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Timing of routine infant vaccinations and risk of food allergy and eczema at one year of age

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33
34 **Short title:** Vaccination timing and risk of food allergy and eczema

35
36
37 **Abstract**

38 **Background:** Epidemiological evidence suggests that routine vaccinations can have non-
39 targeted effects on susceptibility to infections and allergic disease. Such effects may depend
40 on age at vaccination, and a delay in pertussis vaccination has been linked to reduced risk of
41 allergic disease. We aimed to test the hypothesis that delay in vaccines containing diphtheria-
42 tetanus-acellular pertussis (DTaP) is associated with reduced risk of food allergy and other
43 allergic diseases.

44 **Methods:** HealthNuts is a population-based cohort in Melbourne, Australia. 12 month-old
45 infants were skin prick tested to common food allergens, and sensitized infants were offered
46 oral food challenges to determine food allergy status. In this data linkage study, vaccination
47 data for children in the HealthNuts cohort were obtained from the Australian Childhood
48 Immunisation Register. Associations were examined between age at the first dose of DTaP
49 and allergic disease.

50 **Results:** 109 of 4433 children (2.5%) received the first dose of DTaP one month late (delayed
51 DTaP). Overall, delayed DTaP was not associated with primary outcomes of food allergy
52 (adjusted odds ratio (aOR) 0.77; 95% CI 0.36-1.62, p=0.49) or atopic sensitization (aOR
53 0.66; 95% CI 0.35-1.24, p=0.19). Among secondary outcomes, delayed DTaP was associated
54 with reduced eczema (aOR 0.57; 95% CI 0.34-0.97, p=0.04) and reduced use of eczema
55 medication (aOR 0.45; 95% CI 0.24-0.83, p=0.01).

56 **Conclusions:** There was no overall association between delayed DTaP and food allergy,
57 however children with delayed DTaP had less eczema and less use of eczema medication.
58 Timing of routine infant immunizations may affect susceptibility to allergic disease.

59
60 **Study registration:** This observational study was registered with ANZCTR, trial ID
61 ACTRN12614001193662.

62

63 **Key Words**

64 Atopic hypersensitivity, food allergy, eczema, DTaP vaccine, infant

65

66

67 **Introduction**

68 The prevalence of food allergy is rising in industrialized countries and Melbourne, Australia,
69 has the highest reported prevalence of childhood food allergy in the world (1). While some
70 environmental factors have been clearly associated with protection against food allergy,
71 including older siblings, pets ownership, timing of introduction of allergenic food (2), and
72 vitamin D sufficiency (3), the causes for the increasing prevalence of food allergy are largely
73 unknown.

74

75 There is evidence that immunizations given early in life have the potential to deviate the
76 immune system toward a more, or less, allergic phenotype. Bacille Calmette-Guérin (BCG),
77 the live-attenuated tuberculosis vaccine, has been associated with protection against allergic
78 disease (4) and randomized trials are ongoing to test this association (5). In contrast, studies
79 of the associations between inactivated pertussis vaccines and allergic disease have shown
80 conflicting results with statistical heterogeneity (6-9). The only randomised trial of pertussis
81 vaccine and allergy compared both whole cell and acellular pertussis vaccines against a
82 diphtheria-tetanus control vaccine and found no large differences in eczema or other allergic
83 diseases at age 2 ½ years (10). However this study did not include an unvaccinated control
84 group. If other components (diphtheria or tetanus toxoids or adjuvants) contribute a
85 biological effect on allergic disease, the ability of the randomised trial to detect differences
86 between vaccine groups may have been limited (11).

87

88 Two other important issues might impact on these previous observational studies of pertussis
89 vaccines. First, confounding may occur due to factors associated with receipt or refusal of
90 vaccination. Second, heterogeneity of vaccination timing may lead to heterogeneity in study
91 findings, as the age of exposure to the immune modulating effects of vaccines may be
92 important for the resulting immune phenotype (12). Five studies have investigated age of
93 pertussis vaccination and allergic disease (9, 13-16), with three suggesting that delayed
94 vaccination is protective against asthma (15), hay fever (13), and atopic sensitization (14),
95 and two studies showing no association between timing of pertussis vaccination and asthma

96 (9, 16) or eczema (16). Four of these studies investigated the whole cell pertussis vaccine no
97 longer used in most industrialized countries (13-16), only one investigated eczema (16), and
98 none investigated food allergy.

