

Title	Advanced glycation end product intake during pregnancy and offspring allergy outcomes: prospective cohort study
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Publication date	2021-10-05
Original Citation	Venter, C., Pickett , K., Starling, A., Maslin, K., Smith, P. K., Palumbo, M. P., O'Mahony, L., Ben Abdallah, M. and Dabelea, D. (2021) 'Advanced glycation end product intake during pregnancy and offspring allergy outcomes: prospective cohort study', Clinical and Experimental Allergy. doi: 10.1111/cea.14027
Type of publication	Article (peer-reviewed)
Link to publisher's version	10.1111/cea.14027
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Download date	2025-01-10 10:52:25
ltem downloaded from	https://hdl.handle.net/10468/12081



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Article type : Original Article

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Advanced glycation end product intake during pregnancy and offspring allergy outcomes: prospective cohort study Running title: Maternal AGEs intake and offspring allergies

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This article has been accepted for publication and undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the <u>Version of Record</u>. Please cite this article as <u>doi:</u> <u>10.1111/CEA.14027</u>

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#### **Declarations of interest**

Dr. Venter reports grants from Reckitt Benckiser, grants from Food Allergy Research and Education, grants from National Peanut Board, during the conduct of the study; personal fees from Reckitt Benckiser, personal fees from Nestle Nutrition Institute, personal fees from Danone, personal fees from Abbott Nutrition, personal fees from Else Nutrition, and personal fees from Before Brands, outside the submitted work. Dr Smith reports personal fees from the Nestle Nutrition Institute and speaker fee from Danone and Bayer outside of the submitted work.

Dr. O'Mahony reports personal fees from Alimentary Health, grants from GSK, outside the submitted work. The other authors declare no interests.

#### Author contributions:

CV initiated the paper and drafted the first version of the paper. KP performed the data analysis with guidance and review by MP. AS guided on the epidemiological aspects of the study. PS and LOM advised on interpretation of AGEs intake and cytokine data. MAB extracted the allergy data from the electronic medical records. KM assisted with interpretation of the dietary and AGEs intake. DD is the principal investigator and assessed and guided the data analysis plan. All authors reviewed and commented on various drafts of the paper.

### Informed consent

Informed consent was obtained from all subjects involved in the study.

#### Institutional review board

The Healthy Start study protocol was approved by the Colorado Multiple Institutional Review Board. (IRB number: 09-0563; Healthy Start 1; 2009-2014 and Healthy Start 2; 2015-present).

#### Data availability statement

Data available on request from the authors from https://coloradosph.cuanschutz.edu/research-andpractice/centers-programs/lead.

### Funding/ Acknowledgements:

This work was supported by the of Health, grant numbers: R01 DK076648/DK/NIDDK NIH HHS/United States, R01 GM121081/GM/NIGMS NIH HHS/United States, UG3 OD023248/OD/NIH HHS/United States, UH3 OD023248/OD/NIH HHS/United States, R25GM111901-S1, R25GM11190, NIH grant R00ES025817

1 Abstract

#### 2 Background

Associations have been shown between concurrent assessment of dietary intake of AGEs and childhood allergic outcomes. We examined the association between maternal AGEs intake and development of offspring asthma, wheeze, atopic dermatitis, allergic rhinitis, and food allergies, and sought to determine whether intake of AGEs was associated with cord sera cytokines/chemokines.

#### 7 Methods

8 Pregnant women  $\ge$  16 years were recruited in the Healthy Start study, a prospective pre-birth cohort 9 from Colorado (N =1410). The analysis included 962 dyads with adequate diet ( $\ge$ 2 recalls) and 10 allergy outcome details. AGEs intake was estimated for each mother by matching intakes reported 11 using 24-hour dietary recalls during pregnancy to a reference database of commonly consumed 12 foods' AGEs values. Child diagnoses of asthma and allergies up to 8 years were obtained from 13 electronic medical records. Cord sera cytokines and chemokines were analyzed in a subset (N = 14 462) of children.

#### 15 **Results**

The median [IQR] AGEs intake for the overall sample was 11919 kU/day [8293, 16573]. Unadjusted analysis showed a positive association between maternal AGEs intake in pregnancy and rhinitis up to 8 years of age (HR = 1.03; 95% CI: 1.01, 1.06), but the association was attenuated and no longer significant in adjusted models (HR = 1.01; 95% CI: 0.98, 1.04). Both adjusted and unadjusted models showed no associations between AGEs intake in pregnancy and any of the other outcomes (p>0.05). There were no significant associations between any cytokine or chemokine measured and AGEs intake or any of the outcomes studied (p>0.05).

