 to 2.01)
- This evidence review found low quality evidence from 1 study with 6,853 children showing that exposure to mould was protective against acute respiratory infection aOR 0.81 (95%CI 0.67 to 0.99)
- This evidence review found low quality evidence from 1 study with 1,002 infants at risk of asthma showing that exposure to mould was not associated with earache aOR 1.37 (95%CI 0.94 to 2.02)
- This evidence review found moderate quality evidence from 1 study with 813 infants showing that exposure to mould was not associated with
 - o any episode of earache aOR 1.15 (95%Cl 0.87 to 1.99)
 - recurrent earache (four or more episodes separated by 21 days in 1 year) aOR 1.05 (95%CI 0.88 to 1.26)
- · No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people

- People with disabilities
- o Pregnant women

Fungal spore levels (Grade F.2.1.3)

- This evidence review found high quality evidence from 1 study with 499 infants at risk of asthma showing that exposure to damp/mould/fungi was associated with lower respiratory illness aOR 1.86 (95%CI 1.21 to 2.88) for greater than 90th percentile for specific taxon
- This evidence review found low quality evidence from 1 study with 1,233 children with asthma showing that exposure to Cladosporium >148 CFU/m³ was not associated with
 - o cough aOR 0.98 (95%CI 0.54 to 1.80)
 - o wheeze aOR 1.22 (95%CI 0.66 to 2.26)
 - o asthma severity aOR 1.58 (95%CI 0.88 to 2.83)
 - o asthma exacerbations (reported as rescue medication use) aOR 0.69 (95%CI 0.37 to 1.29)
- This evidence review found low quality evidence from 1 study with 1,002 infants at risk of asthma showing that exposure to fungi was not associated with earache in the 1st 6 months of life
 - o aOR 1.27 (95%CI 0.56 to 2.86) for penicillium ≥1000 CFU/m³,
 - aOR 1.09 (95%CI 0.52 to 2.29) for Cladosporium ≥1000 CFU/m³
- This evidence review found moderate quality evidence from 1 study with 1,002 infants at risk of asthma showing that exposure to other mould (not yeast, penicillium or cladosporium) ≥1,000 CFU/m³ was associated with earache in the 1st 6 months of life aOR 3.45 (95%CI 1.36 to 8.76)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - People with disabilities
 - Pregnant women

Formaldehyde (Grade F.2.2)

- This evidence review found moderate quality evidence from 1 study with 2,940 infants showing that exposure to elevated levels of formaldehyde was associated with
 - lower respiratory tract infections aOR 1.32 (95%Cl 1.11 to 1.55) per IQR increase in formaldehyde
 - lower respiratory tract infection with wheeze aOR 1.41 (95%CI 1.14 to 1.74) per IQR increase in formaldehyde
- This evidence review found very low quality evidence from 1 study with 9,808 infants at risk of asthma showing that exposure to elevated levels of formaldehyde was not associated with wheeze
 - aOR 1.11 (95%Cl 0.47 to 2.63) for formaldehyde levels between 12.4 and 16.3 μg/m³ compared to less than 12.4,
 - $\circ~$ aOR 1.21 (95%Cl 0.51 to 2.92) for formaldehyde levels between 16,3 and 20.3 $\mu g/m^3$ compared to less than 12.4,
 - $_{\odot}~$ aOR 1.40 (95%Cl 0.57 to 3.47) for formaldehyde levels between 20.3 and 25.6 $\mu g/m^3$ compared to less than 12.4
 - $\circ~$ aOR 0.67 (95%Cl 0.29 to 1.54) for formaldehyde levels greater than 25.6 $\mu g/m^3$ compared to less than 12.4

- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - People with conditions associated with or exacerbated by indoor air pollution

Allergens

House dust mites (Der p 1 and / or Der f 1) (Grade F.2.3.1)

- This evidence review found low quality evidence from 1 study with 1,233 children with showing that exposure to house dust mite allergens was not associated with asthma exacerbations
 - o aOR 1.19 (95%Cl 0.92 to 1.55) for Der p 1 + Der f 1 >10 μg/g
- This evidence review found high quality evidence from 1 study with 1,879 infants showing that exposure to house dust mite allergens (levels not specified) was associated with wheeze aOR 1.39 (95%Cl 1.12 to 1.73)
- This evidence review found low quality evidence from 1 study with 593 infants not at risk of asthma showing that exposure to house dust mite allergens was not associated with wheeze aOR 0.78 (95%CI 0.55 to 1.13) for Der p 1 + Der f 1 ≥ 2 μg/g
- This evidence review found moderate quality evidence from 1 study with 4,334 children showing that exposure to house dust mite allergens was not associated with wheeze, persistent
 - o aOR 1.3 (95%Cl 0.8 to 2.2) for Der p 1 + Der f 1 ≥0.19 μg/g to <0.4 μg/g,
 - aOR 1.1 (95%CI 0.8 to 1.5) for Der p 1 +Der f 1 0.4 to <2 μg/g,
 - o aOR 0.9 (95%Cl 0.7 to 1.3) for Der p 1 + Der f 1 ≥2 μg/g,
- This evidence review found high quality evidence from 1 study with 4,334 children showing that
 exposure to house dust mite allergens was associated with asthma at 6 years of age or younger
 - aOR 1.4 (95%Cl 1.1 to 1.9) for Der p 1 + Der f 1 0.4 to <2 μg/g
- This evidence review found moderate quality evidence from 1 study with 4,334 children showing that exposure to house dust mite allergens was not associated with asthma at 6 years of age or younger
 - aOR 1.6 (95%Cl 0.9 to 2.6) for Der p 1 + Der f 1 ≥0.19 μg/g to <0.4 μg/g,
 - aOR 1.1 (95%Cl 0.8 to 1.6) for Der p 1 + Der f 1 ?2 μg/g,
- This evidence review found moderate quality evidence from 1 study with 4,334 children showing that exposure to house dust mite allergens was not associated with asthma at older than 6 years
 - o aOR 1.3 (95%Cl 0.8 to 2.3) for Der p 1 + Der f 1 ≥0.19 μ g/g to <0.4 μ g/g,
 - aOR 1.1 (95%Cl 0.8 to 1.6) for Der p 1 +Der f 1 0.4 to <2 μg/g,
 - o aOR 1.0 (95%Cl 0.7 to 1.4) for Der p 1 + Der f 1 ?2 μg/g,
- This evidence review found low quality evidence from 1 study with 593 infants not at risk of asthma showing that exposure to house dust mite allergens was not associated with cough aOR 0.76 (95%CI 0.54 to 1.07) for Der p 1 + Der f 1 ≥ 2 µg/g
- This evidence review found low quality evidence from 1 study with 256 infants at risk of asthma showing that exposure to house dust mite allergens was not associated with wheeze aOR 1.04 (95%Cl 0.60 to 1.80) for Der p 1 + Der f 1 ≥ 2 μg/g

- This evidence review found moderate quality evidence from 1 study with 1,314 children at risk of asthma showing that exposure to house dust mite allergens was not associated with wheeze aOR 1.03 (95%CI 0.52 to 2.04) for Der p 1 + Der f 1 between 0.981 – 240µg/g
- This evidence review found low quality evidence from 1 study with 256 infants at risk of asthma showing that exposure to house dust mite allergens was not associated with cough aOR 1.27 (95%CI 0.75 to 2.15) for Der p 1 + Der f 1 ≥ 2 µg/g
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - o People with disabilities
 - o Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Der p 1 (Grade F.2.3.2)

- This evidence review found moderate quality evidence from 1 study with 1,233 children with asthma showing that exposure to house dust mite allergens was associated with
 - $_{\odot}$ asthma exacerbations (reported as rescue mediation use) aOR 1.47 (95%Cl 1.11 to 1.94) for exposure to Der p 1 >0.10 $\mu g/g$
 - $_{\odot}$ asthma exacerbations (reported as moderate or severe GINA score) aOR 2.93 (95%CI 1.37 to 6.30) for Der p 1 (μ g/g) 2.0 to <10.0 vs <0.10 in main living area
- This evidence review found low quality evidence from 1 study with 1,223 children with asthma showing that exposure to house dust mite allergens was not associated with
 - o cough aOR 1.18 (95%Cl 0.90 to 1.55) for Der p 1 >0.10 μg/g
 - wheeze aOR 1.26 (95%CI 0.95 to 1.67) for Der p 1 >0.10 μg/g
 - \circ asthma exacerbations aOR 1.19 (95%Cl 0.92 to 1.55) for Der p 1 >0.10 μ g/g
- This evidence review found moderate quality evidence from 1 study with 4,334 children showing that exposure to house dust mite allergens was not associated with wheeze, persistent
 - aOR 1.1 (95%Cl 0.7 to 1.8) for Der p 1 ≥0.12 to <0.4 μg/g,
 - o aOR 0.9 (95%CI 0.7 to 1.3) for Der p 1 > 0.4 to $<2 \mu g/g$,
 - aOR 0.8 (95%Cl 0.5 to 1.1) for Der p 1 ≥2 μg/g
- This evidence review found moderate quality evidence from 1 study with 4,334 showing that exposure to house dust mite allergens was not associated with asthma at 6 years of age or younger
 - $\circ~$ aOR 1.4 (95%Cl 0.9, 2.3) for Der p 1 low to < 2 $\mu g/g$
 - o aOR 1.1 (95%Cl 0.8 to 1.6) for Der p 1 0.4 to $< 2 \mu g/g$,
 - o aOR 1.0 (95%Cl 0.7 to 1.5) for Der p 1 > 2 μ g/g,
- This evidence review found moderate quality evidence from 1 study with 4,334 showing that exposure to house dust mite allergens was not associated with asthma at 6 years of age or older
 - \circ aOR 1.1 (95%Cl 0.8 to 1.6) for Der p 1 0.4 to < 2 μ g/g
 - o aOR 0.7 (95%Cl 0.4 to 1.0) for Der p 1 > 2 μ g/g,
 - o aOR 1.4 (95%CI 0.9 to 2.3) for Der p 1 >1.9 μ g/g to <0.4 μ g/g
- This evidence review found moderate quality evidence from 1 study with 1,611 infants showing that exposure to house dust mite allergens was not associated with asthma
 - aOR 0.67 (95%CI 0.40 to 1.12) for Der p 1 0.83 to 6.46 μg/g

- aOR 0.68 (95%Cl 0.37 to 1.25) for Der p 1 >6.46 μg/g
- This evidence review found moderate quality evidence from 1 study with 593 children showing that exposure to house dust mite allergens was not associated with eczema
 - o aOR 1.01 (95%CI 0.53 to 1.92) for Der p 1 0.28 to 0.81 units
 - o aOR 1.37 (95%Cl 0.74 to 2.55) for Der p 1 0.82 to 2.22 units
 - aOR 0.66 (95%Cl 0.34 to 1.29) for Der p 1 2.23 to 7.75 units
 - o aOR 0.71 (95%CI 0.37 to 1.37) for Der p 1 7.76 to 384.97 units
- This evidence review found moderate quality evidence from 1 study with 593 children showing that exposure to house dust mite allergens was not associated with visible flexural dermatitis
 - o aOR 1.17 (95%CI 0.58 to 2.34) for Der p 1 0.28 to 0.81 units
 - o aOR 1.73 (95%Cl 0.87 to 3.46) for Der p 1 0.82 to 2.22 units
 - o aOR 0.88 (95%Cl 0.43 to 1.81) for Der p 1 2.23 to 7.75 units
 - o aOR 0.96 (95%CI 0.47 to 1.94) for Der p 1 7.76 to 384.97 units
- This evidence review found moderate quality evidence from 1 study with 300 children with asthma showing that exposure to house dust mites was associated with
 - o wheeze aOR 3.58 (95%Cl 1.28 to 9.97) for Der p 1 (μg/g) ≥10.0 vs <0.10 in bed
 - o asthma exacerbations (reported as controller medication for 9 months or more) aOR 2.52 (95%Cl 1.17 to 5.42) for Der p 1 (μ g/g) 2.0 to <10.0 vs <0.10 in main living area
 - $_{\odot}$ asthma exacerbations (reported as controller medication for 9 months or more) aOR 2.73 (95%Cl 1.32 to 5.64) for Der p 1 (μ g/g) 2.0 to <10.0 vs <0.10 in main living area
 - o asthma exacerbations (reported as moderate or severe GINA score) aOR 2.55 (95%Cl 1.13 to 5.73) for Der p 1 (μg/g) ≥10.0 vs <0.10 in main living area
 - $_{\odot}$ asthma exacerbations (reported as moderate or severe GINA score) aOR 2.93 (95%CI 1.37 to 6.30) for Der p 1 (µg/g) 2.0 to <10.0 vs <0.10 in bed
 - o asthma exacerbations (reported as controller medication for 9 months or more) aOR 2.16 (95%CI 1.04 to 4.48) for Der p 1 (μ g/g) 2.0 to <10.0 vs <0.10 in bed
- This evidence review found moderate quality evidence from 1 study with 300 children with asthma showing that exposure to house dust mite allergens was not associated with
 - $\circ~$ Wheeze aOR 1.05 (95%Cl 0.38 to 2.84) for Der p 1 (µg/g) 0.10 to <2.0 vs <0.10 in main living area,
 - \circ Wheeze aOR 1.55 (95%Cl 0.52 to 3.85) for Der p 1 (µg/g) 2.0 to <10.0 vs <0.10 in main living area,
 - Wheeze aOR 2.01 (95%Cl 0.75 to 5.19) for Der p 1 (μg/g) >10.0 vs <0.10 in main living area,
 - \circ Wheeze aOR 1.70 (95%Cl 0.65 to 4.22) for Der p 1 (µg/g) 0.10 to <2.0 vs <0.10 in bed,
 - Wheeze aOR 1.60 (95%Cl 0.64 to 4.00) for Der p 1 (μg/g) 2.0 to <10.0 vs <0.10 in bed
 - asthma exacerbations (reported as moderate or severe GINA score) aOR 0.93 (95%CI 0.41 to 2.10) for Der p 1 (μg/g) 0.10 to <2.0 vs <0.10 in main living area
 - o asthma exacerbations (reported as controller medication for 9 months or more) aOR 0.61 (95%Cl 0.27 to 1.35) for Der p 1 (μ g/g) 0.10 to <2.0 vs <0.10 in main living area
 - o asthma exacerbations (reported as controller medication for 9 months or more) aOR 2.17 (95%CI 0.97 to 4.86) for Der p 1 (μ g/g) >0.10 in main living area
 - asthma exacerbations (reported as moderate or severe GINA score) aOR 0.99 (95%CI 0.47 to 2.08) for Der p 1 (μg/g) 0.10 to <2.0 vs <0.10 in bed
 - $_{\odot}$ asthma exacerbations (reported as controller medication for 9 months or more) aOR 1.35 (95%Cl 0.66 to 2.73) for Der p 1 (µg/g) 0.10 to <2.0 vs <0.10 in bed
 - $_{\odot}$ asthma exacerbations (reported as moderate or severe GINA score) aOR 1.19 (95%CI 0.46 to 3.03) for Der p 1 (μ g/g) 10.0 to <2.0 vs <0.10 in bed

- asthma exacerbations (reported as controller medication for 9 months or more) aOR 1.41 (95%Cl 0.57 to 3.46) for Der p 1 (μg/g) ≥10.0 to <2.0 vs <0.10 in bed
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - People with disabilities
 - o Pregnant women

Der f 1 (Grade F.2.3.3)

- This evidence review found low quality evidence from 1 study with 1,233 children with asthma showing that exposure to house dust mite allergens (Der f 1 >2.10 μg/g) was not associated with
 - asthma exacerbations (reported using Asthma Severity Index) aOR 1.28 (95%CI 0.94 to 1.74)
 - o asthma exacerbations (reported as rescue medication use) aOR 1.09 (95%CI 0.78 to 1.51)
- This evidence review found low quality evidence from 1 study with 1,223 children with asthma showing that exposure to house dust mite allergens was not associated with
 - \circ wheeze aOR 0.89 (95%Cl 0.63 to 1.24) for Der f 1 >2.1 μ g/g
 - \circ cough aOR 0.90 (95%Cl 0.65 to 1.25) for Der f 1 >2.1 μ g/g
- This evidence review found moderate quality evidence from 1 study with 4,334 children showing that exposure to house dust mite allergens was not associated with wheeze, persistent
 - aOR 1.2 (95%CI 0.8 to 1.8) for Der f 1 ≥0.07 to <0.4 μg/g,
 - \circ aOR 1.1 (95%Cl 0.8 to 1.5) for Der f 1 0.4 to <2 μ g/g,
 - aOR 1.1 (95%Cl 0.8 to 1.6) for Der f 1 ≥2 μg/g,
- This evidence review found moderate quality evidence from 1 study with 4,334 showing that exposure to house dust mite allergens was not associated with asthma at 6 years of age
 - o aOR 1.2 (95%CI 0.8 to 1.8) for Der f 1 ≥2 μ g/g,
 - aOR 1.2 (96%CI 0.8 to 1.8) for Der f 1 ≥0.07 to < 2 μg/g
 - \circ aOR 1.2 (95%Cl 0.8 to 1.6) for Der f 1 1 0.4 to < 2 μ g/g
- This evidence review found moderate quality evidence from 1 study with 4,334 showing that exposure to house dust mite allergens was not associated with asthma at 6 years of age or older
 - o aOR 1.0 (95%CI 0.7 to 1.6) for Der f 1 > 2 μ g/g,
 - o aOR 1.0 (96%CI 0.7 to 1.6) for Der f 1 0.07 ≥ to < 2 μ g/g
 - \circ aOR 1.0 (95%Cl 0.7 to 1.4) for Der f 1 1 0.4 to < 2 μ g/g
- This evidence review found high quality evidence from 1 study with 4,334 children showing that exposure to house dust mite allergens was not associated with asthma at 6 years of age or older
 - aOR 1.1 (95%CI 0.8 to 1.6) for Der f 1 0.4 to <2 μg/g)
 - o aOR 1.0 (95%CI 0.7 to 1.5) for Der f 1 > 2 μ g/g
 - aOR 1.2 (95%Cl 0.8 to 1.8) for Der f 1 > 0.07 to < 0.4 μg/g)
 - aOR 1.2 (95%CI 0.8 to 1.6) for Der f 1 0.4 to <2 μg/g
- This evidence review found showing moderate quality evidence from 1 study with 442 children
 at risk of developing asthma that exposure to house dust mite allergens (per interquartile
 increase in Der f 1) at 3 months of age was not associated with asthma aOR 0.98 (95%CI 0.91
 to 1.04)

- This evidence review found low quality evidence from 1 study with 560 children at risk of atopy showing that exposure to house dust mite allergens (Der f 1) in first year of life was not associated with wheeze at 3 years aOR 0.92 (95%CI 0.73 to 1.15) per 1-log increase in allergen level.
- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Cat allergens (Fel d 1) (Grade F.2.3.4)

- This evidence review found high quality evidence from 1 study of 260 children showing that exposure to cat allergens (per 10mg increase in Fel d 1) was associated with
 - o bronchial hyper-responsiveness aOR 1.22 (95%CI 1.02 to 1.46)
 - o asthma aOR 1.20 (95%CI 1.01 to 1.43)
- This evidence review found low quality evidence from 1 study with 593 infants not at risk of asthma showing that exposure to cat allergens was not associated with
 - wheeze aOR 0.84 (95%CI 0.57 to 1.24)
 - o cough aOR 0.81 (95%CI 0.56 to 1.17)
- This evidence review found moderate quality evidence from 1 study with 1,233 children with asthma showing that exposure to cat allergens (Fel d 1 > 0.12 µg/g) was associated with
 - o asthma exacerbations (reported as rescue medication use) aOR 1.32 (95%CI 1.01 to 1.74)
 - wheeze aOR 1.39 (95%CI 1.05 to 1.84)
- This evidence review found moderate quality evidence from 1 study with 1,233 children with asthma showing that exposure to cat allergens was not associated with
 - o cough aOR 0.89 (95%CI 0.68 to 1.17) for Fel d 1 >0.12 μg/g
 - o asthma exacerbations (reported as Asthma Severity Index) aOR 1.14 (95%CI 0.88 to 1.47)
- This evidence review found low quality evidence from 1 study with 256 children at risk of asthma showing that exposure to cat allergens was not associated with wheeze aOR 0.64 (95%Cl 0.36 to 1.12) for Fel d 1 >1 μ g/g
- This evidence review found moderate quality evidence from 1 study with 226 children at risk of atopy that exposure to cat allergens was not associated with
 - o wheeze aOR 0.61 (95%CI 0.27 to 1.35)
 - o cough aOR 1.13 (95%CI 0.66 to 1.94)
- This evidence review found moderate quality evidence from 1 study with 1,314 children at risk of asthma showing that exposure to cat allergens was not associated with
 - \circ wheeze aOR 1.47 (95%Cl 0.72 to 1.26) for Fel d 1 >1 μ g/g
 - asthma aOR 1.52 (95%Cl 0.64 to 2.62) for Fel d 1 0.216 to 47μg
- This evidence review found low quality evidence from 1 study with 560 children at risk of atopy showing that exposure to cat allergens in the first year of life was protective against wheeze at 3 years aOR 0.71 (95%CI 0.58 to 0.88) per 1-log increase in allergen level in first year
- This evidence review found high quality evidence from 1 study with 360 showing that exposure to cat allergens was associated with asthma
 - aOR 3.33 (95%Cl 1.72 to 6.45) for Fel d 1 ≥2 μg/gm
 - o aOR 1.20 (95%CI 1.01 to 1.43) per 10 mg increase in Fel d 1
 - o aOR 1.20 (95%CI 1.01 to 1.43), per 10mg increase in Fel d 1

- This evidence review found moderate quality evidence from 1 study with 593 children showing that exposure to cat allergens was not associated with eczema
 - o aOR 1.42 (95%CI 0.72 to 2.81) for Fel d 1 0.45 to 1.04 units,
 - o aOR 1.41 (95%CI 0.71 to 2.79) for Fel d 1 1.05 to 3.33 units,
 - o aOR 1.31 (95%Cl 0.65 to 2.62) for Fel d 1 3.34 to 44.72 units,
 - o aOR 1.41 (95%CI 0.72 to 2.75) for Fel d 1 44.73 to 14151.32 units
- This evidence review found moderate quality evidence from 1 study with 593 children showing that exposure to cat allergens was not associated with visible flexural dermatitis
 - o aOR 1.28 (95%CI 0.64 to 2.56) Fel d1 0.45 to 1.04 units
 - o aOR 0.75 (95%Cl 0.36 to 1.55) Fel d 1 1.05 to 3.33 units
 - o aOR 1.18 (95%CI 0.59 to 2.38) Fel d 1 3.34 to 44.72 units
 - o aOR 0.96 (95%CI 0.48 to 1.91) Fel d 1 44.73 to 14151.32
- This evidence review found high quality evidence from 1 study with 442 children at risk of developing asthma showing that exposure to cat allergens (per interquartile increase in Fel d 1) at 3 months of age was protective against an asthma diagnosis at 7 years aOR 0.78 (95%CI 0.62 to 0.98)
- No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - o Older people
 - o People with disabilities
 - o Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Dog allergens (Can d 1) (Grade F.2.3.6)

- This evidence review found high quality evidence from 1 study with 380 infants at risk of asthma that exposure to dog allergens (Can f 1 >2 μ g/g) was associated with asthma aOR 3.84 (95%CI 1.79 to 8.22)
- This evidence review found low quality evidence from 1 study with 593 infants not at risk of asthma that exposure to dog allergens (Can f 1 ≥1.8µg/g) was not associated with cough aOR 1.11 (95%Cl 0.78 to 1.58)
- This evidence review found low quality evidence from 1 study with 1,233 children with asthma showing that exposure to dog allergens (Can f 1 >1.2 µg/g) was not associated with
 - o asthma exacerbations (reported as Asthma Severity Index) aOR 1.15 (95%Cl 0.83 to 1.58)
 - o asthma exacerbations (reported as rescue medication use) aOR 1.15 (95%Cl 0.83 to 1.62)
 - o cough aOR 1.11 (95%CI 0.80 to 1.56)
- This evidence review found low quality evidence from 1 study with 256 infants at risk of asthma that exposure to dog allergens (Can f 1 >1.8μg/g) was not associated with cough aOR 0.91 (95%CI 0.53 to 1.56)
- This evidence review found low quality evidence from 1 study with 560 children at risk of atopy that exposure to dog allergens (per 1-log increase in allergen level in first year) was not associated with wheeze at 3 years aOR 1.00 (95%CI 0.79 to 1.28)
- This evidence review found moderate quality evidence from 1 study with 442 children at risk of developing asthma showing that exposure to dog allergens (per interquartile increase in Can d 1) at 3 months of age was not associated with an asthma diagnosis at 7 years aOR 0.62 (95%CI 0.37 to 1.03)
- No evidence was identified for the following subgroups of interest

- People living in deprived areas
- o Older people
- o People with disabilities
- o Pregnant women
- People with conditions associated with or exacerbated by indoor air pollution

Nitrogen dioxide (NO₂) (Grade F.2.4)

- This evidence review found high quality evidence from 1 study with 242 children in multi-family housing showing that exposure to elevated levels of NO₂ was associated with wheeze aOR per 9.74 μg/m³ increase in NO₂ aOR 1.52 (95%CI 1.04 to 2.21)
- This evidence review found moderate quality evidence from 1 study with 593 infants at risk of asthma showing that exposure to elevated levels of NO₂ was associated with cough aOR 1.21 (95%CI 1.05 to 1.40) per 4.87 μg/m³ (reported as 10ppb) increase in NO₂
- This evidence review found moderate quality evidence from 1 study with 411 infants at risk of asthma showing that elevated levels of NO₂ were not associated with wheeze
 - \circ aOR 0.66 (95%Cl 0.27 to 1.61) for NO₂ levels between 5.2 to 6.8 μ g/m³ compared to less than 5.2.
 - $\circ~$ aOR 0.80 (95%Cl 0.32 to 2.01) for NO $_2$ levels between 6.8 to 8.6 $\mu g/m^3$ compared to less than 5.2,
 - $\circ~$ aOR 1.15 (95%Cl 0.40 to 3.32) for NO $_2$ levels between 8.6 to 11.7 $\mu g/m^3$ compared to less than 5.2
 - $\circ~$ aOR 0.43 (95%Cl 0.15 to 1.18) for NO $_2$ levels greater than 11.7 $\mu g/m^3$ compared to less than 5.2
- This evidence review found moderate quality evidence from 1 study with 1,342 children with asthma showing that exposure elevated levels of nitrogen dioxide (NO₂) were associated with wheeze
 - o aOR 1.44 (95%CI 1.11 to 1.86) for NO₂ between 18.23 and 29.35 μg/m³ and
 - o aOR 1.53 (95%CI 1.16 to 2.02) for $NO_2 > 29.35 \mu g/m^3$
- This evidence review found low quality evidence from 1 study with 1,342 children with asthma showing that elevated levels of NO₂ were not associated with wheeze aOR 1.15 (95%Cl 0.90 to 1.45) for NO₂ between 12.36 and 18.23 µg/m³ aOR
- This evidence review found moderate quality evidence from 1 study with 486 children in single-family housing showing that elevated levels of NO₂ were not associated with wheeze aOR 0.99 (95%CI 0.71 to 1.38) per 9.74 µg/m³ increase in NO₂
- This evidence review found low quality evidence from 1 study with 1,342 children with asthma that NO₂ levels between 12.36 and 18.23 μg/m³ were not associated with asthma exacerbations aOR 1.15 (95%CI 0.94 to 1.42)
- This evidence review found moderate quality evidence from 1 study with 1,342 children with asthma showing that exposure to elevated levels of NO₂) was associated with asthma exacerbations
 - aOR 1.31 (95%CI 1.04 to 1.66) for NO₂ between 18.23 and 29.35 μg/m³
 - o aOR 1.43 (95%CI 1.08 to 1.88) for $NO_2 > 29.35 \mu g/m^3$
- This evidence review found moderate quality evidence from 1 study with 442 children at risk of developing asthma showing that exposure to NO₂ (per inter quartile increase) at 12 months of age was not associated with asthma diagnosis at 7 years aOR 0.97 (95%CI 0.75 to 1.26)
- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas

- Older people
- People with disabilities
- o Pregnant women
- o People with conditions associated with or exacerbated by indoor air pollution

Polycyclic aromatic hydrocarbons (PAHs) (Grade F.2.5)

- This evidence review found moderate quality evidence from 1 study of 333 infants showing that exposure to elevated levels of PAH (per log unit of PAH concentration in ng/m³) was associated with
 - wheezing or whistling in the chest irrespective of respiratory infection aOR 3.83 (95%CI 1.18 to 12.43)
 - o wheezing without cold aOR 1.96 (95%CI 1.38 to 2.78)
 - o cough aOR 1.72 (95%CI 1.02 to 2.92),
 - o cough without cold aOR 4.80 (2.73 to 8.44)
 - o sore throat aOR 1.27 (95%CI 1.07 to 1.52)
 - o earache aOR 1.82 (95%CI 1.03 to 3.23)
- This evidence review found low quality evidence from 1 study of 333 infants showing that
 prenatal exposure to elevated levels of PAH (per log unit of PAH concentration in ng/m³) was
 not associated with
 - o barking cough aOR 1.12 (95%CI 0.82 to 1.55)
 - o difficult (puffed) breathing aOR 1.23 (95%CI 0.83 to 1.84)
 - o runny or stuffy nose aOR 1.11 (95%CI 0.97 to 1.27)
- This evidence review found moderate quality evidence from 1 study with 257 children showing that postnatal exposure to elevated levels of PAH was associated with wheeze aOR 1.61 (95%CI 1.16 to 2.24)
- This evidence review found low quality evidence from 1 study with 257 children showing that prenatal exposure to elevated levels of PAH was not associated with wheeze aOR 1.40 (95%CI 0.97 to 2.03)
- This evidence review found moderate quality evidence from 1 study with 369 children between 1 and 2 years of age showing that prenatal exposure to elevated levels of PAH was associated with wheeze aOR 1.69 (95%CI 1.52 to 1.88)
- This evidence review found low quality evidence from 1 study with 369 children between 3 and 4 years of age showing that prenatal exposure to elevated levels of PAH was not associated with wheeze aOR 0.96 (95%CI 0.84 to 1.09)
- This evidence review found moderate quality evidence from 1 study with 349 children showing that exposure to elevated levels of pyrene was associated with asthma aOR 1.90 (95%CI 1.13 to 3.20)
- This evidence review found low quality evidence from 1 study with 349 children showing that prenatal exposure to pyrene was not associated with wheeze aOR 1.53 (95%Cl 0.93 to 2.51)
- This evidence review found low quality evidence from 1 study with 349 children showing that prenatal exposure to Σ_8 PAH non-volatile was not associated with wheeze aOR 0.86 (95%CI 0.52 to 1.42)
- This evidence review found low quality evidence from 1 study with 475 premature infants and children at risk of allergies that prenatal exposure to elevated levels of PAH was associated with pulmonary infections
 - aOR 2.1 (95%CI 1.1 to 4.2) for Styrene>2.0 μg/m³

- o aOR 2.4 (95%CI 1.3 to 4.5) for Benzene > 5.6 μ g/m³
- This evidence review found low quality evidence from 1 study with 349 children showing that prenatal exposure to Σ_8 PAH non-volatile was not associated with asthma aOR 0.90 (95%CI 0.52 to 1.56)
- This evidence review found low quality evidence from 1 study with 363 children showing that prenatal exposure to Pyrene was not associated with asthma aRR 0.81 (95%CI 0.59 to 1.12)
- This evidence review found low quality evidence from 1 study with 363 children showing that prenatal exposure to Σ_8 PAH non-volatile was not associated with asthma aRR 0.74 (95%CI 0.46 to 1.18)
- This evidence review found low quality evidence from 1 study with 363 children showing that prenatal exposure to Σ_8 PAH semi-volatile was not associated with asthma aRR 0.82 (95%CI 0.60 to 1.12)
- This evidence review found low quality evidence from 1 study of 333 infants showing that prenatal exposure to elevated levels of PAH (log unit of PAH concentration in ng/m³) was associated with earache aOR 1.82 (95%CI 1.03 to 3.23)
- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - o Older people
 - People with disabilities
 - Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

Particulate Matter

PM_{2.5} (Grade F.2.6.1)

- This evidence review found low quality evidence from 1 study with 905 adults showing that PM_{2.5} from residential heating sources was not associated with cough aOR 0.95 (95%CI 0.72 to 1.29)
- This evidence review found moderate quality evidence from 1 study with 322 infants showing that prenatal exposure to PM_{2.5} was not associated with wheeze, aHR 1.06 (95%CI 0.72 to 1.57)
- This evidence review found moderate quality evidence from 1 study with 408 children showing that exposure to PM_{2.5} (per 8.75 μg/m³ increase) was associated with wheeze aOR 1.51 (95%CI 1.05 to 2.16)
- This evidence review found high quality evidence from 1 study with 103 infants at risk of asthma showing that exposure to PM_{2.5} ≥15µg/m³ was associated with wheeze aOR 4.21 (95%CI 1.36 to 13.03)
- This evidence review found moderate quality evidence from 1 study with 36 children with asthma showing that exposure to PM_{2.5} was associated with
 - \circ wheeze aOR 1.57 (95%CI 1.09 to 2.26) per 17.3 µg/m³ increase in indoor PM_{2.5}
 - wheeze aOR 1.55 (95%Cl 1.05 to 2.28) per 16.5 μg/m³ increase in indoor PM_{2.5} from indoor sources
- This evidence review found moderate quality evidence from 1 study with 150 children with asthma showing that exposure to PM_{2.5} (per 10 µg/m³ increase in PM_{2.5}) was associated with
 - o cough, wheezing or chest tightness aIRR 1.05 (95%CI 1.01 to 1.12)
 - o Asthma symptoms causing children to slow down aIRR 1.04 (95%CI 1.0 to 1.09),
 - o symptoms with running aIRR 1.07 (95%CI 1.02 to 1.11),
 - o nocturnal symptoms aIRR 1.06 (95%CI 1.01 to 1.10),

- o limited speech aIRR 1.07 (95%CI 1.00 to 1.14)
- o rescue medication use aIRR 1.04 (95%CI 1.01 to 1.08)
- This evidence review found moderate quality evidence from 1 study with 411 infants at risk of asthma showing that exposure to PM_{2.5} was not associated with wheeze
 - aOR 1.32 (95%CI 0.53 to 3.27) for PM_{2.5} between 10.6 and 13.2 μg/m³
 - aOR 1.74 (95%CI 0.67 to 4.47) for PM_{2.5} between 13.2 and 16.8 μg/m³,
 - aOR 0.67 (95%CI 0.28 to 1.59) for PM_{2.5} between 16.8 and 24 μg/m³
 - aOR 1.02 (95%Cl 0.41 to 2.57) for PM_{2.5} greater than 24.1 μg/m³
- This evidence review found low quality evidence from 1 study with 36 children with asthma showing that exposure to PM_{2.5} was not associated with
 - o cough aOR 1.22 (95%Cl 0.91 to 1.63) per 17.3 μg/m³ increase in indoor PM_{2.5}
 - cough aOR 1.20 (95%CI 0.88 to 1.64) per 17.6 µg/m³ increase in indoor PM_{2.5} from indoor sources
- No evidence was identified for the following subgroups of interest
 - o People living in deprived areas
 - o Older people
 - o People with disabilities
 - o Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

PM₁₀ (Grade F.2.6.2)

- This evidence review found moderate quality evidence from 1 study with 150 children with asthma showing that exposure to PM₁₀ (per 10 μg/m³ increase in PM₁₀) was associated with
 - o cough, wheezing or chest tightness aIRR 1.06 (95%CI 1.01 to 1.12)
 - o slowdown 1.08 (95%CI 1.02 to 1.14)
 - o nocturnal symptoms aIRR 1.08 (95%CI 1.01 to 1.14),
 - o limited speech aIRR 1.11 (95%CI 1.03 to 1.19)
 - o rescue medication use aIRR 1.06 (95%CI 1.01 to 1.10)
- This evidence review found moderate quality evidence from 1 study with 150 children with asthma showing that exposure to PM₁₀ (per 10 μg/m³ increase in PM₁₀) was not associated with symptoms with running –aIRR 1.00 (95%CI 0.94 to 1.08)
- · No evidence was identified for the following subgroups of interest
 - People living in deprived areas
 - Older people
 - o People with disabilities
 - o Pregnant women
 - o People with conditions associated with or exacerbated by indoor air pollution

The committee's discussion of the evidence

Interpreting the evidence

The outcomes that matter most

The committee noted that pollutants such as NO₂, volatile organic compounds (VOCs), particulate matter (PM) from open solid-fuel fires, polycyclic aromatic hydrocarbons (PAHs) and biological agents such as mould and pet dander are sometimes associated with many symptoms including those affecting the respiratory, cardiovascular and neurological systems

The quality of the evidence

The committee acknowledged the certainty of the evidence was mixed but also noted that this was largely due to different context in each study, such as differences in populations, age, and the myriad of ways of reporting on the same outcome. For example, respiratory symptoms covers different symptoms such coughing, sneezing, wheezing, sinus congestion, phlegm, sore throat, nasal congestion and runny nose. Most of the studies used self-report rather than objective measures for symptoms.

The committee noted that the studies did not adjust for the same confounders and tis limited and i the opportunity to get any pooled estimates of the associations between exposure and symptoms. However, a member of the committee highlighted that where point estimates from different studies showed an association but some of the confidence intervals crossed the line of no effect that the latter is a measure of uncertainty which is reflected in the overall certainty, However the committee recognised the potentially large limitations of the research and discussed the likelihood and impact of bias and imprecision with the experts in some detail and avoided the use of simple heuristics to summarise evidence.

The committee noted that many of the studies included selected populations, such as those with pre-existing conditions such as asthma or those at risk of developing conditions due to a family history. The remaining studies were in unselected populations of infants, children or adults though there were no studies that included older people or people with disabilities. Associations were less common in studies that used unselected populations than in the studies that used selected populations. Overall the committee noted only limited association between exposure and ill health in the healthy population and drafted a research recommendation to look at the health impact of indoor air pollutants, alone or in combination on people's health.

There was limited evidence of pregnancy outcomes of interest, such as low birth weight for gestational age or premature birth, though there were studies including pregnant women.

Benefits and harms

Evidence showed that people with pre-existing conditions for example respiratory or cardiovascular conditions or allergies are particularly affected by indoor air pollutants. While the majority of the evidence showed that indoor pollutants were associated with harms some showed benefits in terms of protecting against poor health. The effects of exposure to poor indoor air quality are generally cough or wheeze, nasal or throat symptoms, and eye irritation.

The committee also noted that women who are pregnant and babies under 12 months are particularly vulnerable to poor health from exposure to some pollutants such as VOC's and particulates. Evidence suggests that exposure to volatile organic compounds (VOCs) during pregnancy was associated with poor health outcomes for the child, for example, cough or wheeze in the first years of life.

The committee noted the importance of recognising signs and symptoms associated with exposure to indoor air pollution was key to action being taken. If these symptoms are associated with poor

indoor air quality, then action can be taken by health care professionals in both managing the symptoms and in referring for appropriate assessment of the property in order for the cause to be identified and remedied. If poor indoor air quality is not identified as a cause of the symptoms, the symptoms are likely to worsen with resulting greater impact on the health of the occupants.

Cost effectiveness and resource use

No cost-effectiveness review was conducted for this question as it was not an effectiveness question.

Other factors the committee took into account

As well as the evidence of associating open solid-fuel fires with poor health symptoms, the committee were also made aware of a Public Health England review in this area. The committee noted that some groups are more vulnerable to exposure to poor indoor air quality as shown in the literature review, with emphasis on the very young, those with or at risk of developing respiratory conditions. The committee noted that many of these groups will be in contact with health care professionals already and those who are social tenants will be in contact with other relevant professionals employed by local authorities.

The committee also accepted topic expert advice that knowledge of the health impact of indoor air pollution was low amongst many professionals such as those health enforcement officers and health care professionals and so the committee drafted recommendations to raise awareness around the populations at increased risk and signs and symptoms associated with indoor air pollution. There was no evidence on how effective it is for local authority and health and social care staff to trigger a referral for a housing assessment. The committee agreed that knowledge of how to do this is key to ensuring action is taken. That way, staff can make every contact count and improve people's health. To this end, the committee recommended that local authorities should raise awareness of the referral pathway for a housing assessment.

The committee also highlighted that local authorities should set up a process that their staff as well as health and social care professionals can use to contact the environmental officer if they have concerns about poor indoor air quality. The local authority should ensure that their staff and health and social care professionals are aware of this process and how to request a housing assessment.

There were discussions around symptoms with strong links to poor air quality and the committee highlighted that if people keep presenting with symptoms (such as cough, wheeze, nasal or throat symptoms) or they are getting worse, then these might be linked to their home environment.

The committee noted that healthcare professionals are more likely than environmental health officers (EHO) to see people with pre-existing conditions and women who are pregnant or have very young children. The committee agreed that this puts them in an ideal position to ask about their home and housing conditions and to give advice on how damp, mould and other pollutants such as house dust mites and VOCs from household sprays can affect their health. It also gives them the opportunity to explain how they can reduce the risks or refer people for a housing assessment if necessary. Though the committee stressed that some healthcare professionals might need training on how poor indoor air quality affects health and how to mitigate it. Also asking about housing conditions and making requests for a housing assessment may lead to an increase in consultation time. The committee suggested that training healthcare professionals on poor indoor air quality and its health effects could be incorporated into general training and professional development programmes.

The committee considered whether the problems associated with poor indoor air quality in urban areas were different to those in rural areas. The limited evidence did not show any significant difference in terms of the health impact but noted that the outdoor sources of air pollution may be different.

Appendix A: Review protocol

Field	Content		
Review question	What signs and symptoms should prompt healthcare professionals to consider exposure to poor indoor air quality at home in people presenting to health services?		
Type of review question	Prognostic type question		
Objective of the review	To identify clinical signs and symptoms that are associated with exposure to poor indoor air quality at home.		
Eligibility criteria – population/disease/condition/issue/domain	People in all dwellings		
Eligibility criteria –prognostic factor	 Prognostic factors Clinical signs / symptoms associated with exposure to indoor air pollutants at home including: Neurological symptoms for example: headache, drowsiness, fatigue, poor concentration, confusion Respiratory symptoms for example: coughing, sneezing, wheezing, sinus congestion, phlegm, sore throat, nasal congestion, runny nose Cardiovascular symptoms for example chest pain, shortness of breath Nausea Eye irritation Signs and symptoms of immune response disorder for example asthma, allergic rhinitis, dermatological conditions for example atopic eczema, psoriasis Pregnancy related for example low birth weight for gestational age, premature birth, infant mortality (but not account of the control of th		
Outcomes and prioritisation	sudden infant death (SID)), stillbirth Risk ratios, odds ratios of exposure to indoor air pollutants at home as defined or reported in the paper		
Eligibility criteria – study design	Inclusion: Prospective and retrospective cohort studies Exclusion: Systematic reviews of observational studies will not be included but may be used as a source of primary studies Cross-sectional studies		
Other inclusion exclusion criteria	Inclusion: • English language only • Published peer-reviewed studies only • Studies conducted in developed economies similar to the UK • Studies conducted from 1970 onwards • Exclusion:		

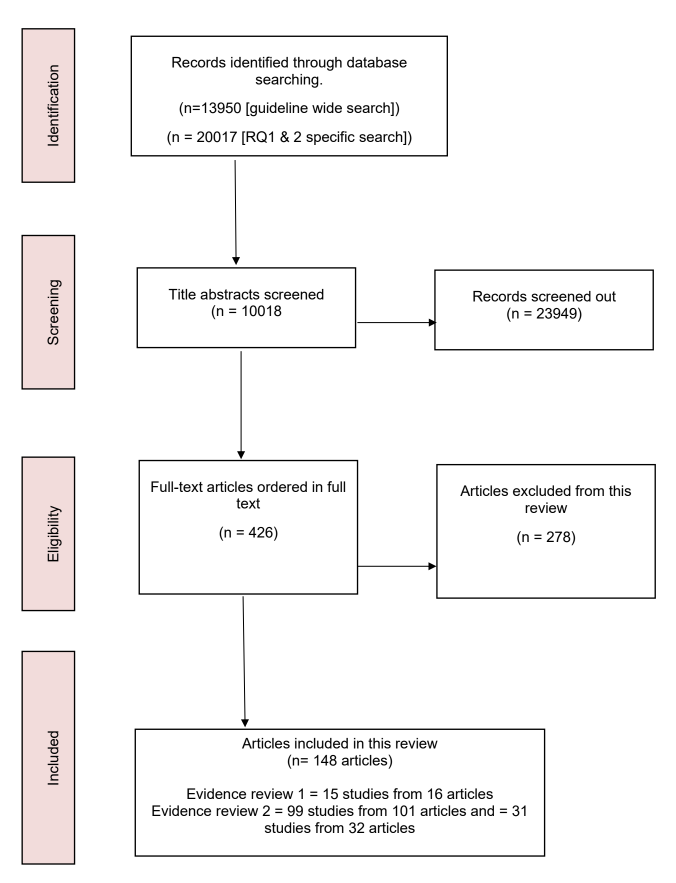
Content
Conference abstract, letter, opinion piece, review articles
• Conference abstract, letter, opinion piece, review articles
Not relevant for this type of review question
All abstracts will be duplicate screened as a reliability check. Any disagreement will be resolved by discussion, or if necessary, a third independent reviewer.
Data extraction and critical appraisal will be checked by a second reviewer. Any disagreements will be resolved by the two reviewers and escalated to a third reviewer if agreement cannot be reached.
The inclusion list will be double checked with PHAC to ensure no studies are excluded inappropriately
A systematic search of relevant databases will be carried out to identify relevant studies and evidence. Appropriate limits will be applied. Database functionality will be
used, where available, to exclude: Non-English language papers
Animal studies
Editorials, letters, news items and commentaries
 Conference abstracts and posters Theses and dissertations
Duplicates
Websites will be browsed or searched to focus on relevant evidence. The bibliographies of relevant reports and findings may also be used to capture evidence.
The following databases will be searched:
MEDLINE and MEDLINE in Process (OVID) Embase (OVID)
Health Management Information Consortium (HMIC) (OVID) Social Policy and Practice (OVID)
CENTRAL (Wiley) Cochrane Database of Systematic Reviews (Wiley) DARE (Wiley)
Greenfile (EBSCO)
NHS EED (legacy database) (Wiley)
EconLit (OVID)
OpenGrey Web of Science
The following websites will be searched:
Google and Google scholar (with appropriate limits and looking specifically for reports or evaluations of interventions related to indoor air quality)

Field	Content
Data management (software)	Where feasible data management will be undertaken using EPPI-reviewer software.
	Quantitative analysis will be performed using R software
	Where appropriate, qualitative data will be summarised using an appropriate qualitative synthesis approach, for example, narrative synthesis.
Methods for assessing bias at outcome/study level	The risk of bias across eligible studies will be assessed using the standard methodology checklist for prognostic studies. For details please see section 6.4 of Developing NICE guidelines: the manual The Grading of Recommendations Assessment, Development and Evaluation (short GRADE) developed by the GRADE working group http://www.gradeworkinggroup.org/ will be used to assess the quality of evidence across outcomes. Where necessary, GRADE will be modified to meet the needs of the review question. GRADE-CERQUAL will be used for qualitative findings.
	, , , , , , , , , , , , , , , , , , ,
Criteria for quantitative synthesis	Data from eligible studies will be extracted for inclusion in evidence tables. For details please see section 6.4 of Developing NICE guidelines: the manual
Methods of quantitative analysis – combining studies and exploring (in)consistency	Data from eligible studies shall be meta-analysed (combined) if studies are judged to be similar enough in terms of population, prognostic factors, outcomes, study design or risk of bias. Where appropriate, inconsistency will be incorporated by performing random-effect analyses If the studies are found to be too heterogeneous to be pooled statistically, a narrative synthesis will be conducted.
Meta-bias assessment – publication bias, selective reporting bias	For details please see section 6.2 of Developing NICE guidelines: the manual.
Confidence in cumulative evidence	For details please see sections 6.4 and 9.1 of Developing NICE guidelines: the manual

Appendix B: Literature search strategies

Please see search strategies here

Appendix C: Public health evidence study selection



Appendix D: Public health evidence tables

D.1.1 Bajeux 2014

Bajeux 2014					
Bibliographic reference	Bajeux E, Cordier S, Garlantézec R et.al (2014) Perinatal exposure to solvents and wheezing, eczema and food allergies at age 2. Occup Environ Med; 71: 636–641.				
Study design	Prospective cohort study				
Objective	To examine the effects of detailed eczema and food allergies in chi				
Setting/Study location	France				
Number of participants	1505 pregnant women				
Selected population	No				
Participant characteristics	Description Sex Male Age (years) reported as maternal Ethnicity Education Primary or secondary education Baccalaureate Higher education SES Building characteristics	-	777 (51.7%) 31.0 (4.2) 220 (14.7%) 278 (18.5%) 1003 (66.8%) Not reported)) 6) d	
Inclusion criteria	Not reported			-	
Exclusion criteria	Not reported				
Type of pollutant/exposure	VOC				
Pollutant/exposure assessment	Prenatal domestic exposure corresponds to exposure, self-reported by mothers at inclusion, to chemical products considered to contain solvents (paint, glues, varnishes, wood treatment products, remover products or diluents) at home in the previous 3 months. Women exposed to at least one product classified the child as prenatally exposed Postnatal exposure was defined by the use, reported by mothers at the 2-year follow-up, of chemical products known to contain solvents (paints, glues, varnishes or solvents themselves) in the home since the child's birth. The use of at least one product classified the child as postnatally exposed.				
Outcome	Wheeze Eczema Food allergies				
Results	Adjusted odds ratios (aORs) and association between solvent exp				
	\	Wheeze	Eczema	Food allergies	
		aOR (95%CI)	aOR (95%CI)	aOR (95%CI)	

Bibliographic reference	Bajeux E, Cordier S, Garlant solvents and wheezing, ecz Environ Med; 71: 636–641.				
	Prenatal domestic exposure	1.34 (0.89, 2.03)	1.13 (0.83, 1.55)	1.11 (0.69, 1.80)	
	Postnatal domestic exposure	1.80 (1.25, 2.59)	1.10 (0.86, 1.42)	1.32 (0.86, 2.03)	
	Exposed prenatally, not exposed postnatally	0.89 (0.34, 2.31)	0.72 (0.35, 1.50)	1.25 (0.41, 3.80)	
	Not exposed prenatally, exposed postnatally	1.66 (1.11, 2.47)	1.03 (0.79, 1.36)	1.28 (0.80, 2.03)	
	Exposed both prenatally and postnatally	2.50 (1.45, 4.33)	1.23 (0.84, 1.82)	1.32 (0.71, 2.46)	
Follow up	2 years				
Risk of bias (Newcastle-Ottawa Scale)					
Source of funding	Government: National Institute for Public Health Surveillance (InVS), the Ministry of Labor, and the French Agency for Food, Environmental and Occupational Health and Safety (ANSES).				
Comments	·	,			

D.1.2 Baker 2006

Dakei 2000					
Bibliographic reference	Baker R J, Hertz-Picciotto I, Dostal M Environmental Tobacco Smoke in Re Czech Children, from Birth to 3 Years 114:1126–1132	lation to Lower Respiratory Illness in			
Study design	Prospective cohort study				
Objective	To evaluate how indoor pollution from hor respiratory health in young children	ome heating may adversely affect			
Setting/Study location	United States				
Number of participants	452 children				
Selected population	No				
Participant characteristics	Description Sex Male Female Age (years) Ethnicity Education (Mother's education in years) 6–10 11 ≥ 12 Unknown Education (Father's education in years) 6–10 11 ≥ 12 Unknown) SES Building characteristics	250 (55.3%) 202 (44.7%) Not reported Not reported 91 (20.1%) 166 (36.7%) 193 (42.7%) 2 (0.4%) 75 (16.6%) 181 (40.0%) 190 (42.0%) 6 (1.3 Not reported			
Inclusion criteria	Only singleton births were included				
Exclusion criteria	Those who had moved to another district Those who were adopted or in social care Those who had died).	•			
Type of pollutant/exposu re	Coal heating				
Pollutant/expos ure assessment	Structured interview				
Outcome	Lower respiratory illness				
Results	Adjusted risk ratios (aRRs) and 95% corbetween coal as primary heating fuel an				
		Lower respiratory illness			
	Heating fuel	aRR (95%CI)			
	Coal	1.45 (1.07, 1.97)			

Bibliographic	Baker R J, Hertz-Picciotto I, Dostal M et.al (2006) Coal Home Heating and Environmental Tobacco Smoke in Relation to Lower Respiratory Illness in Czech Children, from Birth to 3 Years of Age. Environ Health Perspect
reference	114:1126–1132
Follow up	3 years
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • structured interview Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for exposure to environmental tobacco smoke • study controls for additional factors - mother's age, child's sex and year of life, child care attendance, siblings, season, day of the week, and 14-day average temperature.) Outcome Assessment of outcome • record linkage Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for
Source of funding	Overall risk of bias: Low Government: Czech Ministry of Environment (Teplice Program), the U.S. Environmental Protection Agency; the U.S. Agency for International Development, and the Commission of the European Community
Comments	

D.1.3 Bedard 2014

Bibliographic reference	Bedard A, Varraso R, Sanchez M, et al (2014) Cleaning sprays, household help and asthma among elderly women. Respiratory medicine 108(1), 171-80
Study design	Nested case-control study
Objective	To study the relationship between domestic exposure estimates, especially the use of cleaning sprays, and current asthma in elderly women
Setting/Study location	France
Number of participants	570 women
Selected population	Yes – cases selected for asthma

Bibliographic reference	Bedard A, Varraso R, Sanchez M, et al household help and asthma among eld medicine 108(1), 171-80			
Participant characteristics	Description Sex Female Age (years) – Mean (SD)	570 (100%) 68.2 (6.2)		
	Ethnicity Education <high 2-level="" 3-level="" 4-level="" diploma="" diploma<="" high="" or="" school="" td="" to="" university=""><td>Not reported 10.0% 54.2% 17.2%</td></high>	Not reported 10.0% 54.2% 17.2%		
	5-Level university diploma SES Building characteristics	18.6% Not reported Not reported		
Inclusion criteria	Not reported			
Exclusion criteria	Women with missing data for domestic exposure or asthma, and women with non-current asthma were excluded from the analysis. Women with "ever asthma" (according to the main E3N questionnaires) who did not report asthma in the specific respiratory health questionnaire were also excluded			
Type of pollutant/exposure	Domestic use of cleaning sprays			
Pollutant/exposure assessment	Questionnaire			
Outcome	Asthma			
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between domestic self-reported exposure to cleaning products and asthma.			
		Current asthma		
		aOR (95%CI)		
	Home cleaning ≥1 day/week	0.97 (0.65, 1.46)		
	Spray use ≥1 day/week	1.45 (0.94, 2.24)		
	Stratified result for women without househ	old help		
	Weekly use of at least one spray	1.86 (1.04, 3.33)		
Follow up	Not reported	· · · · · · · · · · · · · · · · · · ·		
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • selected group of women with asthma Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • written self-report Demonstration that outcome of interest was not present at start of study • No Comparability Comparability of cohorts on the basis of the design or analysis • study controls for smoking status			

Bibliographic reference	Bedard A, Varraso R, Sanchez M, et al (2014) Cleaning sprays, household help and asthma among elderly women. Respiratory medicine 108(1), 171-80
	 study controls for additional factors - age, education level and BMI Outcome Assessment of outcome record linkage self-report Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts subjects lost to follow up unlikely to introduce bias - description provided of those lost) Overall risk of bias: Moderate (concerns over self-report of exposure)
Source of funding	Government: Mutuelle Generale de l'Education Nationale (MGEN), the French League against Cancer (LNCC), the Gustave Roussy Institute (IGR) and the National Institute for Health and Medical Research
Comments	Women who reported home cleaning at least one day per week were considered as exposed for home cleaning. Frequency of nine types of sprays (furniture, glass cleaning, carpets/rugs/curtains, mopping the floor, oven, ironing, air refreshing, degreasing, insecticide/pesticide/anti-dust mite product) was collected Women who reported the use of at least one type of sprays at least one day per week were considered as exposed for spray use. Women exposed to sprays were classified as either weekly exposed to one spray or weekly exposed to at least two sprays. Information on household help (yes, no) was also recorded.

D.1.4 Belanger 2003

Bibliographic reference	Belanger K, Beckett W, Triche E et.al (2003). Symptoms of wheeze and persistent cough in the first year of life: associations with indoor allergens, air contaminants, and maternal history of asthma. American journal of epidemiology, 158(3), pp.195-202.				
Study design	Prospective coh	ort study			
Objective	and dog allerger	To examine the relationship between exposure to dust mite, cockroach, cat, and dog allergen, gas stoves, wood-burning stoves, and mould with wheeze and persistent cough in early infancy			
Setting/Study location	United States	United States			
Number of participants	849 infants. Index child was 2-4 month old)				
Selected population	Yes – selected as at risk of asthma				
Participants characteristics	Description	No.	Wheeze (%)	Persistent cough (%)	

Bibliographic reference	Belanger K, Beckett W, Triche E et.al (2003). Symptoms of wheeze and persistent cough in the first year of life: associations with indoor allergens, air contaminants, and maternal history of asthma. American journal of epidemiology, 158(3), pp.195-202.					r		
			<30 days	≥30 days	<30 days	≥30 days		
	Sex				,	,-		
	Male	419	50.8	35.6	38.7	17.4		
	Female	430	32.6	8.1	30.5	16.7		
	Age	Not reported	t					
	Ethnicity							
	White	503	33.0	7.9	422	86.8		
	Black	117	33.3	13.7	29.9	15.4		
	Hispanic	188	37.8	15.4	39.9	22.9		
	(Maintenance) medication use	Not reported	d					
	Parental asthma	a and/or atopic	3					
	Mother has asth	ıma						
	No	593	31.7	9.3	32.7	14.8		
	Yes	256	39.4	14.4	38.7	22.3		
	Father has asth	ma						
	No	669	35.0	11.1	35.7	16.4		
	Yes	171	29.8	9.4	30.4	17.5		
	Parental educat	Parental education						
	Mothers educati	on (years)						
	<12	108	43.5	17.6	38.9	25.0		
	12–15	445	32.8	11.5	35.1	17.3		
	≥16	295	32.5	7.5	32.2	13.9		
	Annual family in	come						
	<\$20,000	251	35.1	15.1	33.5	19.9		
	\$20,000 - \$50,000	183	38.2	10.4	36.6	18.0		
	>\$50,000	415	31.6	8.4	34.2	14.9		
	Building characteristics	Not reported	t					
Inclusion criteria		Only mothers who already had a child under 11 years of age with a physician diagnosis of asthma						
Exclusion criteria	Not reported							
Type of pollutant/exposure	Dust mite Pet dander Mould/mildew NO ₂ from gas s							

Bibliographic reference	Belanger K, Beckett W, Triche E et.al (2003). Symptoms of wheeze and persistent cough in the first year of life: associations with indoor allergens, air contaminants, and maternal history of asthma. American journal of epidemiology, 158(3), pp.195-202.					
Pollutant/exposure assessment	Dust samples collected in the index child's bed and in the main living area, usually the site of the highest allergen levels. These samples were analyse for house dust mite and pet allergens.					
	Dust mite: samples analysed for Der p 1 and Der f 1 Pet dander (allergen): Samples analysed for cats (Fel d 1), and dogs (Can f 1). Results are reported in micrograms per gram of fine dust for Der p 1, Der 1, Fel d 1, and Can f 1. Exposure to dust mite and cat and dog allergens was defined as exposure at ≥2 µg/g.					
	living area and the Nitrogen dioxide tube placed in the	ungal spores we ne infant's bedroo (NO ₂) from gas ne main living are	om using a B stove use wa a for 10–14 c	urkard portable a s measured usi lays. The effect	air sampler ng a Palmes of nitrogen	
	billion (ppb).		exposure grea	ater than or equa	al to 10 parts per	
Outcome	Wheeze and per					
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for indoor risk factors for wheeze and persistent cough in the first year of life					
		Children whose mothers had asthma (n=256)		Children whose mothers did not have asthma (n=593)		
		Wheeze	Persistent cough	Wheeze	Persistent cough	
		OR (95%CI)	OR (95%CI)	OR (95%CI)	OR (95%CI)	
	Dust mite (Der p 1 + Der f 1) ≥2µg/g	1.04 (0.60, 1.80)	1.27 (0.75, 2.15)	0.78 (0.55, 1.13)	0.76 (0.54, 1.07)	
	Cat allergen (Fel d 1) ≥1µg/g	0.64 (0.36, 1.12)	1.13 (0.66, 1.94)	0.84 (0.57, 1.24)	0.81 (0.56, 1.17)	
	Dog allergen (Can f 1) ≥1.8µg/g	0.69 (0.39, 1.21)	0.91 (0.53, 1.56)	1.03 (0.71, 1.49)	1.11 (0.78, 1.58)	
	Gas stove	1.03 (0.59, 1.79)	0.79 (0.46, 1.36)	1.28 (0.88, 1.86)	1.52 (1.06, 2.18)	
	Mould/mildew	2.51 (1.37, 4.62)	1.91 (1.07, 3.42)	1.22 (0.80, 1.88)	1.53 (1.01, 2.30)	
	Wood stove	Not estimable	1.04 (0.27, 3.97)	0.76 (0.37, 1.55)	1.68 (0.89, 3.20)	
	NO ₂ >10ppb	Not reported	Not reported	Not reported	1.21 (1.05, 1.40)	
Follow up	12 months					

Bibliographic reference	Belanger K, Beckett W, Triche E et.al (2003). Symptoms of wheeze and persistent cough in the first year of life: associations with indoor allergens, air contaminants, and maternal history of asthma. American journal of epidemiology, 158(3), pp.195-202.
Risk of bias (Newcastle-Ottawa Scale)	Representativeness of the exposed cohort • truly representative of the average infant at risk of asthma in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • objective sampling Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for smoking in the home • study controls for additional factors as follows maternal education, ethnicity, gender, maternal asthma, paternal asthma, maternal allergies, Annual family income, respiratory illness and smoking during pregnancy Outcome Assessment of outcome • self-report (Parent) Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall level of bias - Moderate (concerns over self-report of outcomes)
Source of funding Comments	Government: The National Institute of Environmental Health Sciences. Authors hypothesized that effects of allergens and other environmental factors on respiratory symptoms might differ between infants whose mothers had a history of physician diagnosed asthma (n=256) and children whose mothers did not (n=593). Study found no increased risk of wheeze associated with exposure to house dust mite, cat, or dog allergen. Using the home interview and quarterly questionnaires, days of wheeze and persistent cough reported for each month were summed for 12 months, and the variables were analysed as none, <30 days, or ≥30 days. This was done to distinguish children who had symptoms from those who did not and children with mild symptoms from those with severe symptoms. The choice of 30 days as the cut-off for severe symptoms was made a priori and was based on an asthma severity index developed in this cohort

D.1.5 Belanger 2006

Belanger 2006							
Bibliographic reference	Belanger K, Gent J F, Triche E W et.al (2006). Association of indoor nitrogen dioxide exposure with respiratory symptoms in children with asthma. American journal of respiratory and critical care medicine, 173(3), pp.297-303						
Study design	Prospective coh	Prospective cohort study					
Objective		To examine the associations of indoor NO ₂ exposure with respiratory symptoms among children with asthma.					
Setting/Study location	United States						
Number of participants	728 children yo	unger than 12	year				
Selected population	Yes – all had as	sthma					
Participant	Description	Multi-family h	ousing	Single-family hou	ısing		
characteristics		No.	%	No.	%		
	Sex						
	Male	151	62.4	307	63.2		
	Female	91	37.6	178	36.8		
	Age (years)						
	< 6	161	66.5	310	63.8		
	≥ 6	81	33.5	176	36.2		
	Ethnicity						
	White, Asian, other	68	28.1	422	86.8		
	Black	44	18.2	30	6.2		
	Hispanic	130	53.7	34	7.0		
	Maintenance m	edication use					
	Yes	94	38.8	276	56.8		
	No	148	61.2	210	43.2		
	Parental asthma and/or atopic	Not reported		Not reported			
	Parental education (years)	Not reported		Not reported			
	Annual family income	Not reported		Not reported			
	Building charac	teristics					
	No. rooms in ho	me					
	< 6	203	83.9	85	17.6		
	≥ 6	39	16.1	398	82.4		
	Mould/mildew						
	Yes	95	39.4	215	44.3		
	No	146	60.6	270	55.7		

Bibliographic reference	nitrogen dioxid	de exposure v can journal o	vith respiratory	s). Association of symptoms in characteristical care m	ildren with
	Water leaks				
	Yes	72	29.8	167	34.4
	No	170	70.2	318	65.6
Inclusion criteria	Children younger than 12 yr. old at the time the family enrolled Had active asthma (exhibited respiratory symptoms or used asthma medication within the year before enrolment), and Had lived at the enrolment address for at least 2 months before NO ₂ sampling. If two children in a family met the eligibility criteria, the child with more sever asthma was selected		NO ₂		
Exclusion criteria	Not reported				
Type of pollutant/exposure	NO ₂ from gas s	toves, gas dry	rers		
Pollutant/exposure assessment	NO ₂ was measured in each home using a Palmes tube placed in the main living area for 10 to 14 d after the enrolment visit. NO ₂ levels were dichotomized as less than 20 ppb versus 20 ppb or more. A concentration of 20 ppb was the median concentration of indoor NO ₂ reported for an inner-city population. Exposure data reported separately for the different sources of NO ₂ .			pb or more. A r NO ₂	
Outcome	Respiratory syn breath and ches		ted by wheeze, բ	persistent cough,	shortness of
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for household sources of NO ₂ related to respiratory symptoms				
		Wheeze	Persistent cough	Shortness of breath	Chest tightness
	Multi-family housing	OR (95%CI)	OR (95%CI)	OR (95%CI)	OR (95%CI)
	Gas stove	2.27 (1.15, 4.47)	1.19 (0.66, 2.16)	2.38 (1.12, 5.06)	4.34 (1.76, 10.69)
	Gas dryer	0.78 (0.23, 2.57)	1.19 (0.40, 3.53)	2.39 (0.77, 7.43)	1.09 (0.31, 3.90)
	NO ₂ (per 20 ppb increase)	1.52 (1.04, 2.21)	1.06 (0.75, 1.49)	1.28 (0.85, 1.91)	1.61 (1.04, 2.49)
	Single-family housing	OR (95%CI)	OR (95%CI)	OR (95%CI)	OR (95%CI)
	Gas stove	0.61 (0.35, 1.05)	0.92 (0.55, 1.51)	0.91 (0.50, 1.64)	0.68 (0.34, 1.32)
	Gas dryer	1.02 (0.50, 2.12)	0.98 (0.49, 1.94)	0.93 (0.42, 2.07)	1.41 (0.61, 3.26)
	NO ₂ (per 20 ppb increase)	0.99 (0.71, 1.38)	1.07 (0.78, 1.47)	0.83, 0.52, 1.31)	1.10 (0.78, 1.57)
				nce intervals (Cls usehold occupant	•

Bibliographic reference	nitrogen dioxid	de exposure v can journal o	vith respiratory	i). Association of symptoms in characteristical care m	ildren with
		Wheeze	Persistent cough	Shortness of breath	Chest tightness
		OR (95%CI)	OR (95%CI)	OR (95%CI)	OR (95%CI)
	Multifamily housing	1.52 (1.04, 2.21)	1.06 (0.75, 1.49)	1.28 (0.85, 1.91)	1.61 (1.04, 2.49)
	Single-family housing	0.99 (0.71, 1.38)	1.07 (0.78, 1.47)	0.83 (0.52, 1.31)	1.10 (0.78, 1.57)
Follow up	12 months				
(Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child with asthma in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • objective sampling Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for age • study controls for other factors including ethnicity, maintenance medication use, season of sampling and housing characteristics Outcome Assessment of outcome • self-report Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall level of bias — Moderate (concerns over self-report of outcomes)				
Source of funding Comments	Government: The National Institute of Environmental Health Sciences Season of sampling classified as warmer months [April—October] or cooler months [November–March]; housing characteristics classified as multi- vs.				
	single-family, number of rooms, water leaks, and visible mould. Use of maintenance medication (inhaled steroid, cromolyn, long-acting β2-agonist, or leukotriene inhibitor during the year before enrolment) was examined as a proxy for asthma severity and provided a reasonable alternative to using respiratory symptoms to classify asthma severity. • Study suggests an association between indoor NO ₂ and increased respiratory symptoms among children with asthma. The NO ₂ levels to				
	which particip The levels as housing are s	ants responde sociated with h imilar to the ou	d to are commor ealth effects amout atdoor annual ave	n in homes using ong the children i erage exposure c organization (WH	gas stoves. In multifamily If 21 ppb (40

D.1.6 Belanger 2013

Bibliographic reference		Gent JF et.al (2013). Hou diatric asthma severity. I , 24(2), 320-30.		
Study design	Prospective cohort study			
Objective	To determine the relationship between measured indoor NO_2 and concurrent asthma severity			
Setting/Study location	United states			
Number of participants	1,342 children			
Selected population	Yes – all had asthma			
Participant	Description	No.	%	
characteristics	Sex			
	Male	786	59	
	Female	556	41	
	Age (years)			
	5 – 7	703	52	
	8 – 10	639	48	
	Race/Ethnicity			
	White	538	40	
	African American	260	19	
	Hispanic	477	36	
	Mixed, Other	67	5	
	(Maintenance) medication use			
	No	460	34	
	Yes	882	66	
	Atopic			
	No	451	34	
	Yes	886	66	
	Parental education			
	Mother's education (years	3)		
	< 12	219	16	
	12 – 15	729	55	
	≥ 16	393	29	
	Annual family income	Not reported		
	Building characteristics Not reported			
Inclusion criteria	Aged between 5 to 10 years Had a caregiver who spoke English Had active asthma defined as two or more of the following: physician diagnosis; asthma symptoms within the past 12 months (wheeze, persistent cough, chest tightness, shortness of breath); use of prescription asthma medication within the past 12 months (short-acting rescue medications and		nths (wheeze, persistent prescription asthma	

Bibliographic	nitrogen dioxide and pe	Gent JF et.al (2013). Hou diatric asthma severity. I	
reference	(Cambridge, and Mass.), 24(2), 320-30. maintenance medications including inhaled steroids, systemic steroids,		
	cromolyn, leukotriene inhibitors) Children who had complete information for health outcome measures and		
Exclusion criteria	indoor NO ₂ monitoring Not reported		
Type of	·		
pollutant/exposure	NO ₂ from gas stoves		
Pollutant/exposure assessment		e NO ₂ concentration in rooi rroom) and asleep (bedroo	
	Quintile NO ₂ concentration 6.03 – 8.88, 8.89 – 14.32	on boundaries (in ppb) wer , and > 14.32.	$e \le 4.02, > 4.02 - 6.02,$
Outcome	Paediatric asthma severit Wheeze	y using Global Initiative for	Asthmas guidelines
Results	Adjusted odds ratios (aOl the risk of increased asth	Rs) and 95% confidence in ma severity	tervals (CIs) for NO ₂ and
	NO ₂ exposure	Asthma severity	Wheeze
		OR (95%CI)	OR (95%CI)
	≤ 6.02 (reference)	1.00	1.00
	6.02 – ≤ 8.88	1.15 (0.94, 1.42)	1.15 (0.90, 1.45)
	8.88 – ≤ 14.30	1.31 (1.04, 1.66)	1.44 (1.11, 1.86)
	> 14.30	1.43 (1.08, 1.88)	1.53 (1.16, 2.02)
Follow up		(,)	
Follow up Risk of bias (Newcastle-Ottawa Scale)	12 months Selection Representativeness of the exposed cohort • truly representative of the average child with asthma in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • secure sampling Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for exposure to smoking in the home • study controls for additional factors as follows age, sex, general atopy, season, specific sensitization, exposure to indoor allergens (Der p 1, Der f 1, Fel d 1, Can f 1), race/ethnicity and mother's education Outcome Assessment of outcome • record linkage Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for		

Bibliographic reference	Belanger K, Holford TR, Gent JF et.al (2013). Household levels of nitrogen dioxide and pediatric asthma severity. Epidemiology (Cambridge, and Mass.), 24(2), 320-30.
	Overall level of bias – Low
Source of funding	Government: NIH National Institutes of Environmental Health Sciences
Comments	Sampling seasons were defined by winter and summer solstice and vernal and autumnal equinox.
	Study suggests that increase in NO_2 exposure was associated with increased risk in asthma severity. The levels associated with health effects among the children are well below the outdoor annual average exposure of 21 ppb (40 μ g/m³) recommended by the World Health Organization (WHO)

D.1.7 Bertelsen 2010

Bibliographic reference	Bertelsen R J, Lodrup Carlsen K asthma and early life exposure to beta(1,3)-glucans. Clinical and ex British Society for Allergy and Cl	indoor allergens, experimental allergy	endotoxin and : journal of the	
Study design	Prospective cohort study			
Objective	To determine if exposure to indoor allergens, b(1,3)-glucans and endotoxin in the homes of the children at 2 years of age modified the risk of asthma, BHR and lung function at 10 years of age.			
Setting/Study location	Norway			
Number of participants	260 children			
Selected population	No			
Participant	Description	No.	%	
characteristics	Sex			
	Male	132	50.8	
	Female	128	49.2	
	Race/Ethnicity Not reported			
	Parental asthma	34	13.1	
	SES reported as parental education			
	University	177	68.0	
Inclusion criteria	Not reported			
Exclusion criteria	Not reported			
Type of pollutant/exposure	Allergens			
Pollutant/exposure assessment	The dust sample was collected by the parents according to detailed written instructions. The floor of the living area of the house was vacuumed using a new vacuum cleaner bag, and the collected dust was sent to the Norwegian Institute of Public Health and stored at -20 °C until extraction and analysis.			
Outcome	Asthma			
Results	Adjusted odds ratios (aORs) and 95	5% confidence interva	als (Cls)	
		Asthma	Bronchial hyper- responsiveness	
	Cat allergen (per 10 mg increase)	1.20 (1.01, 1.43)	1.22 (1.02, 1.46)	

Follow up Risk of bias (Newcastle-Ottawa Scale) Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • secure sampling Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability Comparability Comparability Comparability of cohorts on the basis of the design or analysis • study controls for alcohol in pregnancy, parental rhinoconjunctivitis at birth and parental education Outcome Assessment of outcome • Clinical diagnosis Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall level of bias – Low Source of funding Source of funding Source of funding Government: Norwegian Research Council, The Eastern Norway Regional Health Authority, Norwegian Institute of Public Health, The Norwegian Foundation for Health and Rehabilitation, Academic: The University of Oslo, Oslo University Hospital, Professional: The Norwegian Association for Asthma and Allergy, Charity: the Kloster foundation, Voksentoppen BKL, Industry: AstraZeneca, Ullev°al Pharmacia and the Hakon group	Bibliographic reference	Bertelsen R J, Lodrup Carlsen K C. Carlsen K H, et al (2010) Childhood asthma and early life exposure to indoor allergens, endotoxin and beta(1,3)-glucans. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 40(2), 307-16
(Newcastle-Ottawa Scale) Representativeness of the exposed cohort truly representative of the average child in the community Selection of the non-exposed cohort drawn from the same community as the exposed cohort Ascertainment of exposure secure sampling Demonstration that outcome of interest was not present at start of study Yes Comparability Comparability Comparability Comparability of cohorts on the basis of the design or analysis study controls for alcohol in pregnancy, parental rhinoconjunctivitis at birth and parental education Outcome Assessment of outcome Clinical diagnosis Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts complete follow up - all subjects accounted for Overall level of bias – Low Source of funding Government: Norwegian Research Council, The Eastern Norway Regional Health Authority, Norwegian Institute of Public Health, The Norwegian Foundation for Health and Rehabilitation, Academic: The University of Oslo, Oslo University Hospital, Professional: The Norwegian Association for Asthma and Allergy, Charity: the Kloster foundation, Voksentoppen BKL, Industry: AstraZeneca, Ullev°al Pharmacia and the Hakon group	Follow up	8 years
Health Authority, Norwegian Institute of Public Health, The Norwegian Foundation for Health and Rehabilitation, Academic: The University of Oslo, Oslo University Hospital, Professional: The Norwegian Association for Asthma and Allergy, Charity: the Kloster foundation, Voksentoppen BKL, Industry: AstraZeneca, Ullev°al Pharmacia and the Hakon group	(Newcastle-Ottawa	Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • secure sampling Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for alcohol in pregnancy, parental rhinoconjunctivitis at birth and parental education Outcome Assessment of outcome • Clinical diagnosis Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for
	Source of funding	Health Authority, Norwegian Institute of Public Health, The Norwegian Foundation for Health and Rehabilitation, Academic: The University of Oslo, Oslo University Hospital, Professional: The Norwegian Association for Asthma and Allergy, Charity: the Kloster foundation, Voksentoppen BKL,
	Comments	, , , , , , , , , , , , , , , , , , ,

D.1.8 Bhinder 2014

Bibliographic reference	Bhinder S, Chen H, Sato M, et al (2014) Air pollution and the development of post-transplant chronic lung allograft dysfunction (CLAD). American journal of transplantation: official journal of the American Society of Transplantation and the American Society of Transplant Surgeons 14(12), 2749-57
Study design	Retrospective cohort study

development of post-tran (CLAD). American journa American Society of Tran Transplant Surgeons 14(nsplant chronic lung allo al of transplantation : off nsplantation and the Am 12), 2749-57	graft dysfunction icial journal of the erican Society of
To identify relationship between Traffic-related air pollution (TRAP) and outcomes following transplantation in a geographically distinct cohort of lung transplant recipients		
Canada		
397 adults		
Yes – all had a lung transp	olant	
Description	No.	%
Recipient age (mean ± SD), years	46±15	-
Donor age (mean ± SD), years	43±17	-
Transplant indication		
COPD	90	23
Cystic fibrosis	102	26
Idiopathic pulmonary fibrosis	86	22
Pulmonary arterial hypertension	19	5
Bronchiectasis	18	4
Other	82	21
Developed CLAD	185	47
Death	101	25
Not reported		
Inability to geocode pern	nanent home address	
Proximity to traffic		
Authors assessed long-term exposure to traffic-related air pollution (TRAP) using two metrics of distance from major traffic roads: Computed the shortest distances between the patients' residential addresses at the time of transplantation and major traffic roads. Distances were categorized as 0–100, 101–200, 201–1000 and >1000 m. Calculated the total length of major roads that fell within circular buffer regions of a series of radii (200, 300, 500 and 1000 m) from the patients' home addresses		
Post-transplant chronic lung allograft dysfunction (CLAD)		
Adjusted hazard ratios (aHRs) and 95% confidence intervals (Cls) for association between proximity to traffic and chronic lung allograft dysfunction		
	development of post-trair (CLAD). American journal American Society of Trait Transplant Surgeons 14(To identify relationship bet outcomes following transplant recipients Canada 397 adults Yes – all had a lung transplant age (mean ± SD), years Donor age (mean ± SD), years Transplant indication COPD Cystic fibrosis Idiopathic pulmonary fibrosis Pulmonary arterial hypertension Bronchiectasis Other Developed CLAD Death Not reported Permanent home address Inability to geocode permanent home address Inability to geocode permanent home address Inability to traffic Authors assessed long-terming two metrics of distant Computed the shortest dis at the time of transplantatic categorized as 0–100, 101 Calculated the total length regions of a series of radii home addresses Post-transplant chronic lunce Adjusted hazard ratios (al-	outcomes following transplantation in a geographical transplant recipients Canada 397 adults Yes – all had a lung transplant Description No. Recipient age (mean ± 46±15 sD), years Donor age (mean ± SD), 43±17 years Transplant indication COPD 90 Cystic fibrosis 102 Idiopathic pulmonary 86 fibrosis Pulmonary arterial 19 hypertension Bronchiectasis 18 Other 82 Developed CLAD 185 Death 101 Not reported Permanent home address outside of Ontario Inability to geocode permanent home address Missing demographic data Proximity to traffic Authors assessed long-term exposure to traffic-relation using two metrics of distance from major traffic roads categorized as 0–100, 101–200, 201–1000 and >100 calculated the total length of major roads that fell wit regions of a series of radii (200, 300, 500 and 1000 mome addresses) Post-transplant chronic lung allograft dysfunction (Cl. Adjusted hazard ratios (aHRs) and 95% confidence association between proximity to traffic and chronic lung allograft dysfunction (Cl. Adjusted hazard ratios (aHRs) and 95% confidence association between proximity to traffic and chronic lung allograft dysfunction (Cl. Adjusted hazard ratios (aHRs) and 95% confidence association between proximity to traffic and chronic lung allograft dysfunction (Cl. Adjusted hazard ratios (aHRs) and 95% confidence association between proximity to traffic and chronic lung allograft dysfunction (Cl. Adjusted hazard ratios (aHRs) and 95% confidence association between proximity to traffic and chronic lung allograft dysfunction (Cl. Adjusted hazard ratios (aHRs) and 95% confidence association between proximity to traffic and chronic lung allograft dysfunction (Cl. Adjusted hazard ratios (aHRs) and 95% confidence association between proximity to traffic and chronic lung allograft dysfunction (Cl. Adjusted hazard ratios (aHRs) and 95% confidence association between proximity to traffic and chronic lung allograft dysfunction (Cl. Adjusted hazard ratios (aHRs) and proximal and and chronic lung allograft dy

Bibliographic reference	Bhinder S, Chen H, Sato M, et al (20 development of post-transplant chro (CLAD). American journal of transpl American Society of Transplantation Transplant Surgeons 14(12), 2749-5	onic lung allograft dysfunction lantation : official journal of the n and the American Society of	
		CLAD	
		aHR (95%CI)	
	Distance to major roads		
	<100m	1.96 (0.90, 4.29)	
	101–200m	1.97 (0.87, 4.47)	
	201–1000m	1.72 (0.81, 3.65)	
	>1000m	1.00	
	Distance to highways		
	<100m	4.72 (2.13, 10.47)	
	101–200m	2.72 (1.11, 6.65)	
	201–1000m	1.05 (0.70, 1.57)	
	>1000m	1.00	
Follow up	18 years		
	baseline, measured on two separate occasions at least 3 weeks apart. Irreversibility was determined after appropriate treatment for infection, rejection or both. To control for large-scale spatial patterns in CLAD that might be caused by factors other than pollution, authors created an indica variable classifying Ontario into southern and northern regions according the 14 Local Health Integrated Networks of Ontario. Authors used a stratified Cox proportional hazards model with strata defir as region to determine association between proximity to major road and CLAD adjusting for possible confounders.		
Risk of bias (Newcastle-Ottawa	Selection Representativeness of the exposed cohort		
Scale)	 somewhat representative of the average population with bilateral lung transplant in the community Selection of the non-exposed cohort 		
	 no description of the derivation of the non-exposed cohort 		
	Ascertainment of exposure		
	validated measurements used		
	Demonstration that outcome of interest was not present at start of study		
	• Yes		
	Comparability		
	 Comparability of cohorts on the basis of the design or analysis study controls for age at baseline, sex, pre-transplant diagnosis, age and gender of donor, sex matching between donor and recipient, year of transplantation Outcome 		
	Assessment of outcome		
	 Independent assessment with CLAD FEV1 to less than 80% of baseline m least 3 weeks apart 	defined as an irreversible decline in neasured on two separate occasions at	

Bibliographic reference	Bhinder S, Chen H, Sato M, et al (2014) Air pollution and the development of post-transplant chronic lung allograft dysfunction (CLAD). American journal of transplantation: official journal of the American Society of Transplant Surgeons 14(12), 2749-57
	Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall risk of bias: Moderate (associations are based a small number of patients)
Source of funding	Government: the Canadian Foundation for Innovation and the Ontario Ministry for Research and Innovation
Comments	

D.1.9 Bornehag 2005

Bibliographic reference	Bornehag CG, Sundell J, Hägerhed-Engman L, et al. (2005) Association between ventilation rates in 390 Swedish homes and allergic symptoms in children. Indoor Air. 15(4):275-80.
Study design	Nested case control
Objective	To test the hypothesis that a low-ventilation rate in homes is associated with an increased prevalence of asthma and allergic symptoms among children.
Setting/Study location	Varmland, Sweden
Number of dwellings and participants	Number of dwellings: 390 Number of participants: 400 participants (198 symptomatic and 202 non-symptomatic)
Selected population	Yes –cases has respiratory symptoms
Building and Participant characteristics	Building characteristics: Location: unclear Dwelling type: not reported Building age built before 1960, 45.9%; 1961 to 1983, 40.3%; 1984 onwards, 13.9% Type of ownership/tenancy: not reported Type of ventilation: Natural (including kitchen fan), 65.9%; Mechanical exhaust, 23.8%; mechanical exhaust and supply, 10.2% Participant characteristics: Not reported
Inclusion criteria	Cases and controls were selected from children participating in a cohort study. Cases had to have reported at least 2 symptoms of the following symptoms within the last 12 months (at the first follow-up assessment): wheezing without a cold, rhinitis without a cold or eczema Controls had to have reported no symptoms at any follow-up period.