99

100 The HealthNuts cohort is a population-sample of one year old infants recruited to study
101 prevalence and risk factors for food allergy (17). By linking with vaccination data from the
102 Australian Childhood Immunisation Register (ACIR), the HealthNuts cohort was utilized to
103 test the hypothesis that a delay in the first dose of an acellular-pertussis containing vaccine is
104 associated with reduced prevalence of food allergy, eczema, wheeze or bronchiolitis in the
105 first year of life.

106

107 **Methods**

108 **1. The HealthNuts Cohort**

109 The HealthNuts cohort is comprised of 5276 infants who were recruited at immunization
110 clinics across Melbourne, Australia between 2007 and 2011 (17, 18). Parents provided
111 written informed consent, completed an extensive survey of demography and history of
112 allergic diseases, and their infants were examined for eczema and underwent a skin prick test
113 (SPT) to common childhood food allergens (whole hen's egg, peanut, sesame, and shellfish
114 or cow's milk) (1). Children with a SPT reaction to any allergen (SPT wheal \geq 1mm after
115 subtracting the negative control) were invited to attend the oral food challenge clinic (928 out
116 of 1089 (85%) sensitized children attended) along with a random sample of SPT negative
117 controls (n=218, approximately 19% of the cohort who received oral food challenges). We
118 used 1mm as the criterion for invitation to the oral food challenge clinic to ensure no children
119 with potential food allergy were missed. In the oral food challenge clinic SPTs were repeated,
120 blood samples were taken for specific IgE, and infants were given oral food challenges with
121 each food to which they were sensitized (18).

122

123 **2. Immunization exposures and Data linkage**

124 Over the birth years of the cohort, the Australian National Immunisation Program Schedule
125 included a birth dose of hepatitis B vaccine followed by diphtheria-tetanus-acellular pertussis
126 vaccine (DTaP) at two, four and six months of age, usually as part of Infanrix Hexa® (GSK,
127 Boronia, Victoria, Australia) also containing inactivated polio vaccine (IPV), hepatitis B
128 vaccine and Haemophilus influenzae type b vaccine (Supplementary Figure 1). A 13-valent

129 pneumococcal conjugate vaccine (PCV) and oral rotavirus vaccination was also usually co-
130 administered. In Australia, all childhood immunizations are recorded in the ACIR.

131

132 Children were eligible for the present study if they remained in the HealthNuts cohort with
133 up-to-date contact details in May 2014. A letter was sent to the parents of all eligible children
134 outlining the details of the study. Unless parents opted out, data on routine vaccinations were
135 sought from ACIR between October 2014 and March 2015. Children were matched on first
136 name, surname and date of birth with or without postcode; only definite matches were
137 included. All children who received acellular pertussis vaccine also received diphtheria and
138 tetanus components; thus the age at first dose of DTaP was considered the primary exposure
139 irrespective of other vaccines co-administered. Delayed DTaP was defined as the first dose
140 given after 90 days of age (one month late) as per the Australian National Immunisation
141 Program. Children with missing data for the first dose of DTaP but with data for subsequent
142 doses (n=46) were excluded from the primary analyses, as it was likely these children
143 received prior doses of DTaP at an unknown time. Vaccination data was available from
144 personal records for eight of these 46 children as part of HealthNuts age six follow up
145 (currently underway), all of whom received a dose of DTaP prior to the first dose recorded on
146 ACIR (seven were vaccinated on-time and one was delayed).