#### 23 Conclusion

The study showed that maternal AGEs intake was not associated with offspring asthma and allergy outcomes. AGEs exposure during pregnancy may not have the same impact on child development to postnatal exposure.

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#### 29 Key messages

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- AGEs exposure during postnatal life has been associated with multiple adverse health outcomes.
  - Maternal intake of AGEs during pregnancy was not associated with offspring allergies.
  - Maternal intake of AGEs during pregnancy was not associated with cord blood cytokine or chemokine levels.

#### 37 Introduction

Allergic diseases are an increasing public health concern.<sup>1,2</sup> The four major presentations of allergic diseases include asthma, atopic dermatitis, allergic rhinitis, and food allergies. Atopic dermatitis is usually the first manifestation of allergic diseases, followed by food allergies, asthma and allergic rhinitis, a process of allergic disease development referred to as the atopic march.<sup>3</sup> Although allergic diseases share a common immunological profile involving T-helper 2 (Th2) and inflammatory cells and their cytokines and chemokines, each condition may also appear separately with its own individual gene/environment interactions.<sup>4</sup>

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46 Early life factors, including maternal diet during pregnancy, have been studied as a possible risk 47 factor for offspring allergic diseases. However, despite the large number of studies focusing on 48 maternal dietary intake in pregnancy and offspring allergic outcomes, the results do not give clear 49 guidance on which dietary factors to address.<sup>5,6</sup> The European Academy of Allergy and Clinical 50 Immunology also concludes that the role of the maternal diet in the development of offspring allergy 51 outcomes is unclear.<sup>7</sup> Much more progress has been made in terms of the infant diet and allergy 52 prevention, with most international allergy prevention guidelines advising against delaying 53 introduction of food allergens, particularly peanut and egg. 8,7,9,10

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Observational data indicates that increased intake in childhood of foods typical of the Western diet, such as burgers, sugar and high fructose corn syrup,<sup>11</sup> and food preparations, such as frying, may be associated with the increase in allergic diseases.<sup>12</sup> Advanced glycation end products (AGEs) are compounds formed when sugar binds to protein or it is formed via lipoxidation.<sup>11</sup> It is postulated that the Western diet is high in AGEs and via its effect on both Th2 and inflammatory cytokines, may contribute to the increase in allergic outcomes. This is also referred to as the false alarm hypothesis.<sup>12</sup>

Inflammatory cytokines and chemokines include, but are not limited to IL-1, IL-6, IL-8, IL-4, IL-5, IL-13, alarmins (IL-25, IL-33 and TSLP), and TNF- $\alpha$ .<sup>13,14</sup> The Receptor for Advanced Glycation End Products (RAGE) is activated by ligands, including endogenous pathogen associated molecular patterns (PAMPs), and also dietary derived AGEs, which are high in the western diet. The AGEs receptor (RAGE) is linked to up-regulation of TNF- $\alpha$ , IL-1, IL-6 and IL-8.<sup>13</sup> Induction of alarmins (IL25, IL33, TSLP) and the importance of this is reinforced by models of atopic dermatitis and allergic

68 asthma that show RAGE and its activation ligands are central to the development of sensitization 69 and allergic responses.<sup>12,15,16</sup> In addition, higher levels of soluble RAGE (a decoy ligand) is protective 70 against asthma.<sup>17</sup> Further inflammatory mechanisms related to AGEs products are: 1) induction 71 of glycation of intracellular proteins including transcription factors, 2) alteration of the pericellular 72 matrix, resulting in signaling changes and cellular dysfunction, 3) oxidative stress and mitochondrial 73 dysfunction, 4) methylglyoxal – an archetypal AGE binds to lysine on DNA and induces 74 oxidative/nitrogen induced damage and DNA cleavage. Each of the fore-mentioned mechanisms 75 may have more impact on a developing fetus than a child/adult and is worthy of investigation in the 76 context of development of atopic disease. In animal models of food allergy, resveratrol, a well-77 defined anti-oxidant, reduces development of ovalbumin allergy in a cholera-toxin adjuvant model 78 with both reduced sensitization of ovalbumin and a reduction in dendritic cell activation.<sup>18</sup> Sov 79 isoflavones, daidzien and genistein, have been shown to suppress allergic reactions in a murine 80 model of peanut allergy.<sup>19</sup> Furthermore, in-vitro studies of resveratrol have shown a reduction of 81 AGEs induced dendritic cell (DC) maturation, decreased dendritic cell RAGE receptor activation in 82 response to AGE-albumin, and reduced dendritic cell activation to AGE-albumin stimulated DC 83 (cytokines, co-stimulatory cells, mitogen activated protein kinases and NF-kB).<sup>20</sup>

