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Prenatal maternal psychosocial stress and offspring's asthma and allergic disease: a systematic review and meta-analysis

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38 **ABSTRACT**

39

40 **Background:** Prenatal maternal stress may influence offspring's atopic risk through
41 sustained cortisol secretion resulting from activation of the hypothalamic-pituitary-axis (HPA),
42 leading to Th2-biased cell differentiation in the fetus. We undertook a systematic review and
43 meta-analysis investigating the relationship between prenatal maternal psychosocial stress
44 and risk of asthma and allergy in the offspring.

45 **Design:**

46 **Methods:** We searched 11 electronic databases from 1960 to 2016, search the grey
47 literature, and contacted experts in the field. Type of stress indicator included mood
48 disorders, anxiety, exposure to violence, bereavement and socio-economic problems
49 occurring during pregnancy, both objectively or subjectively measured. We included all
50 possible asthma and IgE-mediated allergy outcomes. We conducted random-effects meta-
51 analyses to synthesize the data.

52

53 **Results:** We identified 9,779 papers of which 30 studies (enrolling >6 million participants)
54 satisfied inclusion criteria. The quality of 25 studies was moderate, four were strong, and one
55 was weak. Maternal exposure to any type of stressors was associated with an increased risk
56 of offspring atopic eczema/dermatitis (OR 1.34, 95%CI 1.22-1.47), allergic rhinitis (OR 1.30,
57 95%CI 1.04-1.62), wheeze (OR 1.34, 95%CI 1.16-1.54) and asthma (OR 1.15, 95%CI 1.04-
58 1.27). Exposure to anxiety and depression had strongest effect compared to other stressors.
59 Exposure during the third trimester had the greatest impact compared to first and second
60 trimesters. The increased risk was stronger for early-onset and persistent than for late-onset
61 wheeze. Bereavement of a child (HR 1.28, 95%CI 1.10-1.48) or a spouse (HR 1.40, 95%CI
62 1.03-1.90) increased the risk of offspring asthma.

63

64 **Conclusions:** Exposure to prenatal maternal psychosocial stress was associated with
65 increased risk, albeit modestly, of asthma and allergy in the offspring. The pronounced risk
66 during the third trimester may represent cumulative stress exposure throughout pregnancy
67 rather than trimester-specific effect. Our findings may represent a causal effect or a result of
68 inherent biases in studies, particularly residual confounding.

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73 **Systematic review registration:** PROSPERO (2016:CRD42016036456)

74

75

76 INTRODUCTION

77 The susceptibility to develop asthma and allergy may be established already in utero.¹⁻⁴ The
78 concept of fetal programming has provided important insights on the influence of the
79 intrauterine environment on the development of the fetus and subsequent⁵ risk of chronic
80 conditions later in life.⁶ As adaptive immunity develops prenatally, allergen specific immune
81 responses are demonstrable in newborns^{2,3,7} with umbilical cord blood shown to contain
82 fetally derived immunoglobulin E (IgE).^{1,3}

83

84 One suggested pathway through which prenatal maternal stress may influence the risk of
85 asthma and allergy in the offspring is through the activation of the hypothalamic-pituitary-axis
86 (HPA) in response to external stress.^{8,9} This then causes secretion of hormones by the
87 hypothalamus and pituitary gland in the brain and subsequent stimulation of the release of
88 glucocorticoids, adrenaline and noradrenaline.⁸⁻¹⁰ The release of glucocorticoids leads to
89 increases in cortisol levels.^{8,9} The activities of the HPA and the resultant chemical reactions
90 can be transmitted to the fetus and thus influences development.^{6,11} High levels of cortisol
91 increase airway responsiveness in the offspring and potentiated cell differentiation from T
92 helper cell type 1 (Th1) to T helper cell type 2 (Th2) phenotypes.¹² Maternal stress can also
93 lead to a decrease in the glutathione/glutathione disulfide (GSSG) ratio, leading to oxidative
94 stress and subsequent risk of asthma in the offspring.^{8,9}

95

96 Several epidemiologic studies investigating indicators of prenatal maternal psychosocial
97 stress on the risk of asthma and allergy in the offspring show that maternal exposure to
98 stress may increase the risk of asthma and allergy in the offspring. Although two recent
99 systematic reviews summarized existing studies,^{13,14} a comprehensive synthesis of the
100 underlying evidence is lacking. In the first review, only wheeze and asthma outcomes were
101 considered.¹³ Although the second review included the full spectrum of allergy outcomes, the
102 searches were confined to a limited number of databases and only a narrative synthesis was
103 performed.¹⁴ Since the publication of these reviews, there have been a number of additional
104 studies published. To provide a clearer and comprehensive picture of the underlying
105 evidence, we undertook a systematic review with meta-analysis of studies that have
106 investigated the association between prenatal maternal exposure to psychosocial stress and
107 the risk of asthma and allergy in the offspring. We included the full spectrum of asthma and
108 allergy outcomes and were also interested in understanding whether the type of indicator of
109 prenatal psychosocial stress and timing (trimester) of exposure were important.

110

111 METHODS

112 We published¹⁵ and registered in PROSPERO (registration number:
113 2016:CRD42016036456) the protocol for the review prior to commencement of the
114 systematic review, which outlined the approaches to undertaking the review.

115

116 **Study types**

117 Experimental studies (i.e. randomized-controlled trials, quasi-randomized controlled trials,
118 controlled-clinical trials, controlled before-and-after studies, interrupted time-series designs)
119 and analytical epidemiological studies (cohort, case-control, and cross-sectional studies)
120 were eligible for inclusion. We excluded reviews, case-studies, case-series, and animal
121 studies.

122

123 **Participants**

124 Pregnant women and their offspring of any age.

125

126 **Exposure**

127 We included all indicators of psychosocial stress, mood states, and acute or chronic
128 stressors or negative life events (NLEs), including: anxiety and depression, issues with
129 existing children, exposure to violence, discrimination or prejudice, financial problems,
130 residential move or housing issues, daily stressors or generalized psychological distress,
131 bereavement, natural disasters, separation, divorce or marital problems, involuntary job loss
132 for mother or partner, and homelessness. We included studies with either objectively-
133 measured or subjectively-reported measures of the stress events.

134

135 **Outcomes**

136 Primary outcomes were: asthma, atopic dermatitis/eczema, atopic sensitization, food allergy,
137 allergic rhinitis, urticaria and anaphylaxis. All primary outcomes, with the exception of atopic
138 sensitization, were defined either by physician assessment or by the self-report. Additionally,
139 asthma diagnosis through use of airway function tests including peak expiratory flow [PEF],
140 forced expiratory volume in 1 second [FEV1], forced vital capacity [FVC], forced expiratory
141 flow rate or alternative age appropriate pulmonary function tests [oscillometry or exhaled
142 nitric oxide analysis] were also accepted methods of assessment. Secondary outcomes
143 included: asthma exacerbations, use of asthma medications, hospitalization for asthma,
144 wheeze as defined by self-report or physician diagnosis, and measures of Health Related
145 Quality of Life (HRQoL).

146

147 **Study identification**

148 We searched the following databases from 1960 to the end of 2016: MEDLINE, EMBASE,
149 Cochrane Library, Web of Science, Scopus, Global Health and Cab International; WHO
150 Global Library; Psych INFO, CINAHL, AHMED, National Health Service (NHS) Evidence
151 Health Information Resources, and Google Scholar. The following databases for international
152 conference proceedings were also searched: Conference Proceedings Citation Index via
153 Web of Knowledge and Zetoc via British Library. Reference lists of eligible articles were hand
154 checked for additional citations. International experts in the field were contacted to ask for
155 any relevant studies not captured through our database searches. We also searched the
156 grey literature through Open Grey and The Grey Literature Report. Finally, the following
157 registers were searched to locate ongoing studies: The Cochrane Central Register of
158 Controlled Trials, International Standardized Randomized Control Trial Number Registry,
159 WHO International Clinical Trials Registry Platform, ClinicalTrials.gov, Australian and New
160 Zealand Clinic Trials Registry, and Current Controlled Trials. There was no language
161 restriction. The search strategies were published in our systematic review protocol.¹⁵

162

163 **Study selection**

164 Records retrieved from the databases were exported to Endnote for study screening, de-
165 duplication, and overall management of the retrieved records. Study titles and abstracts were
166 independently screened by two reviewers (CF and BN); any discrepancies were resolved by
167 discussion. The same process was employed for full-text screening. Multiple reports based
168 on the same study were reported as one study. Where data or information was missing from
169 any study, we contacted authors requesting additional information.

170

171 **Data extraction and management**

172 A standardized form was developed and used to extract relevant data from each study. The
173 data extraction form was piloted and revised prior to use in the review and was published
174 alongside the protocol for this study.¹⁵ Data extraction was completed independently by two
175 reviewers (CF and BN) and discrepancies were resolved by discussion.

176

177 **Quality assessment**

178 All included studies were assessed for quality and risk of bias by two independent reviewers
179 (CF and BN) using the Effective Public Health Practice Project (EPHPP) tool
180 (<http://www.ephpp.ca/tools.html>). In addition to a global rating for each study, the EPHPP tool
181 provides individual ratings for six domains of study quality, including appropriateness of study
182 design for the research question, selection bias, exposure assessment, outcome
183 assessment, data analysis, and generalizability of findings. The two reviewers graded the
184 quality of each study with regards to each of these domains. The quality grading derived from

185 each of the domains informed an overall grading for each study. For each study, the grading
186 for the domains individually and the global study rating were assigned one of the three
187 categories of risk of bias: weak, moderate, or strong. Concerning the domain on confounder
188 adjustment, we considered the number and type of potential confounders adjusted in the
189 studies, with close attention to the confounding factors listed in our own causal diagram (see
190 Figure S1).

191

192 **Analysis**

193 We tabulated the features and key findings of the studies in order to provide a descriptive
194 summary of the literature. We employed both narrative and quantitative synthesis of the
195 underlying evidence. The quantitative synthesis (meta-analysis) was undertaken for studies
196 judged to be reasonably homogenous with regards to similarity in the clinical,
197 methodological, and statistical aspects. We used the random-effects meta-analysis approach
198 for this purpose and included the adjusted risk estimates from each study. We included only
199 cohort studies in the meta-analyses and not cross-sectional and case-control studies.
200 Studies reporting odds ratios as effect measures were first converted to risk ratios before
201 combining in meta-analyses with studies reporting risk ratios. Conversion of odds ratios to
202 risk ratios was undertaken using the formulae by Grant, which given as follows: $RR = OR / (1 -$
203 $p_0 + (p_0 \times OR))$; where p_0 is the baseline risk.¹⁶ We quantified heterogeneity between studies
204 using the I^2 test. We performed the following stratified analyses for each outcome: by type of
205 stress indicator and timing (trimester) of exposure during pregnancy. To enhance
206 comparability between studies that categorized any of the exposures as binary, we collapsed
207 estimates from exposure categories in studies that used multiple exposure categories by
208 using the Mantel-Haenszel approach,¹⁷ thus in all analyses we estimated the risk of maternal
209 stress versus no stress: we estimated the role of maternal exposure to any type of stress
210 indicator (i.e. any of the studied stressors – anxiety, depression, bereavement, work-related
211 stress, or NLEs), specific stress indicators, and trimester of exposure in relation to the
212 outcomes. We used funnel plots to evaluate the potential for publication bias and small study
213 effects and calculated the Begg and Egger's test for this purpose.¹⁸ Meta-analysis was
214 undertaken using Stata 14 statistical software.

215

216 **RESULTS**

217 In total, we identified 9779 articles, of which 7110 were included for screening by title and/or
218 abstract after de-duplication. Of these, 7001 were excluded for not meeting the inclusion
219 criteria and 109 articles were assessed for full text screening. A further 77 articles were
220 excluded, leaving 32 articles (based on 30 studies) that met our inclusion criteria for narrative

221 synthesis; 24 papers (based on 22 studies) were included in at least one meta-analysis
222 (Figure 1).¹⁹⁻⁵⁰

223

224 **Study characteristics**

225 The key characteristics of the studies are presented in Table E1. No experimental studies
226 were found; therefore only analytical epidemiological studies were included, which comprised
227 of 27 cohort studies, two case-control studies, and one cross-sectional study. The type of
228 psychosocial stress indicators investigated in the studies included anxiety,^{21-24,27,30,35,36,48,49}
229 depression,^{22-24,30,35,37,41,42} bereavement,^{29,32,38} work-related stress,^{33,36,47,48} and
230 NLEs,^{19,20,25,26,28,31,34,39,40,43-46,49,50} which were usually comprised of a composite of different
231 indicators of stressors. Most studies assessed maternal stress using self-completed
232 validated questionnaire; in a few studies maternal stress was assessed from population
233 registers, particularly stress resulting from bereavement of a family member. Twelve studies
234 assessed the impact of maternal stress on asthma,^{20,27-32,34,37,38,42,46} eight studies on atopic
235 eczema/dermatitis,^{22-24,28,31,44,47,48} ten studies on wheeze,^{20,24-28,30,35,42,43,46} three studies on
236 allergic rhinitis,^{24,28,31} three on atopic sensitization,^{27,31,42} and six studies on cord blood IgE or
237 cytokines^{19,21,36,39,45,50} (Table S1).