147

148 **3. Allergic Disease Outcomes**

149 *Primary outcomes*

150 *Food allergy:* Children were classified as food allergic if they had a SPT wheal ≥ 2 mm (after
151 subtracting the negative control) or specific IgE > 0.35 kU_A/L at the oral food challenge
152 clinic visit and any of the following within two hours of oral food challenge: ~~three~~
153 concurrent noncontact urticaria lasting five minutes or more; perioral or periorbital
154 angioedema; vomiting; or circulatory or respiratory compromise. Only children with
155 reactions to egg, peanut and sesame were considered food allergic for this analysis since
156 challenges were not performed for cow's milk and shellfish. Children were also deemed food
157 allergic (without performing oral food challenges) if they had a positive SPT and a confirmed
158 reaction to egg within the past one month, or to peanut or sesame within the past two months
159 (1). Children with a positive food challenge but negative SPT and specific IgE < 0.35 kU_A/L
160 (Figure 1) were excluded from the food allergy analyses because it was unclear if they had
161 IgE mediated food allergy.

162

163 *Atopic sensitization:* All children with a SPT wheal \geq 2mm (after subtracting the negative
164 control) to egg, peanut or sesame at the community clinic were classified as having atopic
165 sensitization.

166

167 ***Secondary outcomes***

168 *Eczema:* eczema was defined as established diagnosis by a doctor with associated use of
169 treatments (medications, topical steroids or moisturizers), or eczematous rash observed by a
170 trained nurse at the time of recruitment. Children who were diagnosed with eczema before
171 three months of age (prior to first scheduled vaccinations plus one month window period)
172 were excluded as eczema in these children was, by definition, unrelated to vaccination
173 (n=313, 21% of 1474 children who otherwise met criteria for eczema); thus onset of eczema
174 was between three and 12 months of age.

175

176 *Eczema medication:* Use of eczema medication was parent reported and included oral
177 medication or topical steroids (but not moisturizers) to treat an itchy rash at any time in the
178 first year of life. Similar to the eczema outcome, children diagnosed with eczema prior to
179 three months of age were excluded from eczema medication analyses (n=224, 19% of 1209
180 children who otherwise met criteria for use of eczema medication).

181

182 *Wheeze:* 'wheeze ever' reported by parents at one year of age.

183

184 *Bronchiolitis admission:* Hospital admissions for bronchiolitis at any time in the first year of
185 life reported by parents at one year of age.

186

187 **4. Statistical Analyses**

188 Demographic variables were compared between groups using Chi square or Kruskal-Wallis
189 tests. The primary analyses were performed using logistic regression to produce odds ratios
190 (OR) for association between delayed DTaP and allergic disease outcomes. Multivariate
191 analyses were performed adjusting for pre-specified potential confounders: sex
192 (female/male), antibiotic use (categorized as yes/no due to excess missing data on number of
193 antibiotic courses), day-care attendance (yes/no), number of siblings (0, 1-2, \geq 3), birth-
194 country of the parents (both Australian, one or both parents born East Asia, either parent born
195 elsewhere), presence of smokers at home (yes/no), and socio-economic indexes for areas
196 (SEIFA; quintiles) (19). The SEIFA was used to estimate socioeconomic status on the basis

197 of postcode (3, 19) due to incomplete parental income data. Other potential confounders
198 (listed and categorized in Table 1) were not included in multivariate models because their
199 inclusion did not alter any estimate by more than 10%. Analyses were stratified by sex as pre-
200 specified in the analysis plan, and a Wald test for homogeneity of effects was performed on
201 all stratified analyses. A sensitivity analysis was performed designating children with
202 indeterminate food allergy status as non-allergic unless they had a parent reported history
203 consistent with IgE mediated food allergy at any age or a SPT wheal greater than the 95%
204 positive predictive value for food allergy (≥ 4 mm to egg or ≥ 8 mm to peanut or sesame) (20),
205 in which case they were designated as food allergic. A sensitivity analysis for eczema was
206 performed by including all children regardless of age of eczema diagnosis. To investigate for
207 reverse causation between eczema and delayed DTaP, a Cox regression model was
208 constructed including time at risk from birth until age of DTaP vaccination, and including
209 time as exposed from the age of doctor diagnosed eczema; the proportional hazards
210 assumption was not violated ($p=0.30$, Schoenfeld residuals).