84 Based on the underlying inflammatory processes in allergic conditions, particularly asthma and 85 wheeze, we previously studied the association between the maternal diet and asthma/wheeze in the 86 child. We concluded that the inflammatory profile of the maternal diet was not associated with 87 cytokine and chemokine levels at birth. However, the results suggested that a maternal diet that 88 scored higher on the dietary inflammatory index (DII) was associated with increased odds of 89 offspring asthma and/or wheeze by age 4 years.<sup>21</sup> The DII is a complex index that provides an 90 estimate of the inflammatory potential of the diet based on the inflammatory potential of foods and 91 nutrients in single index.<sup>22</sup> AGEs are proteins or lipids that become glycated after exposure to sugars 92 during heating and is considered to be one aspect of an inflammatory diet.<sup>11</sup>

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In our pursuit to find the specific dietary factors in pregnancy that are associated with offspring outcomes, we investigated the role of maternal dietary AGEs intake and offspring asthma, wheeze, atopic dermatitis, allergic rhinitis and food allergy. The primary aim of this study was to assess the associations between maternal AGEs intake during pregnancy and offspring diagnosis of asthma, wheeze, allergic rhinitis, atopic dermatitis, and food allergy up to 8 years. We hypothesized that 99 increased maternal AGEs intake during pregnancy would be associated with an increased rate of 100 development of these outcomes in offspring. The secondary aim of this study was to examine 101 associations between maternal AGEs intake and cord sera levels of cytokines and chemokines. We 102 hypothesized that maternal AGEs intake would be associated with cord sera cytokines and 103 chemokines, and that the effect might be exacerbated by maternal obesity, as both AGEs and 104 obesity are related to inflammatory processes.<sup>12,23</sup>

105

#### 106 Methods

#### 107 Study sample

108 This analysis included data from a longitudinal pre-birth cohort of 1410 mother-child dyads. Pregnant 109 women aged 16 years or older with singleton pregnancies were recruited from obstetrics clinics at 110 the local hospital from 2009 to 2014. The Healthy Start study protocol was approved by the Colorado 111 Multiple Institutional Review Board (IRB number: 09-0563; Healthy Start 1; 2009-2014 and Healthy 112 Start 2: 2015-present) and was registered as an observational study 113 at clinicaltrials gov as NCT02273297. Further details regarding the study have been published 114 elsewhere.23

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At recruitment, the cohort included 1410 mother-child dyads. Following written consent, pregnant women completed questionnaires on medical history at enrollment in early pregnancy and were asked to give consent for the review of offspring electronic medical records up to age 4 years. Additional consent for review of offspring electronic medical records up to age 8 was requested when mothers and children came in for a follow-up visit after age 4.

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122 Participants were excluded from this analysis if they did not provide consent for child medical record 123 review from birth up to 4 years of age (n=9), had offspring who died prior to birth (n=6), or who had 124 insufficient data available to search for the child in the electronic medical records system (n=66). 125 Among the 1329 participants eligible for inclusion in the electronic medical record search, 68 children 126 had no records in the electronic medical record system, resulting in a total of 1261 participants with 127 allergy outcome information. Mothers were asked to complete 1 dietary recall per month during 128 pregnancy. The analytic cohort included 962 mother-offspring dyads that completed  $\geq 2$  dietary 129 recalls over the course of pregnancy and had valid offspring allergy outcome information. For the immunological analysis, data from N=462 mother-offspring dyads with data on cord sera cytokineand chemokine levels were used (Supplemental Figure 1).