238

239 **Risk of bias within studies**

240 Table S2 provides details of the quality grading for the studies. Of the 30 studies graded for
241 quality, four were strong, 25 were moderate, and one study was weak. Whilst all studies
242 scored moderate or strong on exposure and outcome assessment, only one study scored
243 weak on study design as it was based on a cross-sectional data. Six studies were graded
244 weak for selection bias, whereas two studies were graded weak for confounding adjustment;
245 no study was graded strong for selection bias and confounding adjustment.

246

247 **Prenatal stress and offspring asthma**

248 Prenatal maternal exposure to any type of stress indicator was associated with an increased
249 risk of asthma onset (hazard ratio (HR) 1.13, 95%CI 0.98-1.32; $I^2=91.5\%$) and current or ever
250 asthma (RR 1.13, 95%CI 1.03-1.24; $I^2=83.5\%$) in the offspring, although result for asthma
251 onset was not statistically significant (Figure 2, Panel A). Concerning the type of stress
252 indicators, only anxiety was significantly associated with an increased risk of asthma (RR
253 1.28, 95%CI 1.16-1.41; $I^2=0\%$) (Figure 2, Panel B). Concerning the timing of prenatal
254 maternal stress, only exposure during the third trimester was significantly associated with an
255 increased risk of asthma (OR 1.45, 95%CI 1.08-1.94; $I^2=78\%$) for the studies that measured
256 current and ever asthma (Figure 3, Panel B). Bereavement of the death of a child (HR 1.28,

257 95%CI 1.10-1.48; $I^2=0\%$) or of a spouse (HR 1.40, 95%CI 1.03-1.90; $I^2=1.3\%$), but not of a
258 parent or sibling, increased the risk of asthma onset in the offspring (Figure 4).

259

260 **Prenatal stress and offspring atopic eczema/dermatitis**

261 Prenatal maternal exposure to anxiety (RR1.29, 95%CI 0.95-1.76; $I^2=29.2\%$), depression
262 (RR 1.45, 95%CI 1.12-1.89; $I^2=0\%$), NLEs (OR 1.18, 95%CI 0.92-1.51; $I^2=0\%$), and work
263 stress (OR 1.32, 95%CI 1.16-1.50) was associated with increased risk of atopic
264 eczema/dermatitis in the offspring, but only depression and work stress were statistically
265 significant (Figure 5, Panel A). Maternal exposure to stress during the third and any
266 trimester, but not during the second trimester, increased the risk of offspring atopic
267 eczema/dermatitis (Figure 5, Panel B).

268

269 **Prenatal stress and offspring wheeze**

270 Prenatal maternal exposure to anxiety (RR 1.19, 95%CI 1.01-1.39; $I^2=52\%$), depression (OR
271 1.74, 95%CI 1.42-2.13; $I^2=0\%$), and NLEs (RR 1.23, 95%CI 1.00-1.52; $I^2=88.3\%$) (Figure 6,
272 Panel A) was associated with an increased risk of wheeze in the offspring. The increased
273 risk was greater for maternal exposure to stress during the second and third than any
274 trimester (Figure 6, Panel B). When we investigated the impact on different wheezing
275 phenotypes, maternal stress increased the risk of early-onset, late-onset, and persistent
276 wheeze, although the impact on late-onset wheeze was not statistically significant (Figure 7).

277

278 **Prenatal stress and offspring allergic rhinitis**

279 Prenatal maternal exposure to any type of stress indicator was associated with an increased
280 risk of subsequent allergic rhinitis in the offspring (OR 1.36, 95%CI 1.08-1.71; $I^2=43.7\%$)
281 (Figure S2). These results were similar when the study by Hartwig et al was analyzed
282 separately for allergic rhinitis at six and 14 years.³¹

283

284 **Prenatal stress and offspring atopic sensitization**

285 Prenatal maternal exposure to any stress indicator was associated with a decreased risk of
286 atopic sensitization in the offspring (OR 0.92, 95%CI 0.86-0.98; $I^2=0\%$) (Figure 8). These
287 results were similar when the study by Hartwig et al was analyzed separately for atopic
288 sensitization at six and 14 years.³¹ The measurement and definition of atopic sensitization
289 differed between the studies: whilst Cookson²⁷ defined it as ≥ 2 mm weal skin prick test to
290 aeroallergens, Hartwig³¹ and Reyes⁴² were based on IgE measurements of a combination of
291 both food and inhalant allergens.

292

293 **Prenatal stress and cord blood IgE and cytokines**

294 Studies reporting the impact of maternal prenatal stress on cord blood IgE and cytokines
295 were heterogeneous, particularly with regards to the type of outcomes investigated, hence a
296 meta-analysis was not undertaken to pool the results of these studies together. However,
297 across studies, maternal exposure to stress during pregnancy was associated with raised
298 cord blood specific and total IgE.^{36,39,45,50} One study reported an alteration of cord blood
299 cytokine profiles (IL-12, IL-1 β , IL-4, IL-5, IL6, IL-8, and TNF- α) in offspring of mothers
300 exposed to stress during pregnancy.¹⁹

301

302 **Assessment of publication bias**

303 Figure S3 shows the funnel plot evaluating possible publication bias and small study effect:
304 by interpretation, a symmetric funnel plot indicates less likelihood of publication bias
305 influencing the estimates of effect. Indeed, the funnel plot in Figure S3 is modestly
306 symmetric. The associated p-values for Egger's test (where Egger's test of with $P < 0.05$
307 indicating presence of publication bias) were as follows: atopic eczema/dermatitis studies
308 $p = 0.949$; atopic sensitization studies $p = 0.855$; wheeze studies $p < 0.001$; asthma studies
309 $p = 0.828$; and allergic rhinitis studies $p = 0.493$.

310

311 **DISCUSSION**

312 This study provides the most comprehensive and robust synthesis of the evidence to date on
313 the association between prenatal maternal exposure to psychosocial stress and the risk of
314 allergy and asthma in the offspring. The majority of included studies were at moderate risk of
315 bias. Overall, prenatal maternal exposure to any psychosocial stress was associated, albeit
316 modestly, with an increased risk of asthma, atopic eczema/dermatitis, wheeze and its
317 phenotypes, and allergic rhinitis. A decreased risk was seen for atopic sensitization. Although
318 these results were similar for specific indicators of stress, exposure to anxiety and
319 depression had the strongest effects. The third trimester appeared to be more vulnerable
320 period of exposure compared to first and second trimesters. Specific to asthma, maternal
321 bereavement of a child or a spouse, but not of a parent or sibling, increased the risk of
322 asthma in the offspring.

323

324 We undertook an exhaustive search of the literature, covering the major medical and public
325 health databases, which was supplemented through search of grey literature and through
326 contacting experts in the field. The search strategies were implemented and published in
327 order to enhance reproducibility. It is therefore unlikely that we missed any relevant literature,
328 this being confirmed by somewhat symmetric funnel plot on publication bias and small study
329 effect. Two reviewers independently performed each stage of the review, including literature
330 screening, data extraction, and quality appraisal of included studies. We developed,

331 published and registered a detailed protocol¹⁵ prior to undertaking the review, which
332 enhanced the transparency of the review process.

333

334 At the same time of publishing our review protocol, two related systematic reviews were
335 published.^{13,14} By the time we were planning the current review, no protocol was published
336 for those reviews, neither were they registered in PROSPERO; hence our preliminary search
337 did not identify them. Nevertheless, in the first review only asthma and wheeze were
338 outcomes, which limits its scope.¹³ The second review considered all possible asthma and
339 allergic outcomes, but only provided narrative synthesis of the existing literature.¹⁴ We aimed
340 for a comprehensive and exhaustive approach by including the full spectrum of allergy and
341 asthma outcomes and considering whether the type of stressor and trimester of exposure
342 were important. We identified 30 unique studies as against 16 studies in the second review¹⁴
343 and 18 studies as against 10 studies in the first review with regards to asthma/wheeze
344 outcomes.¹³ Whilst the second study¹⁴ performed only narrative synthesis, with careful
345 consideration, we judged several of these studies to be reasonably homogenous to be
346 combined in meta-analyses with respect to exposure and outcome definitions, study design,
347 and statistical analyses. Regardless of the differences in methodological rigor and
348 comprehensiveness, the conclusions from the two previous systematic reviews align with our
349 findings, indicating that prenatal maternal exposure to psychosocial stress was associated
350 with increased risk of asthma and allergic disease in the offspring.

351

352 The majority of studies included in our review were graded as at moderate risk of bias, with
353 only four being graded as strong studies, an indication of the potential for biases across
354 studies, particularly within the domains of selection bias and confounding adjustment. In
355 particular, most studies assessed maternal prenatal stress using self-report questionnaires,
356 usually for recall of previous exposure to stress across several months. No study used both
357 self-report and objective measures at regular intervals, which would provide a more robust
358 and informative understanding of unique and combined contribution of environmental and
359 mechanistic factors involved in the developmental pathway of atopic conditions. Several
360 studies also assessed allergy outcomes using maternal subjective questionnaires and the
361 age of onset of the outcomes was not consistent across all studies. Objective assessment of
362 both maternal prenatal stress and offspring outcomes and within the same time-point will
363 improve the underlying evidence and provide a stronger basis to evaluate whether these
364 findings are causal.¹⁴ The test for heterogeneity was significant for a number of associations,
365 an indication of the variations in methods and measures between studies; however as we did
366 not have sufficient number of studies for each of these associations, we were unable to
367 further investigate the possible reasons for these significant heterogeneity tests.

368 Furthermore, as it was not feasible to perform a formal test between subgroups, it is possible
369 that the associations found within subgroups may be a result of chance.

370

371 Although some studies have examined the role of maternal stress during the postpartum
372 period and the child's exposure during infancy on subsequent risk of allergy and asthma in
373 the child,^{26,51} *a priori*,¹⁵ the underpinning objective of our review was to assess the impact of
374 prenatal stress on offspring's asthma and allergy. This objective aligns within the framework
375 of the fetal programming hypothesis. Within this framework, we assumed that the pathway of
376 prenatal stress influencing offspring asthma and allergy risk may be independent of the
377 effects of postnatal and early life stress, hence we excluded all studies not investigating
378 maternal prenatal stress. However, we cannot rule out the possibility that these findings may
379 also be partly explained by postnatal or early life stress exposures.⁵² Furthermore, whilst the
380 timing of exposure was based on the trimester of assessment of prenatal stress, this single
381 time-point assessment may fail to reflect specific trimester effect, as stressful events may be
382 acute or may chronologically be present throughout pregnancy or may even be an extension
383 of stressful events prior to pregnancy.⁵²

384

385 Regardless of the type of stress indicator, timing of exposure, and the type of allergy and
386 asthma outcomes, by bringing together all the available evidence, the current evidence
387 synthesis shows that prenatal maternal psychosocial stress is associated with an increased,
388 albeit modest, risk of asthma and allergy in the offspring. The findings were particularly more
389 evident for depression and anxiety than for other indicators of prenatal stress. This could
390 reflect that anxiety and depression scores were based on self-assessment and
391 questionnaires. However, it should also be noted that the largest study had a strongly
392 positive result.³² This study also had a strong design due to using a 'natural experiment'
393 design and an objective measure of hospitalized asthma, thereby avoiding the risks of
394 reporting bias and reverse causation.³² In addition, although depression and anxiety reflect
395 mood states, they are robust correlates of psychosocial stress and strongly predispose to
396 stress-related conditions, such as smoking, poor diet, poor sleeping habits, and poor quality
397 of life which may also lead to asthma and allergy in offspring.⁵³

398

399 Whilst our observations may represent causal relationships, they may also be a
400 consequence of over-reporting of offspring disease status by distressed mothers^{22,54} or due to
401 residual confounding in the original studies. The number and type of confounders adjusted
402 varied across included studies. In particular, the omission of key confounders in several
403 studies, including maternal allergic history, pregnancy complications, acid reflux conditions,
404 medication use during pregnancy (e.g. antibiotics and acetaminophen), and pregnancy

405 weight gain indicates sub-optimal confounding adjustment and therefore may impact on the
406 observed risk. One way to test for residual confounding in fetal programming studies is to
407 use family design studies such as sibling studies^{54,55} or to use a paternal negative control. A
408 positive association for paternal exposure to stress during the mother's pregnancy and
409 subsequent offspring asthma or allergy may indicate residual confounding is affecting the
410 prenatal maternal stress and offspring asthma association. A recent study using fathers as a
411 negative control in this way, found that after adjusting for measured confounders there was
412 no evidence for residual confounding.⁵⁶

413
414 One hypothesized pathway through which stress may influence risk of asthma and allergy is
415 that high levels of cortisol resulting from external stress may potentiate cell differentiation
416 from Th1 to Th2 phenotype.¹² Prenatal stress-generated cortisol has been linked to
417 increased airway responsiveness in the offspring in animal models.¹² This indicates that
418 prenatal maternal stress may increase risk of atopic sensitization in the offspring and
419 subsequent allergic disease and asthma. However, our findings did not support this line of
420 reasoning, but showed a decreased risk of atopic sensitization and increased risk of clinical
421 allergic outcomes and asthma. The reason for these differential findings between atopic
422 sensitization and other clinical outcomes is unclear. Other suggested mechanism indicate
423 that prenatal maternal stress may cause epigenetic effects with deoxyribonucleic acid (DNA)
424 methylation and altered gene expression in the placenta,⁵⁷ but this proposal and its
425 implication for the development of allergy and asthma in the offspring has yet to be
426 determined.⁵¹ There is some evidence that prenatal stress exposure can influence the
427 composition of the offspring's intestinal microbiota and may also result in increased
428 susceptibility to asthma.^{58,59}

429

430 **CONCLUSION**

431 Prenatal maternal psychosocial stress – particularly anxiety and depression - was associated
432 with a modest increased risk of asthma and allergy in the offspring. Whilst exposure during
433 the third trimester had the greatest impact compared to the first and second trimesters, this
434 may represent cumulative stress exposure throughout pregnancy rather than a trimester-
435 specific effect. These findings may represent a causal association; they may also result from
436 inherent biases in the included studies, particularly residual confounding. Further studies with
437 objective measures of prenatal stress and optimal adjustment of confounding are required,
438 as well as studies elucidating the likely mechanisms for these associations.