211
212 Children with missing data on potential confounders were excluded from the multivariate
213 analyses. P values less than 0.05 were considered statistically significant. The population
214 sample size of 5000 was derived for recruitment into the original cohort (3). Given that DTaP
215 data were available for 4433 children and 109 (2.5%) had a one month delay in DTaP, the
216 study ultimately had a power of 0.80 to detect a 73% reduction in food allergy associated
217 with delayed vaccination in the unadjusted analysis. Statistical analyses were conducted in
218 Stata version 11 (College Station, Texas, USA).

219

220 **5. Ethics and Consent**

221 The protocol for the original HealthNuts study was approved by the Human Research Ethics
222 Committees at the Office for Children, Government of Victoria, the Department of Human
223 Services, Government of Victoria, and at the Royal Children's Hospital, Melbourne.
224 Approval for linkage with immunization data was granted by the Human Research Ethics
225 Committee at the Royal Children's Hospital, Melbourne. Written information was provided
226 to parents of participants with the option to opt-out. The methodology, outcomes and analysis
227 plan for this observational study were registered prior to data linkage with ANZCTR (trial ID
228 ACTRN12614001193662).

229

230 **Results**

231 **1. Background and demography**

232 Of the 5276 children included in the original cohort, vaccination data was sought for 4834
233 (92%) with current contact details, and complete data on vaccinations were available for 4487
234 (85%) children (Figure 1). There were differences between those with and without
235 vaccination data available, including increased prevalence of food allergy and eczema
236 amongst those with available vaccination data (Supplementary Table 1).

237

238 **2. Vaccination exposure and predictors of delayed vaccination**

239 Overall, 4433 out of 4487 (99%) children were recorded as having received a first dose of
240 DTaP. Of these children, all received at least two doses, and 4402 (99.3%) received all three
241 doses. The first dose of DTaP was co-administered with IPV in 4415 children (99.6%), with
242 hepatitis B vaccine in 4400 (99.3%), with Hib in 4416 (99.6%) (usually as part of *Infanrix*
243 *Hexa®*), with PCV in 4396 (99.2%) and with rotavirus vaccine in 4173 (94.1%).

244

245 109 children (2.5%) received the first dose of DTaP one month late (delayed DTaP). Median
246 age of DTaP was 63 days in the on-time group and 103 days in the delayed DTaP group
247 (Table 1). Vaccine timeliness improved somewhat over the period of the study; 70/2378
248 (2.9%) of children born before 1/1/2009 had delayed DTaP, whereas 39/2055 (1.9%) of
249 children born after 1/1/2009 had delayed DTaP. Factors associated with delayed DTaP were
250 older age at recruitment, lack of attendance at child-care, having siblings, smokers at the
251 home, and never having had artificial formula (Table 1). Of these factors, only siblings was
252 also associated with food allergy (an inverse association, $p=0.04$ for 1-2 siblings and $p=0.001$
253 for ≥ 3 siblings) (2) and only smokers at the home was associated with eczema (an inverse
254 association, $p=0.02$).

255

256 **3. Primary outcomes**

257 There was no significant association between delayed DTaP and food allergy (adjusted odds
258 ratio (aOR) 0.77; 95% CI 0.36-1.62, $p=0.49$, Table 2). There was no overall association
259 between delayed DTaP and atopic sensitization (Table 2). In the pre-planned sex-stratified
260 analyses, females tended to have less atopic sensitization if they had delayed DTaP (aOR
261 0.25; 0.06-1.04, $p=0.06$), and this association tended to be different from the association in
262 males (p for interaction= 0.09 ; Table 2). In a sensitivity analysis assuming food allergy status
263 on the basis of skin prick wheal size amongst those with indeterminate food allergy status, the

264 association between delayed DTaP and food allergy was similar (aOR 0.87; 0.45-1.68,
265 p=0.69).