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#### 133 Maternal daily AGEs intake

134 Maternal dietary intake was measured 2-8 times throughout pregnancy, from the second trimester 135 onward, using the Automated Self-Administered 24-hour dietary recall (ASA24).<sup>24</sup> To estimate AGEs 136 intake from the reported dietary intake, a reference database of commonly consumed foods' AGEs 137 values (in kU/100g) built by Uribarri et al.<sup>11</sup> was used. Food descriptions from the ASA24 diaries 138 were matched to the AGEs database by name and cooking process where possible. Foods without 139 simple matches as well as complex multi-ingredient foods were decomposed into gram equivalents 140for 23 food components specified in My Pyramid,<sup>243</sup> e.g. 1 cup of broccoli = 1 dark-green vegetable 141 and 1 English muffin = 1 grain equivalent. The gram equivalents for each code were estimated from 142 My Pyramid by decomposing food codes using the USDA food and nutrient database for dietary 143 studies 3.0 ingredient list.<sup>256</sup> Average AGEs (kU/g) values were then created from the Uribarri et al.<sup>11</sup> 144 AGEs list for each of the 23 components based on general cooking type (fried, baked, stewed, or 145 fresh), as cooking method is the main determinant of AGEs scores. Components for discretionary 146 fats and sugars were excluded from this step due to difficulties with calculating accurate AGEs 147 scores. Finally, daily dietary AGEs intake was calculated (in kU) based on quantity of the food/food 148 component consumed. Median daily AGEs intake (kU/day) was calculated for each individual based 149 on total number of dietary recalls available due to the right skewed nature of the AGEs distribution. 150 Median daily AGEs were also standardized to amount per daily caloric intake (kU/1000kcal/day) to 151 compare to previous literature.

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#### 153 Child allergy outcomes

154 The outcomes of interest for the present paper were child diagnosis of allergic diseases (asthma, 155 wheeze, allergic rhinitis, atopic dermatitis, food allergy) up to 8 years of age based on data obtained 156 from the electronic medical records. "Any outcome" was defined as diagnosis of one or more of the 157 following diseases: asthma, wheeze, allergic rhinitis, atopic dermatitis, food allergy. Children's 158 medical record data was abstracted for participants who consented to child medical record review 159 and whose records were available in the Epic medical records system, as described previously.<sup>23</sup> In 160 short, the following search terms were used: 1) allergic rhinitis: "allergic rhinitis", "allergic 161 rhinoconjunctivitis", "hay fever", "rhinitis", "seasonal allergies"; 2) asthma: "asthma"; 3) atopic dermatitis/eczema: "atopic dermatitis", "eczema"; 4) wheeze: "wheeze." Search terms related to IgEmediated food allergies and food allergens included: "food allergy", "almond", "cashew", "clam", "crab", "egg", "fin fish", "fish", "milk", "pecan", "peanut", "salmon", "sesame seed", "scallop", "shellfish", "shrimp", "soy", "sunflower seed", "tree nut", "tuna", "wheat". The electronic medical records were reviewed by two clinician researchers, who assigned diagnoses for asthma, wheeze, allergic rhinitis, atopic dermatitis/eczema, and food allergies, after extensive review of medical notes.

For any participant with suspected food allergy, we recorded whether the child had IgE-mediated food allergy, the age of diagnosis, and the age at development of tolerance to the food if tolerance had been developed. Children with Eosinophilic Esophagitis (EoE) or Food Protein induced enterocolitis (FPIES) were not considered to have IgE-mediated food allergies unless they also had co-existing IgE-mediated food allergies.

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#### 174 Cord blood collection

Cord blood samples were obtained at delivery, stored on ice for up to 20 minutes, and processed by
centrifugation. Serum aliquots were stored at 4°C for up to 24 hours before being transported (on ice)
to an 80°C freezer for long-term storage. <sup>267</sup>

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#### 179 Cord sera cytokines and chemokines

180 Under a separate ancillary study (NIH: R00ES025817) focusing on sera analysis from maternal-181 offspring dyads with available sera and cord blood, stored frozen umbilical cord sera extracted at 182 birth was analyzed for a range of cytokines and chemokines. Cord blood inflammatory biomarkers 183 were processed by the University of Colorado Cancer Center Flow Cytometry Shared Resource. 184 Plasma cytokine/chemokine concentrations were determined by multiplex panel immunoassay 185 according to manufacturer's instructions (EMD Millipore Corporation, Billerica, MA 01821). Cytokines 186 including IL-1 $\beta$ , IL-4, IL-6, IL-10, and TNF- $\alpha$ ; and chemokines IL-8 and MCP-1, were measured in 187 units of pg/mL. For each analysis, samples were run in duplicate, and the percent coefficient of 188 variation was computed as a quality control measure. If both replicates were out of range, the value 189 was designated as being below (or above) the limit of detection. Values that were below the lower 190 limit of detection were marked as "out-of-range low" and values that were above the upper limit of 191 detection were marked as "out-of-range high".<sup>21</sup> The cytokines and chemokines studied were 192 previously selected to be analyzed for an NIH grant (R00ES025817) and included those that were 193 related to either air pollution exposure or pregnancy/birth outcomes.