439

440 **CONFLICTS OF INTEREST**

441 The authors declare no competing interest related to this work.

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AUTHORS' CONTRIBUTIONS

BN conceived the idea for this work. It was drafted by CF and BN and was then revised after several rounds of critical comments from AS, ADG, BKB, and CA.

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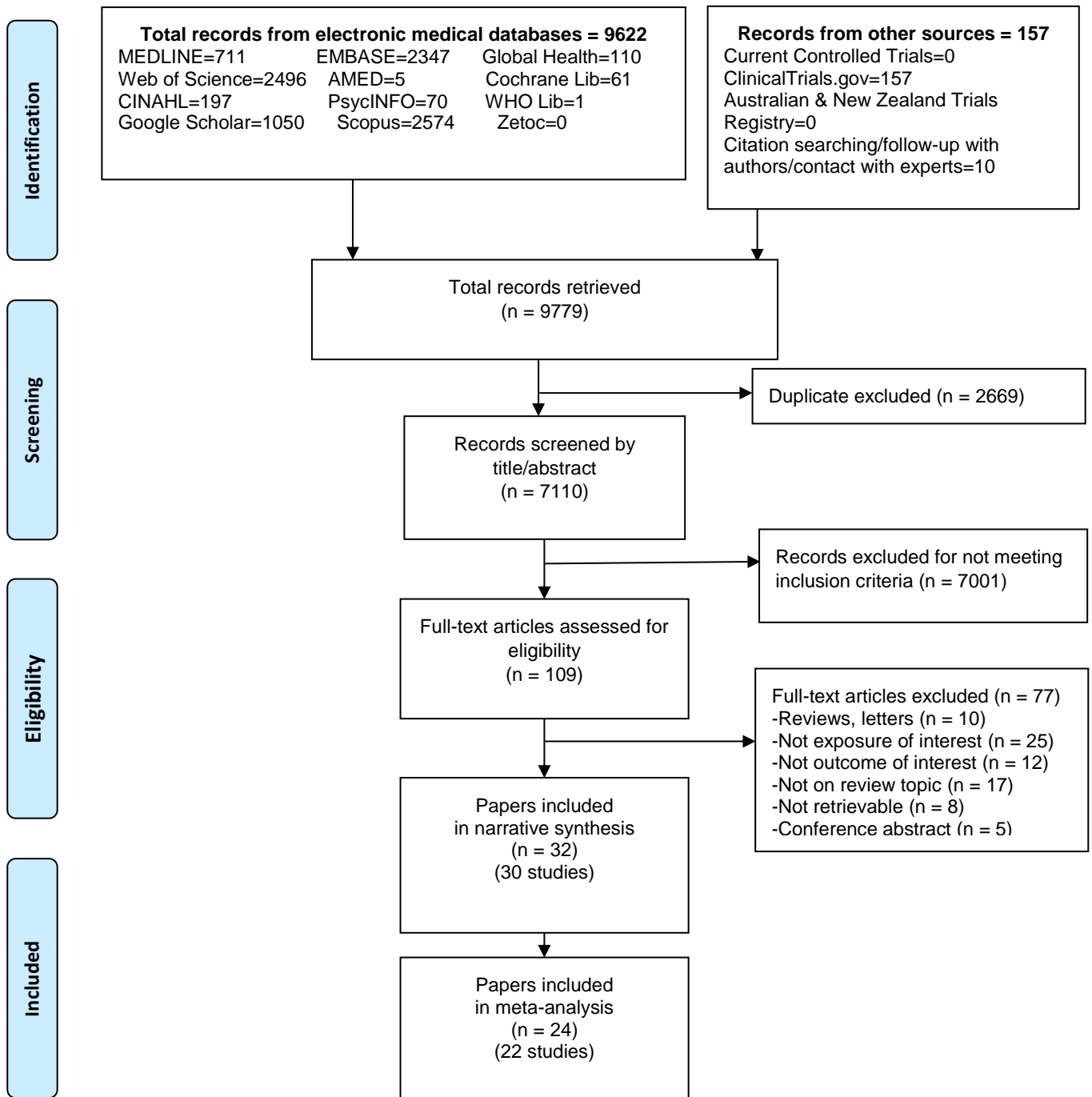
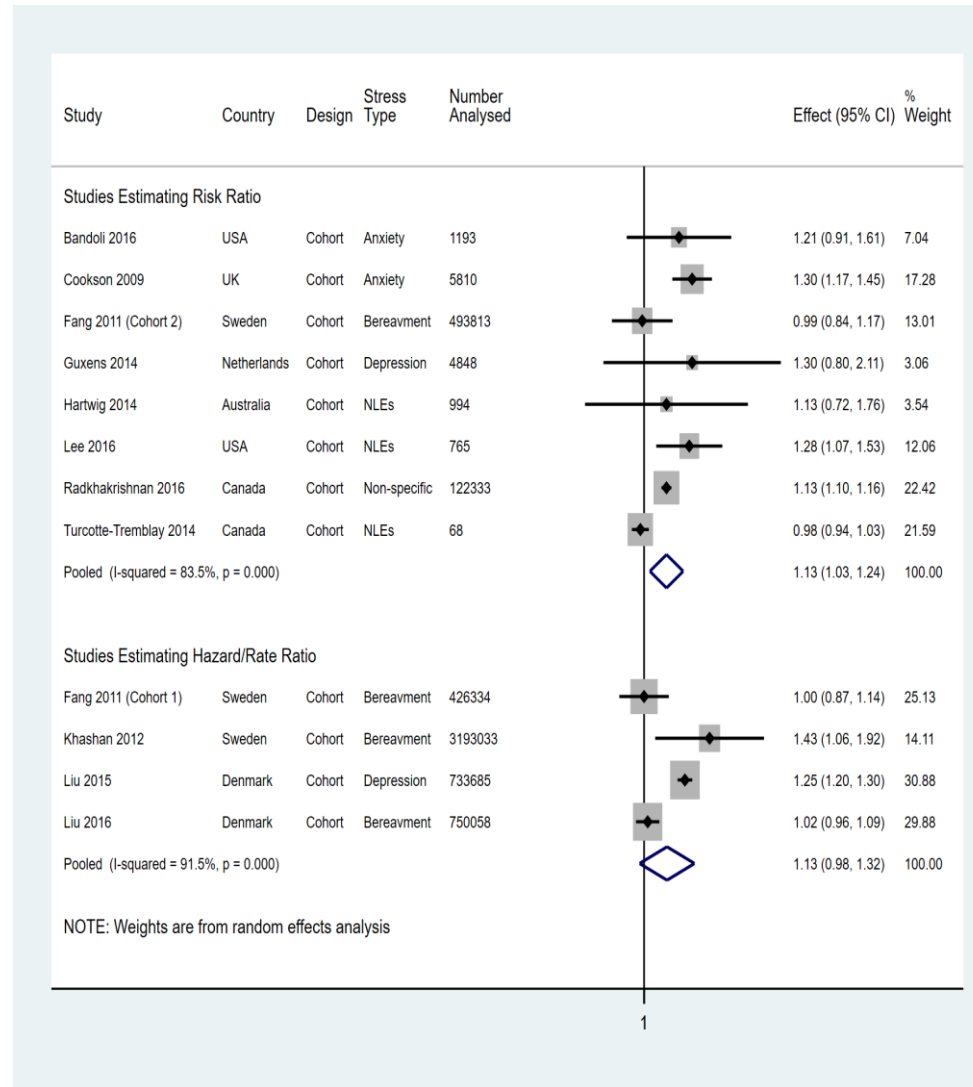


Figure 1. PRISMA flow diagram of studies on the association between maternal prenatal stress and risk of allergy and asthma in the offspring

PANEL A



PANEL B

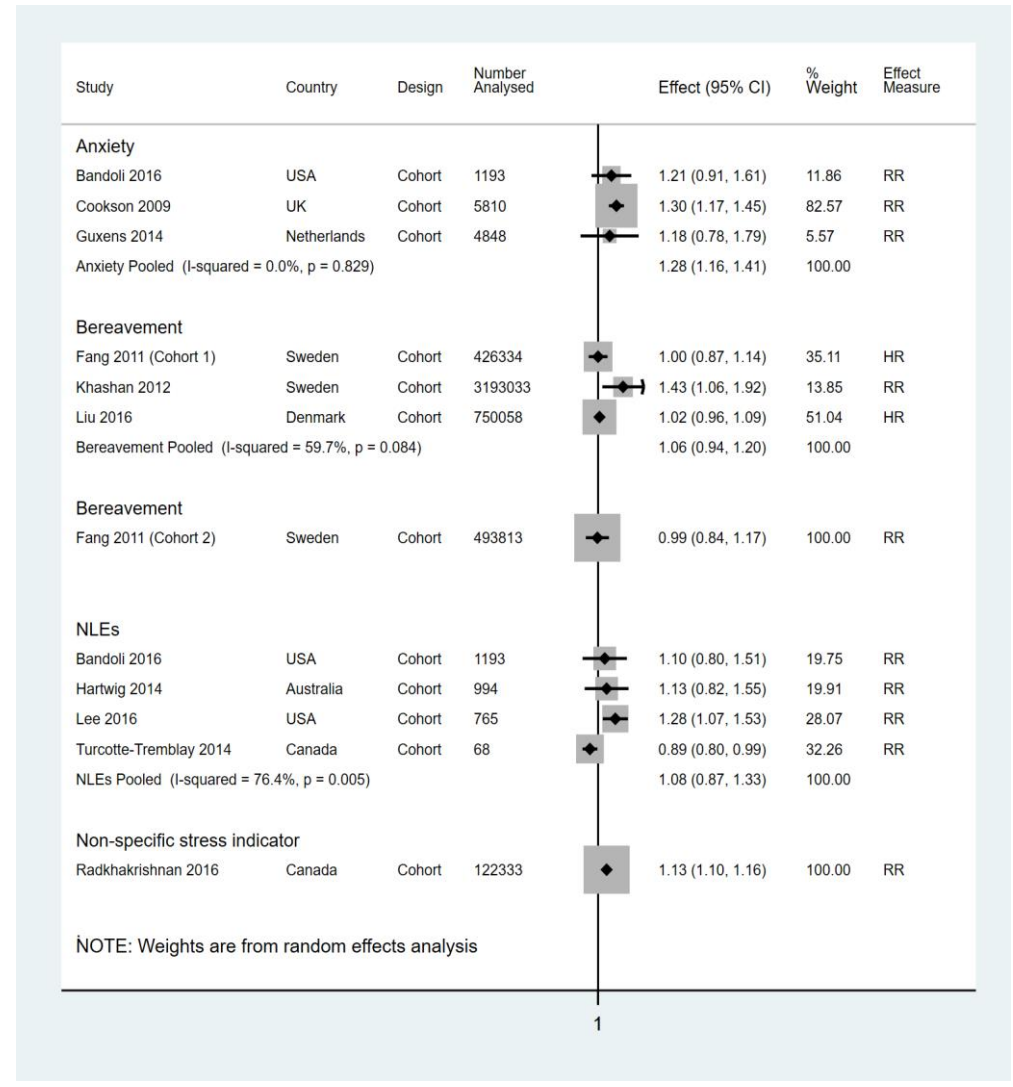
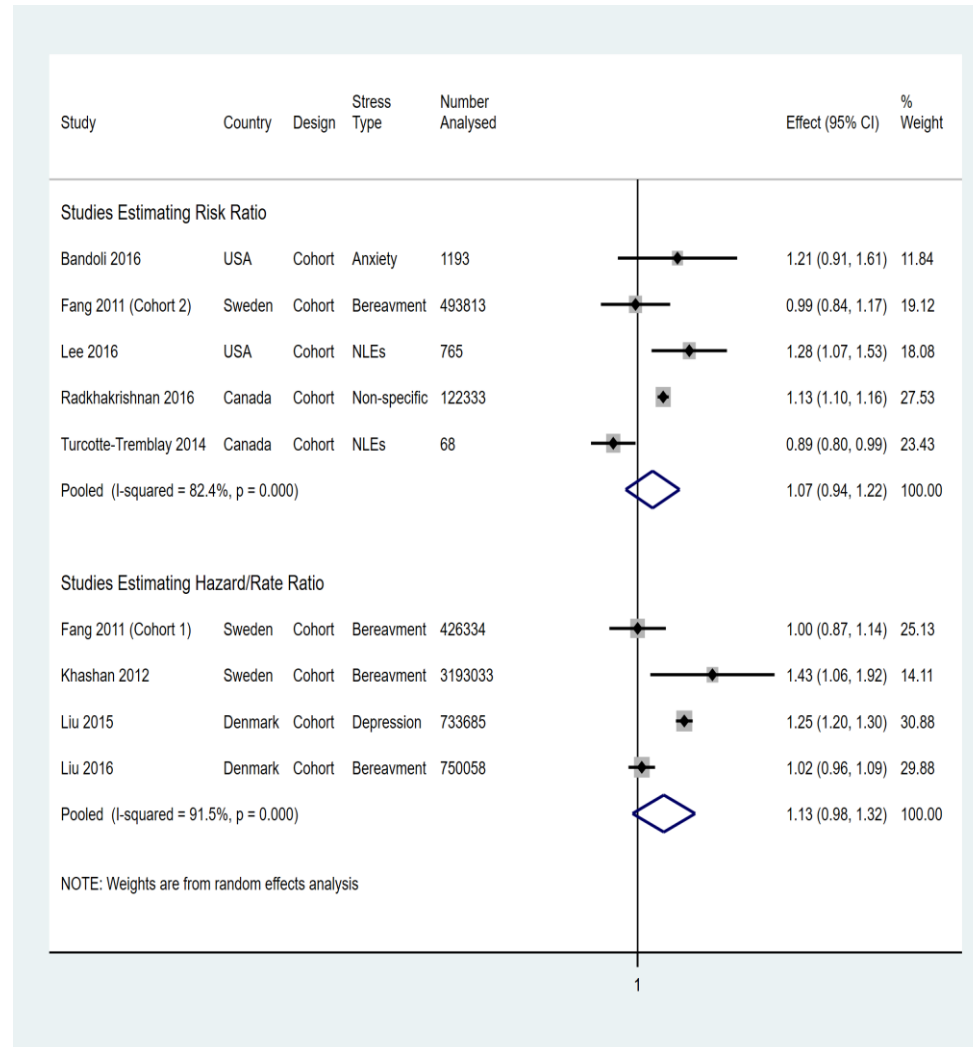


Figure 2. Association between prenatal maternal stress (**Panel A:** any type of stress and **Panel B** by type of stress) and risk of asthma in the offspring. NLEs = Negative life events. The results included Hartwig 2014 and Liu 2016 populations of 14 years and 4-15 years, respectively, as these were not substantially different from the population of 6 years and 0-3 years, respectively, also presented in the studies.