266

267 **4. Secondary outcomes**

268 Children with delayed DTaP had reduced odds of eczema compared to those vaccinated on
269 time (aOR 0.57; 0.34-0.97, p=0.04, Table 3), with a similar magnitude of association in boys
270 and girls (Supplementary Table 2). Similarly there was an association between delayed DTaP
271 and reduced use of eczema medication (aOR 0.45; 0.24-0.83, p=0.01, Table 3). In a
272 sensitivity analysis, including all children regardless of the age of eczema diagnosis made
273 minimal difference to these estimates (eczema: aOR 0.60; 0.37-0.96, p=0.03; eczema
274 medication: aOR 0.47; 0.27-0.82, p=0.008). In *post hoc* analyses, there were no large
275 differences in the association between delayed DTaP and eczema amongst subgroups
276 (Supplementary Table 3). There was no association between doctor-diagnosed eczema and
277 subsequent delayed DTaP (Hazard Ratio 0.98; 0.86-1.11, p=0.70), giving no suggestion of
278 reverse causation. There were no significant associations between delayed DTaP and
279 bronchiolitis admissions or wheeze (Table 3).

280

281 **Discussion**

282 Overall we found no significant associations between delayed DTaP and our primary
283 outcomes of food allergy or atopic sensitization at one year of age. However we found
284 children with their first dose of DTaP containing vaccine delayed by one-month had
285 significantly reduced eczema and reduced use of eczema medication, even after accounting
286 for a variety of potential confounding factors. Additionally, there was some evidence of a
287 differential association by sex, where girls with delayed DTaP tended to have reduced atopic
288 sensitization whereas boys did not. Other vaccinations were invariably co-administered,
289 including diphtheria, tetanus, IPV, hepatitis B, Hib, PCV, rotavirus; thus if the observed
290 associations are causal, it is unclear whether DTaP, another vaccine or a combination of
291 vaccines is responsible.

292

293 This study is the most comprehensive investigation into vaccinations and food allergy to date,
294 and the lack of overall association between DTaP timing and food allergy is reassuring.
295 However this study was limited by statistical power as only 2.5% of children had delayed
296 vaccination, considerably fewer than 4.9% found in a 2001 Australian cohort (21). This
297 improvement in timeliness may be specific to the HealthNuts population or may represent a

298 trend over time, especially given that the recent introduction of rotavirus vaccine has
299 apparently increased timeliness of other co-administered vaccines (22).

300

301 Other important limitations exist. There was a bias towards participation amongst those with
302 food allergy and eczema thus population risk of allergic diseases were slightly overestimated
303 (1). However the overall participation rate was high, and there is no reason to suggest that
304 this participation bias would have affected our analysis of vaccination timing. Over 20% of
305 eczema cases were excluded due to having a diagnosis of eczema prior to age of scheduled
306 vaccination, thus the eczema results presented here pertain only to eczema between three and
307 12 months of age. Importantly, our primary objective was to study timing of the acellular
308 pertussis vaccine, but the almost invariable co-administration of other vaccinations means
309 that we are unable to determine which component may be responsible for modifying risk of
310 eczema. Furthermore, our findings are unable to attribute any risk of allergic disease to
311 vaccination *per se* as all included children were vaccinated.

312

313 By comparing early versus late vaccination, we have eliminated confounding associated with
314 reasons for receipt or refusal of vaccination. However our results have potential to be
315 confounded by reasons for delayed vaccination. Febrile episodes are associated with
316 vaccination delay but fever only after six months of age is linked to protection against
317 allergic disease (23, 24), and it is unlikely that vaccination would be delayed by one month
318 due to fever in many children. Rotavirus vaccine is often withheld when vaccination is
319 delayed because of strict upper age limits for its use (22), but there was no evidence that
320 differential vaccination with rotavirus vaccine affected the results. We investigated many
321 other factors associated with vaccination delay (25) and found no evidence that confounding
322 was responsible for the findings; however we are unable to exclude residual confounding due
323 to the observational design of the study and therefore causation cannot be ascribed. We have
324 not identified any sources of bias that could explain these findings and there was no
325 association between early doctor-diagnosed eczema and subsequent vaccination delay
326 indicating that reverse causation is unlikely.

327

328 The rationale behind this investigation was evidence that delayed administration of
329 diphtheria-tetanus-whole cell pertussis (DTwP) vaccination in infancy might reduce
330 subsequent risk of asthma (15), hay fever (13), and possibly atopic sensitization (14).
331 Additionally, delayed DTwP vaccination provided a survival advantage for girls in a high

332 mortality setting (26), while DTaP-IPV-Hib vaccination was a risk factor for infectious
333 disease hospital admissions in a low mortality setting (27). The World Health Organization
334 has recently acknowledged the importance of the non-targeted (also called non-specific)
335 effects of vaccinations and has called for further research into the epidemiological and
336 underlying immunological mechanisms of such effects (28).