Bibliographic reference	Bornehag CG, Sundell J, Hägerhed-Engman L, between ventilation rates in 390 Swedish homes in children. Indoor Air. 15(4):275-80.				
	All participants would not have built their homes because of moisture problems, changed residence since the first follow-up assessment.				
Exclusion criteria	Not reported				
Building factor/exposure	Ventilation rate				
Building factor/exposure assessment	professional inspectors during the first week of the a questionnaire was sent to parents of all participat questionnaire included questions regarding the chil	Ventilation rates were ascertained using a passive tracer path method by professional inspectors during the first week of the cohort study. At follow-up, a questionnaire was sent to parents of all participating children. The questionnaire included questions regarding the child's and parent's health, asthmatic or allergic symptoms, building characteristics, signs of moisture problems and odours.			
Outcome	Asthma and allergic symptoms				
Results	Building characteristic	Odds ratio (95%CI)			
	Quartile for ventilation rate				
	Third quartile vs. fourth quartile	1.17 (0.57, 2.42)			
	Second quartile vs. fourth quartile	1.35 (0.66, 2.74)			
	First quartile vs. fourth quartile	1.95 (0.94, 4.04)			
Follow up	Not reported				
Newcastle-Ottawa Scale	Selection Representativeness of the exposed cohort • selected group – children between the age of 1 and 6 from Varmland, Sweden Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • Independently assessed (trained inspectors) Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • Analysis was performed adjusting for sex, smoking in the family, and inspector's observations of moisture-related problems. Outcome Assessment of outcome • Self-reported (by parents of participating children) Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts				
	 complete follow up - all subjects accounted for Overall risk of bias: Moderate (Concern over self 	-report of outcomes)			
Source of funding	Government: The Swedish Research council for Environment, Agricultural sciences, and Spatial Planning, Swedish Asthma and Allergy Associations Research foundation, and the Swedish Foundation for Health Care Sciences				

Bibliographic reference	Bornehag CG, Sundell J, Hägerhed-Engman L, et al. (2005) Association between ventilation rates in 390 Swedish homes and allergic symptoms in children. Indoor Air. 15(4):275-80.
Comments	Cases and controls were selected from participants of a cohort study including children between 1 and 6 years old in the county of Varmland in Sweden.

D.1.10 Bowatte 2017

Bibliographic reference	Bowatte G, Lodge C J, Knibbs D L et.al (2017) Traffic-related air pollution (TRAP) exposure is associated with allergic sensitisation, asthma, and poor lung function in middle age. J Allergy Clin Immunol 2017;139:122-9					
Study design	Prospective cohort study	Prospective cohort study				
Objective	•	To determine whether exposure to Traffic-related air pollution (TRAP) in middle age is associated with current asthma and reduced lung function in adults				
Setting/Study location	Australia					
Number of participants	1405 adults					
Selected population	No					
Participant	Description	No.	%			
characteristics	Sex: male	669	49.0			
	Maternal age (years)	Not reported	Not reported			
	Ethnicity	Not reported	Not reported			
	(Maintenance) medication use	Not reported	Not reported			
	Atopy	759	55.8			
	Parental education (Socioeconomic status)					
	Grade 1-9	90	6.6			
	Grade 10-12	525	38.5			
	Trade/apprenticeship	489	35.8			
	University degree or higher	261	19.1			
	Annual family income	Not reported	Not reported			
	Building characteristics	Not reported	Not reported			
Inclusion criteria	Not reported					
Exclusion criteria	Not reported					
Type of pollutant/exposure	Proximity to major roads (living <200 m from a major road)					
Pollutant/exposure assessment	The distance from participants' residences to the nearest major road was calculated in ArcGIS 10.1 (ArcGIS 10.1, Redlands, Calif: Environmental Systems Research Institute). Major roads were defined as public sector mapping agencies Australia transport hierarchy code 301 and 302 for the					

Bowatte G, Lodge C J, Knibbs D L et.al (2017) Traffic-related air pollution (TRAP) exposure is associated with allergic sensitisation, asthma, and poor lung function in middle age. J Allergy Clin Immunol 2017;139:122-9 states of Victoria, Tasmania, Queensland, and New South Wales. Major roads included roads carrying "massive traffic," categorized as freeways, highways, arterial roads, and sub arterial roads. Participants were classified as living in proximity to a major road if their residential address was less than 200 m in straight-line distance from a major road. This cut-off point was chosen on the basis of the known rate of decay observed in levels of major traffic pollutants downwind		
Adjusted odds ratios (aOI association between prox	imity to major roads (living	
	Wheeze	Asthma
	OR (95%CI)	OR (95%CI)
Living <200 m from a major road	1.38 (1.06, 1.80)	1.21 (0.91, 1.59)
Not reported		
At the laboratory visit, participants underwent lung function tests and skin prick tests for allergens and provided blood samples. In addition to laboratory tests, participants completed a detailed interviewer-administered questionnaire. Associations were assessed using logistic regression models. Socioeconomic status (defined using education), smoking status, gas cooking, gas heating, keeping windows open more than 1 hour per week, and rural or urban status (using Accessibility/Remoteness Index of Australia 2006) were included in the regression models. Associations were examined between the exposure variables, asthma, and wheeze		
2006) were included in the regression models. Associations were examined between the exposure variables, asthma, and wheeze Selection Representativeness of the exposed cohort • truly representative of the average population in the community Selection of the non-exposed cohort • no description of the derivation of the non-exposed cohort Ascertainment of exposure • validated measurements used Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for socioeconomic status, smoking status, rural/urban location, gas cooking, gas heating, and open windows Outcome Assessment of outcome • using positive control and a standard technique • spirometry was conducted according to the joint American Thoracic Society and European Respiratory Society guidelines		
	asthma, and poor lung to 2017;139:122-9 states of Victoria, Tasmar roads included roads carry highways, arterial roads, as living in proximity to a 200 m in straight-line districtions on the basis of the traffic pollutants downwing Wheeze, asthma Adjusted odds ratios (aOf association between proximoad) and middle-age who association between proximoad) and middle-age who association between proximoad) and middle-age who association between proximoad Not reported At the laboratory visit, par prick tests for allergens at tests, participants comple questionnaire. Association Socioeconomic status (decooking, gas heating, keet and rural or urban status 2006) were included in the between the exposure vance and rural or urban status 2006) were included in the between the exposure vance of the selection of the non-exposure validated measurement Demonstration that outco. Yes Comparability Comparability of cohorts of study controls for socious location, gas cooking, goot outcome Assessment of outcome Assessment of outcome using positive control are spirometry was conducted and European Respirate	pollution (TRAP) exposure is associated with all asthma, and poor lung function in middle age. J 2017;139:122-9 states of Victoria, Tasmania, Queensland, and New roads included roads carrying "massive traffic," cate highways, arterial roads, and sub arterial roads. Par as living in proximity to a major road if their resident 200 m in straight-line distance from a major road. The chosen on the basis of the known rate of decay obstraffic pollutants downwind. Wheeze, asthma Adjusted odds ratios (aORs) and 95% confidence in association between proximity to major roads (living road) and middle-age wheeze and asthma Wheeze OR (95%CI) Living <200 m from a major road in the laboratory visit, participants underwent lung furick tests for allergens and provided blood samples tests, participants completed a detailed interviewer-questionnaire. Associations were assessed using lo Socioeconomic status (defined using education), sm cooking, gas heating, keeping windows open more thand rural or urban status (using Accessibility/Remot 2006) were included in the regression models. Association Representativeness of the exposed cohort • truly representative of the average population in the Selection Representativeness of the exposed cohort • truly representative of the average population in the Selection of the non-exposed cohort • truly representative of the derivation of the non-exposed Ascertainment of exposure • validated measurements used Demonstration that outcome of interest was not presented the properties of the design of the design of the design of the design of the properties of the design of the prope

Bibliographic reference	Bowatte G, Lodge C J, Knibbs D L et.al (2017) Traffic-related air pollution (TRAP) exposure is associated with allergic sensitisation, asthma, and poor lung function in middle age. J Allergy Clin Immunol 2017;139:122-9
	Adequacy of follow up of cohorts • Follow up not reported Overall risk of bias: Low
Source of funding	Government: Supported by the Centre for Air Quality and Health Research and Evaluation (CAR), a National Health &Medical Research Council Centre of Research Excellence, Australia.
Comments	

D.1.11 Brunekreef 1989

orunekreel 1909					
Bibliographic reference	Brunekreef B, Dockery DW, Speizer FE, et al (1989) Home dampness and respiratory morbidity in children. The American review of respiratory disease 140(5), 1363-7				
Study design	Prospective cohort study				
Objective	To explore the relationship between moisture in the home and respiratory symptoms				
Setting/Study location	United States				
Number of participants	4625 children				
Selected population	No				
Participant	Description	No. (%)			
characteristics	Sex	Not reported			
	Age	7 – 11 years			
	Ethnicity White 4625 (100%)				
	Maternal asthma and/or atopic	Not reported			
	Parental education	Not reported			
	Annual family income	Not reported			
	Building characteristics	Not reported			
Inclusion criteria	Not reported				
Exclusion criteria	Not reported				
Type of pollutant/exposure	Dampness and mould				
Pollutant/exposure assessment	Questionnaire and sampling of air quality (including humidity) done in a random sample of houses				
Outcome	aOR (95%CI) for Respiratory	symptoms			
Results	Wheeze Cough Bronchitis Chest illness	Mould 1.79 (1.44, 2.32) 2.12 (1.64, 2.73) 1.48 (1.17, 1.87) 1.40 (1.11, 1.78)	Dampness 1.23 (1.10, 1.39) 2.16 (1.64, 2.84) 1.32 (1.05, 1.67) 1.52 (1.20, 1.93)		

Bibliographic	Brunekreef B, Dockery DW, S and respiratory morbidity in respiratory disease 140(5), 13	children. The Ameri	
	Lower respiratory illness Asthma Hay fever Non-chest illness	1.57 (1.31, 1.87) 1.27 (0.93, 1.74) 1.57 (1.31, 1.74) 1.40 (1.13, 1.74)	1.68 (1.41, 2.01) 1.42 (1.04, 1.94) 1.55 (1.25, 1.93) 1.55 (1.25, 1.93)
Follow up	12 months		
Scale)	Selection Representativeness of the experimental selection of the non-exposed of the experimental selection of the non-exposed of the exposure	the average child in the cohort anity as the exposed of interest was not present the basis of the design and actors as follows — grantal education outcomes to occur at sects accounted for	sent at start of study or analysis ender, age, height,
Source of funding	Government: National Institute Environmental Protection Agen Industry: Electric Power Resea	ісу	ealth Sciences and
Comments		s. s montoro	

D.1.12 Cable 2014

Bibliographic reference	Cable N, Kelly Y, Bartley M et.al (2014). Critical role of smoking and household dampness during childhood for adult phlegm and cough: a research example from a prospective cohort study in Great Britain. BMJ open, 4(4), pp.e004807.
Study design	Prospective cohort study
Objective	To examine independent associations between childhood exposures to smoking and household dampness, and phlegm and cough in adulthood
Setting/Study location	United Kingdom

Bibliographic reference	Cable N, Kelly Y, Bartley M et.al (2014). Critical role of smoking and household dampness during childhood for adult phlegm and cough: a research example from a prospective cohort study in Great Britain. BMJ open, 4(4), pp.e004807.						
Number of participants		7320 of the British cohort who were born during 1 week in 1970 and had complete data for childhood and adult information					
Selected population	No						
Participants characteristics		Phlegm (n=214)		Cough (n=675)		Phlegm + cough (639)	
		No.	%	No.	%	No.	%
	Sex						
	Male	145	67.76	308	45.6 3	408	63.85
	Female	69	32.24	367	54.3 7	231	36.15
	Age	Not repo	orted	Not reported		Not repo	rted
	Ethnicity	Not repo	orted	Not reported		Not repo	rted
	(Maintenance) medication use	Not reported		Not reported		Not reported	
	Parental asthma and/or atopic	Not reported		Not reported		Not reported	
	Building charact	eristics					
	Household dam	oness at a	nge 10				
	No dampness	178	83.18	568	84.1 5	488	76.37
	Slight to moderate	25	11.68	84	12.4 0	108	16.91
	Marked	11	5.14	23	3.41	41	6.73
	Phlegm at age 1	0					
	Present	12	5.61	27	4.00	28	4.38
	Cough at age 10)					
	Present	31	14.49	97	14.3 7	104	16.28
	Respiratory diffic	culties at l	oirth				
	Present	6	2.34	15	2.22	14	2.19
	Social position of	f origin					
	Professional and managerial	44	20.56	99	14.6 7	88	13.77
	Skilled non- manual	24	11.21	88	13.0 4	66	10.33
	Skilled manual	92	42.99	317	46.9 6	309	48.36

Bibliographic reference	household da	mpness di nple from a	ıring child	2014). Critical r dhood for adul tive cohort stu	t phleg	m and coເ	ıgh: a
	Non-skilled manual + no male head	54	25.23	171	25.3 3	176	27.54
Inclusion criteria	British Cohort S in 1970; data h			tish residents w egularly across			ıg 1 week
Exclusion criteria	Not reported						
Type of pollutant/exposure	Dampness exp	osure at aç	je 10				
Pollutant/exposure assessment	asking them to moderate or ma	rate the pr arked. Only imp; theref	esent state a few pe	s was addressed e of household ople responded esponse was ind	dampne that the	ess as: non eir house w	e, slight, as
Outcome	Adjusted relative phlegm and co		•) and 95% confi ious 3 months fo		ntervals (C	ls
Results		Phlegm o	nly	Cough only	Phlegr	m + cough	
	Slight to moderate dampness	0.82 (0.54	1, 1.27)	0.85 (0.67, 1.09)	1.24 (0	0.99, 1.56)	
	Marked dampness	2.05 (1.07	7, 3.91)	1.26 (0.80,1.99)	2.73 (1	1.88, 3.99)	
Follow up	Not reported						
Risk of bias (Newcastle- Ottawa Scale)	dampness (0.80,1.99)					dy	

Bibliographic reference	Cable N, Kelly Y, Bartley M et.al (2014). Critical role of smoking and household dampness during childhood for adult phlegm and cough: a research example from a prospective cohort study in Great Britain. BMJ open, 4(4), pp.e004807.
	Overall level of bias – High (concerns over self-report of exposure and outcome)
Source of funding	Government: Study funded through the UK Economic and Social Research Council's International Centre for Life Course Studies in Society and Health (ES/J019119/1)
Comments	Association between household dampness and co-occurring phlegm and cough suggest long-term detrimental effects of childhood environmental exposures
	The outcome variable, i.e. patterns of two respiratory symptoms (phlegm and cough), was derived from a well-established questionnaire and not indicative of a particular respiratory disease or lung function.

D.1.13 Carlsten 2010

Bibliographic reference	Carlsten C, Dimich-Ward H, et al (2010) Indoor allergen exposure, sensitization, and development of asthma in a high-risk birth cohort. Pediatric allergy and immunology: official publication of the European Society of Pediatric Allergy and Immunology 21(4 Pt 2), e740-746			
Study design	Prospective cohort study			
Objective	To determine how early and current exposures to house dust mites, household cats or dogs, and their respective allergens predict the development of specific sensitization and asthma at the age of 7 yr in this high-risk birth cohort			
Setting/Study location	Canada			
Number of participants	380 children			
Selected population	Yes – infants at risk of asthma			
Participant characteristics	Description No. Sex Age (years) Ethnicity Education Annual family income Building characteristics	Not reported		
Inclusion criteria	At least one-first degree relative with asthma or 2 first degree relatives with other IgE-mediated allergic diseases (atopic dermatitis, seasonal or perennial allergic rhinitis, or food allergy).			
Exclusion criteria	Not reported			
Type of pollutant/exposure	House dust mite (HDM), cat allergen dog allergen			

Bibliographic reference	Carlsten C, Dimich-Ward H, et al (2010) Indoor allergen exposure, sensitization, and development of asthma in a high-risk birth cohort. Pediatric allergy and immunology: official publication of the European Society of Pediatric Allergy and Immunology 21(4 Pt 2), e740-746		
Pollutant/exposure assessment	Dust samples were collected in households at intervals over year 1 (at 2 wk, 4, 8 and 12 months) and at age 7 using a standard protocol from the following sites: the floor and mattress of the child's bedroom (and of the parents in year 1), the floor of the most commonly used room, and the upholstered furniture in that room		
Outcome	Asthma		
Results	Adjusted odds ratios (aORs) and 95	% confidence intervals (CIs)	
		Asthma	
		aOR (95%CI)	
	HDM	4.81 (2.47, 9.34)	
	Cat allergen	3.33 (1.72, 6.45)	
	Dog allergen	3.84 (1.79, 8.22)	
Follow up	7 years		
Risk of bias (Newcastle-Ottawa Scale)			
Source of funding	Government: Canadian Institutes of Health Research, the British Columbia Lung Association, and the Manitoba Medical Service Foundation.		
Comments	Lung Association, and the Manitoba	ivieuicai Service Fouridation.	

D.1.14 Casas 2012

Bibliographic	Casas L, Tischer C, Tiesler C et.al (2012). Association of gas cooking			
reference	with children's respiratory health: results from GINIplus and LISAplus birth cohort studies. Indoor air, 22(6), pp.476-82.			
Study design	Prospective cohort study			
Objective	To examine the effects of long-term exposure to gas cooking on the onset of asthma and respiratory symptoms			
Setting/Study location	Germany			
Number of participants	5078 children from birth up to the age of 10 years			
Selected population	No			
Participant	Description	No.	%	
characteristics	Sex			
	Male	2590	51	
	Female	2488	49	
	Age (years)	Not reported	Not reported	
	Ethnicity	Not reported	Not reported	
	(Maintenance) medication use	Not reported	Not reported	
	Parental asthma and/or atop	oic		
	Never	1423	28	
	Ever	3457	68	
	Parental education			
	Low	303	6	
	Medium	1314	25.9	
	High	3236	63.7	
	Annual family income	Not reported		
	Building characteristics	Not reported		
Inclusion criteria	Not reported			
Exclusion criteria	Neonates displaying at least one of the following criteria were excluded: Preterm new-borns (< 37 gestational weeks) Low birth weight (< 2500g) Congenital malformation Symptomatic neonatal infection On antibiotic medication Hospitalisation or intensive medical care during neonatal period New-borns from women with immune-related diseases, on long term medication or who suffered from drug and/or alcohol abuse New-borns from parents with nationalities other than German or who were not born in Germany			
Type of pollutant/exposure	Dampness Pet dander NO ₂ from gas cooking			

Bibliographic reference	Casas L, Tischer C, Tiesler C et.al (2012). Association of gas cooking with children's respiratory health: results from GINIplus and LISAplus birth cohort studies. Indoor air, 22(6), pp.476-82.			
Pollutant/exposure assessment	Information on pollutant exposure was taken from questionnaires administered to the parents from birth until the child was 10 years old.			
Outcome	Asthma Persistent wheeze: wheezing episodes before 3 years of age and between 3 and 10 years of age			
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for asthma and persistent wheezing			
		Asthma	Persistent wheeze	
		OR (95%CI)	OR (95%CI)	
	NO ₂ from gas cooking	1.33 (0.88, 2.00)	1.09 (0.76, 1.57)	
	Dampness (ever)	1.16 (0.87, 1.53)	1.11 (0.87, 1.43)	
	Pets at home (ever)	0.69 (0.52, 0.91)	1.05 (0.83, 1.33)	
Follow up	` ,	` '	, ,	
(Newcastle-Ottawa Scale)	From birth to the age of 10 years Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • interview □ Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for parental atopy • study controls for other factors as follows - Sex, exclusive breastfeeding during the first 4 months of life, day care centre attendance in the first 2 years, pets, dampness and mould, parental education and maternal smoking Outcome Assessment of outcome • self-report Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for □ Overall level of bias - High (concerns over self-report of exposure and			
Source of funding	outcomes) Not reported			
Comments	None	·		
Commonto	110/10			

D.1.15 Casas 2013

Casas 2013				
Bibliographic reference	Casas L, Zock JP, Carsin AE, et al (2013) The use of household cleaning products during pregnancy and lower respiratory tract infections and wheezing during early life. International journal of public health 58(5), 757-64			
Study design	Prospective cohort study			
Objective	To evaluate the effects of household use of cleaning products during pregnancy on respiratory symptoms and airway infections during the first year of life			
Setting/Study location	Spain			
Number of participants	2,292 children			
Selected population	No			
Participant characteristics	Description Sex Boys 51.7% Age 1 year Ethnicity Not reported Education (reported as maternal) Primary education or below 23.2% Secondary education 41.5% University education or more 35.3% SES Not reported Building characteristics Not reported			
Inclusion criteria	First trimester of pregnancy age ≥16 years Intention to deliver at the reference hospital Singleton pregnancy Ability to communicate in Spanish			
Exclusion criteria	Not reported			
Type of pollutant/exposure	Domestic cleaning products such as bleach, ammonia, solvents, furniture polishes, glass cleaners, air fresheners, multiuse cleaners, ironing sprays, floor cleaning sprays, oven sprays and carpet sprays			
Pollutant/exposure assessment	Interview			
Outcome	Wheeze Lower respiratory tract infection			
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between cleaning product use, lower respiratory infection (LRTI) and wheezing.			
		LRTI	Wheezing	
	Use during pregnancy	aOR (95%CI)	aOR (95%CI)	
	Categories of products (all cohorts)			
	Furniture polishes, glass cleaners and air fresheners	0.97 (0.75, 1.24)	0.93 (0.73, 1.92)	
	Spray and solvents	1.54 (1.11, 2.14)	1.68 (1.21, 2.35)	

Bibliographic reference	Casas L, Zock JP, Carsin AE, et al (2 cleaning products during pregnancy infections and wheezing during earl health 58(5), 757-64	y and lower respira	ntory tract
	Bleach and ammonia	0.99 (0.62, 1.57)	1.01 (0.78, 1.30)
	Individual products (all cohorts)	, , , , ,	, , , , , ,
	Bleach	0.91 (0.71–1.17)	0.91 (0.72–1.17)
	Ammonia	1.03 (0.82–1.29)	1.00 (0.80–1.26)
	Solvents	1.19 (0.95–1.48)	1.30 (1.03–1.62)
	Furniture polishes	0.99 (0.81–1.22)	1.01 (0.82–1.24)
	Glass cleaners	0.92 (0.72–1.18)	0.94 (0.74–1.20)
	Air fresheners	1.29 (1.03–1.63)	1.09 (0.87–1.37)
	Multiuse cleaners	0.92 (0.74–1.15)	0.91 (0.73–1.13)
	Degreasing products	1.23 (0.90–1.69)	1.32 (0.97–1.79)
	Sprays	1.29 (1.04–1.59)	1.37 (1.10–1.69)
	Timing of use (2 cohorts only n=1157)	·	·
	Spray during pregnancy only	1.19 (0.82–1.73)	1.62 (1.11–2.36)
	Spray after pregnancy only	0.90 (0.53-1.55)	1.34 (0.80–2.24)
	Spray during and after pregnancy	1.43 (0.97–2.13)	1.61 (1.08–2.41)
	Solvents during pregnancy only	0.94 (0.65-1.38)	1.04 (0.71–1.51)
	Solvents after pregnancy only	1.06 (0.67-1.66)	0.87 (0.55-1.37)
	Solvents during and after pregnancy	1.35 (0.73-2.50)	1.81 (0.98-3.37)
	Air fresheners during pregnancy only	1.31 (0.77–2.21)	1.39 (0.85-2.29)
	Air fresheners after pregnancy only	1.85 (1.04-3.30)	1.75 (1.01–3.04)
	Air fresheners during and after pregnancy	1.59 (1.00–2.55)	1.23 (0.79–1.93)
Follow up	1 year		
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average pregnant woman in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • structured interview Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for maternal smoking history • study controls for additional factors - sex, month of birth, parity, breast feeding, day care attendance, maternal age, country of origin of the mother, maternal education, maternal asthma and maternal atopy.) Outcome Assessment of outcome • self-report (maternal)		

Bibliographic reference	Casas L, Zock JP, Carsin AE, et al (2013) The use of household cleaning products during pregnancy and lower respiratory tract infections and wheezing during early life. International journal of public health 58(5), 757-64
	 Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts complete follow up - all subjects accounted for Overall risk of bias: Moderate (concerns over self-report of outcomes)
Source of funding	Government: Instituto de Salud Carlos III, the Conselleria de Sanitat Generalitat Valenciana, the Spanish Ministry of Health, the Generalitat de Catalunya, Obra Social Cajastur, Universidad de Oviedo, Department of Health of the Basque Government, Provincial Government of Gipuzkoa, and Fundacio n Roger Torne
Comments	

D.1.16 Casas 2015

Bibliographic reference	Casas L, Sunyer J, Tischer C, et al (2015) Early-life house dust mite allergens, childhood mite sensitization, and respiratory outcomes. Allergy 70(7), 820-7			
Study design	Prospective cohort study			
Objective	To evaluate the associations of early-life HDM allergen concentrations in indoor dust with respiratory symptoms, and asthma from birth to school age			
Setting/Study location	Spain, Germany, Sweden, the Netherlands			
Number of participants	4334			
Selected population	No			
Participant	Description	No.	%	
characteristics	Sex			
	Male	Not reported	Not reported	
	Female	2094	48.3	
	Age (years)	Not reported	Not reported	
	Ethnicity	Not reported	Not reported	
	Parental education High Medium Low	2331 1279 660	53.8 29.5 15.2	
	Paternal asthma / allergic rhinitis	2076	47.9	
Inclusion criteria	Children with measured HDM allergen levels (Dermatophagoides pteronyssinus (Der p1) or Dermatophagoides farina (Der f1)) in home dust			

	Casas L, Sunyer J, Tischer C, et al (2015) Early-life house dust mite			
Bibliographic reference	allergens, childhood mite sensitization, and respiratory outcomes. Allergy 70(7), 820-7			
	samples collected during early life, and with follow-ups until age 8 years (PIAMA-NHS and BAMSE) or 10 years (INMA-Menorca, LISAplus, and MAS).			
Exclusion criteria	Not reported			
Type of pollutant/exposure	House dust mite	House dust mite		
Pollutant/exposure assessment	Bedroom dust samples were collected using vacuum cleaners equipped with ALK filter holders containing paper filters on the child's mattress (INMA-Menorca, LISAplus, and PIAMA-NHS), bedroom floor (MAS) or parents' mattress (BAMSE), stored at 20°C for up to 6 years,			
Outcome	Asthma, wheeze:			
Results	Adjusted odds ratios (aORs) and 95% confidence int	ervals (Cls)	
	HDM allergen	Der p1	Der f1	
	Persistent wheezing <low ≥Low to <0.4 µg/g 0.4 to <2 µg/g ≥2 µg/g</low 	Reference 1.1 (0.7–1.8) 0.9 (0.7–1.3) 0.8 (0.5–1.1)	Reference 1.2 (0.8–1.8) 1.1 (0.8–1.5) 1.1 (0.8–1.6)	
	Asthma ≤ 6 years <low 0.4="" <0.4="" <2="" asthma="" g="" to="" μg="" ≥2="" ≥low=""> 6 years <low 0.4="" <0.4="" <2="" g="" g<="" td="" to="" μg="" ≥2="" ≥low=""><td>Reference 1.4 (0.9–2.3) 1.1 (0.8–1.6) 1.0 (0.7–1.5) Reference 1.1 (0.6–1.8) 1.1 (0.8–1.6) 0.7 (0.4–1.0)</td><td>Reference 1.2 (0.8–1.8) 1.2 (0.8–1.6) 1.2 (0.8–1.8) Reference 1.0 (0.7–1.6) 1.0 (0.7–1.4) 1.1 (0.7–1.6)</td></low></low>	Reference 1.4 (0.9–2.3) 1.1 (0.8–1.6) 1.0 (0.7–1.5) Reference 1.1 (0.6–1.8) 1.1 (0.8–1.6) 0.7 (0.4–1.0)	Reference 1.2 (0.8–1.8) 1.2 (0.8–1.6) 1.2 (0.8–1.8) Reference 1.0 (0.7–1.6) 1.0 (0.7–1.4) 1.1 (0.7–1.6)	
Follow up	From birth to the age of 10	,		
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • Objective assessment Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for sex, no. of siblings at birth, parental education, maternal smoking during pregnancy, and parental asthma or atopy. Outcome Assessment of outcome • Clinical diagnosis			

Bibliographic reference	Casas L, Sunyer J, Tischer C, et al (2015) Early-life house dust mite allergens, childhood mite sensitization, and respiratory outcomes. Allergy 70(7), 820-7
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	complete follow up - all subjects accounted for
	Overall level of bias – Low
Source of funding	Not reported
Comments	None

D.1.17 Chang 2009

Chang 2009					
Bibliographic reference	Chang J, Delfino R J, Gillen D, et al (2009) Repeated respiratory hospital encounters among children with asthma and residential proximity to traffic. Occupational and environmental medicine 66(2), 90-8				
Study design	Retrospective cohort study				
Objective	To examine the association between neighbourhood traffic burden and repeated acute respiratory illnesses that required emergency department visits and/or hospitalisation for children with a primary or secondary diagnosis of asthma				
Setting/Study location	United States				
Number of participants	3297 children	3297 children			
Selected population	Yes – all had asthma	es – all had asthma			
Participant	Description	Readmission			
characteristics		No.	%	No.	%
	Sex				
	male	1410	56.85	490	59.98
	Female	1070	43.15	327	40.02
	Ethnicity				
	White non-Hispanic	1072	43.23	392	47.98
	White Hispanic	1087	43.83	360	44.06
	Black	66	2.66	17	2.08
	Asian	69	2.78	18	2.20
	Other	124	5.00	21	2.57
	Unknown	62	2.50	9	1.10
	Maternal age (years)	Not reported		Not reported	
	(Maintenance) medication use	Not reported Not reported		d	
	Maternal asthma and/or atopic	Not reported Not reported		d	
	Parental education	Not reported		Not reported	

Bibliographic reference	Chang J, Delfino R J, Gill encounters among child traffic. Occupational and	en with asth	nma and resi	dential proxi	
	Median household (family) income				
	≤\$29 999	180	7.26	64	7.83
	\$30 000-\$39 999	687	27.7	226	27.66
	\$40 000–\$49 999	586	23.63	203	24.85
	\$50 000-\$59 999	441	17.78	150	18.36
	\$60 000 +	586	23.63	174	21.30
	Building characteristics	Not reported	d	Not reported	d
Inclusion criteria	Aged 18 years or younger One or more respiratory hospital encounters for a primary or secondary diagnosis of asthma (ICD-9 493) within the study period Home residence in census block areas located within 13 km of either UCIMC or CHOC (catchment area).				
Exclusion criteria	Not reported				
Type of pollutant/exposure	Neighbourhood traffic expo	sure			
Pollutant/exposure assessment	EZ-Locate (Tele Atlas North America Inc, Boston, MA, USA) was used to geocode residential addresses reported at the first hospital encounter. ArcView GIS was used to calculate three traffic proxies reflecting local traffic-related air pollution exposure levels. For the first traffic metric, authors calculated the shortest distance from each child's primary residence to the nearest major road (arterial road or freeway). Thereafter, a 300-metre buffer was drawn around each child's residence to reflect an "exposure zone" to local traffic-related air pollution. Authors then calculated the total length of major roads within the 300-metre buffer by summing up all arterial road and/or freeway lengths within the 300-metre buffer. Lastly, neighbourhood traffic density was calculated by dividing the total vehicle metres travelled (VMT) within the 300-metre buffer by the area of the buffer				
Outcome	Asthma hospital attendance	ce			
Results	Adjusted hazard ratios (aHRs) and 95% confidence intervals (CIs) for association between residential traffic exposure and repeated hospital encounters for children age 0 to 18 years diagnosed with asthma				
			Repeated ho	spital encour	nters
			aHR (95%CI)	
	Residence distance (metres) to nearest arterial road or freeway				
	< 300		1.00		
	150 – 300		1.21 (1.00, 1	.45)	
	50–150		1.14 (0.95, 1	.37)	
	< 50		1.11 (0.92, 1	.33)	
Follow up	Follow up not reported				
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child the community Selection of the non-exposed cohort				

Bibliographic reference	Chang J, Delfino R J, Gillen D, et al (2009) Repeated respiratory hospital encounters among children with asthma and residential proximity to traffic. Occupational and environmental medicine 66(2), 90-8
	 no description of the derivation of the non-exposed cohort Ascertainment of exposure validated measurements used Demonstration that outcome of interest was not present at start of study Yes Comparability Comparability of cohorts on the basis of the design or analysis study controls for race, age group, gender, insurance status, residence distance to treating hospital and median household income Outcome Assessment of outcome Hospital encounters for a primary diagnosis of asthma Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts Follow up not reported Overall risk of bias: Low
Source of funding	Government: South Coast Air Management District (SCAQMD), through the University of California, Los Angeles, Asthma and Outdoor Air Quality Consortium, the National Institute of Environmental Health Sciences (NIEHS) and US National Institutes of Health (NIH)
Comments	

D.1.18 Cho 2006

Bibliographic reference	Cho SH, Reponen T, LeMasters G, et al (2006) Mold damage in home and wheezing in infants. Annals of Allergy, and Asthma and Immunology 97(4), 539-545		
Study design	Prospective cohort study		
Objective	To examine association of exposure to mould or water damage and HDM with the prevalence of recurrent wheezing in infants at the age of 1 year.		
Setting/Study location	United States		
Number of participants	640 infants		
Selected population	Yes – at risk (due to parental atopy)		
Participant characteristics	Description Sex	No (%)	
	Male	354 (55.3%)	
	Maternal age (years)	Not reported	
	Ethnicity		
	White	519 (81.1%)	
	Paternal asthma and/or atopic	640 (100%)	
	Parental education	Not reported	

Bibliographic reference	Cho SH, Reponen T, LeMasters G, et al (2006) Mold damage in homes and wheezing in infants. Annals of Allergy, and Asthma and Immunology 97(4), 539-545		
	Annual family income <\$20,000 Building characteristics	95 (14.8%) Not reported	
Inclusion criteria	At least one parent was atopic, defined as having allergic symptoms and a positive reaction on a skin prick test (SPT) to at least 1 of 15 common aeroallergens		
Exclusion criteria	Not reported		
Type of pollutant/exposure	Mould from water damage		
Pollutant/exposure assessment	Home inspection - The families were requested not to clean the floor for at least 1 day before the dust sampling. At the visit, a parent was asked to identify the room where the child spent most of his or her daytime, referred to as the child's primary activity room (PAR). Dust samples were collected from flooring materials in the PAR using a vacuum cleaner (Filter Queen Majestic, HMI Industries Inc, Seven Hills, OH) at a flow rate of 800 L/min. Additionally, in the infant's PAR, the infant's bedroom, and the basement, the existence of mouldy odour was recorded using a checklist, and temperature and relative humidity were measured with a thermohygrometer		
Outcome	·		
Results	Adjusted odds ratios (aRRs) and 95% confidence intervals (CIs) for association between mould, house dust mite (HDM) and Recurrent wheeze		
		Recurrent wheeze aRRs (95%CI)	
	Mould class 1 (minor damage)	1.2 (0.9, 1.7)	
	Mould class 2 (major damage)	2.1 (1.2, 3.6)	
	HDM > 2 μ g/g	1.1 (0.8, 1.7)	
Follow up	4 – 5 months		
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • selected group of infants Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • written report (for mould) Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for house dust mite • study controls for additional factor ass follows – mould class and family income Outcome Assessment of outcome • independent blind assessment Was follow-up long enough for outcomes to occur		

Bibliographic reference	Cho SH, Reponen T, LeMasters G, et al (2006) Mold damage in homes and wheezing in infants. Annals of Allergy, and Asthma and Immunology 97(4), 539-545
	 Yes Adequacy of follow up of cohorts complete follow up - all subjects accounted for Overall level of bias - Moderate (concern over self-report of exposure)
Source of funding	Government: National Institute of Environmental Health Sciences
Comments	No
Additional references	Reponen T, Lockey J, Bernstein DI, et al (2012) Infant origins of childhood asthma associated with specific moulds. The Journal of allergy and clinical immunology 130(3), 639-644.e5

D.1.19 Clarke 2015

blance 2013			
Bibliographic reference	Clarke CA, Reynolds P, Oakley-GI, et al (2015) Indicators of microbial- rich environments and the development of papillary thyroid cancer in the California Teachers Study. Cancer epidemiology 39(4), 548-53		
Study design	Prospective cohort study		
Objective	To investigate the association between microbial-rich environments and papill	·	
Setting/Study location	United States		
Number of participants	61,799 women	61,799 women	
Selected population	No		
Participant characteristics	Description Sex	24-22 (42-24)	
	Female	61799 (100%)	
	Age (years) – Mean (SD) Ethnicity	62.7 (12.3)6	
	White	54,473 (88.1%)	
	Non-white	6,913 (11.2%)	
	Unknown	413 (0.7%)	
	Education SES	Not reported	
	Q1, Q2-low SES	12,404 (20.1%)	
	Q3	19,821 (32.1%)	
	Q4-high SES	28,839 (46.7%)	
	Unknown	735 (1.2%)	
	Building characteristics	Not reported	
Inclusion criteria	Not reported		
Exclusion criteria	Not reported		
Type of pollutant/exposure	Pet ownership		

	Clarke CA, Reynolds P, Oakley-Gl, e	et al (2015) Indicators of microbial-		
Bibliographic reference	rich environments and the development of papillary thyroid cancer in the California Teachers Study. Cancer epidemiology 39(4), 548-53			
Pollutant/exposure assessment	Questionnaire			
Outcome	Papillary thyroid cancer			
Results	Adjusted risk ratios (aRRs) and 95% confidence intervals (CIs) for association between pets at home and repeated wheeze			
		Papillary thyroid cancer aRR (95%CI)		
	Lived with a cat or dog	0.77 (0.51, 1.17)		
Follow up	Not reported			
Risk of bias	Selection			
(Newcastle-Ottawa	Representativeness of the exposed co	hort		
Scale)	• truly representative of the average w	oman in the community		
	Selection of the non-exposed cohort			
	drawn from the same community as the exposed cohort			
	Ascertainment of exposure			
	Questionnaire			
	Demonstration that outcome of interes	interest was not present at start of study		
	• Yes			
	Comparability of cohorts on the basis of the design or analysis			
	 study controls for family history of thyroid cancer 			
 study controls for additional factor as follows - race/ethnicipersonal history of BTD, parity, adolescent menstrual cycles to regular cycles, recency of pregnancy, oral contraceptive alcohol use, smoking history) 		escent menstrual cycle length and time		
	Outcome			
	Assessment of outcome			
	record linkage			
		Was follow-up long enough for outcomes to occur		
	• Yes			
	Adequacy of follow up of cohorts			
	 subjects lost to follow up unlikely to introduce bias - or description provided of those lost) 			
	Overall risk of bias - Moderate (cond	cern over self-report of exposure)		
Source of funding	Government: National Cancer Institute	e.		
Comments				

D.1.20 Cole Johnson 2004

Bibliographic reference	Cole Johnson C, Ownby DR, Havstad SL, et al (2004) Family history, dust mite exposure in early childhood, and risk for pediatric atopy and asthma. The Journal of allergy and clinical immunology 114(1), 105-10
Study design	Prospective cohort study

Bibliographic reference	Cole Johnson C, Ownby DR, Havstad SL, et a dust mite exposure in early childhood, and ris asthma. The Journal of allergy and clinical im	sk for pediatric atopy and	
Objective	To investigate the relationship of dust mite allergen exposure during early life to allergic sensitivity and asthma at 6 to 7 years of age.		
Setting/Study location	United States		
Number of participants	428 children		
Selected population	No		
Participant	Description		
characteristics	Sex Male Age (years) Ethnicity White Parental history of allergy, hay fever or asthma Education SES class Building characteristics	201 (49.1%) Up to 7 years 413 (96.5%) 243 (56.8%) Not reported Not reported Not reported	
Inclusion criteria Children born at term (>36 weeks) without complications		ications	
Exclusion criteria			
Type of pollutant/exposure	Allergens		
Pollutant/exposure assessment	Dust samples were obtained by vacuuming a 1-m2 area of floor directly beside the child's bed for 2 minutes and were assayed for Der f 1 and Der by using monoclonal antibody based assays		
Outcome	Asthma		
Results	Adjusted odds ratios (aORs) and 95% confidence	e intervals (CIs)	
	dust mite [Der f 1 + Der p 1] exposure at ≤2 years of age ≥2 μg per gram of house dust,	Asthma	
	Children without a parental history	1.04 (0.36, 3.04)	
	Children with a parental history	1.30 (0.56, 3.03)	
	Children with a maternal history	1.73 (0.59, 5.04)	
	Children with a paternal history	1.77 (0.55, 5.74)	
Follow up	7 years		
Risk of bias (Newcastle-Ottawa Scale) Selection Representativeness of the exposed cohort • truly representative of the average child in the communication of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • objective sampling Demonstration that outcome of interest was not present at the exposed cohort Ascertainment of exposure • Yes Comparability		ed cohort	

Bibliographic reference	Cole Johnson C, Ownby DR, Havstad SL, et al (2004) Family history, dust mite exposure in early childhood, and risk for pediatric atopy and asthma. The Journal of allergy and clinical immunology 114(1), 105-10
	Comparability of cohorts on the basis of the design or analysis
	 study controls for child's sex, first-born status, cord blood IgE, parental education, parental history of allergies and asthma (except where stratified by this variable), and early exposure to household cats or dogs, tobacco smoke, and day-care.
	Outcome
	Assessment of outcome
	clinical diagnosis
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	subjects lost to follow up unlikely to introduce bias
	Overall risk of bias: Low
Source of funding	Government: National Institute of Allergy and Infectious Diseases, National Institute of Environmental Health Sciences. Charity: The Fund for Henry Ford Hospital
Comments	

D.1.21 Cullinan 2004

Bibliographic reference	Cullinan P, MacNeill SJ, Harris JM et al (2004) Early allergen exposure, skin prick responses, and atopic wheeze at age 5 in English children: a cohort study. Thorax 59(10), 855-61	
Study design	Prospective cohort study	
Objective	To test the null hypothesis that early allergen exposure does not influence the development of either specific IgE sensitisation or of associated asthma	
Setting/Study location	United Kingdom	
Number of participants	552 children	
Selected population	No	
Participant	Description	
characteristics	Sex	Not reported
	Age (years)	Up to 8 years
	Ethnicity	Not reported
	Education	Not reported
	SES class	
	1/11	146
	III–V	352
	Building characteristics	Not reported
Inclusion criteria	Not reported	
Exclusion criteria	Not reported	

Bibliographic reference	Cullinan P, MacNeill SJ, Harris JM et al (2004) skin prick responses, and atopic wheeze at accohort study. Thorax 59(10), 855-61		
Type of pollutant/exposure	Dust mite and cat allergen exposure		
Pollutant/exposure assessment	Approximately 8 weeks after birth each baby was visited at home and dust samples were collected from the living room floor. These samples were assayed for concentrations of house dust mite and cat allergen using standard techniques.		
Outcome	Atopic wheeze		
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between dust mite, cat allergen exposure and asthma		
		Atopic wheeze	
		aOR (95%CI)	
	Der p 1 exposure- µg/g dust median (range)		
	Quintile 1=0.1 ((0.02–0.3)	1.00	
	Quintile 2=0.5 (0.3–0.8)	1.35 (0.37, 4.88)	
	Quintile 3=1.3 (0.8–2.2)	2.44 (0.75, 7.92)	
	Quintile 4=4.3 (2.2–7.9)	1.16 (0.30, 4.48)	
	Quintile 5=17.5 (8.0–385.0)	1.71 (0.47, 6.23)	
	Fel d 1 exposure - µg/g dust median (range)		
	Quintile 1=0.2 (0.01–0.5)	1.00	
	Quintile 2=0.7 (0.5–1.1)	1.14 (0.36, 3.65)	
	Quintile 3=1.9 (1.1–3.5)	0.73 (0.21, 2.47)	
	Quintile 4=10.1 (3.6–47.8)	0.88 (0.27, 2.86)	
	Quintile 5=140.5 (47.8–2799.8	0.51 (0.14, 1.81)	
Follow up	5.5 years		
Risk of bias	Selection		
(Newcastle-Ottawa	Representativeness of the exposed cohort		
Scale)	 truly representative of the average child in the community		
	 drawn from the same community as the exposed cohort □ 		
	Ascertainment of exposure		
	objective sampling		
	Demonstration that outcome of interest was not present at start of study		
	 Yes Comparability 		
	Comparability of cohorts on the basis of the design or analysis		
	study controls for maternal allergy		
	 study controls for additional factors – paternal atopy, maternal education, crowding index and paternal age Outcome 		
	Assessment of outcome		
	• record linkage		
	Was follow-up long enough for outcomes to occu	r	
	• Yes		

Bibliographic reference	Cullinan P, MacNeill SJ, Harris JM et al (2004) Early allergen exposure, skin prick responses, and atopic wheeze at age 5 in English children: a cohort study. Thorax 59(10), 855-61
	Adequacy of follow up of cohorts
	• subjects lost to follow up unlikely to introduce bias - small number lost -
	Overall risk of bias: Low
Source of funding	Charity: Colt Foundation
Comments	

D.1.22 de Bilderling 2005

Bibliographic reference	smoking habits and the r	n AJ, Jeff's JA et.al (2005). risk of childhood and adole emiology, 162(6), 513-22.		
Study design	Prospective cohort study			
Objective	To determine whether exposure to indoor pollutant sources in childhood and adolescence was associated with respiratory symptoms			
Setting/Study location	United Kingdom			
Number of participants	1868 children and adolesc	ents		
Selected population	No	No		
Participant	Description	No.	%	
characteristics	Sex	Not reported	Not reported	
	Maternal age (years)	Not reported	Not reported	
	Ethnicity	Not reported	Not reported	
	(Maintenance) medication use	Not reported	Not reported	
	Maternal asthma and/or atopic	Not reported	Not reported	
	Parental education	Not reported	Not reported	
	Annual family income	Not reported	Not reported	
	Building characteristics	Not reported	Not reported	
Inclusion criteria	Not reported			
Exclusion criteria	Not reported			
Type of pollutant/exposu re	NO ₂ from gas heating and cooking appliances			
Pollutant/expos ure assessment	Questionnaire ascertained data on history of atopic disorders (hay fever and eczema), the presence of smokers in the household, paternal occupation, and current smoking habits (by means of a separate confidential smoking questionnaire sent to each adolescent in a separate reply paid envelope).			

smoking habits and the	n AJ, Jeff's JA et.al (2005). Gas cooking and risk of childhood and adolescent wheeze. demiology, 162(6), 513-22.
Postal questionnaire in the same cohort to obtain data on cooking, heating, and smoking habits and personal and lifestyle factors contributing to indoor pollution.	
Childhood wheezing – reported wheezing (past and/or current) was present at any time up to age 7–8 years but not at any time up to age 15–17 years Adolescent wheezing – no reported wheezing at any time up to age 7–8 years but reported wheezing (past and/or current) at age 15–17 years Persistent wheezing – reported wheezing (past and/or current) was present at any time up to age 7–8 years and at any time up to age 15–17 years	
	Rs) and 95% confidence intervals (CIs) for association ating sources, childhood wheezing and persistent
	Adolescent wheezing
	aOR (95%CI)
Any gas for cooking	1.02 (0.77, 1.36)
Gas hob for cooking	0.93 (0.69, 1.26)
Gas hob + pilot light	1.02 (0.72, 1.42)
Gas hob + fan	1.16 (0.78, 1.74)
Gas oven for cooking	0.98 (0.73, 1.33)
Gas oven + pilot light	0.89 (0.63, 1.24)
Gas oven + fan	1.39 (0.78, 2.46)
Gas central heating	0.76 (0.47, 1.23)
Gas fire for heating	0.97 (0.67, 1.39)
7 – 8 years	
Selection Representativeness of the • truly representative of the Selection of the non-expo • drawn from the same concentration and the same concentration are self-report Demonstration that outcomes to the self-report Demonstration that outcomes to the self-report Comparability Comparability of cohorts of the study controls for person	ne average adolescent in the community sed cohort ommunity as the exposed cohort e me of interest was not present at start of study on the basis of the design or analysis nal atopic status factors as follows, gender and social class.)
	smoking habits and the American journal of epice Postal questionnaire in the smoking habits and person Childhood wheezing – repany time up to age 7–8 yeany time up to age 7–8 yeary time up to age

Bibliographic reference	de Bilderling G, Chauhan AJ, Jeff's JA et.al (2005). Gas cooking and smoking habits and the risk of childhood and adolescent wheeze. American journal of epidemiology, 162(6), 513-22.
	Overall level of bias – High (concerns over self-report of outcome and presence of outcome at 7-8 years of age),
Source of funding	Government: United Kingdom Department for Environment, Food and Rural Affairs; the United Kingdom Department of Health; and the Medical Research Council.
Comments	Only data on persistent wheeze from 1987 exposure used due to lack of information on other data points.
	Study suggests increased risk in wheezing and exposure to any gas in childhood and reduced risk with the use of electric storage heating also in childhood.

D.1.23 du Prel 2006

du Prel, X, Kramer U, Behrendt H, et al (2006) Preschool children's health and its association with parental education and individual living conditions in East and West Germany. BMC public health 6, 312 Study design Objective To investigate the associations between health indicators, living conditions and parental educational level as indicator of the social status of 6-year-old children
Objective To investigate the associations between health indicators, living conditions and parental educational level as indicator of the social status of 6-year-old children
parental educational level as indicator of the social status of 6-year-old children
0.4% -/04-1
Setting/Study Germany location
Number of 28888 children participants
Selected No population
Participant Individual characteristics
characteristics Age - Median (Range) 6.3 years (5.6 to 7.1)
Sex
female 49.2%.
Race / ethnicity Not reported
SES Not reported
Inclusion All children born in geographically defined areas criteria
Exclusion Not reported criteria
Type of pollutant / Unfavourable indoor air (defined as oven heated with fossil fuel or cooking with exposure gas)
Pollutant / Questionnaire exposure assessment
Outcome Adjusted odds ratio (95% confidence interval)

Bibliographic reference	du Prel, X , Kramer U, Behrendt H, et al (2006) and its association with parental education an in East and West Germany. BMC public health	d individual living conditions
Results	Unfavourable indoor air (East Germany) Overweight, BMI > 19 kg/m2	0.89 (0.78, 1.01)
	Bronchitis, ever diagnosed More than 4 colds in the last 12 months	1.02 (0.96, 1.09) 1.13 (1.03, 1.23)
	Frequent cough	0.97 (0.86, 1.10)
	Sneeze attacks in the last 12 months	0.92 (0.80, 1.06)
	Allergy, ever diagnosed	1.07 (0.96, 1.18)
	Eczema, ever diagnosed	0.90 (0.83, 0.98)
	Damp housing conditions (East Germany)	
	Overweight, BMI > 19 kg/m2	0.87 (0.70, 1.08)
	Bronchitis, ever diagnosed	1.25 (1.13, 1.37)
	More than 4 colds in the last 12 months	1.41 (1.25, 1.60)
	Frequent cough Sneeze attacks in the last 12 months	1.66 (1.42, 1.95) 1.52 (1.26, 1.83)
	Allergy, ever diagnosed	1.09 (0.93, 1.28)
	Eczema, ever diagnosed	1.15 (1.01, 1.31)
	Unfavourable indoor air (West Germany)	
	Overweight, BMI > 19 kg/m2	1.12 (0.86, 1.47)
	Bronchitis, ever diagnosed	1.15 (1.00, 1.32)
	More than 4 colds in the last 12 months	0.96 (0.79, 1.18)
	Frequent cough Sneeze attacks in the last 12 months	0.88 (0.68, 1.15) 1.21 (0.88, 1.66)
	Allergy, ever diagnosed	0.97 (0.79, 1.19)
	Eczema, ever diagnosed	1.07 (0.87, 1.32)
	Damp housing conditions (West Germany)	4.00 (0.05, 4.00)
	Overweight, BMI > 19 kg/m2	1.26 (0.85, 1.86)
	Bronchitis, ever diagnosed More than 4 colds in the last 12 months	1.30 (1.03, 1.65) 1.62 (1.21, 2.17)
	Frequent cough	2.60 (1.90, 3.55)
	Sneeze attacks in the last 12 months	2.25 (1.52, 3.33)
	Allergy, ever diagnosed	1.20 (0.87, 1.66)
	Eczema, ever diagnosed	1.10 (0.77, 1.57)
Follow up	6 years	
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the consection of the non-exposed cohort • drawn from the same community as the exposed	·

Bibliographic reference	du Prel, X, Kramer U, Behrendt H, et al (2006) Preschool children's health and its association with parental education and individual living conditions in East and West Germany. BMC public health 6, 312
	Ascertainment of exposure
	• structured interview
	Demonstration that outcome of interest was not present at start of study
	• Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	study controls for living conditions
	 study controls for additional factor – age, gender, location and parental education
	Outcome
	Assessment of outcome
	independent assessment
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	complete follow up - all subjects accounted for
	 subjects lost to follow up unlikely to introduce bias - or description provided of those lost)
	Overall risk of bias: Low
Source of funding	Government: Ministries of Environment of North Rhine-Westphalia and Saxony-Anhalt
Comments	

D.1.24 Dales 1991

Bibliographic reference	Dales R E, Burnett R, and Zwanenburg H (1991) Adverse health effects among adults exposed to home dampness and moulds. The American review of respiratory disease 143(3), 505-9	
Study design	Retrospective cohort study	
Objective	To examine the relationship betwee	n dampness and symptoms
Setting/Study location	Canada	
Number of participants	14,799 adults	
Selected population	No	
Participant characteristics	Description Sex Female Age (years) Ethnicity	11472 (83.2%) 33.4 (SD 5.2) years

	Dales R E, Burnett R, and Zwaner	nburg H (1991) Adverse health effects
Bibliographic reference	among adults exposed to home dampness and moulds. The American review of respiratory disease 143(3), 505-9	
	White	13430 (97.4%)
	Education Some post / secondary	5676 (41 2%)
	Other	5676 (41.2%) 6132 (44.5%)
	Annual family income	Not reported
Inclusion criteria	Not reported	
Exclusion criteria	Not reported	
Type of pollutant/exposure	Dampness and mould	
Pollutant/exposure assessment	Questionnaire	
Outcome	Adjusted odds ratios (aORs) and 95 respiratory symptoms and eye irritat	
Results	Upper respiratory symptoms Lower respiratory symptoms Chronic respiratory disease Asthma Eye irritation	aOR (95%CI) 1.50 (1.38, 1.61) 1.62 (1.48, 1.78) 1.45 (1.29, 1.64) 1.56 (1.25, 1.95) 1.63 (1.46, 1.82)
Follow up	12 months	
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average adult Selection of the non-exposed cohort	
	drawn from the same community a Ascertainment of exposure written self report	as the exposed cohort □
	 written self-report Demonstration that outcome of interest was not present at start of study No 	
	Comparability	
	Comparability of cohorts on the basis of the design or analysis	
	study controls for smoking at hom	
	 study controls for additional factors as follows – age, gender, over- crowding, region and occupation 	
	Outcome Assessment of outcome	
	• self-report	
	Was follow-up long enough for outcomes to occur • Yes	
	Adequacy of follow up of cohorts	
	• complete follow up - all subjects a	
	Overall level of bias – High (conce exposure)	erns over self-reporting of outcome and
Source of funding	None reported	

Bibliographic reference	Dales R E, Burnett R, and Zwanenburg H (1991) Adverse health effects among adults exposed to home dampness and moulds. The American review of respiratory disease 143(3), 505-9
Comments	

D.1.25 Diez 2002

DIEZ 2002			
Bibliographic reference	Diez U, Kroessner T, Rehwagen M, et al (2000) Effects of indoor painting and smoking on airway symptoms in atopy risk children in the first year of life - results of the LARS-study. International Journal of Hygiene and Environmental Health 203(1), 23-28		
Study design	Nested case control study		
Objective	To examine the influence of chemical is outcome of atopy-risk children during the	·	
Setting/Study location	Germany		
Number of participants	475 premature and at risk children of a	llergies	
Selected population	Yes – selected for risk for allergies		
Participant characteristics	Description Sex Age Ethnicity Education SES Building characteristics	Not reported Up to 1 year Not reported Not reported Not reported Not reported Not reported	
Inclusion criteria	Children with elevated cord-blood-lgE- Children with two family members suffer Children with birth-weight between 150	ering from atopic diseases,	
Exclusion criteria			
Type of pollutant/exposure	Volatile organic compounds (VOC)		
Pollutant/exposure assessment	After passive sampling of VOC using 3M monitors the substances at the adsorption layers of the monitors were desorhed by means of carbon disulphide. The extracts were analysed qualitatively and quantitatively by capillary gaschromatography The detection limit of the studied components was between 0.1 and 1.0 pg per ml. 6 out of a total of 2.5 quantitatively detected components were further analysed relative to their importance to health effects.		
Outcome	Wheezing Pulmonary infections		
Results	Adjusted odds ratios (aORs) and 95% association between VOC, pulmonary		

	Diez U, Kroessner T, Rehwagen M, et al (2000) Effects of indoor painting and smoking on airway symptoms in atopy risk children in the					
Bibliographic reference	first year of life - results of the LARS-study. International Journal of Hygiene and Environmental Health 203(1), 23-28					
		Pulmonary infections at 6 weeks	Wheezing at 1 year			
	Activity	aOR (95%CI)	aOR (95%CI)			
	Restoration reported by parents	5.6 (1.3, 24.0)	1.9 (1.1, 3.5)			
	Holding an animal		1.8 (1 0, 3.3)			
	Styrene>2.0 µg/m ³	2.1 (1.1, 4.2)				
	Benzene > 5.6 μg/m ³	2.4 (1.3, 4.5)				
Follow up	1 year					
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the expose • selected group of children at ris Selection of the non-exposed coh • drawn from the same communit Ascertainment of exposure • Objective sampling Demonstration that outcome of in • Yes Comparability Comparability of cohorts on the be • study controls for gas cooking • study controls for additional fact and domestic animals Outcome Assessment of outcome • self-report Was follow-up long enough for ou • Yes Adequacy of follow up of cohorts • complete follow up - all subjects Overall risk of bias: Moderate risk	k of allergies ort y as the exposed cohort terest was not present a asis of the design or ana tors - heating, size of the	t start of study alysis e flat, new furniture			
Source of funding	Not reported	Sit (3011001110 0701 3011-11	opert or outcomes)			
Comments						

D.1.26 Diez 2003

Bibliographic	Diez U, Rehwagen M, Rolle-Kampczyk U, et al (2003) Redecoration of apartments promotes obstructive bronchitis in atopy risk infants - Results of the LARS study. International Journal of Hygiene and Environmental Health 206(3), 173-179					
Study design	Prospective cohort study					
,	To examine the effective bronchitis in one- a					tructive
Setting/Study location	Germany					
Number of participants	186					
Selected population	Yes – children at ris	sk of asthma				
characteristics	Description Sex Male 99 (53%) Age Up to 2 years Ethnicity Not reported Education Not reported SES Not reported Building characteristics Not reported					
	Double positive fan Cord blood IgE >0.9 Low birth weight be	9 kU/l				
Exclusion criteria	Not reported					
Type of pollutant/exposure	Volatile organic cor	mpounds (VOC	s): rede	ecoratic	n of apartment	
Pollutant/exposure assessment	Questionnaire					
	Obstructive bronch	itis,				
	Adjusted odds ratio association betwee 2nd year of life and risk children during	n redecoration its effect on ob	of the a	apartme ve bron	ent during pregi	nancy, 1st and
		During 1st year	ar		During 2nd ye	ar
		Obstructive Wheezing bronchitis		zing	Obstructive bronchitis	Wheezing
	Redecoration	aOR (95%CI)	aOR (95%)	CI)	aOR (95%CI)	aOR (95%CI)
	Redecoration during pregnancy	0.6 (0.3, 1.3)	0.7 (0 1.8)	.3,	1.5 (0.5, 4.3)	0.9 (0.3, 2.6)
	Redecoration during 1st year	3.6 (1.4, 9.1)	2.5 (0 7.3)	.9,	1.6 (0.5, 5.1)	1.5 (0.5, 5.1)
	Redecoration during 2nd year	NA	NA		4.3 (1.6, 12.2)	3.7 (1.3, 10.1)
Follow up	2 years					

Bibliographic reference	Diez U, Rehwagen M, Rolle-Kampczyk U, et al (2003) Redecoration of apartments promotes obstructive bronchitis in atopy risk infants - Results of the LARS study. International Journal of Hygiene and Environmental Health 206(3), 173-179
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • selected group of children at risk of asthma Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • self-report Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for exposure to environmental tobacco smoke • study controls for additional factors – pets, dampness Outcome Assessment of outcome • self-report Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for □ Overall risk of bias: High (concerns over self-report of exposure and outcomes)
Source of funding Comments	Government: Ministry of Science and Arts, Germany

D.1.27 Emenius 2003

	Bibliographic reference	Emenius G, Pershagen G, Berglind N, et al (2003) NO ₂ , as a marker of air pollution, and recurrent wheezing in children: a nested case-control study within the BAMSE birth cohort. Occupational and environmental medicine 60(11), 876-81
_	Study design	Nested case-control study

Bibliographic reference	Emenius G, Pershagen G, Berglind N, et al (2003) NO_2 , as a marker of air pollution, and recurrent wheezing in children: a nested case-control study within the BAMSE birth cohort. Occupational and environmental medicine 60(11), 876-81			
Objective	To examine the possible association between NO ₂ exposure and recurrent wheezing during the first two years of life,			
Setting/Study location	Sweden			
Number of participants	540 children			
Selected population	No			
Participant characteristics	Description Sex Male Female Age (years) Ethnicity Education Annual family income Cases Controls Controls Cases Controls Cases Controls Cases Controls Anous Controls Ano			
Inclusion criteria	Participants had to reside in the same	dwelling as when the	y were born	
Exclusion criteria	None reported			
Type of pollutant/exposur e	NO $_2$ reported as quartiles <8.4 μ g/m 3 ; 8.4–11.6 μ g/m 3 ; 11.7–15.6 μ g/m 3 ; and >15.6 μ g/m 3 , respectively			
Pollutant/exposur e assessment	Home inspection and self-report			
Outcome	Recurrent wheezing			
Results	Adjusted odds ratios (aORs) and 95%	confidence intervals ((Cls)	
		Recurrent wheezing		
	Exposure	aORs (95%CI)		
	NO ₂ Quartile 1 <8.4 μg/m ³ Quartile 2 8.4–11.6 μg/m ³ Quartile 3 11.7–15.6 μg/m ³ Quartile 4 >15.6 μg/m ³ Building age 1940–75 1975 onwards	Reference 0.96 (0.52, 1.77) 1.08 (0.57, 2.03) 1.51 (0.81, 2.82) 1.69 (1.01, 2.89) 1.86 (1.05, 3.27)		
Follow up	2 years			
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average infant in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure			

Bibliographic reference	Emenius G, Pershagen G, Berglind N, et al (2003) NO ₂ , as a marker of air pollution, and recurrent wheezing in children: a nested case-control study within the BAMSE birth cohort. Occupational and environmental medicine 60(11), 876-81
	Home inspection including sampling
	written self-report
	Demonstration that outcome of interest was not present at start of study
	• Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	 study controls for gender, heredity, maternal age and smoking, any breast feeding, and building age.
	Outcome
	Assessment of outcome
	• self-report
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	complete follow up - all subjects accounted for □
	Overall level of bias - Moderate (concerns over self-reporting of outcome)
Source of funding	Government: Swedish National Board of Building Research, the Swedish Asthma and Allergy Association, the Swedish Foundation for Health Care Sciences and Allergy Research (Vardalstiftelsen), and the Swedish Environmental Protection Agency
Comments	

D.1.28 Emenius 2004

Bibliographic reference	Emenius G, Svartengren M, Korsgaard J, et al (2004) Indoor exposures and recurrent wheezing in infants: A study in the BAMSE cohort. Acta Paediatrica, and International Journal of Paediatrics 93(7), 899-905
Study design	Nested case-control study
Objective	To examine the relationship between the home environment and the development of recurrent wheezing during infancy
Setting/Study location	Sweden
Number of participants	540 children
Selected population	No

Emenius G, Svartengren M, Korsgaa and recurrent wheezing in infants: A Paediatrica, and International Journal	A study in the BAMSE cohort. Acta	
Description Sex Age (years) Ethnicity Education Annual family income	Not reported Not reported Not reported Not reported Not reported Not reported	
Participants had to reside in the same	dwelling as when they were born	
None reported		
Damp, mould Repainting		
Home inspection and self-report		
Risk of recurrent wheezing		
Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between recurrent wheezing, damp, mould and repainting		
	Recurrent wheezing	
Exposure	aORs (95%CI)	
Any dampness	1.4 (0.9, 2.2)	
Mould odour	2.0 (1.0, 3.9)	
Mould spots on surface material/tile joints in wet areas (shower/bath room)	1.0 (0.5, 1.7)	
Repainting	1.7 (1.1, 2.6)	
2 years		
Selection Representativeness of the exposed cohort • truly representative of the average infant in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • Home inspection including sampling • written self-report Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for maternal smoking • study controls for other additional factor as follows - gender, heredity, breastfeeding and building age. Outcome Assessment of outcome		
	and recurrent wheezing in infants: A Paediatrica, and International Journ Description Sex Age (years) Ethnicity Education Annual family income Participants had to reside in the same None reported Damp, mould Repainting Home inspection and self-report Risk of recurrent wheezing Adjusted odds ratios (aORs) and 95% association between recurrent wheezing Exposure Any dampness Mould odour Mould spots on surface material/tile joints in wet areas (shower/bath room) Repainting 2 years Selection Representativeness of the exposed coetruly representative of the average in Selection of the non-exposed cohort of drawn from the same community as Ascertainment of exposure Home inspection including sampling written self-report Demonstration that outcome of interest Yes Comparability Comparability of cohorts on the basis of study controls for other additional factor breastfeeding and building age. Outcome	

Bibliographic reference	Emenius G, Svartengren M, Korsgaard J, et al (2004) Indoor exposures and recurrent wheezing in infants: A study in the BAMSE cohort. Acta Paediatrica, and International Journal of Paediatrics 93(7), 899-905
	Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for □ Overall level of bias – Moderate (concerns over self-reporting of outcome)
Source of funding	Government: Swedish National Board of Building Research, the Swedish Asthma and Allergy Association, the Swedish Foundation for Health Care Sciences and Allergy Research (Va°rdalstiftelsen), and the Swedish Environmental Protection Agency
Comments	

D.1.29 Emenius 2004 b

Bibliographic reference	Emenius G, Svartengren M, Korsgaard J, et al (2004) Building characteristics, indoor air quality and recurrent wheezing in very young children (BAMSE). Indoor air 14(1), 34-42				
Study design	Nested case-control study				
Objective	To assess the influence of building characteristics and indoor air quality, using objective measurements of ventilation rate and indoor humidity on recurrent wheezing in children up to the age of 2 years.				
Setting/Study location	Sweden				
Number of participants	540 children				
Selected population	Yes – cases selected on basis of recurrent wheezing				
Participant characteristics	Description Sex Male Female Age (years) Ethnicity Education Annual family income Building characteristics Building age: houses built before 1940 houses built between 1940–1975 houses built after 1975 apartment buildings erected after 1940 Single-family homes single-family homes: cellar basement	escription ex ale ale framale ge (years) hnicity ducation nnual family income aliding characteristics uilding age: buses built before 1940 buses built between 1940–1975 buses built after 1975 buses built after 1975 buses built after 1975 buses built dings erected after 1940 Cases Controls 170 189 2 years Not reported Not reported Not reported Not reported 153 (42.6) 153 (42.6) 154 (29.0) 178 (50.1)			

Bibliographic	Emenius G, Svartengren M, Korsgaard J characteristics, indoor air quality and red			
reference	children (BAMSE). Indoor air 14(1), 34-42		· ·	,, ,
	single-family: crawl space/concrete slab	34 (18.9)		61 (17.2)
	single-family homes with exhaust			
	ventilation and crawl space/concrete slab foundation	40 (0.7)		40 (0.7)
	12 (6.1)			13 (3.7)
	Balanced ventilation	85 (47.0) 43 (23.8)		151 (42.1) 78 (21.8)
	Ventilation rate 0.5 ACH	130 (71.8)		240 (67.0)
	Absolute indoor humidity > median, 5.8 g/kg	102 (56.4)		167 (46.7)
	Relative humidity >45%	25 (13.8)		43 (12.0)
	Indoor temperature > median, 21.7°C	93 (51.4)		182 (50.7)
	Windowpane condensation	26(14.4)		26(7.2)
Inclusion criteria	Participants had to reside in the same dwell	` '	they we	, ,
Exclusion criteria	None reported			
Type of	NO ₂			
pollutant/exposur e	Ventilation rate Humidity			
Pollutant/exposur e assessment	Home inspection and self-report			
Outcome	Recurrent wheezing			
Results	Adjusted odds ratios (aORs) and 95% confi	dence interva	als (CIs)	
			Recurrent wheezing	
	Exposure		aORs (95%CI)	
	Single-family homes vs. apartments		2.5 (1.	3, 4.8)
	Houses built before 1940		Referent	
	houses built between 1940–1975		2.3 (1.2, 4.3)	
	houses built after 1975 63		2.5 (1.3, 4.8)	
	apartment buildings erected after 1940 vs before 1940		0.9 (0.6, 1.5)	
	single-family homes: cellar basement		1.5 (0.	•
	single-family: crawl space/concrete slab		2.5 (1.	1, 5.4)
	single-family homes with exhaust ventilation	and crawl	2.0 (1	1 0 0\
	space/concrete slab foundation		3.0 (1.	•
	Exhaust ventilation (vs. natural ventilation) Balanced ventilation (vs. natural ventilation)		1.1 (0. 0.8 (0.	•
	Ventilation rate 0.5 ACH vs. lower		1.3 (0.	•
	Absolute indoor humidity > median, 5.8 g/kg	1	1.7 (1.	•
Relative humidity >45% Indoor temperature > median, 21.7°C			0.8 (0.	•
			0.9 (0.	
	Windowpane condensation		2.2 (1.	
Follow up	2 years			
Risk of bias	Selection			
(Newcastle-	Representativeness of the exposed cohort			
Ottawa Scale)	• truly representative of the average infant i	n the commu	ınity	

Bibliographic reference	Emenius G, Svartengren M, Korsgaard J, et al (2004) Building characteristics, indoor air quality and recurrent wheezing in very young children (BAMSE). Indoor air 14(1), 34-42
	Selection of the non-exposed cohort
	$ullet$ drawn from the same community as the exposed cohort \Box
	Ascertainment of exposure
	Home inspection including sampling
	written self-report
	Demonstration that outcome of interest was not present at start of study
	Yes Comparability
	Comparability Comparability of cohorts on the basis of the design or analysis
	 study controls for gender, heredity, maternal age and smoking, any breast feeding, and building age.
	Outcome
	Assessment of outcome
	• self-report
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts • complete follow up - all subjects accounted for □
	Overall level of bias – Moderate (concerns over self-reporting of outcome)
Source of funding	Government: Swedish National Board of Building Research, the Swedish
cource or furnaling	Asthma and Allergy Association, the Swedish Foundation for Health Care Sciences and Allergy Research (Va°rdalstiftelsen), and the Swedish Environmental Protection Agency
Comments	

D.1.30 Engvall 2001

Bibliographic reference	Engvall K, Norrby C, and Norback D (2001) Sick building syndrome in relation to building dampness in multi-family residential buildings in Stockholm. International archives of occupational and environmental health 74(4), 270-8
Study design	Retrospective cohort study
Objective	To investigate relationship between symptoms and signs of building dampness
Setting/Study location	Sweden
Number of participants	9,808 adults
Selected population	No

Bibliographic reference	Engvall K, Norrby C, and Norback D (2001) Sick building syndrome in relation to building dampness in multi-family residential buildings in Stockholm. International archives of occupational and environmental health 74(4), 270-8	
Participant characteristics	Sex Female Age (years) 18 – 44 45 – 64 > 65 Ethnicity Education Annual family income	5783 (60%) 4904 (51%) 2397 (25%) 2365 (24%) Not reported Not reported Not reported
Inclusion criteria	Not reported	
Exclusion criteria	Not reported	
Type of pollutant/exposure	Dampness	
Pollutant/exposure assessment	Self-administered questionn	aire
Outcome	Risk of eye symptoms, nasa fatigue and facial skin symptoms	I symptoms, throat symptoms, cough, headache, oms
Results	Eye irritation Nasal symptoms Throat symptoms Cough Facial skin symptoms Headache Tiredness	Condensation on windows 3.14 (3.01, 3.27) 2.72 (2.62, 2.81) 3.22 (3.09, 3.35) 2.58 (2.47, 2.70) 2.11 (2.02, 2.20) 3.30 (3.19, 3.43) 2.19 (2.12, 2.25)
	Eye irritation Nasal symptoms Throat symptoms Cough Facial skin symptoms Headache Tiredness	High air humidity in bathroom 2.94 (2.83, 3.05) 1.94 (1.88, 2.01) 3.23 (3.12, 3.25) 2.30 (2.21, 2.40) 2.42 (2.33, 2.51) 3.07 (2.96, 3.17) 2.16 (2.11, 2.22)
	Eye irritation Nasal symptoms Throat symptoms Cough Facial skin symptoms Headache Tiredness	Mouldy odour 3.75 (3.60, 3.92) 2.83 (2.73, 2.93) 3.48 (3.33, 3.62) 3.30 (3.16, 3.46) 2.93 (2.80, 3.06) 3.37 (3.24, 3.51) 2.38 (2.31, 2.46)
	Eye irritation	History of water leakage 1.57 (1.50, 1.65)

Bibliographic reference	Engvall K, Norrby C, and Norback D (2001) Sick building syndrome in relation to building dampness in multi-family residential buildings in Stockholm. International archives of occupational and environmental health 74(4), 270-8		
	Nasal symptoms	1.36 (1.31, 1.41)	
	Throat symptoms	2.18 (2.09, 2.28)	
	Cough	1.52 (1.44, 1.59)	
	Facial skin symptoms	1.56 (1.48, 1.63)	
	Headache	1.27 (1.21, 1.33)	
	Tiredness	1.35 (1.30, 1.39)	
Follow up	5 years		
Risk of bias	Selection		
(Newcastle-Ottawa	Representativeness of the exposed	cohort	
Scale)	• truly representative of the average	adult in the community	
	Selection of the non-exposed cohort		
	• drawn from the same community a	as the exposed cohort	
	Ascertainment of exposure		
	 written self-report 		
	Demonstration that outcome of interest was not present at start of study		
	• No		
	Comparability		
	Comparability of cohorts on the basis of the design or analysis		
	• study controls for type of ventilatio	n	
	• study controls for additional factors	s as follows – age, gender	
	Outcome		
	Assessment of outcome		
	• self-report		
	Was follow-up long enough for outco	omes to occur	
	• Yes		
	Adequacy of follow up of cohorts		
	 complete follow up - all subjects accounted for □ 		
	Overall level of bias – High (conce exposure)	rns over self-reporting of outcome and	
Source of funding	Government: Building Research Foundation, City of Stockholm, Social Board of Welfare and Health, and Swedish National Institute of Public Health		
Comments			
23/11/10/10			

D.1.31 Engvall 2010

9	
	Engvall K, Hult M, Corner R, et al. (2010) A new multiple regression model to identify multi-family houses with a high prevalence of sick building symptoms
Bibliographi	"SBS", within the healthy sustainable house study in Stockholm (3H). Int Arch
c reference	Occup Environ Health 83: 85–94
Study design	Retrospective cohort study

	Engvall K, Hult M, Corner R, et al. (2010) A new multiple regression model to		
Bibliographi c reference	identify multi-family houses with a high prevalence of sick building symptoms "SBS", within the healthy sustainable house study in Stockholm (3H). Int Arch Occup Environ Health 83: 85–94		
Objective	To develop a new model to identify residential buildings with higher frequencies of "sick building systems": a set of non-specific symptoms occurring in a particular building and not caused by specific illness, such as allergy or infection.		
Setting/Stud y location	Sweden		
Number of dwellings and participants	Number of dwellings: 11,160 dwellings in 481 buildings Number of participants: 7,640 adults		
Selected population	No		
Building and Participant characteristic s	Building characteristics: Location: urban Dwelling type: apartments Building age: varied		
	Type of ownership/tenancy: publicly owned, 29%; owned by inhabitant, 52%; private landlord, 19% Participant characteristics:		
	Age: 18 to 44 years, 46%; 45 to 54 years, 32%; >54 years 22% Current smoker: 14% Hay fever: 24% Atopy: 40%		
Inclusion criteria	Apartments in multi-family buildings were included. No further details were provided		
Exclusion criteria	Detached and semi-detached houses, as well as buildings used as nursing homes for the elderly were excluded		
Building factor/expos ure	Type of ownership: self-owned versus rented		
Building factor/expos ure assessment	Data on property ownership was obtained from the Stockholm central building register.		
Outcome	Eye, nasal, throat irritation, and coughing that occupant considered to be caused by the indoor environment of their building/property. For the context of this review these symptoms are considered likely to be caused by air pollutants.		
Results	Odds ratio (95%CI) Building factor Eye Nasal Throat Coughing irritation irritation Rented vs. 2.07 2.07 1.98 1.85 self-owned (1.19, 3.58) (1.33, 3.20) (0.98, 3.97) (0.94, 3.65) *odds ratio (95%CI) Coughing irritation irritation (0.98, 3.97) (0.94, 3.65)		
Follow up	Minimum of 6 months		
Study methods	Methods:		

Bibliographi c reference	Engvall K, Hult M, Corner R, et al. (2010) A new multiple regression model to identify multi-family houses with a high prevalence of sick building symptoms "SBS", within the healthy sustainable house study in Stockholm (3H). Int Arch Occup Environ Health 83: 85–94
	Residences were selected from a central building register by stratified random sampling. Strata were based on building size (number of apartments) and age in 6 classes (based on major changes in building technology). In each selected dwelling, one randomly selected person who was over 18 years and living in the apartment or over 6 months by combining the building register with the civil registration register. A self-administered postal questionnaire (Stockholm Indoor Environmental Questionnaire; SIEQ) was sent to these individuals. The questionnaire assessed demographic traits, medical conditions and building-related allergies or symptoms. Statistical analysis: multivariate logistic regression.
Newcastle- Ottawa Scale	Selection Representativeness of the exposed cohort • truly representative of the average resident in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • secure record (building register) Demonstration that outcome of interest was not present at start of study • No Comparability Comparability of cohorts on the basis of the design or analysis • study controls for age of the building and the number of residences Outcome Assessment of outcome • self-reported Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall risk of bias: Moderate (Concerns over sell-report of outcome)
Source of funding	Government: City of Stockholm, The Swedish Research Council, and Stockholm county council
Comments	No additional comments

D.1.32 Farrow 2003

Bibliographic reference	Farrow A, Taylor H, Northstone K, et al (2003) Symptoms of mothers and infants related to total volatile organic compounds in household products. Archives of Environmental Health 58(10), 633-641
Study design	Prospective cohort study
Objective	Not reported
Setting/Study location	United Kingdom

Bibliographic reference	Farrow A, Taylor H, Northstone K, et al (2003) Symptoms of mothers and infants related to total volatile organic compounds in household products. Archives of Environmental Health 58(10), 633-641			
Number of participants	14,541 pregnant women and 13,971 of their children			
Selected population	No			
Participant	Description			
characteristics	Sex		Not reported	
	Age		Up to 1 year	
	Ethnicity Education		Not reported Not reported	
	SES		Not reported	
	Building characteristics		Not reported	
Inclusion criteria	Expected date of deliver Place of residence within county of Avon, UK	•		
Exclusion criteria	Not reported			
Type of pollutant/exposure	Total volatile organic compounds (TVOCs): Air freshener use Aerosol use			
Pollutant/exposure assessment	Questionnaire and rando each pregnant woman w to 6 months following bir	as monitored		
	VOCs were monitored with TENAX tubes which were exposed for more that 1 month for each of the 12 months of sampling in the main bedroom and living room of the home.			
Outcome	Questionnaire			
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between Infants' Symptoms during the First 6 month Postpartum and Air Freshener and Aerosol Use during pregnancy			
	Frequency of air freshen	er use		
	Symptom	Never/< once per week	Once/week	Daily/most days
		aOR (95%CI)	aOR (95%CI)	aOR (95%CI)
	Diarrhoea (infant)	1.00 Ref	1.20 (1.06, 1.35)	1.10 (0.99, 1.23)
	Vomiting (infant)	1.00 Ref	1.06 (0.93, 1.20)	1.09 (0.97, 1.22)
	Earache (Infant)	1.00 Ref	1.24 (1.02, 1.50)	1.30 (1.09, 1.54)
	Frequency of aerosol use			
	Diarrhoea (infant)	1.00 Ref	1.09 (0.93, 1.28)	1.22 (1.09, 1.36)
	Vomiting (infant)	1.00 Ref	1.17 (1.00, 1.37)	1.14 (1.02, 1.27)
	Earache (infant)	1.00 Ref	1.00 (0.78, 1.29)	1.05 (0.84, 1.25)
	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between Maternal Symptoms of Depression and Headache during the 8 months Following Birth of the Infant and Reported Air Freshener and Aerosol Use during Pregnancy			

Bibliographic reference	Farrow A, Taylor H, No and infants related to t products. Archives of	total volatile o	rganic compounds	s in household
	Frequency of air fresher	ner use	•	
	Symptom	Never/< once per week	Once/week	Daily/most days
		aOR (95%CI)	aOR (95%CI)	aOR (95%CI)
	Depression (mother)	1.00 Ref	1.11 (0.96, 1.29)	1.19 (1.05, 1.36)
	Headache (mother)	1.00 Ref	1.06 (0.94, 1.19)	1.24 (1.11, 1.38)
	Frequency of aerosol us	e		
	Depression (mother)	1.00 Ref	1.06 (0.88, 1.27)	1.03 (0.91, 1.17)
	Headache (mother)	1.00 Ref	1.16 (1.00, 1.35)	1.25 (1.13, 1.39)
	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between Maternal Symptoms 9–21 month after the Infant Was Born That Were Associated Significantly with Reported Air Freshener and Aerosol Use during Pregnancy			
	Frequency of air fresher	ner use		
	Symptom		Once/week	Daily/most days
			aOR (95%CI)	aOR (95%CI)
	Headache (mother)		1.29 (1.14, 1.47)	1.22 (1.09, 1.36)
	Cough or cold (mother)		1.03 (0.87, 1.20)	0.82 (0.72, 0.93)
	Diarrhoea (mother)		1.14 (1.00, 1.31)	1.14 (1.01, 1.28)
	Frequency of aerosol use			
	Headache (mother)		1.35 (1.15, 1.59)	1.21 (1.10, 1.34)
	Influenza (mother)		1.03 (0.85, 1.24)	0.87 (0.77, 0.99)
	Urinary tract infection (m	nother)	1.16 (0.89, 1.52)	1.23 (1.04, 1.45)
Follow up	21 months			
Risk of bias	Selection			
(Newcastle-Ottawa Scale)	 Representativeness of the exposed cohort truly representative of the average mother / child in the community Selection of the non-exposed cohort drawn from the same community as the exposed cohort Ascertainment of exposure questionnaire and samples form randomly selected houses Demonstration that outcome of interest was not present at start of study Yes Comparability Comparability of cohorts on the basis of the design or analysis study controls for number of smokers in the home study controls for any additional factors - education, mother's age, housing tenure, number of children in the home, paid job subsequent to birth of the child, dampness or condensation in the home, mould in the home, type of winter heating fuel.) Outcome 			

Bibliographic reference	Farrow A, Taylor H, Northstone K, et al (2003) Symptoms of mothers and infants related to total volatile organic compounds in household products. Archives of Environmental Health 58(10), 633-641
	 Assessment of outcome self-report Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts complete follow up - all subjects accounted for Overall risk of bias: Moderate (Concerns over self-report of outcomes)
Source of funding	Government: Medical Research Council, the Department of Health, the Department of the Environment, Ministry of Agriculture Fisheries and Food, Industry: Nutricia, and other companies Charity: the Wellcome Trust
Comments	

D.1.33 Franck 2014

Bibliographic reference	Franck U, Weller A, Roder SW, et al (2014) Prenatal VOC exposure and redecoration are related to wheezing in early infancy. Environment international 73, 393-401		
Study design	Prospective cohort		
Objective	To evaluate the impact of prenatal corassociated VOC exposure	mpared to postnatal decoration and	
Setting/Study location	Germany		
Number of participants	629 mother-baby dyads (465 completed)		
Selected population	No		
Participant characteristics	Building characteristics: Location: Dwelling type: Building age: Type of ownership/tenancy Individual characteristics: Age: Current smoker (reported as mother) Hay fever: Atopy: 1 parent 2 parents	Urban Not reported Not reported Not reported So (11.4%) Not reported (229 (49.2%) 81 (17.4%)	

Bibliographic reference	Franck U, Weller A, Roder SW, et al (2014) Prenatal VOC exposure and redecoration are related to wheezing in early infancy. Environment international 73, 393-401					
Inclusion criteria	Pregnant women between the 20th and 34th weeks of gestation					
Exclusion criteria	Mothers with autoimmune diseases					
	M0thers with infections duri	ng the pregnancy				
Type of pollutant/exposure	VOCs as a result of redecoration					
Pollutant/exposure assessment	Passive samples taken over a period of 1 month using a diffusion sampler (OVM 300 from 3M company) placed in the middle of the room at between 1.5 and 2 metres height in the 34th week of gestation and the end of 1st year of life. The sampler was placed in the living room or the sleeping room of the mother during pregnancy and in the child's bedroom and room where the child spent daytime.					
Outcome	AOR (95%CI) for	AOR (95%CI) for				
	Physician-diagnosed obstructive bronchitis, non-obstructive bronchitis or asthma in first 12 months of life					
Dogulto	Parent report of wheezing in					
Results	Any type of redecoration During pregnancy	Recurrent wheeze 2.04 (0.78, 5.28)	Obstructive bronchitis Not reported			
	During pregnancy During 1st year of life New furniture	1.89 (0.71, 5.06)	Not reported			
	During pregnancy	1.94 (0.72, 5.26)	Not reported			
	During 1st year of life Painting	2.26 (0.83, 6.17)	Not reported			
	During pregnancy During 1st year of life Floor covering (wall to wall carpet)	2.35 (0.89, 6.20) 2.53 (0.85, 7.49)	5.46 (1.09, 27.20) Not reported			
	During pregnancy	5.39 (1.75, 16.54)	4.39 (1.01, 19.05)			
	During 1st year of life	4.18 (0.40, 43.70)	Not reported			
	Type of floor covering - pregnancy	Treated wheeze				
	Parquet	5.78 (0.30, 111.08)				
	Laminate	4.46 (1.01, 19.63)				
	Wall to wall carpet	4.57 (1.14, 18.39)				
	PVC Other	24.7 (2.18, 280.39)				
	Adhesive	2.34 (0.19, 29.29)				
	Non-adhesive	7.05 (1.61, 30.92)				
	Type of floor covering - 1st year of life	5.46 (1.44, 20.62)				
	Parquet	No data				
	Laminate	2.44 (0.40, 14.74)				
	Wall to wall carpet PVC	0.98 (0.10, 9.83)				
	Other	51.7 (3.21, 833.2)				
	Outo	1.53 (0.10, 22.36)				

Bibliographic reference		r SW, et al (2014) Prenatal VOC exposure and o wheezing in early infancy. Environment		
	Adhesive Non-adhesive	1.18 (0.06, 22.75) 2.49 (0.57, 10.85)		
Follow up	12 months			
Study methods	Part of Lifestyle and Environmental Factors and their Influence of Newburn Allergy risk (LINA) birth cohort Children were followed up once a year with questionnaire evaluations, clinical examinations and indoor measurements. Univariate and multivariate analyses were performed for binary disease outcomes using logistic regression and adjusted odds ratios (aOR) were calculated.			
Newcastle-Ottawa Scale	outcomes using logistic regression and adjusted odds ratios (aOR) were			
Source of funding	Private - Helmholtz Centre is funded by government)	for Environmental Research GmbH (this centre		
Comments	None			

D.1.34 Gan 2010

Bibliographic reference	Gan WQ, Tamburic L, Davies HW, et al (2010) Changes in residential proximity to road traffic and the risk of death from coronary heart disease. Epidemiology (Cambridge, and Mass.) 21(5), 642-9				
Study design	Prospective cohort		, , ,		
Objective		To explore the association between changes in residential proximity to road traffic and the risk of CHD mortality			
Setting/Study location	Canada				
Number of participants	450,283 adults				
Participant characteristics		No. (%)	No. (%)	No. (%)	
	Description	Not exposed (n= 328,609)	Moved close to traffic (n= 15747)	Consistent Exposure to Traffic (n=52,948)	
	Sex	Not reported	Not reported	Not reported	
	Age (years); mean (SD)	58.7 (10.4)	58.6 (10.2)	61.0 (10.9)	
	Comorbidity				
	Diabetes	6243 (1.9)	331 (2.1)	1324 (2.5)	
	COPD	12158 (1.0)	189 (1.2)	794 (1.5)	
	Hypertensive heart disease	12158 (3.7)	630 (4.0)	2436 (4.6)	
	Any of the above	18402 (5.6)	1008 (6.4)	3812 (7.2)	
	Ethnicity	Not reported	Not reported	Not reported	
	(Maintenance) medication use	Not reported	Not reported	Not reported	
	Maternal asthma and/or atopic	Not reported	Not reported	Not reported	
	Parental education	Not reported	Not reported	Not reported	
	Annual family income	Not reported	Not reported	Not reported	
	Building characteristics	Not reported	Not reported	Not reported	
Inclusion criteria	Registered with the provincial health insurance plan, which provides universal coverage to the resident population Age 45–85 years Without previous diagnosis of CHD				
Exclusion criteria	Not reported				
Type of pollutant/exposure	Proximity to traffic				
Pollutant/exposure assessment	Authors used high-resolution land-use regression models to evaluate exposure levels to traffic-related air pollutants. Using detailed residential history and corresponding monthly concentrations of traffic-related air pollutants during the 5-year exposure period, average concentrations of air pollutants were calculated for each subject.				