PANEL A



PANEL B

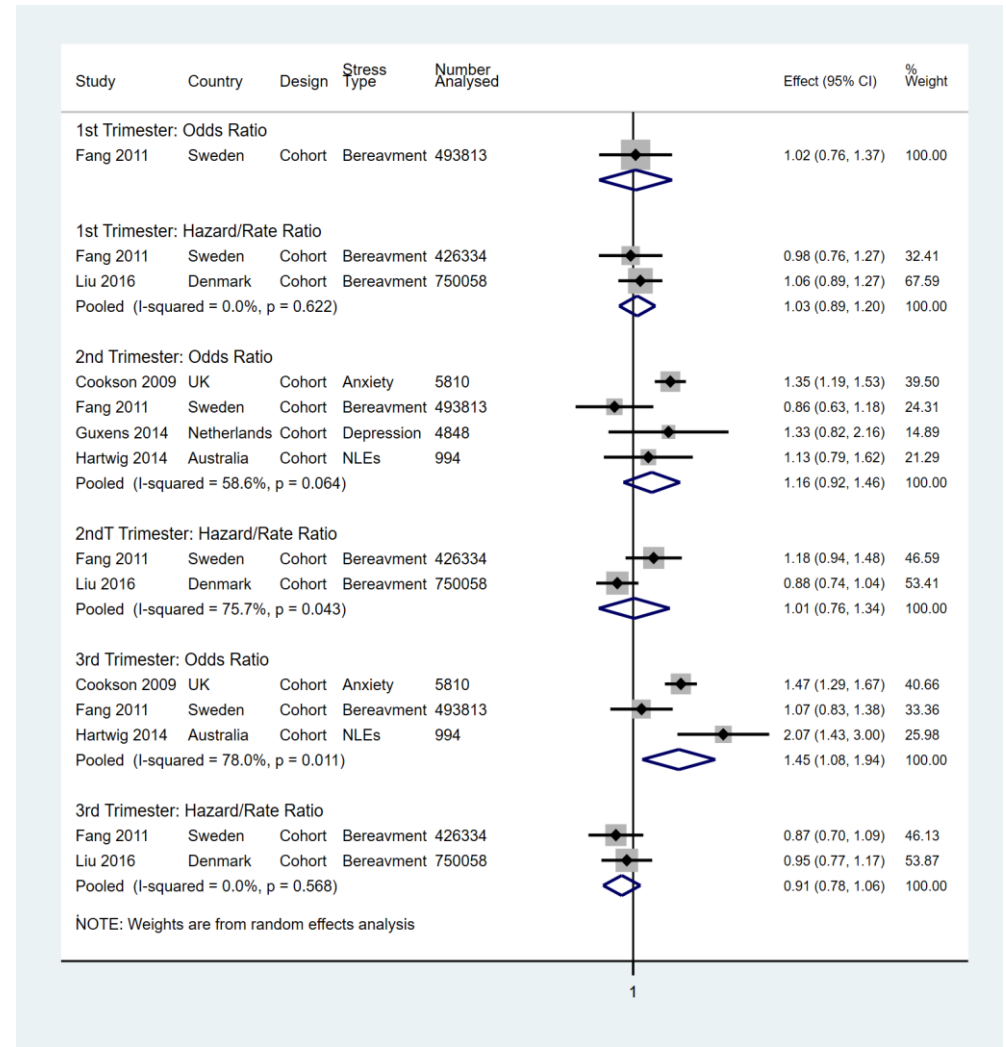


Figure 3. Association between prenatal maternal stress and risk of asthma in the offspring, by timing of exposure during pregnancy: **Panel A:** during any trimester; **Panel B:** at different trimesters. The results included Hartwig 2014 and Liu 2016 population of 14 years and 4-15 years as these were not substantially different from the population of 6 years and 0-3 years also presented in the studies.

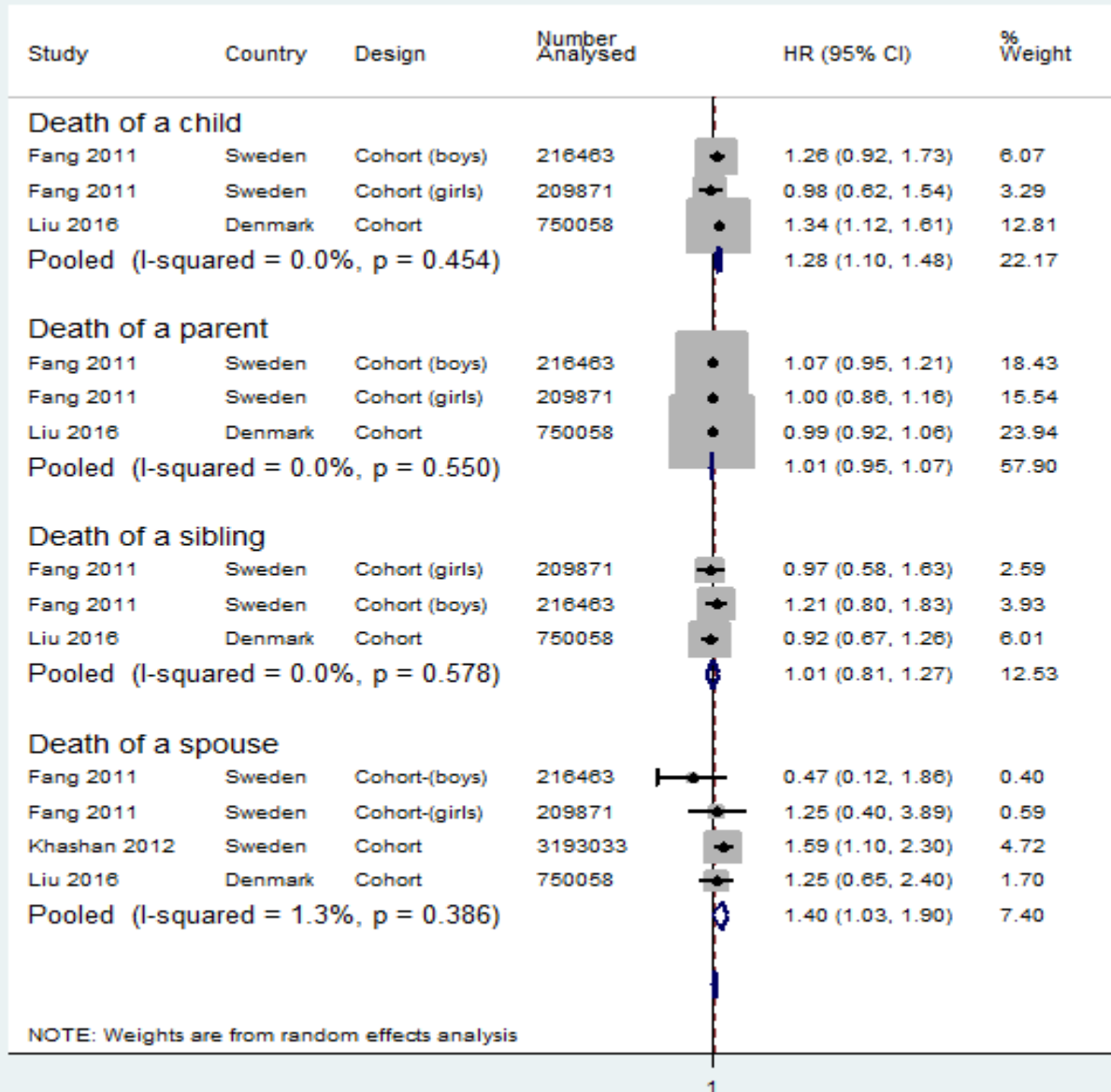
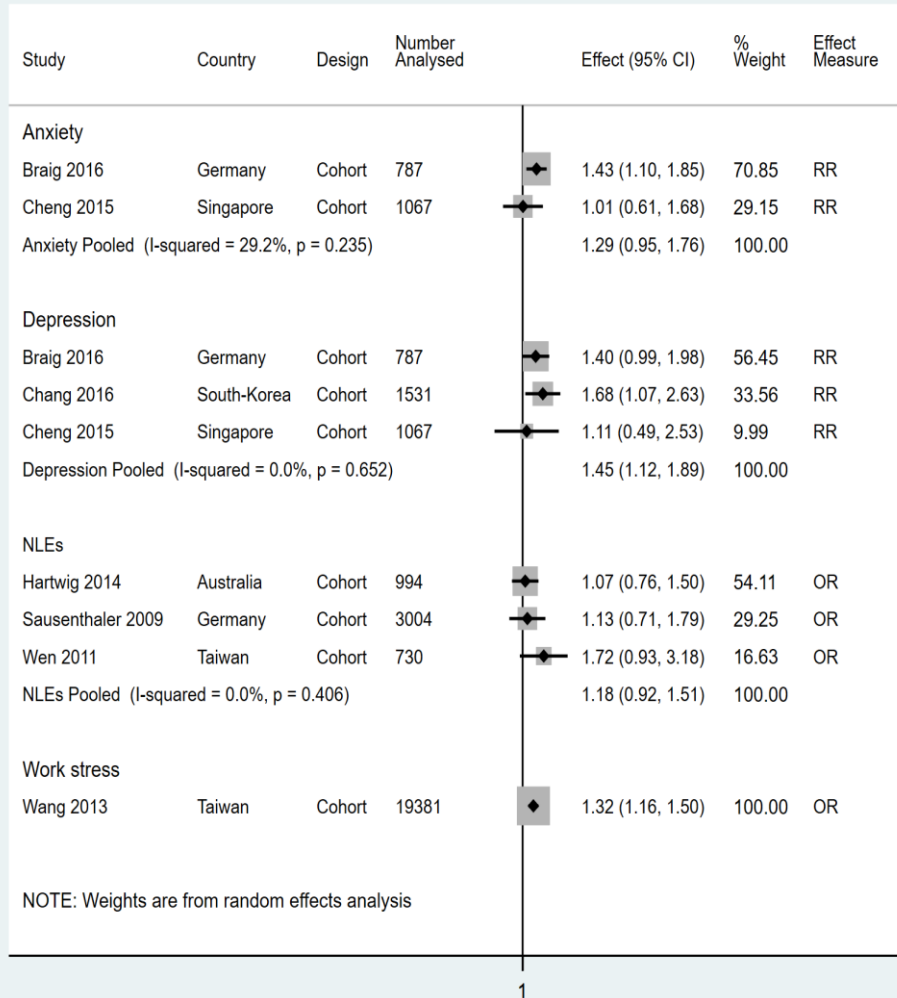


Figure 4. Association between maternal stress resulting from bereavement of a relative and risk of asthma in the offspring, by type relative. The results included Hartwig 2014 and Liu 2016 population of 14 years and 4-15 years as these were not substantially different from the population of 6 years and 0-3 years also presented in the studies.