337

338 Various observational studies of pertussis vaccination and atopic disease have found
339 protection, no association, or increased risk associated with receipt of vaccination (6-9, 29).
340 One international study reported reduction in eczema severity associated with pertussis
341 vaccinations, with an apparent dose-response to total number of vaccinations received (30).
342 Many of these studies did not report age of vaccination and thus heterogeneity in vaccination
343 timing may have led to conflicting results in studies of pertussis vaccination versus no
344 vaccination. No previous studies have examined timing of pertussis vaccination and food
345 allergy, and only one study examined eczema (16). In a large UK cohort, no association
346 between DTwP-IPV timing and eczema was found (16); however vaccination timing was
347 divided into quartiles and age of vaccination was not reported, thus it is unclear if there was
348 any practical difference in vaccination timing between groups. The association between
349 delayed DTaP and eczema in the present study resembles a previous observation where
350 delayed DTwP was associated with reduced risk of childhood asthma (15).

351

352 These findings need to be considered in the context of pertussis disease. In the USA, pertussis
353 has a case fatality of 0.6% in two-month old infants (31), and a single dose of acellular
354 pertussis vaccine is highly protective against pertussis disease and pertussis mortality (31,
355 32). There is a movement to advance the age of the first pertussis vaccine to reduce pertussis
356 morbidity and mortality in infants (33) and the primary immunization series is now
357 encouraged from six weeks of age in Australia. If delayed vaccination is proven beneficial for
358 allergic disease, such benefits would need to be carefully measured against the specific
359 advantage of early vaccination in the relevant population (31, 33) while considering impact
360 on other vaccines in the schedule (22). It should also be noted that Victoria has recently
361 introduced a maternal pertussis immunization program recommended in the third trimester of
362 pregnancy (34), which may have implications on the immune response to the infant dose of
363 pertussis vaccine and any future evaluations on vaccination timing and atopic disease.

364

365 While trained innate immunity and T cell cross-reactivity may explain some non-targeted
366 effects of vaccination in relation to susceptibility to infection (35, 36), mechanisms to explain
367 altered risk of allergic disease are unknown. However T helper (T_H) 2 stimulation by
368 vaccinations could theoretically offer an explanation, as acellular pertussis vaccines are
369 strong T_H2 stimulants (32, 37, 38) and T_H2 polarization is associated with development of
370 food allergy and atopic eczema. Components of the *Bordetella pertussis* cell wall may have
371 an inhibitory role on the development of IgE in relation to other vaccine components (39),
372 suggesting that DTaP, which lacks the cellular components, has greater potential to be
373 allergenic than DTwP. Also promoting a T_H2 response to non-vaccine antigens are
374 pneumococcal vaccines (40) and aluminium adjuvant (41), used in both DTaP and
375 pneumococcal vaccines. Therefore multiple components of the primary immunization series
376 are theoretically capable of initiating a generalized T_H2 bias.

377

378 *Staphylococcus aureus* colonization is implicated in the pathogenesis of infant eczema, as
379 infants with eczema have excessive T_H2 cytokine responses to staphylococcal superantigens
380 (42). Thus it is possible that a T_H2 stimulant such as DTaP or another vaccine component
381 given during a critical window in infancy could have a bystander effect whereby predisposed
382 infants who are colonized with *S. aureus* become sensitized to staphylococcal superantigens
383 and develop eczema, without having the stronger T_H2 bias required to cause other allergic
384 diseases (42).

385

386 In conclusion, delayed DTaP vaccination was not associated with food allergy or atopic
387 sensitization overall, but was associated with less eczema and less use of eczema medication.
388 Additionally there was a borderline association between delayed DTaP and reduced atopic
389 sensitization amongst girls. This finding is consistent with the observation that the non-
390 targeted effects of vaccines tend to be sex differential with females generally being more
391 susceptible than males (43). These results warrant further investigation. Timing of routine
392 immunizations in infancy may affect susceptibility to allergic disease.