Bibliographic reference	Gan WQ, Tamburic L, Davies HW, et al (2010) Changes in residential proximity to road traffic and the risk of death from coronary heart disease. Epidemiology (Cambridge, and Mass.) 21(5), 642-9					
Outcome		Coronary Heart Disease Mortality				
Results	Adjusted odds ratios (aRRs) and 95% confidence intervals (CIs) for association between proximity to traffic and Coronary Heart Disease Mortality					
		Not Exposed to Traffic	Moved Close to Traffic	Consistent Exposure to Traffic		
		RR (95%CI)	RR (95%CI)	RR (95%CI)		
	≤150 m Highway or ≤50 m major road	1.00	1.20 (1.00, 1.43)	1.29 (1.18, 1.41)		
Follow up	5 years					
Study methods	A case of CHD death was defined as a death record in the provincial death registration database with CHD (ICD-9 codes 410–414, 429.2 and ICD-10 codes I20 –I25) as the cause of death. A small proportion of deaths were identified using provincial hospitalization records. To determine the association between residential proximity to traffic (predictor variable) and the risk of CHD mortality (dependent variable), authors first performed bi-variable logistic regression analysis using the non-exposed group as the reference category. Then performed multivariable logistic regression analysis to adjust for confounders. These analyses were repeated for different combinations of road types (highway or major road) and distances (50 or 150 m)					
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average population in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • validated measurements used Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for age, sex, neighbourhood socioeconomic status, and pre-existing comorbidities Outcome Assessment of outcome • from the death registration database and from hospitalization records Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • 8% loss to follow up					
Source of funding	Overall risk of bias: low f funding Government: Health Canada via an agreement with the British					
Source of furfuling	Columbia Centre for Disease Control to the Border Air Quality Study; the Centre for Health and Environment Research at The University of British					

Bibliographic reference	Gan WQ, Tamburic L, Davies HW, et al (2010) Changes in residential proximity to road traffic and the risk of death from coronary heart disease. Epidemiology (Cambridge, and Mass.) 21(5), 642-9
	Columbia, supported by the Michael Smith Foundation for Health Research; the Canadian Institutes of Health Research Frederick Banting and Charles Best Canada Graduate Scholarship and by the Michael Smith Foundation for Health Research Senior Graduate Studentship (to W.G.); and a Michael Smith Foundation for Health Research Senior Scholar Award
Comments	The study cohort was constructed using linked administrative databases that did not include certain important information about individual cardiovascular risk factors (such as active or passive smoking status, body mass index, and individual SES).

D.1.35 Garshick 2003

Gai Silick 2003					
Bibliographic reference	Garshick E, Laden F, Hart JE, et al (2003) Residence near a major road and respiratory symptoms in U.S. Veterans. Epidemiology (Cambridge, and Mass.) 14(6), 728-36				
Study design	Prospective cohort study				
Objective	To assess the relation bet respiratory symptoms in a		vehicle exhaust and		
Setting/Study location	United States				
Number of participants	2628 adults				
Selected population	No				
Participant	Description	No.	%		
characteristics	Sex	Not reported	Not reported		
	Age (years)	Not reported	Not reported		
	Ethnicity	Not reported	Not reported		
	(Maintenance) medication use	Not reported	Not reported		
	Maternal asthma and/or atopic	Not reported	Not reported		
	Education				
	<12th Grade	706	27		
	12th Grade	1011	38		
	>12th Grade	858	33		
	Missing	53	2		
	Annual family income (employment status)				
	Employed	996	38		
	Unemployed	144	5		
	Retired	1458	55		
	Missing	30	1		
	Building characteristics	Not reported	Not reported		
Inclusion criteria	Not reported				
Exclusion criteria	Not reported				

Bibliographic reference	Garshick E, Laden F, Hart JE, et al (2003) Residence near a major road and respiratory symptoms in U.S. Veterans. Epidemiology (Cambridge, and Mass.) 14(6), 728-36			
Type of pollutant/exposure	Proximity to a major road			
Pollutant/exposure assessment	residential address major road and by traffic counts are d	Exposure to motor vehicle exhaust was defined by the distance from each residential address at the time of the questionnaire mailing to the nearest major road and by the average daily traffic count for that road. Average daily traffic counts are defined as the average number of vehicles per weekday based on an average of the counts obtained throughout the year.		
Outcome	Persistent Wheez	e, Chronic Cough, C	hronic Phlegm	
Results		os (aORs) and 95% en distance of reside		
		Persistent Wheeze	Chronic Cough	Chronic Phlegm
	Distance to road (meters)	aOR (95%CI)	aOR (95%CI)	aOR (95%CI)
	≤50	1.31 (1.00, 1.71)	1.24 (0.92, 1.68)	1.18 (0.88, 1.56)
	>50 to 100	0.87 (0.61, 1.25)	0.92 (0.61, 1.39)	1.07 (0.73, 1.56)
	>100 to 200	1.11 (0.83, 1.48)	1.21 (0.87, 1.67)	1.24 (0.91, 1.68)
	>200 to 300	1.11 (0.80, 1.54)	1.30 (0.90, 1.87)	1.23 (0.87, 1.73)
	>300 to 400	1.19 (0.83, 1.72)	1.34 (0.90, 2.01)	1.32 (0.91, 1.94)
	>400	1.00	1.00	1.00
Follow up	Not reported			
Study methods	Outcome was defined by self-report of symptoms. "Chronic cough" was cough on most days for 3 consecutive months or more during the year. "Chronic phlegm" was phlegm on most days for 3 consecutive months or more during the year. "Persistent wheeze" was wheezing with a cold and occasionally apart from colds, or on most days or nights. Authors used a multiple logistic regression model to examine the association of exposure to motor vehicle exhaust with each respiratory symptom independently and to adjust for potential confounders. Exposure to motor vehicle exhaust was examined in 2 ways, by distance to the closest major road and by the average daily traffic count for that road.			
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average veteran population in the community Selection of the non-exposed cohort • no description of the derivation of the non-exposed cohort Ascertainment of exposure • validated measurements used Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for smoking, occupational dust and age Outcome			

Bibliographic reference	Garshick E, Laden F, Hart JE, et al (2003) Residence near a major road and respiratory symptoms in U.S. Veterans. Epidemiology (Cambridge, and Mass.) 14(6), 728-36
	Assessment of outcome • self-report of respiratory symptoms Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall risk of bias: Moderate (potential for response bias from outcome assessment)
Source of funding	Government: NIH/NCI
Comments	Authors reported that study lacks information on duration of residence in each address, and information regarding home exposures to nitrogen oxides from cooking or heating. Also lacks information regarding the health status of non-responders.

D.1.36 Gent 2009

Bibliographic reference	Gent JF, Belanger K, Triche EW, et al (2009) Association of pediatric asthma severity with exposure to common household dust allergens. Environmental Research 109(6), 768-774			
Study design	Prospective cohort			
Objective	To examine the dose respons household dust allergens on	se relationships and health impact of five common disease severity		
Setting/Study location	United States			
Number of participants	300 children			
Selected population	Yes – All had asthma			
Participant	Individual characteristics	l characteristics		
characteristics	Age	Mean (SD): 8.6 (2.0)		
	Sex	Male N (%): 191 (63.7) Female N (%): 109 (36.3)		
	Race / ethnicity	White/Asian/White/Asian/Another N (%): 199 (66.3) Black N (%): 46 (15.3) Hispanic N (%): 55 (18.3)		
	SES	Maternal education N (%): <12 years, 23 (7.7) 12-15 years, 162 (54.2) ≥ 16 years, 114 (33.1)		
	Asthma severity (GINA score), N (%) 0 No symptoms 1 Intermittent 2 Mild persistent	9 (3.0) 135 (45) 58 (19.3)		

Bibliographic reference		sure to co		2009) Association ousehold dust allerge	
	3 Moderate persistent 6		61 (20.3)		
	Building characteris	tics	Not report	ed	
Inclusion criteria	 Asthmatic chil physician-diag age less than asthma sympt 	nosed as 12 years		se in the previous 12 i	months
Exclusion criteria	Not reported				
Type of pollutant / exposure	dust mite (Der p 1) a	allergen			
Pollutant / exposure assessment	ELISA (enzyme-link	ed immur	osorbent as	ssay) method	
Outcome	Daily symptoms and	d medicati	on use		
Results	Adjusted Odds Ratio	o and 95%	6 Confidence	e Intervals	
			e/severe ore* aOR	Wheeze ≥30 days aOR (95%CI)	Controller meds≥9 months aOR (95%CI)
	Main living Area Der p 1 (μg/g) <0.10 0.10 to <2.0 2.0 to<10.0 ≥10.0	2.93 (1.3	11, 2.10) 37, 6.30) 13, 5.73)	1.0 1.05 (0.38,2.84) 1.55 (0.62,3.85) 2.01 (0.78,5.19)	1.0 0.61 (0.27,1.35) 2.52 (1.17,5.42) 2.17 (0.97,4.86)
	[severe persistent	2.73 (1.3 1.19 (0.4 severity s], based o	46,3.08) core (from on the Globa	1.0 1.70 (0.68,4.22) 1.60 (0.64,4.00) 3.58 (1.28,9.97) 0 [no symptoms or most initiative for Asthmasear-long prospective serverses	1.41 (0.57,3.46) edication use] to 4 i (GINA) guidelines
Follow up	1 year		,	31 1	
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness • truly representativ Selection of the non • drawn from the sa Ascertainment of ex • objective measure	e of the a exposed me comm posure	verage child cohort nunity as the	in the community	rt of study

Bibliographic reference	Gent JF, Belanger K, Triche EW, et al (2009) Association of pediatric asthma severity with exposure to common household dust allergens. Environmental Research 109(6), 768-774
	• Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	study controls for general atopy
	 study controls for additional factors - age, gender, ethnicity and mother's education.
	Outcome
	Assessment of outcome
	record linkage
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	complete follow up - all subjects accounted for
	Overall risk of bias: Low
Source of funding	Government: NIH:
Comments	

D.1.37 Gent 2012

Jenit 2012			
Bibliographic reference	Gent JF, Kezik JM, Hill ME, et al (2012) Household mould and dust allergens: exposure, sensitization and childhood asthma morbidity. Environmental research 118, 86-93		
Study design	Prospective cohort study		
Objective	To examine the association between specific allergic status, level of household exposure to specific allergens and asthma severity as measured by days of wheeze, persistent cough, rescue medication use, and an asthma severity score for the month immediately following allergy testing and sample collection.		
Setting/Study location	United States		
Number of participants	1233 children		
Selected population	Yes – children had asthma		
Participant characteristics	Description Sex Male Female Age (years) - Mean (SD) Ethnicity White	726 (58.9%) 507 (41.1%) 7.4 (1.7) 488 (39.6%)	

	Cont IF Kozik	M Lill ME at a	I (2012) Have	sahald mauld	and dust
Bibliographic	Gent JF, Kezik J allergens: expos				
reference	Environmental r	esearch 118, 80	6-93		-
	Black		239 (19	•	
	Hispanic		444 (36	•	
	Mixed and other	.,	62 (5.09	%)	
	Maternal asthma	•			
	Parental education	n Maternal)	211 (17	10/.)	
	12–15		660 (53	•	
	≥16		361 (29	•	
	Annual family inc	ome	Not rep	•	
	Building characte		Not rep	orted	
Inclusion criteria	Children were elig	gible if they were	e age 5–10		
	Had a caregiver v	vho spoke Engli	sh		
	Had active asthm	s within the past	t 12 months (v	vheeze, persis	tent cough,
	chest tightness, s medication within			e or prescription	on asunna
Exclusion criteria	Not reported				
Type of	House dust mite	(HDM)			
pollutant/exposure	Mould				
D. II. d W	Pets				at the Control
Pollutant/exposure assessment	Environmental samples were collected once during the study at the time of the enrolment home visit. Indoor airborne fungal propagules were collected using a Burkard Portable Sampler) in the main living area. The research assistant obtained a single sample using a plate with dichloran-18% glycerol (DG-18) agar, and a sampler air collection rate of 20 liters per minute [L/min] for 1 min. Samples were brought to the study laboratory for incubation at 25 °C for approximately 7 days after which the resulting fungal colonies were identified, enumerated and reported as colony-forming units per cubic meter (CFU/m³).				
Outcome	Wheeze Persistent cough Rescue medication Asthma severity s				
Results					
		Wheeze	Persistent cough	Rescue medication use	Asthma severity score
	Cladosporium >148 CFU/m ³	1.22 (0.66, 2.26)	0.98 (0.54, 1.80)	0.69 (0.37, 1.29)	1.58 (0.88, 2.83)
	HDM Der p 1 >0.10 μg/g	1.26 (0.95, 1.67)	1.18 (0.90, 1.55)	1.47 (1.11, 1.94)	1.19 (0.92, 1.55)
	HDM Der f 1 >2.1 μg/g	0.89 (0.63, 1.24)	0.90 (0.65, 1.25)	1.09 (0.78, 1.51)	1.28 (0.94, 1.74)
	Fel d 1 >0.12 µg/g	1.39 (1.05, 1.84)	0.89 (0.68, 1.17)	1.32 (1.01, 1.74)	1.14 (0.88, 1.47)

Bibliographic reference	Gent JF, Kezik J allergens: expos Environmental re	ure, sensitizat	ion and child		
	Can f 1 >1.2 μg/g	1.53 (1.09, 2.15)	1.11 (0.80, 1.56)	1.15 (0.82, 1.62)	1.15 (0.83, 1.58)
Follow up	1 months				
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativene truly representation of the none of th	tive of the avera on-exposed con- if the derivation exposure urement used at outcome of in cohorts on the boor gender, age, ousing type, small atcome otoms ag enough for out w up of cohorts up - all subjects	age child in the nort of the non-expected was not asis of the designation of the designat	posed cohort t present at sta sign or analysi medication use NO ₂ concentra	s e, general ation, and
Source of funding	Government: Nati	onal Institutes o	of Health:		
Comments					

D.1.38 Habre 2014

Bibliographic reference	Habre R, Moshier E, Castro W, et al (2014) The effects of $PM_{2.5}$ and its components from indoor and outdoor sources on cough and wheeze symptoms in asthmatic children. Journal of exposure science & environmental epidemiology 24(4), 380-7
Study design	Cohort
Objective	To investigate the association of indoor PM _{2.5} of outdoor origin and its components with cough and wheeze symptom scores in asthmatic children

Bibliographic reference	Habre R, Moshier E, Castro W, et al (2014) The effects of $PM_{2.5}$ and its components from indoor and outdoor sources on cough and wheeze symptoms in asthmatic children. Journal of exposure science & environmental epidemiology 24(4), 380-7
Setting/Study location	United States
Number of participants	36
Selected population	Yes (focused on children with asthma)
Participant characteristics	Building characteristics: Location: Urban Dwelling type: Not reported Building age: Not reported Type of ownership/tenancy: Not reported Parental characteristics: Age (Mean & range): 10 years (6 – 15) Smoker in home: Not reported Race: Hispanic> 23 (64%) Black: 12 (36%) Selected: Asthma (36 (100%)
Inclusion criteria	Children aged 6–14 years, with doctor-diagnosed persistent moderate-to-severe asthma defined by at least one of the following: daily use of controller medication for at least 3 months over the past year, use of a bagonist at least four times per month in any one of the past 3 months, or nocturnal awakenings twice a month in the past 3 months. Have slept at the household identified as the primary residence at least five times a week.
Exclusion criteria	More than 1 smoker in the household, family planning to move within the next 6 months, or no access to a phone. Presence of haematological, endocrine, or cardiac condition that required the use of daily medication, Presence of clotting disorder, Presence of severe mental disability that interfered with answering questions or following instructions.
Type of pollutant/exposure	Particulate matter 2.5
Pollutant/exposure assessment	A Multi-Pollutant Sampler (MPS) was placed in participants' living rooms.to collect 7-day integrated $PM_{2.5}$ samples on 37mm Teflon filters. $PM_{2.5}$ mass concentration was measured gravimetrically using standard methods. A Dustrak aerosol monitor (TSI, Model 8520) with a 2.5-mm inlet measured $PM_{2.5}$ mass concentrations at 10-min intervals. Biases in light-scattering devices have been previously described.20 Therefore, linear mixed effects models were used to calibrate the continuous Dustrak readings to weekly gravimetric $PM_{2.5}$ mass concentrations by fitting a random intercept and slope for each of the 13 Dustrak monitors used. After calibration, average daily indoor $PM_{2.5}$ mass concentrations were calculated starting at 0900 hours.
Outcome	Wheeze
Catoonio	

Bibliographic reference	Habre R, Moshier E, Castro W, et al (2014) The effects of PM _{2.5} and its components from indoor and outdoor sources on cough and wheeze symptoms in asthmatic children. Journal of exposure science & environmental epidemiology 24(4), 380-7 Cough Participants or their caretakers were asked to record their asthma cough and wheeze scores in a diary every day, with a score of zero equivalent to none, one as mild, two as moderate, and three as severe.		
Results	per SD increase PM _{2.5} (Indoor) (SD=17.3) PM _{2.5} (indoor sources) (SD=17.6) PM _{2.5} (outdoor sources) (SD=61.) PM _{2.5} (Indoor) PM _{2.5} (indoor sources) PM _{2.5} (outdoor sources)	aOR (95%CI) for cough 1.22 (0.91, 1.63) 1.20 (0.88, 1.64) 1.27 (0.90, 1.77) aOR (95%CI) for wheeze 1.57 (1.09, 2.26) 1.55 (1.05, 2.28) 1.13 (0.75, 1.72)	
Follow up Newcastle-Ottawa Scale	Selection Representativeness of the exposed cohort truly representative of the average child with Selection of the non-exposed cohort drawn from the same community as the extended assertainment of exposure secure record (objective sampling) Demonstration that outcome of interest was as a line of the comparability Comparability Comparability of cohorts on the basis of the comparability of secure of the comparability Outcome Assessment of outcome self-report Was follow-up long enough for outcomes to complete follow up - all subjects accounted Overall assessment: Moderate (self-report	posed cohort not present at start of study design or analysis	
Source of funding	Industry: (Electric Power Research Institute Academic: Harvard School of Public Health		
Comments			

D.1.39 Hagerhed-Engman 2009

Bibliographic reference	Hagerhed-Engman L, Sigsgaard T, Samuelson I et.al (2009) Low home ventilation rate in combination with mouldy odor from the building structure increase the risk for allergic symptoms in children. Indoor air 19(3), 184-92					
Study design	Nested case	Nested case control				
Objective	diseases in p	re-school childre		gnosed asthma a related problems spectors		
Setting/Study location	Sweden					
Number of participants	400 children					
Selected population	No					
Participant	Description	Control		Cases		
characteristics		No.	%	No.	%	
	Sex					
	Male	114	56.4	113	57.1	
	Female	88	43.6	85	42.9	
	Age (years)					
	2-4	74	36.6	77	38.9	
	5-6	80	39.6	85	42.9	
	7-8	48	23.8	36	18.2	
	Ethnicity	Not reported Not reported				
	Parental asthma and/or atopic					
	Atopic symptom in at least one parent	87	43.1	161	81.3	
	Parental education	Not reported		Not reported		
	Annual family income	Not reported		Not reported		
	Building char	Building characteristics				
	Type of build	ing				
	Single family house	172	85.1	161	81.3	
	Row house	11	5.4	12	6.1	
	Multi-family house	19	9.4	25	12.6	
	Type of venti	lation system				
	Natural	147	70.3	124	62.6	
	Exhaust	37	18.3	56	28.3	

	Hagerhed-Eng	ıman L, Sigsg	aard T,	Samuels	on I et.al (2	2009) Low home
Bibliographic	ventilation rate in combination with mouldy odor from the building structure increase the risk for allergic symptoms in children. Indoor air					
reference	19(3), 184-92					
	Balanced 2	23	11.4		18	9.1
	Construction period					
	Before 6 1940	67	33.2		47	23.7
	1941-1960	35	17.3		37	18.7
	1961-1970	25	12.4		28	14.1
	1971-1983	14	21.8		63	31.8
	1984- 3 present	31	15.3		23	11.6
Inclusion criteria	Inclusion criteria for cases Reports of at least two symptoms of either wheezing (without a cold), rhinitis (without a cold) or eczema Inclusion criteria for controls No symptoms For both groups, they had to: Agree to co-operate Not have rebuilt their homes because of moisture problems Not have changed residence since the first questionnaire					
Exclusion criteria	Not reported	Not reported				
Type of pollutant/exposure	Dampness and	Dampness and mould				
Pollutant/exposure assessment	Blood samples were drawn and screened for sensitisation to common allergens and sensitised children were further tested for specific IgE (RAST) for cat, dog, horse, birch-, mugwort- and timothy grass pollen and mould. Samples of dust and air were collected in the children's bedroom and living room for analyses of chemical and biological compounds and measurements of ventilation rate, temperature and relative humidity were made. Each home scored on a three level scale for each index: grade 0 (no remarks), grade 1-2 (mild) and grade 3 (severe)					
Health outcome	Physician diagnosed asthma and allergic diseases (rhinitis and eczema)					
Results	Adjusted odds association bet					
		Asthma		Rhinitis		Eczema
	Mould - Mild	1.30 (0.73,	2.29)	2.23 (1.	17, 4.24)	1.86 (1.04, 3.30)
	Mould - Severe	1.28 (0.60,	2.73)	2.45 (1.	08, 5.54)	1.93 (0.91, 4.12)
	Damp stains - Mild	0.86 (0.47,	1.60)	1.39 (0.	73, 2.67)	0.64 (0.34, 1.20)
	Damp stains - Severe	0.28 (0.5,	1.52)	0.37 (0.	04, 3.43)	0.30 (0.06, 1.57)
	Floor damp - Mild	0.82 (0.28,	2.42)	1.16 (0.	36, 3.76)	0.78 (0.26, 2.34)
	Floor damp - Severe	0.82 (0.05,	12.33)	1.58 (0.	10, 26.14)	1.88 (0.20, 17.29)
Follow up	12 months					

Bibliographic reference	Hagerhed-Engman L, Sigsgaard T, Samuelson I et.al (2009) Low home ventilation rate in combination with mouldy odor from the building structure increase the risk for allergic symptoms in children. Indoor air 19(3), 184-92
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • objective samples Demonstration that outcome of interest was not present at start of study Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for parental smoking • study controls for additional factors – age, gender, type of building, construction period and family history of asthma and allergy Outcome Assessment of outcome • independent blind assessment Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall risk of bias: Low
Source of funding	Study supported by the Swedish Research Council for Environment, Agricultural Sciences and Spatial Planning (Formas), Swedish Asthma and Allergy Association's research Foundation, the Swedish Foundation for Health Care Sciences and Allergy Research, SP Technical Research Institute of Sweden
Comments	Authors suggest that study support the hypothesis that odour from microbiological and/or chemical degradation of building material can be transported into the indoor environment and increase the risk for allergic symptoms among children as well as sensitization.

D.1.40 Hagmolen of Ten Have 2007

Hagmolen of Ten		_	
Bibliographic reference	Hagmolen of Ten Have W, van den Berg NJ, van der Palen J et al (2007) Residential exposure to mould and dampness is associated with adverse respiratory health. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 37(12), 1827-32		
Study design	Prospective cohort study		
Objective	to investigate the association of reported exposure to mould and dampness and respiratory health in a general practice-based population of asthmatic children		
Setting/Study location	Netherlands		
Number of participants	526 children		
Selected population	Yes - all had asthma		
Participant characteristics	Description Sex Female Age (years)- Mean (SD) Ethnicity Education Parental less than 11 years education Annual family income		240 (45.6%) 11 (2.5) Not reported 70 (13.3%) Not reported
Inclusion criteria	If at least two prescriptions of b2-mimetics or an inhaled corticosteroid (ICS) were prescribed in the year before invitation		
Exclusion criteria	FEV1 was<75% predicted.		
Type of pollutant/exposure	Dampness Pets		
Pollutant/exposure assessment	Questionnaire		
Outcome	Risk of airway hyper-responsiveness		
		aOR	(95%CI)
	Mould and dampness, living or child's sleeping room	3.95 ((1.82, 8.57)
	Pet ownership	1.17 ((0.70, 1.94)
Follow up	2 weeks		
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed co • selected group of children with astheta Selection of the non-exposed cohort • drawn from the same community as Ascertainment of exposure • written self-report Demonstration that outcome of interes • No Comparability Comparability of cohorts on the basis	ma the ex	not present at start of study

Bibliographic reference	Hagmolen of Ten Have W, van den Berg NJ, van der Palen J et al (2007) Residential exposure to mould and dampness is associated with adverse respiratory health. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 37(12), 1827-32
Study design	Prospective cohort study
	study controls for smoking status
	 study controls for any additional factors as follows - gender, history of inhalant allergy, history of rhinitis, family history of asthma, the level of parental education, pet ownership, and use of ICS in the previous 4 months, season of study assessment, health care centre, exposure to environmental smoking by parents or household members Outcome Assessment of outcome self-report Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts
	• complete follow up - all subjects accounted for
	Overall level of bias: High (concerns over self-reporting of outcome and exposure)
Source of funding	Industry: GlaxoSmithKline
Comments	

D.1.41 Harris 2007

Bibliographic reference	Harris J M, Williams H C, White C, et al (2007) Early allergen exposure and atopic eczema. The British journal of dermatology 156(4), 698-704		
Study design	Prospective cohort study		
Objective	To analyse allergen exposure years,	and eczema outcomes measured up to age 8	
Setting/Study location	United Kingdom		
Number of participants	593 children		
Selected population	No		
Participant characteristics	Description Sex Maternal age (years) Ethnicity Maternal asthma and/or atopic SES	Not reported Not reported Not reported Not reported Not reported	
Inclusion criteria	Not reported		
Exclusion criteria	Not reported		

Bibliographic reference	Harris J M, Williams H C, Wh and atopic eczema. The Briti				
Type of pollutant/exposure	Dust mite and cat allergen exposure				
Pollutant/exposure assessment	Dust samples were collected from the living room floor 8 weeks after birth. These samples were assayed for concentrations of house dust mite and cat allergen using standard techniques.				
Outcome	Eczema				
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs)				
		Eczema	Visible flexural dermatitis		
		aOR (95%CI)	aOR (95%CI)		
	Quintile of house dust mite exp	oosure (units not rep	oorted)		
	1 (lowest) (0.02 to 0.27)	1.00	1.00		
	2 (0.28 to 0.81)	1.01 (0.53, 1.92)	1.17 (0.58, 2.34)		
	3 (0.82 to 2.22)	1.37 (0.74, 2.55)	1.73 (0.87, 3.46)		
	4 (2.23 to 7.75)	0.66 (0.34, 1.29)	0.88 (0.43, 1.81)		
	5 (highest) (7.76 to 384.97)	0.71 (0.37, 1.37)	0.96 (0.47, 1.94)		
	Quintile of cat allergen exposu	, ,	`		
	1 (lowest) (0.01 to 0.44)	1.00	1.00		
	2 (0.45 to 1.04)	1.42 (0.72, 2.81)	1.28 (0.64, 2.56)		
	3 (1.05 to 3.33)	1.41 (0.71, 2.79)	0.75 (0.36, 1.55)		
	4 (3.34 to 44.72)	1.31 (0.65, 2.62)	1.18 (0.59, 2.38)		
	5 (highest) (44.73 to 14,151.32)	1.41 (0.72, 2.75)	0.96 (0.48, 1.91)		
Follow up	8 years				
Risk of bias	Selection				
(Newcastle-Ottawa	Representativeness of the exp	osed cohort			
Scale)	 truly representative of the ch 	•	1		
	Selection of the non-exposed of				
	drawn from the same community as the exposed cohort				
	Ascertainment of exposure				
	Objective measurement Personalization that subsample of interest was not present at start of study.				
	Demonstration that outcome of interest was not present at start of study				
	Yes Comparability				
	Comparability Comparability of cohorts on the basis of the design or analysis				
	study controls for all variables				
	Outcome				
	Assessment of outcome				
	Clinical diagnosis				
	Was follow-up long enough for outcomes to occur • Yes				
	Adequacy of follow up of cohorts				
	• complete follow up - all subjects accounted for				
	Overall risk of bias: Low				

Bibliographic reference	Harris J M, Williams H C, White C, et al (2007) Early allergen exposure and atopic eczema. The British journal of dermatology 156(4), 698-704
Source of funding	Charity: The Colt foundation
Comments	

D.1.42 Hunt 2011

Bibliographic reference	Hunt A, Crawford JA, Rosenbaum P F, et al (2011) Levels of household particulate matter and environmental tobacco smoke exposure in the first year of life for a cohort at risk for asthma in urban Syracuse, NY. Environment International 37(7), 1196-205
Study design	Birth cohort
Objective	To investigate possible associations between indoor exposures and infant health status (in particular wheezing)
Setting/Study location	United States
Selected population	Yes (selected on based of being at risk of asthma)
Number of participants	103 mother baby-dyads
Participant characteristics	Building characteristics: Location: urban Dwelling type: Not reported Building age: Not reported Type of ownership/tenancy: Not reported Parental characteristics: Age: Not reported Current smoker (mother): 55 (54%) Hay fever: Not reported Atopy: Not reported but all mothers had asthma
Inclusion criteria	Documented history of maternal asthma Expectation of the mother residing in same residence for at least 1 year or an adjacent urban location Infant criteria Gestational age ≥ 37 weeks Birthweight ≥ 2500 g Absence of any major congenital abnormality Singleton birth
Exclusion criteria	None reported
Type of pollutant/exposure	Particulate matter 2.5 ≥15μg/m³
Pollutant/exposure assessment	Particulate matter was collected using size selective Harvard impactors operating at 10 L/min placed in the living room
Outcome	Wheeze defined as Primary-care provided-documented wheezing, reactive airway disease, asthma or bronchiolitis or

Bibliographic reference	Hunt A, Crawford JA, Rosenbaum P F, et al (2011) Levels of household particulate matter and environmental tobacco smoke exposure in the first year of life for a cohort at risk for asthma in urban Syracuse, NY. Environment International 37(7), 1196-205
	Wheeze heard on physical examination by the nurse practitioner or A prescription for bronchodilator, inhaled steroid or steroid pulse prescription document in the medical records
Results	Wheeze PM 2.5 ≥15µg/m³=aOR 4.21 (1.36, 13.03)
Follow up	12 months
Newcastle-Ottawa Scale	Selection Representativeness of the exposed cohort • truly representative of the average infant in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • secure record of objective measurement Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for gender, maternal age and education, season of home visit and presence of carpeting Outcome Assessment of outcome • independent blind assessment or medical records Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall assessment: Low
Source of funding	Government: Research grant (USEPA)
Comments	Particulate matter was primarily sourced from environmental tobacco smoke

D.1.43 Hart 2014

Bibliographic reference	Hart JE, Chiuve S E, Laden F, et al (2014) Roadway proximity and risk of sudden cardiac death in women. Circulation 130(17), 1474-82
Study design	Prospective cohort study
Objective	To determine whether roadway proximity was associated with an increased risk of Sudden Cardiac Death (SCD)
Setting/Study location	United States

Bibliographic reference	Hart JE, Chiuve S E, Lad sudden cardiac death in		dway proximity and risk of 30(17), 1474-82	
Number of participants	107130 women			
Participant	Description	No.	%	
characteristics	Sex	Not reported	Not reported	
	Age (years); mean (SD)	64.3 (10.0)		
	Ethnicity			
	White	100702	94	
	Black	2143	2	
	Other/multiple	5356	5	
	(Maintenance) medication	use		
	Aspirin use			
	<1 times/week	61064	57	
	1–6 times/week	19283	18	
	≥7 times/week	10713	10	
	Maternal asthma and/or atopic	Not reported	Not reported	
	Parental history of MI			
	None	79276	74	
	Before 60 years of age	12855	12	
	After 60 years of age	16070	15	
	Parental education	Not reported	Not reported	
	Annual family income	Not reported	Not reported	
	Building characteristics	Not reported	Not reported	
Inclusion criteria	Women were included in the questionnaires Had at least 1 home addres No cancer (other than non No cardiovascular disease	ess that was geocoded -melanoma skin cancer	to the street-segment level	
Exclusion criteria	No reported			
Type of pollutant/expos ure	Proximity to traffic			
Pollutant/expo sure assessment	traffic exposure. Distance Information System softwa StreetMap Pro 2007 road s Census Feature Class Cowith limited access, division	(in meters) was determ are (ArcGIS 10.2, ESRI, segments were selecte des: A1, primary roads, on between the opposing major, non-interstate hi	d to include the 3 largest US typically interstate highways, g directions of traffic, and ghways and major roads without	
Outcome	Sudden Cardiac Death (SCD)			
Results	Adjusted hazard ratios (albetween proximity to traffic		ce intervals (CIs) for association Death	

Bibliographic reference	Hart JE, Chiuve S E, Laden F, et al (20 sudden cardiac death in women. Circu	
		Sudden Cardiac Death
	Distance	aHRs (95%CI)
	0–49	1.38 (1.04, 1.82)
	50–199	1.17 (0.91, 1.51)
	200–499	1.20 (0.93, 1.53)
	≥500	Reference
Follow up	Over 26 years of follow-up	
Study methods	On all questionnaires, authors inquire about the occurrence of physician diagnosed coronary heart disease (CHD) events, and deaths are identified by reports from next-of-kin or postal authorities or by searches of the National Death Index. SCDs were confirmed by physician review of medical records and next-of-kin reports on the circumstances surrounding the death if not adequately documented in the medical record. Cardiac deaths were considered sudden if the death or cardiac arrest occurred within 1 hour of the onset of symptoms. Time-varying Cox proportional hazards models were used to assess the relationship of outcome with roadway proximity. All models were based on a biennial time scale and were stratified by age in months and time-period.	
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed coho • truly representative of the average femal Selection of the non-exposed cohort • no description of the derivation of the non-exposure • validated measurements used Demonstration that outcome of interest work • Yes Comparability Comparability Comparability of cohorts on the basis of the study controls for age, race, comorbidity blood pressure, stroke, or coronary head Outcome Assessment of outcome • Physician diagnosed Was follow-up long enough for outcomes • Yes Adequacy of follow up of cohorts • no statements Overall risk of bias: low	ale population in the community on-exposed cohort vas not present at start of study the design or analysis ies: incidence of high cholesterol, high art disease
Source of funding	Government: National Institutes of Healt Charity ; American Heart Association	
Comments	Authors reported that measure of exposure true traffic exposures such as noise or powith information on temporal changes in triggering of events. These are expected misclassifications of exposure, which wou	exposures that may be associated with to lead to non-differential

D.1.44 Harville 2018

Bibliographic reference	Harville EW, Rabito FA. (2018) Housing conditions and birth outcomes: The National Child Development Study.			
Study design	Prospective cohort study			
Objective	To examine if there is an association between poor housing/living conditions and undesirable birth outcomes			
Setting/Study location	Unspecified location	ns in the UK		
Number of dwellings and participants	Number of dwellings Number of participa			
Selected population	No			
Building and Participant characteristics	Building characteristics: Location: not reported Dwelling type: detached, 22.3%; semi-detached, 70.7%; apartment or room, 6.2%; caravan/houseboat/mobile home, 0.7% Building age: not reported Type of ownership/tenancy: rented, 15%; owned, 85% Double glazing: 32.3% Central heating: 26.2% Participant characteristics: Age at included pregnancy: <24 years, 4.4%; 25-28 years, 18.7%; 29-30 years, 18.6%; >30 years, 58.2% Smoking during pregnancy: 29.1% BMI at age 33: <20, 10%; 20-24, 53.6%; 25-29, 24.7%; ≥30, 11.7%			
Inclusion criteria	Women participating in a national birth cohort study who had given birth at least once while living in their current property were included. If they reported more than 1 birth, outcomes of the latest one were used.			
Exclusion criteria	Not reported			
Building factor/exposure	Location and severi	ty of mould		
Building factor/exposure assessment	Building factors wer reported questionna	e ascertained by asl iire.	king participants to	complete a self-
Outcome	Low birth weight, pr	eterm birth, small fo	r gestational age	
Results		Odds ratio (95%C	l)	
	Building characteristic	Low birthweight	Preterm birth	Small for gestational age
	Mould			
	Mould anywhere	1.98 (1.13, 3.47)	1.23 (0.69, 2.19)	2.06 (1.25, 3.38)
	Serious mould anywhere	2.42 (1.20, 4.86)	1.60 (0.79, 3.23)	1.89 (0.96, 3.71)
	Mould in bedroom	1.87 (0.68, 5.15)	2.23 (0.94, 5.28)	1.35 (0.53, 3.43)
	Mould in kitchen	2.24 (0.75, 6.66)	0.82 (0.19, 3.52)	1.04 (0.30, 3.58)

Bibliographic reference		o FA. (2018) Housi Development Stud		d birth outcomes:
reference	Serious mould in bedroom	1.47 (0.33, 6.61)	1.82 (0.52, 6.36)	0.85 (0.19, 3.86)
	Serious mould in kitchen	2.25 (0.65, 7.74)	1.12 (0.26, 4.83)	0.48 (0.06, 3.63)
	Renovations			
	Against damp	1.04 (0.52, 2.11)	0.32 (0.11, 0.89)	1.28 (0.69, 2.35)
	Roof	0.80 (0.42, 1.52)	0.25 (0.10, 0.62)	0.97 (0.55, 1.70)
	Gutter	1.15 (0.65, 2.04)	0.55 (0.29, 1.04)	1.08 (0.64, 1.82)
	Point	1.30 (0.67, 2.51)	0.89 (0.45, 1.77)	1.26 (0.68, 2.33)
	Glazing	0.96 (0.57, 1.64)	0.86 (0.52, 1.41)	1.03 (0.64, 1.66)
	Heating	1.25 (0.73, 2.13)	1.15 (0.70, 1.90)	1.27 (0.78, 2.05)
	Garage	1.44 (0.60, 3.48)	1.59 (0.74, 3.43)	1.09 (0.45, 2.60)
	Extension	1.20 (0.67, 2.17)	0.94 (0.53, 1.68)	0.94 (0.54, 1.65)
	Loft	1.88 (0.65, 5.47)	1.34 (0.47, 3.84)	1.02 (0.31, 3.37)
	Wiring	1.01 (0.57, 1.79)	0.81 (0.49, 1.41)	1.07 (0.64, 1.80)
	Plumbing	1.38 (0.78, 2.44)	0.96 (0.55, 1.67)	1.02 (0.59, 1.78)
Follow up	33 years			
Study methods	child development of were obtained from living in the same pro- questionnaire in whistory, renovations of mould. Participandue date, the gestate	roperty. Exposure da ich participants prov and housing condit its were then asked	se participants, 33- given birth at leas ata were taken fror rided information or ions; namely, presor if they had been pro- irthweight of the ba	year follow-up data t once and were n a self-reported n residential ence and severity regnant, estimated
Newcastle-Ottawa Scale	Selection Representativeness • selected group – v Selection of the non	s of the exposed coh women who were pa n-exposed cohort ame community as the xposure	nort articipating in a birt he exposed cohort	ŕ

Bibliographic reference	Harville EW, Rabito FA. (2018) Housing conditions and birth outcomes: The National Child Development Study.
	• Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	 study controls for BMI, education, social class, age, smoking, home ownership, housing type, number of residents, year of birth and time in the house
	Outcome
	Assessment of outcome
	Self-reported
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	complete follow up - all subjects accounted for
	Overall risk of bias: moderate (Concerns over self-report of exposure an outcomes)
Source of funding	Authors reported that no external funding was provided for this study
Comments	

D.1.45 Heinrich 2013

= 0 . 0				
Bibliographic reference	Heinrich J, Thiering E, Rzehak P, et al (2013) Long-term exposure to NO_2 and PM_{10} and all-cause and cause-specific mortality in a prospective cohort of women. Occupational and environmental medicine 70(3), 179-86			
Study design	Prospective cohort study			
Objective		To assess whether long-term exposure to air pollution is associated with all-cause and cause-specific mortality during a period of declining particulate matter concentrations.		
Setting/Study location	Germany			
Number of participants	4800 women			
Selected population	No			
Participant	Description	No.	%	
characteristics	Sex	Not reported	Not reported	
	Maternal age (years)	Not reported	Not reported	
	Ethnicity	Not reported	Not reported	
	(Maintenance) medication use	Not reported	Not reported	
	Maternal asthma and/or atopic	Not reported	Not reported	
	SES			
	Parental education	Not reported	Not reported	
	Annual family income	Not reported	Not reported	

Bibliographic reference	and PM ₁₀ an	d all-cause an	hak P, et al (2013) L d cause-specific mo nd environmental m	ortality in a pros	pective cohort
	Building char	acteristics N	lot reported	Not reported	
Inclusion criteria	Not reported				
Exclusion criteria	Not reported				
Type of pollutant/expos ure	Particulate m	atter (PM), NO	₂ and proximity to the	major road	
Pollutant/expo sure assessment	(TSP) as suri NO ₂ concent levels were n calculated as	Authors used NO_2 and PM_{10} derived from total suspended particulate matter (TSP) as surrogates for air pollution. NO_2 concentrations were measured by means of chemiluminescence, and TSP levels were measured at state routine monitoring sites by β absorption. PM_{10} was calculated as $0.71\times TSP$ for all monitoring sites. The factor of 0.71 was derived from parallel measurements of PM_{10} and TSP at seven monitoring sites in the			
Outcome	All-cause, Ca	ardiopulmonary	, Lung cancer, Respi	ratory	
Results	between all-c	ause, cardiopul	and 95% confidence monary and lung car entrations and distan	ncer mortality, and	d an IQR
		All-cause	Cardiopulmonary	Lung cancer	Respiratory
		aRR (95%CI)	aRR (95%CI)	aRR (95%CI)	aRR (95%CI)
	Distance from	n home to a ma	jor road		
	>50 m	1.00	1.00	1.00	1.00
	≤50 m	1.42 (1.12, 1.79)	1.95 (1.37, 2.77)	0.62 (0.15, 2.60)	3.54 (1.49, 8.40)
	1-year average	ge			
	NO ₂	1.18 (1.07, 1.30)	1.55 (1.30, 1.84)	1.46 (0.92, 2.32)	1.13 (0.71, 1.80)
	PM ₁₀	1.15 (1.04, 1.27)	1.39 (1.17, 1.64)	1.84 (1.23, 2.74)	0.96 (0.60, 1.53)
Follow up	21.9 years				
Study methods	proportional had participants with difference be fatality. For the calculated as participants withe study was	nazards models who passed awa tween the date nose alive at the the difference l who moved during s calculated as	lity and exposure were including adjustmentary, the time in the sture of the baseline crosses and of follow-up, the between the start and follow-up and were the difference between start and place of reside	t for potential condy was calculated -sectional study as time in the study dend of follow-up as subsequently losen the start of follow the start of follow.	founders. For d as the and the date of y was b. For ost, the time in ow-up and the
Risk of bias (Newcastle- Ottawa Scale)	Selection Representati truly repres	veness of the ex	xposed cohort average female popu		

Bibliographic reference	Heinrich J, Thiering E, Rzehak P, et al (2013) Long-term exposure to NO_2 and PM_{10} and all-cause and cause-specific mortality in a prospective cohort of women. Occupational and environmental medicine 70(3), 179-86
	 no description of the derivation of the non-exposed cohort Ascertainment of exposure
	validated measurement used
	Demonstration that outcome of interest was not present at start of study
	• Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	 study controls for age, educational level and smoking status
	Outcome
	Assessment of outcome
	record linkage and death certificates
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	• no statements
	Overall risk of bias: low
Source of funding	Government: North Rhine-Westphalia State Environment Agency (LANUV-NRW) of the Ministry of the Environment and Conservation, Agriculture and Consumer Protection North Rhine-Westphalia (MUNLV), Dusseldorf, Germany.
Comments	

D.1.46 Henderson 2008

Bibliographic reference	Henderson J, Sherriff A, Farrow A, et al (2008) Household chemicals, persistent wheezing and lung function: Effect modification by atopy? European Respiratory Journal 31(3), 547-554		
Study design	Prospective cohort study		
Objective	To investigate the effects of maternal chemical use during pregnancy on wheezing patterns in children whether increasing use is associated with decrements in lung function at age 8.5 yrs.; and 3) whether atopy modifies these associations		
Setting/Study location	United Kingdom		
Number of participants	14,541 pregnant women		
Selected population	No		
Participant characteristics	Description Sex Age (years) (reported as maternal) < 25 years ≥25 years Ethnicity Education	3711 521 (14.0%) 3190 (86.0) Not reported	3451 539 (15.6%) 2912 (84.4) Not reported

Bibliographic reference	Henderson J, Sherriff A, Farrow A, et a persistent wheezing and lung function European Respiratory Journal 31(3), 5	n: Èffect modification	
	None/CSE Vocational O-level A-level Degree Annual family income	487 (13.3%) 315 (8.6%) 1325 (36.3%) 938 (25.7%) 590 (16.1%) Not reported	464 (13.6%) 281 (8.3%) 1194 (35.1%) 902 (26.5%) 563 (16.45%) Not reported
Inclusion criteria	Expected date of delivery between April Place of residence within the three Bristo county of Avon, UK		
Exclusion criteria	Not reported		
Type of pollutant/exposur e	Composite household chemical exposure chemicals	e – VOCs and non V	OCs containing
Pollutant/exposur e assessment	Questionnaire		
Outcome	early-onset transient wheeze, i.e. wheezed at 0–18 months but not at 69–81 months intermediate-onset transient wheeze, i.e. no wheeze at 0–18 months and wheeze at 18–42 months and no wheeze at 69–81 months; early-onset persistent wheeze, i.e. wheeze at 0–18 and 69–81 months; intermediate onset persistent wheeze, i.e.no wheeze at 0–18 months and wheeze at 18–42 and 69–81 months late onset wheeze, i.e. onset of wheeze after 42 months and before 81 months		months and 31 months; 3 months and
Results	Adjusted odds ratios (aORs) and 95% cobetween Composite Household Chemica		
		Composite Ho Exposure	usehold Chemical
		aOR (95%CI)	
	Early-onset transient wheeze Early onset persistent wheeze Intermediate-onset transient wheeze Intermediate-onset persistent wheeze Late onset wheeze	1.07 (0.99–1.1 1.21 (1.08–1.3 1.13 (1.01–1.2 1.11 (0.91–1.3 1.07 (0.88–1.2	88) 28) 36)
Follow up	81 months		
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average pregnant woman in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • questionnaire Demonstration that outcome of interest was not present at start of study • Yes		

Bibliographic reference	Henderson J, Sherriff A, Farrow A, et al (2008) Household chemicals, persistent wheezing and lung function: Effect modification by atopy? European Respiratory Journal 31(3), 547-554
	Comparability Comparability of cohorts on the basis of the design or analysis • study controls for exposure to environmental tobacco smoke • study controls for additional factors - overcrowding in home, highest maternal education level, housing tenure, sex, maternal history of asthma, maternal parity, maternal age at delivery, smoking during pregnancy, month of completion of chemicals questionnaire and maternal hours worked outside home Outcome Assessment of outcome • self-report Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • subjects lost to follow up unlikely to introduce bias - description provided of those lost Overall risk of bias: High (concerns over self-report of exposure and outcomes)
Source of funding	Government: The UK Medical Research Council, Charity: the Wellcome Trust UK Academic: University of Bristol
Comments	The composite household chemical exposure (CHCE) comprises of 11 different products (disinfectant; bleach; carpet cleaner; window cleaner; dry cleaning fluid; aerosols, turpentine/white spirit, air fresheners (spray, stick or aerosol); paint stripper; paint or varnish; and pesticides/insect killers). A simple score for frequency of use of each product was derived: 0 for not at all; 1 for less than once a week; 2 for about once a week; 3 for most days; and 4 for every day. The scores for each product were summed to produce a composite household chemical exposure (CHCE) score for each respondent

D.1.47 Herr 2011

Bibliographic reference	Herr M, Nikasinovic L, Foucault C, et al (2011) Can early household exposure influence the development of rhinitis symptoms in infancy? Findings from the PARIS birth cohort. Annals of Allergy, and Asthma and Immunology 107(4), 303-309
Study design	Prospective cohort
Objective	To investigate risk factors for rhinitis symptoms in infants
Setting/Study location	France
Number of participants	1850 infants
Selected population	No
	Individual characteristics

Bibliographic reference	Herr M, Nikasinovic L, Foucault C, et al (2011) Can early household exposure influence the development of rhinitis symptoms in infancy? Findings from the PARIS birth cohort. Annals of Allergy, and Asthma and Immunology 107(4), 303-309		
Participant characteristics	Age (months – Mean (SD)		19 (2)
	Sex, n (%) Male		925 (50)
	Race / ethnicity		Not reported
	SES, n (%)	·	
	High		1231 (66.5)
	Intermediate		476 (25.7)
	Low		143 (7.7)
	Building characteristics		
	Apartment, %		92
	Gas cooking or heating, n (%)		1,031(56.4)
	Indoor renovation activities in the h	• •	780 (42.7%)
	Indoor renovation activities in the c	hild's bedroom in the	686 (37.5%)
	past year		784 (42.9%)
	Presence of particle-board furniture less than 1 year old in the home		540 (20 5%)
	Presence of particle-board furniture less than 1 year old		540 (29.5%)
	in the child's bedroom		332 (18.2%)
	Presence of moulds in the home		300 (16.4%)
	Use of an air dampener		761 (41.6%)
	Carpet in the bedroom of the child		968 (52.9%)
	New mattress in the baby's bedding Use of an anti-dust mite cover in the baby's bedding		636 (35.0%)
Inclusion criteria	Infants included in the PARIS birth cohort		
Exclusion criteria	Not reported		
Type of pollutant / exposure	VOCs		
Pollutant / exposure assessment	Not reported		
Outcome	Rhinitis symptoms in last 12 months		
Results	Adjusted Odds Ratio and 95% Confidence Intervals		
		Allergic rhinitis	Non-allergic Rhinitis
	Presence of particle board less than 1 year old in child's room	1.09 (0.63, 1.87)	1.87 (1.21, 2.90)
Follow up	18 months		
Risk of bias	Selection		
(Newcastle-	Representativeness of the exposed cohort		
Ottawa Scale)	truly representative of the average infant in the community		

Bibliographic reference	Herr M, Nikasinovic L, Foucault C, et al (2011) Can early household exposure influence the development of rhinitis symptoms in infancy? Findings from the PARIS birth cohort. Annals of Allergy, and Asthma and Immunology 107(4), 303-309
	Selection of the non-exposed cohort
	 drawn from the same community as the exposed cohort
	Ascertainment of exposure
	• no description
	Demonstration that outcome of interest was not present at start of study
	• Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	study controls for environmental tobacco smoke
	 study controls for additional factors including sex, socioeconomic status, duration of maternal breastfeeding and presence of siblings
	Outcome
	Assessment of outcome
	self-report (parent)
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	complete follow up - all subjects accounted for
	Overall risk of bias: High (concerns over self- report of exposure and outcomes)
Source of funding	Government: Social, Childhood and Health Direction (DASES) of the Paris Council;
	Academic: Paris Descartes University;
	Industry: French Health Insurance System; Biochemistry Laboratory of the Groupe Hospitalier Trousseau-La Roche Guyon, Assistance Publique–Hôpitaux de Paris.
Comments	Allergic rhinitis was defined as the combination of rhinitis symptoms with an atopic status.
	Non-allergic rhinitis was defined as the occurrence of rhinitis symptoms in the absence of an atopic status

D.1.48 Herr 2012

Bibliographic reference	Herr M, Just J, Nikasinovic L et.al (2012) Influence of host and environmental factors on wheezing severity in infants: findings from the PARIS birth cohort. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 42(2), 275-83
Study design	Prospective cohort study
Objective	To investigate host and environmental risk factors for the occurrence of wheeze during the first 18 months of life

Bibliographic reference	Herr M, Just J, Nikasinov environmental factors of the PARIS birth cohort. O British Society for Allerg	n wheezing : Clinical and	severity in inf experimental	fants: findings from allergy: journal of the
Setting/Study location	France			
Number of participants	1879 infants			
Selected population	No			
Participant	Description	No.		%
characteristics	Sex			
	Male	938		49.4
	Female	941		50.1
	Maternal age (years)	Not reporte	ed	Not reported
	Ethnicity	Not reporte	ed	Not reported
	(Maintenance) medication use	Not reporte	ed	Not reported
	Parental asthma and/or atopic	348		18.5
	Parental education	Not reporte	ed	Not reported
	Annual family income	Not reporte	ed	Not reported
	Building characteristics	Not reporte	ed	Not reported
Inclusion criteria	Not reported			
Exclusion criteria	Not reported			
Type of pollutant/exposure	Presence of furred pet House dust mite from cleaning habits, carpeted covered floor, age of the bedding and use of anti-dust mite cover on the mattress			
Pollutant/exposure assessment	House dust mite, pets, and mould was analysed using ImmunoCAP Phadiap and Trophatop fx26, fx27 and fx28 with a detection limit set at 0.35 U/mL. Sensitised infant were further investigated to identify the allergen(s) involved.			
Outcome	Wheeze – that required in	haled cortico	steroids and/o	r hospital based care
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between host and environmental risk factors for wheeze in the first 18 months of life			
			Wheeze	
			OR (95%CI)	
	Renovation activities after	birth	1.22 (0.96, 1	.54)
	Presence of a cat in the he	ome	0.65 (0.47, 0.89)	
	House dust mite from carp covered floor	peted	1.39 (1.12, 1	.73)
	Daily use of cleaning spra	у	1.50 (0.97, 2	.32)
Follow up	18 months			

Bibliographic reference	Herr M, Just J, Nikasinovic L et.al (2012) Influence of host and environmental factors on wheezing severity in infants: findings from the PARIS birth cohort. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 42(2), 275-83
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • objective sample Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for Prenatal exposure to tobacco smoke • study controls for any additional factor birth weight, socioeconomic status, duration of maternal exclusive breastfeeding and presence of mould and cockroaches in the home Outcome Assessment of outcome • record linkage Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for □ Overall risk of bias - Low
Source of funding	Government: Social, Childhood and Health Direction (DASES) of the Paris Council.
Comments	

D.1.49 Hjortebjerg 2012

Bibliographic reference	Hjortebjerg D, Andersen A N, Garne E et.al (2012) Non-occupational exposure to paint fumes during pregnancy and risk of congenital anomalies: a cohort study. Environmental health: a global access science source 11, 54
Study design	Prospective cohort study
Objective	to investigate the association between exposure to paint fumes in the residence during the 1st trimester of pregnancy and the risk of congenital anomalies in a prospective cohort
Setting/Study location	Denmark
Number of participants	20103 pregnant women
Selected population	No

Bibliographic reference	Hjortebjerg D, Andersen A N exposure to paint fumes dur anomalies: a cohort study. E science source 11, 54	ring preg	nancy and risk of ental health: a glo	congenital
Participant characteristics	Description Sex o Female Age (years) reported as mater age – Mean (SD) Ethnicity Education SES	nal	Exposed (n=1404) 1404 (100%) 29.2 (4.2%) Not reported Not reported Not reported	Non-exposed (n=18531) 18531 (100%) 29.3 (4.3%) Not reported Not reported Not reported
Inclusion criteria	Building characteristics Able to speak Danish Pregnant and intended to carr	v the prec	Not reported	Not reported
Exclusion criteria	Pregnant and intended to carry the pregnancy to term birth of stillborns women whose children had a diagnosis of chromosomal abnormalities incomplete information on covariates and not the main exposure of interest, paint fumes.			
Type of pollutant/exposure	Volatile organic compounds (VOC) – paint fumes in 1st trimester			
Pollutant/exposure assessment	Interview			
Outcome	Congenital anomalies (via National Hospital Discharge Registry)			
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between paint fumes in the residence during 1st trimester of pregnancy and congenital anomalies			
		All congenital malformations		
		aOR (9	5%CI)	
	All congenital malformations	0.95 (0.	74, 1.21)	
	Nervous system	2.19 (0.	76, 6.32)	
	Eye	,	70, 4.57)	
	Ear, face and neck	2.15 (0.84, 5.55)		
	Congenital heart defects	0.76 (0.39, 1.49)		
	Respiratory system	1.13 (0.	27-4.79)	
	Cloft lin and sloft nalata	1.06 (0.33, 3.46)		
	Cleft lip and cleft palate	1.00 (0.	55, 5. 4 6)	
	·		•	
	Digestive system Abdominal wall defects		15, 2.50)	
	Digestive system	0.61 (0. NA	•	
	Digestive system Abdominal wall defects	0.61 (0. NA 2.16 (1.	15, 2.50)	
	Digestive system Abdominal wall defects Renal	0.61 (0. NA 2.16 (1. 0.83 (0.	15, 2.50) 02-4.58)	
	Digestive system Abdominal wall defects Renal Genital	0.61 (0. NA 2.16 (1. 0.83 (0. 0.82 (0.	15, 2.50) 02-4.58) 48-1.43)	
	Digestive system Abdominal wall defects Renal Genital Limb defects	0.61 (0. NA 2.16 (1. 0.83 (0. 0.82 (0. 1.77 (0.	15, 2.50) 02-4.58) 48-1.43) 54-1.24)	

Bibliographic reference	Hjortebjerg D, Andersen A N, Garne E et.al (2012) Non-occupational exposure to paint fumes during pregnancy and risk of congenital anomalies: a cohort study. Environmental health: a global access science source 11, 54
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average pregnant woman in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • structured interview Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for working with organic solvents at first interview (12th week) • study controls for any additional factors - Maternal age, smoking during 1st trimester, alcohol consumption during 1st trimester, Outcome Assessment of outcome • record linkage Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • subjects lost to follow up unlikely to introduce bias - or description provided of those lost)
Source of funding	Charity: The Danish National Research Foundation; Pharmacy Foundation, the Egmont Foundation, the March of Dimes Birth Defects Foundation, the Augustinus Foundation, and the Health Foundation.
Comments	

D.1.50 Hoffmann 2007

Bibliographic reference	Hoffmann B, Moebus S, Mohlenkamp S, et al (2007) Residential exposure to traffic is associated with coronary atherosclerosis. Circulation 116(5), 489-96
Study design	Prospective cohort study
Objective	To investigate the association of long-term residential traffic exposure and $PM_{2.5}$ exposure with the degree of coronary atherosclerosis in a population-based cohort in Germany
Setting/Study location	Germany
Number of participants	4494 adults

Bibliographic reference	Hoffmann B, Moebus S, Mohlenkamp S, et al (2007) Residential exposure to traffic is associated with coronary atherosclerosis. Circulation 116(5), 489-96		
Selected population	No		
Participant	Description	No.	%
characteristics	Sex (male)	2206	49.1
	Age (years); mean (SD)	60.2 (7.8)	-
	Ethnicity	Not reported	Not reported
	(Maintenance) medication use	Not reported	Not reported
	Maternal asthma and/or atopic	Not reported	Not reported
	Parental education		
	Low	2491	55.4
	Medium	1249	27.8
	High	754	16.8
	Household income		
	<3000 €/month	1554	36.8
	3000 to 5999 €/month	1628	38.6
	≥ 6000 €/month	1041	24.7
	Building characteristics	Not reported	Not reported
Inclusion criteria	Not reported		
Exclusion criteria	Not reported		
Type of pollutant/expos ure	Proximity to traffic		
Pollutant/expo sure assessment	Daily mean values for PM _{2.5} (midpoint of the baseline examination) on a grid of 5 km were estimated with the EURAD dispersion model using input data from official emission inventories, meteorological information, and regional topographical data. The model was validated by comparing the daily model-derived values with measured air pollution data from monitoring sites, showing very good agreement		
Outcome	coronary artery calcification (CAC)		
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between proximity to traffic, PM _{2.5} and coronary artery calcification (CAC)		
	High Traffic Exposure (≤100 m)		
		aOR (95%CI)	
	Coronary artery calcification (CAC)	1.45 (1.15, 1.82)	
Follow up	Not reported		
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average population in the community		n the community

Bibliographic reference	Hoffmann B, Moebus S, Mohlenkamp S, et al (2007) Residential exposure to traffic is associated with coronary atherosclerosis. Circulation 116(5), 489-96
	 Selection of the non-exposed cohort no description of the derivation of the non-exposed cohort Ascertainment of exposure validated measurement used Demonstration that outcome of interest was not present at start of study Yes Comparability Comparability of cohorts on the basis of the design or analysis Study controls for city, area of residence, age, sex, education, smoking, ETS, physical inactivity, waist-to-hip ratio, diabetes, blood pressure, and lipids. Outcome Assessment of outcome medical examination/investigation of CAC Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts no statement Overall risk of bias: low
Source of funding	Charity: Heinz Nixdorf Stiftung
Comments	Authors reported that a lack of a residential history is a limitation to the study. Relocations, a change in traffic patterns, and a change in other anthropogenic emissions (industry, heating with fossil fuels) before the baseline examination might have led to exposure misclassifications

D.1.51 Hunt 2011

Bibliographic reference	Hunt A, Crawford JA, Rosenbaum P F, et al (2011) Levels of household particulate matter and environmental tobacco smoke exposure in the first year of life for a cohort at risk for asthma in urban Syracuse, NY. Environment International 37(7), 1196-205
Study design	Prospective cohort
Objective	To investigate possible associations between indoor exposures and infant health status (in particular wheezing)
Setting/Study location	United States
Number of participants	103 mother baby-dyads
Selected population	Yes (selected on based of being at risk of asthma)
Participant characteristics	Building characteristics: Location: urban Dwelling type: Not reported Building age: Not reported Type of ownership/tenancy: Not reported

Bibliographic reference	Hunt A, Crawford JA, Rosenbaum P F, et al (2011) Levels of household particulate matter and environmental tobacco smoke exposure in the first year of life for a cohort at risk for asthma in urban Syracuse, NY. Environment International 37(7), 1196-205	
	Parental characteristics:	
	Age: Not reported	
	Current smoker (mother): 55 (54%)	
	Hay fever: Not reported	
	Atopy: Not reported but all mothers had asthma	
Inclusion criteria	Documented history of maternal asthma	
	Expectation of the mother residing in same residence for at least 1 year or an adjacent urban location	
	Infant criteria	
	Gestational age ≥ 37 weeks	
	Birthweight ≥ 2500 g	
	Absence of any major congenital abnormality	
	Singleton birth	
Exclusion criteria	None reported	
Type of pollutant/exposure	Particulate matter 2.5 ≥15µg/m³	
Pollutant/exposure assessment	Particulate matter was collected using size selective Harvard impactors operating at 10 L/min placed in the living room	
Outcome	Wheeze defined as	
	Primary-care provided-documented wheezing, reactive airway disease, asthma or bronchiolitis or	
	Wheeze heard on physical examination by the nurse practitioner or	
	A prescription for bronchodilator, inhaled steroid or steroid pulse prescription document in the medical records	
Results	Wheeze	
	PM 2.5 ≥15µg/m³=aOR 4.21 (1.36, 13.03)	
Follow up	12 months	
Newcastle-Ottawa Scale	Selection	
	Representativeness of the exposed cohort	
	truly representative of the average infant in the community	
	Selection of the non-exposed cohort	
	drawn from the same community as the exposed cohort □	
	Ascertainment of exposure	
	• secure record of objective measurement	
	Demonstration that outcome of interest was not present at start of study	
	• Yes	
	Comparability	
	Comparability of cohorts on the basis of the design or analysis	
	 study controls for gender, maternal age and education, season of home visit and presence of carpeting 	
	Outcome	
	Assessment of outcome	
	independent blind assessment or medical records	

Bibliographic reference	Hunt A, Crawford JA, Rosenbaum P F, et al (2011) Levels of household particulate matter and environmental tobacco smoke exposure in the first year of life for a cohort at risk for asthma in urban Syracuse, NY. Environment International 37(7), 1196-205	
	Was follow-up long enough for outcomes to occur	
	• Yes	
	Adequacy of follow up of cohorts	
	complete follow up - all subjects accounted for	
	Overall assessment=Low	
Source of funding	Government: USEPA	
Comments	Particulate matter was primarily sourced from environmental tobacco smoke	

D.1.52 Ibargoyen-Roteta 2007

ibargoyeri-Rolela 2	.001		
Bibliographic reference	Ibargoyen-Roteta N, Aguinaga-Ontoso I, Fernandez-Benitez M et.al (2007) Role of the home environment in rhinoconjuctivitis and eczema in schoolchildren in Pamplona, Spain. Journal of investigational allergology & clinical immunology 17(3), 137-44		
Study design	Prospective cohort study		
Objective	To analyse the possible home-condition risk factors for allergic rhino conjunctivitis, atopic eczema, and severe disease in schoolchildren		
Setting/Study location	Spain		
Number of participants	3360 children		
Selected population	No		
Participant characteristics	Description Sex Age (years)- range Ethnicity Education Annual family income	Not reported 5 – 8 Not reported Not reported Not reported	
Inclusion criteria	Not reported		
Exclusion criteria	Not reported		
Type of pollutant/exposure	Dust, moulds, animal dander,		
Pollutant/exposure assessment	Questionnaire (self-report)		
Outcome	Allergic Rhino conjunctivitis		
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between allergic rhinoconjuctivitis damp and mould		

Bibliographic reference	Ibargoyen-Roteta N, Aguinaga-Ontoso I, Fernandez-Benitez M et.al (2007) Role of the home environment in rhinoconjuctivitis and eczema in schoolchildren in Pamplona, Spain. Journal of investigational allergology & clinical immunology 17(3), 137-44			
		Allergic Rhinoconjuctivitis		
	Moisture on walls	1.90 (1.01, 3.56)		
	Moulds on walls	1.34 (0.64, 2.79)		
	Single glass window	1.52 (1.03, 2.23)		
	Double-glazed window	1.83 (1.26, 2.66)		
Follow up	7 years			
Risk of bias	Selection			
(Newcastle-Ottawa	Representativeness of the ex	kposed cohort		
Scale)	truly representative of the average child			
	Selection of the non-exposed cohort			
		munity as the exposed cohort □		
	Ascertainment of exposure			
	written self-report			
	 Demonstration that outcome of interest was not present at start of study No Comparability Comparability of cohorts on the basis of the design or analysis 			
	study controls for age			
	study controls for additional factor as follows – sex and language			
	Outcome	ů ů		
	Assessment of outcome			
	• self-report			
	Was follow-up long enough f	or outcomes to occur		
	Yes Adequacy of follow up of cohorts			
	complete follow up - all sub			
	Overall level of bias – High (concerns over self-reporting of outcome and exposure)			
Source of funding	Government: Navarre Depa	rtment of Health		
Comments				

D.1.53 lossifova 2009

Bibliographic reference	lossifova YY, Reponen T, Ryan PH, et al (2009) Mold exposure during infancy as a predictor of potential asthma development. Annals of allergy, asthma & immunology: official publication of the American College of Allergy, Asthma, and & Immunology 102(2), 131-7
Study design	Prospective cohort study
Objective	To examine how exposure to mould in infancy predicts the risk of future asthma

Bibliographic reference	infancy as a predictor of po	tential asth	al (2009) Mold exposure during ima development. Annals of allergy, ition of the American College of 02(2), 131-7
Setting/Study location	United States		
Number of participants	483 children		
Selected population	No		
Participant	Individual characteristics		
characteristics	Age	Not reporte	ed
	Sex Male Female	206 277	
	Race / ethnicity Black 77 Other 406		
	SES	Not reporte	ed
Inclusion criteria	Infants born between October 2001 and July 2003 in the Greater Cincinnati / Northern Kentucky area		
Exclusion criteria	Not reported		
Type of pollutant / exposure	Mould		
Pollutant / exposure assessment	The extent of home mould and water damage was categorized as none, low (mouldy odour or moisture damage or visible mould area<0.2 m2), and high (moisture damage and visible mould area ≥ 0.2 m2).		
Outcome	Adjusted odds ratio and 95%	confidence	intervals
Results	Wheeze in children with atopy Visible mould (low vs none) 1.86 (0.86, 4.00) Visible mould (high vs none) 6.16 (1.38, 27.44)		
Follow up	2 years		
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort		
	drawn from the same community as the exposed cohort Ascertainment of exposure		
	 structured interview Demonstration that outcome of interest was not present at start of study Yes 		
	Comparability Comparability of cohorts on the basis of the design or analysis study controls for maternal smoking		

Bibliographic reference	lossifova YY, Reponen T, Ryan PH, et al (2009) Mold exposure during infancy as a predictor of potential asthma development. Annals of allergy, asthma & immunology: official publication of the American College of Allergy, Asthma, and & Immunology 102(2), 131-7
	 study controls for additional factors - race, number of siblings in the household, lower respiratory tract symptoms, and upper respiratory tract symptoms Outcome Assessment of outcome independent assessment Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts complete follow up - all subjects accounted for Overall risk of bias: Low
Source of funding	Government: National Institute of Environmental Health Sciences, Academic: National Institute for Occupational Safety and Health Training Program of the University of Cincinnati Education and Research Center
Comments	

D.1.54 Jaakkola 2005

Additional and the second and the se			
Bibliographic reference	Jaakkola JJ, Hwang BF, and Jaakkola N (2005) Home dampness and molds, parental atopy, and asthma in childhood: A six-year population-based cohort study. Environmental Health Perspectives 113(3), 357-61		
Study design	Prospective cohort study		
Objective	To assess the relation between indica development of asthma later in life.	tors of exposure to moulds and	
Setting/Study location	Finland		
Number of participants	1,916 children		
Selected population	No		
Participant characteristics	Age (at baseline) 1 2 3 4 5 6–7 Sex Male Female	324 (16.3) 301 (15.2) 318 (16.0) 333 (16.8) 314 (15.8) 394 (19.9) 983 (49.6%) 1,001 (50.5%)	

Bibliographic reference	Jaakkola JJ, Hwang BF, and Jaakkola N (2005) Home dampness and molds, parental atopy, and asthma in childhood: A six-year population-based cohort study. Environmental Health Perspectives 113(3), 357-61			
	Race/ Ethnicity SES (reported as parental education) No professional Trade school College or university	Not reported 369 (18.7) 523 (26.5) 1,085 (54.9)		
Inclusion criteria	Children living in the city of Espoo in F Born between January 1, 1984, and D			
Exclusion criteria	Children with an asthma diagnosis at l asthma were available	paseline or for whom no details on		
Type of pollutant/exposure	Dampness and mould			
Pollutant/exposure assessment	Authors used indicators of exposure (mould odour, visible mould, moisture and water damage) defined from the answers to structured questions at baseline and at follow up			
Health outcome	Asthma			
Results	Adjusted incident rate ratios (aIRRs) and 95% confidence intervals (CIs)			
	Mould odour Visible mould Moisture in the surfaces Water damage	2.44 (1.07, 5.60) 0.65 (0.24, 1.72) 0.92 (0.54, 1.54) 1.01 (0.45, 2.26)		
Follow up	6 years			
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed co truly representative of the average of Selection of the non-exposed cohort drawn from the same community as Ascertainment of exposure structured interview Demonstration that outcome of interest Yes Comparability Comparability Comparability of cohorts on the basis study controls for second-hand tobat study controls for additional factor as breastfeeding, parents' highest educe maternal smoking in pregnancy, gas feathery pets at home and type of day Outcome Assessment of outcome self-report Was follow-up long enough for outcome	the exposed cohort the ex		

Bibliographic reference	Jaakkola JJ, Hwang BF, and Jaakkola N (2005) Home dampness and molds, parental atopy, and asthma in childhood: A six-year population-based cohort study. Environmental Health Perspectives 113(3), 357-61
	 Yes Adequacy of follow up of cohorts subjects lost to follow up unlikely to introduce bias - description provided of those lost) Overall risk of bias=High (concerns of self-report of exposure and outcome)
Source of funding	Government: Ministry of the Environment, the National Agency for Welfare, and Health and the Medical Research Council of the Academy of Finland, Charity: Yrjö Jahnsson Foundation.
Comments	

D.1.55 Jaakkola 2010

Jaakkola JJ. K, Hwang B, and Jaakkola M S (2010) Home Dampness and moulds as Determinants of Allergic Rhinitis in Childhood: A 6-Year, Population-based Cohort Study. American Journal of Epidemiology 172(4), 451-459				
Prospective co	ohort study			
To assess the relationship between exposure to moulds and dampness in dwellings and the risk of developing allergic rhinitis in childhood up to 14 years of age				
Finland				
1,863 children				
No				
Description	Baseline		6 – yeaı	r cohort
	No.	%	No.	%
Sex				
Male	1,258	49.0	983	49.6
Female	1,310	51.0	1,001	50.5
Age (years)				
1	424	16.5	324	16.3
2	405	15.8	301	15.2
	and moulds a Year, Populat Epidemiology Prospective of To assess the dwellings and years of age Finland 1,863 children No Description Sex Male Female Age (years) 1	and moulds as Determinants Year, Population-based Coh Epidemiology 172(4), 451-45 Prospective cohort study To assess the relationship bet dwellings and the risk of devel years of age Finland 1,863 children No Description Baseline No. Sex Male 1,258 Female 1,310 Age (years) 1 424	and moulds as Determinants of Allergic Rhinitis Year, Population-based Cohort Study. American Epidemiology 172(4), 451-459 Prospective cohort study To assess the relationship between exposure to mo dwellings and the risk of developing allergic rhinitis i years of age Finland 1,863 children No Description Baseline No. Sex Male 1,258 49.0 Female 1,310 51.0 Age (years) 1 424 16.5	and moulds as Determinants of Allergic Rhinitis in Child Year, Population-based Cohort Study. American Journal Epidemiology 172(4), 451-459 Prospective cohort study To assess the relationship between exposure to moulds and dwellings and the risk of developing allergic rhinitis in childhouse years of age Finland 1,863 children No Description Baseline No. % No. Sex Male 1,258 49.0 983 Female 1,310 51.0 1,001 Age (years) 1 424 16.5 324

Bibliographic reference	Jaakkola JJ. K, Hwang B, and Jaakkola M S (2010) Home Dampness and moulds as Determinants of Allergic Rhinitis in Childhood: A 6-Year, Population-based Cohort Study. American Journal of Epidemiology 172(4), 451-459					
Study design	Prospective co	spective cohort study				
	3	410	16.0	318	16.0	
	4	400	15.6	333	16.8	
	5	415	16.2	314	15.8	
	6-7	514	20.0	394	19.9	
	Ethnicity	Not reported		Not reported		
	Maintenance medication use	Not reported		Not rep	Not reported	
	Parental asthma and/or atopic	Not reported		Not rep	orted	
	Parental educ	ation (years)				
	Nonprofessi onal	498	19.5	369	18.7	
	Trade school	663	25.9	523	26.5	
	College or university	1,395	54.6	1,085	54.9	
	Annual family income	Not reported Not reported			orted	
	Building chara	cteristics				
	Gas stove					
	Yes	86	3.4	62	3.1	
	No	2,469	96.6	1,913	96.9	
Inclusion criteria	Born between	Children living in the city of Espoo in Finland Born between January 1, 1984, and December 31, 1989 Children who did not have physician-diagnosed rhinitis				
Exclusion criteria	Not reported					
Type of pollutant/exposure	Dampness and mould					
Pollutant/exposure assessment	Authors used indicators of exposure (mould odour, visible mould, moisture and water damage) defined from the answers to structured questions at baseline and at follow up					
Health outcome	New cases of	allergic rhinitis				
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for dampness and mould and incidence of allergic rhinitis					
	Exposure	Allergic rhinitis				
	Water damage Moisture on th Visible mould	· · · · · · · · · · · · · · · · · · ·				

Bibliographic reference	Jaakkola JJ. K, Hwang B, and Jaakkola M S (2010) Home Dampness and moulds as Determinants of Allergic Rhinitis in Childhood: A 6-Year, Population-based Cohort Study. American Journal of Epidemiology 172(4), 451-459
Study design	Prospective cohort study
Follow up	6 years
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • structured interview Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for second-hand tobacco smoke • study controls for additional factor as follows - Age, gender, duration of breastfeeding, parents' highest education, single parent or guardian, maternal smoking in pregnancy, gas cooking, presence of hairy or feathery pets at home and type of day care Outcome Assessment of outcome • self-report Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • subjects lost to follow up unlikely to introduce bias - description provided of those lost) Overall risk of bias=High (concerns of self-report of exposure and outcome)
Source of funding	Government: Supported by the Ministry of the Environment, the National Agency for Welfare and Health, the Medical Research Council of the Academy of Finland and the Yrjo Jahnsson Foundation and the Medical Research Council of the Academy of Finland
Comments	Evidence suggests increase risk of allergic rhinitis to damp and mould, with a higher risk of allergic rhinitis related to a longer duration of exposure.

D.1.56 Jedrychowski 2005

Bibliographic reference	Jedrychowski W, Galas A, Pac A, et al (2005) Prenatal ambient air exposure to polycyclic aromatic hydrocarbons and the occurrence of respiratory symptoms over the first year of life. European journal of epidemiology 20(9), 775-82
Study design	Prospective cohort study
Objective	To test the hypothesis that infants with higher levels of prenatal exposure to PAHs may be at greater risk of developing respiratory symptoms.