PANEL A



PANEL B

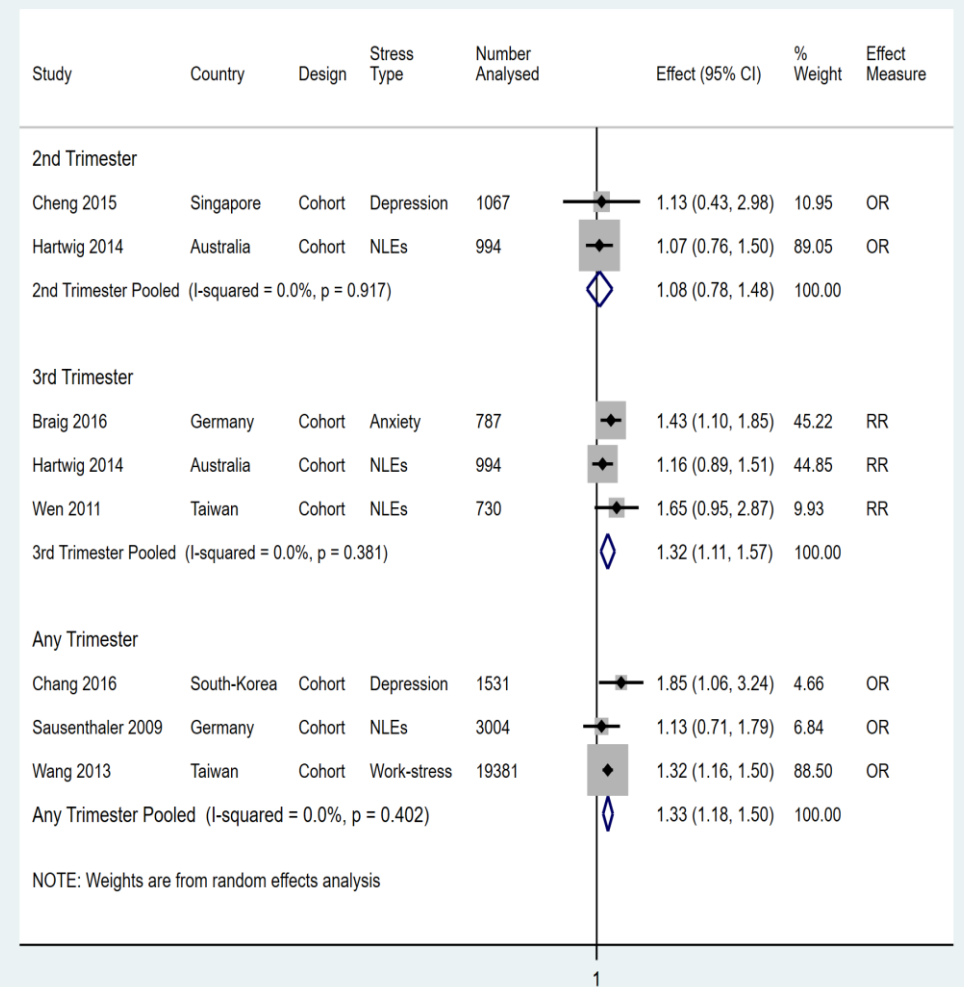
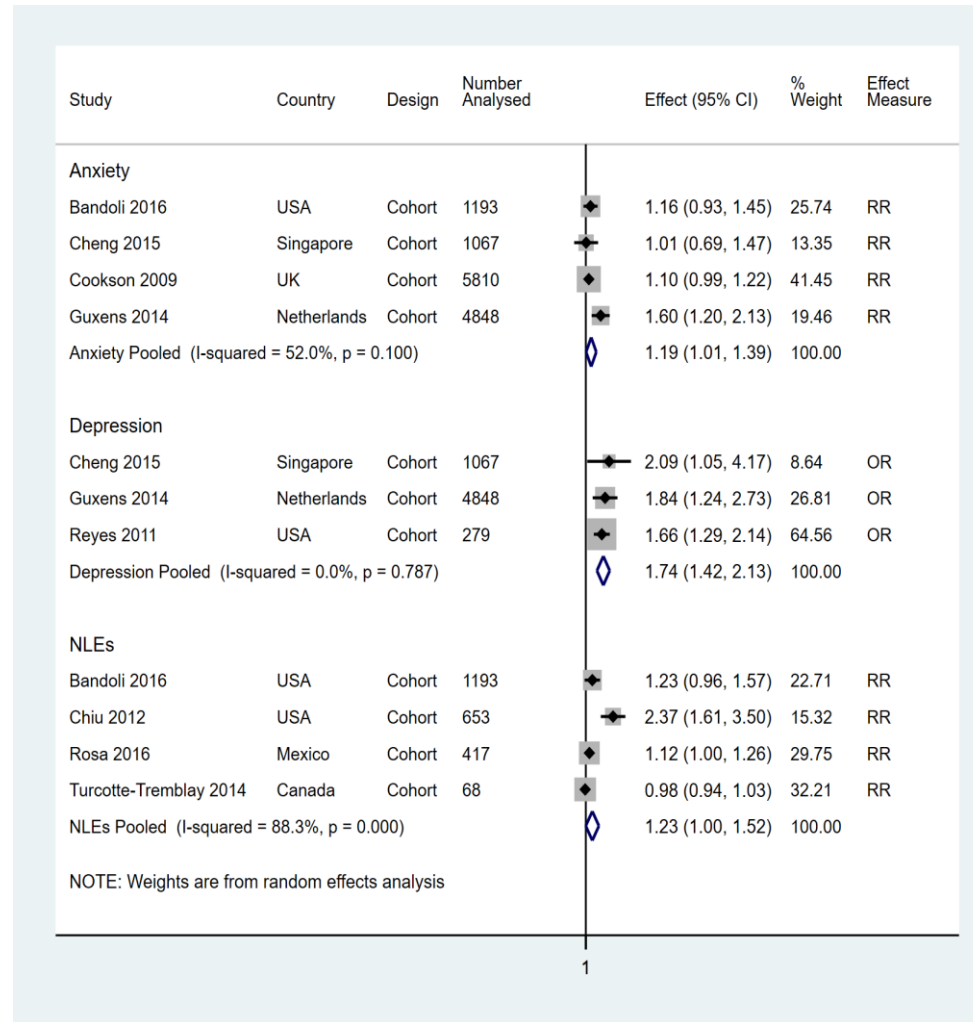


Figure 5. Association between maternal prenatal stress and risk of atopic eczema/dermatitis in the offspring, by type of stress (**Panel A**) and timing of exposure during pregnancy (**Panel B**). NLEs = negative live events. No major differences when Hartwig 2014's 6-year-olds and 14-year-olds were analysed separately, hence we presented the results for 6-year-olds

PANEL A



PANEL B

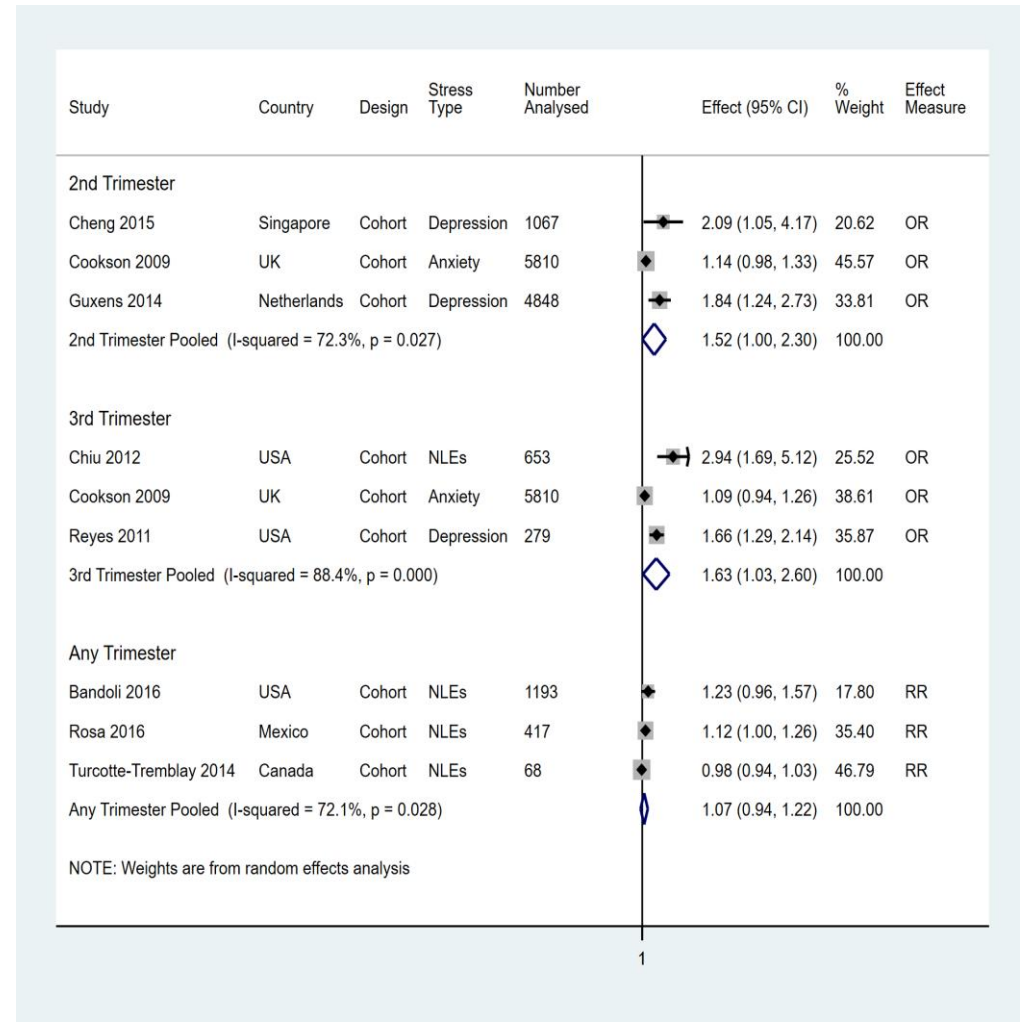


Figure 6. Association between maternal prenatal stress and risk of wheeze in the offspring, by type of stress (**Panel A**) and timing of exposure during pregnancy (**Panel B**). NLEs = negative live events. No major differences when Cookson 2009's wheeze at 6-18 months and 69-81 months were analysed separately, hence we present the results for the 6-18 month age group

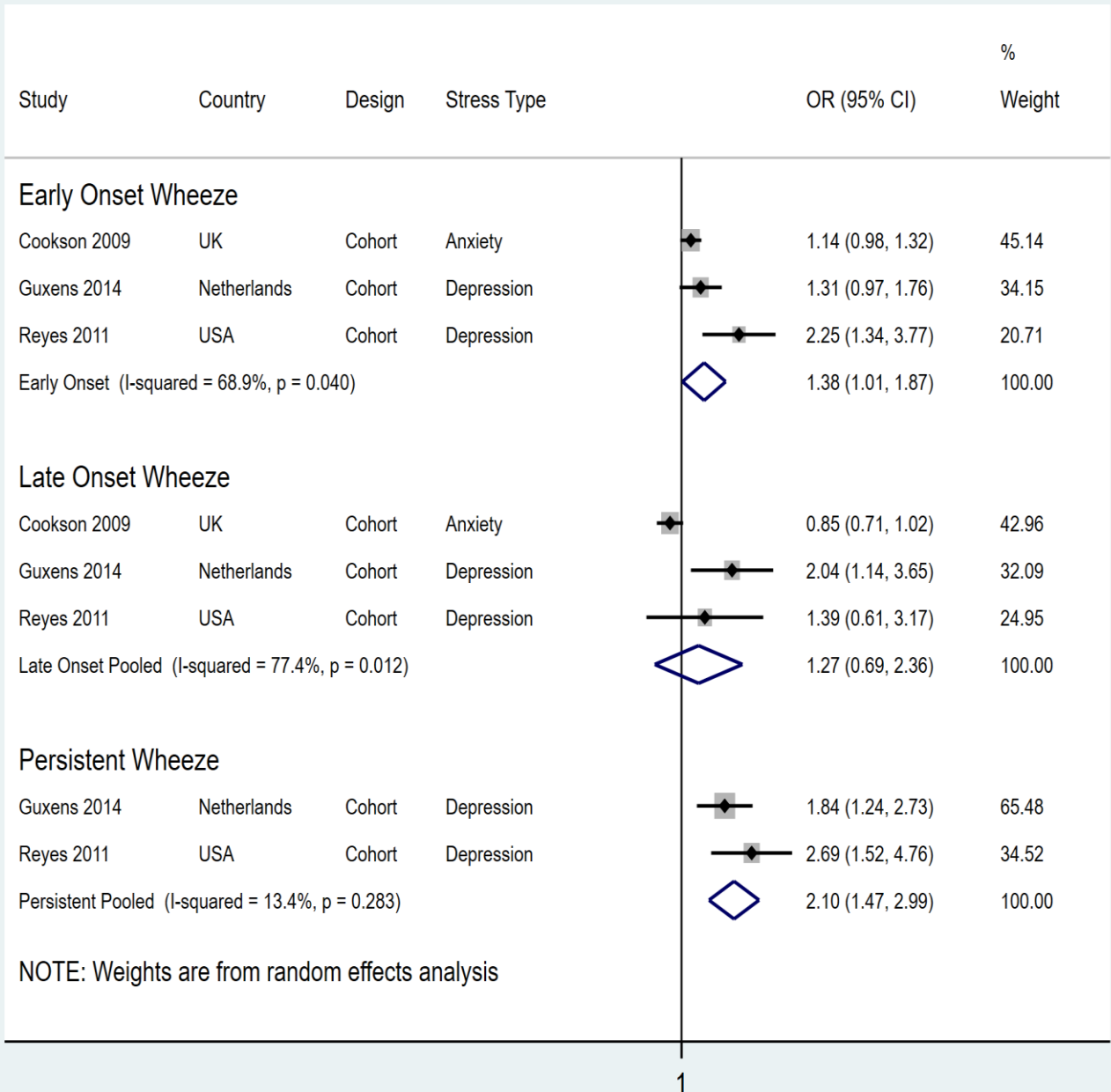


Figure 7. Association between maternal prenatal stress and risk of wheeze phenotypes in the offspring