393

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412

413 **Author Contributions**

414 All authors interpreted the results, critically reviewed and approved the final manuscript as
415 submitted. NK conceptualised and designed this data-linkage study, coordinated collection of
416 vaccination data, analysed the data and drafted the manuscript. JJK is a co-investigator for the
417 original HealthNuts study, designed this data-linkage study, and assisted with statistical
418 analyses. NWC coordinated vaccination data collection. SB assisted with data management
419 and processing. LCG is a co-investigator for the original HealthNuts study and assisted with
420 statistical analyses. AJL, MLKT, MW and A-LP are all co-investigators of the original
421 HealthNuts study. SCD is a co-investigators of the original HealthNuts study and assisted
422 with study design and statistical analyses. KJA is the principle investigator of the original
423 HealthNuts study and conceptualised and designed this data-linkage study.

424

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430

431 **Conflict of Interest Statement**

432 None of the authors have conflicts of interest to declare.

433

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552

553 **Figure legends**

554 Figure 1 – HealthNuts cohort and participant flow for analysis of vaccination timing
555 ACIR=Australian Childhood Immunisation Register. *Unable to determine food allergy
556 status due to failure to attend food challenge clinic (n=230), inconclusive food challenge
557 (n=18), incomplete food challenge to all foods to which infant was sensitized (n=22), or
558 positive food challenge in absence of atopic sensitization (n=27).

559

560 Supplementary Figure 1 – Australian National Immunisation Program Schedule over the birth
561 years of the HealthNuts cohort. Hep B=hepatitis B vaccine, DTaP=diphtheria-tetanus-
562 acellular pertussis, IPV=inactivated polio vaccine; Hib=*Haemophilus influenzae* type B
563 vaccine; PCV=7-valent pneumococcal vaccine; MMR=measles-mumps-rubella vaccine; Men
564 C=meningococcal C vaccine.

565

566

567 Table 1 – Comparison of demography between children with on-time and delayed 1st dose of DTaP

		n	1 st dose DTaP		p
		4433	< 1 month late (n=4324)	≥ 1 month late (n=109)	
Sex	Female	4433	2155/4324 (50%)	45/109 (41%)	0.08
SEIFA	median (10 th -90 th centile)	4428	1057 (987-1108)	1057 (987-1108)	0.79
Ever had antibiotics		4289	2136/4183 (51%)	51/106 (48%)	0.55
Attends child care		4338	1231/4231 (29%)	18/107 (17%)	0.006
Siblings	None	4386	2150/4278 (50%)	20/108 (19%)	
	1-2	4386	1418/4278 (33%)	46/108 (43%)	
	3 or more	4386	710/4278 (17%)	42/108 (39%)	<0.001
Parent country of birth	Any parent East Asia	4308	469/4201 (11%)	17/107 (16%)	
	Both parents Australia	4308	2557/4201 (61%)	67/107 (63%)	
	Any parent other country	4308	1175/4201 (28%)	23/107 (21%)	0.16
Smoker at the home		4403	905/4296 (21%)	38/107 (36%)	<0.001
Maternal smoking during pregnancy		4311	183/4207 (4%)	8/104 (8%)	0.10
Atopic family history		4433	2942/4324 (68%)	77/109 (70%)	0.57
Birthweight (grams) n=4349	Median (10 th -90 th centile)	4349	3420 (2730-4060)	3432 (2855-4150)	0.39
Premature (<37 weeks gestation)		4163	244/4066 (6%)	6/97 (6%)	0.94
Born in winter months		4433	1107/4324 (26%)	27/109 (25%)	0.84
Breastfed > 6 months of age		4229	2436/4129 (59%)	55/100 (55%)	0.42
Ever had artificial formula		4114	3197/4011 (80%)	73/103 (71%)	0.03