Bibliographic reference	Jedrychowski W, Galas A, Pac A, et exposure to polycyclic aromatic hyd respiratory symptoms over the first epidemiology 20(9), 775-82	Irocarbons a	and the occurrence of	
Setting/Study location	Poland			
Number of participants	333 infants			
Selected population	No			
Participant characteristics	Description Sex Male Age Maternal allergy Ethnicity Education Annual family income	N (%) 168 (50.5% Not reporte 81 (24.3%) Not reporte Not reporte Not reporte	d d d	
Inclusion criteria	Non-smoking women Ages 18–35 years Singleton pregnancies Free from chronic diseases such as diabetes and hypertension			
Exclusion criteria	Not reported			
Type of pollutant/exposure	Polycyclic aromatic hydrocarbons (PAH)			
Pollutant/exposure assessment	Monitoring of personal PAH inhalation was carried out in all pregnant women for over a 48-hour period during the second trimester of pregnancy.			
Outcome	Runny or stuffy nose, Ear infections (otitis media) Sore throat, Cough with or without cold, barking cough, Difficult (puffed) breathing, Wheezing or whistling in the chest irrespective of respiratory infection, Wheezing without cold.			
Results	Adjusted risk ratios (aHRs) and 95% coof PAH concentration in ng/m³)	onfidence into	ervals (Cls) (per log unit	
			aRR (95%CI)	
	Runny or stuffy nose 1.11 (0.97–1.27) Ear infections (otitis media) 1.82 (1.03–3.23) Sore throat, 1.27 (1.07–1.52) Cough 1.72 (1.02–2.92) Cough without cold, 4.80 (2.73–8.44) Barking cough, 1.12 (0.82–1.55) Difficult (puffed) breathing, 1.23 (0.83–1.84) Wheezing or whistling in the chest irrespective of respiratory infection, 1.96 (1.38–2.78)		1.82 (1.03–3.23) 1.27 (1.07–1.52) 1.72 (1.02–2.92) 4.80 (2.73–8.44) 1.12 (0.82–1.55) 1.23 (0.83–1.84) 3.83 (1.18–12.43)	
Follow up	2 years			
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average pregnant woman			

Bibliographic reference	Jedrychowski W, Galas A, Pac A, et al (2005) Prenatal ambient air exposure to polycyclic aromatic hydrocarbons and the occurrence of respiratory symptoms over the first year of life. European journal of epidemiology 20(9), 775-82
	 Selection of the non-exposed cohort drawn from the same community as the exposed cohort □ Ascertainment of exposure written self-report Demonstration that outcome of interest was not present at start of study Yes Comparability Comparability of cohorts on the basis of the design or analysis study controls for gender of child, child's birth weight, season of birth, ETS in postnatal period, mother's allergy, mother's education level, moulds at home. Outcome Assessment of outcome self-report Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts complete follow up - all subjects accounted for Overall assessment=Moderate (concerns over self-report of outcomes)
Source of funding	Government: National Institute of Environmental Health Sciences Charity: The Gladys and Roland Harriman Foundation N. York
Comments	

D.1.57 Jedrychowski 2010

Bibliographic reference	Jedrychowski WA, Perera FP, Maugeri U, et al (2010) Intrauterine exposure to polycyclic aromatic hydrocarbons, fine particulate matter and early wheeze. Prospective birth cohort study in 4-year olds. Pediatric allergy and immunology: official publication of the European Society of Pediatric Allergy and Immunology 21(4 Pt 2), e723-32
Study design	Prospective cohort
Objective	To examine relationship between prenatal exposure to PAH compounds to the onset and frequency of wheezing in early childhood.
Setting/Study location	Poland
Number of participants	369
Selected	No

Bibliographic reference	Jedrychowski WA, Perera FP, Maugeri U, et al (2010) Intrauterine exposure to polycyclic aromatic hydrocarbons, fine particulate matter and early wheeze. Prospective birth cohort study in 4-year olds. Pediatric allergy and immunology: official publication of the European Society of Pediatric Allergy and Immunology 21(4 Pt 2), e723-32		
Participant characteristics	Building characteristics: Location: Dwelling type: single family home Multi-family home Building age: Type of ownership/tenancy: Individual characteristics: Age (range) Gender: Male (%) Girls (%) Smoker in home: Race Not reported Not reported 100 (85.6% Not reported		
Inclusion criteria	women 18-35 years of age claimed to be non-smokers, singleton pregnancies, without illicit drug use and HIV infection, free from chronic diseases such as diabetes or hypertension, and resided in Krakow for at least one year prior to pregnancy		
Exclusion criteria	None reported		
Type of pollutant/exposure	Polycyclic aromatic hydrocarbons (PAH's) (>0.250 adducts per 108 nucleotides) Particulate matter 2.5 Damp / Mould		
Pollutant/exposure assessment	Prenatal exposure to PAHs were measured by PAH-DNA adducts in umbilical cord blood. The level of PAH-DNA adducts in the cord blood is assumed to reflect the cumulative dose of PAHs absorbed by the fetus over the prenatal period. Monitoring of personal of fine particles (PM _{2.5}) was carried out in all pregnant women over a 48-hour period during the second trimester of pregnancy. The women were instructed by the trained staff member as how to use personal monitor and asked to carry the monitoring device during the daytime hours for two consecutive days and place it by their bed at night. On the second day the air monitoring staff assistant and interviewer visited the woman's home to change the battery-pack and to complete the questionnaire on the household characteristics.		
Outcome	Wheeze		
Results	Damp / mould Cord blood PAH-adducts	IRR (95%CI) for wheeze in years 1 & 2 1.429 (1.265, 1.614) 1.686 (1.517, 1.875) 1.377 (1.252, 1.514)	

Bibliographic reference	Jedrychowski WA, Perera FP, Maugeri U, et al (2010) Intrauterine exposure to polycyclic aromatic hydrocarbons, fine particulate matter and early wheeze. Prospective birth cohort study in 4-year olds. Pediatric allergy and immunology: official publication of the European Society of Pediatric Allergy and Immunology 21(4 Pt 2), e723-32	
	Particulate matter2.5 (median prenatal 35.4 µg/m³) Damp / mould Cord blood PAH-adducts Particulate matter2.5 (median prenatal 35.4 µg/m³)	IRR (95%CI) for wheeze in years 3 & 4 1.669 (1.390, 2.005) 0.956 (0.836, 1.093) 1.063 (0.923, 1.223)
Follow up	5 years	
Newcastle-Ottawa Scale	, ,	
Source of funding	Government: National Institute of Environmental Health Sciences (NIEHS); Charity; the Lundin Foundation; the Gladys T. and Roland Harriman Foundation.	
Comments	Part of a larger cohort Data from years 3-4 used in GRADE table	

J

D.1.58 Jedrychowski 2011

Jedrychowski 201			
Bibliographic reference	Jedrychowski W, Spengler JD, Maugeri U, et al (2011) Joint effect of prenatal exposure to fine particulate matter and intake of Paracetamol (Acetaminophen) in pregnancy on eczema occurrence in early childhood. The Science of the total environment 409(24), 5205-9		
Study design	Prospective cohort study		
Objective	To assess the role of very low prenatal exposure to Paracetamol in the occurrence of eczema symptoms in early childhood and assess the possible interaction with prenatal exposure to particulate pollutants		
Setting/Study location	Poland		
Number of participants	322 infants		
Selected population	No		
Participant characteristics	Description Sex Male 159 (49.4%) Female 163 (50.6%) Age (years)- Mean (SD) Maternal 27.82 (3.39) Ethnicity Not reported Education Elementary Elementary 28 (8.7%) Medium 77 (23.9%) Higher 217 (67.4%) Annual family income Not reported		
Inclusion criteria	Non-smoking women Ages 18–35 years Singleton pregnancies Free from chronic diseases such as diabetes and hypertension		
Exclusion criteria	Not reported		
Type of pollutant/exposure	PM _{2.5} , Damp/mould		
Pollutant/exposure assessment	Questionnaire (Damp / Mould) The woman was asked to wear the backpack monitor during the daytime hours for 2 consecutive days and to place the monitor near the bed at night. During the morning of the second day, the air monitoring staff-person and interviewer visited the woman's home to change the battery pack and administer the full questionnaire. A Personal Environmental Monitoring Sampler (PEMS) was used to measure particle mass. The PEMS is designed to achieve the particle target size of ≤ 2.5 μm at a flow rate of 4.0 liters per minute (LPM) with an array of 10 impactor nozzles. Flow rates are calibrated (with filters in place) using a bubble meter prior to the monitoring and are checked again with a change of the battery pack on the second day and at the conclusion of the monitoring. Pumps operated continuously at 2 LPM over the 48-hour period.		
Outcome	Eczema	·	

Bibliographic reference	Jedrychowski W, Spengler JD, Maugeri U, et al (2011) Joint effect of prenatal exposure to fine particulate matter and intake of Paracetamol (Acetaminophen) in pregnancy on eczema occurrence in early childhood. The Science of the total environment 409(24), 5205-9		
Results	Adjusted hazard ratios (aHRs) and 95% confidence intervals (CIs) for association between prenatal $PM_{2.5}$, damp/mould and occurrence of eczema in early childhood		
		Occurrence of eczema	
		aHR (95%CI)	
	Prenatal PM _{2.5}	1.06 (0.72, 1.57)	
	Damp/mould house	1.22 (1.07, 1.40)	
Follow up			
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average pregnant woman		
	Selection of the non-exposed cohort		
	• drawn from the same community as the exposed cohort		
	Ascertainment of exposure		
	 written self-report Demonstration that outcome of interest was not present at start of study 		
	• Yes	ion that outsome of interest was not present at start of stady	
	Comparability		
	Comparability of cohorts on the basis of the design or analysis		
	 study controls for prenatal and postnatal exposure to environmental tobacco smoke study controls for additional factor as follows – maternal education, maternal atopy, gender of child, presence of older siblings, breastfeeding practice, and the presence of moulds in the household). 		
	Outcome		
	Assessment of outcome		
	• self-report		
	Was follow-up long enough for outcomes to occur		
	• Yes		
	Adequacy of follow up of cohorts		
	 complete follow up - all subjects accounted for Overall assessment=Moderate (concerns over self-report of outcomes) 		
Source of funding		te of Environmental Health Sciences	
·	Charity: The Gladys and Roland Harriman Foundation		
Comments			

D.1.59 Jung 2012

Julig 2012	1 1/11 W B 1			4 1 1
Bibliographic reference	Jung KH, Yan B, Moors K, et al (2012) Repeated exposure to polycyclic aromatic hydrocarbons and asthma: effect of seroatopy. Annals of allergy, asthma & immunology: official publication of the American College of Allergy, Asthma, and & Immunology 109(4), 249-54			
Study design	Prospective cohort study			
Objective	To examine whether the associations between repeated early pyrene and ∑8PAH non-volatile exposure and asthma would differ between nonatopic and atopic children			
Setting/Study location	United States			
Number of participants	349 children			
Selected population	No			
Participant characteristics	Individual characteristics: Age- Gender: Not repo		Not reported 201 (53%)	
	Race		Not reported	
	Education			
	Maternal high school or greater degree Smoking		238 (63%)	
	Prenatal ETS		115 (31%)	
	Postnatal ETS		161 (43%)	
Inclusion criteria	Not reported			
Exclusion criteria	Not reported			
Type of pollutant/exposure	Polycyclic aromatic hydrocarbons (PAHs): pyrene and Σ8PAH non-volatile			
Pollutant/exposure assessment	Prenatal PAH (pyrene and Σ8 PAH non-volatile) exposure was measured from 48-hour personal air monitoring between 1998 and 2006, and PAH exposure at 5 to 6 years of age was measured from 2-week residential indoor monitoring between 2005 and 2011			
Outcome	Asthma Wheeze Visits to emergency department			
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between PAH, asthma, wheeze and emergency department visits			
		Asthma	Wheeze	Emergency department visits
		aOR (95%CI)	aOR (95%CI)	aOR (95%CI)
	Pyrene	1.90 (1.13, 3.20)	1.53 (0.93, 2.51)	1.21 (0.71, 2.09)
	Σ8PAH non- volatile	0.90 (0.52, 1.56)	0.86 (0.52, 1.42)	0.82 (0.46, 1.45)
Follow up	6 years			

Bibliographic reference	Jung KH, Yan B, Moors K, et al (2012) Repeated exposure to polycyclic aromatic hydrocarbons and asthma: effect of seroatopy. Annals of allergy, asthma & immunology: official publication of the American College of Allergy, Asthma, and & Immunology 109(4), 249-54
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • Objective sampling Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for exposure to environmental tobacco smoke • study controls for any additional factors as follows – maternal ethnicity, sex, maternal asthma, maternal education and cold/influenza season Outcome Assessment of outcome • independent blind assessment self-report Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall risk of bias – Moderate (concerns over self-report of some outcomes)
Source of funding	Government: National Institutes of Health; Environmental Protection Agency; Charity: The Educational Foundation of America; the John & Wendy Neu Family Foundation; the New York Community Trust; and the Trustees of the Blanchette Hooker Rockefeller Fund.
Comments	

D.1.60 Jung 2012 b

Bibliographic reference	Jung KH, Hsu SI, Yan B, et al (2012) Childhood exposure to fine particulate matter and black carbon and the development of new wheeze between ages 5 and 7 in an urban prospective cohort. Environment international 45, 44-50
Study design	Prospective cohort study
Objective	To examine associations between pollutant levels and subsequent new onset of respiratory symptoms and indoor allergen specific immunoglobulin (Ig) E at age 7 years after controlling for known covariates.

Bibliographic reference	Jung KH, Hsu SI, Yan B, et al (2012) Childr particulate matter and black carbon and th wheeze between ages 5 and 7 in an urban Environment international 45, 44-50	e development of new	
Setting/Study location	United States		
Number of participants	408 children		
Selected population	No		
Participant characteristics	Description Gender Girls 192 (55%) Age Not reported Ethnicity (maternal) 221 (63%) Dominican 128 (37%) African American Education (maternal) high school or greater degree 221 (64%) SES Not reported Building characteristics Not reported		
Inclusion criteria	Not reported		
Exclusion criteria	Not reported		
Type of pollutant/exposure	PM _{2.5}		
Pollutant/exposure assessment	Indoor air monitors were placed in a room where the child spent most of his or her time (mostly the room where the child sleeps).		
Outcome	New wheeze (no wheeze up to 5 years and wheeze in past 12 months at 6 or 7 years of age		
Results	Adjusted odds ratios (aORs) and 95% confide association between $PM_{2.5}$ and wheeze incidents		
		Wheeze	
		aOR (95%CI)	
	PM _{2.5} Per IQR increase (8.75 μg/m ³ ,)	1.51 (1.05, 2.16)	
Follow up	12 months		
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure		

Bibliographic reference	Jung KH, Hsu SI, Yan B, et al (2012) Childhood exposure to fine particulate matter and black carbon and the development of new wheeze between ages 5 and 7 in an urban prospective cohort. Environment international 45, 44-50
	 objective sampling Demonstration that outcome of interest was not present at start of study Yes Comparability Comparability of cohorts on the basis of the design or analysis study controls for prenatal and postnatal environmental tobacco smoke exposure study controls for additional factors - ethnicity, sex, maternal education, maternal asthma, cold/flu season, residential monitoring conducted prior to age 6,) Outcome Assessment of outcome self-report Was follow-up long enough for outcomes to occur No Adequacy of follow up of cohorts complete follow up - all subjects accounted for Overall risk of bias: Moderate (concerns over self-report of outcome)
Source of funding	Government: NIH Charity: The Educational Foundation of America, the John & Wendy Neu Family Foundation, the New York Community Trust, and the Trustees of the Blanchette Hooker Rockefeller Fund.
Comments	

D.1.61 Jung 2014

Bibliographic reference	Jung KH, Perzanowski M, Rundle A, et al (2014) Polycyclic aromatic hydrocarbon exposure, obesity and childhood asthma in an urban cohort. Environmental research 128, 35-41	
Study design	Prospective cohort study	
Objective	To examine whether obesity may modify the effects of age 5–6 year PAH exposure, and semi volatile and alkylated PAHs in particular, on asthma in 5–7 year old inner-city children	
Setting/Study location	United States	
Number of participants	363 children	
Selected populations	No	
Participant characteristics	Individual characteristics: Age-years Gender: Female Race	6 – 7 164 (53%) Not reported

Bibliographic reference		obesity and	et al (2014) Polycyclic aromatic childhood asthma in an urban , 35-41
	Education Maternal high school or g degree Smoking Prenatal ETS		193 (62%) 100 (32%)
	Postnatal ETS		142 (46%)
Inclusion criteria	Not reported		
Exclusion criteria	Not reported		
Type of pollutant/exposure	Polycyclic aromatic hydronon-volatile	carbons (PAI	Hs): Σ8PAH semi-volatile and Σ8PAH
Pollutant/exposure assessment	Indoor air monitors were por her time for two weeks		om where the child spent most of his ugh 6.
Outcome	Asthma		
Results			6 confidence intervals (CIs) for on and asthma at age 5 years
		Asthma	
		aRR (95%CI)	
	Pyrene	0.81 (0.59–1.12)	
	Σ8PAH semi-volatile	0.82 (0.60–1.12)	
	Σ8PAH non-volatile	0.74 (0.46–	1.18)
Follow up	2 years		
Risk of bias (Newcastle-Ottawa Scale)	,		the exposed cohort t was not present at start of study of the design or analysis nood exposure to environment tobacco hnicity, sex, maternal education, sidential monitoring conducted at age opic, prenatal pyrene, PM _{2.5} and

Bibliographic reference	Jung KH, Perzanowski M, Rundle A, et al (2014) Polycyclic aromatic hydrocarbon exposure, obesity and childhood asthma in an urban cohort. Environmental research 128, 35-41
	 complete follow up - all subjects accounted for Overall risk of bias – Moderate (Concerns over self-report of outcomes)
Source of funding	Government: National Institutes of Health, Environmental Protection Agency Charity: The Educational Foundation of America, the John & Wendy Neu Family Foundation, the New York Community Trust, and the Trustees of the
0	Blanchette Hooker Rockefeller Fund
Comments	

D.1.62 Karvonen 2015

Bibliographic reference	Karvonen A M, Hyvarinen A, Korppi M et.al (2015). Moisture damage and asthma: a birth cohort study. Paediatrics, 135(3), pp. e598-606.		
Study design	Prospective cohort study		
Objective	To prospectively evaluate whether inspector-observed moisture damage with or without visible mould in the home in infancy is associated with the development of new physician diagnosed asthma and with respiratory tract symptoms and atopic sensitization up to the age of 6 years		
Setting/Study location	Finland		
Number of participants	398 children		
Selected population	No		
	Description	No.	%

Bibliographic reference	Karvonen A M, Hyvarin and asthma: a birth col				
Participants	Sex			. ,,	
characteristics	Male	Not re	ported	Not reporte	ed
	Female	Not re	ported	Not reporte	ed
	Age	Not re	ported	Not reporte	ed
	Ethnicity	Not re	ported	Not reporte	ed
	(Maintenance) medication use	Not re	ported	Not reporte	ed
	Parental asthma and/or atopic	Not re	ported	Not reporte	ed
	Parental education	Not re	ported	Not reporte	ed
	Annual family income	Not re	ported	Not reporte	ed
	Building characteristics	Not re	ported	Not reporte	ed
	A family could participate in the study: Only with 1 child Mothers living on a farm with livestock, in rural areas or suburban areas Age ≥ 18 years Singleton pregnancy Delivery in a hospital No plans to move from the study area and spoke Finnish language			an areas	
Exclusion criteria	Premature delivery (< 37 weeks of gestation) Home delivery Congenital abnormalities in the infants and failure to obtain cord blood samples				
Type of pollutant/exposure	Moisture damage with m	ould			
Pollutant/exposure assessment	Sign of excess moisture graded by using a 6-point "need to repair" estimation scale and area damaged was measured. If there were several moisture-damaged locations in a given room or area, the areas of damage with the same need for repair estimation were totalled. Presence of mould odour or visible mould was recorded for each damage observation. Mould growth only on silicone sealants in the kitchen or in the bathroom was classified as no mould. The cut-off level to define atopic sensitization to inhalant allergens was 0.70 kU/L at the age of 6 years.				
Health outcome	Asthma: Incidence of asthma ever Wheezing, nocturnal cough and sensitisation to inhalant allergens				
Results	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	g aa	wheezing	Nocturnal cough	Asthma
	Minor moisture damage or without mould spots in child's main living area		1.16 (0.69, 1.94)	1.07 (0.71,1.59)	1.31 (0.72, 2.36)
	Major moisture damage moisture damage with vis		1.69 (0.88, 3.24)	1.27 (0.77, 2.09)	1.33 (0.60, 2.98)

Bibliographic reference	Karvonen A M, Hyvarinen A, Korppi M et.al (2015). Moisture damage and asthma: a birth cohort study. Paediatrics, 135(3), pp. e598-606.
	mould in child's main living area
Follow up	6 years
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • home inspection by trained engineers Demonstration that outcome of interest was not present at start of study • Yes Comparability
	Comparability of cohorts on the basis of the design or analysis • study controls for smoking during pregnancy • study controls for additional factors as follows - study cohort, farming
Source of funding	status, gender, maternal history of allergic diseases and number of siblings Outcome Assessment of outcome • independent blind assessment Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • subjects lost to follow up unlikely to introduce bias - description provided of those lost) Overall assessment=Low
Source of funding	Government: Study supported by research grants from the European Union (grant QLK4-CT-2001-00250); the Graduate School in Environmental Health (SYTYKE); EVO and VTR funding; the Farmers' Social Insurance Institution (Mela); the Academy of Finland (grant 139021); the Juho Vainio Foundation; the Finnish Cultural Foundation; and the Finnish National Institute for Health and Welfare.
Comments	A combined variable ("moisture damage or mould in the child's main living areas") was created by using information regarding signs of moisture damage and visible mould in the child's bedroom, the living room, or kitchen. Moisture damage and mould in early infancy in the child's bedroom, living room, or kitchen were associated with asthma development. Atopic children may be more susceptible than non-atopic children to the harmful effects of moisture damage and mould growth. Study suggests association between moisture damage in the living rooms, child's bedrooms, and kitchens with the risk of physician-diagnosed asthma ever, persistent asthma, and respiratory symptoms Associations with asthma ever were strongest for moisture damage with visible mould in the child's bedroom and in the living room.

D.1.63 Kingsley 2015

Bibliographic reference	Kingsley S L, Eliot M N, Whitsel E A et.al (2015) Residential proximity to major roadways and incident hypertension in post-menopausal women. Environ Res. 2015 October; 142: 522–528.			
Study design	Prospective cohort study			
Objective	To assess the association between residential distance to nearest major roadway and the risk of incident hypertension in the Women's Health Initiative (WHI) Clinical Trial (CT) cohorts			
Setting/Study location	United States			
Number of participants	38,360 women between 50 a	and 79 yea	rs of age	
Participant	Description	No.		%
characteristics	Sex	All female	Э	All female
	Age (years); mean (SD)	61.6 ± 6.9	9	-
	Ethnicity			
	White, Non-Hispanic	32529		84.8
	Black, Non-Hispanic	2647		6.9
	Hispanic/Latino	1765		4.6
	Asian or Pacific Islander	805		2.1
	Other	422		1.1
	Cases/selected population	Not repor	ted	Not reported
	Socio-economic status (maternal education)			
	<college degree<="" td=""><td>21060</td><td></td><td>54.9</td></college>	21060		54.9
	College graduate	14116		36.8
	Building characteristics	Not repor	ted	Not reported
Inclusion criteria	Not reported			
Exclusion criteria	Participants with hypertension at baseline defined as a systolic blood pressure (SBP) ≥140 mmHg, a diastolic blood pressure (DBP) ≥90 mmHg			
	Self-reported use of antihypertensive medication at baseline, or use of an antihypertensive medication as determined at baseline via medical inventory			
Type of pollutant/exposure	Residential proximity to major	or roadway	S	
Pollutant/exposure assessment	Major roadways were defined as those with US census feature class codes A1 (primary highway with limited access), A2 (primary road without limited access), or A3 (secondary and connecting roads). A1 and A2 roadways include interstate highways and US highways, which typically contain a mix of car and truck traffic moving at higher speeds, and A3 roadways include state highways and other major arteries, which typically have lower traffic counts moving at slower average speeds.			
Outcome	Incident hypertension			
Results	Adjusted hazard ratios (aHRs) and 95% confidence intervals (CIs) for association between categories of residential distance to nearest major roadway and incident hypertension			
			Incident hyper	rtension
	Residential Distance to Near Major Roadway (metres)	rest	HR (95%CI)	

Bibliographic reference	Kingsley S L, Eliot M N, Whitsel E A major roadways and incident hypertenerical Environ Res. 2015 October; 142: 522	ension in post-menopausal women.
	≤50	1.09 (0.95, 1.24)
	>50-200	1.02 (0.94, 1.10)
	>200-400	1.04 (0.97, 1.11)
	>400-1000	1.03 (0.98, 1.08)
	>1000	1.00 (Ref.)
Follow up	Median of 7.9 years	· ,
Study methods	Incident hypertension was defined as a mmHg, or a first self-report of medication pressure was measured at clinical cent standardized procedures after participal Authors used stratified Cox proportional hypertension associated with living ≤50 from nearest major roadway compared	on prescribed for hypertension. Blood res by trained personnel using ints had been seated for 5 minutes. Il hazards models to estimate incident 0, >50-200, >200-400, >400-1000 m
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed col truly representative of the average por Selection of the non-exposed cohort no description of the derivation of the Ascertainment of exposure validated measurement used Demonstration that outcome of interest Yes Comparability Comparability of cohorts on the basis of study controls for age, race, smoking education, household income, employ participation in the hormone replacent Outcome Assessment of outcome measured at clinical centres by trained procedures Was follow-up long enough for outcome Yes Adequacy of follow up of cohorts subjects lost to follow up unlikely to int Overall risk of bias: low	non-exposed cohort was not present at start of study f the design or analysis status, alcohol consumption, yment status, high cholesterol, nent therapy trial ed personnel using standardized es to occur
Source of funding	Government: National Institute of Envi NIH, National Heart, Lung, and Blood In Department of Health and Human Serv Academic: Brown University.	nstitute (NHLBI), NIH, U.S.
Comments	Tioudonno. Brown Oniversity.	

D.1.64 Koloski 2015

Bibliographic reference	Koloski N, Jones M, Weltman M, et al (2015) Identification of early environmental risk factors for irritable bowel syndrome and dyspepsia. Neurogastroenterology and motility 27(9), 1317-1325		
Study design	Prospective study		
Objective	To assess the role of a range of early environmental factors in IBS and functional dyspepsia		
Setting/Study location	Australia		
Number of participants	767 adults		
Selected population	No		
Participant	Individual characteristics		
characteristics	Age - Mean (SD)	59.9 (11.5)
	Sex Female	48.2%	
	Race / ethnicity	Not reporte	ed
	SES	Not report	ed
Inclusion criteria	Random sample of population who had taken part in 2 sirveys 1997 and 2011		
Exclusion criteria	Not reported		
Type of pollutant / exposure	Pets		
Pollutant / exposure assessment	Self-report		
Outcome	Adjusted odds ratio and 95%	confidence	intervals
Results	Irritable bowel syndrome		
	Pet exposure		1.47 (0.83, 2.61)
	Herbivore pet		2.09 (1.19, 3.67)
	Carnivore pet		1.58 (0.90,2.76)
	Omnivore pet		0.97 (0.26, 3.59)
	Functional dyspepsia		
	Pet exposure		1.69 (0.86, 3.36)
	Herbivore pet		2.34 (1.24, 4.45)
	Carnivore pet 2.04 (1.03, 4.03)		
	Omnivore pet		0.98 (0.21, 4.50)
Follow up			
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average adult) in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort		
		•	

Bibliographic reference	Koloski N, Jones M, Weltman M, et al (2015) Identification of early environmental risk factors for irritable bowel syndrome and dyspepsia. Neurogastroenterology and motility 27(9), 1317-1325
	Ascertainment of exposure
	Questionnaire
	Demonstration that outcome of interest was not present at start of study
	• Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	study controls for age
	• study controls for additional factor – gender and frequency of walking
	Outcome
	Assessment of outcome
	independent assessment
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	complete follow up - all subjects accounted for
	Overall risk of bias: Low
Source of funding	Non funding declared
Comments	

D.1.65 Korppi 2008

Bibliographic reference	Korppi M, Hyvarinen M, Kotaniemi-Syrjanen A et al (2008) Early exposure and sensitization to cat and dog: different effects on asthma risk after wheezing in infancy. Pediatric allergy and immunology: official publication of the European Society of Pediatric Allergy and Immunology 19(8), 696-701			
Study design	Cohort study			
Objective	To evaluate associations between exposure and sensitization to cats or dogs in infancy and later asthma and allergy in children hospitalized for wheezing at<24 months of age			
Setting/Study location	Finland	Finland		
Number of participants	100 children			
Selected population	Yes - children requiring hospitalisation	on for wheeze		
Participant characteristics	Description Sex Age (months) – range Ethnicity Education SES	Not reported 1 - 23 Not reported Not reported Not reported		

Bibliographic reference	risk after wheezing in infancy. Pe	and dog: different effects on asthma
Inclusion criteria	Presence of wheezing and respirator during an acute respiratory tract infe	
Exclusion criteria	Not reported	
Type of pollutant/exposure	Exposure to furry pets	
Pollutant/exposure assessment	Parent report	
Outcome	Physician-diagnosed persistent child	dhood asthma
Results		aOR (95%CI)
	Exposure to cats	0.26 (0.03, 2.42)
	Exposure to dogs	0.20 (0.02, 1.78)
Follow up	12.3 years (median)	
Scale	Representativeness of the exposed • selected group of children Selection of the non-exposed cohort • drawn from the same community at Ascertainment of exposure • self-report (parent) • no description Demonstration that outcome of inter • Yes Comparability Comparability Comparability of cohorts on the basi • study controls for parental history • study controls for additional factor RSV aetiology of bronchiolitis.) Outcome Assessment of outcome • independent blind assessment Was follow-up long enough for outcome • Yes Adequacy of follow up of cohorts • complete follow up - all subjects a	t as the exposed cohort est was not present at start of study is of the design or analysis of asthma s – atopic dermatitis in infancy and omes to occur
Source of funding	Not reported	oncerns over self-report of exposure)
Comments		
Commonto		

D.1.66 Larsson 2009

Larsson 2009			
Bibliographic reference	Larsson M, Weiss B, Janson S, et al (2009) Associations between indoor environmental factors and parental-reported autistic spectrum disorders in children 6-8 years of age. NeuroToxicology 30(5), 822-831		
Study design	Cohort		
Objective	To determine the associations between ASD in children aged 6-8 years and a number of environmental factors, including exposure conditions when they were 1-3 years of age and during pregnancy and the first year of life.		
Setting/Study location	Sweden		
Number of participants	4779 children		
Selected population	No		
Participant characteristics	Building characteristics: Dwelling type: Single family home Multi-family home Building age: Type of ownership/tenancy: : Age of child 6 years 7 years 8 years Current smoker Any Mother Father	N (%) 4090 (85.6) 546 (11.4) Not reported Not reported 135 (2.8) 3240 (67.8) 1332 (27.9) 909 (19.0) 605 (12.7) 398 (8.3)	
Inclusion criteria	Provided data at two survey timepoint	ts	
Exclusion criteria	Not reported		
Type of pollutant/exposure	Phthalates in house dust (PVC flooring	g used as proxy)	
Pollutant/exposure assessment	Self-report of use of PVC flooring Condensation on window		
Outcome	Autism spectrum disorder, (ASD)		
Results	PVC flooring in child's room (1.5 cm) in parents 'room (1.5 cm) in parent's room $(1.5 \text$		
Follow up	5 years		
Newcastle-Ottawa Scale	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort		

Bibliographic reference	Larsson M, Weiss B, Janson S, et al (2009) Associations between indoor environmental factors and parental-reported autistic spectrum disorders in children 6-8 years of age. NeuroToxicology 30(5), 822-831
	 drawn from the same community as the exposed cohort Ascertainment of exposure written self-report Demonstration that outcome of interest was not present at start of study Yes Comparability Comparability of cohorts on the basis of the design or analysis study controls for sex, age of the child (6, 7, or 8 years old), any smoking in mother (smoking during pregnancy and/or smoking during the child's first year and/or current smoking vs. No smoking in the mother), asthma in the child 2000 (no vs. yes), financial insecurity expressed as problems with paying bills (no, yes, no reply), and condensation on the inside of the windows in the child's room during winter time as a proxy for low ventilation (no, 1-5 cm, >5 cm) Outcome Assessment of outcome self-report (parent) Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts Unlikely to introduce bias Overall assessment=Moderate (concerns over parental report of
	outcome)
Source of funding	Government: Swedish Research Council for Environment, Agricultural Sciences and Spatial Planning (Formas), Swedish Asthma and Allergy Association's Research Foundation, and the Swedish Foundation for Health Care Sciences and Allergy Research.
Comments	None

D.1.67 Larsson 2010

Bibliographic reference	Larsson M, Hagerhed-Engman L, Kolarik B et al (2010) PVC as flooring material and its association with incident asthma in a Swedish child cohort study. Indoor air 20(6), 494-501
Study design	Cohort study
Objective	To examine the association between exposure to PVC flooring in the child's or parent's bedroom in the homes of children aged $1-3$ years and the incidence of asthma, rhinitis and eczema in the follow 5 years
Setting/Study location	Sweden
Number of participants	2779 children
Selected population	No

Bibliographic reference	Larsson M, Hagerhed-Engman L, I flooring material and its association Swedish child cohort study. Indoor	on with incident asthma in a
Participant characteristics	Building characteristics: Location: urban Dwelling type: Not reported Building age: Not reported Type of ownership/tenancy: Not reported Parental characteristics: Age: Not reported Current smoker (mother): 55 (54%) Hay fever: Not reported Atopy: Not reported but all mothers h	
Inclusion criteria		
Exclusion criteria	Diagnosis of asthma at baseline	
Type of pollutant/exposure	Phthalates in house dust (PVC floori	ng used as proxy)
Pollutant/exposure assessment	Self-report of use of PVC flooring	
Outcome	Doctor diagnosed asthma (incident)	
Results	PVC flooring in child's bedroom PVC flooring in parent's bedroom Multi-family house	aOR (95%CI) 1.52 (0.99, 2.35) 1.46 (0.96, 2.23) 1.48 (0.86, 2.57)
Follow up	5 years	
Newcastle-Ottawa Scale	Selection Representativeness of the exposed of truly representative of the average Selection of the non-exposed cohort of drawn from the same community at Ascertainment of exposure written self-report Demonstration that outcome of interest of the Yes Comparability Comparability Comparability of cohorts on the basis of study controls for smoking in the hole of study controls for any additional factorization that outcome Assessment of outcome record linkage Was follow-up long enough for outcome Yes Adequacy of follow up of cohorts Unlikely to introduce bias Overall assessment=Low	child in the community as the exposed cohort est was not present at start of study as of the design or analysis ome ctor s

Bibliographic reference	Larsson M, Hagerhed-Engman L, Kolarik B et al (2010) PVC as flooring material and its association with incident asthma in a Swedish child cohort study. Indoor air 20(6), 494-501
Source of funding	Government: Swedish Research Council for Environment, Agricultural Sciences and Spatial Planning (Formas), Swedish Asthma and Allergy Association's Research Foundation, and the Swedish Foundation for Health Care Sciences and Allergy Research.
Comments	None

D.1.68 Lau 2000

_au 2000				
Bibliographic reference	Lau S, Illi S, Sommerfeld C, et al (2000) Early exposure to house-dust mite and cat allergens and development of childhood asthma: A cohort study. Lancet 356(9239), 1392-1397			
Study design	Prospective cohort study			
Objective	To examine the relationship development of asthma at 7			posure and the
Setting/Study location	Germany			
Number of participants	1314 children			
Selected population	Yes – selected as being at h	nigh risk fo	r asthma	
Participant characteristics	Description Sex Age (years) Ethnicity Education SES Building characteristics		Not reported Not reported Not reported Not reported Not reported Not reported	
Inclusion criteria	At risk of asthma			
Exclusion criteria	Not reported			
Type of pollutant/exposure	House dust mite and cat allergen			
Pollutant/exposure assessment	Parents collected dust samples according to detail ed instructions			
Outcome	Wheezing Physician-diagnosed asthma			
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between exposure to house dust mite, cat allergen, wheeze and asthma			
		Current	wheeze	Asthma diagnosed by a doctor
		aOR (95	5%CI)	aOR (95%CI)
	Cat allergen exposure (Fel d1)			
	0.216 – 47μg/g	1.47 (0.7	72, 1.26)	1.52 (0.64, 2.62)
	Mite allergen exposure (Der	p1 + Der	f1)	

Bibliographic reference	Lau S, Illi S, Sommerfeld C mite and cat allergens and cohort study. Lancet 356(9	development of childhoo	
	0.981 - 240µg/g	1.03 (0.52, 2.04)	0.72 (0.26, 2.00)
Follow up	7 years		
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the experimental selection of the non-exposed of the experimental drawn from the same community. Ascertainment of exposure objective sampling Demonstration that outcome yes. Comparability Comparability Comparability of cohorts on the study controls for family his ostudy controls for social state. Outcome Assessment of outcome outcome independent blind assessment of outcome Assessment of outcome outcome outcome outcome. Assessment of outcome outcome outcome outcome. Adequacy of follow up of controls for social state. Overall risk of bias – Low	at high risk of asthma d cohort munity as the exposed coho of interest was not present the basis of the design or a story of atopy atus nent or outcomes to occur	t at start of study
Source of funding	Government: German Minis	try of Research and Educa	tion
Comments			

D.1.69 Le Moual 2012

Bibliographic reference	Le Moual N, Varraso R, Siroux V, et al(2012) Domestic use of cleaning sprays and asthma activity in females. The European respiratory journal 40(6), 1381-9
Study design	Prospective cohort study
Objective	To assess, the associations between home cleaning, particularly the use of household cleaning sprays, and asthma activity
Setting/Study location	France
Number of participants	683 adult women
Selected population	Yes – only women with asthma and the first degree relatives included but this study focuses on 683 women
Participant characteristics	Description

Bibliographic reference	Le Moual N, Varraso sprays and asthma journal 40(6), 1381-	activity in female		
Telefelle	Sex	,		
	Female		683 (100%)	
	Age (years) Mean (S	D)	43.8 (15.5)	
	Ethnicity		Not reported	
	Education			
	Primary		154 (22.6%)	
	Secondary		177 (26.0%)	
	University		350 (51.4\$)	
	SES		Not reported	
	Building characteristi	cs	Not reported	
Inclusion criteria	Had detailed informa sprays, was collected		nestic exposures, in	particular to
Exclusion criteria	Not reported			
Type of pollutant/exposure	Domestic use of clea	ning sprays		
Pollutant/exposure assessment	Current domestic exposures (last 12 months) were based on 24 domestic exposure variables including nine cleaning tasks and 15 cleaning agents			
	Exposure to sprays v sprays (furniture, glas refreshing, other use	ss-cleaning, carpe	t, mopping the floor	
Outcome	Asthma			
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between domestic self-reported exposure to cleaning products and asthma.			
		Current asthma	Controlled asthma	Poorly controlled asthma
	Home cleaning ≥1 day/week	1.34 (0.87, 2.05)	1.12 (0.66– 1.90)	1.50 (0.88–2.52)
	1 type of spray used ≥1 day/week	0.68 (0.44, 1.04)	0.67 (0.38– 1.18)	0.65 (0.38–1.12)
	≥ 2 types of sprays used ≥1 day/week	1.67 (1.08, 2.56)	1.32 (0.75– 2.34)	2.04 (1.25–3.32)
Follow up	12 months			
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of selected group of which selection of the non- of drawn from the same ascertainment of expension of the non- of drawn from the same ascertainment of expension of the non- of drawn from the same ascertainment of expension of the non- of drawn from the non- of drawn fr	vomen exposed cohort ne community as t oosure utcome of interest	he exposed cohort was not present at	·

Bibliographic reference	Le Moual N, Varraso R, Siroux V, et al(2012) Domestic use of cleaning sprays and asthma activity in females. The European respiratory journal 40(6), 1381-9
	study controls for smoking habits study controls for smoking habits
	 study controls for any additional factors - age, , body mass index and occupational exposure.
	Outcome
	Assessment of outcome
	• self-report
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	complete follow up - all subjects accounted for
	Overall risk of bias: High (Concerns over self-report of exposure and outcomes)
Source of funding	Government: French Agency of Health Safety, Environment and Work; National Research Agency - Health Environment, Health-Work Program Industry: Merck Sharp & Dohme; Hospital Program of Clinical Research (PHRC)-Paris
Comments	

D.1.70 Li 2006

LI 2006	
Bibliographic reference	Li R, Weller E, Dockery DW, et al. (2006) Association of indoor nitrogen dioxide with respiratory symptoms in children: application of measurement error correction techniques to utilize data from multiple surrogates. J Expo Sci Environ Epidemiol 16(4): 342-50.
Study design	Prospective cohort study
Objective	To evaluate the effect of indoor nitrogen dioxide exposure on the annual risk of lower respiratory tract symptoms
Setting/Study location	Different communities, USA
Number of	Number of dwellings: 1,137
dwellings and participants	Number of participants: 1,137 children
Selected population	No
Building and Participant characteristics	Building characteristics: Location: unclear Dwelling type: not reported Building age: not reported Type of ownership/tenancy: not reported Participant characteristics: Age: not reported Smokers living in the property: 13% Allergies: not reported Parental history of asthma: 13%

Bibliographic reference	Li R, Weller E, Dockery DW, et al. (2 dioxide with respiratory symptoms measurement error correction techn surrogates. J Expo Sci Environ Epic	in children: application of niques to utilize data from multiple
Inclusion criteria	Households of children between 7 and communities across the USA were inc	
Exclusion criteria	Not reported	
Building factor/exposure	Gas stove with a pilot light, gas stove wood stove, number of rooms in the ho	•
Building factor/exposure assessment	Building factors were ascertained by a reported questionnaire.	sking participants to complete a self-
Outcome	Lower respiratory tract symptoms (not	specified/defined)
Results	Building characteristic	Odds ratio (95%CI)
	Gas stove, no pilot	0.68 (0.42, 1.10)
	Gas stove, pilot	1.54 (0.94, 2.25)
	Stove heater	1.61 (1.05, 2.47)
	Fan	0.93 (0.81, 1.07)
	Wood stove	0.91 (0.66, 1.25)
	Kerosene heater	1.41 (0.96, 2.07)
	Per room increase in the household	0.99 (0.92, 1.06)
Follow up	Up to 2.5 years	
	A respiratory symptom questionnaire was administered at the time of enrolment. A year later, a second questionnaire and pulmonary function examination were administered. Between 12 and 18 months later, a final health questionnaire was administered. A third of the population was selected by stratified random sampling (accounting for smoking, and mai source of nitrogen dioxide) to have their household air quality monitored. Based on residential indoor air quality, an annual average nitrogen dioxid measurement was obtained. Statistical analysis: multivariate logistic regression was performed to associations	
Newcastle-Ottawa Scale	Selection Representativeness of the exposed cohort • no description of the derivation of the cohort Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • self-reported Demonstration that outcome of interest was not present at start of study • No Comparability Comparability of cohorts on the basis of the design or analysis • study controls for single marital status, higher education status, parental history of bronchitis or emphysema, parental history of asthma, gender, age, and the total packs of cigarette smoking inside the child's home Outcome	

Bibliographic reference	Li R, Weller E, Dockery DW, et al. (2006) Association of indoor nitrogen dioxide with respiratory symptoms in children: application of measurement error correction techniques to utilize data from multiple surrogates. J Expo Sci Environ Epidemiol 16(4): 342-50.
	Assessment of outcome
	• self-reported
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	complete follow up - all subjects accounted for
	Overall risk of bias: high (Concerns over self-report of exposure and outcome
Source of funding	Government: National Institute of Environmental Health Sciences (NIEHS)
Comments	None

D.1.71 Li 2016

Bibliographic reference	Li W, Dorans K S, Wilker E H et al. (2016) Residential Proximity to Major Roadways, Fine Particulate Matter, and Adiposity: The Framingham Heart Study. Obesity 24, 2593-2599			
Study design	Prospective cohort study			
Objective	To examine the associations of residential-based estimates of ambient $PM_{2.5}$ exposure and proximity to the nearest major roadway with body mass index (BMI) and MDCT-based measures of abdominal adiposity			
Setting/Study location	United States			
Number of participants	2,372 adults			
Selected population	No			
Participant characteristics	Description	No.	%	
	Sex	Not reported	Not reported	
	Age (years); mean (SD)	53.9	11.8	
	Ethnicity			
	Cases/selected population/comorbidity			
	Cardiovascular disease	192	8.1	
	Diabetes	173	7.3	
	Socio-economic status			
	Education			
	<high school<="" td=""><td>41</td><td>1.7</td></high>	41	1.7	
	High school	521	22.0	
	Some college	771	32.5	
	College graduate	1,039	43.8	
	Building characteristics	Not reported	Not reported	
Inclusion criteria	Men aged ≥ 35 years old Women were aged ≥ 40 years old and not pregnant Because of physical constraints of the scanner, all participants weighed <350 lbs (160 kg)			

Bibliographic reference	Li W, Dorans K S, Wilker E H et al. (2016) Residential Proximity to Major Roadways, Fine Particulate Matter, and Adiposity: The Framingham Heart Study. Obesity 24, 2593-2599		
Study design	Prospective cohort study		
Exclusion criteria			
Type of pollutant/exposure	PM 2.5 and proximity to the nearest major roadway		
Pollutant/exposure assessment	Authors used a spatial-temporal model to estimate $PM_{2.5}$ concentrations at a 1 X 1 km2 resolution based on residential addresses.		
Outcome	Obesity		
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association of distance to a major roadway and PM _{2.5} with adiposity measures. We defined major roadways as primary highways with limited access (A1), primary roads without limited-access (A2), or secondary and connecting roads (A3). And we estimated residential distance to the nearest major roadway based on the geocoded addresses.		
	, ,	Obesity	
		OR (95%CI)	
	Closer to a major roadway	1.10 (0.97; 1.25)	
	2003 annual average PM _{2.5}	1.01 (0.92,1.12)	
Follow up	Not reported	, , ,	
Study methods	Both standing height and weight were measured without shoes according to a standardized protocol. Height was recorded to the nearest 1/4 inch, and weight was recorded to the nearest pound (rounded up if ≥0.5 pound). Body mass index (BMI) was calculated as weight (kg)/ height (m)2 Authors fit multivariable linear regression models for continuous BMI, subcutaneous adipose tissue (SAT), and visceral adipose tissue (VAT), and multivariable logistic regression models for a binary indicator of obesity (BMI ≥30 kg/m2) adjusting for confounding factors		
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average adult population in the community Selection of the non-exposed cohort • no description of the derivation of the non-exposed cohort Ascertainment of exposure • validated measurement used Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for age, sex, smoking, alcohol intake, educational level; physical activity, medication use and household income Outcome Assessment of outcome • Validated measurement used Was follow-up long enough for outcomes to occur • Yes		

Bibliographic reference	Li W, Dorans K S, Wilker E H et al. (2016) Residential Proximity to Major Roadways, Fine Particulate Matter, and Adiposity: The Framingham Heart Study. Obesity 24, 2593-2599
Study design	Prospective cohort study
	Adequacy of follow up of cohorts • no statement Overall risk of bias: low
Source of funding	Government: USEPA , National Heart, Lung, and Blood Institute of the National Institutes of Health, National Institutes of Environmental Health Sciences
Comments	

D.1.72 Lindgren 2013

Linagren 2013				
Bibliographic reference	Lindgren A, Stroh E, Bjork J, et al (2013) Asthma incidence in children growing up close to traffic: a registry-based birth cohort. Environmental health: a global access science source 12, 91			
Study design	Prospective cohort study			
Objective	to investigate if children growing up close to high traffic intensity are at higher risk of developing asthma or other obstructive respiratory disease in early childhood			
Setting/Study location	Sweden			
Number of participants	7898			
Selected population	No			
Participant	Description	No.	%	
characteristics	Sex			
	Male	3784	49	
	Female	3996	51	
	Missing	118		
	Maternal age (years) Not reported			
	Ethnicity	Not reported		
	Parental allergy Yes No Missing	3177 3751 970	46 54	
	Parental education > 12 years > 9 – 12 years > ≤ 9 years > Missing	5612 1792 297 197	73 23 4	
	Annual family income	Not reported		
	Building characteristics Owned house Tenant-owned apartment	2783 2242	36 29	

Bibliographic reference	Lindgren A, Stroh E, Bjork J, et al (2013) Asthma incidence in children growing up close to traffic: a registry-based birth cohort. Environmental health: a global access science source 12, 91					
	Rented apartment Other Missing		2616 101 156		34 1	
Inclusion criteria	Children born in Scania Mothers registered as living in the municipalities Malmö, Svedala, Vellinge or Trelleborg					
Exclusion criteria	Not registered in S Address for birth y No Child Health Ca	ear not r	•			
Type of pollutant/exposure	Proximity to traffic					
Pollutant/exposure assessment	To assess exposure to traffic, we identified the road with the heaviest traffic intensity within 100 m of the residence. Traffic intensity was categorized as "no road", "road with 0–2880 cars/day", "2880–8640 cars/day", "8640–14400 cars/day", and "≥14400 cars/day", based upon daily (24-hour) mean levels.					
Outcome	Asthma, bronchioli	itis, obstı	ructive bron	chitis		
Results	Adjusted hazard ra	atios (aH	Rs) and 95°	% confidence in	nterva	ls (CIs)
		Bronch	iolitis	Obstructive bronchitis		Asthma
		aHR (9	5%CI)	aHR (95%CI)	١	aHR (95%CI)
	Heaviest road ≤100 m, birth address					
	0-8640 cars/day	1.00		1.00		1.00
	≥8640 cars/day	0.7 (0.6	•	1.0 (0.9,1.2)		0.7 (0.6, 0.9)
	Heaviest road ≤100 m, never moved					
	0-8640 cars/day	1.00		1.00		1.00
	≥8640 cars/day	0.7 (0.6	6, 0.9)	1.0 (0.8,1.2)		0.7 (0.6, 0.9)
Follow up	Up to 6 years					
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • Objective report Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability Comparability of cohorts on the basis of the design or analysis • study controls for sex, birth weight, smoking during pregnancy, environmental tobacco smoke (ETS), mold at home, parental allergy, furred pets at home, breastfeeding, parental origin, parental education, problems to pay bills, type of housing, and birth year. Outcome					

Bibliographic reference	Lindgren A, Stroh E, Bjork J, et al (2013) Asthma incidence in children growing up close to traffic: a registry-based birth cohort. Environmental health: a global access science source 12, 91
	Assessment of outcome
	record linkage
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	complete follow up - all subjects accounted for
	Overall risk of bias: Low
Source of funding	Government: Swedish Council for Working Life and Social Research
	Academic: Faculty of Medicine, Lund University, Sweden.
Comments	

D.1.73 Litonjua 2002

Litorijaa 2002			
Bibliographic reference	Litonjua AA, Milton DK, Celedon JC, et al (2002) A longitudinal analysis of wheezing in young children: The independent effects of early life exposure to house dust endotoxin, allergens, and pets. Journal of Allergy and Clinical Immunology 110(5), 736-742		
Study design	Prospective cohort study		
Objective	to examine the longitudinal association of exposure to HDE, allergen levels, and the presence of a dog in the home and wheezing over a 4- year period.		
Setting/Study location	United States		
Number of participants	226 children		
Selected population	Yes –family history of atopy		
Participant characteristics	Description Sex Female Male Age (years) – Median (range) Ethnicity White Black Hispanic Asian Education SES (by percentage of households below poverty,) <10% 10%-<20% ≥20% Building characteristics	117 (51.8%) 109 (48.2%) 2.87 (1.10-4.99) 186 (82.3%) 18 (8.0%) 13 (5.8%) 9 (4.0%) Not reported 167 (73.9%) 46 (20.4%) 13 (5.8%) Not reported	

Bibliographic reference	Litonjua AA, Milton DK, Celedon JC, et al (2002) A longitudinal analysis of wheezing in young children: The independent effects of early life exposure to house dust endotoxin, allergens, and pets. Journal of Allergy and Clinical Immunology 110(5), 736-742		
Inclusion criteria	Parents allergic to house dust or house animals, or mould	e dust mites, cockroaches, pollens,	
Exclusion criteria	Not reported		
Type of pollutant/exposure	Pets at home		
Pollutant/exposure assessment	Objective sampling of house dust for a	llergens	
Outcome	Wheeze		
Results	Adjusted odds ratios (aORs) and 95% association between pets at home and		
		Repeated wheeze	
		aOR (95%CI)	
	Dog in the home	0.12 (0.01-0.97)	
	Fel d 1 ≥1 µg/g	0.61 (0.27-1.35)	
Follow up	4 years		
(Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • selected group of children at risk of asthma Selection of the non-exposed cohort • drawn from the same community as the exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • Objective sampling Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for maternal asthma • study controls for any additional factors as follows - maternal age, sex, prematurity, area of residence, and clustering of outcomes. Outcome Assessment of outcome • self-report Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for		
Source of funding	Overall risk of bias – Moderate (concerns over self-report of outcomes) Government: National Institute of Allergy and Infectious Disease; National Institute for Environmental Health Science		
Comments	modele of Environmental Florida Colonido		

D.1.74 Lynch 2014

-yrich 2014			
Bibliographic reference	Lynch S V, Wood R A, Boushey H et exposure to allergens and bacteria curban children. The Journal of allerg 593-601.e12	on recurrent wheeze and atopy in	
Study design	Prospective cohort study		
Objective	To determine whether early-life exposure to house dust obtained from innercity homes is associated with development of allergic sensitization and wheezing		
Setting/Study location	United States		
Number of participants	560 children		
Selected population	Yes – high risk for atopy		
Participant characteristics	Description Sex Male Age (years) – Median (range) Ethnicity Black Hispanic Other Education (mother complete high school) SES by household income) <\$15,000 Building characteristics Residence in an area with more than 2 level	240 (51%) 333 (71%) 92 (20%) 42 (9%) 273 (55%) 321 (69%) Not reported 20% of residents below the poverty	
	Mother or father with allergic rhinitis, ed Birth at 34 weeks' gestation or later	czema, and/or asthma	
Exclusion criteria	Not reported		
Type of pollutant/exposure	Pets and dust mite		
Pollutant/exposure assessment	Household dust samples from the living room (chair or sofa and floor) and child's bedroom (mattress and floor) were collected		
Outcome	Aeroallergen sensitization - defined by a wheal 3mm or more, larger than that elicited by the saline control on skin prick testing or a specific IgE level of 0.35 kU/L or greater. Recurrent wheeze was defined as parental report of at least 2 wheezing		
	episodes, with at least 1 episode occurring in the third year. Eczema was defined as a score of 1.0 or greater on the Eczema Area and Severity Index14 at age 3 years.		

Bibliographic reference	Lynch S V, Wood R A, Boushey H et.al (2014) Effects of early-life exposure to allergens and bacteria on recurrent wheeze and atopy in urban children. The Journal of allergy and clinical immunology 134(3), 593-601.e12		
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between pets at home and repeated wheeze		
		Recurrent wheeze	
	for 1-log increase in allergen level.	aOR (95%CI)	
	Fel d 1	0.71 (0.58, 0.88)	
	Can f 1	1.00 (0.79, 1.28)	
	Der f 1	0.92 (0.73, 1.15)	
Follow up	2 years		
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed co • selected group of children in inner cir Selection of the non-exposed cohort • drawn from the same community as Ascertainment of exposure • Objective sampling Demonstration that outcome of interes • Yes Comparability Comparability Comparability of cohorts on the basis of • study controls for environmental tobal • study controls for additional factor as Outcome Assessment of outcome • self-report Was follow-up long enough for outcome • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accomputations • Comparation of the properties of the exposure	the exposed cohort It was not present at start of study of the design or analysis acco smoke is follows, age, sex, and stress hes to occur	
Source of funding	Government: National Institute of Allergy and Infectious Diseases, National Institutes of Health, National Center for Advancing Translational Sciences, National Institutes of Health		
Comments			

D.1.75 McConnell 2002

Bibliographic reference	McConnell R, Berhane K, Gilliland F et.al (2002) Indoor risk factors for asthma in a prospective study of adolescents. Epidemiology 13(3), 288-295
Study design	Prospective cohort study

Bibliographic reference	McConnell R, Berhane R asthma in a prospective 288-295		2) Indoor risk factors for Epidemiology 13(3),	
Objective	To determine the association of indoor exposures with the development of asthma among adolescents and children entering adolescence			
Setting/Study location	United States			
Number of participants	3535 children with no history of asthma			
Selected population	No			
Participant	Description	No.	%	
characteristics	Sex			
	Male	396	23.6	
	Female	387	20.8	
	Age (years)			
	< 9	236	21.7	
	9 - 11	279	23.0	
	> 11	241	21.7	
	Ethnicity			
	White	520	25.1	
	Black	30	17.1	
	Asian	17	9.0	
	Hispanic	190	19.8	
	Other	26	25.0	
	(Maintenance) medication use	Not reported	Not reported	
	Parental asthma and/or atopic	Not reported	Not reported	
	Parental education (years)	Not reported	Not reported	
	Annual family income	Not reported	Not reported	
	Building characteristics	Not reported	Not reported	
Inclusion criteria	Children from schools in r populations	neighbourhoods with sta	ble middle income	
Exclusion criteria	Children diagnosed with asthma History of cystic fibrosis Severe chest injury or chest surgery			
Type of pollutant/exposure	Pet dander Mould/mildew Gas stove			
Pollutant/exposure assessment	Information on characteristics of the child's home environment was collected in a baseline questionnaire. This includes the presence of water damage, mould or mildew and the use of combustion sources.			
Health outcome	Asthma defined by wheezing			
Results	Adjusted relative risks (aRRs) and 95% confidence intervals (CIs) for indoor exposures and the risk of asthma by baseline history of wheeze			

Bibliographic reference			(2002) Indoor risk factors for ents. Epidemiology 13(3),
		Asthma with wheeze	Asthma with no wheeze
		aRR (95%CI)	aRR (95%CI)
	Any pet	1.10 (0.60, 2.00)	1.60 (1.00, 2.50)
	Water damage	0.80 (0.50, 1.40)	1.4 (0.90, 2.00)
	Mould/mildew	0.60 (0.40, 0.90)	1.10 (0.80, 1.60)
	Wood fire	0.90 (0.60, 1.50)	0.90 (0.60, 1.30)
	Gas stove	1.20 (0.70, 2.00)	1.3 (0.80, 2.00)
Follow up	5 years		
Risk of bias (Newcastle-Ottawa Scale)	truly representative Selection of the non- drawn from the san Ascertainment of exp Questionnaire and Demonstration that of Yes Comparability Comparability Comparability of cohe study controls for far a community of resid health insurance placempared with mid Outcome Assessment of outcome Assessment of outcome Assessment of outcome Yes Adequacy of follow up complete follow up Overall risk of bias	ne community as the exposure objective samples taker utcome of interest was a orts on the basis of the orts amily history of asthma ny additional factor — go ence, , child's history of an, and high or low soci dle-income families. me nough for outcomes to or p of cohorts - all subjects accounted - Low	posed cohort not present at start of study design or analysis ender, age, race and ethnicity, allergy, membership in a loeconomic status (SES)
Source of fulfulling		n Sciences, the Environi and Blood Institute	mental Protection Agency, the
Comments	Authors suggest that		n and potentially remediable nts.

D.1.76 McConnell 2006

Bibliographic reference	McConnell R, Berhane K, Yao L et.al (2006) Traffic, susceptibility, and childhood asthma. Environmental health perspectives 114(5), 766-72			
Study design	Prospective cohort study			
Objective	To examine characteristics that might increase childhood susceptibility to the effects of traffic-related air pollution			
Setting/Study location	United States			
Number of participants	5,341 children			
Selected population	No			
Participant	Description	No.	%	
characteristics	Sex (male)	2,425	51	
	Age (years)	Not reported	Not reported	
	Ethnicity			
	North American Indian	44	0.93	
	Asian	170	3.6	
	Black	197	4.2	
	Hispanic white	2,617	55	
	Non-Hispanic white	1,682	35	
	Other	32	0.67	
	Cases/selected population/comorbidity	Not reported	Not reported	
	Socio-economic status			
	Parental education			
	< 12th grade	982	22	
	Grade 12	880	20	
	Some post-high school	1,681	38	
	Four years of college	512	11	
	Some postgraduate	417	9.3	
	Building characteristics			
	Water damage	653	14	
	Mould or mildew	1,068	25	
Inclusion criteria	Not reported			
Exclusion criteria	Not reported			
Type of pollutant/exposure	Proximity to a major road			
Pollutant/exposure assessment	Authors estimated distance of each participant's residence to the nearest major road, including freeways, other highways, and arterial roads. Participant residence addresses were standardized, and their locations were geocoded to 13 m perpendicular to the side of the adjacent road, using the Tele Atlas Multinet road network data			
Outcome	Asthma and wheeze			
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between proximity to a major road, asthma and wheeze			

Bibliographic reference	McConnell R, Berhane K, Yao L et.al (2006) Traffic, susceptibility, and childhood asthma. Environmental health perspectives 114(5), 766-72				
		Lifetime asthma	Prevalent asthma	Current wheeze	
	Major road distance (metres)	aOR (95%CI)	aOR (95%CI)	aOR (95%CI)	
	> 300	1.00	1.00	1.00	
	150-300	0.92 (0.73-1.15)	1.04 (0.82, 1.33)	1.02 (0.82, 1.27)	
	75–150	1.06 (0.82-1.36)	1.33 (1.02, 1.72)	1.30 (1.02, 1.66)	
	< 75	1.29 (1.01–1.66)	1.50 (1.16, 1.95)	1.40 (1.09, 1.78)	
Follow up	5 years				
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • no description of the derivation of the non-exposed cohort Ascertainment of exposure • written self-report (questionnaire) Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for child's age, sex, race, community, and language of questionnaire completion Outcome Assessment of outcome • record linkage • self-report Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • subjects lost to follow up unlikely to introduce bias Overall risk of bias: Low				
Source of funding	Government: National Institute of Environmental Health Science the U.S. Environmental Protection Agency, (the Southern California Particle centre)], the South Coast Air Quality Management District, the National Heart, Lung, and Blood Institute Charity: Hastings Foundation.				
Comments					

D.1.77 McCormack 2009

Bibliographic reference	McCormack MC, Breysse PN, Matsui EC, et al (2009) In-Home Particle Concentrations and Childhood Asthma Morbidity. Environmental Health Perspectives 117(2), 294-298
Study design	Prospective cohort

	14 0 1 140 B 511 11 1 1			
Bibliographic reference	McCormack MC, Breysse PN, Matsui EC, et al (2009) In-Home Particle Concentrations and Childhood Asthma Morbidity. Environmental Health Perspectives 117(2), 294-298			
Objective	To investigate the effect of in-home coarse and fine PM on respiratory symptoms, rescue medication use, and acute health care use among preschool asthmatic children			
Setting/Study location	United States			
Number of participants	150			
Selected population	Yes – all had asthma			
Participant characteristics	Description Sex Age (years); mean (Range) Ethnicity African American 91 Caucasian Other Cases/selected population/comorbidity Socio-economic status (reported as Caregiver education level) Eighth grade/some high school High school Some college	(58%) 4.4 (2–6) (91%) (5%) (4%) (38%) (43%) (19%)		
Inclusion criteria	Participants had to report a physician diagnosis of asthma and had to have symptoms of asthma and/or medication use for asthma in the previous 6 months. Other inclusion criteria were age between 2 and 6 years and residence within one of nine contiguous ZIP codes within East Baltimore.			
Exclusion criteria	Not reported			
Type of pollutant/expo sure	P2,5 and PM 10.0			
Pollutant/expo sure assessment	A trained environmental technician completed home visits. Environmental monitoring was carried out at baseline and at 3 and 6 months. At each time period, integrated air sampling in the child's bedroom over a 3-day period was performed.			
Outcome				
Results	Adjusted Incident rate ratios (aHRs) and	95% confidence intervals (CIs)		
	PM _{2.5} –10 (per 10 μg/m³ increase)	alRR (95%CI)		
	Cough, wheezing, chest tightness	1.06 (1.01, 1.12)		
	Asthma symptoms causing children to slow down	1.08 (1.02, 1.14)		
	Symptoms with running	1.00 (0.94, 1.08)		
	Nocturnal symptoms	1.08 (1.01, 1.14)		
	Limited speech	1.11 (1.03, 1.19)		
	Rescue medication use	1.06 (1.01, 1.10)		

Bibliographic reference	McCormack MC, Breysse PN, Matsui Concentrations and Childhood Asthm Perspectives 117(2), 294-298		
	PM _{2.5} (per 10 μg/m ³ increase)	aIRR (95%CI)	
	Cough, wheezing, chest tightness	1.03 (0.99, 1.07)	
	Asthma symptoms causing children to slow down	1.04 (1.0, 1.09)	
	Symptoms with running	1.07 (1.02,1.11)	
	Nocturnal symptoms	1.06 (1.01, 1.10)	
	Limited speech	1.07 (1.00, 1.14)	
	Rescue medication use	1.04 (1.01, 1.08)	
Follow up			
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed coho • truly representative of the average chill community Selection of the non-exposed cohort • no description of the derivation of the representation of the derivation of the representation of t	d with asthma population in the non-exposed cohort was not present at start of study the design or analysis at education level, season, indoor fine PM, sto occur	
Source of funding	Government: National Institute of Environmental Health Sciences (NIEHS); U.S. Environmental Protection Agency Academic: Johns Hopkins NIEHS Center in Urban Environmental Health		
Comments			

D.1.78 Mahalingaiah 2014

Bibliographic reference	Mahalingaiah S, Hart JE, Laden F, et al (2014) Air pollution and risk of uterine leiomyomata. Epidemiology (Cambridge, and Mass.) 25(5), 682-8				
Study design	Prospective cohort study				
Objective	To examine if air pollution exposure is associated with the occurrence of uterine leiomyomata				
Setting/Study location	United States				
Number of participants	85251 women				
Participant	Description	No. (%)			
characteristics	Sex				
	Female	85251 (100%)			
	Age (years); mean (SD)	42.6 (5.3%)			
	Ethnicity Caucasian	(24%)			
	SES	Not reported			
Inclusion criteria	alive at the given questionnaire cycle, premenopausal, free of cancer (other than non-melanoma skin cancer), had no history of infertility, had intact uteri, and did not have a diagnosis of uterine leiomyomata prior to 1993				
Exclusion criteria	More than 1 home address in continental US				
Type of pollutant/expo sure	Proximity to traffic				
Pollutant/expo sure assessment	Authors calculated distance to road at each residential address as a proxy for all exposures related to traffic. Distance to road (in meters) for all available nurses' addresses was determined using geographic information system (GIS) software (ArcGIS, version 9.2; ESRI, Redlands, CA) and the ESRI StreetPro 2007 data layer.				
Outcome	uterine leiomyomata				
Results	Adjusted hazard ratios (aHRs) and 95% co	onfidence intervals (CIs)			
		Uterine leiomyomata			
	Distance to A1–A3 roadway (metres)	aHR (95%CI)			
	0–50 51–199 > 200	1.01 (0.93, 1.09) 1.04 (0.98, 1.11) Reference			
	Distance to A1–A2 roadway (metres)				
	0–50 51–199 > 200	1.00 (0.80, 1.25) 1.02 (0.91, 1.15) Reference			
Follow up	14 years				
	•				

Bibliographic reference	Mahalingaiah S, Hart JE, Laden F, et al (2014) Air pollution and risk of uterine leiomyomata. Epidemiology (Cambridge, and Mass.) 25(5), 682-8
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort a) truly representative of the average female population in the community Selection of the non-exposed cohort b) no description of the derivation of the non-exposed cohort Ascertainment of exposure c) validated measurement used Demonstration that outcome of interest was not present at start of study d) Yes Comparability Comparability of cohorts on the basis of the design or analysis e) study controls for age, race, region, body mass index, smoking status, parity, oral contraceptive use, age at menarche Outcome Assessment of outcome f) self-report Was follow-up long enough for outcomes to occur g) Yes Adequacy of follow up of cohorts h) people lost to follow up unlikely to introduce bias Overall risk of bias: low
Source of funding	Government: National Institute of Child Health and Human Development, National Cancer Institute, National Institute for Environmental Health Sciences Academic: Boston University Department of Obstetrics and Gynecology, and the Massachusetts Institute of Technology Center for Environmental Health Sciences Translational Pilot Project Program
Comments	

D.1.79 Mahalingaiah 2016

Bibliographic reference	Mahalingaiah S, Hart J E, Laden F, et al (2016) Adult air pollution exposure and risk of infertility in the Nurses' Health Study II. Human reproduction (Oxford, and England) 31(3), 638-47
Study design	Prospective cohort study
Objective	To assess the relation between incident infertility and air pollution exposures as measured by exposure to PM less as well as traffic-related exposure measured by distance to road

Bibliographic reference	Mahalingaiah S, Hart J E, Ladand risk of infertility in the Nu (Oxford, and England) 31(3), 6	rses' Health Study II.		
Setting/Study location	United States			
Number of participants	36 294 women			
Participant	Description No. %			
characteristics	Sex	All female	All female	
	Age (years); mean (SD) Ethnicity	38.7 (4.7)	-	
	Caucasian	107115.6	92	
	Cases/selected population/comorbidity	Not reported	Not reported	
	Socio-economic status	Not reported	Not reported	
	Building characteristics	Not reported	Not reported	
Inclusion criteria	Not reported			
criteria	Over 45 years of age No longer responded to questionnaires Had undergone a hysterectomy or tubal ligation Had previously been diagnosed with cancer (other than skin cancer) Were under 45 years of age and menopausal Had a partner who had undergone a vasectomy Had previously reported infertility			
Type of pollutant/expo sure	Proximity to traffic and particulate matter (PM)			
Pollutant/expo sure assessment	Authors calculated distance to road at each residential address as a proxy for all exposures related to traffic. Distance to road (in meters) for all available nurses' addresses was determined using geographic information system (GIS) software (ArcGIS, version 9.2; ESRI, Redlands, CA) and the ESRI StreetPro 2007 data layer.			
Outcome	Infertility			
Results	Adjusted hazard ratios (aHRs) a between proximity to traffic PM		tervals (CIs) for association	
		Infertility		
		aHR (95%	CI)	
	Distance to A1–A3 roadway (metres)			
	0–199 1.11 (1.02, 1.20)			
	200+	Ref		
	PM cumulative average exposu	re (Per 10 mg/m³ incre	ease)	
	PM ₁₀	1.06 (0.99	, 1.13)	
	PM _{2.5} – 10	1.10 (0.99	, 1.22)	
	PM _{2.5}	1.05 (0.93		

Bibliographic reference	Mahalingaiah S, Hart J E, Laden F, et al (2016) Adult air pollution exposure and risk of infertility in the Nurses' Health Study II. Human reproduction (Oxford, and England) 31(3), 638-47
Follow up	14 years
Study methods	On the baseline questionnaire and each follow-up questionnaire, women were asked to report if they had attempted to become pregnant for at least 1 year without success, the age at which this occurred and, if known, the reason or reasons for the infertility. Time-varying Cox proportional hazards models were used to assess the association of incidence of overall infertility or specific reasons for infertility with exposure to roadway proximity or each size fraction of PM. Authors examined possible confounding by numerous a priori selected risk factors for infertility or predictors of exposure including
Risk of bias	Selection
(Newcastle-	Representativeness of the exposed cohort
Ottawa Scale)	 truly representative of the average female population in the community Selection of the non-exposed cohort
	 no description of the derivation of the non-exposed cohort
	Ascertainment of exposure
	validated measurement used
	Demonstration that outcome of interest was not present at start of study
	• Yes
	Comparability Comparability of cohorts on the basis of the design or analysis
	• study controls for age, race, region, body mass index, smoking status, parity, oral contraceptive use, age at menarche
	Outcome Assessment of outcome
	self-report
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	people lost to follow up unlikely to introduce bias
	Overall risk of bias: low
Source of funding	Government: Reproductive Scientist Development Program, and the Building Interdisciplinary Research Careers in Women's Health, National Institute of Child Health and Human Development and the Massachusetts Institute of Technology Centre for Environmental Health Sciences Translational Pilot Project Program, National Cancer Institute, National Institute for Environmental Health Sciences, Eunice Kennedy Shriver National Institute of Child Health and Human Development Academic: Boston University CTS, Boston University Department of Obstetrics and Gynaecology, S.A.M
Comments	and Syndocology, O.A.W
Comments	

D.1.80 Mommers 2005

Mommers 2005						
Bibliographic reference	Mommers M, Jongmans-Liedekerken A W, Derkx R et.al (2005) Indoor environment and respiratory symptoms in children living in the Dutch-German borderland. International journal of hygiene and environmental health 208(5), 373-81					
Study design	Nested case-co	ontrol				
	To investigate the role of indoor environmental risk factors on respiratory symptoms in 7-8 year old children living in the Dutch-German borderland					
Setting/Study location	Germany and The Netherlands					
Number of participants	1562 children					
Selected population	No					
•	Description	German chil	dren	Dutch	children	
characteristics		Control	Cases	Contr	ol	Cases
		No. %	No.%	No. %)	No. %
	Sex	Not reported	Not reported	Not re	eported	Not reported
	Age (years)	Not reported	Not reported	Not re	eported	Not reported
	Ethnicity	Not reported	Not reported	Not re	eported	Not reported
	Maintenance medication use	Not reported	Not reported	Not re	eported	Not reported
	Parental asthma and/or atopic	Not reported	Not reported	Not re	eported	Not reported
	Parental education	Not reported	Not reported	Not re	eported	Not reported
	Annual family income	Not reported	Not reported	Not re	eported	Not reported
	Building charac	cteristics				
	Pets	93 (46.5)	106 (56.1))	182 (46)	197 (50.6)
	Presence of mould or damp spots	21 (10.6)	44 (23.8)		54 (13.7)	83 (21.2)
	Coal, wood, gas or oil for heating	8 (4.4)	13 (7.9)		10 (2.7)	12 (3.4)
	Double glazing or door and window seals as insulating measures	151 (74.4)	131 (70.4)		372 (93.5)	356 (90.4)
	Gas cooking	4 (2)	7 (3.8)		254 (67.9)	268 (71.1)

Bibliographic reference	Mommers M, Jongmans-Liedekerken A W, Derkx R et.al (2005) Indoor environment and respiratory symptoms in children living in the Dutch-German borderland. International journal of hygiene and environmental health 208(5), 373-81			
Inclusion criteria	Inclusion criteria for cases Asthmatic symptoms (reported wheezing and attacks of shortness of breath with wheezing in the past 12 months Coughing (reported coughing in the morning or during the day or evening, in the autumn and winter and coughing daily for about 3 months a year Inclusion criteria for controls No symptoms			
Exclusion criteria	Not reported			
Type of pollutant/exposure	Pet dander from	cooking and unvented g	as appliances	
Pollutant/exposure assessment	Assessed using corresponding questions from the International Study of Asthma and Allergies in Childhood (ISAAC) questionnaire: For pet keeping and mould or damp spots categories including duration of exposure were constructed (i.e. never exposed, exposed for a short period, exposed for a long period, and always exposed). Insulation included double glazing or door and window seals in the living room, child's bedroom, bathroom or kitchen. Heating was defined as favourable when central heating or electricity was used and unfavourable when coal, wood, gas or oil was used. Socio-economic status (SES) was defined as low, middle or high according to			
Health outcome	, , ,	el of education of the fat nptoms and coughing		
Results	Adjusted odds		confidence intervals (CIs) for association nptoms and coughing	
		Asthmatic symptoms	Coughing	
		OR (95%CI)	OR (95%CI)	
	Gender (male vs. female)	2.25 (1.63, 3.12)	1.30 (0.98, 1.72)	
	Pets			
	Short period in the past	1.21 (0.81, 1.82)	1.56 (1.10, 2.20)	
	Long period in the past	1.32 (0.83, 2.10)	1.10 (0.72, 1.68)	
	Always	2.18 (1.39, 3.42)	1.64 (1.09, 2.46)	
	Mould or damp spots			
	Short period	1.97 (1.21, 3.22)	2.03 (1.32, 3.14)	
	Long period	2.98 (1.10, 8.28)	3.25 (1.35, 8.28)	
	Always	0.76 (0.21, 2.57)	1.24 (0.40, 3.88)	
	Gas cooking w	ith cooker hood		
	Used daily	0.94 (0.60, 1.46)	0.93 (0.64,1.36)	

Bibliographic reference	environment	and respiratory sympto erland. International jo	n A W, Derkx R et.al (2005) Indoor oms in children living in the Dutch- urnal of hygiene and environmental
	Used regularly	1.27 (0.61, 2.64)	1.49 (0.80, 2.78)
	Not used	1.25 (0.69, 2.26)	1.25 (0.74, 2.11)
	Water heating		
	Unvented gas geyser	3.01 (1.21, 7.56)	1.74 (0.74, 4.12)
	Vented gas geyser	1.33 (0.83, 2.14)	1.28 (0.85, 1.94)
	Heating		
	Unfavourable vs. favourable	0.93 (0.34, 2.37)	1.52 (0.72, 3.23)
	Socio-econom	ic status (SES)	
	Middle vs. high	1.43 (1.00, 2.04)	1.53 (1.12, 2.10)
	Low vs. high	3.32 (1.88, 5.93)	3.37 (2.01, 5.71)
Follow up	12 months		
Risk of bias (Newcastle- Ottawa Scale)			

Bibliographic reference	Mommers M, Jongmans-Liedekerken A W, Derkx R et.al (2005) Indoor environment and respiratory symptoms in children living in the Dutch-German borderland. International journal of hygiene and environmental health 208(5), 373-81
Source of funding	Government: Study was financially supported by European Union, the Euregio Maas-Rhine, the Land Northrhine-West-phalia, the province of Limburg and the counties of Heinsberg, Midden-Limburg and Westelijke Mijnstreek
Comments	Though authors did not measure indoor NO_2 levels directly but used the presence of gas appliances as a proxy.