194

#### 195 Covariate data

196 Data regarding maternal race/ethnicity, parity, maternal history of allergy (asthma and/or allergic 197 rhinitis), and age of introduction of solid foods were obtained through self-reported questionnaires. 198 Maternal history of asthma was assessed using the following question, "Has a health professional 199 such as a doctor, physician assistant, or nurse practitioner ever told you that you have asthma?" 200 Maternal history of allergic rhinitis was assessed using the following guestion, "Has a health 201 professional such as a doctor, physician assistant, or nurse practitioner ever told you that you have 202 hay fever, seasonal allergies or allergic rhinitis?" Mothers were considered to have a history of 203 allergy if they answered yes to one or both of the preceding questions. Pre-pregnancy weight was 204 obtained from either medical records or self-reported early in pregnancy. Maternal height was 205 measured at the first research visit via stadiometer. Pre-pregnancy body mass index (BMI) was 206 calculated using pre-pregnancy weight (kg) divided by height (m) squared. Pre-pregnancy BMI was 207 categorized as follows: lean (BMI <25 kg/m<sup>2</sup>), overweight (BMI 25-29.99 kg/m<sup>2</sup>), and obese (BMI  $\geq$ 30 208 kg/m<sup>2</sup>).<sup>24</sup> Observed gestational weight gain was calculated as the difference between the last 209 available weight recorded during pregnancy and the pre-pregnancy weight.<sup>278</sup> Gestational weight 210 gain was categorized as less than recommended, within the recommended range, or more than 211 recommended (excessive weight gain) based on pre-pregnancy BMI categories, as described by the 212 2009 Institute of Medicine (IOM) guidelines.<sup>289</sup> Information on total caloric intake during pregnancy 213 (kcal/day) was obtained using repeated 24-hour recalls, as described above. Breastfeeding duration 214 was computed as breastmilk months, a product of breastfeeding duration and intensity, using feeding 215 information reported by mothers at the 18 months postnatal interview. For exclusively breastfed 216 infants, breastmilk months is equivalent to the duration of breastfeeding (e.g., 8 months of 217 breastfeeding = 8 breastmilk months). For infants fed both breastmilk and formula, breastmilk months 218 is the duration of exclusive breastfeeding plus the weighted duration of mixed feeding (e.g., 4 months 219 of exclusive breastfeeding + 2 months of 50% breastmilk and 50% formula = 5 breastmilk months.) 220 For infants fed formula exclusively, breastmilk months is 0.<sup>30</sup>

221

#### 222 Statistical Analysis

223 Descriptive statistics were calculated for maternal and offspring characteristics, including means and 224 standard deviations or median and interquartile range (IQR) for continuous variables dependent on 225 distribution. Categorical variables were presented as frequencies and percentages. Demographic

226 variables were compared between those with or without available cytokine data using statistical 227 tests, including t-tests for normally distributed continuous variables, Wilcoxon rank sum tests for non-228 normally distributed continuous variables, and chi-square tests for categorical variables. We 229 computed cumulative incidence of the medical record verified respiratory and allergy outcomes up to 2301 year, 2 years, 3 years, and 4 years of age to describe the burden of these diseases in the cohort. 231 Hypothesis testing was conducted at an alpha level of 0.05. In the final models, we reported, where 232 appropriate, beta estimates, hazard ratios, 95% confidence intervals, and p-values and for the 233 associations.