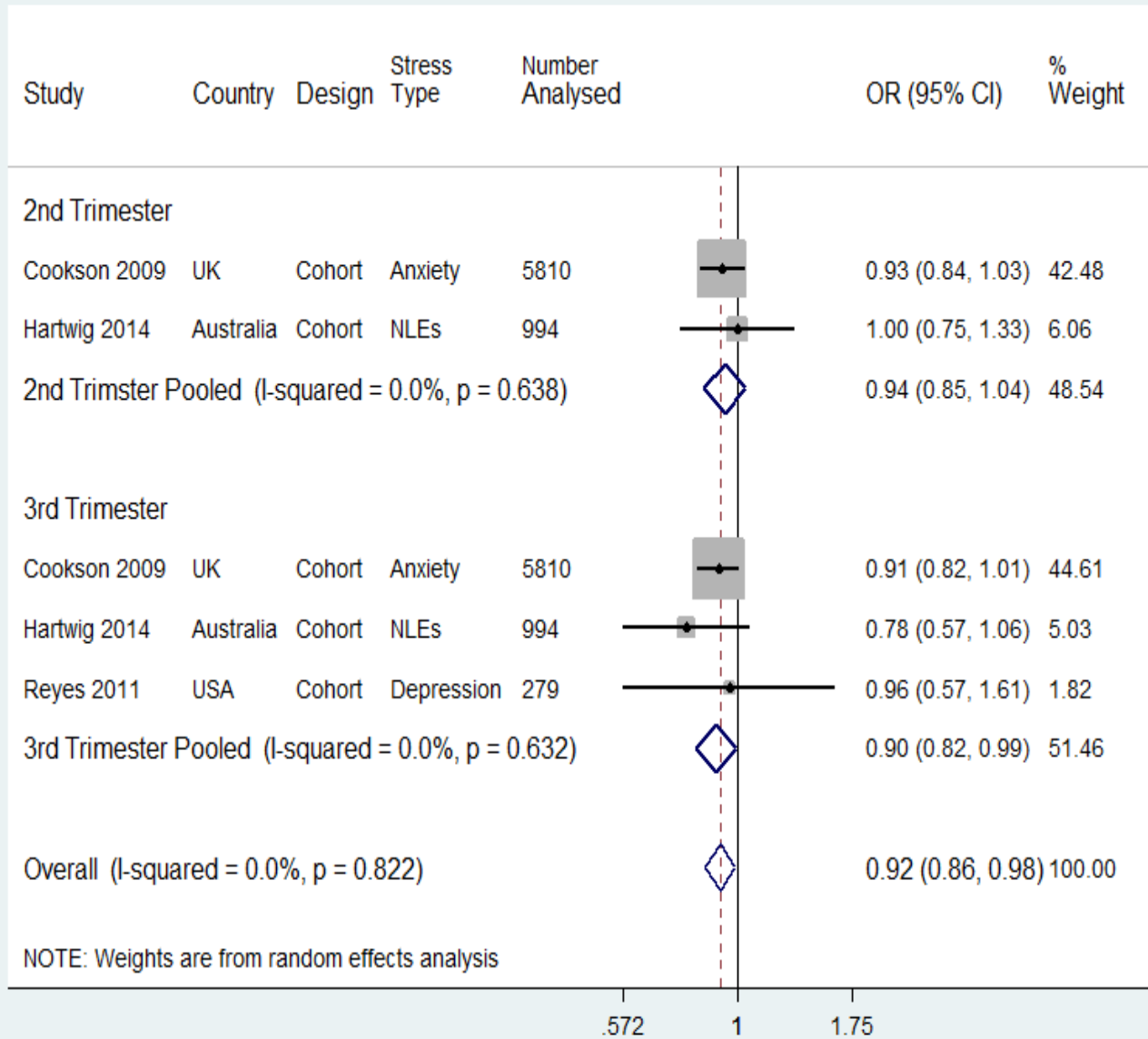


Figure 8. Association between maternal prenatal stress and atopic sensitisation in the offspring, by timing of exposure during pregnancy: No major differences when Hartwig 2014's 6-year-olds and 14-year-olds were analysed separately, hence we presented the results for 6-year-olds

Table S1: Main characteristics, key results, and overall risk of bias of studies on the association between maternal psychosocial stress during pregnancy and risk of allergy and asthma in the offspring

Reference, country; and study design	Study population N (maternal-child; source of study population)		Age of children/ follow-up years	Type of prenatal stress and assessment	Outcomes and assessment	Key results	Overall risk of bias assessment
	Number recruited	Number analysed					
Andersson et al 2016; USA; Prospective cohort study	50	43	At birth	Maternal reported anxiety and negative life events (NLEs) during the 2 nd trimester using Self-reported using Perceived Stress Scale (PSS-14); State-Trait anxiety inventory (STAI); Life Experience Interview	Cord blood cytokines: IL-12, IL-1 β , IL-4, IL-5, IL6, IL-8, and TNF- α	Prenatal maternal stress increased the risk of alteration of cord blood cytokine profiles in the offspring	Moderate
Bandoli et al 2016; USA; Prospective cohort study	6,347	1,193	2.3-5.8 years	Maternal reported anxiety, chronic stress, and NLEs (loss of car/job/home, serious arguments with partner, close acquaintance with health, drug or legal problems, death of a relative, threatened with physical harm, discrimination due to race/ethnicity) at any time during pregnancy	Maternal reported wheeze and asthma using the ISAAC questionnaire	Maternal anxiety and current asthma: adjusted Moderate vs none: RR 1.22 (0.73-2.05) Somewhat vs none: RR 1.07 (0.67-1.71) Very much vs none: RR 1.39 (0.83-2.35) Maternal chronic stress and current asthma: adjusted Moderate vs low: RR 0.75 (0.49-1.14) High vs low: RR 1.11 (0.67-1.86) Maternal NLEs and current asthma: adjusted 1 NLE vs none: RR 1.21 (0.80-1.83) 2+ NLEs vs none: RR 0.96 (0.58-1.57) Results also given in the paper for ever and current wheeze	Moderate
Bidaki et al 2011; Iran; Prospective cohort study	320	290	At birth	Maternal reported anxiety. Assessed using the Holme-Rahe stress questionnaire at the 3 rd trimester of pregnancy	Cord blood IgE	No estimate of effect computed only descriptive data presented in the paper	Moderate
Braig et al 2016; Germany; Prospective cohort study	2,610	787	Up to 2 years	Maternal questionnaire-reported chronic stress, anxiety, and depression symptoms during the 3 rd trimester. Also maternal hair cortisol	Maternal questionnaire-reported atopic dermatitis (AD). Dermatologist confirmation using	Maternal chronic stress and AD symptoms: adjusted Quartile 2 vs quartile 1: RR 1.3 (0.8-1.9) Quartile 3 vs quartile 1: RR 1.4 (0.9-2.1) Quartile 4 vs quartile 1: RR 1.5 (1.0-2.3) Maternal depression and AD symptoms : adjusted High vs low: RR 1.4 (1.0-2.0)	Moderate

					SCORAD	Maternal anxiety and AD symptoms: adjusted Quartile 2 vs quartile 1: RR 1.4 (0.9-2.1) Quartile 3 vs quartile 1: RR 1.4 (0.9-2.2) Quartile 4 vs quartile 1: RR 1.5 (0.9-2.4) Maternal hair cortisol and AD symptoms: adjusted Quartile 2 vs quartile 1: RR 1.0 (0.7-1.5) Quartile 3 vs quartile 1: RR 0.9 (0.6-1.4) Quartile 4 vs quartile 1: RR 0.8 (0.5-1.2) Results also given in the paper for parent-reported, paediatrician-reported and strictly defined AD	
Chang et al 2016; South Korea; Prospective cohort study	Cohort 1: 2,021 Cohort 2: 2,150	Cohort 1: 973 Cohort 2: 1,531	Up to 5 years	Maternal questionnaire-reported stress: Cohort 1: depression and anxiety at 3 rd trimester Cohort 2: depression any time during pregnancy	AD symptoms Cohort 1: diagnosis by pediatric allergist Cohort 2: maternal report using ISAAC questionnaire	Cohort 1: maternal depression and AD: adjusted HR 1.31 (1.02-1.69) Cohort 1: maternal anxiety and AD: adjusted HR 1.41 (1.06-1.89) Cohort 2: maternal depression and AD: adjusted OR 1.85 (1.06-3.25)	Moderate
Cheng et al 2015; Singapore; Prospective cohort study	1,152	1,067	Up to 1 year	Maternal questionnaire-reported depression (using Edinburgh Postnatal Depression Scale) and anxiety (State-Trait Anxiety Inventory) during 2 nd trimester	Maternal reported doctor-diagnosed wheeze, allergic rhinitis, and atopic eczema	Maternal depression: adjusted Wheeze: OR 2.09 (1.05-4.19) Allergic rhinitis: OR 1.13 (0.54-2.36) Atopic eczema: OR 1.13 (0.43- 2.99) Maternal anxiety state: adjusted Wheeze: OR 1.01 (0.63-1.61) Allergic rhinitis: OR 1.82 (1.17-2.82) Atopic eczema: OR 1.01 (0.56-1.81) Maternal anxiety trait: adjusted Wheeze: OR 1.23 (0.78-1.94) Allergic rhinitis: OR 1.70 (1.10-2.61) Atopic eczema: OR 1.02 (0.58-1.81)	Moderate
Chiu et al 2012; Chiu et al 2014; USA; Prospective cohort study	989	653 for Chiu 2012 ; 708 for Chiu 2014	Up to 2 years	Maternal reported 11 domains of NLEs – e.g. financial, relationships, violence, other housing issues, and discrimination and prejudice; exposure to community violence (ECV). Assessed through self-report at 3 rd trimester	Maternal questionnaire-reported wheeze in the offspring up to 2 years of age	NLEs and wheeze: adjusted 1-2 vs none: OR 1.95 (0.76-5.00) 3-4 vs none: OR 3.55 (1.38-9.15) 5+ vs none: OR 3.79 (1.39-10.3) ECVs scores and wheeze: adjusted Medium vs low: OR 1.34 (0.71-2.52) High vs low: OR 1.95 (1.13-3.36)	Moderate
Cookson et al 2009; UK; Prospective cohort study	14,062	5,810	Up to 7.5 years	Maternal anxiety reported using the Crown-Crisp	Maternal questionnaire-	Anxiety at 18 weeks and current asthma: adjusted 2 nd vs 1 st quart. OR 1.24 (1.00-1.55)	Moderate

				Experimental Index. Assessed at 18 and 32 weeks pregnancy	reported doctor-diagnosed current asthma and consultation for wheeze. Specific IgE sensitisation; and methacholine-based bronchohyper-responsiveness (BHR)	3 rd vs 1 st quart. OR 1.32 (1.07-1.63) 4 th vs 1 st quart. OR 1.53 (1.22-1.93) Anxiety at 32 weeks and current asthma: adjusted 2 nd vs 1 st quart. OR 1.36 (1.09-1.71) 3 rd vs 1 st quart. OR 1.42 (1.14-1.77) 4 th vs 1 st quart. OR 1.65 (1.30-2.08) Results also given in the paper for wheeze at 18 and 81 months; atopic and non-atopic asthma, asthma + BHR; BHR, and atopic sensitisation	
de Marco et al 2012; Italy; Cross-sectional study	3,907	3,758	3-14 years (mean 8.5 years)	Maternal NLEs (mourning, loss of own or husband's job, separation, divorce). Self-reported questionnaire. Assessed for events occurring during any trimester	Maternal questionnaire-reported ever asthma, wheeze, atopic eczema, and allergic rhinitis	Exposure to any NLEs: adjusted Asthma: OR 1.71 (1.02-2.89) Wheeze: OR 1.41 (1.03-1.94) Atopic eczema: OR 1.53 (1.11-2.10) Allergic rhinitis: OR 1.75 (1.08-2.84)	Moderate
Fang et al 2011; Sweden; Retrospective cohort study	Cohort 1 (born 2004-2008): 449,363 Cohort 2 (born 1997-2002): 514,261	Cohort 1: 426,334 Cohort 2: 493,813	Cohort 1: up to 4 years Cohort 2: 7-12 years	Maternal bereavement. Assessed from population register. For bereavement ≤1 year before pregnancy and at different trimesters	Cohort 1: incident asthma Cohort 2: current asthma Assessed from disease register	Cohort 1: bereavement and asthma onset :adjusted Any time during pregnancy: HR 1.00 (0.87-1.14) 1 st trimester: HR 0.98 (0.76-1.27) 2 nd trimester: HR 1.18 (0.94-1.48) 3 rd trimester: HR 0.87 (0.70-1.09) Cohort 2: bereavement and current asthma: adjusted Any time during pregnancy: OR 0.99 (0.84-1.17) 1 st trimester: OR 1.02 (0.76-1.37) 2 nd trimester: OR 0.86 (0.63-1.18) 3 rd trimester: OR 1.07 (0.84-1.39) Estimates reported in the paper also separately for boys and girls; also by relative type, and by cause of death	Strong
Guxens et al 2014; Netherlands; Prospective cohort study	8,880	4,848	Up to 6 years	Maternal reported depression and anxiety symptoms during the 2 nd trimester	Maternal questionnaire-reported doctor-diagnosed ever asthma at 6 years and wheeze at 4 years	Exposure to anxiety + depression: adjusted Ever asthma: OR 1.45 (0.91-2.31) Early wheeze (≤3 yrs): OR 1.23 (0.89-1.69) Late wheeze (at 4 yrs): OR 1.94 (1.04-3.60) Persistent wheeze (1-4 yrs): OR 2.15 (1.47-3.13) Exposure to depression: adjusted Ever asthma: OR 1.33 (0.82-2.16) Early wheeze (≤3 yrs): OR 1.31 (0.97-1.76) Late wheeze (at 4 yrs): OR 2.04 (1.14-3.64) Persistent wheeze (1-4 yrs): OR 1.84 (1.24-2.72) Exposure to anxiety: adjusted	Moderate