D.1.81 Morgenstern 2007

Bibliographic reference	Morgenstern V, Zutavern A, Cyrys J, et al (2007) Respiratory health and individual estimated exposure to traffic-related air pollutants in a cohort of young children. Occupational and environmental medicine 64(1), 8-16			
Study design	Prospective cohort study			
Objective	To estimate long-term exposure to traffic-related air pollutants on an individual basis and to assess adverse health effects using a combination of air pollution measurement data, data from geographical information systems (GIS) and questionnaire data			
Setting/Study location	Germany			
Number of participants	3577 children			
Selected population	No			
Participant	Description	No.	%	
characteristics	Sex (female) sex	1489	52.4	
	Age (years); mean (SD)	Not reported	Not reported	
	Ethnicity	Not reported	Not reported	
	Cases/selected population/comorbidity	Not reported	Not reported	
	Socio-economic status (maternal education)			
	<12 grades	925	29.7	
	≥12 grades	1853	59.5	
	Building characteristics			
	Home dampness	218	7.1	
	Indoor moulds	934	30.3	
Inclusion criteria	Not reported			
Exclusion criteria	Not reported			
Type of pollutant/expos ure	Particulate matter (PM), NO ₂ and p	proximity to the main roa	d	

Bibliographic reference	Morgenstern V, Zutavern A, Cyrys J, et individual estimated exposure to traffic young children. Occupational and envi	related air pollutants in a cohort of
Pollutant/expos ure assessment	All particulate matter and NO_2 measurements were made during 2-week intervals. The air was sampled for 15 min every 2 h for a total of approximately 42 hours per sampling period. The collection time was recorded by an electronic timer. For traffic data, circular buffers with radii of 50, 100, 250, 500, 1000, 2500 and 5000 m were created around the coordinates of interest and intersected with the road network. As it was not feasible to measure personal exposure to the traffic-related air pollutants NO_2 , $PM_{2.5}$ and $PM_{2.5}$ absorbance for all study subjects, exposure modelling was used.	
Outcome	Asthma, allergic symptoms and respirator	y infections
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between distance to major road, PM, NO ₂ , asthma, allergic symptoms and respiratory infections. aORs of symptoms associated with interquartile range of air pollution variables	
	Distance to main road<50m	
	Wheeze	1.14 (0.92, 1.42)
	Cough without infection	0.74 (0.55, 1.00)
	Dry cough at night	0.84 (0.61, 1.16)
	Asthmatic/spastic/ obstructive bronchitis	1.12 (0.88, 1.44)
	Respiratory infections	1.03 (0.86, 1.23)
	Sneezing, runny/stuffed nose	1.10 (0.87, 1.39)
Follow up	2 years	
Study methods	All data on health outcomes and potential confounding variables were obtained through questionnaires that were completed by the parents. The association between exposure and health outcomes was tested by multiple logistic regression, with adjustment for potential confounding factors. In addition, authors looked at the association between living close to major roads and the health effects. The cut-off for the variable "living close to major road" was based on the hypothesis that the largest contribution from large streets to air pollution is expected at short distances.	
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • no description of the derivation of the non-exposed cohort Ascertainment of exposure • validated measurement used Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for sex, parental atopy, maternal education, siblings, environmental tobacco smoke at home, use of gas for cooking, home dampness, indoor moulds and keeping pets	

Bibliographic reference	Morgenstern V, Zutavern A, Cyrys J, et al (2007) Respiratory health and individual estimated exposure to traffic-related air pollutants in a cohort of young children. Occupational and environmental medicine 64(1), 8-16
	Outcome
	Assessment of outcome
	questionnaires/self-report
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	subjects lost to follow up unlikely to introduce bias
	overall risk of bias: Moderate (concerns over self-report of outcomes)
	moderate risk: potential for response bias for outcome assessment
Source of funding	Not reported
Comments	

D.1.82 Morgenstern 2008

norgenstern z				
Bibliographic reference	Morgenstern V, Zutavern A, Cyrys J, et al (2008) Atopic diseases, allergic sensitization, and exposure to traffic-related air pollution in children. American journal of respiratory and critical care medicine 177(12), 1331-7			
Study design	Prospective cohort study	Prospective cohort study		
Objective		To assess the relationship between individual-based exposure to traffic-related air pollutants and allergic disease outcomes in a prospective birth cohort study during the first 6 years of life		
Setting/Study location	Germany			
Number of participants	5921 children			
Selected population	No			
Participant	Description	No.	%	
characteristics	Sex (female) sex	1,486	51.6	
	Age (years); mean (SD)	Not reported	Not reported	
	Ethnicity	Not reported	Not reported	
	Cases/selected population	Not reported	Not reported	
	Socio-economic status (m	aternal education)		
	<12 grades	1,909	70.1	
	≥12 grades	1,074	39.4	
	Building characteristics			
	Home dampness	89	3.1	
	Indoor moulds	415	15.1	
Inclusion criteria	Not reported			
Exclusion criteria	Not reported			

Bibliographic reference	Morgenstern V, Zutavern A, Cyrys J, et al (2008) Atopic diseases, allergic sensitization, and exposure to traffic-related air pollution in children. American journal of respiratory and critical care medicine 177(12), 1331-7	
Type of pollutant/expos ure	Particulate matter (PM), NO ₂ and proximity to the main road	
Pollutant/expos ure assessment	All particulate matter and NO_2 measurements were made during 2-week intervals. The air was sampled for 15 min every 2 h for a total of approximately 42 hours per sampling period. The collection time was recorded by an electronic timer. For traffic data, circular buffers with radii of 50, 100, 250, 500, 1000, 2500 and 5000 m were created around the coordinates of interest and intersected with the road network. As it was not feasible to measure personal exposure to the traffic-related air pollutants NO_2 , $PM_{2.5}$ and $PM_{2.5}$ absorbance for all study subjects, exposure modelling was used.	
Outcome	Asthma, Hay fever, eczema	
Results	Adjusted odds ratios (aORs) and 95% co	nfidence intervals (Cls)
	Distance to main road<50m	aOR (95%CI)
	Asthma	1.66 (1.01, 2.59)
	Hay Fever	1.16 (0.67, 2.00)
	Eczema	0.96 (0.72, 1.11)
Follow up	6 years	
Study methods	All data on health outcomes and potential confounding variables were obtained through questionnaires that were completed by the parents. Parents were asked the following: "Has a physician diagnosed any of the following diseases during the past year of life: asthmatic/spastic/ obstructive bronchitis, asthma, hay fever, allergic/eczema?" If the parents selected yes, the child was defined to have "physician-diagnosed disease," which was the primary outcome parameter. The association between exposure and health outcomes was tested by multiple logistic regression, with adjustment for potential confounding factors. In addition, authors looked at the association between living close to major roads and the health effects. The cut-off for the variable "living close to major road" was based on the hypothesis that the largest contribution from large streets to air pollution is expected at short distances.	
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • no description of the derivation of the non-exposed cohort Ascertainment of exposure • validated measurement used Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for sex, parental atopy, maternal education, siblings, environmental tobacco smoke at home, use of gas for cooking, home dampness, indoor moulds and keeping pets	

Bibliographic reference	Morgenstern V, Zutavern A, Cyrys J, et al (2008) Atopic diseases, allergic sensitization, and exposure to traffic-related air pollution in children. American journal of respiratory and critical care medicine 177(12), 1331-7
	Outcome
	Assessment of outcome
	• questionnaires/self-report
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	subjects lost to follow up unlikely to introduce bias
	overall risk of bias: Moderate (potential for recall bias for outcome assessment)
Source of funding	Government: BMU (for the Institut fu"r Umweltmedizinische Forschung; FKZ 20462296) and the Federal Ministry for Education, Science, Research, and Technology (no. 01EG9705/2 and 01EG9732). The determination of specific IgE antibodies was financially supported by the Child Health Foundation (Stiftung Kindergesundheit).
Comments	

D.1.83 Nenna 2017

Nemia Zum			
Bibliographic reference	Nenna R, Cutrera R, Frass associated with bronchiol Disease 11(10), 393-401		
Study design	Case control study		
Objective	To examine whether exposul was associated with acute be		d outdoor pollutants
Setting/Study location	Italy		
Number of participants	416 infants		
Selected population	Yes (selected on case hosp	italised for bronchiolitis)	
Participant characteristics	Building characteristics: Location: Urban Dwelling type: Apartment Building age: Before 1990 Type of ownership/tenancy: Individual characteristics: Age (Median & range): Gender: Male (%) Smoker in home: Race	Cases Not reported 75.0% 63.9% Not reported 2 months; (0.5–12) 118 (55.4%) Not reported Not reported 213 (100%)	Controls Not reported 71.7% 53.0% Not reported 12 months (0.5–36) 116 (54.5%) Not reported Not reported O (0%)
Inclusion criteria	Inclusion criteria for cases were a diagnosis of bronchiolitis, without neonatal respiratory disorders or other chronic diseases. Inclusion criteria for controls were no respiratory diseases, and a medical history negative for lower respiratory tract diseases and neonatal respiratory disorders.		
Exclusion criteria	None reported		
Type of pollutant/exposure	Use of seed oil for cooking Number of cohabitants		

Bibliographic reference	Nenna R, Cutrera R, Frassanito A (2017) Modifiable risk factors associated with bronchiolitis. Therapeutic Advances in Respiratory Disease 11(10), 393-401	
Pollutant/exposure assessment	Self-reported questionnaire	
Outcome	Bronchiolitis - defined as the first episode of acute lower respiratory tract infection characterized by the presence of auscultator crackles, in infants aged ≤12 months	
Results	Use of seed oil for cooking Number of cohabitants ≥ 4	aOR (95%CI) 1.82 (1.206; 2.741) 1.748 (1.364; 2.132]
Follow up	Unclear	
Newcastle-Ottawa Scale	1.748 (1.364; 2.132]	
Source of funding	No funding reported (This research received no specific grant from any funding agency in the public, commercial or not-for-profit sectors).	
Comments	Not all pollutants measured reported	

D.1.84 Norback 2013

Bibliographic reference	Norback D, Zock J P, Plana E, et al (2013) Mould and dampness in dwelling places, and onset of asthma: The population-based cohort ECRHS. Occupational and Environmental Medicine 70(5), 325-331
Study design	Prospective cohort study
Objective	To investigate new onset of asthma in the ECRHS II in relation to self-reported as well as o2013 (observed building dampness and indoor moulds in the dwelling,

dwelling places, and onset of asth	nma: The populatio	n-based cohort
11 countries in Europe and two outside Europe (Melbourne in Australia and Portland in USA)		
7104 adults		
No		
Description Sex Age (years)- Ethnicity Education Annual family income	Not reported Not reported Not reported Not reported Not reported	
Not reported		
Not report		
Dampness and mould		
Questionnaire		
Asthma Bronchial hyper-responsiveness		
Adjusted risk ratios (aRRs) and 95% confidence intervals (CIs) for association between dampness, moulds, asthma and bronchial hyper-		
	Asthma	Asthma and BHR
	aRR (95%CI)	aRR (95%CI)
Any visible mould	1.15 (0.71, 1.85)	1.74 (0.68, 4.45)
Any damp spots	1.49 (1.00, 2.22)	1.88 (0.84, 4.22)
Reported window condensation in winter in any room	1.07 (0.75, 1.53)	1.43 (0.67, 3.07)
Between 5.9 and 11.7 years		
Selection Representativeness of the exposed cohort • truly representative of the average person in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • written self-report Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for smoking status • study controls for additional factor as follows - age, sex, and centre		
	dwelling places, and onset of asth ECRHS. Occupational and Enviror 11 countries in Europe and two outs Portland in USA) 7104 adults No Description Sex Age (years)- Ethnicity Education Annual family income Not reported Not report Dampness and mould Questionnaire Asthma Bronchial hyper-responsiveness Adjusted risk ratios (aRRs) and 95% association between dampness, moresponsiveness (BHR) Any visible mould Any damp spots Reported window condensation in winter in any room Between 5.9 and 11.7 years Selection Representativeness of the exposed • truly representative of the average Selection of the non-exposed cohort • drawn from the same community and Ascertainment of exposure • written self-report Demonstration that outcome of interesponsibility Comparability Comparability of cohorts on the basice study controls for smoking status	Portland in USA) 7104 adults No Description Sex Not reported Age (years)- Ethnicity Rot reported Annual family income Not reported Not

Bibliographic reference	Norback D, Zock J P, Plana E, et al (2013) Mould and dampness in dwelling places, and onset of asthma: The population-based cohort ECRHS. Occupational and Environmental Medicine 70(5), 325-331
	Assessment of outcome
	• self-report
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	 complete follow up - all subjects accounted for
	Overall level of bias – High (concerns over self-report of exposure and outcomes)
Source of funding	Government: European Commission
Comments	

D.1.85 O'Connor 2017

0 00111101 2017			
Bibliographic reference	O'Connor GT, Lynch SV, Bloomberg GR, et al (2017) Early-life home environment and risk of asthma among inner-city children. Journal of Allergy and Clinical Immunology, 141(6) 1468 - 75		
Study design	Prospective cohort study		
Objective	To examine exposures in the prenatal period and fir allergens and microbes in house dust, as potential ryears		
Setting/Study location	United States		
Number of participants	442 children		
Selected population	Yes – at risk of asthma		
Participant	Age	Not reported	
characteristics	Sex Male	226 (51%)	
	Race / ethnicity Black Hispanic Mixed / Other	318 (72%) 87 (20%) 37 (8%)	
	SES Reported as maternal education Less than high school High school More than high school	183 (42%) 151 (34%) 107 (24%)	
Inclusion criteria	Pregnant women aged 18 years or older a history of asthma, allergic rhinitis, or eczema, in the mother or father		
Exclusion criteria	Not reported		
Type of pollutant / exposure	House dust NO ₂		

Bibliographic reference	O'Connor GT, Lynch SV, Bloomberg GR, et al (2017) Early environment and risk of asthma among inner-city childre Allergy and Clinical Immunology , 141(6) 1468 - 75	
Pollutant / exposure assessment	Home visits to collect environmental data and specimens beg visits 3 months after birth and in the second and third years o house dust collection Indoor nitrogen dioxide concentration was measured during the production of the p	f life that included
Outcome	with a modified diffusion filter sampler Adjusted odds ratio and 95% confidence intervals	
Results	Exposure at 3 months Allergens (mg/g) House dust (Der f 1) per interquartile increase in exposure. Cat (Fel d 1) per interquartile increase in exposure. Dog (Can f 1) increase from the 25th to the 85th percentile. Exposure at 1 year Nitrogen dioxide (per interquartile increase in exposure.	Asthma at 7 years 0.98 (0.91, 1.04) 0.78 (0.62, 0.98) 0.62 (0.37, 1.03) 0.97 (0.75, 1.26)
Follow up	7 years	(**************************************
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • objective sampling Demonstration that outcome of interest was not present at state • Yes Comparability Comparability of cohorts on the basis of the design or analysite • study controls for maternal asthma • study controls for additional factors – sex and race Outcome Assessment of outcome • independent assessment Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall risk of bias: Low	·
Source of funding	Government: National Institute of Allergy and Infectious Dise Institutes of Health (NIH), National Center for Research Reso Center for the Advancement of Translational Research/NIH	
Comments		

D.1.86 Ostro 1993

Bibliographic reference Situdy design Objective Setting/Study location Number of participants Selected Population Characteristics Selected Population Participant Characteristics Age Sex Male Race / ethnicity White SES reported as educational level High school graduate degree 10.6 4-year college degree 10.6 4-year college degree 10.6 4-year college degree 9.0 Inclusion criteria Covina, or Azusa Exclusion criteria Type of pollutant / exposure assessment Outcome Adjust odds ratio and 95% confidence intervals for Respiratory illness (upper or lower) Follow up Risk of bias (Newcastle-Ottawa Scale) Follow up Risk of bias (Newcastle-Ottawa Scale) Find the participant on the same community as the exposed cohort juding for solution of exposure k, written self-report	JSHO 1333			
Objective To examine exposure to poor air quality and respiratory illness Setion/Study location Number of participants Selected No population Participant characteristics Age Not reported Sex Male Race / ethnicity White 89% SES reported as educational level High school graduate 2-year college degree 10.6 4-year college degree 17.1 Postgraduate degree 9.0 Inclusion criteria Covina, or Azusa Exclusion criteria Type of pollutant / exposure assessment Outcome Adjust odds ratio and 95% confidence intervals for Respiratory illness (upper or lower) Results Lower RTI aOR (95%CI) 1.23 (1.03, 1.47) 1.06 (0.94, 1.18) Follow up Gas stove Follow up Gas stove Cottana Scale) Follow up Gas stove Selection Representativeness of the exposed cohort i) truly representative of the average adult in the community Selection of the non-exposed cohort i) drawn from the same community as the exposed cohort Ascertainment of exposure		morbidity among adults i	n southern California. Ame	
Setting/Study location Number of participants Selected population Participant Age Not reported Sex Male Assec / ethnicity White SES reported as educational level High school graduate 42.4 2-year college degree 10.6 4-year college degree 9.0 Inclusion criteria Families who had at least one child in elementary school and resided in Glendora. Covina, or Azusa Exclusion criteria Gas stove Pollutant / exposure Adjust odds ratio and 95% confidence intervals for Respiratory illness (upper or lower) Results Lower RTI QNG (95%CI) Gas stove Follow up Risk of bias (Newcastle-Ottawa Scale) Representativeness of the exposed cohort i) truly representative of the average adult in the community Selection of the non-exposed cohort Ascertainment of exposure Ascertainment of exposure Setting/Stable Not reported Representativeness of the exposed cohort Ascertainment of exposure Ascertainment of exposure	Study design	Prospective cohort study		
location Number of participants Selected population Participant characteristics Age Sex Male Age Sex Male Age Sex Male Age Ses reported as educational level High school graduate 2-year college degree 10.6 4-year college degree 17.1 Postgraduate degree 9.0 Postgraduate 9.0 Postgraduate 9.0 Postgraduate 9.0 Postgraduate 9.0 Postgraduate 9.0 Postgraduate 9.0 Postgra	Objective	To examine exposure to po	oor air quality and respiratory	illness
participants Selected population Participant characteristics Age Sex Male Age Age Sex Male Age Sex Male Age Sex Male Age Age Sex Male Age Age Age Age Age Age Age Age Age Ag	•	United States		
Participant characteristics Age		321 adults		
characteristics Sex Male Race / ethnicity White SES reported as educational level High school graduate 2-year college degree 4-year college degree 4-year college degree 9.0 Inclusion criteria Exclusion criteria Type of pollutant / exposure assessment Outcome Adjust odds ratio and 95% confidence intervals for Respiratory illness (upper or lower) Results Lower RTI aOR (95%CI) Gas stove Follow up Risk of bias (Newcastle-Ottawa Scale) Representativeness of the exposed cohort j) truly representative of the average adult in the community Selection of the non-exposed cohort Ascertainment of exposure Ascertainment of exposure		No		
Male Race / ethnicity White 89% SES reported as educational level High school graduate 2-year college degree 4-year college degree 4-year college degree 9.0 Inclusion criteria Exclusion criteria Type of pollutant / exposure Pollutant / exposure assessment Outcome Adjust odds ratio and 95% confidence intervals for Respiratory illness (upper or lower) Results Lower RTI aOR (95%CI) 3OR (95%CI) 1.23 (1.03, 1.47) Follow up Risk of bias (Newcastle- Ottawa Scale) Representativeness of the exposed cohort J) drawn from the same community as the exposed cohort Ascertainment of exposure		Age		Not reported
White SES reported as educational level High school graduate 2-year college degree 4-year college degree 10.6 4-year college degree 9.0 Inclusion criteria Exclusion criteria Type of pollutant / exposure Pollutant / exposure assessment Outcome Adjust odds ratio and 95% confidence intervals for Respiratory illness (upper or lower) Results Lower RTI aOR (95%CI) Gas stove Follow up Risk of bias (Newcastle- Ottawa Scale) White SES reported as educational level 42.4 42.4 22-year college degree 10.6 47.1 9.0 9.0 Selmilies who had at least one child in elementary school and resided in Glendora, Covina, or Azusa Self-report Self-report Lower RTI aOR (95%CI) 1.23 (1.03, 1.47) 1.06 (0.94, 1.18) Follow up Risk of bias (Newcastle- Ottawa Scale) Ottawa Scale) In truly representativeness of the exposed cohort i) truly representative of the average adult in the community Selection of the non-exposed cohort j) drawn from the same community as the exposed cohort Ascertainment of exposure	characteristics			48%
High school graduate 2-year college degree 4-year college degree Postgraduate degree P		•		89%
Covina, or Azusa Exclusion criteria Type of pollutant / exposure Pollutant / exposure assessment Outcome Adjust odds ratio and 95% confidence intervals for Respiratory illness (upper or lower) Results Lower RTI Upper RTI aOR (95%CI) aOR (95%CI) Gas stove 1.23 (1.03, 1.47) 1.06 (0.94, 1.18) Follow up 6 months Risk of bias (Newcastle-Ottawa Scale) Ottawa Scale) Ottawa from the same community as the exposed cohort Ascertainment of exposure		High school graduate 2-year college degree 4-year college degree	al level	10.6 17.1
Type of pollutant / exposure Pollutant / exposure Pollutant / exposure assessment Outcome Adjust odds ratio and 95% confidence intervals for Respiratory illness (upper or lower) Results Lower RTI Upper RTI aOR (95%CI) aOR (95%CI) Gas stove 1.23 (1.03, 1.47) 1.06 (0.94, 1.18) Follow up 6 months Risk of bias (Newcastle-Ottawa Scale) Ottawa Scale) Selection Representative of the average adult in the community Selection of the non-exposed cohort j) drawn from the same community as the exposed cohort Ascertainment of exposure		Families who had at least one child in elementary school and resided in Glendora,		
pollutant / exposure Pollutant / exposure assessment Outcome Adjust odds ratio and 95% confidence intervals for Respiratory illness (upper or lower) Results Lower RTI Upper RTI aOR (95%CI) aOR (95%CI) Gas stove 1.23 (1.03, 1.47) 1.06 (0.94, 1.18) Follow up 6 months Risk of bias (Newcastle-Ottawa Scale) Ottawa Scale) Representativeness of the exposed cohort i) truly representative of the average adult in the community Selection of the non-exposed cohort j) drawn from the same community as the exposed cohort Ascertainment of exposure		Not reported		
exposure assessment Outcome Adjust odds ratio and 95% confidence intervals for Respiratory illness (upper or lower) Results Lower RTI Upper RTI aOR (95%CI) aOR (95%CI) Gas stove 1.23 (1.03, 1.47) 1.06 (0.94, 1.18) Follow up 6 months Risk of bias (Newcastle-Ottawa Scale) Representativeness of the exposed cohort i) truly representative of the average adult in the community Selection of the non-exposed cohort j) drawn from the same community as the exposed cohort Ascertainment of exposure	pollutant /	Gas stove		
Results Lower RTI aOR (95%CI) Gas stove 1.23 (1.03, 1.47) Follow up 6 months Risk of bias (Newcastle-Ottawa Scale) Ottawa Scale) Representativeness of the exposed cohort i) truly representative of the average adult in the community Selection of the non-exposed cohort j) drawn from the same community as the exposed cohort Ascertainment of exposure	exposure	Self-report		
Follow up Risk of bias (Newcastle-Ottawa Scale) Gas stove Accertainment of exposure aOR (95%CI) 1.23 (1.03, 1.47) 1.06 (0.94, 1.18) 1.06 (0.94, 1.18) ACR (95%CI) 1.06 (0.94, 1.18) 1.06 (0.94, 1.18)	Outcome		confidence intervals for Resp	piratory illness (upper or
Risk of bias (Newcastle- Ottawa Scale) Selection Representativeness of the exposed cohort i) truly representative of the average adult in the community Selection of the non-exposed cohort j) drawn from the same community as the exposed cohort Ascertainment of exposure	Results	Gas stove	aOR (95%CI)	aOR (95%CI)
(Newcastle-Ottawa Scale) Representativeness of the exposed cohort i) truly representative of the average adult in the community Selection of the non-exposed cohort j) drawn from the same community as the exposed cohort Ascertainment of exposure	Follow up	6 months		
	(Newcastle-	Representativeness of the exposed cohort i) truly representative of the average adult in the community Selection of the non-exposed cohort j) drawn from the same community as the exposed cohort Ascertainment of exposure		

Bibliographic reference	Ostro B D, Lipsett M J, Mann J K, et al (1993) Air pollution and respiratory morbidity among adults in southern California. American journal of epidemiology 137(7), 691-700
	Demonstration that outcome of interest was not present at start of study
	I) Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	m) study controls for presence of chronic respiratory disease.
	n) study controls for additional factor - sex
	Outcome
	Assessment of outcome
	o) self-report – daily diary
	Was follow-up long enough for outcomes to occur
	p) Yes
	Adequacy of follow up of cohorts
	q) complete follow up - all subjects accounted for
	Overall risk of bias: Low
Source of funding	Not reported
Comments	

D.1.87 Pettigrew 2004

Bibliographic reference	Pettigrew M M, Gent J F, Triche E W, Belanger K O, Bracken M B, and Leaderer B P (2004) Association of early-onset otitis media in infants and exposure to household mould. Paediatric and Perinatal Epidemiology 18(6), 441-447		
Study design	Prospective cohort study		
Objective	To examine the relationship between levels of h media among a cohort of infants at high risk for		
Setting/Study location	United States		
Number of participants	1002		
Selected population	Yes – at high risk of asthma		
Participant characteristics	Description Sex o Male o Female Age (years)- Mean (SD) Maternal Ethnicity o White o Black o Hispanic Education o <high .="" college<="" diploma="" high="" o="" school="" some="" td=""><td>398 408 Not reported 533 103 170 98 419</td></high>	398 408 Not reported 533 103 170 98 419	

Bibliographic reference	Pettigrew M M, Gent J F, Triche E Leaderer B P (2004) Association and exposure to household moul Epidemiology 18(6), 441-447	of early-ons	et otitis media in infants
	o College / Higher		289
Inclusion criteria	Women with at least one other child	d with physic	ian-diagnosed asthma
Exclusion criteria	Not reported		
Type of pollutant/exposure	Mould		
Pollutant/exposure assessment	Objective sampling - airborne mould samples were collected from the main living area of the home Fungi were identified to the genus level and recorded in colony forming units (CFU) per cubic metre		
Outcome	First episode of otitis media <6 mor	nths of age	
Results		aOR ((95%	6CI)
	Mould	1.37 (0.94,	, 2.02)
	Penicillium Undetectable 0 CFU/m³ Low 1–499 CFU/m³ Medium 500–999 CFU/m³ High ≥1000 CFU/m³ Cladosporium Undetectable 0 CFU/m³ Low 1–499 CFU/m³ Medium 500–999 CFU/m³ High ≥1000 CFU/m³ 'Other' mould Undetectable 0 CFU/m³ Low 1–499 CFU/m³	Reference 0.75 [0.52, 1.89 [0.67, 1.27 [0.56, Reference 1.04 [0.70, 0.92 [0.48, 1.09 [0.52, Reference 1.21 [0.84,	1.08] 5.30] 2.86] 1.56] 1.79] 2.29]
	Medium 500–999 CFU/m ³	0.72 [0.29, 3.45 [1.36,	-
	High ≥1000 CFU/m ³		
Follow up	6 months		
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • selected group of at risk children Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • Objective sampling Demonstration that outcome of interest was not present at start of study • Yes Comparability		

Bibliographic reference	Pettigrew M M, Gent J F, Triche E W, Belanger K O, Bracken M B, and Leaderer B P (2004) Association of early-onset otitis media in infants and exposure to household mould. Paediatric and Perinatal Epidemiology 18(6), 441-447	
	Comparability of cohorts on the basis of the design or analysis	
	study controls for smoke exposure	
	• study controls for additional factors as follows , ethnicity.)	
	Outcome	
	Assessment of outcome	
	• self-report	
	Was follow-up long enough for outcomes to occur	
	• Yes	
	Adequacy of follow up of cohorts	
	$ullet$ complete follow up - all subjects accounted for \Box	
	Overall level of bias – Moderate (concerns over self-report of outcomes)	
Source of funding	Government: National Institute of Environmental Health Sciences.	
Comments		

D.1.88 Pettigrew 2004 b

Bibliographic reference	Pettigrew MM, Gent JF, Triche EW, et al (2004) Infant otitis media and the use of secondary heating sources. Epidemiology (Cambridge, and Mass.) 15(1), 13-20	
Study design	Prospective cohort study	
Objective	To assess the effect of environmental exposures from secondary home heating sources on otitis media and recurrent otitis media on infants in the first year of life.	
Setting/Study location	United States	
Number of participants	813 infants	
Selected population	No	
Participant	Description	No. (%)
characteristics	Sex	
	Male	(52%)
	female	(48%)
	Maternal age (years)	Not reported
	Ethnicity	Not reported
	Maternal asthma and/or atopic	80 (9%)
	SES	Not reported
	Annual family income	Not reported
	Building characteristics	Not reported
Inclusion criteria	Mothers who were delivering babies at 7 hospitals in Connecticut and 5 hospitals in Virginia between 1993 and 1996.	

Bibliographic reference		the EW, et al (2004) Infant otitis media and the ources. Epidemiology (Cambridge, and Mass.)
Exclusion criteria	Smoking in the household	
Type of pollutant/exposu re	Secondary heating sources Air conditioning Pets Mould	
Pollutant/expos ure assessment	Interviews	
Outcome	Clinical diagnosis of otitis med Recurrent otitis media defined by at least 21 days) in one ye	d as 4 or more episodes of otitis media (separated
Results	Adjusted odds ratios (aORs) a	and 95% confidence intervals (CIs)
	Any exposure	Recurrent otitis media
	Fireplace Wood stove Kerosene heater Air conditioning Reported mould Cat or dog Any daily use Fireplace Wood stove Kerosene heater Air conditioning Reported mould	0.99 (0.58, 1.72) 1.22 (0.66, 2.23) 0.94 (0.50, 1.78) 0.52 (0.27, 1.03) 1.15 (0.67, 1.99) 0.76 (0.47, 1.26) Episode of otitis media 1.14 (0.90, 1.45) 1.08 (0.85, 1.38) 0.91 (0.67, 1.25) 0.93 (0.77, 1.11) 1.05 (0.88, 1.26)
	Cat or dog	1.06 (0.90, 1.25)
Follow up	12 months	
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort r) truly representative of the average infant Selection of the non-exposed cohort s) drawn from the same community as the exposed cohort Ascertainment of exposure t) written self-report Demonstration that outcome of interest was not present at start of study u) Yes Comparability Comparability of cohorts on the basis of the design or analysis v) study controls for gas stove use w) study controls for additional factors as follows - number of children in household, multifamily dwelling, history of allergies, education, race, state of residence Outcome	

Bibliographic reference	Pettigrew MM, Gent JF, Triche EW, et al (2004) Infant otitis media and the use of secondary heating sources. Epidemiology (Cambridge, and Mass.) 15(1), 13-20
	Assessment of outcome
	x) self-report (maternal)
	Was follow-up long enough for outcomes to occur
	y) Yes
	Adequacy of follow up of cohorts
	z) complete follow up - all subjects accounted for \square
	Overall level of bias – Low
Source of funding	Government: National Institute of Environmental Health Sciences
Comments	

D.1.89 Pindus 2016

Bibliographic reference	Pindus M, Orru H, Maasikmets M, et al (2016) Association between health symptoms and particulate matter from traffic and residential heating - Results from RHINE III in Tartu. Open Respiratory Medicine Journal 10, 58-69		
Study design	Prospective cohort study		
Objective	To investigate potential effects of traffic particles on respiratory and cardiac hea		
Setting/Study location	Estonia		
Number of participants	905		
Selected population	No		
Participant characteristics	Description Gender Male Female Age years (mean) Ethnicity Education Basic Secondary Higher SES Building characteristics	362 (40.0%) 543 (60.0%) 50 Not reported 46 (5.2%) 454 (51.4%) 383 (43.4%) Not reported Not reported	
Inclusion criteria	Not reported	1 - 1 - 2 - 2	
Exclusion criteria	Not reported		

	Pindus M, Orru H, Maasikmets M, et al (2016) Association between health symptoms and particulate matter from traffic and residential		
Bibliographic reference	heating - Results from RHINE III in Tartu. Open Respiratory Medicine Journal 10, 58-69		
Type of pollutant/exposure	Particulate matter (PM) from traffic and residential heating		
Pollutant/exposure assessment	Concentrations of $PM_{2.5}$ and PM_{10} for the years 2009-2012 were calculated for grid size of 100x100 m across Tartu using a Eulerian air quality dispersion model part of the AirViro Air Quality Management System.		
	Household $PM_{2.5}$ emissions (g/s) were calculated according to the size (m2 of each's homes heated area		
Outcome			
Results	Adjusted odds ratios (aORs) and 98 association between PM, respirator		
		Residential heating induced PM _{2.5}	
		aOR (95%CI)	
	Cough	0.95 (0.72, 1.29)	
	Wheeze without cold	1.14 (0.75, 1.73)	
	Asthma	1.16 (0.60, 2.19)	
	Allergic rhinitis	0.63 (0.42, 0.94)	
	Breathlessness	0.97 (0.64, 1.48)	
	Chest tightness	1.05 (0.72, 1.51)	
	Cardiac disease	0.92 (0.60, 1.39)	
	Hypertension	0.78 (0.54, 1.12)	
	Stroke	0.85 (0.27, 2.71)	
	Heart infarction or angina pectoris	0.67 (0.28, 1.56)	
Follow up	12 months	(0.20, 1.00)	
Risk of bias	Selection		
(Newcastle-Ottawa	Representativeness of the exposed	cohort	
Scale)	truly representative of the average		
	Selection of the non-exposed cohort		
	drawn from the same community as the exposed cohort		
	Ascertainment of exposure		
	Modelled exposure		
	Demonstration that outcome of interest was not present at start of study		
	• Yes		
	Comparability of cohorts on the basis of the design or analysis		
	Comparability of cohorts on the basis of the design or analysis • study controls for ETS (environmental tobacco smoke) at home		
	 study controls for ETS (environmental tobacco smoke) at nome study controls for any additional factors - gender, age, body mass index (BMI), education level, and smoking history 		
	Outcome		
Assessment of outcome			
	• self-report		
	Was follow-up long enough for outcomes to occur		

Bibliographic reference	Pindus M, Orru H, Maasikmets M, et al (2016) Association between health symptoms and particulate matter from traffic and residential heating - Results from RHINE III in Tartu. Open Respiratory Medicine Journal 10, 58-69
	 Yes Adequacy of follow up of cohorts complete follow up - all subjects accounted for □ Overall risk of bias: Moderate (concerns over self-report of outcomes)
Source of funding	Government: The Estonian Ministry of Education and Research Charity: The Estonian Science Foundation.
Comments	

D.1.90 Ponsonby 2001

Bibliographic reference Study design Objective	Ponsonby A L, Dwyer T, Kemp A et.al (2001). A prospective study of the association between home gas appliance use during infancy and subsequent dust mite sensitization and lung function in childhood. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology, 31(10), pp.1544-52. Prospective cohort study To examine the relationship between domestic gas appliance, use during infancy and childhood and the development of house dust mite (HDM) sensitization and asthma			
Setting/Study location	Australia			
Number of participants	456 children			
Selected population	No			
Participant characteristics	Description	No.	%	
	Sex	Not reported	Not reported	
	Maternal age (years)	Not reported	Not reported	
	Ethnicity	Not reported	Not reported	
	(Maintenance) medication use	Not reported	Not reported	
	Maternal asthma and/or atopic	Not reported	Not reported	
	Parental education	Not reported	Not reported	
	Annual family income	Not reported	Not reported	
	Building characteristics	Not reported	Not reported	
Inclusion criteria	Multiple births			
Exclusion criteria	Not reported			
Type of pollutant/exposure	NO ₂ from home gas appliance House dust mite (HDM)			

Bibliographic reference	Ponsonby A L, Dwyer T, Kemp A et.al (2001). A prospective study of the association between home gas appliance use during infancy and subsequent dust mite sensitization and lung function in childhood. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology, 31(10), pp.1544-52.			
Pollutant/exposure assessment	Home gas appliance use defined as the positive report of gas cooking or gas heater use in the living room. Skin prick testing (SPT) was used to assess the cutaneous reaction to exposure to house dust mites. Weal allergen reactions of 3 mm or greater at 15 minutes were classified as positive			
Outcome	HDM sensitisation Asthma			
Results	Adjusted relative risk (aRR) and 95% confidence intervals (CIs) for association between home gas cooking, home gas appliance use, asthma and HDM sensitisation during infancy			
		Asthma		
		RR (95%CI)		
	Home gas cooking	Not reported		
	Gas heaters	Not reported		
	Home gas appliance use	1.30 (0.74, 2.29)		
	Adjusted relative risk (aRR) and 95% confidence intervals (CIs) for association between asthma and HDM sensitisation during infancy			
		Asthma		
		RR (95%CI)		
	HDM sensitisation	1.65 (1.32, 2.06)		
Follow up	8 years	, , , , , , , , , , , , , , , , , , ,		
Risk of bias (Newcastle-Ottawa Scale)				

Bibliographic reference	Ponsonby A L, Dwyer T, Kemp A et.al (2001). A prospective study of the association between home gas appliance use during infancy and subsequent dust mite sensitization and lung function in childhood. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology, 31(10), pp.1544-52.
	Adequacy of follow up of cohorts
	$ullet$ complete follow up - all subjects accounted for \square
	Overall level of bias – Moderate (concerns over self-report of exposure)
Source of funding	Charity: Asthma Foundation of Tasmania for equipment loan.
	Government: The Tasmanian Infant Health Survey was supported by the US National Institutes of Health Grant, the Tasmanian State Government, the Australian Rotatory Health Research Fund, the National Health and Medical Research Council of Australia, the National Sudden Infant Death Syndrome Council of Australia, Health, Zonta International Industry: The Tasmanian Infant Health Survey was supported by Wyeth Pharmaceuticals
Comments	Authors suggest that that indoor gas appliance use was associated with an increased risk of allergic sensitisation in children. Study survey reported the use of bottled gas for heaters of portable or fixed type. Thus, results pertain to gas combustion heaters rather than modern ducted gas central heating

D.1.91 Power 2015

OWCI Z010			
Power MC, Kioumourtzoglou M-A, Hart JaE, et al (2015) The relation between past exposure to fine particulate air pollution and prevalent anxiety: observational cohort study. BMJ (Clinical research ed.) 350, h1111			
Prospective cohort study			
	To determine whether higher past exposure to particulate air pollution is associated with prevalent high symptoms of anxiety.		
United States			
71 271 women			
Description	No.	%	
Sex (female) sex	All female	All female	
Age (years); mean (SD)	Not reported	Not reported	
Ethnicity	Not reported	Not reported	
Cases/selected population	Not reported	Not reported	
Socio-economic status (educ	cation)		
Registered nurse	44 907	63.0	
Bachelor's degree	13 368	18.8	
Master's degree or PhD	6607	9.3	
Missing	6389	9.0	
	between past exposure to anxiety: observational coherospective cohort study To determine whether higher associated with prevalent higher associated higher associa	between past exposure to fine particulate air pollular anxiety: observational cohort study. BMJ (Clinical Prospective cohort study To determine whether higher past exposure to particulassociated with prevalent high symptoms of anxiety. United States 71 271 women Description No. Sex (female) sex All female Age (years); mean (SD) Ethnicity Cases/selected population Not reported Socio-economic status (education) Registered nurse 44 907 Bachelor's degree 13 368 Master's degree or PhD 6607	

Bibliographic	Power MC, Kioumourtzoglou M-A, Har between past exposure to fine particul	late air pollution and prevalent	
reference	anxiety: observational cohort study. B Building characteristics Not reporte		
Inclusion	Not reported	Not reported	
criteria			
Exclusion criteria	Not reported		
Type of pollutant/expos ure	Proximity to traffic and particulate matter (PM)		
Pollutant/expos ure assessment	Using geographic information software (A computed distance from the residential awith a street level geocoding match to the A1 (limited access to primary roads with a travel, that is, interstate highways), A2 (p and major roads without access restriction typically with more than two lanes) roads. Authors used spatiotemporal prediction in exposure to particulate matter <10 µm (P matter) in aerodynamic diameter at the relevel geocoding match for each participare each participant.	ddress of each participant up to 500 m, e nearest US census feature class code defined exits and divided directions of rimary major, non-interstate highways ns), or A3 (smaller, secondary roads, segment. nodels yielding monthly estimates of PM) and <2.5 µm (PM or fine particulate esidential address with at least a zip code	
Outcome	High symptoms of anxiety		
Results	Adjusted odds ratios (aORs) and 95% co between PM and high symptoms of anxie		
		High symptoms of anxiety	
		aOR (95%CI)	
	≤ 50m from motorway	1.01,(0.95, 1.08)	
Follow up	Not reported		
Study methods	The Crown-Crisp index phobic anxiety scale, one of six scales from the Crown-Crisp experiential index, is a measure of anxiety symptom levels and was included in the questionnaire. This scale has been shown to differentiate between people with general anxiety or phobias from those with other psychiatric conditions and healthy comparison participants and has been used in population based research. Authors used logistic regression models to estimate the association between each exposure and high anxiety symptoms (Crown-Crisp index phobic anxiety scale score ≥6) and adjusted for possible confounders		
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average female population in the community Selection of the non-exposed cohort • no description of the derivation of the non-exposed cohort Ascertainment of exposure • validated measurements used Demonstration that outcome of interest was not present at start of study • Yes Comparability		
	- 1,		

Bibliographic reference	Power MC, Kioumourtzoglou M-A, Hart JaE, et al (2015) The relation between past exposure to fine particulate air pollution and prevalent anxiety: observational cohort study. BMJ (Clinical research ed.) 350, h1111			
	Comparability of cohorts on the basis of the design or analysis			
	• study controls for socioeconomic status, education, husband's education, age, employment status, physical activity			
	Outcome			
	Assessment of outcome			
	validated anxiety scale used			
	Was follow-up long enough for outcomes to occur			
	• Yes			
	Adequacy of follow up of cohorts			
	subjects lost to follow up unlikely to introduce bias			
	Overall risk of bias: low			
Source of funding	Government: National Institute of Environmental Health Sciences, National Institute of Aging			
Comments				

D.1.92 Puett 2014

Bibliographic reference	Puett RC, Hart JE, Yanosky JD, et al (2014) Particulate matter air pollution exposure, distance to road, and incident lung cancer in the nurses' health study cohort. Environmental health perspectives 122(9), 926-32		
Study design	Prospective cohort study		
Objective	To examine the relation of lung cancer incidence with long-term residential exposures to ambient particulate matter and residential distance to roadway, as a proxy for traffic related exposures		
Setting/Study location	United States		
Number of participants	103,650 women		
Selected population	No		
Participant	Description	No.	%
characteristics	Sex (female) sex	All female	All female
	Age (years); mean (SD)	67.0 (8.3)	-
	Ethnicity	Not reported	Not reported
	Cases/selected population	Not reported	Not reported
	Socio-economic status (education)	Not reported	Not reported
	Building characteristics	Not reported	Not reported
Inclusion criteria	Not reported		
Exclusion criteria	Previous diagnosis of cancer (except for non-melanoma skin cancer) before follow-up Did not have information for the exposures of interest		

Bibliographic reference	Puett RC, Hart JE, Yanosky JD, et al (2014) Particulate matter air pollution exposure, distance to road, and incident lung cancer in the nurses' health study cohort. Environmental health perspectives 122(9), 926-32			
Type of pollutant/expos ure	Proximity to major road			
Pollutant/expos ure assessment	Authors calculated distance to road at each address as a proxy for traffic related exposures. Distance to the nearest road (meters) was determined using geographic information system (GIS) software (ArcGIS, version 9.3; ESRI, Redlands, CA) and the ESRI Street map Pro2007 data set. Authors calculated the shortest distances to the following road classes as defined by the U.S. Census Bureau (2001): A1 (primary roads, typically interstate highways, with limited access, division between the opposing directions of traffic, and defined exits), A2 (primary major, non-interstate highways and major roads without access restrictions), and A3 (smaller, secondary roads, usually with more than two lanes). Prediction models were used to determine PM surfaces for each month and each PM size fraction			
Outcome	Lung cancer incidence			
Results	Adjusted hazard ratios (aHRs) and 95% of between proximity to major road, PM and			
		Lung cancer incidence		
		aHR (95%CI)		
	Residential proximity to a major road (metres)			
	≥200	Reference		
	50 – 199	0.73 (0.51, 1.04)		
	0 – 49	2.01 (1.06, 3.80)		
Follow up	•	2.0 . (
Study methods	Lung cancers were self-reported by the participants or next of kin or were identified from death certificates; and first reports were subsequently confirmed with medical records by physicians blinded to exposure status. However, because lung cancers were well reported in this cohort, we included any primary report reconfirmed by the participant where pathological reports were not available. Time-varying Cox proportional hazards models were used to assess the relationship of incident lung cancer with residential distance to road and exposure to PM _{2.5} , PM ₁₀ , or PM _{2.5} –10 adjusting for possible confounders			
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average female population in the community Selection of the non-exposed cohort • no description of the derivation of the non-exposed cohort Ascertainment of exposure • validated measurements used Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for age, time period, geographic region, BMI, alcohol consumption, physical activity, overall diet quality, smoking status Outcome			

Bibliographic reference	Puett RC, Hart JE, Yanosky JD, et al (2014) Particulate matter air pollution exposure, distance to road, and incident lung cancer in the nurses' health study cohort. Environmental health perspectives 122(9), 926-32
	Assessment of outcome
	record linkage
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	subjects lost to follow up unlikely to introduce bias
	overall risk of bias: low
Source of funding	Government: National Institutes of Health
Comments	

D.1.93 Pujades-Rodriguez 2009

Bibliographic reference	Pujades-Rodriguez M, McKeever T, Lewis S et.al (2009) Effect of traffic pollution on respiratory and allergic disease in adults: cross-sectional and longitudinal analyses. BMC pulmonary medicine 9, 42				
Study design	Prospective coh	Prospective cohort study			
Objective	To determine eff adults	ect of traffic pol	lution on respira	atory and allergio	disease in
Setting/Study location	United Kingdom				
Number of participants	2644 adults				
Selected population	No	No			
Participant characteristics		Cross-sectional analysis (N=2599)		Longitudinal analysis (N=1329)	
	Description	No.	%	No.	%
	Sex (female)	1300	50.0	670	50.4
	Age (years); mean (SD)	Not reported	Not reported	Not reported	Not reported
	Ethnicity	Not reported	Not reported	Not reported	Not reported
	Cases/selecte d population	Not reported	Not reported	Not reported	Not reported
	Socio- economic status (education)	Not reported	Not reported	Not reported	Not reported
	Building characteristics	Not reported	Not reported	Not reported	Not reported
Inclusion criteria	Not reported				
Exclusion criteria	Not reported				
Type of pollutant/exposure	Proximity to traffic and traffic related NO ₂				

	Pujades-Rodriguez M, McKeever T, Lewis S et.al (2009) Effect of traffic				
Bibliographic reference	pollution on respiratory and allergic disease in adults: cross-sectional and longitudinal analyses. BMC pulmonary medicine 9, 42				
Pollutant/exposure assessment	Authors calculated the shortest distance (in metres) between each address location and the nearest major road, defined as a motorway (freeway), or 'A' or 'B' class road (principal road as classified by UK Department for Transport), using Geographical Information System (GIS) software (ArcGIS 9.0). To compute our modelled NO ₂ variable, we linked each home location grid reference to a high resolution map of modelled traffic-related NO ₂ using ArcGIS.				
Outcome	Respiratory and	allergic outcor	mes		
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between residential proximity to a main road and respiratory and allergic outcomes				
		Wheezing in the last year	COPD	Bronchial hyper responsiveness	Allergic sensitisation
		aOR (95%CI)	aOR (95%CI)	aOR (95%CI)	aOR (95%CI)
	≤150 m	0.86 (0.68, 1.08)	0.97 (0.68, 1.37)	0.92 (0.68, 1.24)	0.87 (0.70, 1.07)
	>150 m	1.00	1.00	1.00	1.00
	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between modelled NO ₂ level and respiratory and allergic outcomes				
	<33.92	1	1	1	1
	33.92 – 34.23	1.03 (0.76, 1.39)	1.09 (0.68, 1.73)	1.08 (0.73, 1.60)	0.98 (0.74, 1.30)
	34.23 – 34.73	0.86 (0.63, 1.16)	0.95 (0.60, 1.52)	0.95 (0.64, 1.41)	1.02 (0.77, 1.35)
	34.73 – 36.79	0.84 (0.63, 1.14)	0.91 (0.57, 1.45)	1.03 (0.70, 1.54)	0.97 (0.73, 1.28)
	>36.79	0.88 (0.66, 1.19)	1.07 (0.68, 1.68)	0.81 (0.54, 1.21)	0.94 (0.72, 1.24)
Follow up					
Study methods	Respiratory outcomes were self-reported from questionnaires. Allergen skin sensitisation, defined as a response to any of the allergens tested at least 3 mm greater than the saline control response in the presence of a positive histamine control; and high total IgE, defined as a concentration above 100 kU/l. Multiple logistic regression analyses were carried out to assess the effect of distance and modelled NO ₂ level on each outcome, adjusting for possible confounders.				
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average population in the community				

Bibliographic reference	Pujades-Rodriguez M, McKeever T, Lewis S et.al (2009) Effect of traffic pollution on respiratory and allergic disease in adults: cross-sectional and longitudinal analyses. BMC pulmonary medicine 9, 42
	Selection of the non-exposed cohort
	 no description of the derivation of the non-exposed cohort
	Ascertainment of exposure
	validated measurement used
	Demonstration that outcome of interest was not present at start of study • Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	• study controls for age, sex, smoking status and deprivation score
	Outcome
	Assessment of outcome
	clinical investigation
	• self-report
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	subjects lost to follow up unlikely to introduce bias
	Overall risk of bias: Moderate: response bias from self-reported respiratory outcomes
Source of funding	Charity: British Lung Foundation, Asthma UK.
	Academic: The Institute of Clinical Research (University of Nottingham)
Comments	

D.1.94 Raaschou-Nielsen 2010

itaasciiou-iticiscii			
Bibliographic reference	Raaschou-Nielsen O, Hermansen M N, Loland L, et al (2010) Long-term exposure to indoor air pollution and wheezing symptoms in infants. Indoor Air 20(2), 159-167		
Study design	Prospective cohort study		
Objective	To study whether air pollutants alter underlying bronchial hyperresponsiveness, in addition to its previously demonstrated role as a trigger of symptoms		
Setting/Study location	Denmark		
Number of participants	411		
Selected population	Yes – infants at risk of asthma		
Participant characteristics	Description Sex Not reported Age (months); Ethnicity Not reported Socio-economic status Building characteristics Not reported Not reported		

Bibliographic reference		mansen M N, Loland L, et al (2010) Long-term llution and wheezing symptoms in infants.			
Inclusion criteria	Infants born to mothers with	n asthma			
Exclusion criteria	Not reported	Not reported			
Type of pollutant/exposure	PM _{2.5} NO ₂ Formaldehyde	NO_2			
Pollutant/exposure assessment	NO ₂ , and formaldehyde were measured in the children's bedrooms away from windows and doors, preferably at about 1.5 m above the floor. Measurements were performed up to three times during the first 18 months of life, for 10 weeks on each occasion. NO ₂ , and formaldehyde samplers were given to the parents, with comprehensive instructions on how to start and stop the measurements. PM _{2.5} was measured over 1-week periods at the same location in the children's bedrooms. Trained personnel initiated and concluded each measurement				
Outcome	Wheezing				
Results	Adjusted odds ratios (aORs	s) and 95% confidence intervals (CIs)			
	PM _{2.5} (μg/m³) Q1 (<10.6) Q2 (10.6–13.2) Q3 (13.2–16.8) Q4 (16.8–24.1) Q5 (>24.1) NO ₂ (μg/m³) Q1 (<5.2) Q2 (5.2–6.8) Q3 (6.8–8.6) Q4 (8.6–11.7) Q5 (>11.7) Formaldehyde (μg/m³) Q1 (<12.4) Q2 (12.4–16.3) Q3 (16.3–20.3) Q4 (20.3–25.6) Q5 (>25.6)	aOR (95%CI) Reference 1.32 (0.53, 3.27) 1.74 (0.67, 4.47) 0.67 (0.28, 1.59) 1.02 (0.41, t2.57) Reference 0.66 (0.27, 1.61) 0.80 (0.32, 2.01) 1.15 (0.40, 3.32) 0.43 (0.15, 1.18) Reference 1.11 (0.47, 2.63) 1.21 (0.51, 2.92) 1.40 (0.57, 3.47) 0.67 (0.29, 1.54)			
Follow up	18 months				
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the population in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort				

Bibliographic reference	Raaschou-Nielsen O, Hermansen M N, Loland L, et al (2010) Long-term exposure to indoor air pollution and wheezing symptoms in infants. Indoor Air 20(2), 159-167
	Ascertainment of exposure Objective measurement Demonstration that outcome of interest was not present at start of study Yes Comparability Comparability of cohorts on the basis of the design or analysis Study controls for sex, area of residence, education of mother and (log-transformed) baseline lung function. Outcome Assessment of outcome Self-report (parent) Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts subjects lost to follow up unlikely to introduce bias Overall risk of bias: Moderate (concern over self-report of outcomes)
Source of funding	Government: The Danish Ministry of the Interior and Health Research Centre for Environmental Medicine. Charity: the Pharmacy Foundation of 1991; the Lundbeck Foundation; The Augustino Foundation; Ronald McDonald House Charities; the Danish Medical Research Council; The Danish Pediatric Asthma Center; Direktør, cand.pharm. K. Gad Andersen og Hustrus Familiefond; Aage Bangs Fond; Danish Lung Association; Kai Lange og Gunhild Kai Langes Fond; Direktør Ib Henriksens Fond; Gerda og Aage Henschs Fond; Rosalie Petersens Fond; Hans og Nora Buchards Fond; Dagmar Marshalls Fond; Foundation of Queen Louise¢ Children Hospital; the Danish Hospital Foundation for Medical Research, Region of Copenhagen, the Faroe Island, and Greenland; Gangsted Fond; Højmosega° rd-Legatet; Fonden til Lægevidenskabens Fremme; A.P. Møller og Hustru Chastine Mc-Kinney Møllers Fond til almene Formaal; Industry: AstraZenaca; LEOpharma; Pharmacia-Pfizer and Yamanouchi Pharma.
Comments	

D.1.95 Rice 2015

	Rice MB, Ljungman PL, Wilker EH, et al (2015) Long-term exposure to traffic emissions and fine particulate matter and lung function decline
Bibliographic reference	in the Framingham heart study. American journal of respiratory and critical care medicine 191(6), 656-64
Study design	Prospective cohort study

Bibliographic reference	Rice MB, Ljungman traffic emissions ar in the Framingham critical care medici	nd fine heart	particulate study. Amer	matter and l	ung fu	nction decline	
Objective	To determine if expo changes in lung fund					with longitudinal	
Setting/Study location	United States						
Number of participants	6,339						
Participant	Description		No.		%		
characteristics	Sex (male) sex		2700		42.6		
	Age (years); mean (S	SD)	50.4 (12.4)		-		
	Ethnicity		Not reported	t	Not re	eported	
	Cases/selected population		Not reported	d	Not re	eported	
	Socio-economic state (education)	us					
	<high school<="" td=""><td></td><td>114</td><td></td><td>1.8</td><td></td></high>		114		1.8		
	High school		1179		18.6		
	Some college		1807		28.5		
	College graduate school		3157		49.8		
	Missing education		82		1.3		
	Building characteristics		Not reported	t	Not re	Not reported	
Inclusion criteria	Not reported						
Exclusion criteria	Not reported						
Type of pollutant/exposure	Proximity to traffic ar	nd PM ₂	2.5				
Pollutant/exposure assessment	Distance to major roa home address at the road (U.S. Census F	time o	f the examin				
	Daily estimates of $PM_{2.5}$ at home address were derived from a model using moderate resolution imaging spectroradiometer satellite-derived aerosol optical thickness measurements at a 10310-km spatial resolution across the Northeast and then resolved to a specific location within a 50350-m grid using land-use terms.				rived aerosol lution across the		
Outcome							
Results		Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between proximity to traffic, PM, respiratory and cardiac conditions					
		Asthma	a	Wheeze in F 12 month	Past	Chronic Cough (>3 mo/yr)	
		OR (95	5%CI)	OR (95%CI))	OR (95%CI)	
	Distance to roadway	•	•	` `		•	
			0.95, 1.46)	1.02 (0.84,	1.25)	1.22 (0.89, 1.66)	
	100 to <200	1.35 (1	.06, 1.72)	0.89 (0.70,	-	0.89 (0.61, 1.30)	

Bibliographic reference	Rice MB, Ljungman PL, Wilker EH, et al (2015) Long-term exposure to traffic emissions and fine particulate matter and lung function decline in the Framingham heart study. American journal of respiratory and critical care medicine 191(6), 656-64
	200 to <400 1.26 (1.01, 1.58) 0.94 (0.76, 1.16) 1.17 (0.84, 1.63)
Follow up	16 years
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average population in the community Selection of the non-exposed cohort • no description of the derivation of the non-exposed cohort Ascertainment of exposure • validated measurements used Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for sex, age, height, weight, education, median household income Outcome Assessment of outcome • questionnaires • self-report no description Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • subjects lost to follow up unlikely to introduce bias overall risk of bias: moderate: possible recall bias from self-reported
Source of funding	outcomes Government: Environmental Protection Agency the National Institute for Environmental Health Sciences and the NHLBI
Comments	

D.1.96 Roda 2013

1100a 2013	Roda C, Guihenneuc-Jouyaux C, et of nocturnal dry cough in infancy:					
Bibliographic reference	domestic exposure to formaldehyd Environmental research 123, 46-51					
Study design	Prospective cohort					
Objective	To examine whether formaldehyde had during first year of life	as an impact on nocturnal dry cough				
Setting/Study location	France	France				
Number of participants	2898 infants					
Selected population	No					
Participant	Sex					
characteristics	Male	1486 (51.3%)				
	female	1412 (48.7%)				
	Maternal age (years)	Not reported				
	Ethnicity SES	Not reported				
	High	905 (65.7%)				
	Intermediate	770 (26.6%)				
	Low	223 (7.7%)				
	Annual family income	Not reported				
	Building characteristics	Not reported				
Inclusion criteria	Healthy full-term infants					
Exclusion criteria	None					
Type of pollutant/exposure	Formaldehyde					
Pollutant/exposure assessment	Formaldehyde was measured using a bedroom for 7 days and an annual ex	passive sample placed in the infant's posure level calculated.				
Outcome	Nocturnal dry cough apart from cold o	or chest infection				
Results	Formaldehyde exposure	aOR (95%CI)				
	With parental history of allergy	1.14 (0.88, 1.49)				
	Without parental history of allergy Gas heating	1.45 (1.08, 1.96)				
	With parental history of allergy	0.78 (0.56, 1.09)				
	Without parental history of allergy Used mattress	1.01 (0.69, 1.46)				
	With parental history of allergy	1.47 (1.00, 2.17)				
	Without parental history of allergy 1.22 (0.80, 1.88)					
Follow up	12 months					
Newcastle-Ottawa Scale	Selection Representativeness of the exposed cohort • truly representative of the average infant in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort t					
	Ascertainment of exposure					

Bibliographic reference	Roda C, Guihenneuc-Jouyaux C, et al (2013) Environmental triggers of nocturnal dry cough in infancy: new insights about chronic domestic exposure to formaldehyde in the PARIS birth cohort. Environmental research 123, 46-51
	Objective sampling)
	Demonstration that outcome of interest was not present at start of study
	• Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	study controls for environmental tobacco smoke
	 study controls for other factors as follows – SES, gender, breastfeeding, number of episodes of lower respiratory infections.
	Outcome
	Assessment of outcome
	• self-report
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	\bullet subjects lost to follow up unlikely to introduce bias - description provided of those lost) \Box
	Overall level of bias – Moderate (concerns over self-report of outcomes)
Source of funding	Government: Paris Council, French National Agency for Food, Environment and Occupational health safet6y (Anses), French Institute for Public health Surveillance (InVS)
Comments	

D.1.97 Samet 1993

Bibliographic reference	Samet J M, Lambert W E, Skipper B J et.al (1993) Nitrogen dioxide and respiratory illnesses in infants. The American review of respiratory disease 148(5), 1258-65				
Study design	Prospective col	nort study			
Objective		To test the hypothesis that exposure to NO ₂ indoors increases the incidence and severity of respiratory infections during the first 18 months of life			
Setting/Study location	United states	United states			
Number of participants	1,205 infants	1,205 infants			
Selected population	No				
Participant	Description	Gas stove		Electric stove	
characteristics		No.	%	No.	%
	Sex	Sex			
	Male	489	51.4	138	54.5

Bibliographic reference		esses in infa		(1993) Nitrogen can review of re		
	Female	463	48.6	115	45.5	
	Age (years)	Not reported		Not reported		
	Ethnicity	·				
	Hispanic	380	39.9	78	30.8	
	Non-Hispanic white	494	51.9	166	65.6	
	Other	78	8.2	9	3.6	
	Maintenance medication use	Not reported		Not reported		
	Parental asthma and/or atopic	Not reported		Not reported		
	Parental educat	ion				
	Maternal (years)				
	≤ 12	372	39.1	56	22.1	
	13 – 15	327	34.3	85	33.6	
	≥ 16	253	26.6	112	44.3	
	Annual family income (\$)					
	< 10, 000	117	12.3	11	4.4	
	10, 000- 19,000	227	23.8	30	12.0	
	20, 000- 29,000	226	23.7	53	20.9	
	30, 000- 39,000	186	19.5	60	23.7	
	≥ 40,000	197	20.7	99	39.0	
	Building charact	eristics				
	Single family					
	Unattached	686	72.1	205	81.0	
	Single family					
	Attached	59	6.2	10	4.0	
	Multifamily	108	11.3	35	13.8	
	Mobile home	96	10.1	3	1.2	
Inclusion criteria	Healthy term births Non-smoking mother and no other family member smoking inside the home Caring for the child at home Telephone in the residence Mother older than 18 years of age and English speaking No plans to move from study area					
Exclusion criteria	Not reported					

Bibliographic reference	Samet J M, Lambert W E, Skipper B J et.al (1993) Nitrogen dioxide and respiratory illnesses in infants. The American review of respiratory disease 148(5), 1258-65			
Type of pollutant/exposure	NO ₂ from cooking appliances with electric stove as reference category			
Pollutant/exposure assessment	NO ₂ concentrations were obtained with passive diffusion samplers (Palmes tubes). In homes with gas stoves, the child's bedroom was monitored every 2-week year round; during the colder seasons, additional 2-week measurements were made every other month in the kitchen and the activity room. In homes with electric stoves, the child's bedroom was monitored every other 2-week cycle year round. Consecutive 2-week measurements of outdoor concentrations were obtained at 11 monitoring sites			
Health outcome	of any: runny or s	stuffy nose, wet coug	sence of at least two lh, dry cough, wheez wo consecutive symp	e, or trouble
Results		tios (aORs) and 95% idence of respiratory	confidence intervals illness (RI)	s (CIs) for NO ₂
		All RI	Wet cough	Wheezing
		aOR (95%CI)	aOR (95%CI)	aOR (95%CI)
	Gas stove	0.98 (0.90, 1.07)	0.94 (0.82, 1.07)	0.84 (0.64, 1.09)
Follow up	18 months	,	, , ,	, , ,
Risk of bias (Newcastle-Ottawa Scale)	truly represental Selection of the record linkage self-report Was follow-up lor yes Adequacy of follow-up lor selection of the record linkage truly representation the record linkage Adequacy of follow-up lor yes	con-exposed cohort same community as exposure urement at outcome of interes cohorts on the basis for parental atopy and for other factors as for , income, breastfeed ting and season utcome and assessment or enough for outcor ow up of cohorts follow up unlikely to	nfant in the commun the exposed cohort st was not present at of the design or ana d asthma ollows - Age, gender, ling, maternal educat	start of study lysis ethnicity, birth tion, maternal

Bibliographic reference	Samet J M, Lambert W E, Skipper B J et.al (1993) Nitrogen dioxide and respiratory illnesses in infants. The American review of respiratory disease 148(5), 1258-65
Source of funding	Government: Research was conducted under contract to the Health Effects Institute (HEI), an organisation funded by the U.S. Environmental Protection Agency (EPA)
	Industry: Research was conducted under contract to the Health Effects Institute (HEI), an organisation funded by automobile manufacturers, and the Gas Research Institute (GRI).
Comments	Study suggests that NO_2 exposure from gas stove does not adversely affect the respiratory health of children during the first 18 months of life.
Additional references	Samet JM, Marbury MC, and Spengler JD (1987) Health Effects and Sources of Indoor Air Pollution. Part I. American Review of Respiratory Disease 136(6), 1486-1508

D.1.98 Sbihi 2016

Bibliographic reference	Sbihi Hind, Tamburic Lillian, Koehoorn Mieke, and Brauer Michael (2016) Perinatal air pollution exposure and development of asthma from birth to age 10 years. The European respiratory journal 47(4), 1062-71					
Study design	Prospective coho	Prospective cohort study				
Objective		To examine whether perinatal air pollution exposure affected asthma onset during "pre-school and "school age" periods in a population-based birth cohort				
Setting/Study location	Canada	Canada				
Number of participants	68195 children					
Participant		Pre-school		School age		
characteristics		Cases	Control	Cases	Control	
	Description	No. (%)	No. (%)	No. (%)	No. (%)	
	Sex (male) sex	4302 (62)	21478 (62)	5097 (59)	32642 (51)	
	Age (maternal, years); mean (SD)	31.2±5.09	31.5±5.06	31.5±5.12	31.4±5.06	
	Ethnicity	Not reported	Not reported	Not reported	Not reported	
	Cases/selected population	Not reported	Not reported	Not reported	Not reported	

Bibliographic reference	Sbihi Hind, Tam (2016) Perinatal birth to age 10 y	air polluti	ion e	xposure and d	evelo	pment of	asthma from
	Socio-economic status (maternal post-secondary education quartiles)					artiles)	
	1 (lowest)	1882 (27))	7670 (22)	195	9 (22)	13982 (22)
	2	1622 (23)		7973 (23)	199	6 (23)	14365 (23)
	3	1781 (26))	9407 (27)	227	0 (27)	17371 (27)
	4	1663 (24))	9571 (28)	235	2 (28)	17825 (28)
	Building characteristics	Not repor	ted	Not reported	Not	reported	Not reported
Inclusion criteria	Not reported						
Exclusion criteria	Not reported						
Type of pollutant/exposure	Proximity to traffi	С					
Pollutant/exposure assessment	Exposure to air pollution for each cohort member was assigned at their residential six-digit postal code(s), which corresponds to one block-face in urban areas (typically 100–150 m), by three different approaches: land use regression (LUR) models, interpolation of regulatory monitoring data (BC Ministry of Environment and Metro Vancouver) and proximity measures.						
Outcome	Asthma						
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between proximity to traffic and asthma onset						
	Asthma pre-school			Asthma school age			
		aOR (95%CI)			aOR (95°	%CI)	
	Within 50 m of hi		1.25	5 (1.04, 1.49)		0.81 (0.5	5, 1.19)
	Within 150 m of road	major	ajor 1.03 (0.98, 1.09)			1.04 (0.9	2, 1.16)
Follow up	10 years						
Study methods	Asthma diagnoses were identified from physician billing and hospital discharge records, obtained from the BC Ministry of Health. Using a validated case definition of asthma, children with a minimum of two primary-care physician diagnoses or one hospital admission in a rolling 12-month period were identified as asthma cases. Each asthma case was randomly matched to five controls and analysed with nested conditional logistic regression models adjusting for possible confounders						
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average population in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • validated measurement used Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis						

Bibliographic reference	Sbihi Hind, Tamburic Lillian, Koehoorn Mieke, and Brauer Michael (2016) Perinatal air pollution exposure and development of asthma from birth to age 10 years. The European respiratory journal 47(4), 1062-71
	 study controls for breastfeeding status at the time of discharge, parity, maternal education, household income, gestational length and birthweight Outcome Assessment of outcome record linkage physician billing Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts subjects lost to follow up unlikely to introduce bias
	Overall risk of bias: low
Source of funding	Government: Health Canada via an agreement with the British Columbia Centre for Disease Control. Additional support was provided by the Centre for Health and Environment Research at the University of British Columbia, funded by the Michael Smith Foundation for Health Research. H. Sbihi was funded by a Canadian Institutes of Health Research Banting and Best doctoral award
Comments	

D.1.99 Sherriff 2005

Sherrin 2005					
Bibliographic reference	Sherriff A, Farrow A, Golding J, and Henderson J (2005) Frequent use of chemical household products is associated with persistent wheezing in pre-school age children. Thorax 60(1), 45-9				
Study design	Prospective cohort study				
Objective	To examine the effect of prenatal exposure to multiple chemical agents on patterns of wheeze (never wheezed, transient early wheeze, persistent wheeze, late onset wheeze) during the first 3.5 years of life				
Setting/Study location	United Kingdom				
Number of participants	14,541 pregnant women				
Selected population	No				
Participant characteristics	Description Sex Age Ethnicity Education	Not reported Up to 4 years Not reported			
Inclusion criteria	Expected date of delivery between April 1, 1991 and December 31, 1992 Place of residence within the three Bristol-based health districts of the former county of Avon, UK				
Exclusion criteria	Not reported				
Type of pollutant/exposure	Total chemical burden (TCB) score				

Bibliographic reference	Sherriff A, Farrow A, Golding J, and He chemical household products is assoc pre-school age children. Thorax 60(1),	iated with persistent wheezing in			
Pollutant/exposure assessment	Questionnaire				
Outcome	Wheeze in child				
Results	Early-onset transient wheeze Early onset persistent wheeze Late onset wheeze	TCB burden during pregnancy 1.01 (0.99 to 1.02) 1.06 (1.03 to 1.09) 1.02 (0.98 to 1.06)			
Follow up	4 years				
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohor • selected group of pregnant women Selection of the non-exposed cohort • drawn from the same community as the Ascertainment of exposure • questionnaire Demonstration that outcome of interest water of the exposure • Yes Comparability Comparability Comparability of cohorts on the basis of the • study controls for weekend exposure to months • study controls for any additional factors pregnancy, maternal history of asthma, home, sex, contact with pets, damp how maternal educational attainment, housing outside home, month of returning chemical duration of breastfeeding.) Outcome Assessment of outcome • self-report Was follow-up long enough for outcomes • Yes Adequacy of follow up of cohorts • complete follow up - all subjects account Overall risk of bias: High (concerns over outcomes)	exposed cohort as not present at start of study ne design or analysis environmental tobacco smoke at 6 -, maternal smoking during maternal parity, crowding in the using, maternal age at delivery, ng tenure, hours mother worked ical usage questionnaire, and to occur			
Source of funding	Government: The Department of Health, and the Department of the Environment. Medical Research Council, Charity: the Wellcome Trust, Academic: University of Bristol,				
Comments	The composite household chemical exposure (CHCE) comprises of 11 different products (disinfectant; bleach; carpet cleaner; window cleaner; dry cleaning fluid; aerosols, turpentine/white spirit, air fresheners (spray, stick or aerosol); paint stripper; paint or varnish; and pesticides/insect killers). A simple score for frequency of use of each product was derived: 0 for not at all; 1 for less than once a week; 2 for about once a week; 3 for most days;				

Bibliographic reference	Sherriff A, Farrow A, Golding J, and Henderson J (2005) Frequent use of chemical household products is associated with persistent wheezing in pre-school age children. Thorax 60(1), 45-9
	and 4 for every day. The scores for each product were summed to produce a composite household chemical exposure (CHCE) score for each respondent

D.1.100 Shmuel 2017

Bibliographic reference Study design Objective	Shmuel S, White AJ, and Sandler DP (2017) Residential exposure to vehicular traffic-related air pollution during childhood and breast cancer risk. Environmental Research 159, 257-263 Prospective cohort study United States and Puerto Rico						
Setting/Study location		u Puerto Kico					
Number of participants	42,934 adults						
Selected population	No						
Participant		Cases		Non-cases			
characteristics	Description	No.	%	No.	%		
	Sex	Not reported	Not reported	Not reported	Not reported		
	Age (years)						
	<50	464	23	11,499	28		
	50 - <55	359	18	8,028	20		
	55 - <60	402	20	8,196	20		
	60 - <65	377	19	6,217	15		
	65+	426	21	6,966	17		
	Ethnicity						
	Non-Hispanic, White	1,760	87	34,623	85		
	Non-Hispanic, Black	143	7	3,413	8		
	Hispanic	67	3	1,866	5		
	Other	58	3	1,004	2		
	Cases/selected population						
	Socio-economic	status (educatio	on)				
	High School or Less	1,076	53	22,088	54		
	Some College	409	20	7,726	19		

Bibliographic reference	Shmuel S, White vehicular traffic cancer risk. Env	-related air po	llution d	uring o	childhood and	
	Bachelor's Degree	336	17		6,752	17
	Graduate Degree	207	10		4,340	11
	Building characteristics	Not reported	Not rep	orted	Not reported	Not reported
Inclusion criteria	Participants with one sister who had been diagnosed with breast cancer but had not been diagnosed with breast cancer themselves at the time of enrolment					
Exclusion criteria	Not reported					
Type of pollutant/exposure	Proximity to traffi	С				
Pollutant/exposure assessment	Participants completed a Computer-Assisted Telephone Interview in which they reported information on characteristics of their longest lived residence before age 14, including information on nearby roads and exposure to traffic. Participants were asked about the number of lanes, presence of a median or barrier dividing the road ('yes'/'no'), and traffic volume during rush hour ('very light,' 'imoderate,' 'heavy,' 'very heavy,' which were combined as 'light,' 'moderate,' and 'heavy' for most analyses) for their residential road					
Outcome	Breast cancer					
Results	Adjusted hazard ratios (aHRs) and 95% confidence intervals (Cls) for association between proximity to traffic and breast cancer					
	Distance of residence to nearest road and number of lanes on intersecting					
				Total breast cancer		
				aHR (95%CI)		
	100 ft. +			REF		
	Within 100 ft. 1–2 Lanes			0.9 (0.8, 1.0)		
	Within 100 ft. 3+	Lanes		1.1 (0.9, 1.4)		
Follow up	6.3 years					
Study methods	Incident breast cancer diagnoses were ascertained from annual health updates and biennial/triennial questionnaires that participants completed during follow-up. Women who reported a diagnosis during follow-up were asked for consent to review their medical records for confirmation and for diagnostic and treatment details. Cox proportional hazards models were used to estimate the association between characteristics of the primary childhood residence and incident breast cancer adjusting for possible confounders					
Risk of bias (Newcastle-Ottawa Scale)	Selection			1		
	questionnaires on residential proximity to traffic Demonstration that outcome of interest was not present at start of study					

Bibliographic reference	Shmuel S, White AJ, and Sandler DP (2017) Residential exposure to vehicular traffic-related air pollution during childhood and breast cancer risk. Environmental Research 159, 257-263
	Yes Comparability Comparability of cohorts on the basis of the design or analysis study controls for age, race/ethnicity and highest level of education attained in the household at age 13 Outcome Assessment of outcome record linkage and treatment details self-report Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts subjects lost to follow up unlikely to introduce bias Overall risk of bias: moderate: possibility of over or underestimating traffic exposure stemming from self-reporting traffic exposure
Source of funding	Study was supported by the Intramural Research Program of the NIH, National Institute of Environmental Health Sciences [Z01- ES044005]; and the NIEHS grant [T32ES007018]
Comments	

D.1.101 Shu 2013

311u 2013	
Bibliographic reference	Shu H, Jönsson BA, Larsson M, et al. (2006) PVC flooring at home and development of asthma among young children in Sweden, a 10-year follow-up. Indoor Air. 24(3): 227-35. doi: 10.1111/ina.12074.
Study design	Prospective cohort study
Objective	To investigate whether PVC flooring in the home of children between 1 and 5 years old is associated with the development of asthma at 5- and 10-year follow-up investigations.
Setting/Study location	Varmland, Sweden
Number of dwellings and participants	Number of dwellings: Not reported Number of participants: 3,228 children
Building and Participant characteristics	Building characteristics: Location: unclear Dwelling type: single family, 80.9%; attached or semi-attached, 8.6%; flat/apartment/multifamily, 8.3%; other, 2.2% Building age: not reported Type of ownership/tenancy: not reported Participant characteristics:

development of asthma among	young children in Sv	veden, a 10-year		
Sex: 50.3% male				
Age: not reported				
Smokers in the family: mother, 10.1%; father, 9.3%				
-				
Preschool children in the county o further details were provided.	f Varmland in Sweden	, were included. No		
Not reported				
PVC flooring vs. other flooring				
Building factors were ascertained reported questionnaire.	by asking participants	to complete a self-		
Doctor diagnosed asthma				
Building characteristic	Odds ratio (95%CI)			
	5 years	10 years		
Child's bedroom				
PVC vs. other flooring material	1.50 (0.91, 2.47)	1.54 (1.06, 2.23)		
PVC vs. wood flooring material	1.54 (1.06, 2.23)	1.37 (0.92, 2.04)		
Parent's bedroom				
PVC vs. other flooring material	1.71 (1.05, 2.80)	2.04 (1.41, 2.94)		
PVC vs. wood flooring material	1.60 (1.29, 2.81)	1.90 (1.29, 2.81)		
10 years				
Selection Representativeness of the exposed cohort • selected group – preschool children in the county of Varmland in Sweden Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • self-reported Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study adjusts for age, sex, allergies in family, single parent households, smoking in the family, and housing type. Outcome Assessment of outcome • self-reported but the questionnaire question was asked in such a way, it is unlikely to have introduced bias Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts				
	development of asthma among follow-up. Indoor Air. 24(3): 227 Sex: 50.3% male Age: not reported Smokers in the family: mother, 10 Asthma or allergies in the family: Preschool children in the county of further details were provided. Not reported PVC flooring vs. other flooring Building factors were ascertained reported questionnaire. Doctor diagnosed asthma Building characteristic Child's bedroom PVC vs. other flooring material PVC vs. wood flooring material O years Selection Representativeness of the expose selected group – preschool child Selection of the non-exposed cohe drawn from the same community Ascertainment of exposure self-reported Demonstration that outcome of interposite study adjusts for age, sex, allergy smoking in the family, and house Outcome Assessment of outcome self-reported but the questionna unlikely to have introduced bias Was follow-up long enough for out Yes Adequacy of follow up of cohorts	Age: not reported Smokers in the family: mother, 10.1%; father, 9.3% Asthma or allergies in the family: 57% Preschool children in the county of Varmland in Sweden further details were provided. Not reported PVC flooring vs. other flooring Building factors were ascertained by asking participants reported questionnaire. Doctor diagnosed asthma Building characteristic Odds ratio (95%CI) 5 years Child's bedroom PVC vs. other flooring material 1.50 (0.91, 2.47) PVC vs. wood flooring material 1.54 (1.06, 2.23) Parent's bedroom PVC vs. other flooring material 1.71 (1.05, 2.80) PVC vs. wood flooring material 1.60 (1.29, 2.81) 10 years Selection Representativeness of the exposed cohort • selected group – preschool children in the county of V Selection of the non-exposed cohort • drawn from the same community as the exposed cohord • selected group – preschool children in the county of V Selection of the same community as the exposed cohord • selected group – the same community as the exposed cohord • self-reported Demonstration that outcome of interest was not present • Yes Comparability Comparability of cohorts on the basis of the design or an exposed subject of the family, and housing type. Outcome Assessment of outcome • self-reported but the questionnaire question was askerunlikely to have introduced bias Was follow-up long enough for outcomes to occur		

Bibliographic reference	Shu H, Jönsson BA, Larsson M, et al. (2006) PVC flooring at home and development of asthma among young children in Sweden, a 10-year follow-up. Indoor Air. 24(3): 227-35. doi: 10.1111/ina.12074.
Source of funding	Government: Swedish Research Council for Environment, Agricultural Sciences and Spatial Planning Charity: Swedish Asthma and Allergy Associations Research Foundation
Comments	None

D.1.102 Smith 2000

Bibliographic reference	Sorensen M, Andersen A-M, and Raaschou-Nielsen O (2010) Non- occupational exposure to paint fumes during pregnancy and fetal growth in a general population. Environmental research 110(4), 383-7					
Study design	Prospective cohort study					
Objective	during pregnancy a	To investigate the association between exposure to paint fumes in the residence during pregnancy and birth weight, small for gestational age (SGA) and preterm births in a national prospective birth cohort.				
Setting/Study location	Denmark					
Number of participants	19,000 women					
Selected population	No					
		Exposed to pa	int fumes	Not exposed to	paint fumes	
Participant	Description	No.	%	No.	%	
characteristics	Sex	All female	All female	All female	All female	
	Age (years); mean (SD)	29.0 (4.3)	-	29.6 (4.3)	-	
	Ethnicity	Not reported	Not reported	Not reported	Not reported	
	Cases/selected population	Pregnant women	Pregnant women	Pregnant women	Pregnant women	
	Socio-economic status (education)	Not reported	Not reported	Not reported	Not reported	
	Building characteristics	Not reported	Not reported	Not reported	Not reported	
Inclusion criteria	Women who gave birth to live-born singletons Available information on birth weight Not occupationally exposed to organic solvents					
Exclusion criteria	Not reported					
Type of pollutant/expos ure	Exposure to paint fumes					
Pollutant/expo sure assessment	questions regarding	During the second prenatal telephone interview, participants were asked questions regarding the use of paint in their residence. They were asked if any painting had been done in their residence during pregnancy and if so, what rooms				

Bibliographic reference	Sorensen M, Andersen A-M, and Raaschou-Nielsen O (2010) Non- occupational exposure to paint fumes during pregnancy and fetal growth in a general population. Environmental research 110(4), 383-7		
	had been painted, if they painted "furniture, floor, radiator and/or woodwork" or "wall and/or ceiling", and when the painting was done. Furthermore, the women were asked if they were present in the room for two or more hours during painted.		
Outcome	Small for gestational age (SGA)	and preterm birth	
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between exposure to paint fumes in the residence during pregnancy and small for gestational age (SGA) and preterm birth		
		Small for gestational age	Preterm birth
		aOR (95%CI)	aOR (95%CI)
	Not exposed to paint fumes	1.00	1.00
	Exposed to paint fumes	0.89 (0.81, 0.98)	0.95 (0.82, 1.11)
Follow up	6 years		
Study methods	ods SGA births were defined as those with birth weights below the 10th percentile of the cohort, stratified by sex, for each week of gestation. Preterm birth was defined as birth before the 37th week of gestation. Authors used general and multiple linear regressions models (proc GLM, SAS) to test for associations between exposure to paint fumes and birth weight and		
Risk of bias (Newcastle- Ottawa Scale)	Authors used general and multiple linear regressions models (proc GLM, SAS) to		

Bibliographic reference	Sorensen M, Andersen A-M, and Raaschou-Nielsen O (2010) Non- occupational exposure to paint fumes during pregnancy and fetal growth in a general population. Environmental research 110(4), 383-7
Source of funding	Government: The Danish Agency for Science, Technology and Innovation. Charity: The Danish National Research Foundation, Pharmacy Foundation, the Egmont Foundation, the March of Dimes Birth Defects Foundation, the Augustinus Foundation, and the Health Foundation.
Comments	

D.1.103 Sorensen 2010

Bibliographic reference	Sorensen M, Andersen A-M, and Raaschou-Nielsen O (2010) Non- occupational exposure to paint fumes during pregnancy and fetal growth in a general population. Environmental research 110(4), 383-7				
Study design	Prospective cohort	study			
Objective	during pregnancy a	To investigate the association between exposure to paint fumes in the residence during pregnancy and birth weight, small for gestational age (SGA) and preterm births in a national prospective birth cohort.			
Setting/Study location	Denmark	Denmark			
Number of participants	19,000 women	19,000 women			
Selected population	No				
		Exposed to pa	int fumes	Not exposed to	paint fumes
Participant	Description	No.	%	No.	%
characteristics	Sex	All female	All female	All female	All female
	Age (years); mean (SD)	29.0 (4.3)	-	29.6 (4.3)	-
	Ethnicity	Not reported	Not reported	Not reported	Not reported
	Cases/selected population	Pregnant women	Pregnant women	Pregnant women	Pregnant women
	Socio-economic status (education)	Not reported	Not reported	Not reported	Not reported
	Building characteristics	Not reported	Not reported	Not reported	Not reported
Inclusion criteria	Women who gave birth to live-born singletons Available information on birth weight Not occupationally exposed to organic solvents				
Exclusion criteria	Not reported				

Bibliographic	Sorensen M, Andersen A-M, a occupational exposure to pair	nt fumes during preg	gnancy and fetal growth in
reference	a general population. Environmental research 110(4), 383-7		
Type of pollutant/expos ure	Exposure to paint fumes		
Pollutant/expo sure assessment	During the second prenatal telephone interview, participants were asked questions regarding the use of paint in their residence. They were asked if any painting had been done in their residence during pregnancy and if so, what rooms had been painted, if they painted "furniture, floor, radiator and/or woodwork" or "wall and/or ceiling", and when the painting was done. Furthermore, the women were asked if they were present in the room for two or more hours during painted.		
Outcome	Small for gestational age (SGA)	and preterm birth	
Results	Adjusted odds ratios (aORs) and between exposure to paint fume gestational age (SGA) and preter	s in the residence du	
		Small for gestational age	Preterm birth
		aOR (95%CI)	aOR (95%CI)
	Not exposed to paint fumes	1.00	1.00
	Exposed to paint fumes	0.89 (0.81, 0.98)	0.95 (0.82, 1.11)
Follow up	6 years		
Study methods	the cohort, stratified by sex, for each week of gestation. Preterm birth was as birth before the 37th week of gestation. Authors used general and multiple linear regressions models (proc GLM test for associations between exposure to paint fumes and birth weights)		n. Preterm birth was defined models (proc GLM, SAS) to
Risk of bias (Newcastle- Ottawa Scale)	adjusted for confounding factors. Selection Representativeness of the exposed cohort • truly representative of the average female population in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • structured interview • self-report Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • Study controls for smoking in 30th pregnancy week, maternal age, maternal pre-pregnancy BMI, parity (nulliparous, uniparous or multiparous) and occupational status. Outcome Assessment of outcome • Information on birth weight and gestational age was obtained from the Danish National Birth Register and the Danish National Discharge Registry, respectively. Gestational age was recorded by midwives at birth Was follow-up long enough for outcomes to occur		

Bibliographic reference	Sorensen M, Andersen A-M, and Raaschou-Nielsen O (2010) Non- occupational exposure to paint fumes during pregnancy and fetal growth in a general population. Environmental research 110(4), 383-7
	 Yes Adequacy of follow up of cohorts subjects lost to follow up unlikely to introduce bias Overall risk of bias: Low
Source of funding	Government: The Danish Agency for Science, Technology and Innovation. Charity: The Danish National Research Foundation, Pharmacy Foundation, the Egmont Foundation, the March of Dimes Birth Defects Foundation, the Augustinus Foundation, and the Health Foundation.
Comments	

D.1.104 Stark 2003

5tark 2005			
Bibliographic reference	Stark PC, Burge HA, Ryan LM, et al (2003) Fungal levels in the home and lower respiratory tract illnesses in the first year of life. American journal of respiratory and critical care medicine 168(2), 232-7		
Study design	Prospective cohort study		
Objective	Too determine if exposure to fungi is a in the first year of life.	associated with lower respiratory illness	
Setting/Study location	United States		
Number of participants	499 infants		
Selected population	Yes – parental history of asthma or all	ergy	
Participant characteristics	Description Sex Male Female Age (years)- Maternal 18 -<30 30 to<33 33 to<36 36 to<46 Ethnicity White Back Hispanic Asian Other Education Annual family income	233 266 Not reported 124 125 125 125 375 59 30 28 7 Not reported Not reported	
Inclusion criteria	Not reported		
Exclusion criteria	Not reported		
Type of pollutant/exposure	Water damage or mould/mildew		

Bibliographic reference	Stark PC, Burge HA, Ryan LM, et al (2003) Fungal levels in the home and lower respiratory tract illnesses in the first year of life. American journal of respiratory and critical care medicine 168(2), 232-7		
Pollutant/exposure assessment	Indoor air samples were collected from each home using a Burkard culture plate		
	Sequential duplicate 1-min air samples were collected in the bedroom 1–1. m above the area of the floor demarcated for dust collection. After sampling the Petri plates were returned to the laboratory on the same day for incubation High fungal levels defined as > 90th percentile or specific taxon		
Outcome	Lower respiratory illness		
Results		LRI	
		aRR (95%CI)	
	Water damage or mould/mildew	1.34 (0.99, 1.82)	
	High fungal levels	1.86 (1.21, 2.88)	
Follow up	1 year		
Risk of bias (Newcastle-Ottawa Scale)	, , ,		
Source of funding	Overall level of bias – Low Government: This study was supported by National Institutes of Health		
Comments	Government. This study was suppor	Ted by Ivalional Institutes of Health	
Comments			

D.1.105 Stark 2005

Stark PC, Celedon JC, Chew GL, et al (2005) Fungal levels in the home and allergic rhinitis by 5 years of age. Environmental health perspectives 113(10), 1405-9		
Prospective cohort study		
To evaluate whether high fungal levels were independently associated with doctor-diagnosed allergic rhinitis in the first 5 years of life.		
United States		
405 children (< 5 years)		
Yes – parental history of asthma or	allergy	
Sex Male 210 Female 195 Age (years)- Maternal Not reported Ethnicity White 309 African American 46 Hispanic 18 Asian 26 Other 6 Education Not reported		
Residence inside route 128 (a highway encircling the Boston metropolitan area) Maternal age ≥ 18 years History of hay fever, asthma, or allergies in either parent Maternal ability to speak English or Spanish		
Mould/mildew		
Indoor air samples were collected from each home using a Burkard culture plate Sequential duplicate 1-min air samples were collected in the bedroom 1–1.5 m above the area of the floor demarcated for dust collection. After sampling, the Petri plates were returned to the laboratory on the same day for incubation High fungal levels defined as > 90th percentile or specific taxon		
Allergic rhinitis		
Adjusted hazard ratios (aHRs) and 95% confidence intervals (CIs) for association between exposure to house dust mite, cat allergen, dog allergen and asthma		
	Allergic rhinitis	
	aHR (95%CI)	
Water damage or mould/mildew in year	1.66 (0.88, 3.15)	
	and allergic rhinitis by 5 years of perspectives 113(10), 1405-9 Prospective cohort study To evaluate whether high fungal lev doctor-diagnosed allergic rhinitis in the United States 405 children (< 5 years) Yes – parental history of asthma or Description Sex Male Female Age (years)- Maternal Ethnicity White African American Hispanic Asian Other Education Annual family income Residence inside route 128 (a highwarea) Maternal age ≥ 18 years History of hay fever, asthma, or alled Maternal ability to speak English or Mould/mildew Indoor air samples were collected frolate Sequential duplicate 1-min air samplem above the area of the floor demant the Petri plates were returned to the incubation High fungal levels defined as > 90th Allergic rhinitis Adjusted hazard ratios (aHRs) and sassociation between exposure to he and asthma	

Bibliographic reference	Stark PC, Celedon JC, Chew GL, et al (2005) Fungal levels in the home and allergic rhinitis by 5 years of age. Environmental health perspectives 113(10), 1405-9
Follow up	
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • selected group of children at high risk of asthma Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • Objective sampling Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability Comparability of cohorts on the basis of the design or analysis • study controls for any maternal allergy • study controls for additional factors as follows - male sex, African-American race, fall date of birth, and maternal IgE Outcome Assessment of outcome • independent blind assessment Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall level of bias – Low risk
Source of funding	Government: This study was supported by National Institutes of Health
Comments	

D.1.106 Thacher 2017

Bibliographic reference	Thacher J D, Gruzieva O, Pershagen G, et al (2017) Mold and dampness exposure and allergic outcomes from birth to adolescence: data from the BAMSE cohort. Allergy 72(6), 967-974	
Study design	Prospective cohort study	
Objective	To assess whether exposure to mould or dampness during infancy influences the risk of asthma or rhinitis in children followed prospectively from birth to adolescence.	
Setting/Study location	Sweden	
Number of participants	3798 children	
Selected population	No	
Participant	Age	Not reported
characteristics	Sex	

exposure and allergic outcomes	from birth to adolesce	
Male		1593
Race / ethnicity		Not reported
SES reported as working status Manual worker Non-manual worker		488 2655
Children born in selected areas of S November 1996	Stockholm County betw	een February 1994 and
Not reported		
Mould Dampness		
Self-report		
Asthma and rhinitis at 1–16 years of age were based on symptoms reported by parents from questionnaires and were defined as follows:		
Asthma—four or more episodes of wheeze in the last 12 months or one or more episode of wheeze in the last 12 months in combination with inhaled steroids Rhinitis—eye or nose symptoms following exposure to allergens in the last 12 months and/or a doctor's diagnosis of allergic rhinitis (18)		ith inhaled steroids
Exposure	Asthma aOR (95%CI)	Rhinitis aOR (95%CI)
No mould or dampness indicator 1 indicator 2 indicators 3 indicators	Reference 1.16 (0.93, 1.44) 1.37 (1.01, 1.86) 1.73 (1.10, 2.74)	Reference 1.03 (0.87, 1.22) 1.18 (0.92, 1.52) 1.23 (0.82, 1.85)
16 years		
Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • written self-report Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for presence of parental smoking during infancy, • study controls for additional factor - sex, socioeconomic status, parental allergic disease, maternal smoking during pregnancy, maternal age<26 years, and presence of siblings. Outcome		
	exposure and allergic outcomes BAMSE cohort. Allergy 72(6), 967 Male Race / ethnicity SES reported as working status Manual worker Non-manual worker Children born in selected areas of S November 1996 Not reported Mould Dampness Self-report Asthma and rhinitis at 1–16 years of parents from questionnaires and we asthma—four or more episodes of episode of wheeze in the last 12 mc Rhinitis—eye or nose symptoms for months and/or a doctor's diagnosis Exposure No mould or dampness indicator 1 indicator 2 indicators 3 indicators 16 years Selection Representativeness of the exposed of truly representative of the average selection of the non-exposed cohor of drawn from the same community Ascertainment of exposure • written self-report Demonstration that outcome of inte of yes Comparability Comparability Comparability of cohorts on the base of study controls for presence of parents of study controls for additional factor disease, maternal smoking during presence of siblings.	Race / ethnicity SES reported as working status Manual worker Non-manual worker Children born in selected areas of Stockholm County between November 1996 Not reported Mould Dampness Self-report Asthma and rhinitis at 1–16 years of age were based on syparents from questionnaires and were defined as follows: Asthma—four or more episodes of wheeze in the last 12 nepisode of wheeze in the last 12 months in combination with Rhinitis—eye or nose symptoms following exposure to alle months and/or a doctor's diagnosis of allergic rhinitis (18). Exposure Exposure Asthma Asthma AoR (95%CI) No mould or dampness indicator 1 indicator 2 indicators 3 indicators 1.37 (1.01, 1.86) 3 indicators 1.73 (1.10, 2.74) 16 years Selection Representativeness of the exposed cohort • truly representative of the average child in the communit Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • written self-report Demonstration that outcome of interest was not present at Pyes Comparability Comparability Comparability Comparability Comparability of cohorts on the basis of the design or anal study controls for additional factor - sex, socioeconomic disease, maternal smoking during pregnancy, maternal appresence of siblings. Outcome

Bibliographic reference	Thacher J D, Gruzieva O, Pershagen G, et al (2017) Mold and dampness exposure and allergic outcomes from birth to adolescence: data from the BAMSE cohort. Allergy 72(6), 967-974
	 clinical diagnosis Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts complete follow up - all subjects accounted for Overall risk of bias: Low
Source of funding	Government: The Swedish Research Council, The Swedish Heart and Lung Foundation, The Swedish Research Council for Working Life and Social Welfare, The Swedish Asthma and Allergy Association Research Foundation, The Swedish Research Council Formas, Stockholm County Council, and the European Commission
Comments	Mould or dampness indicators=Mould odour, visible mould, or dampness damage.