Early introduction of egg into diet	≤6 months of age	4248	1007/4142 (24%)	39/106 (37%)	0.003
Cat at the house	No	4427	3585/4318 (83%)	88/109 (81%)	
	Outside	4427	134//4318 (3%)	4/109 (4%)	
	Inside	4427	599/4318 (14%)	17/109 (16%)	0.82
Dog at the house	No	4430	2945/4321 (68%)	84/109 (77%)	
	Outside	4430	459/4321 (11%)	5/109 (5%)	
	Inside	4430	917/4321 (21%)	20/109 (18%)	0.07
Date of recruitment (dd/mm/yy);	Median (10 th -90 th centile)	4433	07/12/09 (07/10/08–24/03/11)	23/09/09 (21/10/08–24/03/11)	0.15
Age at recruitment (days)	Median (10 th -90 th centile)	4433	378 (365-411)	391 (352-448)	<0.001
Age of 1 st dose of DTaP (days)	Median (10 th -90 th centile)	4433	63 (56-73)	103 (91-216)	<0.001

568

569 DTaP=diphtheria-tetanus-acellular pertussis vaccination; SEIFA = Socio-Economic Indexes for Areas (17), included in
570 multivariate models as quintiles. Atopic family history defined as any 1st degree relative eczema, asthma, hay fever. Data
571 are n/N (%) unless otherwise specified. n indicates number of participants with complete data.

572 Table 2 – Association between timing of 1st dose of DTaP, food allergy and atopic sensitisation

1 st dose DTaP		OR (95% CI)	P	n	aOR (95% CI)	p
	Food allergy					
All children						
< 1 month late	463/4041 (11%)					
≥ 1 month late	9/103 (9%)	0.74 (0.37-1.48)	0.39	3846	0.77 (0.36-1.62)	0.49
Females						
< 1 month late	206/1998 (10%)					
≥ 1 month late	1/41 (2%)	0.22 (0.03-1.59)	0.14		0.23 (0.03-1.73)	0.15
Males						
< 1 month late	256/2027 (13%)					
≥ 1 month late	8/62 (13%)	1.02 (0.48-2.18)	0.95		1.15 (0.50-2.63)	0.74
P for same effect in males and females			0.15			0.15

	Atopic sensitisation					
All children						
< 1 month late	734/4175 (18%)					
≥ 1 month late	13/107 (12%)	0.65 (0.36-1.16)	0.15	3971	0.66 (0.35-1.24)	0.19
Females						
< 1 month late	348/2070 (17%)					
≥ 1 month late	2/44 (5%)	0.24 (0.06-0.98)	0.05		0.25 (0.06-1.04)	0.06
Males						
< 1 month late	385/2087 (18%)					
≥ 1 month late	11/63 (17%)	0.94 (0.48-1.81)	0.84		1.00 (0.49-2.05)	0.99
P for same effect in males and females			0.09			0.09

573

574 DTaP=diphtheria-tetanus-acellular pertussis vaccination; OR=univariate odds ratio; aOR=odds ratio adjusted for sex, SEIFA
575 quintile, siblings, antibiotic use, child-care attendance, smokers at the home, parent country of birth.. n indicates number
576 of participants included in multivariate analyses. Results in **bold** indicate p<0.05.

577

578

579 Table 3 – Association between timing of 1st dose of DTaP and secondary outcomes

	1 st dose DTaP		OR (95% CI)	p	n	aOR (95% CI)	p
	< 1 month late	≥ 1 month late					
Eczema	1127/3735 (30%)	19/99 (19%)	0.55 (0.33-0.91)	0.02	3550	0.57 (0.34-0.97)	0.04
Eczema medication	961/3838 (25%)	12/99 (12%)	0.41 (0.22-0.76)	0.004	3725	0.45 (0.24-0.83)	0.01
Bronchiolitis admission	112/4160 (3%)	4/105 (4%)	1.43 (0.52-3.96)	0.49	4038	1.17 (0.41-3.33)	0.77
Wheeze ever	677/3786 (18%)	20/100 (20%)	1.15 (0.70-1.89)	0.59	3691	1.16 (0.69-1.95)	0.57

580

581 DTaP=diphtheria-tetanus-acellular pertussis vaccination; OR=univariate odds ratio; aOR=odds ratio adjusted for sex, SEIFA
582 quintile, siblings, antibiotic use, childcare attendance, smokers at the home, and parent country of birth. n indicates
583 number of participants included in multivariate analyses. Results in **bold** indicate p<0.05.