						<p>Ever asthma: OR 1.19 (0.76-1.86) Early wheeze (≤ 3 yrs): OR 1.17 (0.88-1.55) Late wheeze (at 4 yrs): OR 1.81 (1.05-3.12) Persistent wheeze (1-4 yrs): OR 1.72 (1.22-2.43)</p>	
Hartwig et al 2014; Australia; Prospective cohort study	2,860	994	14 years	Maternal reported 10 NLEs including separation or divorce, marital problems, problems with children, pregnancy problems, experience of involuntary job loss, partner experienced involuntary job loss, money problems, a residential move, death of a close relative, and death of a close friend. Assessed for events occurring ≤ 18 and 18-34 weeks of pregnancy	Maternal questionnaire reported current asthma, allergic rhinitis, and atopic eczema. At 6 and 14 years Atopic sensitisation also assessed at 6 and 14 years	<p>Exposure to NLEs by 18 weeks: adjusted Asthma at 6 years: 1 vs no NLEs: OR 1.30 (0.86-1.95) 2 vs no NLEs: OR 1.21 (0.72-2.04) 3+ vs no NLEs: OR 1.73 (0.87-3.44) Asthma at 14 years: 1 vs no NLEs: OR 1.12 (0.67-1.87) 2 vs no NLEs: OR 1.08 (0.56-2.07) 3+ vs no NLEs: OR 1.26 (0.54-2.91) Exposure to NLEs by 34 weeks: adjusted Asthma at 6 years: 1 vs no NLEs: OR 1.10 (0.72-1.66) 2 vs no NLEs: OR 1.34 (0.80-2.24) 3+ vs no NLEs: OR 0.99 (0.47-2.08) Asthma at 14 years: 1 vs no NLEs: OR 2.24 (1.33-3.75) 2 vs no NLEs: OR 1.96 (1.01-3.79) 3+ vs no NLEs: OR 1.81 (0.74-4.46) Estimated also reported in the paper for allergic rhinitis, atopic eczema, and atopic sensitisation and by maternal history of asthma</p>	Moderate
Khashan et al 2012; Sweden; Retrospective cohort study	3,290,141	3,193,033	Up to 2-34 years	Bereavement (death of a spouse or child) during the 1 st , 2 nd and 3 rd trimester. Assessed from population register	Asthma hospitalisation and asthma hospitalisation plus other related outcomes (bronchitis, COPD, allergic rhinitis, atopic dermatitis, ALRI). Assessed from national patient register	<p>Death of spouse or child: adjusted Asthma hospitalisation: RR 1.43 (1.06-1.92) Asthma hospitalisation + other related outcomes: RR 1.40 (1.14-1.72) Death of spouse only: adjusted Asthma hospitalisation: RR 1.59 (1.10-2.30) Asthma hospitalisation + other related outcomes: RR 1.64 (1.29-2.10)</p>	Strong
Larsen et al 2014; Denmark; Prospective cohort study	100,418	32,271	Up to 7 years	Psychosocial job strain/stress. Assessed using questionnaire and telephone interviews. Assessed for events	Maternal reported current asthma and atopic dermatitis (AD). Assessed using	<p>Work stress and asthma + AD: adjusted High vs low strain: OR 1.11 (0.88-1.40) Active vs low strain: OR 1.09 (0.95-1.25) Passive vs low strain: OR 1.10 (0.91-1.34)</p>	Moderate

				during 2 nd trimester	ISAAC-based questionnaire	Results also given in the paper for having asthma and no AD and for having AD and no asthma	
Lee et al 2016; USA; Prospective cohort study	989	765	Up to 6 years	Maternal questionnaire-reported NLEs in 11 domains (eg, financial, legal, career, relationships, home safety, community safety, medical issues pertaining to self, medical issues pertaining to others, home issues, authority, and prejudice) any time during pregnancy.	Maternal reported clinician-diagnosed asthma	NLEs and asthma onset: adjusted All 1-2 NLEs vs none: OR 1.05 (0.59-1.88) 3-4 NLEs vs none: OR 1.60 (0.88-2.92) ≥5 NLEs vs none: OR 2.02 (1.05-3.87) Continuous: OR 1.31 (1.07-1.60) Boys: Continuous: OR 1.38 (1.06-1.79) Girls: Continuous: OR 1.17 (0.84-1.63) Results also given in paper for mutual adjustment for prenatal and postnatal maternal stress	
Lefevre et al 2011; France; Case-control study	Cases: 142 Controls: 142	Cases: 138 Controls: 109	<2 years	Maternal questionnaire-reported anxiety and depression symptoms occurring any time during pregnancy	Maternal reported wheeze and objective measures of eosinophilia and IgE sensitisation	Depression and wheeze: adjusted OR 1.55 (0.12-19.8) Anxiety and wheeze: adjusted OR 1.98 (0.69-5.68)	Moderate
Lin et al 2004; Taiwan; Prospective cohort study	353	334	At birth	Maternal questionnaire-reported stress (comprising nervousness, exhaustion, anxiety, tiredness, working stress, and discouragement) at any time during pregnancy	Cord blood IgE sensitisation	Maternal stress and cord blood IgE sensitisation: adjusted OR 7.7 (1.1- 58.9) Maternal nervousness and cord blood IgE sensitisation: adjusted Occasionally vs never/seldom: OR 1.1 (0.5-2.5) Regularly vs never/seldom: OR 4.0 (1.3-12.8)	Moderate
Liu et al 2015; Denmark; Retrospective cohort study	755,358	733,685	Up to 3 years	Maternal depression and use of antidepressant drugs throughout pregnancy. Assessed from population register	Asthma onset assessed from medication prescription database	Maternal depression vs none: adjusted HR 1.25 (1.20-1.30) Use of antidepressant vs non-use: adjusted HR 1.25 (1.18-1.33)	Strong
Liu et al 2015; Denmark; Retrospective cohort study	755,358	750,058	Up to 15 years	Maternal bereavement throughout pregnancy. Assessed from population register	Asthma onset assessed from medication prescription database	Maternal bereavement vs none: adjusted 0-3 years: HR 1.04 (1.00-1.07) 4-15 years: HR 1.02 (0.96-1.09) Timing of maternal bereavement: adjusted 0-3 years: 1 st trimester: HR 1.05 (0.95-1.15); 2 nd trimester: HR 0.99 (0.91-1.08); 3 rd trimester: HR 1.06	Strong

						(0.96-1.17) 4-15 years: 1st trimester: HR 1.06 (0.89-1.27); 2nd trimester: HR 0.88 (0.74-1.04); 3 rd trimester: HR 0.95 (0.77-1.16) Results also given in the paper for type of relative death and cause of death and also stratified by maternal asthma history	
Peters et al 2012; USA; Prospective cohort study	500	403	At birth	Maternal questionnaire-reported NLEs (financial, legal, career, relationships, medical, safety in community and home, difficulty with authority, and discrimination) during 3 rd trimester	Total cord blood IgE sensitisation using CAP fluorescent enzyme immunoassay	NLEs score and log cord blood IgE in all women: β 0.10 (0.03-0.16) NLEs score and log cord blood IgE in atopic women: β 0.13 (0.02-0.24) NLEs score and log cord blood IgE in all women: β 0.05 (-0.03-0.14)	Moderate
Polloni et al 2015; Italy; Case-control study	Cases: 67 Controls: not indicated	Cases: 59 Controls: 59	Mean 8.3 years	Maternal questionnaire-reported NLEs: bereavement, divorce or separation, financial problem, and anxiety symptoms at any time during pregnancy	Food allergy assessed via IgE, skin prick test, clinical evaluation	Only descriptive results are given in paper	Weak
Radhakrishnan et al 2016; Canada; Retrospective cohort study	122,333	122,333	12 years	Maternal use of mental health services any time during the 12 months preceding child birth. Assessed from health administrative records	Physician diagnosed asthma based on health administrative records	Maternal stress and asthma: adjusted OR 1.16 (1.12-1.20)	Moderate
Reyes et al 2011; USA; Prospective cohort study	727	279	Up to 5 years	Maternal questionnaire-reported demoralisation during the 3 rd trimester	Maternal questionnaire-reported wheeze. Serum total and specific IgE sensitisation to inhalant allergens	Unit increase in maternal demoralisation score: adjusted Any wheeze at 5 yrs: OR 1.66 (1.29-2.14) Transient wheeze (3-30 mo): OR 2.25 (1.34-3.76) Late onset wheeze (>30 mo): OR 1.39 (0.61-3.17) Persistent wheeze (3-5 yrs): OR 2.69 (1.52-4.76) Specific IgE sensitisation: OR 0.96 (0.57-1.60)	Moderate
Rosa et al 2016; Mexico; Prospective cohort study	815	417 Boys: 211 Girls: 206	Up to 4 years	Maternal questionnaire-reported NLEs in 11 domains (financial, legal, career, relationship,	Maternal reported ever and current wheeze using the ISAAC	Unit increase in NLEs scores and ever wheeze: adjusted All: RR 1.08 (1.00-1.16) Boys: RR 1.12 (1.02-1.24)	Moderate

				home safety, neighbourhood safety, medical issues (self and others), home, prejudice, and authority) during the 2 nd or 3 rd trimester	questionnaire	Girls: RR 1.03 (0.92-1.15) Unit increase in NLEs scores and current wheeze: adjusted All: RR 1.12 (1.00-1.26) Boys: RR 1.11 (0.96-1.28) Girls: RR 1.11 (0.93-1.34)	
Sausenthaler et al 2009; Germany; Prospective cohort study	3097	3004	Up to 6 years	Indicators of psychological, social, and serological stress at any time during pregnancy. Assessed from maternity cards	Maternal questionnaire-reported atopic eczema	Any stress and atopic eczema: adjusted Atopic eczema by 1 year: OR 1.24 (0.72-2.13) Atopic eczema by 2 years: OR 1.48 (0.95-2.30) Atopic eczema by 3 years: OR 1.06 (0.66-1.70) Atopic eczema by 4 years: OR 1.06 (0.67-1.68) Atopic eczema by 5 year: OR 1.21 (0.76-1.91) Atopic eczema by 6 year: OR 1.13 (0.71-1.79)	Moderate
Sternthal et al 2009; USA; Prospective cohort study	500	478	At birth	Maternal questionnaire-reported exposure to interpersonal trauma during the 3 rd trimester	Total cord blood IgE sensitisation using CAP fluorescent enzyme immunoassay	Interpersonal trauma vs none: adjusted OR 2.19 (0.89-5.38)	Moderate
Turcotte-Tremblay et al 2014; Canada; Prospective cohort study	224	68	11-12 years	Maternal questionnaire-reported posttraumatic stress disorder occurring at any time during pregnancy	Maternal questionnaire-reported doctor-diagnosed current asthma, use of asthma medication and wheeze	Maternal stress and wheeze: adjusted Girls only: OR 1.11 (1.01-1.23) Girls and boys: OR 0.97 (0.91-1.03) Maternal stress and asthma: adjusted Girls only: OR 1.09 (1.00-1.19) Boys only: OR 0.88 (0.78-0.99) Girls and boys: OR 0.88 (0.78-0.99) Maternal stress and asthma medication: adjusted Girls only: OR 1.12 (1.01-1.25) Girls and boys: OR 0.95 (0.87-1.02)	Moderate
Wang et al 2013; Taiwan; Prospective cohort study	24,200	19,381	Up to 3 years	Maternal questionnaire-reported work stress occurring at any time during pregnancy	Maternal questionnaire-reported parent-perceived and doctor-diagnosed AD	Work stress and parent-perceived AD: adjusted Intermediate vs low/no stress: OR 1.20 (1.00-1.46) High vs low/no stress: OR 1.43 (1.20-1.70) Work stress and doctor-diagnosed AD: adjusted Intermediate vs low/no stress: OR 1.22 (1.05-1.41) High vs low/no stress: OR 1.34 (1.16-1.54)	Moderate
Wen et al 2011; Taiwan; Prospective cohort study	1,264	730	Up to 2 years	Maternal questionnaire-reported indicators of stress (vitality, vigour, happiness, anxiety, discouragement,	Maternal reported AD via telephone	Maternal stress and AD: adjusted Medium vs low/no stress: OR 1.10 (0.40-2.80) High vs low/no stress: OR 2.30 (1.10-5.30)	Moderate

				nervousness, tiredness, exhaustion, and work stress) occurring during the 3 rd trimester			
Wood et al 2011; USA; Prospective cohort study	1,850	515	1 year	Maternal questionnaire-reported exposure to work stress, community violence, anxiety, and depression during the 2 nd and 3 rd trimester	Maternal questionnaire-reported wheeze and atopic eczema. IgE sensitisation	All estimates presented in figures. Higher maternal stress scores appeared associated with increased risk of wheeze and atopic eczema	Moderate
Wright et al 2010; USA; Prospective cohort study	1,853	560	At birth	Maternal questionnaire-reported difficult life circumstances, economic strain, neighbourhood /block conditions, perceived community violence and housing worries.	Cord blood cytokine responses	Only p-values presented in paper, hence results difficult to interpret	Moderate