D.1.107 Tiesler 2015

Bibliographic reference	Tiesler C M. T, Thiering E, Tischer C et.al (2015) Exposure to visible mould or dampness at home and sleep problems in children: Results from the LISAplus study. Environmental research 137, 357-63				
Study design	Prospective cohort study				
Objective	To investigate the association between reported current visible mould or dampness at home and sleep problemsin10-year-oldchildren.				
Setting/Study location	Germany				
Number of participants	1719 children				
Selected population	No				
Participant characteristics	Description Sex Age (years) Ethnicity Education Annual family income Building characteristics Owner occupancy Tenancy Crowding index <1 (low)	Not reported Not reported Not reported Not reported Not reported Very reported Not reported 2916 (52%) 2724 (48%) 336 (6%)			

Bibliographic reference	Tiesler C M. T, Thiering E, Tischer C et.al (2015) Exposure to visible mould or dampness at home and sleep problems in children: Results from the LISAplus study. Environmental research 137, 357-63						
	1-<2 (medium) 2+ (high) Building age	1345	4430 (72%) 1345 (2%) Not reported2				
Inclusion criteria	Healthy, full-term neonates born in four German cities (Munich, Leipzig, Wesel and Bad Honnef)						
Exclusion criteria	preterm birth (maturity <37 gestational weeks), low birth weight (<2,500 g), congenital malformation, symptomatic neonatal infection, antibiotic medication, hospitalisation or intensive medical care during neonatal period. new-borns from mothers with immune-related diseases (autoimmune disorders, diabetes, hepatitis B), on long-term medication or who abuse drugs and/or alcohol new-borns from parents with a nationality other than German or who were not born in Germany						
Type of pollutant/exposure	Visible mould or dampness						
Pollutant/exposure assessment	Self-assessment						
Outcome	Sleep problems						
Results		Any sleep problems	Problems to fall asleep	Problems sleeping through the night	Sleep time <9hours		
	Visible mould	1.70 (1.13, 2.54)	1.50 (0.97, 2.33)	1.91 (0.89, 4.13)	1.67 (1.06, 2.65)		
	Damp	2.59 (1.26, 5.31)	1.39 (0.57, 3.40)	4.30 (1.41, 13.13)	0.69 (0.21, 2.28)		
	Visible mould/dampnes s at home	1.80(1.22, 2.66)	1.50 (0.98, 2.30)	2.36 (1.15, 4.84)	1.60 (1.02, 2.51)		
Follow up	10 years						
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • written self-report Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for sex						

Bibliographic reference	Tiesler C M. T, Thiering E, Tischer C et.al (2015) Exposure to visible mould or dampness at home and sleep problems in children: Results from the LISAplus study. Environmental research 137, 357-63
	 study controls for any additional factors as follows – study centre, sex, parental education level and bedroom sharing Outcome Assessment of outcome self-report Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts complete follow up - all subjects accounted for Overall risk of bias – High (concerns over self-report of exposure and
Source of funding	outcomes) Government: Federal Ministry for Education, Science, Research and Technology, Helm-holtz Zentrum Munich (former GSF), Helmholtz Centre for Environmental Research – UFZ, Leipzig,
Comments	e. E. T. Edipeig,

D.1.108 Tin Tin 2016

Bibliographic reference	Tin Tin S, Woodward A, Saraf R et.al (2016) Internal living environment and respiratory disease in children: findings from the Growing Up in New Zealand longitudinal child cohort study. Environmental health: a global access science source 15(1), 120			
Study design	Prospective cohort study			
Objective	To investigate the frequency and pattern of exposure to specific home environmental risk factors and to provide updated evidence of the impact of these exposures on the risk of hospital admission with ARIs during the first five years of life.			
Setting/Study location	New Zealand			
Number of participants	6853 children			
Selected population	No			
Participant characteristics	Description Sex Age (years)- Ethnicity Education Annual family income	Not reported Not reported Not reported Not reported Not reported		
Inclusion criteria	Pregnant women had to be resident within a geographical region defined by three contiguous District Health Board (DHB) regions in the northern part of the country (Auckland, Counties-Manukau and Waikato), Have an estimated delivery date between 25 April 2009 and 25 March 2010			
Exclusion criteria	Not reported			
Type of pollutant/exposure	Mould or mildew			

Bibliographic	Tin Tin S, Woodward A, Saraf R et.al and respiratory disease in children: New Zealand longitudinal child coho	findings fort study.	rom the Growing Up in	
reference	global access science source 15(1),	120		
	Gas heater			
	Over-crowding			
Dallada ada a	tenure			
Pollutant/exposure assessment	Questionnaire			
Outcome	Hospitalization for acute respiratory in	fections		
Results	Adjusted hazard ratios (aHRs) and 95% confidence intervals (CIs) for association between internal living environment and hospital admission for an acute respiratory infection (ARI) during the first five years of life			
			ARI	
			aHR (95%CI)	
	Housing tenure Tenancy		1.00 (0.87, 1.16)	
	Crowding index 2+ (high)		1.07 (0.91, 1.26)	
	Heating the house Yes		0.87 (0.72, 1.05)	
	Gas heater only		1.64 (1.29-2.09)	
	Unflued gas heater		1.48 (1.05-2.09)	
	Gas heater as well as other forms of heating		0.82 (0.68-0.99)	
	Unflued gas heater as well as other forms of heating		0.73 (0.57-0.94)	
	Heating used in the room where child s	leeps		
	No heating			
	Yes	0.82 (0.6	7, 1.00)	
	No	1.00		
	Dampness of the house			
	Never or hardly ever	1.00		
	Not very often	0.96 (0.8	3-1.13)	
	Quite often	1.13 (0.9	•	
	Always or almost always	1.15 (0.8	9, 1.50)	
	Heavy condensation in the room where child sleeps at night			
	Never or hardly ever	· ·		
	Not very often	1.01 (0.8	6-1.17)	
	Quite often	1.05 (0.8	8-1.27)	
	Always or almost always	1.00 (0.7	7, 1.31)	
	Mould or mildew in the walls or ceilings in the room where child sleeps at night in the past two weeks			
	Yes 0.81 (0.67, 0.99)			
Follow up	5 years			
Risk of bias (Newcastle-Ottawa	castle-Ottawa Representativeness of the exposed cohort			
Scale)				

	Tin Tin S Woodward A Saraf P et al (2016) Internal living environment
Bibliographic	Tin Tin S, Woodward A, Saraf R et.al (2016) Internal living environment and respiratory disease in children: findings from the Growing Up in New Zealand longitudinal child cohort study. Environmental health:
reference	global access science source 15(1), 120
	drawn from the same community as the exposed cohort
	Ascertainment of exposure
	self-report □
	Demonstration that outcome of interest was not present at start of study • Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	study controls for maternal history of asthma
	 study controls for additional factors as follows – Maternal factors (age, ethnicity, education, area of residence, neighbourhood deprivation, prepregnancy BMI, pre-pregnancy self-rated health, history use of supplements, maternal smoking, parity and pregnancy planning) and child factors (gender, gestation, birth-weight, season of birth, proxy-rated health at 9 months, health or developmental problems, feeding practices, time spent outdoors and child immunisation)
	Outcome
	Assessment of outcome
	• record linkage
	Was follow-up long enough for outcomes to occur • Yes
	Adequacy of follow up of cohorts
	complete follow up - all subjects accounted for
	Overall risk of bias – Moderate (concerns over self-report of exposure)
Source of funding	Government: New Zealand Ministries of Social Development, Health, Education, Justice and Pacific Island Affairs; the former Ministry of Science Innovation and the former Department of Labour (now both part of the Ministry of Business, Innovation and Employment); the former Ministry of Women's Affairs (now the Ministry for Women); the Department of Corrections; the Families Commission (now known as the Social Policy Evaluation and Research Unit); Te Puni Kokiri; New Zealand Police; Sport New Zealand; the Housing New Zealand Corporation; and the former Mental Health Commission, The University of Auckland and Auckland UniServices Limited. Health Research Council of New Zealand, Statistics New Zealand, the Office of the Children's Commissioner and the Office of Ethnic Affairs.
Comments	

D.1.109 Torrent 2007

Bibliographic reference	Torrent M, Sunyer J, Garcia R, et al (2007) Early-life allergen exposure and atopy, asthma, and wheeze up to 6 years of age. American journal of respiratory and critical care medicine 176(5), 446-53
Study design	Prospective cohort study
Objective	To assess the prospective relationship between exposure to aeroallergens in early life and the development of specific sensitization, wheeze, and asthma up to 6 years of age in non-selected populations

Bibliographic reference	Torrent M, Sunyer J, Garcia R, et atopy, asthma, and wheeze up to respiratory and critical care medi	6 years of age. Amer			
Setting/Study location	Spain & UK				
Number of participants	1611 infants				
Selected population	No				
Participant characteristics	Description	No. (%)			
CHARACTERISTICS	Sex Ethnicity Annual family income Building characteristics	Not reported Not reported Not reported Not reported			
Inclusion criteria	Mothers delivering babies				
Exclusion criteria	Smoking in the household Infant death or adoption Maternal age <19 years Non-English speaking mother Prior participation Plans to move out of study area Having a multiple gestation Having no address or phone number				
Type of pollutant/exposu re	Allergens SES NO ₂				
Pollutant/expos ure assessment	Questionnaire and home inspection				
Outcome	Asthma Wheeze				
Results	Adjusted odds ratios (aORs) and 95	% confidence intervals	s (Cls)		
		Asthma	Persistent wheeze		
		aOR (95%CI)	aOR (95%CI)		
	SES (maternal social class) Skilled nonmanual Skilled manual Unskilled	0.56 (0.26–1.21) 0.74 (0.42–1.30) 0.75 (0.40–1.43)	0.49 (0.19–1.26) 0.94 (0.49–1.79) 1.52 (0.76–3.03)		
	NO ₂ , 10 μ g/m ³ 1.53 (0.87–2.70) Not reported		Not reported		
	Der p1 concentration 0.83–6.46 μg/g >6.46 μg/g Fel d1 concentration	0.67 (0.40–1.12) 0.68 (0.37–1.25)	0.59 (0.32–1.08) 0.74 (0.38–1.46)		
0.25–1.39 μg/g 1.59 (0.75–3.36) 0.73 (0.3 >1.39 μg/g 2.6 1(.27–5.34) 1.56 (0.7)					

Bibliographic reference	Torrent M, Sunyer J, Garcia R, et al (2007) Early-life allergen exposure and atopy, asthma, and wheeze up to 6 years of age. American journal of respiratory and critical care medicine 176(5), 446-53
Follow up	6 years
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average infant Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • written self-report Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for maternal atopy and maternal asthma, home crowding Outcome Assessment of outcome • self-report (wheeze • physician diagnosis (asthma) Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall level of bias – Low
Source of funding	Government: Spanish Ministry of Health, European Community Charity: The COLT Foundation
Comments	

D.1.110 Triche 2002

Bibliographic reference	Triche EW, Belanger K, Beckett W, et al (2002) Infant respiratory symptoms associated with indoor heating sources. American Journal of Respiratory and Critical Care Medicine 166(8), 1105-1111		
Study design	Prospective cohort study		
Objective	To examine the effect of secondary home heating on respiratory symptoms		
Setting/Study location	United States		
Number of participants	890 infants		
Selected population	No		
	Description	No. (%)	

Bibliographic reference	Triche EW, Belanger K, Beckett W, et al (2002) Infant respiratory symptoms associated with indoor heating sources. American Journal of Respiratory and Critical Care Medicine 166(8), 1105-1111			
Participant characteristics	Sex Male female	466 (52%) 423 (48%)		
	Maternal age (years) Not reported			
	Ethnicity White or Asian Black or Hispanic	696 (78%) 193 (21%)		
	Maternal asthma and/or atopic 80 (9%)			
	Parental education High school or less Some college College graduate or higher	High school or less 261 (29%) Some college 325 (39%)		
	Annual family income	Not reported		
	Building characteristics	Not reported		
Inclusion criteria	Mothers delivering babies			
Exclusion criteria	Smoking in the household Infant death or adoption Maternal age <19 years Non-English speaking mother Prior participation Plans to move out of study area Having a multiple gestation Having no address or phone number			
Type of pollutant/exposu re	Heating sources: NO ₂ Fireplace (FP) Gas space heater (GH) Kerosene heater (KH) Wood stove (WS)			
Pollutant/expos ure assessment	Questionnaire and home ins	pection		
Outcome	Respiratory symptoms (diary	')		
Results	Adjusted risk ratios (aRRs) a between Indoor heating sour			
		Wheeze	Cough	
		aRR (95%CI)	aRR (95%CI)	
	Average per day FP use	0.25 (0.04, 1.43)	0.99 (0.81, 1.21)	
	Average per day WS use	1.08 (0.87, 18.39)	1.10 (1.02, 1.19)	
	Average per day GH use	1.25 (1.05, 1.50)	0.94 (0.75, 1.18)	
	Average per day KH use	0.90 (0.64, 1.25)	1.01 (0.93, 1.10)	
Follow up	12 months			
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the ex	xposed cohort		

Bibliographic reference	Triche EW, Belanger K, Beckett W, et al (2002) Infant respiratory symptoms associated with indoor heating sources. American Journal of Respiratory and Critical Care Medicine 166(8), 1105-1111
	 truly representative of the average infant Selection of the non-exposed cohort drawn from the same community as the exposed cohort Ascertainment of exposure written self-report Demonstration that outcome of interest was not present at start of study Yes Comparability Comparability of cohorts on the basis of the design or analysis study controls for gas stove use study controls for additional factors as follows - number of children in household, multifamily dwelling, history of allergies, education, race, state of residence Outcome Assessment of outcome self-report (maternal) Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts complete follow up - all subjects accounted for
Source of funding Comments	Overall level of bias – Low Government: National Institute of Environmental Health Sciences

D.1.111 Triche 2005

Bibliographic reference	Triche E W, Belanger K, Bracken M B, Beckett W S, Holford T R, Gent J F, McSharry J E, and Leaderer B P (2005) Indoor heating sources and respiratory symptoms in non-smoking women. Epidemiology 16(3), 377-384		
Study design	Prospective cohort study		
Objective	To examine the effects of secondary	y heating source use on respiratory symptoms.	
Setting/Study location	United States		
Number of participants	888 mothers of infants		
Selected population	No		
Participant	Description	No. (%)	
characteristics	Sex	888 (100%)	
	Maternal age (years)	Not reported	
	Ethnicity	693 (78%)	
	White or Asian	195 (22%)	

Bibliographic reference	Triche E W, Belanger K, Bracken M B, Beckett W S, Holford T R, Gent J F, McSharry J E, and Leaderer B P (2005) Indoor heating sources and respiratory symptoms in non-smoking women. Epidemiology 16(3), 377-384				
	Black or Hispanic				
	Maternal asthm	a and/or atopic	Not reported	I	
	Parental educat High school or I Some college College graduat	ess	302 (34%) 257 (29%) 329 (37%)		
	Annual family in	icome	Not reported	I	
	Building charac	teristics	Not reported		
Inclusion criteria	Had an infant cl	hild			
Exclusion criteria	Smoking in the household Infant death or adoption Maternal age <19 years Prior participation Plans to move out of study area Having a multiple gestation Having no address or phone number				
Type of pollutant/expo sure	Heating sources: Fireplace (FP) Gas space heater (GH) Kerosene heater (KH) Wood stove (WS)				
Pollutant/expo sure assessment	Palmes tubes were used to passively monitor indoor concentrations of NO ₂ . SO ₂ concentrations were measured using a passive monitor consisting of a 37-mm diameter polystyrene sampling cassette with a washed glass fiber treated filter coated with a 2% sodium carbonate solution placed at the bottom. At home interview, the research assistant placed the monitors in the main living area of the home and instructed respondents on their use. Monitors were exposed in the home for 2 weeks.				
Outcome	Respiratory syn	nptoms			
Results	•	` '	95% confidence in and respiratory s	` '	association
	Lower respiratory symptoms				
		Wheezing	Chest tightness	Laryngitis	Phlegm
		aRR (95%CI)	aRR (95%CI)	aRR (95%CI)	aRR (95%CI)
	Fire place use	1.07 (0.97, 1.18)	1.05 (0.99, 1.12)	1.02 (0.94, 1.10)	1.04 (0.99, 1.09)
	Gas space heater use	1.03 (0.94, 1.13)	1.01 (0.96, 1.07)	0.93 (0.79, 1.10)	0.96 (0.88, 1.05)
	Kerosene Heater Use	1.06 (1.01, 1.11)	1.02 (0.99, 1.05)	1.01 (0.97, 1.04)	0.98 (0.93, 1.03)
	Wood Stove Use	0.97 (0.91, 1.04)	1.01 (0.98, 1.03)	1.00 (0.97, 1.02)	1.00 (0.99, 1.02)

Bibliographic reference	Triche E W, Belanger K, Bracken M B, Beckett W S, Holford T R, Gent J F, McSharry J E, and Leaderer B P (2005) Indoor heating sources and respiratory symptoms in non-smoking women. Epidemiology 16(3), 377-384						
		Upper respiratory symptoms					
		Cough	Runny/stuffy nose	Sore throat			
	Fire place use	1.05 (1.01, 1.09)	0.99 (0.95, 1.04)	1.04 (1.00, 1.08)			
	Gas space heater use	1.00 (0.97, 1.04)	0.99 (0.95, 1.03)	0.99 (0.95, 1.04)			
	Kerosene Heater Use	1.01 (0.99, 1.03)	1.01 (0.99, 1.03)	1.00 (0.97, 1.02)			
	Wood Stove Use	1.01 (0.99, 1.02)	1.01 (0.99, 1.02)	1.00 (0.99, 1.02)			
Follow up	12 months						
Risk of bias (Newcastle- Ottawa Scale)							
Source of	Overall level of bia Government: Nation		mental Health Scienc	es			
funding							
Comments							

D.1.112 Virtanen 2014

Bibliographic reference	Virtanen SM, Takkinen HM, Nwaru BI, et al (2014) Microbial exposure in infancy and subsequent appearance of type 1 diabetes mellitus-associated autoantibodies: a cohort study. JAMA paediatrics 168(8), 755-63
Study design	Prospective cohort study

To investigate whether contacts with animals or other microbial exposit during infancy are associated with the development of clinical or preclitype 1 diabetes Setting/Study location Number of participants Selected population Participant characteristics Male 2646 (52.4%) Female 4497 (47.6%) Age (years) − Maternal < 25 25 − 29 30 − 34 ≥ 35 Ethnicity Education SES (Maternal professional level) None 157 (5.1%) Professional education/course 808 (26.2%) Secondary professional education University Missing 54				
Number of participants				
participants No Participant characteristics Description Sex Male 2646 (52.4%) Female 1497 (47.6%) Age (years) – Maternal < 25				
Participant characteristics Description Sex Male 2646 (52.4%) Female 1497 (47.6%) Age (years) – Maternal 355 (14.8%) 25 – 29 1099 (35.0%) 30 – 34 973 (31.0%) ≥ 35 605 (19.2%) Ethnicity Not reported Education SES (Maternal professional level) None 157 (5.1%) Professional education/course 808 (26.2%) Secondary professional education 1434 (46.4%) University 690(22.3%)				
characteristics Sex Male 2646 (52.4%) Female 1497 (47.6%) Age (years) – Maternal 355 (14.8%) 25 – 29 1099 (35.0%) 30 – 34 973 (31.0%) ≥ 35 605 (19.2%) Ethnicity Not reported Education SES (Maternal professional level) None 157 (5.1%) Professional education/course 808 (26.2%) Secondary professional education 1434 (46.4%) University 690(22.3%)				
Building characteristics Not reported				
Inclusion criteria Infants with increased genetic susceptibility to HLA antigen-DQB1 type diabetes	: 1			
Exclusion criteria Not reported				
Type of Indoor pets pollutant/exposure	Indoor pets			
Pollutant/exposure Questionnaire assessment				
Outcome Clinical or pre-clinical Type 1 diabetes				
Results Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between pets indoor and repeated wheeze Clinical Type 1 diabetes				
aOR (95%CI)				
Indoor dog 0.40 (0.14, 1.14)				
Indoor cat 1.34 (0.58, 3.10)				
Follow up				

Bibliographic reference	Virtanen SM, Takkinen HM, Nwaru BI, et al (2014) Microbial exposure in infancy and subsequent appearance of type 1 diabetes mellitus-associated autoantibodies: a cohort study. JAMA paediatrics 168(8), 755-63
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average infant in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • questionnaire Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for genetic risk • study controls for any additional factor - sex, genetic risk according to HLA-family history of diabetes mellitus mode of delivery place of birth , parental asthma or allergic rhinitis, maternal professional educational level, maternal age, home municipality urbanization level, and the presence of asthma and atopic eczema in the child by the age of 5 year Outcome Assessment of outcome • clinical diagnosis Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall risk of bias: Moderate (concern over self-report of exposure)
Source of funding	Government: Academy of Finland, Centre of Excellence in Molecular Systems Immunology and Physiology Research 2012-17); the Prevaller Consortium; EU Biomed 2 Program Charity: the European Foundation for the Study of Diabetes (EFSD/Novo Nordisk Partnership and ESFD/Juvenile Diabetes Research Foundation/Novo Nordisk Programme); the Foundation for Pediatric Research; the Tampere Tuberculosis Foundation; the Juho Vainio Foundation; the Yrjö Jahnsson Foundation; Medical Research Funds, the Juvenile Diabetes Research Foundation; Novo Nordisk Foundation; Academic:Competitive Research Funding of the Tampere University Hospital Turku University Hospital and Oulu University Hospital;
Comments	

D.1.113 Weinmann 2017

Bibliographic reference	Weinmann T, Gerlich J, Heinrich S et.al (2017) Association of household cleaning agents and disinfectants with asthma in young German adults. Occup Environ Med 2017; 74:684–690.
Study design	Prospective cohort study

Bibliographic reference	Weinmann T, Gerlich J, Heinrich S et.al (2017) Association of household cleaning agents and disinfectants with asthma in young German adults. Occup Environ Med 2017; 74:684–690.				
Objective	To investigate the potential association of the private use of household cleaning sprays and disinfectants with asthma incidence in young adults in the transition from school to working life.				
Setting/Study location	Germany				
Number of participants	1695 young adults				
Selected population	Unclear				
Participant characteristics	Description Sex Female 938 (55.6%) Age (years) Mean (SD) 21.8 (0.75) Ethnicity Not reported Education Not reported SES High 1103 (65.1%)			d d 6)	
Inclusion criteria	Building characteristics		Not reported		
Exclusion criteria	Not reported				
Type of pollutant/exposure	Not reported Domestic use of cleaning: Spray use Disinfectant use				
Pollutant/exposure assessment	Questionnaire				
Outcome	Doctor diagnosis of asthma and either wheezing without cold or use of asthma medication within the last 12 months				
Results	Adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for association between spray/disinfectant use and incident asthma versus never asthma, persistent asthma versus never asthma				
		Incident asth	nma	Persistent asthma	
	Household spray use	aOR (95%0	CI)	aOR (95%CI)	
	Low use	0.70 (0.23, 2	2.06)	0.66 (0.34,1.28)	
	Medium use	0.78 (0.26, 2	•	0.65 (0.32, 1.29)	
	High use	2.79 (0.84, 9	•	0.55 (0.18, 1.63)	
	Disinfectant use	aOR (95%0	•	aOR (95%CI)	
	Low/medium use	1.55 (0.51, 4	•	1.79 (0.82, 3.91)	
	High use	(2.79 1.14, 6	5.83)	1.70 (0.79, 3.65)	
		Incident whe	eezing	Persistent wheezing	
	Household spray use	a OR (95%0	•	aOR (95%CI)	
	Low use	1.53 (0.88, 2	•	1.02 (0.65,1.61)	
	medium use	1.34 (0.75, 2.39)		0.97 (0.60, 1.57)	
		, ,	,	, , ,	

Bibliographic reference	Weinmann T, Gerlich J, Heinrich S et.al (2017) Association of household cleaning agents and disinfectants with asthma in young German adults. Occup Environ Med 2017; 74:684–690.				
	High use	1.71 (0.80, 3.67)	1.24 (0.65, 2.39)		
	Disinfectant use	aOR (95%CI)	aOR (95%CI)		
	Low/ medium use	1.08 (0.60, 1.98)	1.22 (0.74, 2.01)		
	High use	0.79 (0.40, 1.56)	0.98 (0.56, 1.70)		
Follow up	12 months				
	High use 0.79 (0.40, 1.56) 0.98 (0.56, 1.70)				
Source of funding	outcomes) Not reported				
Comments	Hot reported				
Commond					

D.1.114 Weinmayr 2015

Bibliographic reference	Weinmayr G, Hennig F, Fuks K et.al (2015) Long-term exposure to fine particulate matter and incidence of type 2 diabetes mellitus in a cohort study: effects of total and traffic-specific air pollution. Environmental health: a global access science source 14, 53
Study design	Prospective cohort study
Objective	
Setting/Study location	Germany

Bibliographic reference	Weinmayr G, Hennig F, Fuks K et.al (2015) Long-term exposure to fine particulate matter and incidence of type 2 diabetes mellitus in a cohort study: effects of total and traffic-specific air pollution. Environmental health: a global access science source 14, 53				
Number of participants	3607 adults (45–75 years)				
Participant characteristics	Description	No. (%)		No. (%)	
	Sex (male)	1540 (47)		185 (56)	
	Age (years); mean (SD)	58.8 (7.6)		60.5 (7.5)	
	Ethnicity	Not reported		Not reported	
	Cases/selected population	Not selected		Not selected	
	Socio-economic status (e	ducation)			
	Highest: ≥18 years	426 (13)		20 (6)	
	High: 14–17	753 (23)		70 (21)	
	Middle 11–13 years	1801 (55)		209 (63)	
	Low: ≤10 years	327 (10)		33 (10)	
	Building characteristics (dampness and mould)	Not reported		Not reported	
Inclusion criteria	Not reported				
Exclusion criteria	Not reported				
Type of pollutant/exposure	Particulate matter (PM) and road proximity				
Pollutant/exposure assessment	PM ₁₀ and PM _{2.5} concentrations were estimated with the European Air Pollution Dispersion and Chemistry Transport Model (EURAD-CTM) on a spatial resolution of 1 km2 grid cells. As additional traffic exposure we used the distance to the next road with a traffic density higher than the 80 %-percentile (26062 vehicles/day) in the study region				
Outcome	Diabetes incidence				
Results	Adjusted odds ratios (aRRs) and 95% confidence intervals (CIs) for association between distance to major road, PM and diabetes incidence (aRRs are presented for an increase of 1 μ g/m³).				
		·		Diabetes incidence	
			aRR (95%	RR (95%CI)	
	PM ₁₀		1.05 (1.00, 1.10)		
	PM _{2.5} 1.03 (0.95, 1.12)			i, 1.12)	
	Distance to major road (>200 m reference)				
	≤ 100		1.37 (1.04, 1.81)		
	>100-200		0.77 (0.57, 1.04)		
Follow up	Mean follow-up time 5.1 y	ears			
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average population in the community Selection of the non-exposed cohort			the community	
	drawn from the same community as the exposed cohort				

Bibliographic reference	Weinmayr G, Hennig F, Fuks K et.al (2015) Long-term exposure to fine particulate matter and incidence of type 2 diabetes mellitus in a cohort study: effects of total and traffic-specific air pollution. Environmental health: a global access science source 14, 53
	Ascertainment of exposure
	validated measurement used
	Demonstration that outcome of interest was not present at start of study
	• Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	 study controls for age, gender, lifestyle variables, BMI, individual and neighbourhood SES
	Outcome
	Assessment of outcome
	 self-reported physician diagnosis or incident intake of an anti-diabetic drug during follow-up
	Was follow-up long enough for outcomes to occur • Yes
	Adequacy of follow up of cohorts
	subjects lost to follow up unlikely to introduce bias
	Overall risk of bias: low
Source of funding	Government: German Ministry of Education and Science and from the German Research Council
	Charity: Heinz Nixdorf Foundation
Comments	

D.1.115 Wesselink 2017

1000011111K 20 11	VC35CHIR 2017					
Bibliographic reference	Wesselink A K, Carwile J L, Fabian M P et.al (2017) Residential Proximity to Roadways and Ischemic Placental Disease in a Cape Cod Family Health Study. Int. J. Environ. Res. Public Health 2017, 14, 682					
Study design	Retrospective cohort study					
Objective	To examine the association between exposure to traffic-related air pollution and the risk of ischemic placental disease and other obstetric conditions with a placental aetiology.					
Setting/Study location	United states					
Number of participants	3309 pregnant women					
Participant		Ischemic Placental Disease				
characteristics		Yes (n =270)	No (n=3039)			
	Description	No. (%)	No. (%)			
	Sex (male)					
	Age (maternal; years); mean (SD)	26.6 (4.5)	27.7 (4.6)			
	Ethnicity					
	White	256 (94.8)	2956 (97.3)			

Bibliographic reference	Proximity to F	Roadways and	Fabian M P et.al Ischemic Placer Environ. Res. Pu	ntal Disease in	a Cape Cod
	Cases/selected population	d N	ot reported	Not repo	orted
	Socio-economi	ic status (mater	nal education)		
	Less than high	school 2	(0.7)	39 (1.3)	
	High school gra	aduate 5	5 (20.4)	569 (18.	7)
	Some college	10	05 (38.9)	1047 (34	4.5)
	Four-year collegraduate or mo		08 (40.0)	1384 (45	5.5)
	Building charac	cteristics N	ot reported	Not repo	orted
Inclusion criteria	Not reported				
Exclusion criteria	Pregnancies with an unknown outcome Ectopic pregnancies Elective abortions Pregnancy losses at <27 weeks' gestation Multiple births Foetuses with major birth anomalies Pregnancies with an unknown date of the last menstrual period Pregnancies at addresses that could not be geocoded Pregnancies with an incalculable perchloroethylene (PCE) exposure				
Type of pollutant/exposure	Traffic related air pollution				
Pollutant/exposure assessment	Road data were obtained from Topologically Integrated Geographic Encoding and Referencing System (TIGER) files for Barnstable County (which includes Cape Cod) from the 1990 U.S. census website for each of the eight study towns. Major roadways, defined as A1 (primary highways with limited access including roads like interstate highways), A2 (primary roads without limited access like state and local highways that connect cities and towns), and A3 (smaller secondary roads that may connect smaller towns) road segments. Authors used ArcGIS to calculate two metrics of traffic exposure: (a) the shortest Euclidean distance between each residence and the closest major roadway and (b) the length of major roadways within 200 and 500 m buffers around each residence				
Outcome	Preeclampsia, placental abruption, small for gestational age (SGA) and stillbirth				
Results	Adjusted odds ratios (aRRs) and 95% confidence intervals (CIs) for association between traffic exposure and preeclampsia, placental abruption, small for gestational age (SGA) and stillbirth				
		Preeclampsia	Placental Abruption	SGA	Stillbirth
		RR (95%CI)	RR (95%CI)	RR (95%CI)	RR (95%CI)
	Distance from	closest A1-A3	road (m)		
	≥ 200	Reference	Reference	Reference	Reference
	100–199	0.89 (0.37, 2.17)	1.34 (0.54, 3.30)	0.81 (0.55, 1.19)	2.02 (0.65, 6.30)

Bibliographic reference	Wesselink A K, Carwile J L, Fabian M P et.al (2017) Residential Proximity to Roadways and Ischemic Placental Disease in a Cape Cod Family Health Study. Int. J. Environ. Res. Public Health 2017, 14, 682				
	<100	0.46 (0.16, 1.29)	1.75 (0.82, 3.76)	0.91 (0.63, 1.31)	1.71 (0.56, 5.23)
Follow up	Not reported				
Risk of bias (Newcastle-Ottawa Scale)	truly represe Selection of the drawn from the Ascertainment validated measurement validated validat	easurement used self-report that outcome of of cohorts on the self-report at the self-report that outcome of the self-report of the self-report that outcome at the self-report that outcome t	erage population cohort inity as the exponent inity as the exponent interest was not be basis of the degree at pregnancy outcomes to occuts kely to introduce	esed cohort of present at star sign or analysis and year of present cur	egnancy
Source of funding		National Institute nd Human Deve		tal Health, Natio	onal Institute of
Comments					

D.1.116 White 2017

Bibliographic reference	White AJ, and Sandler DP (2017) Indoor Wood-Burning Stove and Fireplace Use and Breast Cancer in a Prospective Cohort Study. Environmental Health Perspectives 125, 1-7
Study design	Prospective cohort study
Objective	To evaluate the risk of breast cancer in relation to indoor heating and cooking practices.
Setting/Study location	United States and Puerto Rico
Number of participants	50,884 women at risk of breast cancer

Bibliographic reference	White AJ, and Sandler DP (2017) Indoor Use and Breast Cancer in a Prospective Health Perspectives 125, 1-7			
Selected population	No			
Participant characteristics	Description Sex Age (years) Mean *SD) at baseline Ethnicity Non-Hispanic white Education Less than high school degree High school degree or equivalent Some college, no degree Associate degree 4-y degree Master's degree Doctoral degree SES (reported as income) <20,000USD 20,000−49,999USD 50,000−99,999USD 100,000−199,999USD ≥200,000 USD Building characteristics	No indoor woodburning stove/fireplace 18; 017 (100%) 54.6 (9.2) 13,543(75.2%) 412 (2.3%) 3,136 (17.4%) 3,949 (21.9%) 2,827 (15.7%) 4,248 (23.6%) 2,884 (16.0%) 558 (3.1%) 1,443 (8.3%) 5,060 (29.0%) 7,299 (41.8%) 3,112 (17.8%) 535 (3.1%) Not reported	Indoor wood- burning stove/fireplace 29; 495 (100%) 55.7 (8.7) 26,274(89.1%) 167 (0.6%) 3,440 (11.7%) 5,308 (18.0%) 3,903 (13.2%) 8,667 (29.4%) 6,592 (22.4%) 1,414 (4.8%) 666 (2.4%) 4,372 (15.5%) 11,343 (40.2%) 9,031 (32.0%) 2,833 (10.0%) Not reported	
Inclusion criteria	No personal history of breast cancer, Living in the United States or Puerto Rico Being between 35–74 y of age Having a sister who had been previously diagnosed with breast cancer			
Exclusion criteria	Not reported			
Type of pollutant/exposu re	Wood burning stove/fireplace Gas stove/fireplace			
Pollutant/expos ure assessment	Questionnaire			
Outcome	Breast cancer			
Results	Adjusted hazard ratios (aHRs) and 95% of association between Indoor heating/cook breast cancer	` ,		
	Breast cancer			
		aHR (95%CI)		
	Indoor wood burning stove/fireplace	1.11 (1.01,1.22)		
	Indoor wood-burning stove/fireplace fuel			
	Fuel - Wood	1.09 (0.98,1.21)		
	Fuel gas	1.15 (1.00,1.32)*		
	Fuel – artificial logs	0.98 (0.85,1.12)		

Bibliographic reference	White AJ, and Sandler DP (2017) Indoo Use and Breast Cancer in a Prospective Health Perspectives 125, 1-7			
	Main source of heating			
	Gas	1.09 (0.98,1.21)		
	Fuel oil	1.13 (0.97,1.32)		
	Propane	0.83 (0.64,1.07)		
	Wood	1.09 (0.82,1.45)		
	Other	0.90 (0.63,1.27)		
Follow up	Mean 64 years			
Risk of bias (Newcastle- Ottawa Scale)	Selection Representativeness of the exposed cohor • truly representative of the average wom Selection of the non-exposed cohort • drawn from the same community as the Ascertainment of exposure • written self-report Demonstration that outcome of interest written self-report Comparability Comparability Comparability Comparability of cohorts on the basis of the • study controls for hormone replacement • study controls for any additional factors annual household income, parity, use of menopause and BMI Outcome Assessment of outcome • record linkage Was follow-up long enough for outcomes • Yes Adequacy of follow up of cohorts • complete follow up - all subjects account Overall risk of bias: Low	exposed cohort as not present at start of study the design or analysis t therapy use at enrolment - race, education, marital status, f oral contraceptives,), age at to occur		
Source of	Government: National Institute of Environ	nmental Health Sciences		
funding				
Comments				

D.1.117 Willers 2006

Bibliographic reference	Willers S M, Brunekreef B, Oldenwening M et.al (2006) Gas cooking, kitchen ventilation, and asthma, allergic symptoms and sensitization in young childrenthe PIAMA study. Allergy 61(5), 563-8
Study design	Prospective cohort study

Bibliographic reference	Willers S M, Brunekreef B, Oldenwening M et.al (2006) Gas cooking, kitchen ventilation, and asthma, allergic symptoms and sensitization in young childrenthe PIAMA study. Allergy 61(5), 563-8			
Objective	To investigate the effect of kitchen ventilation (while cooking) on the relationship between gas cooking, combustion product dispersal, and respiratory and allergic outcomes in children			
Setting/Study location	The Netherlands			
Number of participants	3148 children			
Selected population	Yes – children selected as sample of 'low risk'	s high risk d	lue to maternal	atopy and a random
Participant	Description	No.		%
characteristics	Sex	Not repor	ted	Not reported
	Maternal age (years)	Not repor	ted	Not reported
	Ethnicity	Not repor	rted	Not reported
	(Maintenance) medication use	Not repor	ted	Not reported
	Maternal asthma and/or atopic	Not repor	ted	Not reported
	Parental education	Not reported		Not reported
	Annual family income	Not reported		Not reported
	Building characteristics	Not repor	ted	Not reported
Inclusion criteria	Not reported			
Exclusion criteria	Not reported			
Type of pollutant/exposure	NO ₂ from gas cooking			
Pollutant/exposure assessment	Questionnaire containing questions about cooking and ventilation habits to evaluate the relationships between indoor air pollution, asthma and allergic diseases.			
Outcome	Asthma Wheeze Nasal symptoms Eczema			
Results	•	justed odds ratios (aORs) and 95% confidence intervals (CIs) for sociation between gas cooking and respiratory and atopic outcomes		
			Gas cooking OR (95%CI)	
	Eczema		0.97 (0.74, 1.26)	
	Asthma		1.50 (0.90, 2.49)	
	Nasal symptoms		1.34 (1.06, 1.71)	
	Wheezing		0.99 (0.74, 1.3	32)
Follow up	12 months			
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort truly representative of the average child in the community Selection of the non-exposed cohort			

Bibliographic reference	Willers S M, Brunekreef B, Oldenwening M et.al (2006) Gas cooking, kitchen ventilation, and asthma, allergic symptoms and sensitization in young childrenthe PIAMA study. Allergy 61(5), 563-8
	 drawn from the same community as the exposed cohort Ascertainment of exposure
	 self-report Demonstration that outcome of interest was not present at start of study Yes
	Comparability Comparability of cohorts on the basis of the design or analysis study controls for smoker in the home
	 study controls for other factors as follows - Gender, dampness in the home, allergy or asthma in the parents, presence of older siblings, pets, pregnancy duration, education level of the mother and breast feeding
	Outcome Assessment of outcome
	independent blind assessment
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	• complete follow up - all subjects accounted for □
	Overall level of bias – Moderate (concerns over self-report of exposure)
Source of funding	Industry: Study supported by a grant from Gasuine Trade & supply
Comments	The chance of accumulation of combustion products (CACP) was used besides the distinction between using gas or electricity for cooking and aimed at reducing the misclassification of exposure.
	Authors suggest that study provides only limited evidence that combustion from gas cooking are associated with increased reporting of respiratory and allergic symptoms in young children.

D.1.118 Zhang 2016

Bibliographic reference	Zhang Z, Laden F, Forman J P et.al (2016) Long-Term Exposure to Particulate Matter and Self-Reported Hypertension: A Prospective Analysis in the Nurses' Health Study. Environmental health perspectives 124(9), 1414-20			
Study design	Prospective cohort study			
Objective	To examine the association of hypertension incidence with long-term residential exposures to ambient particulate matter (PM) and residential distance to roadway			
Setting/Study location	United States			
Number of participants	121,700 adult females			
Selected population	No			
Participant	Description	No.	%	
characteristics	Sex	All female	All female	

Bibliographic reference	Zhang Z, Laden F, Forman J P et.al (2016) Long-Term Exposure to Particulate Matter and Self-Reported Hypertension: A Prospective Analysis in the Nurses' Health Study. Environmental health perspectives 124(9), 1414-20			
	Age (years); mean (SD)	60.39 (8.62)		_
	Ethnicity			
	White	71136		95
	Black	749		1
	Asian	749		1
	Other	2995		4
	Cases/selected population	Not reported	d	Not reported
	Socio-economic status (h	usband's edu	cation)	
	Less than high school	2995		4
	High school	20966		28
	More than high school	31450		42
	Building characteristics	Not reported	d l	Not reported
Inclusion criteria	Not reported			
Exclusion criteria	Not reported			
Type of pollutant/exposure	Proximity to traffic and exposures to $PM_{10},PM_{2.5}$, and $PM_{2.5}$ –10.			
assessment	Geographic information system (GIS)-based spatio-temporal models were used to predict monthly exposures to PM ₁₀ and PM _{2.5} for each participant. Authors calculated distance to roads (in meters) for each residential address using GIS (ArcGIS, version 9.2; ESRI). A1 (primary roads, typically interstate highways, with limited access, division between the opposing directions of traffic, and defined exits), A2 (primary major, non-interstate highways and major roads without access restrictions), or A3 (smaller, secondary roads, usually with more than two lanes).			M _{2.5} for each participant. or each residential address y roads, typically interstate e opposing directions of interstate highways and
Outcome	Incident hypertension			
Results	Adjusted hazard ratios (al association between hypermatter exposures	•		e intervals (CIs) for g/m³ increase in particulate
			Incident h	ypertension
			aHR (95%	SCI)
	PM ₁₀		1.02 (1.00), 1.04)
	PM _{2.5} -10		1.03 (1.00, 1.06)	
	PM _{2.5}		1.01 (0.98	3, 1.05)
	Adjusted hazard ratios (aHRs) and 95% confidence intervals (CIs) for association between hypertension and roadway proximity			
	Distance (metres)		Incident h	ypertension
	≥ 200		1.00 (Referent)	
	100–199		0.96 (0.88, 1.05)	
	0–99		1.01 (0.88	3, 1.15)
Follow up	24 months			

	Zhang Z, Laden F, Forman J P et.al (2016) Long-Term Exposure to
Bibliographic reference	Particulate Matter and Self-Reported Hypertension: A Prospective Analysis in the Nurses' Health Study. Environmental health perspectives 124(9), 1414-20
Study methods	Participants were considered to have hypertension if they reported
Study methods	hypertension on the questionnaire ("physician diagnosis of high blood pressure"). In a validation study (n=100) using medical records to confirm systolic or diastolic BP > 140 or > 90 mmHg, respectively, agreement between the medical record and self-report was nearly 100%. Time-varying Cox proportional hazards models were used to model the relationship of incidence of hypertension to roadway proximity and predicted PM _{2.5} , PM ₁₀ , and PM _{2.5} –10 exposure measures adjusting for possible confounders.
Risk of bias	Selection
(Newcastle-Ottawa	Representativeness of the exposed cohort
Scale)	truly representative of the average population in the community
	Selection of the non-exposed cohort
	no description of the derivation of the non-exposed cohort
	Ascertainment of exposure
	validated measurement used Demonstration that outcome of interest was not present at start of study.
	Demonstration that outcome of interest was not present at start of study • Yes
	Comparability
	Comparability of cohorts on the basis of the design or analysis
	study controls for age, race, body mass index (BMI), Dietary, Approaches to Stop Hypertension (DASH) diet score, alcohol consumption, smoking status, physical activity, family history of hypertension, menopausal status, nonnarcotic analgesic intake, statin use, diabetes, individual-level socioeconomic status Outcome
	Assessment of outcome
	record linkage
	• self-report
	Was follow-up long enough for outcomes to occur
	• Yes
	Adequacy of follow up of cohorts
	subjects lost to follow up unlikely to introduce bias
	Overall risk of bias: low
Source of funding	Government: National Institutes of Health Professional: the American Heart Association Academic: China Scholarship Council (CSC), and the Zhejiang University Research Centre for Air Pollution and Health.
Comments	

D.1.119 Zhou 2013

2110u 2013					
Bibliographic reference	Zhou C, Baiz N, Zhang T, et al (2013) Modifiable exposures to air pollutants related to asthma phenotypes in the first year of life in children of the EDEN mother-child cohort study. BMC public health 13, 506				
Study design	Prospective co	hort study			
Objective		To study the impacts of the in utero and first year of life exposures of asthma phenotypes in the first year of life.			
Setting/Study location	France	France			
Number of participants	1,765 mother-o	child pairs			
Selected population	No				
Participant characteristics	Description Sex Male Female Age (years)- M Ethnicity Education (Ma Less than high High school College/Univer Annual family i Low(≤1500) Middle(1501–3 High(>3001)	school sity or more ncome (Euro)	918 647 30.64 (4.81) Not reported 104(5.89) 683(38.70) 950(53.82) 262(14.84) 1001(56.71) 492(27.88)	647 30.64 (4.81) Not reported 104(5.89) 683(38.70) 950(53.82) 262(14.84) 1001(56.71)	
Inclusion criteria	Not reported				
Exclusion criteria	Not reported				
Type of pollutant/exposure	Proximity to traffic Dampness Heating Pets				
Pollutant/exposure assessment	Questionnaire				
Outcome	Asthma, Wheeze Bronchiolitis				
Results	•	` '	5% confidence interval offic and respiratory of	,	
		Asthma	Wheezing	Bronchiolitis	
		aOR (95%CI)	aOR (95%CI)	aOR (95%CI)	
	Traffic- related air pollution	1.71 (1.08, 2.72)	1.47 (1.09,1.97)	1.18 (0.90,1.55)	
	Dampness	2.19 (1.06, 4.53)	2.12 (1.30, 3.46)	1.32 (0.80, 2.18)	

Diblio green bis	pollutants rela	ated to asthma pher	13) Modifiable expo-	ear of life in	
Bibliographic reference	506	e EDEN Motner-chii	d cohort study. BMC	public nealth 13,	
Study design	Prospective cohort study				
	Contact with cats	0.27 (0.08, 0.86)	0.94 (0.61,1.46)	0.69 (0.47,1.03)	
	Domestic wood heating	0.97(0.37,2.50)	0.53 (0.27,1.03)	0.63 (0.38,1.06)	
Follow up					
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average child in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • structured interview Demonstration that outcome of interest was not present at start of study • Yes Comparability Comparability of cohorts on the basis of the design or analysis • study controls for family history • study controls for additional factors including age, gender, family income, maternal age and maternal education Outcome Assessment of outcome • Doctor diagnosed Was follow-up long enough for outcomes to occur • Yes Adequacy of follow up of cohorts				
Source of funding	Overall level of bias – Low Government: Fondation pour la Recherche médicale (FRM); French Ministry of Research: INSERM Nutrition Research Program; French Ministry of Health Perinatality, French Agency for Environmental Security(AFFSET); French National Insitute for Population Health Surveillance (INVS); Paris-sud Unversity; French National Institute for Health Education (INPES); Nestlé, Mutuelle Générale de l'Education Nationale (MGEN); French Speaking Association for the Study of Diabetes and Metabolism (ALFEDIAM); National Agency for Research (ANR) and the fellowship of Erasmus Mundus External Cooperation Window (EM ECW) for China,				
Comments					

D.1.120 Zock 2007

20CK 2007									
Bibliographic reference	Zock JP, Plana E, Ja sprays and adult as journal of respirator	thma: an internat	tional longitudinal	study. American					
Study design	Prospective cohort study								
Objective	To investigate the risk of new-onset asthma in relation to the use of common household cleaners								
Setting/Study location	10 European countrie	es							
Number of participants	3503 adults								
Selected population	No								
Participant characteristics	Description Sex Female Age (years) Mean (ra Ethnicity Education SES Building characteristi	Sex Female 951 (27.1%) Age (years) Mean (range) 42.6 (28 to57) Ethnicity Not reported Education Not reported SES Not reported							
Inclusion criteria	Not reported								
Exclusion criteria	having had nocturnal	People with asthma (those who had reported a history of asthma and/or having had nocturnal attacks of shortness of breath in the last 12 months, and/or wheeze when not having a cold in the last 12 months)							
Type of pollutant/exposure	Household cleaning s	sprays							
Pollutant/exposure assessment	Questionnaire								
Outcome	Physician-diagnosed	d asthma and whee	eze						
Results	Adjusted risk ratios (aRRs), hazard ratios (aHRs) and 95% confidence intervals (CIs) for association between the use of cleaning products at least weekly and the incidence of asthma								
		Asthma attack and/or nocturnal shortness of breath	Current Wheeze	Physician- diagnosed asthma					
		aRR (95%CI)	aRR (95%CI)	aHR (95%CI)					
	Any spray	1.49 (1.12, 1.99)	1.39 (1.06, 1.80)	1.28 (0.78, 2.09)					
	Any perfumed or scented product	1.09 (0.78, 1.50)	1.11 (0.83, 1.49)	1.29 (0.74, 2.26)					
	Frequency of use								
	Use of spray(s) 1 to 3 d/wk	1.36 (0.99,1.89)	1.55 (1.17, 2.06)	0.93 (0.51, 1.67)					
	Use of spray(s) 4 to 7 d/wk	1.75 (1.21,2.54)	1.08 (0.73,1.59)	2.11 (1.15, 3.89)					
	One type of spray used > 1 d/wk	1.37 (0.99, 1.90)	1.25 (0.92, 1.69)	0.97 (0.53, 1.77)					

Bibliographic reference	sprays and adult as	Zock JP, Plana E, Jarvis D, et al (2007) The use of household cleaning sprays and adult asthma: an international longitudinal study. American journal of respiratory and critical care medicine 176(8), 735-41							
	Two types of spray used > 1 d/wk	1.45 (0.92, 2.27)	1.63 (1.10, 2.41)	1.47 (0.70, 3.06)					
	Three or more types of spray used > 1 d/wk	2.40 (1.47, 3.91)	1.80 (1.11, 2.94)	2.96 (1.33, 6.56)					
	Individual products								
	Washing powders	1.10 (0.75, 1.63)	1.28 (0.91, 1.81)	0.82 (0.43, 1.54)					
	Liquid multiuse cleaning products	0.94 (0.64,1.38)	0.97 (0.70, 1.35)	0.98 (0.52, 1.86)					
	Polishes, waxes	1.12 (0.71,1.76)	1.19 (0.77, 1.85)	1.42 (0.68, 2.97)					
	Bleach	1.22 (0.83, 1.80)	1.30 (0.90, 1.87)	1.10 (0.56, 2.17)					
	Ammonia	1.40 (0.87,2.23)	1.31 (0.81, 2.13)	0.92 (0.33, 2.59)					
	Decalcifiers, acids	1.06 (0.70, 1.61)	1.18 (0.77, 1.80)	0.25 (0.06, 1.04)					
	Solvents, stain removers	1.54 (0.94, 2.53)	2.00 (1.30, 3.07)	0.48 (0.12, 1.97)					
	Furniture sprays	1.49 (0.99, 2.23)	1.46 (0.98, 2.19)	2.46 (1.26, 4.80)					
	Glass-cleaning sprays	1.35 (0.98,1.85)	1.49 (1.12, 2.00)	1.43 (0.84, 2.44)					
	Sprays for carpets, rugs, curtains	1.24 (0.47, 3.21)	0.80 (0.26, 2.41)	0.80 (0.11, 5.93)					
	Sprays for mopping the floor	1.05 (0.59, 1.85)	1.03 (0.59,1.79)	0.93 (0.30, 2.85)					
	Oven sprays	0.87 (0.33, 2.28)	1.24 (0.57, 2.69)	0.63 (0.09, 4.64)					
	Ironing sprays	1.66 (0.92, 3.00)	1.05 (0.48, 2.30)	1.51 (0.46, 4.96)					
	Air-refreshing sprays	1.71 (1.22, 2.39)	1.36 (0.98, 1.88)	1.46 (0.78, 2.70)					
Follow up	Mean 8.9 years								
Risk of bias (Newcastle-Ottawa Scale)	Selection Representativeness of the exposed cohort • truly representative of the average adult in the community Selection of the non-exposed cohort • drawn from the same community as the exposed cohort Ascertainment of exposure • questionnaire Demonstration that outcome of interest was not present at start of study • Yes								
	Comparability								

Zock JP, Plana E, Jarvis D, et al (2007) The use of household cleaning **Bibliographic** sprays and adult asthma: an international longitudinal study. American reference journal of respiratory and critical care medicine 176(8), 735-41 Comparability of cohorts on the basis of the design or analysis • study controls for smoking status study controls for any additional factor sex, age, cleaning job, and study center Outcome Assessment of outcome self-report Was follow-up long enough for outcomes to occur Yes Adequacy of follow up of cohorts • complete follow up - all subjects accounted for Overall risk of bias: High r(Concerns over self-report of exposure and outcomes) Source of funding Government: Albacete: Fondo de Investigaciones Santarias (FIS) Hospital Universitario de Albacete, Consejeria de Sanidad; Antwerp: FWO (Fund for Scientific Research)-Flanders Belgium University of Antwerp, Flemish Health Ministry; Barcelona: SEPAR, Public Health Service CIRIT, Red Respira ISCII; Basel: Swiss National Science Foundation, Swiss Federal Office for Education and Science, Swiss National Accident Insurance Fund (SUVA), USC NIEHS; Bergen: Norwegian Research Council, Norwegian Asthma and Allergy Association (NAAF), Glaxo Wellcome AS, Norway Research Fund; Erfurt: GSF-National Research Centre for Environment and Health, Deutsche Forschungsgemeinschaft (DFG); Galdakao: Basque Health Department; Goteborg: Swedish Heart Lung Foundation, Swedish Foundation for Health Care Sciences and Allergy Research, Swedish Asthma and Allergy Foundation, Swedish Cancer and Allergy Foundation; Grenoble: Program Hospitalier de Recherche Clinique-DRC de Grenoble 2000 no. 2610, Ministry of Health, Direction de la Recherche Clinique, Ministere de l'Emploi et de la Solidarite, Direction Generale de la Sante, CHU de Grenoble, Comite des Maladies Respiratoires de l'Isere; Hamburg: GSF-National Research Centre for Environment and Health, DFG; Ipswich and Norwich: Asthma UK (formerly known as National Asthma Campaign); Huelva: FIS Oviedo: FIS); Paris: Ministere de l'Emploi et de la Solidarite, Direction Generale de la Sante, UCB-Pharma (France), Aventis (France), Glaxo France, Program Hospitalier de Recherche Clinique-DRC de Grenoble 2000 no. 2610, Ministry of Health, Direction de la Recherche Clinique, CHU de Grenoble; Pavia: GlaxoSmithKline Italy, Italian Ministry of University and Scientific and Technological Research (MURST), local university funding for research 1998 and 1999 (Pavia, Italy); Tartu: Estonian Science Foundation; Turin: ASL 4 Regione Piemonte (Italy), AO CTO/ICORMA Regione Piemonte (Italy), MURST; GlaxoSmithKline Italy; Umea°: Swedish Heart Lung Foundation, Swedish Foundation for Health Care Sciences and Allergy Research, Swedish Asthma and Allergy Foundation, Swedish Cancer and Allergy Foundation; Uppsala: Swedish Heart Lung Foundation, Swedish Foundation for Health Care Sciences and Allergy Research, Swedish Asthma and Allergy Foundation, Swedish Cancer and Allergy Foundation; Verona: University of Verona; MURST; GlaxoSmithKline Italy. Comments

Appendix E: Forest plots

No forest plots were created for this review.

Appendix F:GRADE tables

F.1 Association between sources of pollutants and health outcomes

F.1.1 Sources of NO₂ and health outcomes

F.1.1.1 Gas heating

Gas nealing									
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Cough									
Gas heating -	infants with pare	ental history of a	llergy						
Roda 2013	Prospective cohort	Serious ^a	NA ^b	Not serious ^c	Serious ^d	None	2898	0.78 (0.56, 1.09)	LOW
Gas heating –	Infants without	parental history	of allergy						
Roda 2013	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	None	2898	1.01 (0.69, 1.46)	LOW
Wheeze									
Gas central he	eating								
De Bilderling 2005	Prospective cohort	Very serious ^e	NAb	Not serious ^c	Serious ^d	None	1868	0.76 (0.47, 1.23)	VERY LOW
Gas fire for he	ating								
De Bilderling 2005	Prospective cohort	Very seriouse	NA ^b	Not serious ^c	Serious ^d	None	1868	0.97 (0.67, 1.39)	VERY LOW
Breast cance	•								
Heating fuel=g	as – Women at	risk of breast ca	ncer						
White 2017	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Not serious ^g	None	50884	aHR 1.15 (1.00,1.32)	HIGH

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Main source	of heating=gas	s – Women at r	isk of breast can	cer					
White 2017	Prospective cohort	Not serious ^f	NA ^b	Not serious ^c	Serious ^d	None	50884	aHR 1.09 (0.98,1.21)	MODERATE

- (a) Serious concerns over self-report of outcomes
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) Serious concerns as findings are not statistically significant (95%Cls cross line of no effect)
 (e) Very serious concerns over self-report of outcome and presence of outcome at 7-8 years of age
 (f) No concerns over risk of bias
- (g) No concerns as findings are statistically significant (95%Cls do not cross line of no effect)

F.1.1.2 Gas space heater

as space ne									
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Cough									
Infants									
Triche 2002	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	890	0.94 (0.75, 1.18)	MODERATE
Mothers of infa	ants								
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	888	1.00 (0.97, 1.04)	MODERATE
Asthma with	wheeze								
McConnell 2002	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	3535	1.20 (0.70, 2.00)	MODERATE
Wheeze	Wheeze								
Infants									

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Triche 2002	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^e	None	890	1.25 (1.05, 1.50)	HIGH
Mothers of inf	ants								
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	888	1.03 (0.94, 1.13)	MODERATE
Chest tightne	ess								
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	888	1.01 (0.96, 1.07)	MODERATE
Laryngitis									
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	888	0.93 (0.79, 1.10)	MODERATE
Phlegm									
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	888	0.96 (0.88, 1.05)	MODERATE
Runny / stuff	y nose								
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	888	0.99 (0.95, 1.03)	MODERATE
Sore throat	Sore throat								
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	888	0.99 (0.95, 1.04)	MODERATE

⁽a) No concerns over risk of bias

⁽b) Not applicable as only one study included (c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%Cls cross line of no effect) (e) No concerns as findings are statistically significant (95%Cls do not cross line of no effect)

F.1.1.3 Gas for cooking

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Asthma	Design	Dias	inconsistency	munectness	Imprecision	Other	Number	Stateu)	Quality
Gas stove									
McConnell 2002	Prospecti ve cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	3535	1.3 (0.80, 2.00)	MODERATE
Casas 2012	Prospecti ve cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	5078	1.33 (0.88, 2.00)	MODERATE
Wheeze									
Gas for cookir	ng								
de Bilderling 2005	Prospecti ve cohort	Very serious ^e	NA ^b	Not serious ^c	Serious ^d	None	1868	1.02 (0.77, 1.36)	VERY LOW
Casas 2012	Prospecti ve cohort	Very serious ^e	NAb	Not serious ^c	Serious ^d	None	5078	1.09 (0.76, 1.57)	VERY LOW
Samet 1993	Prospecti ve cohort	Very serious ^e	NAb	Not serious ^c	Serious ^d	None	1205	0.84 (0.64, 1.09)	VERY LOW
Gas stove (ch	ildren whose	mother had	asthma)						
Belanger 2003	Prospecti ve cohort	Serious ^f	NAb	Not serious ^c	Serious ^d	None	849	1.03 (0.59, 1.79)	LOW
Gas stove (ch	ildren whose	mother did n	ot have asthma)						
Belanger 2003	Prospecti ve cohort	Seriousf	NAb	Not serious ^c	Serious ^d	None	849	1.28 (0.88, 1.86)	LOW
Cough									
Gas cooking v	with hood (da	ily)							
Mommers 2005	Nested case control	Very serious ^e	NAb	Not serious ^c	Serious ^d	None	1191	0.93 (0.64,1.36)	VERY LOW
Gas cooking v	with hood (reg	gularly)							
Mommers 2005	Nested case control	Very serious ^e	NAb	Not serious ^c	Serious ^d	None	1191	1.49 (0.80, 2.78)	VERY LOW

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Gas stove								,	<u> </u>
Samet 1993	Prospecti ve cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	1205	0.94 (0.82, 1.07)	MODERATE
Gas stove (ch	ildren whose	mother had a	asthma)						
Belanger 2003	Prospecti ve cohort	Seriousf	NAb	Not serious ^c	Serious ^e	None	849	0.79 (0.46, 1.36)	LOW
Gas stove (ch	ildren whose	mother did n	ot have asthma)						
Belanger 2003	Prospecti ve cohort	Seriousf	NAb	Not serious ^c	Not serious ^g	None	849	1.52 (1.06, 2.18)	MODERATE
Nasal sympto	oms								
Gas stove									
Willers 2006	Prospecti ve cohort	Seriousf	NA ^b	Not serious ^c	Not serious ^g	None	3148	1.34 (1.06, 1.71)	MODERATE
Respiratory i	llness								
Gas stove									
Samet 1993	Prospecti ve cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	1205	0.98 (0.90, 1.07)	MODERATE
Gas stove (rep	ported as low	er respiratory	/ tract infections)						
Ostro 1993	Prospecti ve cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^g	None	321	1.23 (1.03, 1.47)	HIGH
Gas stove (rep	ported as Up	per respirator	y tract infections)						
Ostro 1993	Prospecti ve cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	321	1.06 (0.94, 1.18)	MODERATE
Eczema									
Gas stove									
Willers 2006	Prospecti ve	Serious ^f	NAb	Not serious ^c	Serious ^d	None	3148	0.97 (0.74, 1.26)	LOW
Shortness of	breath								
Gas stove (mu	ulti-family hou	using)							

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Belanger 2006	Prospecti ve	Serious ^f	NAb	Not serious ^c	Not serious ^g	None	728	2.38 (1.12, 5.06)	MODERATE
Gas stove (sir	ngle-family ho	using)							
Belanger 2006	Prospecti ve	Serious ^f	NAb	Not serious ^c	Serious ^d	None	728	0.91 (0.50, 1.64)	LOW
Chest tightne	ess								
Gas stove (Mi	ulti-family hou	ısing)							
Belanger 2006	Prospecti ve cohort	Serious ^f	NAb	Not serious ^c	Not serious ^g	None	728	4.34 (1.76, 10.69)	MODERATE
Gas stove (Si	Gas stove (Single-family housing)								
Belanger 2006	Prospecti ve cohort	Serious ^f	NAb	Not serious ^c	Serious ^d	None	728	0.68 (0.34, 1.32)	LOW

- (a) No concerns over risk of bias
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) Serious concerns as findings are not statistically significant (95%CIs cross line of no effect
- (e) Very serious concerns over risk of bias due to concerns over self-report of outcomes and of exposure (f) Serious concerns over risk of bias due to self-report of outcomes
- (g) No concerns as findings are statistically significant (95%Cls do not cross line of no effect)

F.1.1.4 Other gas appliance

thor guo upp									
No of studies	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisio n	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Asthma							'		
Home Gas ap	pliance								
Ponsonby 2001	Prospecti ve cohort	Serious ^a	NA ^b	Not serious ^c	Seriousd	None	456	1.30 (0.74, 2.29)	LOW
Cough									
Unvented gas	geyser for wa	ater heating							
Mommers 2005	Nested case control	Very serious ^e	NAb	Not serious ^c	Serious ^d	None	1191	1.74 (0.74, 4.12)	VERY LOW
Vented gas ge	eyser for wate	r heating							
Mommers 2005	Nested case control	Very serious ^e	NAb	Not serious ^c	Serious ^d	None	1191	1.28 (0.85, 1.94)	VERY LOW
Shortness of	breath								
Gas dryer (mu	ulti-family hou	sing)							
Belanger 2006	Prospectiv e cohort	Serious ^a	NA ^b	Not serious ^c	Serious ^d	None	728	2.39 (0.77, 7.43)	LOW
Gas stove (sir	ngle-family ho	using)							
Belanger 2006	Prospectiv e cohort	Serious ^a	NA ^b	Not serious ^c	Serious ^d	None	728	0.91 (0.50, 1.64)	LOW
Chest tightne	ess								
Gas dryer (mu	ulti-family hom	ne)							
Belanger 2006	Prospectiv e cohort	Serious ^a	NA ^b	Not serious ^c	Serious ^d	None	728	1.09 (0.31, 3.90)	LOW
Gas dryer (Sir	ngle-family ho	using)							
Belanger 2006	Prospectiv e cohort	Serious ^a	NA ^b	Not serious ^c	Serious ^d	None	728	1.41 (0.61, 3.26)	LOW

⁽a) Serious concerns over risk of bias due to self-report of outcomes (b) Not applicable as only one study included

- (c) No concerns over directness
- (d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)
 (e) Very serious concerns over risk of bias due to concerns over self-report of outcomes and of exposure

F.1.2 Sources of PM and health outcomes

F.1.2.1 Fireplace

No of								Adjusted relative effect (aOR 95%Cl unless	
studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	stated)	Quality
Cough									
Fireplace for h	eating - Infants								
Triche 2002	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	890	0.99 (0.81, 1.21)	MODERATE
Fireplace for h	eating – Mother	s of infants							
Triche 2005	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Not serious ^e	None	888	1.05 (1.01, 1.09)	HIGH
Wheeze									
Fire place for h	neating - Infants								
Triche 2002	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	890	0.25 (0.04, 1.43)	MODERATE
Fireplace for h	eating – Mother	s of infants							
Triche 2005	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	888	1.07 (0.97, 1.18)	MODERATE
Chest tightne	ss								
Fire place as h	neating – Mother	s of infants							
Triche 2005	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	888	1.05 (0.99, 1.12)	MODERATE
Laryngitis									
Fire place hea	ting as heating -	- Mothers of infa	ints						

No of	D i	District China		Latination		011	Nambar	Adjusted relative effect (aOR 95%Cl unless	0
studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	stated)	Quality
Triche 2005	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Seriousd	None	888	1.02 (0.94, 1.10)	MODERATE
Phlegm									
Fire place as h	eating – Mothe	rs of infants							
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	888	1.04 (0.99, 1.09)	MODERATE
Runny / stuffy	/ nose								
Fire place as h	neating – Mothe	rs of infants							
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	888	0.99 (0.95, 1.04)	MODERATE
Sore throat									
Fire place as h	eating – Mothe	rs of infants							
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not seriouse	None	888	1.04 (1.00, 1.08)	HIGH
Breast cancer	r								
Indoor wood bu	urning stove/fire	eplace – Women	at risk of breast c	ancer					
White 2017	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^e	None	50884	aHR 1.11 (1.01,1.22)	HIGH
Heating fuel=w	vood – Women	at risk of breast	cancer						
White 2017	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	50884	aHR 1.09 (0.98,1.21)	MODERATE
Main source of	f heating=wood	– Women at risl	of breast cancer						
White 2017	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	50884	aHR 1.09 (0.82,1.45)	MODERATE
Wheeze witho	out cold								
PM _{2.5} source f	from residential	wood combustic	n heating						
Pindus 2016	Prospective cohort	Seriousf	NA ^b	Not serious ^c	Serious ^d	None	905	1.14 (0.75, 1.73)	LOW
Allergic rhinit	is								

No of	D	Distriction				Other	Namehan	Adjusted relative effect (aOR 95%Cl unless	0
studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	stated)	Quality
		wood combustio							
Pindus 2016	Prospective cohort	Serious ^f	NAb	Not serious ^c	Not serious ^e	None	905	0.63 (0.42, 0.94)	MODERATE
Breathless									
PM _{2.5} source	from residential	wood combustic	on heating						
Pindus 2016	Prospective cohort	Serious ^f	NAb	Not serious ^c	Seriousd	None	905	0.97 (0.64, 1.48)	LOW
Chest tightne	ess								
PM _{2.5} source	from residential	wood combustic	on heating						
Pindus 2016	Prospective cohort	Serious ^f	NAb	Not serious ^c	Serious ^d	None	905	1.05 (0.72, 1.51)	LOW
Cardiac disea	ise								
PM _{2.5} source	from residential	wood combustic	on heating						
Pindus 2016	Prospective cohort	Serious ^f	NAb	Not serious ^c	Seriousd	None	905	0.92 (0.60, 1.39)	LOW
Hypertension	ı								
PM _{2.5} source	from residential	wood combustic	on heating						
Pindus 2016	Prospective cohort	Serious ^f	NAb	Not serious ^c	Serious ^d	None	905	0.78 (0.54, 1.12)	LOW
Stroke									
PM _{2.5} source t	from residential	wood combustic	on heating						
Pindus 2016	Prospective cohort	Seriousf	NAb	Not serious ^c	Seriousd	None	905	0.85 (0.27, 2.71)	LOW
Heart infarction	on or angina p	ectoris							
		wood combustic	on heating						
Pindus 2016	Prospective cohort	Seriousf	NAb	Not serious ^c	Serious ^d	None	905	0.67 (0.28, 1.56)	LOW
Otitis media									

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Any episode									
Pettigrew 2004 b	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	813	1.14 (0.90, 1.45)	MODERATE
Otitis media, F	Recurrent (4 or n	nore episodes of	otitis media (sepa	rated by at least	: 21 days) in one	year)			
Pettigrew 2004 b	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	813	0.99 (0.58, 1.72)	MODERATE

- (a) No concerns over risk of bias
- (b) Not applicable as only one study included(c) No concerns over directness

- (d) Serious concerns as findings are not statistically significant (95%Cls cross line of no effect) (e) No concerns as findings are statistically significant (95%Cls do not cross line of no effect)
- (f) Serious concerns over risk of bias due to self-report of outcomes

F.1.2.2 Wood Stove

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Otitis media									
Any episode									
Pettigrew 2004 b	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	813	1.22 (0.66, 2.23)	MODERATE
Otitis media, F	Recurrent (4 or m	ore episodes of	otitis media (sepa	rated by at least	21 days) in one	year)			
Pettigrew 2004 b	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	813	1.08 (0.85, 1.38)	MDOERATE

- (a) No concerns over risk of bias
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) Serious concerns as findings are not statistically significant (95%Cls cross line of no effect)

F.1.2.3 Heating or cooking fuel

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Bronchitis (ever		Mok of blus	, cy	mancomess	Imprecision	Other	Humber	Stateay	Quality
East Germany	alagilocca,								
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	28888	1.02 (0.96, 1.09)	MODERATE
West Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^e	None	28888	1.15 (1.00, 1.32)	HIGH
More than 4 cold	s in last 12 mo	onths							
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^e	None	28888	1.13 (1.03, 1.23)	HIGH
West Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	28888	0.96 (0.79, 1.18)	MODERATE
Frequent cough									
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	28888	0.97 (0.86, 1.10)	MODERATE
West Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	28888	0.88 (0.68, 1.15)	MODERATE
Sneeze attacks i	n the last 12 m	nonths							
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	28888	0.92 (0.80, 1.06)	MODERATE
West Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	28888	1.21 (0.88, 1.66)	MODERATE

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Allergy (ever dia	gnosed)								
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	28888	1.07 (0.96, 1.18)	MODERATE
West Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	28888	0.97 (0.79, 1.19)	MODERATE
Eczema (ever dia	agnosed)								
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	28888	0.90 (0.83, 0.98)	HIGH
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^e	None	28888	1.07 (0.87, 1.32)	MODERATE
Overweight (BMI	> kg/m2)								
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	28888	0.89 (0.78, 1.01)	MODERATE
West Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	28888	1.12 (0.86, 1.47)	MODERATE
Heating fuel=woo	d – Women at ı	risk of breast ca	ncer						
White 2017	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	None	50884	aHR 1.09 (0.98,1.21)	MODERATE

⁽a) No concerns over risk of bias

⁽b) Not applicable as only one study included(c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%Cls cross line of no effect)
(e) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

F.1.2.4 Coal heating

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Lower respira	atory tract infec	tions							
Baker 2006	Prospective cohort	Not serious ^e	NA ^b	Not serious ^c	Not serious ^d	None	452	aHR 1.45 (1.07, 1.97)	HIGH

- (a) No concerns over risk of bias
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

F.1.2.5 Artificial logs

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Breast cance	r								
Heating fuel=a	artificial logs – W	omen at risk of	breast cancer						
White 2017	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	50884	aHR 0.98 (0.85,1.12)	MODERATE

- (a) No concerns over risk of bias
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) Serious concerns as findings are not statistically significant (95%CIs cross line of no effect)

F.1.2.6 Fuel oil

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Breast cance	r								
Main source o	f heating=fuel oi	l – Women at ris	sk of breast cancer						
White 2017	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	50884	aHR 1.13 (0.97,1.32)	MODERATE

- (a) No concerns over risk of bias
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) Serious concerns as findings are not statistically significant (95%Cls cross line of no effect)

F.1.2.7 Cooking oil

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Bronchiolitis									
Use of seed o	il for cooking								
Nenna 2017	Case control	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	416	1.82 (1.21; 2.74)	VERY LOW

- (a) Very serious due to concerns over self-report of exposure and outcomes
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

F.1.2.8 Paraffin (Kerosene) heating

•	a. a	Juliu,								
	No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
		200.9	111011 01 10100			prodiction	••		ountou,	4
	Cough									

No of	Docima	Dials of him		In diversion on	Incompanie in m	Othor	Normala are	Adjusted relative effect (aOR 95%Cl unless	Overlife.
studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	stated)	Quality
Triche 2002	iter for heating -		NA ^b	Net corious	Seriousd	Nama	890	1.01 (0.93, 1.10)	MODERATE
Triche 2002	Prospective cohort	Not serious ^a	INA	Not serious ^c	Serious	None	890	1.01 (0.93, 1.10)	MODERATE
Kerosene hea	ter for heating –	Mothers of infar	nts						
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	888	1.01 (0.99, 1.03)	MODERATE
Wheeze									
Kerosene hea	ter for heating -	Infants							
Triche 2002	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	890	0.90 (0.64, 1.25)	MODERATE
Kerosene hea	iter for heating –	Mothers of infar	nts						
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	888	1.06 (1.01, 1.11)	HIGH
Respiratory t	ract infections								
Kerosene hea	ter as heating -	Outcomes repor	ted as Lower respi	iratory tract sym	ptoms				
Li 2006	Prospective cohort	Very serious ^a	NA ^b	Not serious ^c	Serious ^d	None	1137	1.41 (0.96, 2.07)	VERY LOW
Chest tightne	ess								
Kerosene hea	iter as heating –	Mothers of infar	nts						
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	888	1.02 (0.99, 1.05)	MODERATE
Laryngitis									
Kerosene hea	ter – Mothers of	infants							
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	888	1.01 (0.97, 1.04)	MODERATE
Phlegm									
Kerosene hea	iter as heating –	Mothers of infar	nts						
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	888	0.98 (0.93, 1.03)	MODERATE

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Runny / stuffy	y nose								
Kerosene hea	ter as heating –	Mothers of infar	nts						
Triche 2005	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Seriousd	None	888	1.01 (0.99, 1.03)	MODERATE
Sore throat									
Kerosene hea	ter as heating –	mothers of infar	nts						
Triche 2005	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	888	1.00 (0.97, 1.02)	MODERATE
Otitis media									
Any episode									
Pettigrew 2004 b	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	813	0.94 (0.50, 1.78)	MODERATE
Otitis media, F	Recurrent (4 or n	nore episodes of	f otitis media (sepa	arated by at leas	t 21 days) in one	e year)			
Pettigrew 2004 b	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	813	0.91 (0.67, 1.25)	MDOERATE

⁽a) No concerns over risk of bias

⁽b) Not applicable as only one study included (c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%Cls cross line of no effect)
(e) Very serious concerns due to self-report of exposure and outcomes
(f) No concerns as findings are statistically significant (95%Cls do not cross line of no effect)

F.1.3 Sources of allergens

F.1.3.1 Pets

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Wheeze									
Pets at home									
Casas 2012	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	None	5078	1.05 (0.83, 1.33)	LOW
Cat at home									
Herr 2012	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Not serious ^e	None	1879	0.65 (0.47, 0.89)	HIGH
Zhou 2013	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Serious ^d	None	1765	0.94 (0.61,1.46)	MODERATE
Dog in the hor	ne								
Litonjua 2002	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	None	226	0.12 (0.01, 0.97)	MODERATE
Asthma									
Pet ownership	in childhood								
Casas 2012	Prospective cohort	Serious ^a	NA ^b	Not serious ^c	Not serious ^e	None	5078	0.69 (0.52, 0.91)	MODERATE
McConnell 2002	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	None	3535	1.60 (1.00, 2.50)	MODERATE
Contact with c	ats								
Zhou 2013	Prospective cohort	Not seriousf	NAb	Not serious ^h	Not serious ^e	None	1765	0.27 (0.08, 0.86)	HIGH
Exposure to ca	ats (children wit	th wheeze at b	aseline)						
Korppi 2008	Prospective cohort	Serious ^a	NAb	Not serious ^h	Serious ^d	None	100	0.26 (0.03, 2.42)	LOW
Exposure to d	ogs (children wi	i t h wheeze at b	paseline)						

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Korppi 2008	Prospective cohort	Serious ^a	NA ^b	Not serious ^h	Seriousd	None	100	0.20 (0.02, 1.78)	LOW
Any pet (child	ren with wheez	e at baseline)							
McConnell 2002	Prospective cohort	Not serious ^g	NAb	Not serioush	Seriousd	None	3535	1.10 (0.60, 2.00)	MODERATE
Bronchiolitis									
Contact with c	ats								
Zhou 2013	Prospective cohort	Not serious ^g	NAb	Not serious ^c	Seriousd	None	1765	0.69 (0.47,1.03)	MODERATE
Papillary thyr	oid cancer								
Pet ownership	in childhood								
Clarke 2015	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	61799	aRR 0.77 (0.51, 1.17)	LOW
Airway hyper	-responsivene	ess							
Pet ownership									
Hagmolen of Ten Have 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	526	1.17 (0.70, 1.94)	VERY LOW
Type 1 diabet	tes (clinical or	pre-clinical)							
Indoor dog									
Virtanen 2014	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	3143	0.40 (0.14, 1.14)	LOW
Indoor c at									
Virtanen 2014	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	3143	1.34 (0.58, 3.10)	LOW
Irritable bowe	el syndrome								
Pet exposure									
Koloski 2015	Prospective cohort	Not serious ^f	NA ^b	Not serious ^h	Seriousd	None	767	1.47 (0.83, 2.61)	MODERATE

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Herbivore pet								·	
Koloski 2015	Prospective cohort	Not serious ^f	NAb	Not serious ^h	Not seriouse	None	767	2.09 (1.19, 3.67)	HIGH
Carnivore pet									
Koloski 2015	Prospective cohort	Not serious ^f	NAb	Not serious ^h	Seriousd	None	767	1.58 (0.90, 2.76)	MODERATE
Omnivore pet									
Koloski 2015	Prospective cohort	Not serious ^f	NAb	Not serious ^h	Seriousd	None	767	0.97 (0.26, 3.59)	MODERATE
Functional dy	/spepsia								
Any pet expos	ure								
Koloski 2015	Prospective cohort	Not serious ^f	NA ^b	Not serious ^h	Seriousd	None	767	1.69 (0.86, 3.36)	MODERATE
Herbivore pet									
Koloski 2015	Prospective cohort	Not serious ^f	NAb	Not serious ^h	Not seriouse	None	767	2.34 (1.24, 4.45)	HIGH
Carnivore pet									
Koloski 2015	Prospective cohort	Not serious ^f	NAb	Not serious ^h	Not seriouse	None	767	2.04 (1.03, 4.03)	HIGH
Omnivore pet									
Koloski 2015	Prospective cohort	Not serious ^g	NAb	Not serious ^h	Seriousd	None	767	0.98 (0.21, 4.50)	MODERATE
Otitis media									
Any episode									
Pettigrew 2004 b	Prospective cohort	Not serious ^f	NA ^b	Not serious ^c	Serious ^d	None	813	0.76 (0.47, 1.26)	MODERATE
Otitis media, F	Recurrent (4 or	more episodes	of otitis media (se	eparated by at le	east 21 days) in	one year)			
Pettigrew 2004 b	Prospective cohort	Not serious ^f	NA ^b	Not serious ^c	Serious ^d	None	813	1.06 (0.90, 1.25)	MODERATE

- (a) Serious concerns over risk of bias due to concerns over self-report of exposure
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) Serious concerns as findings are not statistically significant (95%Cls cross line of no effect)
- (e) No concerns as findings are statistically significant (95%CIs do not cross the line of no effect)
- (f) No concerns over risk of bias

F.1.3.2 Carpet flooring

zai pet nooriii	9								
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Wheeze									
During pregna	incy								
Franck 2014	Cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	465	5.39 (1.75, 16.54)	HIGH
During first year	ar of life)								
Franck 2014	Cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	465	4.18 (0.40, 43.70)	MODERATE
Obstructive b	ronchitis								
During pregna	incy								
Franck 2014	Cohort	Not serious ^a	NAb	Not serious ^c	Not seriousd	None	465	4.39 (1.01, 19.05)	HIGH

- (a) No concerns over risk of bias
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
- (e) Serious concerns as findings are not statistically significant (95%CIs cross line of no effect)

Second-hand mattress F.1.3.3

Joodina mama									
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Cough									
Used mattress	- infants with pa	arental history of	allergy						
Roda 2013	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^d	None	2898	1.47 (1.00, 2.17)	MODERATE
Used mattress	- infants with no	parental histor	y of allergy						
Roda 2013	Prospective cohort	Serious ^a	NA ^b	Not serious ^c	Serious ^e	None	2898	1.22 (0.80, 1.88)	LOW

- (a) Serious concerns over risk of bias due to self-report of outcomes
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect) (e) Serious concerns as findings are not statistically significant (95%Cls cross line of no effect)

F.1.4 Sources of dampness and health outcomes

F.1.4.1 High air humidity in the bathroom

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Cough									
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	2.30 (2.21, 2.40)	LOW
Nasal symptoms									
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	1.94 (1.88, 2.01)	LOW
Throat symptom	S								

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	3.23 (3.12, 3.25)	LOW
Facial skin symp	otoms								
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	2.42 (2.33, 2.51)	LOW
Headache									
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	3.07 (2.96, 3.17)	LOW
Tiredness									
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	2.16 (2.11, 2.22)	LOW
Eye irritation									
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	2.94 (2.83, 3.05)	LOW

⁽a) Very serious concerns over risk of bias due to self-report of outcomes and exposures

F.1.4.2 Condensation on windows

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Cough									
Engvall 2001	Retrospecti ve cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	9808	2.58 (2.47, 2.70)	LOW
Asthma									

⁽b) Not applicable as only one study included (c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Norback 2013	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriouse	None	7104	aRR 1.07 (0.75, 1.53)	VERY LOW
Asthma and airv	vay hyper-resp	onsiveness							
Norback 2013	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriouse	None	7104	aRR 1.43 (0.67, 3.07)	VERY LOW
Acute respirator	y infection								
Heavy condensat	tion in the room	where child slee	eps at night (not	very often)					
Tin Tin 2016	Prospective cohort	Serious ^f	NA ^b	Not serious ^c	Serious ^e	None	6853	1.00 (0.86, 1.17)	LOW
Heavy condensat	tion in the room	where child slee	eps at night (qui	te often)					
Tin Tin 2016	Prospective cohort	Serious ^f	NAb	Not serious ^c	Seriouse	None	6853	1.05 (0.88, 1.27)	LOW
Heavy condensat	tion in the room	where child slee	eps at night (Alw	ays or almost al	ways)				
Tin Tin 2016	Prospective cohort	Serious ^f	NAb	Not serious ^c	Seriouse	None	6853	1.00 (0.77, 1.31)	LOW
Nasal symptoms	5								
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	2.72 (2.62, 2.81)	LOW
Throat symptom	ıs								
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	3.22 (3.09, 3.35)	LOW
Facial skin sym	otoms								
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	2.11 (2.02, 2.20)	LOW
Headache									
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	3.30 (3.19, 3.43)	LOW
Tiredness									

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	2.19 (2.12, 2.25)	LOW
Eye irritation									
Engvall 2001	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	3.14 (3.01, 3.27)	LOW
Autistic spectrui	n disorders								
Condensation on	windows (1-5	cm) in child's roo	om						
Larsson 2009	Cohort	Not serious ^g	NAb	Not serious ^c	Serious ^e	None	4779	1.35 (0.71, 2.57)	MODERATE
Condensation on	windows (> 5 c	m) in child's roo	m						
Larsson 2009	Cohort	Not serious ^g	NAb	Not serious ^c	Not seriousd	None	4779	2.05 (1.03, 4.10)	HIGH
Condensation on	windows (1-5	cm) in parent's r	oom						
Larsson 2009	Cohort	Not serious ^g	NAb	Not serious ^c	Serious ^e	None	4779	1.52 (0.84, 2.73)	MODERATE
Condensation on	windows (> 5 c	m) in parent's ro	oom						
Larsson 2009	Cohort	Not serious ^g	NA ^b	Not serious ^c	Not serious ^d	None	4779	2.03 (1.08, 3.82)	HIGH

⁽a) Very serious concerns over risk of bias due to self-report of outcomes and exposures

F.1.4.3 Moisture on walls / surfaces

N	lo of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
A	Allergic rhinocon	junctivitis								

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

⁽e) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

⁽f) Serious concerns over risk of bias due to self-report of exposure

⁽g) No concerns over risk of bias

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Ibargoyen- Roteta 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	3360	1.90 (1.01, 3.56)	LOW
Asthma									
Jaakkola 2005	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Serious ^e	None	1916	0.92 (0.54, 1.54)	VERY LOW

- (a) Very serious concerns over risk of bias due to self-report of outcomes and exposures
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
 (e) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

F.1.4.4 History of water leakage

inotory or man								Adjusted relative effect (aOR	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	95%Cl unless stated)	Quality
Cough									
Engvall 2001	Retrospectiv e cohort	Very serious ^a	NAb	Not serious ^c	Not seriousd	None	9808	1.52 (1.44, 1.59)	LOW
Nasal sympto	ms								
Engvall 2001	Retrospectiv e cohort	Very serious ^a	NAb	Not serious ^c	Not seriousd	None	9808	1.36 (1.31, 1.41)	LOW

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Throat sympt	oms								
Engvall 2001	Retrospectiv e cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	2.18 (2.09, 2.28)	LOW
Facial skin sy	mptoms								
Engvall 2001	Retrospectiv e cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	1.56 (1.48, 1.63)	LOW
Headache									
Engvall 2001	Retrospectiv e cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	1.27 (1.21, 1.33)	LOW
Tiredness									
Engvall 2001	Retrospectiv e cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	2.19 (2.12, 2.25)	LOW
Eye irritation									
Engvall 2001	Retrospectiv e cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	1.57 (1.50, 1.65)	LOW

⁽a) Very serious concerns over risk of bias due to self-report of outcomes and exposures
(b) Not applicable as only one study included
(c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

F.1.4.5 Water damage

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Allergic rhinitis									
Jaakkola 2010	Prospective cohort	Very serious ^a	NAb	Not seriouse	Not serious ^d	None	1863	2.06 (1.35, 3.13)	LOW
Asthma									
Jaakkola 2005	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriouse	None	1916	1.01 (0.45, 2.26	VERY LOW
Lower respirator	y illness								
Reported as water	r damage or m	ould / mildew							
Stark 2003	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	499	1.34 (0.99, 1.82)	MODERATE

- (a) Very serious concerns over risk of bias due to self-report of outcomes and exposures
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect) (e) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)
- (f) No concerns over risk of bias

F.1.4.6 Damp condition

Damp Condition									
No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Bronchitis (ever	diagnosed)								
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	28888	1.25 (1.13, 1.37)	HIGH
West Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	28888	1.30 (1.03, 1.65)	HIGH

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
More than 4 cold	ds in last 12 m	onths							
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	28888	1.41 (1.25, 1.60)	HIGH
West Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	28888	1.62 (1.21, 2.17)	HIGH
Frequent cough									
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	28888	1.66 (1.42, 1.95)	HIGH
West Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	28888	2.60 (1.90, 3.55)	HIGH
Sneeze attacks	in the last 12 m	nonths							
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	28888	1.52 (1.26, 1.83)	HIGH
West Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	28888	2.25 (1.52, 3.33)	HIGH
Allergy (ever dia	agnosed)								
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	28888	1.09 (0.93, 1.28)	MODERAT
West Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	28888	1.20 (0.87, 1.66)	MODERATI
Eczema (ever di	agnosed)								

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
East Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	28888	1.15 (1.01, 1.31)	HIGH
West Germany									
Du Prel 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	28888	1.10 (0.77, 1.57)	MODERATE

- (a) No concerns over risk of bias
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
- (e) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

F.1.5 Source of VOCs

F.1.5.1 Parquet flooring

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Wheeze									
During pregna	ncy								
Franck 2014	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	465	5.78 (0.30, 111.08)	MODERATE

- (a) No concerns over risk of bias
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

F.1.5.2 Laminate flooring

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Wheeze									
Laminate floor	ing (during preg	nancy)							
Franck 2014	Cohort	Not serious ^a	NAb	Not serious ^c	Not seriousd	None	465	4.46 (1.01, 19.63)	HIGH
Laminate floor	ing (during 1st y	ear in life)							
Franck 2014	Cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	465	2.44 (0.40, 14.74)	MODERATE

⁽a) No concerns over risk of bias

F.1.5.3 PVC flooring

ve nooning											
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality		
Wheeze											
PVC flooring (during pregnanc	y)									
Franck 2014	Cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	465	24.7 (2.18, 280.39)	HIGH		
PVC flooring (during 1st year i	n life)									
Franck 2014	Cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	465	51.7 (3.21, 833.2)	HIGH		
Asthma											
PVC vs. other	flooring materia	l in child's bedro	om (5 years)								
Shu 200	Prospective cohort	Seriouse	NA ^b	Not serious ^c	Serious ^f	None	3228	1.50 (0.91, 2.47)	LOW		
PVC vs. Wood	d flooring materia	al in child's bedro	oom (5 years)								

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
(e) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Shu 200	Prospective cohort	Serious ^e	NAb	Not serious ^c	Not serious ^d	None	3228	1.54 (1.06, 2.23)	MODERATE
PVC vs. other	flooring materia	al in parent's bed	room (5 years)						
Shu 200	Prospective cohort	Seriouse	NAb	Not serious ^c	Not serious ^d	None	3228	1.71 (1.05, 2.80)	MODERATE
PVC vs. wood	flooring materia	al in parent's bed	lroom (5 years)						
Shu 200	Prospective cohort	Seriouse	NAb	Not serious ^c	Not serious ^d	None	3228	1.60 (1.29, 2.81)	MODERATE
PVC vs. other	flooring materia	al in child's bedro	oom (10 years)						
Shu 200	Prospective cohort	Seriouse	NAb	Not serious ^c	Not serious ^d	None	3228	1.54 (1.06, 2.23)	MODERATE
PVC vs. Woo	d flooring materi	al in child's bedr	oom (10 years)						
Shu 200	Prospective cohort	Serious ^e	NAb	Not serious ^c	Seriousf	None	3228	1.37 (0.92, 2.04)	LOW
PVC vs. other	flooring materia	al in parent's bed	lroom (10 years)						
Shu 200	Prospective cohort	Seriouse	NAb	Not serious ^c	Not serious ^d	None	3228	2.04 (1.41, 2.94)	MODERATE
PVC vs. Woo	d flooring materi	al in parent's be	droom (10 years)						
Shu 200	Prospective cohort	Seriouse	NAb	Not serious ^c	Not serious ^d	None	3228	1.90 (1.29, 2.81)	MODERATE
PVC flooring i	n child's bedroo	m							
Larsson 2010	Cohort	Not serious ^a	NAb	Not serious ^c	Seriousf	None	2779	1.52 (0.99, 2.35)	MODERATE
PVC flooring i	n parent's bedro	oom							
Larsson 2010	Cohort	Not serious ^a	NAb	Not serious ^c	Seriousf	None	2779	1.48 (0.86, 2.57)	MODERATE
Autistic spec	trum disorders								
PVC flooring	n child's bedroo	m							

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Larsson 2009	Cohort	Not serious ^a	NA ^b	Not serious ^c	Seriousf	None	4779	1.19 (0.71, 2.00)	MODERATE
PVC flooring in	n parent's bedro	om							
Larsson 2009	Cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^f	None	4779	1.59 (0.97, 2.61)	MODERATE

- (a) No concerns over risk of bias
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
- (e) Serious concerns over risk of bias due to self-report of outcomes
- (f) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

F.1.5.4 New furniture

new furniture									
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Recurrent wh	eeze								
During pregna	ncy								
Franck 2014	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	465	1.94 (0.72, 5.26)	MODERATE
During 1st year	r of life								
Franck 2014	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	465	2.26 (0.83, 6.17)	MODERATE

- (a) No concerns over risk of bias
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

F.1.5.5 Home products - Air fresheners

			Inconsisten					Adjusted relative effect (aOR 95%Cl unless	
No of studies	Design	Risk of bias	су	Indirectness	Imprecision	Other	Number	stated)	Quality
Asthma									
Any perfumed	or scented prod	luct							
Zock 2007	Prospective cohort	Very serious ^a	NA ^b	Not serious ^c	Serious ^d	None	3503	1.29 (0.74, 2.26)	VERY LOW
Air-refreshing s	prays								
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	3503	1.46 (0.78, 2.70)	VERY LOW
Wheeze									
Air fresheners									
Casas 2013	Prospective cohort	Seriouse	NAb	Not serious ^c	Serious ^d	None	2292	1.09 (0.87, 1.37)	LOW
Air fresheners	during pregnan	cy only							
Casas 2013	Prospective cohort	Seriouse	NAb	Not serious ^c	Serious ^d	None	2292	1.39 (0.85, 2.29)	LOW
Air fresheners	after pregnancy	only							
Casas 2013	Prospective cohort	Seriouse	NAb	Not serious ^c	Not serious ^f	None	2292	1.75 (1.01, 3.04)	MODERATE
Air fresheners	during and after	pregnancy							
Casas 2013	Prospective cohort	Seriouse	NAb	Not serious ^c	Serious ^d	None	2292	1.23 (0.79, 1.93)	LOW
Any perfumed	or scented prod	luct							
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	3503	1.11 (0.83, 1.49)	VERY LOW
Air-refreshing s	prays								

No of Market	P	Piological Control	Inconsisten	L. din dan a		011	N	Adjusted relative effect (aOR 95%Cl unless	0
No of studies	Design	Risk of bias	су	Indirectness	Imprecision	Other	Number	stated)	Quality
Zock 2007	Prospective cohort	Very serious ^a	NA ^b	Not serious ^c	Seriousd	None	3503	1.36 (0.98, 1.88)	VERY LOW
Lower respira	tory tract infec	tions							
Air fresheners									
Casas 2013	Prospective cohort	Seriouse	NAb	Not serious ^c	Not serious ^f	None	2292	1.29 (1.03, 1.63)	MODERATE
Air fresheners	during pregnan	cy only							
Casas 2013	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	None	2292	1.31 (0.77, 2.21)	LOW
Air fresheners	after pregnancy	only							
Casas 2013	Prospective cohort	Seriouse	NA ^b	Not serious ^c	Not serious ^f	None	2292	1.85 (1.04, 3.30)	MODERATE
Air fresheners	during and after	pregnancy							
Casas 2013	Prospective cohort	Seriouse	NA ^b	Not serious ^c	Not serious ^f	None	2292	1.59 (1.00, 2.55)	MODERATE
Diarrhoea									
Air freshener u	se once a week	during pregnancy	/ – Diarrhoea in	infants					
Farrow 2003	Prospective cohort	Seriouse	NAb	Not serious ^c	Not serious ^f	None	13971	1.20 (1.06, 1.35)	MODERATE
Air freshener u	se most days d	uring pregnancy –	Diarrhoea in inf	ants					
Farrow 2003	Prospective cohort	Serious ^e	NAb	Not serious ^c	Seriousd	None	13971	1.10 (0.99, 1.23)	LOW
Air freshener o	nce a week dur	ing pregnancy – C	oiarrhoea in moth	ners 9 to 21 mor	nths after birth				
Farrow 2003	Prospective cohort	Serious ^e	NAb	Not serious ^c	Not serious ^f	None	14541	1.14 (1.00, 1.31)	MODERATE
Air freshener m	nost days during	g pregnancy – Dia	rrhoea in mother	s 9 to 21 month	s after birth				
Farrow 2003	Prospective cohort	Serious ^e	NAb	Not serious ^c	Not serious ^f	None	14541	1.14 (1.01, 1.28)	MODERATE
Vomiting									

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Air freshener u	se once a week	during pregnancy	/ – vomiting in in	nfants					
Farrow 2003	Prospective cohort	Serious ^e	NAb	Not serious ^c	Serious ^d	None	13971	1.06 (0.93, 1.20)	LOW
Air freshener u	se most days d	uring pregnancy –	vomiting in infa	nts					
Farrow 2003	Prospective cohort	Seriouse	NAb	Not serious ^c	Seriousd	None	13971	1.09 (0.97, 1.22)	LOW
Earache									
Air freshener u	se once a week	during pregnancy	/ – earache in in	fants					
Farrow 2003	Prospective cohort	Seriouse	NAb	Not serious ^c	Not serious ^f	None	14541	1.24 (1.02, 1.50)	MODERATE
Air freshener u	se most days d	uring pregnancy –	earache in infa	nts					
Farrow 2003	Prospective cohort	Serious ^e	NAb	Not serious ^c	Not serious ^f	None	14541	1.30 (1.09, 1.54)	MODERATE
Depression									
Air freshener u	se once a week	during pregnancy	/ – depression ir	n mothers 8 mon	ths after birth				
Farrow 2003	Prospective cohort	Seriouse	NAb	Not serious ^c	Seriousd	None	14541	1.11 (0.96, 1.29)	LOW
Air freshener u	se most days d	uring pregnancy –	depression in n	nothers 8 months	s after birth				
Farrow 2003	Prospective cohort	Serious ^e	NAb	Not serious ^c	Not serious ^f	None	14541	1.19 (1.05, 1.36)	MODERATE
Headache									
Air freshener u	se once a week	during pregnancy	/ – headache in	mothers 8 month	hs after birth				
Farrow 2003	Prospective cohort	Serious ^e	NAb	Not serious ^c	Seriousd	None	14541	1.06 (0.94, 1.19)	LOW
Air freshener u	se once a week	during pregnancy	/ – headache in	mothers 9 to 21	months after bir	th			
Farrow 2003	Prospective cohort	Serious ^e	NAb	Not serious ^c	Not serious ^f	None	14541	1.29 (1.14, 1.47)	MODERATE
Air freshener u	se most days d	uring pregnancy –	headache in me	others 8 months	after birth				

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Farrow 2003	Prospective cohort	Seriouse	NAb	Not serious ^c	Not serious ^f	None	14541	1.24 (1.11, 1.38)	MODERATE
Air freshener u	se most days di	uring pregnancy –	headache in mo	others 9 to 21 mo	onths after birth				
Farrow 2003	Prospective cohort	Serious ^e	NA ^b	Not serious ^c	Not serious ^f	None	14541	1.22 (1.09, 1.36)	MODERATE
Cough or cold									
Air freshener u	se once a week	during pregnancy	/ - cough/cold in	mothers 9 to 21	months after bi	rth			
Farrow 2003	Prospective cohort	Seriouse	NAb	Not serious ^c	Serious ^d	None	14541	1.03 (0.87, 1.20)	LOW
Air freshener u	se most days di	uring pregnancy –	cough/cold in m	others 9 to 21 m	onths after birth				
Farrow 2003	Prospective cohort	Seriouse	NAb	Not serious ^c	Serious ^d	None	14541	0.82 (0.72, 0.93)	LOW

⁽a) Very serious risk of bias due to concerns over self-report of outcomes and exposure

F.1.5.6 Home products - Cleaning sprays

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Asthma									
1 type of spray u	sed ≥1 day/wee	k							
Le Moual 2012	Prospective cohort	Very serious ^a	NA ^b	Not serious ^c	Serious ^d	None	683	0.68 (0.44, 1.04)	VERY LOW
≥ 2 types of spra	ys used ≥1 day/	week							

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%CIs cross the line of no effect)

⁽e) Serious risk of bias due to concerns over self-report of outcomes

⁽f) No concerns as findings are statistically significant (95%CIs do not cross the line of no effect)

No of otudios	Decima	Disk of his	Inconsisten	In alive of money		Othor	Number	Adjusted relative effect (aOR 95%Cl unless	Overlife.
No of studies	Design	Risk of bias	су	Indirectness	Imprecision	Other	Number	stated)	Quality
Le Moual 2012	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^e	None	683	1.67 (1.08, 2.56)	LOW
Household spr	ay – Low use								
Weinmann 2017	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	1895	0.70 (0.23, 2.06)	VERY LOW
Household spr	ay – Medium use	e							
Weinmann 2017	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	1895	0.78 (0.26, 2.36)	VERY LOW
Household spr	ay – High use								
Weinmann 2017	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	1895	2.79 (0.84, 9.20)	VERY LOW
Any spray									
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	3503	1.28 (0.78, 2.09)	VERY LOW
Use of spray(s) 1 to 3 d/wk								
Zock 2007	Prospective cohort	Very serious ^e	NAb	Not serious ^c	Seriousd	None	3503	0.93 (0.51, 1.67)	VERY LOW
Use of spray(s) 4 to 7 d/wk								
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not seriouse	None	3503	2.11 (1.15, 3.89)	LOW
Spray use ≥1 c	lay/week								
Bedard 2014	Nested case- control	Serious ^f	NAb	Not serious ^c	Seriousd	None	570	1.45 (0.94, 2.24)	LOW
One type of sp	ray used > 1 d/w	'k							
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	3503	0.97 (0.53, 1.77)	VERY LOW
Two types of s	pray used > 1 d/	wk							
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	3503	1.47 (0.70, 3.06)	VERY LOW

No of studios	Davis.	District China	Inconsisten	L. P.		Oller	Newskar	Adjusted relative effect (aOR 95%Cl unless	Qualita:
No of studies	Design	Risk of bias	су	Indirectness	Imprecision	Other	Number	stated)	Quality
	types of spray					1			
Zock 2007	Prospective cohort	Very serious ^a	NA ^b	Not serious ^c	Not serious ^e	None	3503	2.96 (1.33, 6.56)	LOW
Furniture spray	/S								
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not seriouse	None	3503	2.46 (1.26, 4.80)	LOW
Glass-cleaning	sprays								
Zock 2007	Prospective cohort	Very serious ^a	NA ^b	Not serious ^c	Serious ^d	None	3503	1.43 (0.84, 2.44)	VERY LOW
Sprays for carp	oets, rugs, curta	nins							
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	3503	0.80 (0.11, 5.93)	VERY LOW
Sprays for mor	pping the floor								
Zock 2007	Prospective cohort	Very serious ^a	NA ^b	Not serious ^c	Seriousd	None	3503	0.93 (0.30, 2.85)	VERY LOW
Oven sprays									
Zock 2007	Prospective cohort	Very serious ^e	NA ^b	Not serious ^c	Seriousd	None	3503	0.63 (0.09, 4.64)	VERY LOW
Ironing sprays									
Zock 2007	Prospective cohort	Very serious ^e	NA ^b	Not serious ^c	Seriousd	None	3503	1.51 (0.46, 4.96)	VERY LOW
Wheeze									
Sprays									
Casas 2013	Prospective cohort	Serious ^g	NA ^b	Not serious ^c	Not seriouse	None	2292	1.37 (1.10, 1.69)	MODERATE
Spray during p	regnancy only								
Casas 2013	Prospective cohort	Serious ^g	NA ^b	Not serious ^c	Not seriouse	None	2292	1.62 (1.11, 2.36)	MODERATE
Spray after pre	gnancy only								

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Casas 2013	Prospective cohort	Serious ^g	NA ^b	Not serious ^c	Seriousd	None	2292	1.34 (0.80, 2.24)	LOW
Spray during a	nd after pregna	ancy							
Casas 2013	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^d	None	2292	1.61 (1.08, 2.41)	MODERATE
Household spra	ay - Low use								
Weinmann 2017	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	633	1.53 (0.88, 2.65)	VERY LOW
Household spra	ay -medium us	е							
Weinmann 2017	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	633	1.34 (0.75, 2.39)	VERY LOW
Household spra	ay -High use								
Weinmann 2017	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	633	1.71 (0.80, 3.67)	VERY LOW
Daily use of cle	aning sprays								
Herr 2012	Prospective cohort	Not serious ^g	NAb	Not serious ^c	Seriousd	None	1879	1.50 (0.97, 2.32)	MODERATE
Asthma attack	and/or noctu	rnal shortness of	breath						
Any spray									
Zock 2007	Prospective cohort	Very serious ^a	NA ^b	Not serious ^c	Not seriouse	None	3503	1.49 (1.12, 1.99)	LOW
Use of spray(s) 1 to 3 d/wk								
Zock 2007	Prospective cohort	Very serious ^a	NA ^b	Not serious ^c	Seriousd	None	3503	1.36 (0.99,1.89)	VERY LOW
Use of spray(s) 4 to 7 d/wk								
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not seriouse	None	3503	1.75 (1.21,2.54)	LOW
One type of sp	ray used > 1 d/	wk							

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	3503	1.37 (0.99, 1.90)	VERY LOW
Two types of s	oray used > 1 d/	wk							
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	3503	1.45 (0.92, 2.27)	VERY LOW
Three or more	types of spray u	sed > 1 d/wk							
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^e	None	3503	2.40 (1.47, 3.91)	LOW

- (a) Very serious risk of bias due to concerns over self-report of outcomes and exposure
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)
 (e) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
 (f) Serious risk of bias due to concerns over self-report of exposure
 (g) Serious risk of bias due to concerns over self-report of outcomes

F.1.5.7 Home products - Solvents

nome products - contents											
No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality		
Asthma											
Solvents, stain removers											
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	3503	0.48 (0.12, 1.97)	VERY LOW		

			Inconsisten					Adjusted relative effect (aOR 95%CI unless	
No of studies	Design	Risk of bias	су	Indirectness	Imprecision	Other	Number	stated)	Quality
Wheeze									
Solvents									
Solvents, stain re									
Zock 2007	Prospective cohort	Very serious ^a	NA ^b	Not serious ^c	Not serious ^e	None	3503	2.00 (1.30, 3.07)	LOW
Lower respirato	ry tract infectio	ns							
Spray and solver	nts								
Casas 2013	Prospective cohort	Serious ^f	NAb	Not serious ^c	Not seriouse	None	2292	1.54 (1.11, 2.14)	MODERATE
Solvents									
Casas 2013	Prospective cohort	Seriousf	NAb	Not serious ^c	Seriousd	None	2292	1.19 (0.95, 1.48)	LOW
Wheeze									
Solvent - Expose	ed prenatally, not	exposed postna	tally						
Bajeux 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	1505	0.89 (0.34, 2.31)	VERY LOW
Casas 2013	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	1157	1.04 (0.71, 1.51)	VERY LOW
Solvent - Not exp	posed prenatally,	exposed postna	tally						
Bajeux 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not seriouse	None	1505	1.66 (1.11, 2.47)	LOW
Casas 2013	Prospective cohort	Serious ^f	NAb	Not serious ^c	Seriousd	None	1157	0.87 (0.55, 1.37)	LOW
Solvent - Expose	ed both prenatally	and postnatally							
Bajeux 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not seriouse	None	1505	2.50 (1.45, 4.33)	LOW
Casas 2013	Prospective cohort	Serious ^f	NAb	Not serious ^c	Seriousd	None	1157	1.81 (0.98, 3.37)	LOW
Solvents									

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Casas 2013	Prospective cohort	Seriousf	NAb	Not serious ^c	Not seriouse	None	2292	1.30 (1.03, 1.62)	MODERATE
Solvents, stain re	emovers								
Zock 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^e	None	3503	2.00 (1.30, 3.07)	LOW
Eczema									
Prenatal and not	postnatal solven	t exposure							
Bajeux 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	1505	0.72 (0.35, 1.50)	VERY LOW
Postnatal and no	t prenatal solven	it exposure							
Bajeux 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	1505	1.03 (0.79, 1.36)	VERY LOW
Prenatal and pos	tnatal solvent ex	posure							
Bajeux 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	1505	1.23 (0.84, 1.82)	VERY LOW
Food allergies									
Exposed prenata	lly, not exposed	postnatally							
Bajeux 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	1505	1.25 (0.41, 3.80)	VERY LOW
Not exposed prei	natally, exposed	postnatally							
Bajeux 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriousd	None	1505	1.28 (0.80, 2.03)	VERY LOW
Exposed both pre	enatally and post	tnatally							
Bajeux 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	1505	1.32 (0.71, 2.46)	VERY LOW

⁽a) Very serious risk of bias due to concerns over self-report of outcomes and exposure

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%CIs cross the line of no effect)
(e) No concerns as findings are statistically significant (95%CIs do not cross the line of no effect)
(f) Serious risk of bias due to concerns over self-report of outcomes

F.1.5.8 Home products – Aerosols

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Diarrhoea						'			,
Aerosol use or	ice a week durii	ng pregnancy – [Diarrhoea in infan	ts					
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	13971	1.09 (0.93, 1.28)	LOW
Aerosol use da	ily or most days	s during pregnan	cy – Diarrhoea in	infants					
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not seriouse	None	13971	1.22 (1.09, 1.36)	MODERATE
Vomiting									
Aerosol use or	ice a week durii	ng pregnancy – v	omiting in infants						
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not seriouse	None	14541	1.17 (1.00, 1.37)	MODERATE
Aerosol use da	nily or most days	s during pregnan	cy – vomiting in ir	nfants					
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	None	14541	1.14 (1.02, 1.27)	MODERATE
Earache									
Aerosol use or	ice a week durii	ng pregnancy – e	arache in infants						
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	13971	1.00 (0.78, 1.29)	LOW
Aerosol use da	ily or most day	s during pregnan	cy – earache in ir	nfants					
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	13971	1.05 (0.84, 1.25)	LOW
Depression									
Aerosol use or	ice a week durii	ng pregnancy – c	lepression in mot	hers 8 months at	fter birth				
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	14541	1.06 (0.88, 1.27)	LOW
Aerosol use da	aily or most days	s during pregnan	cy – depression i	n mothers 8 mor	ths after birth				

No of Market	Parallers.	District the same	Inconsisten			011	Nambar	Adjusted relative effect (aOR 95%CI unless	0.21%
No of studies	Design	Risk of bias	су	Indirectness	Imprecision	Other	Number	stated)	Quality
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	serious ^d	None	14541	1.03 (0.91, 1.17)	LOW
Headache									
Aerosol use or	nce a week durin	g pregnancy – h	eadache in mothe	ers 8 months aft	er birth				
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	None	14541	1.16 (1.00, 1.35)	MODERATE
Aerosol use or	ice a week durin	g pregnancy – h	eadache in mothe	ers 9 to 21 mont	hs after birth				
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not seriouse	None	14541	1.35 (1.15, 1.59)	MODERATE
Aerosol use m	ost days during p	oregnancy – hea	dache in mothers	8 months after	birth				
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	None	14541	1.25 (1.13, 1.39)	MODERATE
Aerosol use m	ost days during p	oregnancy – hea	dache in mothers	9 to 21 months	after birth				
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not seriouse	None	14541	1.21 (1.10, 1.34)	MODERATE
Influenzas									
Aerosol use or	nce a week durin	g pregnancy –In	fluenza in mother	s 9 to 21 months	s after birth				
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	None	14541	1.03 (0.85, 1.24)	LOW
Aerosol use me	ost days during p	oregnancy – influ	uenza in mothers	9 to 21 months	after birth				
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not seriouse	None	14541	0.87 (0.77, 0.99)	MODERATE
Urinary tract i	nfection								
Aerosol use or	nce a week durin	g pregnancy –U	TI in mothers 9 to	21 months after	r birth				
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	None	14541	1.16 (0.89, 1.52)	LOW
Aerosol use m	ost days during p	oregnancy – UTI	in mothers 9 to 2	1 months after b	oirth				
Farrow 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not seriouse	None	14541	1.23 (1.04, 1.45)	MODERATE

- (a) Serious risk of bias due to concerns over self-report of outcomes
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect) (e) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

F.1.5.9 Paint

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Congenital an	omalies								
All congenital r	malformations - F	Paint fumes in th	e residence durin	g the 1st trimest	er of pregnancy				
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	20103	0.95 (0.74, 1.21)	MODERATE
Nervous system	m - Paint fumes	in the residence	during the 1st trin	nester of pregna	incy				
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	20103	2.19 (0.76, 6.32)	MODERATE
Eye - Paint fun	nes in the reside	nce during the 1	st trimester of pre	gnancy					
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	20103	1.79 (0.70, 4.57)	MODERATE
Ear, face and r	neck - Paint fume	es in the residen	ce during the 1st	trimester of preg	ınancy				
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	20103	2.15 (0.84, 5.55)	MODERATE
Congenital hea	art defects - Pain	t fumes in the re	sidence during th	e 1st trimester o	f pregnancy				
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	20103	0.76 (0.39, 1.49)	MODERATE
Respiratory sys	stem - Paint fum	es in the resider	nce during the 1st	trimester of preg	gnancy				

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Seriousd	None	20103	1.13 (0.27, 4.79)	MODERATE
Cleft lip and cle	eft palate - Pair	nt fumes in the res	sidence during the	e 1st trimester of	pregnancy				
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	20103	1.06 (0.33, 3.46)	MODERATE
Digestive syste	em - Paint fume	es in the residence	e during the 1st tr	imester of pregn	ancy				
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	20103	0.61 (0.15, 2.50)	MODERATE
Renal - Paint f	umes in the res	sidence during the	1st trimester of p	oregnancy					
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not seriouse	None	20103	2.16 (1.02, 4.58)	HIGH
Genital - Paint	fumes in the re	sidence during th	e 1st trimester of	pregnancy					
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	20103	0.83 (0.48, 1.43)	MODERATE
Limb defects -	Paint fumes in	the residence dur	ring the 1st trimes	ster of pregnancy	/				
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Seriousd	None	20103	0.82 (0.54, 1.24)	MODERATE
Muscula and s	keletal - Paint f	umes in the resid	ence during the 1	st trimester of p	regnancy				
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Seriousd	None	20103	1.77 (0.75, 4.16)	MODERATE
Other malform	ation - Paint fur	mes in the resider	nce during the 1st	trimester of pre	gnancy				
Hjortebjerg 2012	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	20103	1.24 (0.62, 2.46)	MODERATE
Obstructive b	ronchitis in fir	st year of life (Pl	nysician diagnos	sis)					
Painting during	g pregnancy								
Franck 2014	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not seriouse	None	465	5.46 (1.09, 27.20)	HIGH
Recurrent wh	eeze								
Painting during	g pregnancy								

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Franck 2014	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	465	2.35 (0.89, 6.20)	MODERATE
Painting during	1 st year in life								
Franck 2014	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	465	2.53 (0.85, 7.49)	MODERATE
Small for gest	ational age								
Exposure to pa	int fumes during	g pregnancy							
Sorensen 2010	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^e	None	19000	0.89 (0.81, 0.98)	HIGH
Preterm birth									
Exposure to pa	int fumes during	g pregnancy							
Sorensen 2010	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	19000	0.95 (0.82, 1.11)	MODERATE

⁽a) No concerns over risk of bias

F.1.5.10 Any type of redecoration

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Recurrent w	heeze								

⁽b) Not applicable as only one study included(c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect) (e) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Any type of re-	decoration during	g pregnancy							
Franck 2014	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	465	2.04 (0.78, 5.28)	MODERATE
Any type of re	decoration during	g 1 st year of life							
Franck 2014	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	None	465	1.89 (0.71, 5.06)	MODERATE
Redecoration	after birth in first	18th months of	life						
Herr 2012	Prospective cohort	Not serious ^e	NA ^b	Not serious ^c	Serious ^d	None	1879	1.22 (0.96, 1.54)	MODERATE
Pulmonary in	fections								
Restoration (p	arent report)								
Diez 2002	Nested case- control	Serious ^e	NA ^b	Not serious ^c	Not serious ^f	None	475	5.6 (1.3, 24.0)	LOW

⁽a) No concerns over risk of bias

F.1.6 Building characteristics and health outcomes

F.1.6.1 Building age

		Risk of						Adjusted relative effect (aOR 95%Cl unless	
No of studies	Design	bias	Inconsistency	Indirectness	Imprecision	Other	Number	stated)	Quality
Recurrent whee	eze								
1940–75									

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

⁽e) Serious concerns over risk of bias due to self-report of outcomes

⁽f) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Emenius 2003	Nested case control	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	540	1.69 (1.01, 2.89)	MODERATE
1975 onwards									
Emenius 2003	Nested case control	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	540	1.86 (1.05, 3.27)	MODERATE

⁽g) No concerns over risk of bias

F.1.6.2 Dwelling size

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Lower respira	atory tract infec	tions							
Per room incre	ease in the hous	ehold – Outcom	e reported as Low	er respiratory tra	ct infections				
Li 2006	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	1137	0.99 (0.92, 1.06)	VERY LOW

⁽a) Very serious concerns over risk of bias due to self-report of outcomes and exposures

⁽a) Not applicable as only one study included

⁽b) No concerns over directness

⁽c) No concerns as findings are statistically significant (95%CIs do not cross the line of no effect)

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

F.1.6.3 Central air-conditioning

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Asthma									
Reponen 2011	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	176	0.3 (0.14, 0.83)	HIGH
Otitis media									
Any episode									
Pettigrew 2004 b	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^e	None	813	0.52 (0.27, 1.03)	MODERATE
Otitis media, F	Recurrent (4 or n	nore episodes of	f otitis media (sepa	rated by at leas	t 21 days) in one	e year)			
Pettigrew 2004 b	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriousd	None	813	0.93 (0.77, 1.11)	MDOERATE

⁽a) No concerns over risk of bias

F.1.6.4 Ventilation rate

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Asthma and a	allergic sympto	ms							
Ventilation rate	e - Third quartile	vs. fourth quart	ile						
Bornehag 2005	Nested case- control	Serious ^a	NA ^b	Not serious ^c	Serious ^d	None	400	1.17 (0.57, 2.42)	LOW
Ventilation rate	e - Second quart	tile vs. fourth qua	artile						
Bornehag 2005	Nested case- control	Serious ^a	NA ^b	Not serious ^c	Serious ^d	None	400	1.35 (0.66, 2.74)	LOW

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

⁽e) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Ventilation rate	e - First quartile	vs. fourth quartil	е						
Bornehag 2005	Nested case- control	Serious ^a	NA ^b	Not serious ^c	Serious ^d	None	400	1.95 (0.94, 4.04)	LOW

⁽a) Serious concerns over risk of bias due to self-report of outcomes(b) Not applicable as only one study included(c) No concerns over directness

F.1.6.5 Proximity to traffic - Traffic intensity

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Bronchiolitis									
≥8640 cars/day	≤100 m, birth ad	ldress							
Lindgren 2013	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	7898	aHR 0.7 (0.6, 0.9)	HIGH
≥8640 cars/day	≤100 m, birth ad	ldress (never r	moved)						
Lindgren 2013	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	7898	aHR 0.7 (0.6, 0.9)	HIGH
Obstructive bro	onchitis								
≥8640 cars/day	≤100 m, birth ad	Idress							
Lindgren 2013	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	7898	aHR 1.0 (0.9,1.2)	MODERATE
≥8640 cars/day	≤100 m, birth ad	ldress (never r	moved)						
Lindgren 2013	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^e	None	7898	aHR 1.0 (0.8,1.2)	MODERATE
Asthma									
≥8640 cars/day	≤100 m, birth ad	Idress							

⁽d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Lindgren 2013	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	7898	aHR 0.7 (0.6, 0.9)	HIGH
≥8640 cars/day	≤100 m, birth ad	ldress (never i	moved)						
Lindgren 2013	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	7898	aHR 0.7 (0.6, 0.9)	HIGH

⁽a) No concerns over risk of bias

F.1.6.6 Located within 50 m of major traffic

Quality asse	ssment							Adjusted relative	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	effect (aOR 95%CI unless stated)	Quality
Wheeze									
Morgenster n 2007	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	none	3577	1.14 (0.92, 1.42)	LOW
Garshick 2003	Prospective cohort	Seriousª	NAb	Not serious ^c	Not seriouse	none	2628	1.31 (1.00, 1.71)	MODERATE
Anxiety sym	ptoms								
Power 2015	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	none	71 271	1.01 (0.95, 1.08)	MODERATE
Cough withou	ut infection								
Morgenster n 2007	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	none	3577	0.74 (0.55, 1.00)	LOW
Chronic cou	gh								

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
(e) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

Quality asse	ssment							Adjusted relative	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	effect (aOR 95%CI unless stated)	Quality
Garshick 2003	Prospective cohort	Seriousª	NAb	Not serious ^c	Seriousd	none	2628	1.24 (0.92, 1.68)	LOW
Chronic phle	egm								
Garshick 2003	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	none	2628	1.18 (0.88, 1.56)	LOW
Dry cough a	t night								
Morgenster n 2007	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	none	3577	0.84 (0.61, 1.16)	LOW
Asthmatic/s	pastic/ obstruc	tive bronchi	tis						
Morgenster n 2007	Prospective cohort	Seriousª	NAb	Not serious ^c	Serious ^d	none	3577	1.12 (0.88, 1.44)	LOW
Respiratory	infections								
Morgenster n 2007	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	none	3577	1.03 (0.86, 1.23)	LOW
Sneezing, ru	inny/stuffed no	se							
Morgenster n 2007	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	none	3577	1.10 (0.87, 1.39)	LOW
Asthma									
Morgenster n 2008	prospective cohort	Serious ^a	NAb	Not serious ^c	Not seriouse	none	5921	1.66 (1.01, 2.59)	MODERATE
Sbihi 2016	prospective - cohort	Not serious ^f	NAb	Not serious ^c	Not serious ^e	none	68195	1.25 (1.04, 1.49)	HIGH
Asthma requ	uiring hospitali	sations							
Chang 2009	prospective - cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	none	3297	aHR 1.11 (0.92, 1.33)	MODERATE
Hay fever									
Morgenster n 2008	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	none	5921	1.16 (0.67, 2.00)	LOW
Eczema									

Quality asse	essment				Adjusted relative				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	effect (aOR 95%CI unless stated)	Quality
Morgenster n 2008	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	none	5921	0.96 (0.72, 1.11)	LOW
Coronary Ho	eart Disease (C	HD) Mortalit	y/sudden cardiac	death					
Gan 2010	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Not serious ^e	none	450,283	aRR 1.29 (1.18, 1.41)	HIGH
Hart 2014	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Not serious ^e	none	107130	aHR 1.38 (1.04, 1.82)	HIGH
Obesity									
Li 2016	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	none	2372	1.10 (0.97; 1.25)	MODERATE
Lung cance	r incidence								
Puett 2014	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Not serious ^e	none	103,650	aHR 2.01 (1.06, 3.80)	HIGH
Uterine leio	myomata								
Within 50 metres of an US A1- A3 roadway									
Mahalingai ah 2014	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	none	85251	1.01 (0.93, 1.09)	MODERATE

⁽a) Serious concerns over risk of bias due to self-report of outcomes

F.1.6.7 Located within 75 m of major traffic

No of Risk of Inconsistenc effect (aOR 95%CI	Quality asse	ssment			Adjusted relative					
		Design		Inconsistenc y	Indirectness	Imprecision	Other	Number	effect (aOR 95%CI	Quality

⁽b) Not applicable as only one study included(c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)
(e) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
(f) No concerns over risk of bias

Quality asse	essment							Adjusted relative	
No of studies	Design	Risk of bias	Inconsistenc y	Indirectness	Imprecision	Other	Number	effect (aOR 95%CI unless stated)	Quality
McConnell 2006	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	none	5341	1.29 (1.01, 1.66)	HIGH
Wheeze									
McConnell 2006	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Not serious ^d	none	5341	1.40 (1.09, 1.78)	HIGH

⁽a) No concerns over risk of bias

F.1.6.8 Located within 100 m of major traffic

Quality asse	ssment							Adjusted relative	
No of studies	Design	Risk of bias	Inconsistenc y	Indirectness	Imprecision	Other	Number	effect (aOR 95%CI unless stated)	Quality
Chronic lung	allograft dysfund	tion (CLAD)							
Bhinder 2014	Retrospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^d	none	397	aHR 4.72 (2.13, 10.47)	MODERATE
Coronary arte	ery calcification (CAC)							
Hoffmann 2007	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Not serious ^d	none	4494	1.45 (1.15, 1.82)	HIGH
Asthma									
Rice 2015	Prospective cohort	Seriousf	NAb	Not serious ^c	Serious ^g	none	6,339	1.18 (0.95, 1.46)	LOW
Wheeze									
Rice 2015	Prospective cohort	Serious ^f	NAb	Not serious ^c	Serious ^g	none	6,339	1.02 (0.84, 1.25)	LOW
Chronic coug	h								

⁽b) Not applicable as only one study included (c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

Quality asse	essment							Adjusted relative	
No of studies	Design	Risk of bias	Inconsistenc y	Indirectness	Imprecision	Other	Number	effect (aOR 95%CI unless stated)	Quality
Rice 2015	Prospective cohort	Seriousf	NAb	Not serious ^c	Serious ^g	none	6,339	1.22 (0.89, 1.66)	LOW
Diabetes inci	dence								
Weinmayr 2015	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Not serious ^d	none	3607	aRR 1.37 (1.04, 1.81)	HIGH
Preeclampsia	а								
Wesselink 2017	Retrospective cohort	Seriousf	NAb	Not serious ^c	Serious ^g	none	3309	aRR 0.46 (0.16, 1.29)	LOW
Placental Ab	ruption								
Wesselink 2017	Retrospective cohort	Seriousf	NAb	Not serious ^c	Serious ^g	none	3309	aRR 1.75 (0.82, 3.76)	LOW
Small for ges	stational age (SG	A)							
Wesselink 2017	Retrospective cohort	Seriousf	NAb	Not serious ^c	Serious ^g	none	3309	aRR 0.91 (0.63, 1.31)	LOW
Stillbirth									
Wesselink 2017	Retrospective cohort	Seriousf	NAb	Not serious ^c	Serious ^g	none	3309	aRR 1.71 (0.56, 5.23)	LOW
Incident hype	ertension								
Zhang 2016	Prospective cohort	Not serious ^e	NA ^b	Not serious ^c	Serious ^g	none	121,700	aHR 1.01 (0.88, 1.15)	MODERATE

⁽a) Serious concerns over risk of bias due to small sample size

⁽b) Not applicable as only one study included(c) No concerns over directness

⁽d) No concerns over directiness
(d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
(e) No concerns over risk of bias
(f) Serious concerns over risk of bias due to self-report of outcomes
(g) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

F.1.6.9 Located within 150 m of major traffic

Quality asse	essment							Adjusted relative	
No of studies	Design	Risk of bias	Inconsistenc y	Indirectness	Imprecision	Other	Number	effect (aOR 95%Cl unless stated)	Quality
Wheezing in	the last year								
Pujades- Rodriguez 2009	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	none	2644	0.86 (0.68, 1.08)	LOW
COPD									
Pujades- Rodriguez 2009	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	none	2644	0.97 (0.68, 1.37)	LOW
Bronchial hyp	per responsivene	ess							
Pujades- Rodriguez 2009	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	none	2644	0.92 (0.68, 1.24)	LOW
Allergic sens	itisation								
Pujades- Rodriguez 2009	Prospective cohort	Serious ^a	NA ^b	Not serious ^c	Serious ^d	none	2644	0.87 (0.70, 1.07)	LOW
Asthma									
Sbihi 2016	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Serious ^d	Pre-school age	68195	1.03 (0.98, 1.09)	MODERATE
Sbihi 2016	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Serious ^d	School age	68195	1.04 (0.92, 1.16)	MODERATE

⁽a) Serious concerns over risk of bias due to self-report of outcomes

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

⁽e) No concerns over risk of bias

F.1.6.10 Located within 200 m of major traffic

Quality asse	essment							Adjusted relative	
No of studies	Design	Risk of bias	Inconsistenc y	Indirectness	Imprecision	Other	Number	effect (aOR 95%CI unless stated)	Quality
Infertility									
Mahalingai ah 2016	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	none	36294	aHR 1.11 (1.02, 1.20)	HIGH
Asthma									
Bowatte 2017	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^e	none	1405	1.21 (0.91, 1.59)	MODERATE
Wheeze									
Bowatte 2017	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Not serious ^d	none	1405	1.38 (1.06, 1.80)	HIGH

⁽a) No concerns over risk of bias

F.1.7 Individual characteristics and health outcomes

F.1.7.1 Tenancy status

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Cough									
Rented vs owr	ner								

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

⁽e) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Engvall 2010	Retrospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	None	7640	1.85 (0.94, 3.65)	LOW
Eye irritation									
Rented vs owr	ner								
Engvall 2010	Retrospective cohort	Serious ^a	NAb	Not serious ^c	Not seriouse	None	7640	2.07 (1.19, 3.58)	MODERATE
Nasal irritatio	n								
Rented vs owr	ner								
Engvall 2010	Retrospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	None	7640	2.07 (1.33, 3.20)	MODERATE
Throat irritation	on								
Rented vs owr	ner								
Engvall 2010	Retrospective cohort	Serious ^a	NA ^b	Not serious ^c	Serious ^d	None	7640	1.98 (0.98, 3.97)	LOW

⁽a) Serious concerns over risk of bias due to self-report of outcomes

F.1.7.2 Household occupant density

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Bronchiolitis									
Number of col	nabitants ≥ 4								
Nenna 2017	Case control	Very serious ^a	NAb	Not serious ^c	Not seriousd	None	416	1.75 (1.36; 2.13)	VERY LOW

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%Cls cross line of no effect) (e) No concerns as findings are statistically significant (95%Cls do not cross line of no effect)

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Newly diagno	sed asthma								
Multi-family ho	ouse								
Larsson 2010	Cohort	Not serious ^e	NA ^b	Not serious ^c	Serious ^f	None	2779	1.48 (0.86, 2.57)	MODERATE

⁽a) Very serious concerns over risk of bias due over self-report of exposure and outcomes

F.1.7.3 Socio-economic status

30CiO-econon	iic status								
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Cough									
Middle vs high	n socio-economic	c status							
Mommers 2005	Nested case control	Very serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	1191	1.53 (1.12, 2.10)	VERY LOW
Low vs high s	ocio-economic s	tatus							
Mommers 2005	Nested case control	Very serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	1191	3.37 (2.01, 5.71)	VERY LOW

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect) (e) No concerns over risk of bias

⁽f) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Asthma									
Middle vs high	socio-economic	status – outcor	ne reported as ast	hmatic symptom	ıs				
Mommers 2005	Nested case control	Very serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	1191	1.43 (1.00, 2.04)	VERY LOW
Middle vs high	socio-economic	status – outcor	ne reported as ast	hmatic symptom	ıs				
Mommers 2005	Nested case control	Very serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	1191	3.32 (1.88, 5.93)	VERY LOW

⁽a) Serious concerns over risk of bias due over self-report of outcomes

F.2 Association between level of exposure to pollutants and health outcomes

F.2.1 Dampness / Mould

F.2.1.1 Damp

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Cough									
Brunekreef 1989	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	4625	2.16 (1.64, 2.84)	LOW
Reported as slight	to moderate d	ampness							

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%CIs do not cross the line of no effect)

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Cable 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Serious ^e	None	7320	aRR 0.85 (0.67, 1.09)	VERY LOW
Reported as mark	ed dampness								
Cable 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriouse	None	7320	aRR 1.26 (0.80, 1.99)	VERY LOW
Major moisture da	mage or any m	noisture damage	with visible mo	uld in child's mai	in living area				
Karvonen 2015	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Seriouse	None	398	1.27 (0.77, 2.09)	MODERATE
Cough and phleg	jm								
Slight to moderate	dampness								
Cable 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^e	Not serious ^d	None	7320	1.24 (0.99, 1.56)	LOW
Marked dampness	6								
Cable 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^e	Not serious ^d	None	7320	2.73 (1.88, 3.99)	LOW
Asthma									
Brunekreef 1989	Prospective cohort	Not serious ^f	NAb	Not serious ^e	Not serious ^d	None	4625	1.42 (1.04, 1.94)	HIGH
Casas 2012	Prospective cohort	Not serious ^f	NAb	Not seriouse	Seriouse	None	5078	1.16 (0.87, 1.53)	MODERATE
Zhou 2013	Prospective cohort	Not serious ^f	NAb	Not serious ^e	Not serious ^d	None	1765	2.19 (1.06, 4.53)	HIGH
Norback 2013	Prospective cohort	Not serious ^f	Do	Not serious ^e	Not serious ^d	None	7104	aRR 1.49 (1.00, 2.22)	HIGH
Reported as damp	oness and mou	ıld							
Dales 1991	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	14799	1.56 (1.25, 1.95)	LOW
Minor moisture da	mage with or w	vithout mould sp	ots in child's ma	ain living area					
Karvonen 2015	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Seriouse	None	398	1.31 (0.72, 2.36)	MODERATE

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Major moisture da	amage or any m	noisture damage	with visible mo	uld in child's mai	_	'			
Karvonen 2015	Prospective cohort	Not serious ^g	NAb	Not serious ^c	Seriouse	None	398	1.33 (0.60, 2.98)	MODERATE
Airway hyper-res	sponsiveness								
Hagmolen of Ten Have 2007	Prospective cohort	Very serious ^a	NAb	Not serious ^e	Not serious ^d	None	528	3.95 (1.82, 8.57)	LOW
Allergic rhinitis									
Water damage or	mould/mildew	in year							
Stark 2005	Prospective cohort	Not serious ^e	NAb	Not serious ^e	Seriouse	None	405	1.66 (0.88, 3.15)	MODERATE
Acute respirator	y infection req	uiring hospital	isation						
Infrequent dampn	ess in the hous	e							
Tin Tin 2016	Prospective cohort	Serious ^g	NAb	Not serious ^c	Seriouse	None	6853	0.95 (0.82, 1.11)	LOW
Frequent dampne	ess in the house	:							
Tin Tin 2016	Prospective cohort	Serious ^g	NAb	Not serious ^c	Seriouse	None	6853	1.08 (0.91, 1.29)	LOW
Always dampness	s in the house								
Tin Tin 2016	Prospective cohort	Serious ^g	NAb	Not serious ^c	Seriouse	None	6853	1.07 (0.84, 1.37)	LOW
Rhinitis									
Reported as mild	damp stains								
Hagerhed- Engman 2009	Nested case control	Not serious ^f	NAb	Not serious ^c	Seriouse	None	400	1.39 (0.73, 2.67)	LOW
Reported as seve	ere damp stains								
Hagerhed- Engman 2009	Nested case control	Not serious ^f	NAb	Not serious ^c	Seriouse	None	400	0.37 (0.04, 3.43)	LOW

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Reported as mild f		KISK OI DIAS	Су	munectness	IIIIprecision	Other	Nullibel	umess stateuj	Quality
Hagerhed- Engman 2009	Nested case control	Not serious ^f	NAb	Not serious°	Serious ^e	None	400	1.16 (0.36, 3.76)	LOW
Reported as sever	e floor damp								
Hagerhed- Engman 2009	Nested case control	Not serious ^f	NAb	Not serious ^c	Seriouse	None	400	1.58 (0.10, 26.14)	LOW
Bronchitis									
Brunekreef 1989	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	4625	1.32 (1.05, 1.67)	LOW
Bronchiolitis									
Zhou 2013	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Seriouse	None	1765	1.32 (0.80, 2.18)	MODERATE
Chest illness									
Brunekreef 1989	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	4625	1.52 (1.20, 1.93)	LOW
Chronic respirato	ory disease								
Reported as damp	ness and mou	ld							
Dales 1991	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	14799	1.45 (1.29, 1.64)	LOW
Lower respiratory	y illness								
Brunekreef 1989	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	4625	1.68 (1.41, 2.01)	LOW
Reported as water	damage or mo	ould / mildew							
Stark 2003	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Seriouse	None	499	1.34 (0.99, 1.82)	MODERATE
Lower respiratory	y symptoms								
Reported as damp	ness and mou	ld							

			Inconsisten					Adjusted relative effect (aOR 95%CI	
No of studies	Design	Risk of bias	су	Indirectness	Imprecision	Other	Number	unless stated)	Quality
Dales 1991	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	14799	1.62 (1.48. 1.78)	LOW
Upper respirator	ry symptoms								
Reported as dam	pness and mou	ld							
Dales 1991	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	14799	1.50 (1.38, 1.61)	LOW
Phlegm									
Reported as sligh	it to moderate d	ampness							
Cable 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriouse	None	7320	0.82 (0.54, 1.27)	VERY LOW
Reported as marl	ked dampness								
Cable 2014	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	7320	2.05 (1.07, 3.91)	LOW
Any sleep proble	ems								
Reported as dam	pness at home	/ visible mould							
Tiesler 2015	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	1719	1.80 (1.22, 2.66)	LOW
Problems falling	asleep								
Reported as dam	pness at home	/ visible mould							
Tiesler 2015	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Seriouse	None	1719	1.50 (0.98, 2.30)	VERY LOW
Problems sleepi	ng throughout	night							
Reported as dam	pness at home	/ visible mould							
Tiesler 2015	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	1719	2.36 (1.15, 4.84)	LOW
Sleep<9 hours									
Reported as dam	pness at home	/ visible mould							
Tiesler 2015	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	1719	1.60 (1.02, 2.51)	LOW

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Wheeze									
Brunekreef 1989	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Not serious ^d	None	4625	1.23 (1.10, 1.39)	HIGH
Casas 2012	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Seriouse	None	5078	1.11 (0.87, 1.43)	MODERATE
Zhou 2013	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	1765	2.12 (1.30, 3.46)	HIGH
Reported as damp	o / mould								
Jedrychowski 2010	Prospective cohort	Seriouse	NAb	Not serious ^c	Not serious ^d	None	369	IRR 1.67 (1.39, 2.01)	MODERATE
Reported as damp	o / mould								
Jedrychowski 2011	Prospective cohort	Serious ^h	NAb	Not serious ^c	Not serious ^d	None	322	aHR 1.22 (1.07, 1.40)	MODERATE
Eye irritation									
Reported as damp	oness and mou	ild							
Dales 1991	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	14799	1.63 (1.46, 1.82)	LOW
Non-chest illness	S								
Brunekreef 1989	Prospective cohort	Very serious ^a	NAb	Not serious ^c	Not serious ^d	None	4625	1.55 (1.25, 1.93)	LOW

⁽a) Very serious concerns over risk of bias due over self-report of exposure and outcomes (b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
(e) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

⁽f) No concerns over risk of bias

⁽g) Serious concerns over risk of bias due to self-report of exposure

F.2.1.2 Mould

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Cough									
Belanger 2003	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	593	1.53 (1.01, 2.30)	HIGH
Brunekreef 1989	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	4625	2.12 (1.64, 2.73)	HIGH
Engvall 2001	Retrospecti ve cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	9808	3.30 (3.16, 3.46	HIGH
At risk children									
Belanger 2003	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	256 ^g	1.91 (1.07, 3.42)	HIGH
Asthma									
McConnell 2002	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	3535	aRR 1.10 (0.80, 1.60)	MODERATE
Brunekreef 1989	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	4625	1.27 (0.93, 1.74)	MODERATE
Norback 2013	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	7104	aRR 1.15 (0.71, 1.85)	MODERATE
Reported as moul	d odour								
Jaakkola 2005	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	1916	2.44 (1.07, 5.60)	LOW
Reported as visibl	e mould								
Jaakkola 2005	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Seriouse	None	1916	0.65 (0.24, 1.72)	VERY LOW
Any indicator of m	ould or damp (1 of Mould odou	ır, visible mould	or dampness da	amage)				
Thacher 2017	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	3798	1.16 (0.93, 1.44)	MODERATE

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
2 indicators of mo			-				110	,	- Lauring
Thacher 2017	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	3798	1.37 (1.01, 1.86)	HIGH
3 indicators of mo	uld or damp (3	of Mould odour,	visible mould, o	or dampness dar	mage)				
Thacher 2017	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	3798	1.73 (1.10, 2.74)	HIGH
Asthma (childrer	with wheeze	at baseline)							
McConnell 2002	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	3535	0.60 (0.40, 0.90)	HIGH
Allergic rhinocor	njunctivitis								
Reported as moul	d on walls								
Ibargoyen- Roteta 2007	Prospective cohort	Very serious ^f	NAb	Not seriouse	Seriouse	None	3360	1.34 (0.64, 2.79)	VERY LOV
Allergic rhinitis									
Reported as moul	d on walls								
Jaakkola 2010	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	1863	1.73 (1.27, 2.38)	LOW
Reported as visibl	e mould								
Jaakkola 2010	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	1863	1.98 (1.32, 2.99)	LOW
Acute respiratory	y infection								
Mould or mildew in	n the walls or c	eilings in the roo	om where child s	sleeps at night in	the past two we	eeks			
Tin Tin 2016	Prospective cohort	Serious ^g	NAb	Not serious ^c	Not serious ^d	None	6853	0.81 (0.67, 0.99)	MODERAT
Rhinitis									
Any indicator of m	ould or damp (1 of Mould odou	ır, visible mould,	or dampness da	amage)				
Thacher 2017	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	3798	1.03 (0.87, 1.22)	MODERAT

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Thacher 2017	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^e	None	3798	1.18 (0.92, 1.52)	MODERATE
3 indicators of mor	uld or damp (3	of Mould odour,	visible mould, o	or dampness dar	mage)				
Thacher 2017	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^e	None	3798	1.23 (0.82, 1.85)	MODERATE
Bronchitis									
Brunekreef 1989	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	4625	1.48 (1.17, 1.87)	LOW
Chest illness									
Brunekreef 1989	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	4625	1.40 (1.11, 1.78)	LOW
Otitis media									
Pettigrew 2004	Prospective cohort	Serious ^g	NAb	Not serious ^c	Seriouse	None	1002	1.37 (0.94, 2.02)	LOW
Hay fever									
Brunekreef 1989	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	4625	1.57 (1.31, 1.74)	LOW
Lower respiratory	y illness								
Brunekreef 1989	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	4625	1.57 (1.31, 1.87)	LOW
Nasal symptoms									
Reported as mould	dy odour								
Engvall 2001	Retrospecti ve cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	9808	2.83 (2.73, 2.93)	LOW
Throat symptoms	8								
Reported as mould	dy odour								
Engvall 2001	Retrospecti ve cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	9808	3.48 (3.33, 3.62)	LOW
Facial skin symp	toms								

								Adjusted relative effect	
No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	(aOR 95%CI unless stated)	Quality
Reported as mou	_							,	
Engvall 2001	Retrospecti ve cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	9808	2.93 (2.80, 3.06)	LOW
Headache									
Reported as mou	ldy odour								
Engvall 2001	Retrospecti ve cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	9808	3.37 (3.24, 3.51)	LOW
Any sleep proble	ems								
Tiesler 2015	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	1719	1.70 (1.13, 2.54)	LOW
Problems falling	asleep								
Reported as visib	le mould								
Tiesler 2015	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriouse	None	1719	1.50 (0.97, 2.33)	LOW
Problems sleepi	ng throughout	night							
Reported as visib	le mould								
Tiesler 2015	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Serious ^e	None	1719	1.91 (0.89, 4.13)	VERY LOW
Sleep<9 hours									
Reported as visib	le mould								
Tiesler 2015	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	1719	1.67 (1.06, 2.65)	LOW
Tiredness									
Reported as mou	ldy odour								
Engvall 2001	Retrospecti ve cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	9808	2.38 (2.31, 2.46)	LOW
Wheeze									
Reported as mou	ld / mildew								

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Belanger 2003	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Seriouse	None	593	1.22 (0.80, 1.88)	VERY LOW
Brunekreef 1989	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	4625	1.79 (1.44, 2.32)	LOW
Karvonen 2015	Prospective cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	398	1.34 (0.90, 2.01)	LOW
Reported as moule	d / mildew (in a	at risk children)							
Belanger 2003	Prospective cohort	Serious ^g	NAb	Not serious ^c	Not serious ^d	None	256	2.51 (1.37, 4.62)	MODERATE
Reported as visible	e mould (low v	s none) in childr	en with atopy						
lossifova 2009	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	483	1.86 (0.86, 4.00)	MODERATE
Reported as visible	e mould (high	vs none) in child	ren with atopy						
lossifova 2009	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	483	6.16 (1.38, 27.44)	HIGH
Eczema									
Reported as mild i	mould								
Hagerhed- Engman 2009	Nested case control	Not serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	400	1.86 (1.04, 3.30)	MODERATE
Reported as sever	re mould								
Hagerhed- Engman 2009	Nested case control	Not serious ^a	NA ^b	Not serious ^c	Seriouse	None	400	1.93 (0.91, 4.12)	LOW
Eye irritation									
Reported as moule	dy odour								
Engvall 2001	Retrospecti ve cohort	Very serious ^f	NAb	Not serious ^c	Not serious ^d	None	9808	3.75 (3.60, 3.92)	LOW
Non-chest illness	3								

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Brunekreef 1989	Prospective cohort	Very serious ^f	NA ^b	Not serious ^c	Not serious ^d	None	4625	1.40 (1.13, 1.74)	LOW
Otitis media									
Any episode									
Pettigrew 2004 b	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^e	None	813	1.15 (0.67, 1.99)	MODERATE
Otitis media, Recu	ırrent (4 or mor	e episodes of o	titis media (sepa	rated by at least	21 days) in one	e year)			
Pettigrew 2004 b	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Seriouse	None	813	1.05 (0.88, 1.26)	MDOERATE

- (a) No concerns over risk of bias
- (b) Not applicable as only one study included
- (c) No concerns over directness
- (d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
- (e) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)
- (f) Very serious concerns over risk of bias due over self-report of exposure and outcomes (g) Serious concerns over risk of bias due to self-report of outcomes

F.2.1.3 Fungal spore levels

ı unga	i spore leve	713									
No of	f studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality	
Coug	gh in childrei	n with asthma	l								
Clado	osporium >14	8 CFU/m ³									

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Gent 2012	Prospective cohort	Seriousª	NAb	Not serious ^c	Seriousd	None	1233	0.98 (0.54, 1.80)	LOW
Asthma severity	in children wi	th asthma							
Cladosporium >1	48 CFU/m ³								
Gent 2012	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	1233	1.58 (0.88, 2.83)	LOW
Rescue medicat	ion use in chile	dren with asthr	ma						
Cladosporium >1	48 colony formi	ng units (CFU)	m³						
Gent 2012	Prospective cohort	Seriousª	NAb	Not serious ^c	Seriousd	None	1233	0.69 (0.37, 1.29)	LOW
Lower respirato	ry illness in inf	ants at risk of	asthma						
> 90th percentile	for specific taxo	on							
Stark 2003	Prospective cohort	Not seriouse	NAb	Not serious ^c	Not serious ^f	None	499	1.86 (1.21, 2.88)	HIGH
Wheeze in child	ren with asthm	ia							
Cladosporium >1	48 CFU/m ³								
Gent 2012	Prospective cohort	Seriousª	NAb	Not serious ^c	Seriousd	None	1233	1.22 (0.66, 2.26)	LOW
Otitis media in 1	st 6 months of	life in infants a	t risk of asthm	а					
Penicillium ≥1000) CFU/m ³								
Pettigrew 2004 (US)	Prospective cohort	Seriousª	NAb	Not serious ^c	Seriousd	None	1002	1.27 (0.56, 2.86)	LOW
Cladosporium ≥1	000 CFU/m ³								
Pettigrew 2004 (US)	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	1002	1.09 (0.52, 2.29)	LOW
Other mould (not	yeast, penicilliu	ım or cladospori	um) ≥1000 CFU	/m³					
Pettigrew 2004 (US)	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^f	None	1002	3.45 (1.36, 8.76)	MODERA

⁽a) Serious concerns over risk of bias due over self-report of outcomes

- (b) Not applicable as only one study included(c) No concerns over directness
- (d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)
- (e) No concerns over risk of bias
- (f) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

F.2.2 Formaldehyde

Officialdeflyde									
No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Wheezing in infa	nts at risk of a	asthma							
Formaldehyde bet	tween 12.4 and	d 16.3 µg/m³ cor	npared to < 12.4	in infants at risk	of asthma				
Raaschou- Nielsen 2010	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	9808	1.11 (0.47, 2.63)	VERY LOW
Formaldehyde bet	tween 16.3 and	d 20.3 µg/m³ cor	npared to < 12.4	in infants at risk	of asthma				
Raaschou- Nielsen 2010	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	9808	1.21 (0.51, 2.92)	VERY LOW
Formaldehyde bet	tween 20.3 and	d 25.6 µg/m³ cor	npared to < 12.4	in infants at risk	of asthma				
Raaschou- Nielsen 2010	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	9808	1.40 (0.57, 3.47)	VERY LOW
Formaldehyde > 2	25.6 μg/m³ com	pared to < 12.4	in infants at risk	of asthma					
Raaschou- Nielsen 2010	Retrospecti ve cohort	Very serious ^a	NAb	Not serious ^c	Serious ^d	None	9808	0.67 (0.29, 1.54)	VERY LOW
Lower respirator	y tract infection	ons							
per inter-quartile r	ange increase								
Roda 2011	Prospective cohort	Serious ^e	NAb	Not serious ^c	Not serious ^f	None	2940	1.32 (1.11, 1.55)	MODERATE

No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Lower respirator	ry tract infection	on with wheeze							
per inter-quartile ı	range increase								
Roda 2011	Prospective cohort	Serious ^e	NAb	Not serious ^c	Not serious ^f	None	Not reported	1.41 (1.14, 1.74)	MODERATE

⁽a) Very serious concerns over risk of bias due over self-report of exposure and outcomes

F.2.3 Allergens

F.2.3.1 House dust mite allergens (Der p 1 + Der f 1)

No of studies	Design	Risk of bias	Inconsistenc y	Indirectnes s	Imprecisio	Other	Numbe r	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Asthma exacerb	oations								
House dust mite	(HDM) > 10µg	g/g							
Gent 2012	Prospectiv e cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	None	1233	1.19 (0.92, 1.55)	LOW

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%CIs cross the line of no effect)
(e) Serious concerns over risk of bias due to self-report of outcomes
(f) No concerns as findings are statistically significant (95%CIs do not cross the line of no effect)

No of studies	Design	Risk of bias	Inconsistenc y	Indirectnes s	Imprecisio n	Other	Numbe r	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Asthma ≤ 6 yea	rs of age								
Casas 2015	Prospectiv e cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	≥0.19 µg/g to <0.4 µg/g	4334	1.6 (0.9, 2.6)	MODERATE
Casas 2015	Prospectiv e cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^f	0.4 to <2 µg/g	4334	1.4 (1.1, 1.9)	HIGH
Casas 2015	Prospectiv e cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	≥2 µg/g	4334	1.1 (0.8, 1.6)	MODERATE
Asthma > 6 year	irs of age								
Casas 2015	Prospectiv e cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	≥0.19 µg/g to <0.4 µg/g	4334	1.3 (0.8, 2.3)	MODERATE
Casas 2015	Prospectiv e cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	0.4 to <2 µg/g	4334	1.1 (0.8, 1.6)	MODERATE
Casas 2015	Prospectiv e cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	≥2 µg/g	4334	1.0 (0.7, 1.4)	MODERATE
Wheeze									
HDM									
Herr 2012	Prospectiv e cohort	Not serious ^e	NAb	Not serious ^c	Not serious ^f	None	1879	1.39 (1.12, 1.73)	HIGH
HDM 0.981 - 24	0μg/g								
Lau 2000	Prospectiv e cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	None	1314	1.03 (0.52, 2.04)	MODERATE
Reported as per	sistent wheez	e							
Casas 2015	Prospectiv e cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	≥0.19 µg/g to <0.4 µg/g	4334	1.3 (0.8, 2.2)	MODERATE
Casas 2015	Prospectiv e cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	0.4 to <2 µg/g	4334	1.1 (0.8, 1.5)	MODERATE
Casas 2015	Prospectiv e cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	≥2 µg/g	4334	0.9 (0.7, 1.3)	MODERATE
HDM (Der p 1 +			(infants)						

No of studies	Design	Risk of bias	Inconsistenc y	Indirectnes s	Imprecisio n	Other	Numbe r	Adjusted relative effect (aOR 95%CI unless stated)	Quality	
Belanger 2003	Prospectiv e cohort	Serious ^a	NA ^b	Not serious ^c	Serious ^d	None	256	1.04 (0.60, 1.80)	LOW	
Dust mite (Der p	Dust mite (Der p 1 + Der f 1) ≥ 2 μg/g (for not at risk infants)									
Belanger 2003	Prospectiv e cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	None	593	0.78 (0.55, 1.13)	LOW	
Cough										
HDM (Der p 1 + Der f 1) ≥ 2 μg/g (for at risk infants)										
Belanger 2003	Prospectiv e cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	None	256	1.27 (0.75, 2.15)	LOW	
HDM (Der p 1 + Der f 1) ≥ 2 μg/g (for not at risk infants)										
Belanger 2003	Prospectiv e cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	None	593	0.76 (0.54, 1.07)	LOW	

⁽a) Serious concerns over risk of bias due over self-report of outcomes

F.2.3.2 House dust mite allergens (Der p 1)

No of studies	Design	Risk of bias	Inconsistenc y	Indirectnes s	Imprecisio	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Asthma exacerbations									
Der p 1 >0.10 μg/g (reported as Asthma Severity Index)									
Gent 2012	Prospectiv e cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	1233	1.19 (0.92, 1.55)	LOW

⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%CIs cross the line of no effect)

⁽e) No concerns over risk of bias

⁽f) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

		Risk of	Inconsistenc	Indirectnes	Imprecisio			Adjusted relative effect (aOR 95%CI	
No of studies	Design	bias	у	s	n ·	Other	Number	unless stated)	Quality
Der p 1 >0.10 μg	/g (reported a	as rescue med	ication use)						
Gent 2012	Prospectiv e cohort	Serious ^a	NAb	Not serious ^c	Not seriouse	None	1233	1.47 (1.11, 1.94)	MODERATE
Der p 1 (µg/g) 0.	10 to <2.0 vs	<0.10 in main	living area (repor	ted as Moderat	e/severe GINA	score)			
Gent 2009	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Seriousd	None	300	0.93 (0.41, 2.10)	MODERATE
Der p 1 (µg/g) 0.	10 to <2.0 vs	<0.10 in main	living area (repor	ted as Controll	er meds≥9 mor	iths)			
Gent 2009	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Seriousd	None	300	0.61 (0.27,1.35)	MODERATE
Der p 1 (µg/g) 2.	0 to <10.0 vs	<0.10 in main	living area (repor	ted as Moderat	e/severe GINA	score)			
Gent 2009	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Not seriouse	None	300	2.93 (1.37, 6.30)	HIGH
Der p 1 (µg/g) 2.	0 to <10.0 vs	<0.10 in main	living area (repor	rted as Controll	er meds≥9 mor	iths)			
Gent 2009	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Not seriouse	None	300	2.52 (1.17,5.42)	HIGH
Der p 1 (μg/g) ≥1	0.0 vs <0.10	in main living	area (reported as	Moderate/seve	ere GINA score)			
Gent 2009	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Not seriouse	None	300	2.55 (1.13, 5.73)	HIGH
Der p 1 (μg/g) ≥1	0.0 to <2.0 vs	s <0.10 in mai	n living area (repo	orted as Contro	ller meds≥9 mc	onths)			
Gent 2009	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Seriousd	None	300	2.17 (0.97,4.86)	MODERATE
Der p 1 (μg/g) 0.	10 to <2.0 vs	<0.10 in bed (reported as Mode	erate/severe GI	NA score)				
Gent 2009	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	300	0.99 (0.47, 2.08)	MODERATE
Der p 1 (μg/g) 0.	10 to <2.0 vs	<0.10 in bed (reported as Cont	roller meds≥9 n	nonths)				
Gent 2098	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Seriousd	None	300	1.35 (0.66,2.73)	MODERATE
Der p 1 (µg/g) 2.	0 to <10.0 vs	<0.10 in bed (reported as Mode	erate/severe GI	NA score)				

No of studies	Design	Risk of bias	Inconsistenc	Indirectnes s	Imprecisio	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Gent 2009	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Not seriouse	None	300	2.73 (1.32,5.64)	HIGH
Der p 1 (µg/g) 2.	0 to <10.0 vs	<0.10 in bed (reported as Cont	roller meds≥9 m	nonths)				
Gent 2009	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Not serious ^e	None	300	2.16 (1.04,4.48)	HIGH
Der p 1 (µg/g) ≥1	10.0 vs <0.10	in bed (reporte	ed as Moderate/s	evere GINA sco	ore)				
Gent 2009	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	300	1.19 (0.46,3.08)	MODERATE
Der p 1 (µg/g) ≥1	10.0 to <2.0 vs	s <0.10 in bed	(reported as Cor	troller meds≥9	months)				
Gent 2009	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	300	1.41 (0.57,3.46)	MODERATE
Asthma									
Der p 1 for asthn	na at ≤ 6 years	S							
Casas 2015	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Seriousd	≥1.9 µg/g to <0.4 µg/g	4334	1.4 (0.9, 2.3)	MODERATE
Casas 2015	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	0.4 to <2 μg/g	4334	1.1 (0.8, 1.6)	MODERATE
Casas 2015	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	≥2 µg/g	4334	1.0 (0.7, 1.5)	MODERATE
asthma at > 6 ye	ars								
Casas 2015	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	≥1.9 µg/g to <0.4 µg/g	4334	1.1 (0.6, 1.8)	MODERATE
Casas 2015	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	0.4 to <2 μg/g	4334	1.1 (0.8, 1.6)	MODERATE
Casas 2015	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Serious ^d	≥2 µg/g	4334	0.7 (0.4, 1.0)	MODERATE
Der p1 0.83–6.46	β μg/g								
Torrent 2007	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	1611	0.67 (0.40, 1.12)	MODERATE
Der p1 >6.46 μg/	/g								

No of studies	Design	Risk of bias	Inconsistenc	Indirectnes s	Imprecisio	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Torrent 2007	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	None	1611	0.68 (0.37, 1.25)	MODERATE
Wheeze									
Der p 1 >0.10 μg	ı/g								
Gent 2012	Prospectiv e cohort	Serious ^a	NA ^b	Not serious ^c	Seriousd	None	1233	1.26 (0.95, 1.67)	LOW
Der p 1 for persis	stent wheeze								
Casas 2015	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	≥0.12 to <0.4 µg/g	4334	1.1 (0.7, 1.8)	MODERATE
Casas 2015	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Seriousd	> 0.4 to <2 µg/g	4334	0.9 (0.7, 1.3)	MODERATE
Casas 2015	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Seriousd	≥2 µg/g	4334	0.8 (0.5, 1.1)	MODERATE
Der p 1 (µg/g) 0.	10 to <2.0 vs	<0.10 in main	living area						
Gent 2009	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Seriousd	None	300	1.05 (0.38,2.84)	MODERATE
Der p 1 (µg/g) 2.	0 to <10.0 vs	<0.10 in main	living area						
Gent 2009	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Seriousd	None	300	1.55 (0.62,3.85)	MODERATE
Der p 1 (μg/g) ≥1	10.0 vs <0.10	in main living	area						
Gent 2009	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	300	2.01 (0.78,5.19)	MODERATE
Der p 1 (µg/g) 0.	10 to <2.0 vs	<0.10 in bed							
Gent 2009	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	300	1.70 (0.68,4.22)	MODERATE
Der p 1 (µg/g) 2.	0 to <10.0 vs	<0.10 in bed							
Gent 2009	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	300	1.60 (0.64,4.00)	MODERATE
Der p 1 (μg/g) ≥1	10.0 vs <0.10	in bed							

No of studies	Design	Risk of bias	Inconsistenc y	Indirectnes s	Imprecisio n	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Gent 2009	Prospectiv e cohort	Not serious ^f	NA ^b	Not serious ^c	Not seriouse	None	300	3.58 (1.28, 9.97)	HIGH
Cough									
HDM Der p 1 >0.	10 μg/g								
Gent 2012	Prospectiv e cohort	Serious ^a	NA ^b	Not serious ^c	Serious ^d	None	1233	1.18 (0.90, 1.55)	LOW
Eczema									
HDM allergen qu	intiles compai	red to lowest of	quintile (0·02–0·2	27)					
Harris 2007	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	Quintile 2 (0·28 -0·81)	593	1.01 (0.53, 1.92)	MODERATE
Harris 2007	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	Quintile 3 (0·82–2·22)	593	1.37 (0.74, 2.55)	MODERATE
Harris 2007	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	(Quintile 4 (2·23–7·75)	593	0.66 (0.34, 1.29)	MODERATE
Harris 2007	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	Quintile 5 (7·76–384·97)	593	0.71 (0.37, 1.37)	MODERATE
Visible flexural	dermatitis								
House dust mite	allergen quint	iles compared	to lowest quintil	e (0·02–0·27)					
Harris 2007	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	Quintile 2 (0·28– 0·81)	593	1.17 (0.58, 2.34)	MODERATE
Harris 2007	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	Quintile 3 (0·82–2·22)	593	1.73 (0.87, 3.46)	MODERATE
Harris 2007	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	Quintile 4 (2·23–7·75)	593	0.88 (0.43, 1.81)	MODERATE
Harris 2007	Prospectiv e cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	Quintile 5 (7·76–384·97)	593	0.96 (0.47, 1.94)	MODERATE

⁽a) Serious due to concerns over self-report of exposure(b) Not applicable as only one study included(c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect) (e) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)

(f) No concerns over risk of bias

F.2.3.3 House dust mite allergens (Der f 1)

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Asthma exacer	bations								
Der f 1 >2.1 μg/g	g (reported as	rescue medio	cation use)						
Gent 2012	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	1233	1.09 (0.78, 1.51)	LOW
Der f 1 >2.1 μg/g	g (reported as	Asthma Seve	erity Index)						
Gent 2012	Prospective cohort	Serious ^e	NAb	Not serious ^c	Seriousd	None	1233	1.28 (0.94, 1.74)	LOW
Asthma									
Der f 1 for asthm	na at ≤ 6 years								
Casas 2015	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	≥0.07 to <0.4 µg/g	4334	1.2 (0.8, 1.8)	MODERATE
Casas 2015	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	0.4 to <2 μg/g	4334	1.2 (0.8, 1.6)	MODERATE
Casas 2015	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	≥2 µg/g	4334	1.2 (0.8, 1.8)	MODERATE
Der f 1 for asthm	na at > 6 years								
Casas 2015	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	≥Low to <0.4 µg/g	4334	1.0 (0.7, 1.6)	MODERATE
Casas 2015	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	0.4 to <2 μg/g	4334	1.0 (0.7, 1.4)	MODERATE
Casas 2015	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	≥2 µg/g	4334	1.1 (0.7, 1.6)	MODERATE
Der f 1 exposure	e at 3 months f	or asthma at	7 years in children	n at risk of asthm	na (per interqua	artile increase in ex	posure)		
O'Connor 2017	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	None	442	0.98 (0.91, 1.04)	MODERATE

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Wheeze									
Der f 1 per 1-log	increase in all	ergen level ir	n first year of life fo	or wheeze at 3 y	ears in childrer	at risk of asthma			
Lynch 2014	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	560	0.92 (0.73, 1.15)	LOW
Der f 1 for persis	tent wheeze								
Casas 2015	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	≥0.07 to <0.4 µg/g	4334	1.2 (0.8, 1.8)	MODERATE
Casas 2015	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	0.4 to <2 μg/g	4334	1.1 (0.8, 1.5)	MODERATE
Casas 2015	Prospective cohort	Not serious ^e	NAb	Not serious ^c	Seriousd	≥2 µg/g	4334	1.1 (0.8, 1.6)	MODERATE
Der f 1 >2.1 μg/g	3								
Gent 2012	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	1233	0.89 (0.63, 1.24)	LOW
Cough									
Der f 1 >2.1 μg/g)								
Gent 2012	Prospective cohort	Serious ^a	NAb	Not serious ^c	Seriousd	None	1233	0.90 (0.65, 1.25)	LOW

⁽a) Serious concerns over risk of bias due to self-report of exposure
(b) Not applicable as only one study included
(c) No concerns over directness
(d) Serious concerns as findings are not statistically significant (95%CIs cross the line of no effect)
(e) No concerns over risk of bias

F.2.3.4 Cat allergens (Fel d 1)

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Asthma								,	
Fel d 1 ≥2 µg/gm	1								
Carlsten 2010	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	380	3.33 (1.72, 6.45)	HIGH
Fel d 1 >0.12 µg	/g (asthma rep	orted as Ast	hma Severity Inde	ex)					
Gent 2012	Prospective cohort	Seriouse	NA ^b	Not serious ^c	Seriousf	None	1233	1.14 (0.88, 1.47)	LOW
Fel d 1 >0.12 µg	/g (asthma rep	orted as res	cue medication us	e)					
Gent 2012	Prospective cohort	Seriouse	NA ^b	Not serious ^c	Not serious ^d	None	1233	1.32 (1.01, 1.74)	MODERATE
Fel d 1 0.216 - 4	l7μg/g								
Lau 2000	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Seriousf	None	1314	1.52 (0.64, 2.62)	MODERATE
Wheeze									
Fel d 1 >1 µg/g i	n infants at ris	k of asthma	or atopy						
Belanger 2003	Prospective cohort	Seriouse	NA ^b	Not serious ^c	Seriousf	None	256	0.64 (0.36, 1.12)	LOW
Litonjua 2002	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Seriousf	None	226	0.61 (0.27, 1.35)	MODERATE
Lau 2000	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Seriousf	None	1314	1.47 (0.72, 1.26)	MODERATE
Fel d 1 per 1-log	increase in all	lergen level i	n first year for whe	eeze at 3 years a	at risk of asthm	а			
Lynch 2002	Prospective cohort	Seriouse	NA ^b	Not serious ^c	Seriousf	None	560	0.71 (0.58, 0.88)	LOW
Fel d 1 >1 µg/g i	n infants not a	t risk of asth	ma						
Belanger 2003	Prospective cohort	Seriouse	NA ^b	Not serious ^c	Seriousf	None	593	0.84 (0.57, 1.24)	LOW
Children with ast	thma								

No of atualisa	Danima	Risk of	la canciata no c	lu dina stu a sa	Immunoining	Othor	Number	Adjusted relative effect (aOR 95%CI	Overlife.
No of studies	Design	bias	Inconsistency	Indirectness		Other	Number	unless stated)	Quality
Gent 2012	Prospective cohort	Seriouse	NA ^b	Not serious ^c	Not serious ^d	None	1233	1.39 (1.05, 1.84)	MODERATE
Fel d 1 exposure	at 3 months f	or asthma at	7 years in childre	n at risk of asth	ma (per interqu	artile increase in ex	oosure)		
O'Connor 2017	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	442	0.78 (0.62, 0.98)	HIGH
Cough									
Infants at risk of	asthma								
Belanger 2003	Prospective cohort	Serious ^e	NA ^b	Not serious ^c	Serious ^f	None	256	1.13 (0.66, 1.94)	LOW
Infants not at risl	k of asthma								
Belanger 2003	Prospective cohort	Serious ^e	NAb	Not serious ^c	Serious ^f	None	593	0.81 (0.56, 1.17)	LOW
Children with ast	thma								
Gent 2012	Prospective cohort	Serious ^e	NAb	Not serious ^c	Not serious ^d	Fel d 1 >0.12 μg/g	1233	0.89 (0.68, 1.17)	MODERATE
Asthma									
Cat allergen (per	r 10 mg increa	se)							
Bertelsen 2010	Prospective cohort	Not serious ^g	NAb	Not serious ^c	Not serious ^d	None	260	1.20 (1.01, 1.43),	HIGH
Bronchial hype	rresponsiven	ess							
Cat allergen (per	10 mg increa	se)							
Bertelsen 2010	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	260	1.22 (1.02, 1.46)	HIGH
Eczema									
Cat allergen exp	osure – quintil	es compared	to lowest quintile	e (0·01–0·44)					
Harris 2007	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	Quintile 2 (0·45– 1·04)	593	1.42 (0.72, 2.81)	MODERATE
Harris 2007	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	Quintile 3 (1·05– 3·33)	593	1.41 (0.71, 2.79)	MODERATE

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Harris 2007	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	Quintile 4 (3·34– 44·72))	593	1.31 (0.65, 2.62)	MODERATE
Harris 2007	Prospective cohort	Not serious ^g	NA ^b	Not serious ^c	Serious ^d	Quintile 5 (44·73– 14151·32)	593	1.41 (0.72, 2.75)	MODERATE
Visible flexural	dermatitis								
Cat allergen exp	osure – quintil	es compared	to lowest quintile	e (0·01–0·44)					
Harris 2007	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	Quintile 2 (0·28– 0·81)	593	1.28 (0.64, 2.56)	MODERATE
Harris 2007	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	Quintile 3 (0·82– 2·22)	593	0.75 (0.36, 1.55)	MODERATE
Harris 2007	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	Quintile 4 (2·23–7·75)	593	1.18 (0.59, 2.38)	MODERATE
Harris 2007	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^d	Quintile 5 (7·76– 384·97)	593	0.96 (0.48, 1.91)	MODERATE

⁽a) No concerns over risk of bias

F.2.3.5 Dog allergens (Can f 1)

Dog allergens		Risk of						Adjusted relative effect (aOR 95%CI	
No of studies	Design	bias	Inconsistency	Indirectness	Imprecision	Other	Number	unless stated)	Quality
Asthma									
Can f 1 ≥ 2 µg/g									

⁽b) Not applicable as only one study included(c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect) (e) Serious due to concerns over self-report of exposure

⁽f) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%CI unless stated)	Quality
Carlsten 2010	Prospective cohort	Not serious ^a	NAb	Not serious ^c	Not serious ^d	None	380	3.84 (1.79, 8.22)	HIGH
Can f 1 >1.2 µg/g	g (asthma repo	orted as Ast	hma Severity Index	()					
Gent 2012	Prospective cohort	Serious ^e	NAb	Not serious ^c	Seriousf	None	1233	1.15 (0.83, 1.58)	LOW
Can f 1 >1.2 µg/g	g (asthma repo	orted as res	cue medication use	e)					
Gent 2012	Prospective cohort	Seriouse	NAb	Not serious ^c	Seriousf	None	1233	1.15 (0.82, 1.62)	LOW
Can d 1 exposur	e at 3 months	for asthma	at 7 years in childre	en at risk of asth	ıma (per interqu	uartile increase in e	xposure)		
O'Connor 2017	Prospective cohort	Not serious ^a	NA ^b	Not serious ^c	Serious ^f	None	442	0.62 (0.37, 1.03)	MODERATE
Cough									
Dog allergen (Ca	an f 1) ≥1.8µg/g	g (for at risk	infants)						
Belanger 2003	Prospective cohort	Seriouse	NAb	Not serious ^c	Serious ^f	None	256	0.91 (0.53, 1.56)	LOW
Dog allergen (Ca	an f 1) ≥1 for 8	μg/g (for no	t at risk infants)						
Belanger 2003	Prospective cohort	Seriouse	NAb	Not serious ^c	Seriousf	None	593	1.11 (0.78, 1.58)	LOW
Can f 1 >1.2 µg/g	g								
Gent 2012	Prospective cohort	Serious ^e	NAb	Not serious ^c	Seriousf	None	1233	1.11 (0.80, 1.56)	LOW
Wheeze									
Can f 1 per 1-log	increase in al	lergen level	in first year for wh	eeze at 3 years	at risk of asthm	na			
Lynch 2002	Prospective cohort	Seriouse	NA ^b	Not serious ^c	Seriousf	None	560	1.00 (0.79, 1.28)	LOW