Table S2: Domain specific quality assessment of studies on the association between maternal psychosocial stress during pregnancy and risk of allergy and asthma in the offspring

Reference; country	Overall risk of bias assessment	Risk of bias assessment for study components				
		Study design	Exposure assessment	Outcome assessment	Selection bias	Confounding
Andersson et al 2016; USA	Moderate	Strong	Moderate	Strong	Moderate	Moderate
Bandoli et al 2016; USA	Moderate	Moderate	Moderate	Moderate	Weak	moderate
Bidaki et al 2011; Iran	Moderate	Strong	Moderate	Strong	Moderate	Weak
Braig et al 2016; Germany	Strong	Strong	Strong	Strong	Moderate	Moderate
Chang et al 2016; South Korea	Moderate	Strong	Moderate	Moderate	Moderate	Moderate
Cheng et al 2015; Singapore	Moderate	Strong	Moderate	Moderate	Moderate	Moderate
Chiu et al 2012; Chiu et al 2014; USA	Moderate	Strong	Moderate	Moderate	Moderate	Moderate
Cookson et al 2009; UK	Moderate	Strong	Moderate	Moderate	Moderate	Moderate
de Marco et al 2012; Italy	Moderate	Weak	Moderate	Moderate	Moderate	Moderate
Fang et al 2011; Sweden	Strong	Strong	Strong	Strong	Moderate	Moderate
Guxens et al 2014; Netherlands	Moderate	Strong	Moderate	Moderate	Moderate	Moderate
Hartwig et al 2014; Australia	Moderate	Strong	Moderate	Moderate	Weak	Moderate
Khashan et al 2012; Sweden	Strong	Strong	Strong	Strong	Moderate	Moderate
Larsen et al 2014; Denmark	Moderate	Strong	Moderate	Moderate	Moderate	Moderate
Lee et al 2016; USA	Moderate	Strong	Moderate	Moderate	Moderate	Moderate
Lefevre et al 2011; France	Moderate	Moderate	Moderate	Moderate	Weak	Moderate
Lin et al 2004; Taiwan	Moderate	Strong	Moderate	Strong	Moderate	Moderate
Liu et al 2015; Liu et al <i>in press</i> ; Denmark	Strong	Strong	Strong	Strong	Moderate	Moderate

Peters et al 2012; USA	Moderate	Strong	Moderate	Strong	Moderate	Moderate
Polloni et al 2015; Italy	Weak	Moderate	Moderate	Moderate	Weak	Weak
Radhakrishnan et al; Canada	Moderate	Strong	Moderate	Moderate	Moderate	Moderate
Reyes et al 2011; USA	Moderate	Strong	Moderate	Moderate	Weak	Moderate
Rosa et al 2016; Mexico	Moderate	Strong	Moderate	Moderate	Moderate	Moderate
Sausenthaler et al 2009; Germany	Moderate	Strong	Strong	Moderate	Moderate	Moderate
Sternthal et al 2009; USA	Moderate	Strong	Moderate	Strong	Moderate	Moderate
Turcotte-Tremblay et al 2014; Canada	Moderate	Strong	Moderate	Moderate	Weak	Moderate
Wang et al 2013; Taiwan	Moderate	Strong	Moderate	Moderate	Moderate	Moderate
Wen et al 2011; Taiwan	Moderate	Strong	Moderate	Moderate	Moderate	Moderate
Wood et al 2011; USA;	Moderate	Strong	Moderate	Moderate	Moderate	Moderate
Wright et al 2010; USA	Moderate	Strong	Moderate	Strong	Moderate	Moderate

The overall risk assessment was based on the component risk assessments (i.e., on the suitability of the study design for the research question, validity of exposure and outcome assessments, potential for selection bias, adjustment for confounding factors).

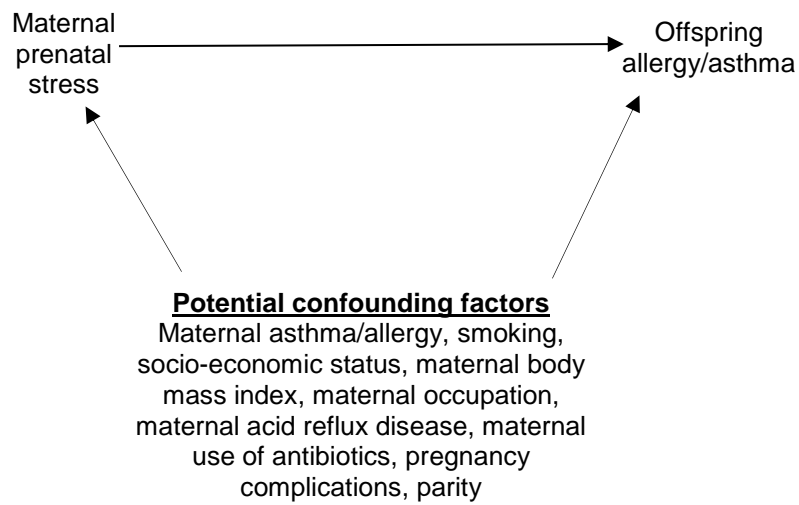


Figure S1 Causal diagram describing the association between maternal prenatal stress and risk of allergy and asthma in the offspring

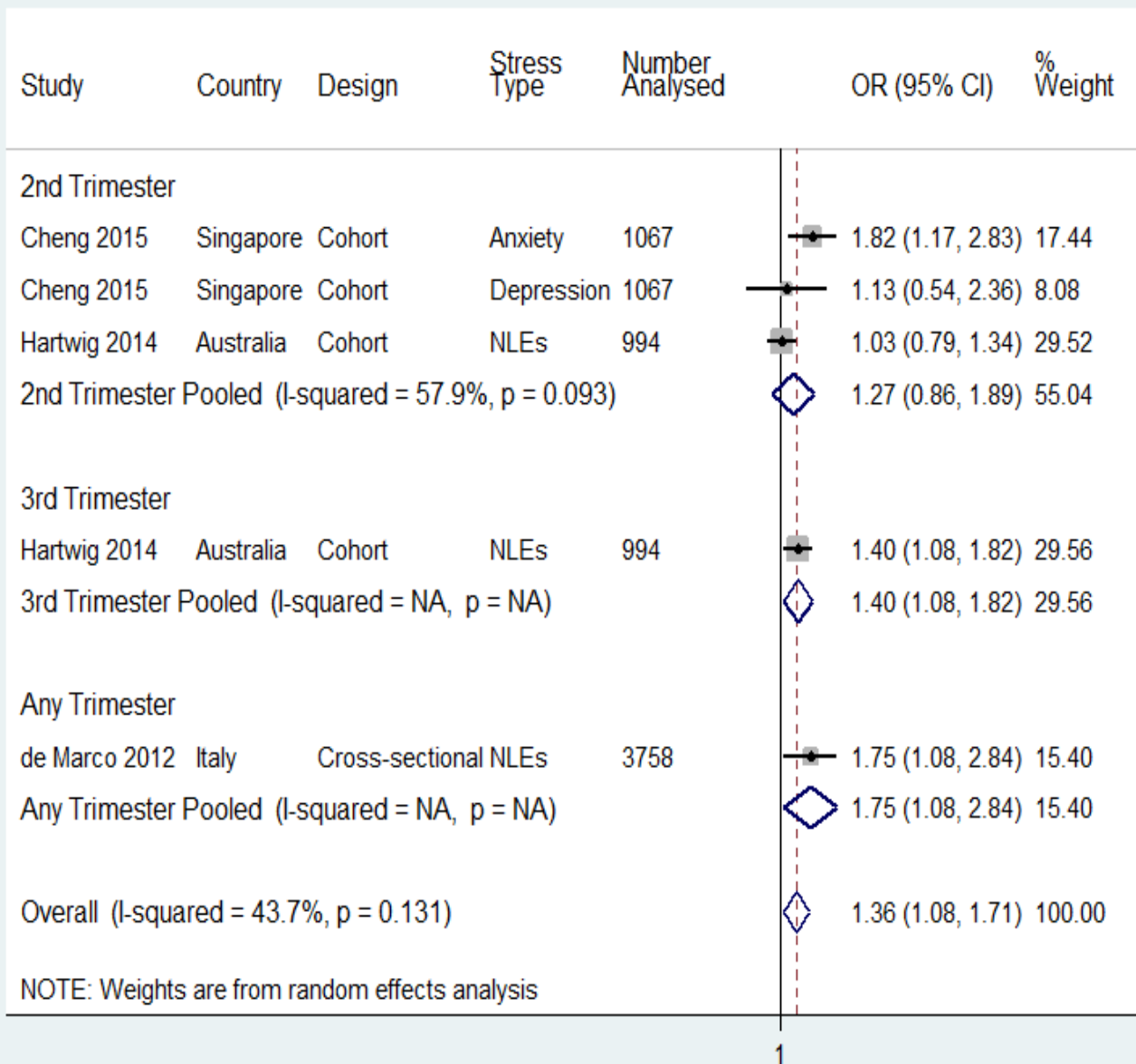


Figure S2. Association between maternal prenatal stress and risk of allergic rhinitis in the offspring, by timing of exposure during pregnancy: No major differences when Hartwig 2014's 6-year-olds and 14-year-olds were analysed separately, hence we presented the results for 14-year-olds

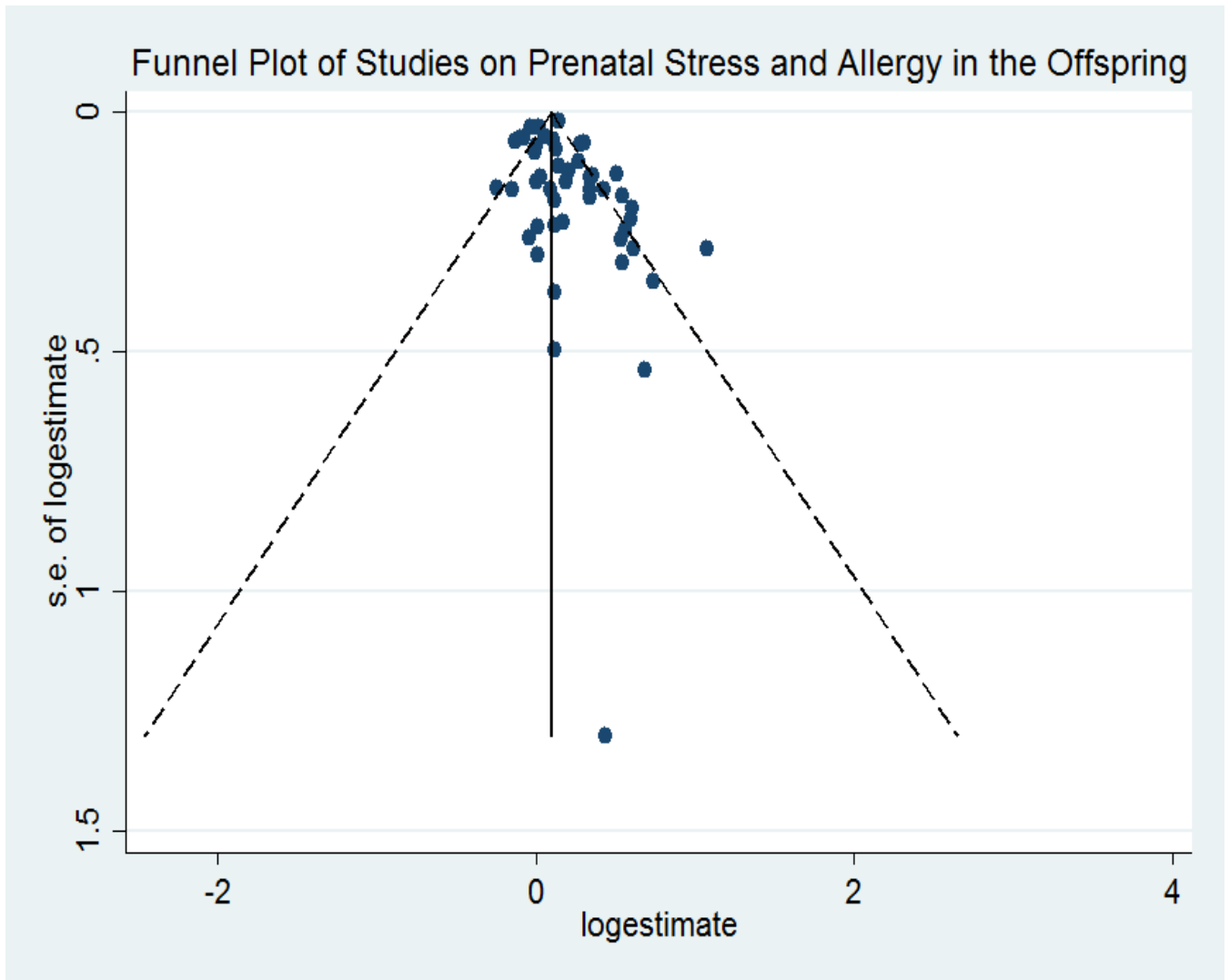


Figure S3. Funnel plot investigating the presence of publication bias or small study effect in studies on the association between maternal prenatal stress and risk of allergy and asthma in the offspring