⁽a) No concerns over risk of bias(b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
(e) Serious due to concerns over self-report of exposure
(f) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

F.2.4 NO₂

No of studies	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisio n	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Asthma exace	erbations								
Belanger 2013	Prospecti ve cohort	Serious ^a	NA ^b	Not serious ^c	Serious ^d	6.02 ppb to 8.88 ppb	1342	1.15 (0.94, 1.42)	LOW
Belanger 2013	Prospecti ve cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	>8.88 ppb to 14.30 ppb	1342	1.31 (1.04, 1.66)	MODERATE
Belanger 2013	Prospecti ve cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	> 14.30 ppb	1342	1.43 (1.08, 1.88)	MODERATE
NO ₂ exposure	at 12 months	for asthma	at 7 years in ch	ildren at risk o	f asthma (per i	nterquartile increas	e in exposu	ire)	
O'Connor 2017	Prospecti ve cohort	Not serious ^a	NAb	Not serious ^c	Serious ^d	None	442	0.97 (0.75, 1.26)	MODERATE
Wheeze									
Belanger 2013 ^a	Prospecti ve cohort	Serious ^a	NAb	Not serious ^c	Seriousd	6.02 - 8.88 ppb	1342	1.15 (0.90, 1.45)	LOW
Belanger 2013 ^a	Prospecti ve cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	8.89- 14.30 ppb	1342	1.44 (1.11, 1.86)	MODERATE
Belanger 2013 ^a	Prospecti ve cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	> 14.30 ppb	1342	1.53 (1.16, 2.02)	MODERATE
NO ₂ 5.2 to 6.8	µg/m³ compa	ared to < 5.2	in infants at ris	k of asthma					
Raaschou- Nielsen 2010	Cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	411	0.66 (0.27, 1.61)	MODERATE
NO ₂ 6.8 to 8.6	µg/m³ compa	ared to < 5.2	in infants at ris	k of asthma					
Raaschou- Nielsen 2010	Cohort	Not serious ^f	NAb	Not serious ^c	Serious ^d	None	411	0.80 (0.32, 2.01)	MODERATE

No of studies	Design	Risk of bias	Inconsiste ncy	Indirectne ss	Imprecisio n	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
NO ₂ 8.6 to 11.	7 μg/m³ com	oared to < 5.2	2 in infants at r	isk of asthma					
Raaschou- Nielsen 2010	Cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	411	1.15 (0.40, 3.32)	MODERATE
NO ₂ > 11.7 μg/	m³ compare	d to < 5.2 in i	nfants at risk o	f asthma					
Raaschou- Nielsen 2010	Cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	411	0.43 (0.15, 1.18)	MODERATE
NO ₂ (per 20pp	b increase in	multi-family	home)						
Belanger 2006	Prospecti ve cohort	Not serious ^f	NAb	Not serious ^c	Not serious ^e	None	242	1.52 (1.04, 2.21)	HIGH
NO ₂ (per 20pp	b increase in	single family	/ home)						
Belanger 2006	Prospecti ve cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	486	0.99 (0.71, 1.38)	MODERATE
Cough									
> 10 ppb (outc	ome reported	d as persister	nt cough)						
Belanger 2003	Prospecti ve cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	None	593 ^g	1.21 (1.05, 1.40)	MODERATE

⁽a) Serious concerns over risk of bias due over self-report of outcomes

F.2.5 PAH



⁽b) Not applicable as only one study included

⁽c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%CIs cross the line of no effect)

⁽e) No concerns as findings are statistically significant (95%CIs do not cross the line of no effect)

⁽f) No concerns over risk of bias

								Adjusted relative effect (aOR	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	95%Cl unless stated)	Quality
Cord blood PA		neeze at 1 – 2 ye	<u> </u>					,	_
Jedrychowski 2010	Prospectiv e cohort	Seriousa	NAb	Not serious ^c	Not serious ^d	None	369	1.69 (1.52, 1.88)	MODERATE
Cord blood PA	H-adducts – wh	neeze at 3 – 4 ye	ears of age						
Jedrychowski 2010	Cohort	Serious ^a	NAb	Not serious ^c	Seriouse	None	369	0.96 (0.84, 1.09)	LOW
per log unit of l	PAH concentrat	tion in ng/m³) for	wheezing or whis	tling in the chest	irrespective of	respiratory	/ infection,		
Jedrychowski 2005	Cohort	Serious ^a	NAb	Not serious ^c	Not serious ^d	None	333	3.83 (1.18, 12.43)	MODERATE
per log unit of l	PAH concentrat	tion in ng/m³) for	wheezing without	cold					
Jedrychowski 2005	Cohort	Serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	333	1.96 (1.38, 2.78)	MODERATE
Prenatal PAH	exposure								
Jedrychowski 2014	Cohort	Seriousa	NA ^b	Not serious ^c	Seriouse	None	257	1.40 (0.97, 2.03)	LOW
Postnatal PAH	exposure								
Jedrychowski 2014	Cohort	Serious ^a	NA ^b	No serious ^c	Not serious ^d	None	257	1.61 (1.16, 2.24)	MODERATE
Pyrene									
Jung 2012	Prospectiv e cohort	Serious ^a	NAb	Not serious ^c	Seriouse	None	349	1.53 (0.93, 2.51)	LOW
Σ8PAH non-vo	latile								
Jung 2012	Prospectiv e cohort	Serious ^a	NAb	Not serious ^c	Serious ^e	None	349	0.86 (0.52, 1.42)	LOW
Asthma									
Pyrene									
Jung 2012	Prospective cohort	Serious ^a	NA ^b	No serious ^c	Not serious ^d	None	349	1.90 (1.13, 3.20)	MODERATE
Σ8PAH non-vo	latile								

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Jung 2012	Prospective cohort	Serious ^a	NAb	No serious ^c	Seriouse	None	349	0.90 (0.52, 1.56)	LOW
Pyrene									
Jung 2014	Prospective cohort	Serious ^a	NA ^b	No serious ^c	Serious ^e	None	363	aRR 0.81 (0.59– 1.12)	LOW
Σ8PAH non-vo	latile							·	
Jung 2014	Prospective cohort	Serious ^a	NAb	No serious ^c	Seriouse	None	363	aRR 0.74 (0.46– 1.18)	LOW
Σ8PAH semi-v	olatile								
Jung 2014	Prospective cohort	Serious ^a	NAb	No serious ^c	Serious ^e	None	363	aRR 0.82 (0.60– 1.12)	LOW
Runny or stuf	fy nose								
per log unit of I	PAH concentra	tion in ng/m ³							
Jedrychowsk i 2005	Cohort	Serious ^a	NA ^b	Not serious ^c	Seriouse	None	333	1.11 (0.97, 1.27)	LOW
Earache (otitis	s media)								
per log unit of I	PAH concentra	tion in ng/m³							
Jedrychowski 2005	Cohort	Serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	333	1.82 (1.03, 3.23)	MODERATE
Sore throat									
per log unit of I	PAH concentra	tion in ng/m³							
Jedrychowski 2005	Cohort	Serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	333	1.27 (1.07, 1.52)	MODERATE
Cough									
per log unit of I	PAH concentra	tion in ng/m³							
Jedrychowski 2005	Cohort	Serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	333	1.72 (1.02, 2.92)	MODERATE
Cough withou	t cold								
per log unit of I	PAH concentra	tion in ng/m³							

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Jedrychowski 2005	Cohort	Serious ^a	NAb	Not serious ^c	Not serious ^d	None	333	4.80 (2.73, 8.44)	MODERATE
Barking cougl	n								
per log unit of F	PAH concentrat	ion in ng/m³							
Jedrychowski 2005	Cohort	Seriousª	NAb	Not serious ^c	Serious ^e	None	333	1.12 (0.82, 1.55)	LOW
Difficult (puffe	ed) breathing,								
per log unit of F	PAH concentrat	ion in ng/m³							
Jedrychowski 2005	Cohort	Serious ^a	NAb	Not serious ^c	Seriouse	None	333	1.23 (0.83, 1.84)	LOW
Pulmonary inf	ections								
Styrene>2.0 µg	_J /m ³								
Diez 2002	Nested case-control	Serious ^a	NAb	Not serious ^c	Not serious ^d	None	475	2.1 (1.1, 4.2)	LOW
Benzene > 5.6	μg/m³								
Diez 2002	Nested case-control	Serious ^a	NAb	Not serious ^c	Not serious ^d	None	475	2.4 (1.3, 4.5)	LOW

⁽a) Serious concerns over risk of bias due over self-report of outcomes(b) Not applicable as only one study included(c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect) (e) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

F.2.6 Particulate matter

F.2.6.1 PM_{2.5}

VI2.5									
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Cough									
Indoor PM _{2.5} ((Indoor) per 17.3	3 μg/m³ increase	e						
Habre 2014	Prospective cohort	Seriousª	NAb	Not serious ^c	Seriousd	None	36	1.22 (0.91, 1.63)	LOW
Indoor PM _{2.5} ((indoor sources)) per 17.6 μg/m ³	increase						
Habre 2014	Prospective cohort	Seriousª	NAb	Not serious ^c	Serious ^d	None	36	1.20 (0.88, 1.64)	LOW
Pindus 2016	Prospective cohort	Serious ^a	NAb	Not serious ^c	Serious ^d	None	905	0.95 (0.72, 1.29)	LOW
Cough, whee	zing or chest ti	ightness in chil	dren with asthma						
Indoor PM _{2.5} (per 10 µg/m³ ind	crease)							
McCormack 2009	Prospective cohort	Serious ^a	NA ^b	Not serious ^c	Not serious ^e	None	150	1.06 (1.01, 1.12)	MODERATE
Wheeze									
PM _{2.5} (Indoor)) per 17.3 μg/m ³	³ increase							
Habre 2014	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	None	36	1.57 (1.09, 2.26)	MODERATE
PM _{2.5} (indoor	sources) per 17	.6 μg/m³ increas	se						
Habre 2014	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not seriouse	None	36	1.55 (1.05, 2.28)	MODERATE
PM _{2.5} ≥15µg/r	m^3								
Hunt 2011	Prospective cohort	Not serious ^f	NAb	Not serious ^c	Not seriouse	None	103	4.21 (1.36, 13.03)	HIGH
PM _{2.5} (prenata	al exposure)								

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Jedrychowski 2011	Prospective cohort	Seriousª	NAb	Not serious ^c	Serious ^d	None	322	1.06 (0.72, 1.57)	LOW
PM _{2.5} per 8.75	μg/m³ increase								
Jung 2012 b	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	None	408	1.51 (1.05, 2.16)	MODERATE
PM _{2.5} 10.6–13.	2 μg/m³ in infan	ts at risk of asth	nma						
Raaschou- Nielsen 2010	Cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	411	1.32 (0.53, 3.27)	MODERATE
PM _{2.5} 13.2–16.	8 μg/m³ in infan	ts at risk of asth	nma						
Raaschou- Nielsen 2010	Cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	411	1.74 (0.67, 4.47)	MODERATE
PM _{2.5} 16.8–24	μg/m³.in infants	at risk of asthm	na						
Raaschou- Nielsen 2010	Cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	411	0.67 (0.28, 1.59)	MODERATE
PM _{2.5} >24.1 μg	/m³ in infants at	risk of asthma							
Raaschou- Nielsen 2010	Cohort	Not serious ^f	NAb	Not serious ^c	Seriousd	None	411	1.02 (0.41, 2.57)	MODERATE
Slowdown in	children with a	sthma							
Indoor PM _{2.5} (p	oer 10 µg/m³ inc	rease)							
McCormack 2009	Prospective cohort	Seriousª	NAb	Not serious ^c	Not seriouse	None	150	1.04 (1.0, 1.09)	MODERATE
Symptoms wi	th running in c	hildren with as	thma						
Indoor PM _{2.5} (p	er 10 µg/m³ inc	rease)							
McCormack 2009	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^e	None	150	1.07 (1.02,1.11)	MODERATE
Nocturnal syn	nptoms in child	lren with asthr	na						
Indoor PM _{2.5} (p	er 10 μg/m³ inc	rease)							
McCormack 2009	Prospective cohort	Serious ^a	NA ^b	Not serious ^c	Not serious ^e	None	150	1.06 (1.01, 1.10)	MODERATE

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
Limited speed	ch in children w	vith asthma							
Indoor PM _{2.5} (per 10 µg/m³ inc	rease)							
McCormack 2009	Prospective cohort	Serious ^a	NA ^b	Not serious ^c	Not serious ^e	None	150	1.07 (1.00, 1.14)	MODERATE
Rescue medi	cation use in cl	nildren with ast	hma						
Indoor PM _{2.5} (per 10 µg/m³ inc	rease)							
McCormack 2009	Prospective cohort	Serious ^a	NA ^b	Not serious ^c	Not serious ^e	None	150	1.04 (1.01, 1.08)	MODERATE

⁽a) Serious concerns over risk of bias due over self-report of outcomes (b) Not applicable as only one study included

PM² F.2.6.2

۲	IVI ₁₀									
	No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
	Cough, wheezing or chest tightness in children with asthma									
	Indoor PM ₁₀ (per 10 μg/m³ increase)									

⁽c) No concerns over directness

⁽d) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)
(e) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
(f) No concerns over risk of bias

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Number	Adjusted relative effect (aOR 95%Cl unless stated)	Quality
McCormack 2009	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^d	None	150	1.06 (1.01, 1.12)	MODERATE
Slowdown in	children with a	sthma							
Indoor PM ₁₀ (p	er 10 µg/m³ inci	rease)							
McCormack 2009	Prospective cohort	Serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	150	1.08 (1.02, 1.14)	MODERATE
Symptoms w	ith running in c	hildren with as	thma						
Indoor PM ₁₀ (p	er 10 µg/m³ inci	rease)							
McCormack 2009	Prospective cohort	Serious ^a	NA ^b	Not serious ^c	Serious ^e	None	150	1.00 (0.94, 1.08)	LOW
Nocturnal syr	nptoms in chile	dren with asthn	na						
Indoor PM ₁₀ (p	er 10 µg/m³ inci	rease)							
McCormack 2009	Prospective cohort	Serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	150	1.08 (1.01, 1.14)	MODERATE
Limited speed	ch in children v	vith asthma							
Indoor PM ₁₀ (p	er 10 µg/m³ inci	rease)							
McCormack 2009	Prospective cohort	Serious ^a	NA ^b	Not serious ^c	Not serious ^d	None	150	1.11 (1.03, 1.19)	MODERATE
Rescue medi	cation use in cl	hildren with ast	hma						
Indoor PM ₁₀ (p	er 10 µg/m³ inci	rease)							
McCormack 2009	Prospective cohort	Serious ^a	NAb	Not serious ^c	Not serious ^d	None	150	1.06 (1.01, 1.10)	MODERATE

⁽a) Serious concerns over risk of bias due over self-report of exposure(b) Not applicable as only one study included(c) No concerns over directness

⁽d) No concerns as findings are statistically significant (95%Cls do not cross the line of no effect)
(e) Serious concerns as findings are not statistically significant (95%Cls cross the line of no effect)

FINAL						
Association b	etween e	exposure	levels	and he	alth ou	utcome



Appendix G: Economic evidence study selection

No economic evidence review was carried out for this review

Appendix H: Health economic evidence profiles

No economic evidence review was carried out for this review

Appendix I: Health economic analysis

No economic evidence modelling was carried out for this review

Excluded studies

I.1 Public health studies

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	STUDY	REASON FOR EXCLUSION
1.	Abbing-Karahagopian V, van der Gugten, AC, van der Ent et al (2012) Effect of endotoxin and allergens on neonatal lung function and infancy respiratory symptoms and eczema. Pediatric Allergy and Immunology 23(5), 448-455	Study is concerned with bacterial endotoxins
2.	Alderton LE, Spector LG, Blair CK et al (2006) Child and maternal household chemical exposure and the risk of acute leukemia in children with Down's syndrome: a report from the Children's Oncology Group. American journal of epidemiology 164(3), 212-21	Case control study and have included cohort studies of chemical exposure
3.	Aldous M B, Holberg C J, Wright A L, et al (1996) Evaporative cooling and other home factors and lower respiratory tract illness during the first year of life. Group Health Medical Associates. American journal of epidemiology 143(5), 423-30	Study is concerned with evaporative cooling.
4.	Amigou Al, Sermage-FC, Orsi L, et al (2011) Road traffic and childhood leukemia: the ESCALE study (SFCE). Environmental health perspectives 119(4), 566-72	Case control study and have included cohort studies of proximity to traffic
5.	Andersen Z J, Ravnskjer L, Andersen K K, et al (2017) Long- term exposure to fine particulate matter and breast cancer incidence in the Danish nurse cohort study. Cancer Epidemiology Biomarkers and Prevention 26(3), 428-430	Study does not provide data on proximity to traffic
6.	Annesi-Maesano I, Norback D, Zielinski J, et al (2013) Geriatric study in Europe on health effects of air quality in nursing homes (GERIE study) profile: objectives, study protocol and descriptive data. Multidisciplinary Respiratory Medicine. 21;8(1):7	Protocol for a study
7.	Araki A, Kanazawa A, Kawai T, et al (2012) The relationship between exposure to microbial volatile organic compound and allergy prevalence in single-family homes. Science of the Total Environment 423, 18-26	Country not similar to UK
8.	Arif AA, and Shah SM (2007) Association between personal exposure to volatile organic compounds and asthma among US adult population. International archives of occupational and environmental health 80(8), 711-9	Cross-sectional study
9.	Baccarelli Andrea, Martinelli Ida, Pegoraro Valeria, et al (2009) Living near major traffic roads and risk of deep vein thrombosis. Circulation 119(24), 3118-24	Case control study and have included cohort studies of proximity to traffic
10.	Bailey H D, De Klerk , N H, Fritschi L, et al (2011) Refuelling of vehicles, the use of wood burners and the risk of acute lymphoblastic leukaemia in childhood. Paediatric and Perinatal Epidemiology 25(6), 528-539	Case control study and have included cohort studies of heating fuel
11.	Bailey HD, Metayer C, Milne E, et al (2015) Home paint exposures and risk of childhood acute lymphoblastic leukemia: findings from the Childhood Leukemia International Consortium. Cancer Causes and Control 26(9), 1257-1270	Case-control study and have included cohort studies of VOC
12.	Bailey HD, Milne E, de Klerk , NH, et al (2011) Exposure to house painting and the use of floor treatments and the risk of	Case control study and have included

	STUDY	REASON FOR EXCLUSION
	childhood acute lymphoblastic leukemia. International journal of cancer 128(10), 2405-14	cohort studies of VOC
13.	Bakolis I, Heinrich J, Zock J P et al (2015) House dust-mite allergen exposure is associated with serum specific IgE but not with respiratory outcomes. Indoor air 25(3), 235-44	Cross-sectional study
14.	Balmes J R, Cisternas M, Quinlan P J, et al (2014) Annual average ambient particulate matter exposure estimates, measured home particulate matter, and hair nicotine are associated with respiratory outcomes in adults with asthma. Environmental Research 129, 1-10	Cross-sectional study
15.	Barry A C, Mannino D M, Hopenhayn C et al (2010) Exposure to indoor biomass fuel pollutants and asthma prevalence in Southeastern Kentucky: results from the Burden of Lung Disease (BOLD) study. The Journal of asthma: official journal of the Association for the Care of Asthma 47(7), 735-41	Cross-sectional study
16.	Batlles Garrido, J, Torres-Borrego J, Bonillo Perales, A, et al. 2010. "Prevalence and factors linked to atopic eczema in 10-and 11-year-old schoolchildren. Isaac 2 in Almeria, Spain". Allergologia et immunopathologia 38(4):174-80.	Cross-sectional study
17.	Baxter LK, Clougherty JE, Laden F et al (2007) Predictors of concentrations of nitrogen dioxide, fine particulate matter, and particle constituents inside of lower socioeconomic status urban homes Journal of exposure science & environmental epidemiology 17(5), 433-44	Cross-sectional study
18.	Baxter LK, Clougherty JE, Paciorek CJ, et al (2007) Predicting residential indoor concentrations of nitrogen dioxide, fine particulate matter, and elemental carbon using questionnaire and geographic information system based data. Atmospheric Environment 41(31), 6561-6571	Cross-sectional study
19.	Beamer PI, Lothrop N, Lu Z et al (2016) Spatial clusters of child lower respiratory illnesses associated with community-level risk factors. Pediatric pulmonology 51(6), 633-42	Study concerned with spatial analysis and not on poor indoor air quality
20.	Beckett WS, Gent JF, Naeher LP, et al (2006) Peak expiratory flow rate variability is not affected by home combustion sources in a group of nonsmoking women. Archives of Environmental and Occupational Health. ;61(4):176-82	Cross sectional study
21.	Behbod B, Sordillo JE, Hoffman EB et al (2015) Asthma and allergy development: contrasting influences of yeasts and other fungal exposures. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 45(1), 154-63	Study is concerned with fungal concentration and diversity
22.	Behbod B, Sordillo JE, Hoffman EB, et al (2013) Wheeze in infancy: protection associated with yeasts in house dust contrasts with increased risk associated with yeasts in indoor air and other fungal taxa. Allergy 68(11), 1410-8	Study is concerned with fungal concentration and diversity
23.	Bennett CM, Dharmage SC, Matheson M et al (2010) Ambient wood smoke exposure and respiratory symptoms in Tasmania, Australia. The Science of the total environment 409(2), 294-9	Study is concerned with respiratory symptoms and outdoor wood smoke
24.	Bentayeb M, Billionnet C, Baiz N et al (2013) Higher prevalence of breathlessness in elderly exposed to indoor	Cross-sectional study

		REASON FOR
	STUDY	EXCLUSION
	aldehydes and VOCs in a representative sample of French dwellings. Respiratory medicine 107(10), 1598-607	
25.	Bentayeb M, Norback D, Bednarek M et al (2015) Indoor air quality, ventilation and respiratory health in elderly residents living in nursing homes in Europe. The European respiratory journal 45(5), 1228-38	Cross-sectional study
26.	Bjornsson E, Norback D, Janson C, et al. 1995. "Asthmatic symptoms and indoor levels of micro-organisms and house dust mites". Clinical and Experimental Allergy 25(5):423-431.	Case-control study and we have cohorts on allergens
27.	Blount RJ, Pascopella L, Catanzaro DG, et al (2017) Traffic-Related Air Pollution and All-Cause Mortality during Tuberculosis Treatment in California. Environmental health perspectives 125(9), 097026	Study does not report data that can be used
28.	Bornehag CG, Sundell J, Weschler CJ, et al (2004) The association between asthma and allergic symptoms in children and phthalates in house dust: a nested case-control study. Environmental health perspectives 112(14), 1393-7	Nested case-control and we have cohort evidence on this topic
29.	Bothwell J E, McManus L, Crawford VL et al (2003) Home heating and respiratory symptoms among children in Belfast, Northern Ireland. Archives of environmental health 58(9), 549-53	Cross-sectional study
30.	Brown T, Dassonville C, Derbez M et al (2015) Relationships between socioeconomic and lifestyle factors and indoor air quality in French dwellings. Environmental research 140, 385- 96	Cross-sectional survey
31.	Brunekreef B, Smit J, de Jongste J, et al (2002) The prevention and incidence of asthma and mite allergy (PIAMA) birth cohort study: design and first results. Pediatric allergy and immunology: official publication of the European Society of Pediatric Allergy and Immunology 13 Suppl 15, 55-60	Studies do not have any results that can be used
32.	Brussee JE, Smit HA, van Strien , RT, et al (2005) Allergen exposure in infancy and the development of sensitization, wheeze, and asthma at 4 years. The Journal of allergy and clinical immunology 115(5), 946-52	Study report of on risk in terms in terms of categories but reports medians of each category not the range
33.	Bundy K W, Gent J F, Beckett W et al (2009). Household airborne Penicillium associated with peak expiratory flow variability in asthmatic children. Annals of allergy, asthma & immunology: official publication of the American College of Allergy, Asthma, and & Immunology, 103(1), pp.26-30.	Cross-sectional study
34.	Canova C, Jarvis D, Walker S et al (2013). Systematic review of the effects of domestic paints on asthma related symptoms in people with or without asthma. The Journal of asthma: official journal of the Association for the Care of Asthma, 50(10), pp.1020-30.	Systematic review. Checked references for possible includes
35.	Carlos-Wallace FM, Zhang L, Smith MT, et al (2016) Parental, In Utero, and Early-Life Exposure to Benzene and the Risk of Childhood Leukemia: A Meta-Analysis. American journal of epidemiology 183(1), 1-14	Systematic review. Checked references for possible includes
36.	Casas L, Tischer C, Wouters I M et al (2013) Early life microbial exposure and fractional exhaled nitric oxide in school-age children: a prospective birth cohort study.	Study is concerned with bacterial endotoxins

		REASON FOR
	STUDY Environmental health: a global access science source, 12,	EXCLUSION
	pp.103.	
37.	Casas L, Torrent M, Zock J-P, et al (2013) Early life exposures to home dampness, pet ownership and farm animal contact and neuropsylhological development in 4 year old children: a prospective birth cohort study. International journal of hygiene and environmental health 216(6), 690-7	Study do not report on outcomes of interest
38.	Chen CM, Sausenthaler S, Bischof W, et al (2010) Perinatal exposure to endotoxin and the development of eczema during the first 6 years of life. Clinical and experimental dermatology 35(3), 238-44	Study is concerned with bacterial endotoxins
39.	Chew GL, Rogers C, Burge HA, et al (2003) Dustborne and airborne fungal propagules represent a different spectrum of fungi with differing relations to home characteristics. Allergy 58(1), 13-20	Cross-sectional analysis of cohort data
40.	Cho SH, Reponen T, Bernstein DI, et al (2006) The effect of home characteristics on dust antigen concentrations and loads in homes. Science of the Total Environment 371(1-3), 31-43	Cross-sectional analysis of cohort data
41.	Colt JS, Hartge P, Davis S, et al (2007) Hobbies with solvent exposure and risk of non-Hodgkin lymphoma. Cancer causes & control: CCC 18(4), 385-90	Case control study
42.	Crawford J A, Rosenbaum P F, Anagnost S E et al (2015) Indicators of airborne fungal concentrations in urban homes: understanding the conditions that affect indoor fungal exposures. The Science of the total environment 517, 113-24	Study concerned with fungal diversity and fungal concentration
43.	Cuijpers C E, Swaen G M, Wesseling G et al (1995) Adverse effects of the indoor environment on respiratory health in primary school children. Environmental research 68(1), 11-23	Cross-sectional study
44.	Custovic A, Simpson B M, Simpson A, et al (2003) Current mite, cat, and dog allergen exposure, pet ownership, and sensitization to inhalant allergens in adults. The Journal of allergy and clinical immunology 111(2), 402-7	Cross-sectional study
45.	Dales R, Miller D, Ruest K, et al (2006) Airborne endotoxin is associated with respiratory illness in the first 2 years of life. Environmental health perspectives 114(4), 610-4	Study is concerned with bacterial endotoxins
46.	Dallongeville A, Le Cann P , Zmirou-Navier D et al (2015) Concentration and determinants of molds and allergens in indoor air and house dust of French dwellings. The Science of the total environment 536, 964-72	Study concerned with fungal diversity and fungal concentration. Not on risk factors.
47.	Daniel AB, Shah H, Kamath Asha, et al (2012) Environmental tobacco and wood smoke increase the risk of Legg-Calve-Perthes disease. Clinical orthopaedics and related research 470(9), 2369-75	Country not similar to UK
48.	Dannemiller KC, Gent JF, Leaderer BP et al (2016) Influence of housing characteristics on bacterial and fungal communities in homes of asthmatic children. Indoor air 26(2), 179-92	Study interested in housing characteristics and microbial ecology
49.	Dannemiller KC, Gent JF, Leaderer BP, and Peccia Jordan (2016) Indoor microbial communities: Influence on asthma severity in atopic and nonatopic children. The Journal of allergy and clinical immunology 138(1), 76-83.e1	Study is concerned with atopic status and asthma severity

		REASON FOR
	STUDY	EXCLUSION
50.	Dannemiller KC, Mendell MJ, Macher JM et al (2014) Next- generation DNA sequencing reveals that low fungal diversity in house dust is associated with childhood asthma development. Indoor air 24(3), 236-47	Study concerned with fungal diversity and asthma development
51.	Danysh HE, Zhang K, Mitchell LE, et al (2016) Maternal residential proximity to major roadways at delivery and childhood central nervous system tumors. Environmental research 146, 315-22	Case control study and have cohort study on proximity to traffic
52.	de Bilderling G , Mathot M, Agustsson S (2008). Early skin sensitization to aeroallergens. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology, 38(4), pp.643-8.	Study is concerned with early skin testing to aeroallergens and not on indoor pollutants
53.	De Roos, AJ, Koehoorn M, Tamburic L, et al (2014) Proximity to traffic, ambient air pollution, and community noise in relation to incident rheumatoid arthritis. Environmental health perspectives 122(10), 1075-80	Case control study
54.	Dean T, Venter C, Pereira B, et al (2007) Patterns of sensitization to food and aeroallergens in the first 3 years of life. The Journal of allergy and clinical immunology 120(5), 1166-71	Study has no adjustment for confounders
55.	DellaValle CT, Deziel NC, Jones RR, et al (2016) Polycyclic aromatic hydrocarbons: determinants of residential carpet dust levels and risk of non-Hodgkin lymphoma. Cancer causes & control: CCC 27(1), 1-13	Case control study and have cohort study on
56.	Deshmukh JS, Motghare DD, Zodpey SP et al (1998) Low birth weight and associated maternal factors in an urban area. Indian pediatrics 35(1), 33-36	Study is concerned with exposure to tobacco as a risk factor for low birth weight
57.	Dharmage S, Bailey M, Raven J et al (1999) Prevalence and residential determinants of fungi within homes in Melbourne, Australia. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 29(11), 1481-9	Cross-sectional study
58.	Dharmage S, Bailey M, Raven J, et al. 1999. "Residential characteristics influence Der p 1 levels in homes in Melbourne, Australia". Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 29(4):461-9.	Cross-sectional study
59.	Diette B G, Hansel N N, Buckley T J et al (2007) Home indoor pollutant exposures among inner-city children with and without asthma. Environmental health perspectives, 115(11), pp.1665-9.	Cohort study without adjustment for confounding variables
60.	Dong G H, Qian Z, Liu M M et al (2014) Ambient air pollution and the prevalence of obesity in Chinese children: The seven northeastern cities study. Obesity 22(3), 795-800	Country not similar to UK
61.	Dorans KS, Wilker EH, (2017) Residential proximity to major roads, exposure to fine particulate matter and aortic calcium: the Framingham Heart Study, a cohort study. BMJ open 7(3), e013455	Study is concerned with markers for aortic calcification

		REASON FOR
	STUDY	EXCLUSION
62.	Dorans KS, Wilker EH, Li W, et al (2016) Residential Proximity to Major Roads, Exposure to Fine Particulate Matter, and Coronary Artery Calcium. Arteriosclerosis, Thrombosis, and and Vascular Biology 36(8), 1679-85	Study is concerned with markers for aortic calcification
63.	Douwes J, Doekes G, Heinrich J, et al (2004) Endotoxin and $\beta(1\rightarrow 3)$ -Glucan in House Dust and the Relation with Home Characteristics: A Pilot Study in 25 German Houses. Indoor Air 8(4), 255-263	Cross-sectional study
64.	Edwards S C, Jedrychowski W, Butscher M et al (2010) Prenatal exposure to airborne polycyclic aromatic hydrocarbons and children's intelligence at 5 years of age in a prospective cohort study in Poland. Environmental Health Perspectives 118(9), 1326-1331	Study is concerned with outdoor and indoor air pollution and data are not presented separately by source of pollutant
65.	Eiffert S, Noibi Y, Vesper S, et al (2016) A Citizen-Science Study Documents Environmental Exposures and Asthma Prevalence in Two Communities. Journal of environmental and public health, 2016, pp.1962901.	Cross-sectional study
66.	Eisner MD, and Blanc PD (2003) Gas stove use and respiratory health among adults with asthma in NHANES III. Occupational and Environmental Medicine 60(10), 759-764	Cross-sectional study
67.	Emond A M, Howat P, Evans J A, and Hunt L (1997) The effects of housing on the health of preterm infants. Paediatric and perinatal epidemiology 11(2), 228-39	Case control study and have cohort study on preterm, gas ovens, gas stoves and overcrowding
68.	Engvall K, Norrby C, and Norback D (2001) Asthma symptoms in relation to building dampness and odour in older multifamily houses in Stockholm. The international journal of tuberculosis and lung disease: the official journal of the International Union against Tuberculosis and Lung Disease 5(5), 468-77	Cross-sectional study
69.	Engvall K, Norrby C, Bandel J, et al (2001) Development of a Multiple Regression Model to Identify Multi-Family Residential Buildings with a High Prevalence of Sick Building Syndrome (SBS). Indoor Air 10(2), 101-110	Cross-sectional study
70.	Engvall K, Norrby C, and Norback D (2003) Ocular, nasal, dermal and respiratory symptoms in relation to heating, ventilation, energy conservation, and reconstruction of older multi-family houses. Indoor air 13(3), 206-11	Cross-sectional study
71.	Engvall K, Norrby C, and Norback Dan (2002) Ocular, airway, and dermal symptoms related to building dampness and odors in dwellings. Archives of environmental health 57(4), 304-10	Cross-sectional study
72.	Erdmann CA, and Apte MG (2004) Mucous membrane and lower respiratory building related symptoms in relation to indoor carbon dioxide concentrations in the 100-building BASE dataset. Indoor air 14 Suppl 8, 127-34	Study concerned with indoor air quality in the workplace
73.	Farooq U, Joshi M, Nookala V, et al (2010) Self-reported exposure to pesticides in residential settings and risk of breast cancer: a case-control study. Environmental health: a global access science source 9, 30	Case control study and have cohort study on pesticides

		REASON FOR
	STUDY	EXCLUSION
74.	Filippini T, Heck JE, Malagoli C, et al (2015) A review and meta-analysis of outdoor air pollution and risk of childhood leukemia. Journal of environmental science and health. Part C, and Environmental carcinogenesis & ecotoxicology reviews 33(1), 36-66	Systematic review and not relevant to this guideline
75.	Finn P W, Boudreau J O, He H, et al (2000) Children at risk for asthma: Home allergen levels, lymphocyte proliferation, and wheeze. Journal of Allergy and Clinical Immunology 105(5), 933-942	Study does not report complete data
76.	Fleisch AF, Rifas-Shiman SL, Koutrakis P, et al (2015) Prenatal exposure to traffic pollution: associations with reduced fetal growth and rapid infant weight gain. Epidemiology (Cambridge, and Mass.) 26(1), 43-50	Study not concerned with proximity to traffic
77.	Fleisch A F, Luttmann-Gibson H, Perng W, et al (2017) Prenatal and early life exposure to traffic pollution and cardiometabolic health in childhood. Pediatric obesity 12(1), 48-57	Study concerned with markers for cardio-metabolic health
78.	Freedman DM, Stewart P, Kleinerman RA, et al (2001) Household solvent exposures and childhood acute lymphoblastic leukemia. American journal of public health 91(4), 564-7	Case control study and have cohort study on solvents
79.	Gauderman WJ, Vora H, McConnell R, et al (2007) Effect of exposure to traffic on lung development from 10 to 18 years of age: a cohort study. Lancet (London, and England) 369(9561), 571-7	Odds/risk ratios not reported
80.	Gauderman WJ, Avol E, Lurmann F, et al . 2005. "Childhood asthma and exposure to traffic and nitrogen dioxide". Epidemiology (Cambridge, and Mass.) 16(6):737-43.	Study does not present data in a way that can be re-used
81.	Gehring U, Bischof W, Fahlbusch B, et al (2002) House dust endotoxin and allergic sensitization in children. American Journal of Respiratory and Critical Care Medicine 166(7), 939- 944	Study is concerned with bacterial endotoxins
82.	Gehring U, Bolte G, Borte M et al (2001) Exposure to endotoxin decreases the risk of atopic eczema in infancy: a cohort study. The Journal of allergy and clinical immunology 108(5), 847-54	Study is concerned with bacterial endotoxins
83.	Gehring U, Heinrich J, Hoek G et al (2007) Bacteria and mould components in house dust and children's allergic sensitisation. The European respiratory journal 29(6), 1144-53	Case control study and have cohort study on house dust.
84.	Gent J F, Ren P, Belanger K et al (2002). Levels of household mould associated with respiratory symptoms in the first year of life in a cohort at risk for asthma. Environmental health perspectives, 110(12), pp.A781-6.	Study concerned with the microbiological component/diversity of mould
85.	Ghosh R, Amirian E, Dostal M, Sram R J, et al (2011) Indoor coal use and early childhood growth. Archives of Pediatrics and Adolescent Medicine 165(6), 492-497	Study reports on decrease z scores not adjusted OR / RR
86.	Gillespie J, Wickens K, Siebers R, et al (2006) Endotoxin exposure, wheezing, and rash in infancy in a New Zealand birth cohort. The Journal of allergy and clinical immunology 118(6), 1265-70	Study is concerned with bacterial endotoxins

		REASON FOR
	STUDY	EXCLUSION
87.	Godish T (1990) Residential formaldehyde: Increased exposure levels aggravate adverse health effects. Journal of Environmental Health 53(3), 34-37	Study without adjustment for confounding variables
88.	Greenop KR, Peters S, Fritschi L, et al (2014) Exposure to household painting and floor treatments, and parental occupational paint exposure and risk of childhood brain tumors: results from an Australian case-control study. Cancer causes & control: CCC 25(3), 283-91	Case control study and have cohort study on painting
89.	Greenop KR, Hinwood AL, Fritschi L, et al (2015) Vehicle refuelling, use of domestic wood heaters and the risk of childhood brain tumours: Results from an Australian case-control study. Pediatric blood & cancer 62(2), 229-234	Case control study and have cohort on factors of interest
90.	Gross I, Heinrich J, Fahlbusch B, et al (2000) Indoor determinants of Der p 1 and Der f 1 concentrations in house dust are different. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 30(3), 376-82	Cross-sectional analysis of cohort data
91.	Gunnbjornsdottir M I, Franklin K A, Norback D, et al (2006) Prevalence and incidence of respiratory symptoms in relation to indoor dampness: the RHINE study. Thorax 61(3), 221-5	Cross sectional study
92.	Gunnbjornsdottir M I, Norback D, Plaschke P, et al (2003) The relationship between indicators of building dampness and respiratory health in young Swedish adults. Respiratory medicine 97(4), 302-7	Cross sectional study
93.	Guxens M, Aguilera I, Ballester F et al (2012) Prenatal exposure to residential air pollution and infant mental development: modulation by antioxidants and detoxification factors. Environmental health perspectives 120(1), 144-9	Study is concerned with outdoor air pollution
94.	Hagerhed-Engman L, Bornehag CG, and Sundell J (2009) Building characteristics associated with moisture related problems in 8,918 Swedish dwellings International journal of environmental health research 19(4), 251-65	Cross sectional study
95.	Halterman J S, Lynch K A, Conn K M et al (2009) Environmental exposures and respiratory morbidity among very low birth weight infants at 1 year of life. Archives of disease in childhood 94(1), 28-32	Odds/risk ratios for pre-specified pollutants not reported
96.	Harris MH, Gold DR, Rifas-Shiman SL, et al (2015) Prenatal and Childhood Traffic-Related Pollution Exposure and Childhood Cognition in the Project Viva Cohort (Massachusetts, USA). Environmental health perspectives 123(10), 1072-8	Study concerned with markers for cognition
97.	Heinrich J, Topp R, Gehring U, et al (2005) Traffic at residential address, respiratory health, and atopy in adults: the National German Health Survey 1998. Environmental research 98(2), 240-9	Cross sectional study
98.	Herbarth O, Fritz G J, Rehwagen M (2006) Association between indoor renovation activities and eczema in early childhood. International journal of hygiene and environmental health 209(3), 241-7	Cross sectional study

	STUDY	REASON FOR EXCLUSION
99.	Hernberg S, Sripaiboonkij P, Quansah R, et al (2014). Indoor molds and lung function in healthy adults. Respiratory Medicine. 2014 108(5):677-84	Cross sectional study
100.	Hinwood A L, Callan A C, Heyworth J (2014) Polychlorinated biphenyl (PCB) and dioxin concentrations in residential dust of pregnant women. Environmental science. Processes & impacts 16(12), 2758-63	Cross sectional study
101.	Holm S M, Balmes J, Gillette D, et al (2018) Cooking behaviors are related to household particulate matter exposure in children with asthma in the urban East Bay Area of Northern California. PLoS ONE 13(6), e0197199	Study not present usable data
102.	Horick N, Weller E, Milton D K et al (2006) Home endotoxin exposure and wheeze in infants: correction for bias due to exposure measurement error. Environmental health perspectives 114(1), 135-40	Study is concerned with bacterial endotoxins
103.	Houot J, Marquant F, Goujon S, et al (2014) Residential Proximity to Heavy-Traffic Roads, Benzene Exposure, and Childhood Leukemia-The GEOCAP Study, 2002-2007. American Journal of Epidemiology 182(8), 685-693	Case control study and have cohort study on proximity to traffic
104.	Huss K, Adkinson N F, Jr, Eggleston P A et al (2001). House dust mite and cockroach exposure are strong risk factors for positive allergy skin test responses in the Childhood Asthma Management Program. The Journal of allergy and clinical immunology, 107(1), pp.48-54.	Cross sectional study
105.	Hwang B F, Liu I P, and Huang T P (2011) Molds, parental atopy and pediatric incident asthma. Indoor Air 21(6), 472-478	Country not similar to UK
106.	lossifova Y, Reponen T, Sucharew H et al (2008) Use of (1-3)-beta-d-glucan concentrations in dust as a surrogate method for estimating specific fungal exposures. Indoor air 18(3), 225-32	Study is concerned with bacterial endotoxins
107.	lossifova YY, Reponen T, Bernstein DI, et al (2007) House dust (1-3)-beta-D-glucan and wheezing in infants. Allergy 62(5), 504-13	Study is concerned with bacterial endotoxins
108.	Jaakkola M S, Quansah R, Hugg T T, (2013) Association of indoor dampness and molds with rhinitis risk: A systematic review and meta-analysis. Journal of Allergy and Clinical Immunology 132(5), 1099	Systematic review. Checked references for possible includes
109.	Jaakkola MS, Nordman H, Piipari R, et al (2002) Indoor dampness and molds and development of adult-onset asthma: A population-based incident case-control study. Environmental Health Perspectives 110(5), 543-547	Case control study and have cohort study on damp
110.	Jaakkola JJ, Oie L, Nafstad P, et al (1999) Interior surface materials in the home and the development of bronchial obstruction in young children in Oslo, Norway. American journal of public health 89(2), 188-92	Case control study
111.	Jacob B, Ritz B, Gehring U, et al. (2002) Indoor exposure to molds and allergic sensitization. Environmental Health Perspectives. 110(7):647-53	Case control study and have cohort study on damp
112.	Jarvis D, Chinn S, Luczynska C, et al (1997) The association of family size with atopy and atopic disease. Clinical and experimental allergy 27(3), 240-245	Cross sectional study

	STUDY	REASON FOR EXCLUSION
113.	Jarvis D, Zock JP, Heinrich J, et al (2007) Cat and dust mite allergen levels, specific IgG and IgG4, and respiratory symptoms in adults. The Journal of allergy and clinical immunology 119(3), 697-704	Study concerned with exposure to pets and sensitization
114.	Jedrychowski W A, Perera F P, Maugeri U et al (2012) Prohypertensive effect of gestational personal exposure to fine particulate matter. Prospective cohort study in non-smoking and non-obese pregnant women. Cardiovascular toxicology 12(3), 216-25	Study does not report adjusted ratios for risk
115.	Jedrychowski W, Maugeri U, Mroz E, et al. 2012. "Fractional exhaled nitric oxide in healthy non-asthmatic 7-year olds and prenatal exposure to polycyclic aromatic hydrocarbons: nested regression analysis". Pediatric pulmonology 47(11):1131-9.	Study concerned with markers of illness
116.	Jedrychowski W, Maugeri U, Jedrychowska-Bianchi I et al (2002) The effect of house dust mite sensitization on lung size and airway caliber in symptomatic and nonsymptomatic preadolescent children: a community-based study in Poland. Environmental health perspectives 110(6), 571-4	Cross sectional study
117.	Jedrychowski WA, Perera FP, Spengler JD, et al (2013) Intrauterine exposure to fine particulate matter as a risk factor for increased susceptibility to acute broncho-pulmonary infections in early childhood. International journal of hygiene and environmental health 216(4), 395-401	Study reports on risk factors for increased susceptibility to respiratory infections
118.	Jedrychowski W, Maugeri U, Jedrychowska-Bianchi I et al (2005) Effect of indoor air quality in the postnatal period on lung function in pre-adolescent children: a retrospective cohort study in Poland. Public health 119(6), 535-41	Study concerned with a combination of ETS and household heating with no separate data reported
119.	Jedrychowski W, Maugeri U, Perera F, et al (2011) Cognitive function of 6-year old children exposed to mold-contaminated homes in early postnatal period. Prospective birth cohort study in Poland. Physiology & behavior 104(5), 989-95	Study is concerned with duration of exposure
120.	Jedrychowski WA, Maugeri , Spengler J, et al (2013) Dose- dependent relationship between prenatal exposure to fine particulates and exhaled carbon monoxide in non-asthmatic children. A population-based birth cohort study. International journal of occupational medicine and environmental health 26(1), 73-82	Study is concerned with (exhaled Carbon Monoxide) Eco markers
121.	Jedrychowski W, Maugeri U, Zembala M, et al (2007). Risk of wheezing associated with house-dust mite allergens and indoor air quality among three-year-old children. Kraków inner city study. International Journal of Occupational Medicine and Environmental Health. 20(2):117-26	Cross sectional study
122.	Jedrychowski W, Flak E, Mroz E, et al (2008) Modulating effects of maternal fish consumption on the occurrence of respiratory symptoms in early infancy attributed to prenatal exposure to fine particles. Annals of nutrition & metabolism 52(1), 8-16	Study reports on risk for the number of days with symptoms
123.	Jedrychowski WA, Perera FP, Majewska R, et al (2015) Depressed height gain of children associated with intrauterine exposure to polycyclic aromatic hydrocarbons (PAH) and	Study does not report data that can be used.

	OTUDY.	REASON FOR
	heavy metals: the cohort prospective study. Environmental	EXCLUSION
	research 136, 141-7	
124.	Johansen J D, Andersen T F, Thomsen L K, et al. 2000. "Rash related to use of scented products. A questionnaire study in the Danish population. Is the problem increasing?" Contact dermatitis 42(4):222-6.	Cross sectional study
125.	Johansson E, Reponen T, Vesper S et al (2013) Microbial content of household dust associated with exhaled NO in asthmatic children. Environment international 59, 141-7	Study is concerned with bacterial endotoxins
126.	Just A C, Whyatt R M, Miller R L et al (2012) Children's urinary phthalate metabolites and fractional exhaled nitric oxide in an urban cohort. American journal of respiratory and critical care medicine 186(9), 830-7	Cross sectional study
127.	Karr C J, Rudra C B, Miller K A et.al (2009) Infant exposure to fine particulate matter and traffic and risk of hospitalization for RSV bronchiolitis in a region with lower ambient air pollution. Environmental research 109(3), 321-7	Case-control study
128.	Karvonen A M, Hyvarinen A, Gehring U, et al (2012) Exposure to microbial agents in house dust and wheezing, atopic dermatitis and atopic sensitization in early childhood: a birth cohort study in rural areas. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 42(8), 1246-56	Study does not have a multi-variate analysis
129.	Karvonen AM, Hyvarinen A, Roponen M et al. 2009. "Confirmed moisture damage at home, respiratory symptoms and atopy in early life: a birth-cohort study". Pediatrics 124(2):e329-38.	Conference abstract with insufficient detail to assess risk of bias
130.	Kato I, Koenig KL, Watanabe-Meserve H, et al (2005) Personal and occupational exposure to organic solvents and risk of non-Hodgkin's lymphoma (NHL) in women (United States). Cancer causes & control: CCC 16(10), 1215-24	Case control study and have cohort study on solvents
131.	Kidon MI, Chiang WC, Liew WK, et al. (2005) Sensitization to dust mites in children with allergic rhinitis in Singapore: does it matter if you scratch while you sneeze?. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 35(4), 434-40	Country not similar to UK
132.	Kilpelainen M, Koskenvuo M, Helenius H et al (2001) Wood stove heating, asthma and allergies. Respiratory medicine 95(11), 911-6	Cross sectional study
133.	Kingsley SL, Eliot MN, Whitsel EA, et al (2016) Maternal residential proximity to major roadways, birth weight, and placental DNA methylation. Environment international 92-93, 43-9	Study reports results that cannot be disaggregated to distance to road.
134.	Kirjavainen PV, Taubel M, Karvonen AM, et al (2016) Microbial secondary metabolites in homes in association with moisture damage and asthma. Indoor air 26(3), 448-456	Study does not report on outcomes of interest
135.	Kwon J H, Kim E, Chang M et al (2015) Indoor total volatile organic compounds exposure at 6 months followed by atopic dermatitis at 3 years in children. Pediatric allergy and immunology: official publication of the European Society of Pediatric Allergy and Immunology 26(4), 352-8	Country not similar to UK

	OTUDY.	REASON FOR
400	STUDY	EXCLUSION
136.	Langer S, Ramalho O, Le Ponner et al (2017) Perceived indoor air quality and its relationship to air pollutants in French dwellings. Indoor air 27(6), 1168-1176	Study concerned with perceived air quality
137.	Langer S, and Beko G (2013) Indoor air quality in the Swedish housing stock and its dependence on building characteristics. Building & Environment 69, 44-54	Study does not report data that can be used.
138.	Langer S, Ramalho O, Derbez M et al (2016) Indoor environmental quality in French dwellings and building characteristics. Atmospheric Environment 128, 82-91	Study does not report data that can be used.
139.	Leaderer BP, Belanger K, Triche E, et al (2002) Dust mite, cockroach, cat, and dog allergen concentrations in homes of asthmatic children in the Northeastern United States: Impact of socioeconomic factors and population density. Environmental Health Perspectives 110(4), 419-425	Cross sectional study
140.	Lee K, Yanagisawa Y, Spengler JD et al (1996) Classification of House Characteristics in a Boston Residential Nitrogen Dioxide Characterization Study. Indoor Air 6(3), 211-216	Study does not adjust for confounding variables
141.	Levy J I, Welker-Hood L K, Clougherty J E et al (2004) Lung function, asthma symptoms, and quality of life for children in public housing in Boston: a case-series analysis. Environmental health: a global access science source 3(1), 13	Study not concerned with indoor air quality
142.	Lin S, Jones R, Munsie J P, Nayak S G, Fitzgerald E F, and Hwang S A (2012) Childhood asthma and indoor allergen exposure and sensitization in Buffalo, New York. International journal of hygiene and environmental health 215(3), 297-305	Study does not present adjusted OR / RR
143.	Lindfors A, Wickman M, Hedlin G, et al (1995) Indoor environmental risk factors in young asthmatics: a case-control study. Archives of disease in childhood 73(5), 408-12	Study does not present adjusted OR / RR
144.	Lipfert F W, Zhang J, and Wyzga R E (2000) Infant mortality and air pollution: a comprehensive analysis of U.S. data for 1990. Journal of the Air & Waste Management Association (1995) 50(8), 1350-66	Study is concerned with outdoor and indoor air pollution with no disaggregation of data
145.	Litonjua AA, Carey VJ, Burge HA, et al (2001) Exposure to cockroach allergen in the home is associated with incident doctor-diagnosed asthma and recurrent wheezing. Journal of Allergy and Clinical Immunology 107(1), 41-47	Study addressing cockroach allergen.
146.	Liu X, Tan L, Yu I T et al (2018) Household cleaning products and the risk of allergic dermatitis: a prospective cohort study with primary-school children. Journal of the European Academy of Dermatology and Venereology 32(4), 624-631	Country not similar to UK
147.	Llanora G V, Ming L J, Wei L M, Van Bever , and H P S (2012) House dust mite sensitization in toddlers predict persistent wheeze in children between eight to fourteen years old. Asia Pacific Allergy 2(3), 181-186	Country not similar to UK
148.	Lodge CJ, Lowe AJ, Gurrin LC, et al (2011) House dust mite sensitization in toddlers predicts current wheeze at age 12 years. The Journal of allergy and clinical immunology 128(4), 782-788.e9	Study is concerned with sensitization as a risk factor

		REASON FOR
	STUDY	EXCLUSION
149.	Lowe L A, Woodcock A, Murray C S et al (2004) Lung function at age 3 years: effect of pet ownership and exposure to indoor allergens. Archives of paediatrics & adolescent medicine 158(10), 996-1001	Study without adjustment for confounding variables
150.	Lu Y, Lin S, Lawrence W R et al (2018). Evidence from SINPHONIE project: Impact of home environmental exposures on respiratory health among school-age children in Romania. The Science of the total environment, 621, pp.75-84.	Cross sectional study
151.	Ma Xiaomei, Buffler Patricia A, Gunier Robert B, Dahl Gary, Smith Martyn T, Reinier Kyndaron, and Reynolds Peggy (2002) Critical windows of exposure to household pesticides and risk of childhood leukemia. Environmental health perspectives 110(9), 955-60	Case control study
152.	Martins P, Valente J, Papoila A L et al (2012) Combined effect of air pollution and house dust mite exposure over the airways. Revista Portuguesa de Imunoalergologia 20(1), 47-57	Study concerned with air pollution with no separate data for indoor pollutants
153.	Matheson M C, Dharmage S C, Forbes A B, et al. 2003. Residential characteristics predict changes in Der p 1, Fel d 1 and ergosterol but not fungi over time". Clinical and experimental allergy journal of the British Society for Allergy and Clinical Immunology 33(9):1281-8.	Study does not present numeric data that can be used
154.	Matsui EC, Eggleston PA, Buckley TJ, et al (2006) Household mouse allergen exposure and asthma morbidity in inner-city preschool children. Annals of allergy, asthma & immunology: official publication of the American College of Allergy, Asthma, and & Immunology 97(4), 514-20	Study does not report outcome data for all groups
155.	Matsui E C (2014) Environmental exposures and asthma morbidity in children living in urban neighbourhoods. Allergy 69(5), 553-8	Non –systematic overview
156.	Matulonga B, Rava M, Siroux V, et al (2016) Women using bleach for home cleaning are at increased risk of non-allergic asthma. Respiratory medicine 117, 264-71	Cross sectional study
157.	Mazenq J, Dubus J, Gaudart J et al (2017) City housing atmospheric pollutant impact on emergency visit for asthma: A classification and regression tree approach. Respiratory medicine 132, 1-8	Study concerned with outdoor air pollution
158.	McGuinn Laura A, Voss Robert W, Laurent Cecile A, Greenspan Louise C, Kushi Lawrence H, and Windham Gayle C (2016) Residential proximity to traffic and female pubertal development. Environment international 94, 635-641	Odds/risk ratios not reported
159.	Mendy A, Wilkerson J, Salo P M, Cohn R D, Zeldin D C, and Thorne P S (2018) Endotoxin predictors and associated respiratory outcomes differ with climate regions in the U.S. Environment International 112, 218-226	Cross sectional study
160.	Metayer C, Colt JS, Buffler PA, et al (2013) Exposure to herbicides in house dust and risk of childhood acute lymphoblastic leukemia. Journal of exposure science & environmental epidemiology 23(4), 363-70	Case control study and have cohort study on pesticides
161.	Merrett Tg, Burr Ml, Butland Bk, et al (1988) Infant feeding and allergy: 12-month prospective study of 500 babies born into	Study does not report risk as ratios

		REASON FOR
	STUDY	EXCLUSION
	allergic families. Review 53 refs. Annals of allergy 61(6 (Pt 2)), 13-20	
162.	Moran S E, Strachan D P, Johnston I D et al (1999). Effects of exposure to gas cooking in childhood and adulthood on respiratory symptoms, allergic sensitization and lung function in young British adults. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology, 29(8), pp.1033-41.	Study does not report on adjusted data on odds ratio/risk ratio
163.	Morris K, Morgenlander M, Coulehan J L, Gahagen S, and Arena V C (1990) Wood-burning stoves and lower respiratory tract infection in American Indian children. American journal of diseases of children (1960) 144(1), 105-8	Case control study and we have e cohort study on this topic
164.	Moshammer H, Fletcher T, Heinrich J, et al (2010) Gas cooking is associated with small reductions in lung function in children. The European respiratory journal, 36(2), pp.249-54.	Cross sectional study
165.	Munir A K. M, Bjorksten B, Einarsson R, et al (1995) Mite allergens in relation to home conditions and sensitization of asthmatic children from three climatic regions. Allergy: European Journal of Allergy and Clinical Immunology 50(1), 55-64	Cross sectional study
166.	Nafstad P, Jaakkola J J. K, Skrondal A et al (2005) Day care centre characteristics and children's respiratory health. Indoor air 15(2), 69-75	Study concerned with outdoor air quality
167.	Nafstad P, Oie L, Mehl R, et al (1998) Residential dampness problems and symptoms and signs of bronchial obstruction in young Norwegian children. American journal of respiratory and critical care medicine 157(2), 410-4	Case control study and have cohort study on dampness
168.	Narayan S, Liew Z, Paul K, et al(2013) Household organophosphorus pesticide use and Parkinson's disease. International journal of epidemiology 42(5), 1476-85	Case control study and have cohort study on pesticides
169.	Nguyen T, Lurie M, Gomez M (2010) The National Asthma SurveyNew York State: association of the home environment with current asthma status. Public health reports (Washington, and D.C.: 1974) 125(6), 877-87	Cross sectional study
170.	Nicolaou N, Yiallouros P, Pipis S, et al (2006) Domestic allergen and endotoxin exposure and allergic sensitization in Cyprus. Pediatric allergy and immunology: official publication of the European Society of Pediatric Allergy and Immunology 17(1), 17-21	Case control study and have cohort data on allergen exposure
171.	Norback D, Bjornsson E, Janson C, et al (1999) Current asthma and biochemical signs of inflammation in relation to building dampness in dwellings. The international journal of tuberculosis and lung disease: the official journal of the International Union against Tuberculosis and Lung Disease 3(5), 368-76	Case control study and have cohort study on damp
172.	Norback D, Lampa E, and Engvall K (2014) Asthma, allergy and eczema among adults in multifamily houses in Stockholm (3-HE study)associations with building characteristics, home environment and energy use for heating. PloS one 9(12), e112960	Cross sectional study
173.	Norback D, Zock J P, Plana E, et al (2017) Building dampness and mold in European homes in relation to climate, building characteristics and socio-economic status: The European	Cross sectional study

	STUDY	REASON FOR EXCLUSION
	Community Respiratory Health Survey ECRHS II. Indoor air 27(5), 921-932	
174.	Norback D, Zock J-P, Plana E, et al (2011) Lung function decline in relation to mould and dampness in the home: the longitudinal European Community Respiratory Health Survey ECRHS II. Thorax 66(5), 396-401	Study concerned with lung function not symptoms
175.	Oudin A, Segersson D, Adolfsson R, et al. 2018. "Association between air pollution from residential wood burning and dementia incidence in a longitudinal study in Northern Sweden". PLoS ONE 13(6):e0198283.	Study is concerned with indoor and outdoor pollution
176.	Park D-U, Choi Y-Y, Ahn J-J, et al (2015) Relationship between Exposure to Household Humidifier Disinfectants and Risk of Lung Injury: A Family-Based Study. PloS one 10(5), e0124610	Country not similar to UK
177.	Park JH, Gold DR, Spiegelman DL, et al (2001) House dust endotoxin and wheeze in the first year of life. American journal of respiratory and critical care medicine 163(2), 322-8	Study is considered with bacterial endotoxin
178.	Paulin L M, Williams D L, Peng R et al (2017). 24-h Nitrogen dioxide concentration is associated with cooking behaviors and an increase in rescue medication use in children with asthma. Environmental research, 159, pp.118-123.	Study does not reported results in a way that can be used
179.	Pekkanen J, Hyvarinen A, Haverinen-Shaughnessy U, et al (2007) Moisture damage and childhood asthma: A population-based incident case-control study. European Respiratory Journal 29(3), 509-515	Case control study and have cohort study on damp
180.	Perera Frederica P (2009) Prenatal airborne polycyclic aromatic hydrocarbon exposure and child IQ at age 5 years. Pediatrics 124(2),	Odds/risk ratios not reported
181.	Perry TT, Wood RA, Matsui EC, et al (2006) Room-specific characteristics of suburban homes as predictors of indoor allergen concentrations. Annals of Allergy, and Asthma and Immunology 97(5), 628-635	Cross sectional study
182.	Perzanowski MS, Chew GL, Divjan A, et al (2013) Early-life cockroach allergen and polycyclic aromatic hydrocarbon exposures predict cockroach sensitization among inner-city children. The Journal of allergy and clinical immunology 131(3), 886-93	Study reports on risk factors for sensitization
183.	Perzanowski MS, Ronmark E, James HR, et al (2016) Relevance of specific IgE antibody titer to the prevalence, severity, and persistence of asthma among 19-year-olds in northern Sweden. The Journal of allergy and clinical immunology 138(6), 1582-1590	Study reports on risk factors for sensitization
184.	Perzanowski MS, Miller RL, Thorne PS, et al (2006) Endotoxin in inner-city homes: associations with wheeze and eczema in early childhood. The Journal of allergy and clinical immunology 117(5), 1082-9	Study is considered with bacterial endotoxin
185.	Peters J L, Levy J I, Rogers C A, et al (2007) Determinants of allergen concentrations in apartments of asthmatic children living in public housing. Journal of Urban Health 84(2), 185-197	Cross sectional study
186.	Phipatanakul W, Celedon JC, Raby BA, et al (2004) Endotoxin exposure and eczema in the first year of life. Pediatrics 114(1), 13-8	Study is considered with bacterial endotoxin

	STUDY	REASON FOR EXCLUSION
187.	Phipatanakul W, Gold DR, Muilenberg M, Sredl DL, Weiss ST, and Celedon JC (2005) Predictors of indoor exposure to mouse allergen in urban and suburban homes in Boston. Allergy 60(5), 697-701	Cross sectional study
188.	Pogoda J M, and Preston-Martin S (1997) Household pesticides and risk of pediatric brain tumors. Environmental health perspectives 105(11), 1214-20	Case control study and have cohort study data n pesticides
189.	Ponsonby AL, Dwyer T, Kemp A, et al (2003) The use of mutually exclusive categories for atopic sensitization: A contrasting effect for family size on house dust mite sensitization compared with ryegrass sensitization. Pediatric Allergy and Immunology 14(2), 81-90	Study reports on risk factors for sensitization
190.	Poynter JN, Richardson M, Roesler M, et al (2017) Chemical exposures and risk of acute myeloid leukemia and myelodysplastic syndromes in a population-based study. International journal of cancer 140(1), 23-33	Study concerned with occupational exposure to chemicals
191.	Quansah R, Jaakkola MS, Hugg TT, et al (2012) Residential dampness and molds and the risk of developing asthma: a systematic review and meta-analysis. PloS one 7(11), e47526	Systematic review. Checked references for possible includes
192.	Rabito F A, Carlson J, Holt E W, et al. 2011. "Cockroach exposure independent of sensitization status and association with hospitalizations for asthma in inner-city children". Annals of Allergy, and Asthma and Immunology 106(2):103-109.	Cross sectional study
193.	Ramagopal M, Wang Z, Black K, et al (2014) Improved exposure characterization with robotic (PIPER) sampling and association with children's respiratory symptoms, asthma and eczema. Journal of exposure science & environmental epidemiology 24(4), 421-7	Cross sectional study
194.	Rauh VA, Chew GR, and Garfinkel RS (2002) Deteriorated housing contributes to high cockroach allergen levels in innercity households. Environ Health Perspect. 110 (Suppl 2): 323–327.	Cross sectional analysis of cohort data
195.	Reding KW, Young MT, Szpiro AA, H et al (2015) Breast Cancer Risk in Relation to Ambient Air Pollution Exposure at Residences in the Sister Study Cohort. Cancer Epidemiology Biomarkers & Prevention 24(12), 1907-1909	Study is not concerned with indoor air
196.	Ren P, Jankun TM, Belanger K, et al (2001) The relation between fungal propagules in indoor air and home characteristics. Allergy 56(5), 419-24	Cross sectional analysis of cohort data
197.	Rios P, Bailey H D, Lacour B, et al (2017) Maternal use of household pesticides during pregnancy and risk of neuroblastoma in offspring. A pooled analysis of the ESTELLE and ESCALE French studies (SFCE). Cancer Causes and Control 28(10), 1125-1132	Pooled analysis of 2 case-control studies
198.	Rokoff LB, Koutrakis P, Garshick E, et al (2017) Wood Stove Pollution in the Developed World: A Case to Raise Awareness Among Pediatricians. Current problems in pediatric and adolescent health care 47(6), 123-141	Systematic review. Checked references for possible includes
199.	Rosenbaum PF, Crawford JA, Anagnost SE et al (2010) Indoor airborne fungi and wheeze in the first year of life among a	Study is concerned with bacterial endotoxin

	STUDY	REASON FOR EXCLUSION
	cohort of infants at risk for asthma. Journal of exposure science & environmental epidemiology 20(6), 503-15	
200.	Rosenfeld L, Chew GL, Rudd R, et al (2011) Are building-level characteristics associated with indoor allergens in the household? Journal of urban health: bulletin of the New York Academy of Medicine 88(1), 14-29	Cross sectional analysis of cohort data
201.	Ruckart PZ, Bove FJ, Shanley E 3rd, et al (2015) Evaluation of contaminated drinking water and male breast cancer at Marine Corps Base Camp Lejeune, North Carolina: a case control study. Environmental health: a global access science source 14, 74	Study is not concerned with indoor air pollution
202.	Sahlberg B, Gunnbjornsdottir M, Soon A et al (2013) Airborne moulds and bacteria, microbial volatile organic compounds (MVOC), plasticizers and formaldehyde in dwellings in three North European cities in relation to sick building syndrome (SBS). The Science of the total environment 444, 433-40	Cross sectional study
203.	Salo P M, Wilkerson J, Rose K M, et al (2018) Bedroom allergen exposures in US households. Journal of Allergy and Clinical Immunology 141(5), 1870	Cross sectional study
204.	Sapkota A, Zaridze D, Szeszenia-Dabrowska N et al (2013) Indoor air pollution from solid fuels and risk of upper aerodigestive tract cancers in central and eastern Europe. Environmental research 120, 90-5	Case-control study and have cohort studies on heating fuel
205.	Scelo G, Metayer C, Zhang L, et al (2009) Household exposure to paint and petroleum solvents, chromosomal translocations, and the risk of childhood leukemia. Environmental health perspectives 117(1), 133-9	Case control study and have cohort studies on paint
206.	Schenker MB, Samet JM, and Speizer FE (1983) Risk factors for childhood respiratory disease. The effect of host factors and home environmental exposures. The American review of respiratory disease 128(6), 1038-43	Study does not report results that can be re-used
207.	Schindler C, Keidel D, Gerbase MW, et al (2009) Improvements in PM ₁₀ exposure and reduced rates of respiratory symptoms in a cohort of Swiss adults (SAPALDIA). American journal of respiratory and critical care medicine 179(7), 579-87	Study does not report results that can be re-used
208.	Seo S, Han Y, Kim J, Choung J T, et al (2014) Infrared camera-proven water-damaged homes are associated with the severity of atopic dermatitis in children. Annals of Allergy, and Asthma and Immunology 113(5), 549-555	Country not similar to UK
209.	Sharpe R A, Bearman N, Thornton C R, et al (2015) Indoor fungal diversity and asthma: A meta-analysis and systematic review of risk factors. Journal of Allergy and Clinical Immunology 135(1), 110-122	Systematic review. Checked references for possible includes
210.	Sharpe R A, Thornton C R, Tyrrell J, et al 2015. Variable risk of atopic disease due to indoor fungal exposure in NHANES 2005-2006. Clinical and Experimental Allergy 45(10):1566-1578.	Cross sectional study
211.	Sharpe RA, Thornton CR, Nikolaou V, et al (2015) Higher energy efficient homes are associated with increased risk of doctor diagnosed asthma in a UK subpopulation. Environment international 75, 234-44	Cross sectional study

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	STUDY	EXCLUSION
212.	Sharpe RA, Thornton CR, Nikolaou V, et al (2015) Fuel poverty increases risk of mould contamination, regardless of adult risk perception & ventilation in social housing properties. Environment International 79, 115-129	Cross sectional study
213.	Shenassa ED, Daskalakis C, Liebhaber A, et al (2007) Dampness and mold in the home and depression: An examination of mold-related illness and perceived control of one's home as possible depression pathways. American Journal of Public Health 97(10), 1893-1899	Cross sectional study
214.	Shorter C, Crane J, Pierse N, et al (2017) Indoor visible mold and mold odor are associated with new-onset childhood wheeze in a dose-dependent manner. Indoor Air 28(1), 6-15	Case control study and have cohorts on this topic
215.	Singh U, Levin L, Grinshpun SA et al (2011) Influence of home characteristics on airborne and dust borne endotoxin and beta-D-glucan. Journal of environmental monitoring: JEM 13(11), 3246-53	Study is concerned in bacterial endotoxins
216.	Slater ME, Linabery AM, Spector LG, et al (2011) Maternal exposure to household chemicals and risk of infant leukemia: a report from the Children's Oncology Group. Cancer causes & control: CCC 22(8), 1197-204	Case control study and have cohort studies on chemicals
217.	Smedje G, Wang J, Norback D, et al (2017) SBS symptoms in relation to dampness and ventilation in inspected single-family houses in Sweden. International archives of occupational and environmental health 90(7), 703-711	Cross sectional study
218.	Smith B J, Nitschke M, Pilotto L S, et al (2000) Health effects of daily indoor nitrogen dioxide exposure in people with asthma. European Respiratory Journal 16(5), 879-885	Study does not use regression analysis to identify sources of NO ₂
219.	Sordillo JE, Hoffman EB, Celedon JC, et al (2010) Multiple microbial exposures in the home may protect against asthma or allergy in childhood. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 40(6), 902-10	Study is concerned in bacterial endotoxins
220.	Sordillo J E, Alwis UK, Hoffman E, et al. 2011. "Home characteristics as predictors of bacterial and fungal microbial biomarkers in house dust". Environmental health perspectives 119(2):189-95.	Study concerns with microbial biomarkers in house dust
221.	Spilak MP, Madsen AM, Knudsen SM et al(2015) Impact of dwelling characteristics on concentrations of bacteria, fungi, endotoxin and total inflammatory potential in settled dust. Building & Environment 93, 64-71	Cross sectional study
222.	Sporik R, Holgate ST, Platts-Mills TA, et al (1990) Exposure to house-dust mite allergen (Der p I) and the development of asthma in childhood. A prospective study. The New England journal of medicine 323(8), 502-7	Study does not report results that can be re-used
223.	Squance M L, Reeves G, Attia J, et al (2015) Self-reported Lupus flare: Association with everyday home and personal product exposure. Toxicology Reports 2, 880-888	Case control study and have cohort studies on personal products
224.	Stankovic A, Nikolic M, and Arandjelovic M (2011) Effects of indoor air pollution on respiratory symptoms of non-smoking	Country not similar to UK

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	women in Nis, Serbia. Multidisciplinary respiratory medicine 6(6), 351-5	EXCLUSION
225.	Strachan D P (1988) Damp housing and childhood asthma: validation of reporting of symptoms. BMJ (Clinical research ed.) 297(6658), 1223-6	Cross sectional study
226.	Strachan D P, and Carey I M (1995) Home environment and severe asthma in adolescence: a population based case-control study. BMJ (Clinical research ed.) 311(7012), 1053-6	Case control study
227.	Strumylaite L, and Kregzdyte R (2006) Household gas cooking and respiratory health in preschool children. Family Medicine and Primary Care Review 8(1), 21-25	Cross sectional study
228.	Taha AA. ER, Etewa SE, Abdel-Rahman SA, et al (2018) House dust mites among allergic patients at the Allergy and Immunology Unit, Zagazig University: an immunologic and serologic study. Journal of Parasitic Diseases 42(3), 405-415	Country not similar to UK
229.	Takeda M, Saijo Y, Yuasa M et al (2009) Relationship between sick building syndrome and indoor environmental factors in newly built Japanese dwellings. International archives of occupational and environmental health 82(5), 583-93	Country not similar to UK
230.	Tavernier G O. G, Fletcher G D, Francis H C et al (2005) Endotoxin exposure in asthmatic children and matched healthy controls: results of IPEADAM study. Indoor air 15 Suppl 10, 25- 32	Cross sectional study
231.	Tavernier G, Fletcher G, Gee I et al (2006) IPEADAM study: indoor endotoxin exposure, family status, and some housing characteristics in English children. The Journal of allergy and clinical immunology 117(3), 656-62	Cross-sectional study
232.	Tetreault L F, Doucet M, Gamache P, et al (2016) Childhood exposure to ambient air pollutants and the onset of asthma: An administrative cohort study in Quebec. Environmental Health Perspectives 124(8), 1276-1282	Study is not concerned with indoor air pollution
233.	Thorn J, Brisman J, and Toren K. 2001. "Adult-onset asthma is associated with self-reported mold or environmental tobacco smoke exposures in the home". Allergy: European Journal of Allergy and Clinical Immunology 56(4):287-292.	Case-control study and we have cohort studies on mould
234.	Tischer C G, Gref A, Standl M, et al (2013) Glutathione-S-transferase P1, early exposure to mould in relation to respiratory and allergic health outcomes in children from six birth cohorts. A meta-analysis. Allergy 68(3), 339-46	Systematic review. Checked references for possible includes
235.	Tischer C, Chen C M, and Heinrich J (2011) Association between domestic mould and mould components, and asthma and allergy in children: a systematic review. The European respiratory journal 38(4), 812-24	Systematic review. Checked references for possible includes
236.	Tischer C, Casas L, Wouters IM, et al (2015) Early exposure to bio-contaminants and asthma up to 10 years of age: results of the HITEA study. The European respiratory journal 45(2), 328-37	Study is concerned in bacterial endotoxins
237.	Tischer C G, Hohmann C, Thiering E, et al (2011) Meta- analysis of mould and dampness exposure on asthma and allergy in eight European birth cohorts: an ENRIECO initiative. Allergy 66(12), 1570-9	Systematic review. Checked references for possible includes

	STUDY	REASON FOR EXCLUSION
238.	Tischer C, Weikl F, Probst AJ, et al (2016) Urban Dust Microbiome: Impact on Later Atopy and Wheezing. Environmental health perspectives 124(12), 1919-1923	Study concerned with fungal ddiversity
239.	Trevillian LF, Ponsonby AL, Dwyer T, et al (2003) An association between plastic mattress covers and sheepskin underbedding use in infancy and house dust mite sensitization in childhood: a prospective study. Clinical and experimental allergy: journal of the British Society for Allergy and Clinical Immunology 33(4), 483-9	Study concerned with sensitization
240.	Trupin L, Balmes J R, Chen H et al (2010) An integrated model of environmental factors in adult asthma lung function and disease severity: a cross-sectional study. Environmental health: a global access science source 9, 24	Cross sectional study
241.	Turunen M, Iso-Markku K, Pekkonen M, et al (2017) Statistical associations between housing quality and health among Finnish households with children - Results from two (repeated) national surveys. Science of the Total Environment 574, 1580-1587	Study does not report longitudinal data
242.	Ulrik CS, Backer V, Hesse B, et al (1996) Risk factors for development of asthma in children and adolescents: findings from a longitudinal population study. Respiratory medicine 90(10), 623-30	Study does not report on prognostic factors
243.	van Rossem L, Rifas-Shiman SL, Melly SJ, et al (2015) Prenatal air pollution exposure and newborn blood pressure. Environmental health perspectives 123(4), 353-9	Study is not concerned with indoor air pollution
244.	Venn A J, Cooper M, Antoniak M et al (2003) Effects of volatile organic compounds, damp, and other environmental exposures in the home on wheezing illness in children. Thorax 58(11), 955-60	Case-control study and have cohort studies on VOC
245.	Vesper SJ, McKinstry C, Haugland RA, et al (2007) Relative moldiness index as predictor of childhood respiratory illness. Journal of exposure science & environmental epidemiology 17(1), 88-94	Study does not report on risk as an outcome
246.	Viegi G, Paoletti P, Carrozzi L, et al (1991) Effects of home environment on respiratory symptoms and lung function in a general population sample in north Italy. The European respiratory journal 4(5), 580-6	Cross sectional study
247.	Vilcekova S, Apostoloski I Z, Meciarova L et al (2017) Investigation of Indoor Air Quality in Houses of Macedonia. International journal of environmental research and public health 14(1),	Country not similar to UK
248.	Volk HE, Hertz-Picciotto I, Delwiche L, et al (2011) Residential proximity to freeways and autism in the CHARGE study. Environmental health perspectives 119(6), 873-7	Case control study and have cohort studies on proximity to traffic
249.	Volk HE, Lurmann F, Penfold B, et al (2013) Traffic-related air pollution, particulate matter, and autism. JAMA psychiatry 70(1), 71-7	Study is not concerned with proximity to traffic
250.	Volkmer R E, Ruffin R E, Wigg N R et al (1995) The prevalence of respiratory symptoms in South Australian preschool children. II. Factors associated with indoor air quality. Journal of paediatrics and child health 31(2), 116-20	Cross-sectional study

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251.	Wallace J, D'Silva L, Brannan J, et al . 2011. "Association	Study concerned
	between proximity to major roads and sputum cell counts". Canadian respiratory journal 18(1):13-8.	with markers of illness
252.	Wang J, Cozen W, Thorne PS, et al (2013) Household endotoxin levels and the risk of non-Hodgkin lymphoma. Cancer causes & control: CCC 24(2), 357-64	Study is concerned in bacterial endotoxins
253.	Wang J, Engvall K, Smedje G, et al (2014) Rhinitis, asthma and respiratory infections among adults in relation to the home environment in multi-family buildings in Sweden. PloS one 9(8), e105125	Cross sectional study
254.	Wang J, Engvall K, Smedje G, et al (2017) Current wheeze, asthma, respiratory infections, and rhinitis among adults in relation to inspection data and indoor measurements in single-family houses in Sweden-The BETSI study. Indoor air 27(4), 725-736	Cross sectional study
255.	Wang L, Hu W, Guan Q et al (2018). The association between cooking oil fume exposure during pregnancy and birth weight: A prospective mother-child cohort study. The Science of the total environment, 612, pp.822-830.	Country not similar to the UK
256.	Ward MH, Colt JS, Deziel NC, et al (2014) Residential levels of polybrominated diphenyl ethers and risk of childhood acute lymphoblastic leukemia in California. Environmental health perspectives 122(10), 1110-6	Case control study and have cohort studies on VOC
257.	Ward MH, Colt JS, Metayer C, et al (2009) Residential exposure to polychlorinated biphenyls and organochlorine pesticides and risk of childhood leukemia. Environmental health perspectives 117(6), 1007-13	Case control study and have cohort studies on VOC
258.	Ware J H, Dockery D W, Spiro A, 3rd , Speizer F E, Ferris B G, and Jr (1984) Passive smoking, gas cooking, and respiratory health of children living in six cities. The American review of respiratory disease 129(3), 366-74	Study does not report results that can be re-used
259.	Webb E, Blane D, de Vries , and Robert . 2013. "Housing and respiratory health at older ages". Journal of epidemiology and community health 67(3):280-5.	Study concerned with indicators of poor respiratory health
260.	Wegienka G, Johnson CC, Havstad S, et al (2010) Indoor pet exposure and the outcomes of total IgE and sensitization at age 18 years. Journal of Allergy and Clinical Immunology 126(2), 274	Study did not adjust for confounders
261.	White A J, Teitelbaum SL, Stellman S D, et al (2014) Indoor air pollution exposure from use of indoor stoves and fireplaces in association with breast cancer: a case-control study. Environmental Health: A Global Access Science Source 13(1), 135-158	Case control study and have cohort studies on heating
262.	White AJ, Bradshaw PT, Herring AH, et al (2016) Exposure to multiple sources of polycyclic aromatic hydrocarbons and breast cancer incidence. Environment International 89, 185-192	Case control study and have cohort studies on PAH
263.	Wickens K, Douwes J, Siebers R, et al (2003) Determinants of endotoxin levels in carpets in New Zealand homes. Indoor air 13(2), 128-35	Study is concerned with endotoxins

	STUDY	REASON FOR EXCLUSION
264.	Wilhelm M, and Ritz B (2003) Residential proximity to traffic and adverse birth outcomes in Los Angeles county, California, 1994-1996. Environmental health perspectives 111(2), 207-16	Case control study and we have cohort studies on proximity to traffic
265.	Wilker Elissa H, Martinez-Ramirez Sergi, Kloog Itai et.al (2016) Fine Particulate Matter, Residential Proximity to Major Roads, and Markers of Small Vessel Disease in a Memory Study Population. Journal of Alzheimer's disease: JAD 53(4), 1315-23	Study concerned with markers of disease
266.	Williamson IJ, Martin CJ, McGill G, et al (1997) Damp housing and asthma: a case-control study. Thorax 52(3), 229-34	Case control study and have cohort studies on damp
267.	Wilson J, Dixon SL, Breysse P, et al (2010) Housing and allergens: a pooled analysis of nine US studies. Environmental research 110(2), 189-98	Systematic review. Checked references for possible includes
268.	Wong G W. K, Brunekreef B, Ellwood P et al (2013) Cooking fuels and prevalence of asthma: a global analysis of phase three of the International Study of Asthma and Allergies in Childhood (ISAAC). The Lancet. Respiratory medicine 1(5), 386-94	Data not reported separately for countries similar to the UK
269.	Xu X, and Wang L (1993) Association of indoor and outdoor particulate level with chronic respiratory illness. American Review of Respiratory Disease 148(6 I), 1516-1522	Country not similar to the UK
270.	Yang A, Janssen NA, Brunekreef B, et al (2016) Children's respiratory health and oxidative potential of $PM_{2.5}$: the PIAMA birth cohort study. Occupational and environmental medicine 73(3), 154-60	Study did not measure indoor air quality
271.	Yang S I, Kim B J, Kim H B, et al (2015) Prenatal particulate matter/tobacco smoke increases infants' respiratory infections: COCOA study. Allergy, and Asthma and Immunology Research 7(6), 573-582	Country not similar to UK
272.	Zacharasiewicz A, Zidek T, Haidinger G et al (1999) Indoor factors and their association to respiratory symptoms suggestive of asthma in Austrian children aged 6-9 years. Wiener klinische Wochenschrift 111(21), 882-6	Cross-sectional study
273.	Zejda J E, and Kowalska M. (2003). Risk factors for asthma in school childrenresults of a seven-year follow-up. Central European journal of public health, 11(3), pp.149-54.	Study does not report on adjusted data on odds ratio/risk ratio
274.	Zhang G, Spickett J, Lee A H, et al. 2006. Ever eczema and itchy rash in relation to domestic environments in primary school children. Indoor and Built Environment 15(6):535-541.	Cross sectional study
275.	Zhao Zhiqing, Lin Faying, Wang Bennett, Cao Yihai, Hou Xu, and Wang Yangang (2016) Residential Proximity to Major Roadways and Risk of Type 2 Diabetes Mellitus: A Meta-Analysis. International journal of environmental research and public health 14(1),	Systematic review. Checked references for possible includes
276.	Zock JP, Plana E, Anto JM, et al (2009) Domestic use of hypochlorite bleach, atopic sensitization, and respiratory symptoms in adults. Journal of Allergy and Clinical Immunology 124(4), 731	Pollutant not of interest

	STUDY	REASON FOR EXCLUSION
277.	Zota AR, Aschengrau A, Rudel RA, et al (2010) Self-reported chemicals exposure, beliefs about disease causation, and risk of breast cancer in the Cape Cod Breast Cancer and Environment Study: a case-control study. Environmental health: a global access science source 9, 40	Case control study
278.	Zota A, Adamkiewicz G, Levy JI, et al (2005) Ventilation in public housing: implications for indoor nitrogen dioxide concentrations. Indoor air 15(6), 393-401	Cross sectional study

I.2 Economic studies

No economic evidence review was carried out for this review

Appendix J: Research recommendation

J.1.1 Health impact of air pollutants at home

What is the health impact of exposure to individual air pollutants alone or combined with each other in the home?

Population	Adults and children
Prognostic factors, exposure	Health impact of exposure to individual and combined air pollutants Respiratory health effects Allergic health effects Cardiac health effects Pregnancy related health effects Cancer health effects
Outcomes	Adjusted risk ratios and odd ratios reported for health risk associated with prognostic factor(s) and the Indoor air pollutants
Study design	Cohort study design with multivariate analysis adjusting for variables that might confound results. For example, ingress of outdoor air pollution
Timeframe	At least 1 year follow up

Rationale: People spend up to 90% of their lives indoors and 60% of that time at home. To minimize the health risks from pollutants occurring in homes, exposures to these pollutants should be controlled. Exposure to individual or combined pollutants is very common in practice. Research into these and on pollutants combined with other stressors such as noise could help better assess priorities for regulation and interventions. Also evidence about harms, both in the short and longer term, that may be associated with these pollutants would improve understanding and educate people