234

Associations between maternal AGEs intake during pregnancy and offspring diagnosis of asthma,
 wheeze, allergic rhinitis, atopic dermatitis, and food allergy

237 Cox proportional hazards models were fit to examine the associations between maternal daily AGEs 238 intake (kU) and development of child allergic diseases (asthma, atopic dermatitis/eczema, food 239 allergy, allergic rhinitis, wheeze, and any allergy) up to age 8 years. A literature review<sup>29</sup> identified a 240 set of covariates used in previous publications seeking to identify associations between maternal diet 241 during pregnancy and offspring allergy and respiratory outcomes. Two models were performed for 242 the outcome: a base adjusted model 1, and an expanded adjusted model 2. Adjusted Model 1 was 243 the base model with factors most commonly associated with allergy outcomes<sup>2,30</sup> and included child 244 race/ethnicity, child sex, nulliparity, gestational smoking, and energy intake during pregnancy. 245 Adjusted Model 2 was based on factors previously included when studying maternal diet and 246 childhood allergy outcomes.<sup>29</sup> These included all variables in model 1, and additionally: mode of 247 delivery, maternal history of allergy, breastfeeding duration, age of introduction of solid foods, 248 maternal pre-pregnancy BMI category (Supplemental Figure 2). We constructed a directed acyclic 249 graph to represent hypothesized causal relationships and confirmed that none of the covariates were 250 either intermediates or colliders. To test the hypothesis that the effect of maternal AGEs intake on 251 allergy onset may be exacerbated by maternal obesity and excessive gestational weight gain, we 252 included interaction terms between maternal AGEs intake with pre-pregnancy BMI category and IOM 253 qestational weight gain category into Adjusted model 2. Non-significant interactions (p > 0.05) were 254 removed from the final adjusted models. The follow-up age was different for each child. This 255 occurred for one of two reasons: 1) consent was only given to review child medical records from birth 256 up to age 4 years, but not from age 4 to 8 years; or 2) consent was given to review child medical 257 records from birth up to age 8 years, but the child had not yet reached 8 years of age at the time of the electronic medical record search (e.g. the child was only 6 years old when the search was conducted). The Cox proportional hazards modeling approach allowed us to censor participants at the latest follow-up age for which they had available electronic medical record data, due to one of the two reasons described. For children with multiple recorded diagnoses of a disease outcome, their age at their first diagnosis was used. We checked that the assumption of proportional hazards was met for the "any outcome" prior to interpreting the results of these models. A sensitivity analysis was also performed dividing AGEs intake into quartiles to test for non-linear associations.

Associations between maternal AGEs intake and cord sera levels of cytokines and chemokines.

266 To examine the associations between maternal intake of AGEs and cytokines, two separate 267 modeling strategies were used depending on the detection rate of the values. For each of the 268 cytokines and chemokines, we determined the number and percentage of values that were outside 269 the detection range. In addition, we calculated the median and IQR for each of the cytokines and 270 chemokines. Cord sera cytokines and chemokines with <20% of values outside the detection range 271 were treated as continuous variables (IL-6, IL-8, TNF- $\alpha$ , MCP-1).<sup>1</sup> When the cytokines or chemokines 272 were treated as continuous variables, values below the limit of detection were assigned values equal 273 to half the lowest value observed on the standard curve<sup>28 29</sup> and values above the limit of detection 274 were assigned values equal to 1.5 times the highest value on the standard curve. Cord sera 275 cytokines and chemokines with ≥20% of values outside the detection range were treated as 276 categorical variables and dichotomized as detectable or not detectable (IL-1 $\beta$ , IL-4).

277 For chemokines with <20% of values outside of the detection range, separate general linear 278 univariate models were fit. Prior to fitting models for each of the cytokines and chemokines, the 279 values of these cytokines and chemokines (IL-6, IL-8, TNF- $\alpha$ , MCP-1) were natural log transformed 280to account for the positively skewed distributions . For cytokines with ≥20% values outside of the 281 detection range (IL-1 $\beta$ , IL-4), separate logistic regression models were fit to estimate the odds that 282 the value of the cytokine or chemokine was detectable. For both cytokines and chemokines with 283 <20% or <20% of values outside the detection range, unadjusted models included maternal intake of 284 AGEs as the only predictor. Adjusted models were fit for each cytokine/chemokine, with the 285 covariates and hypothesized interactions tested determined a priori. The adjusted models for the 286 inflammatory cytokines (IL-1 $\beta$ , IL-6, TNF- $\alpha$ ) and chemokines (IL-8, MPC-1) included nulliparity, child 287 race/ethnicity, child sex, pre-pregnancy BMI category, IOM gestational weight gain category, and 288 total caloric intake (kcal/day) as covariates (Supplemental Figure 3). The adjusted models for the T-289 regulatory cytokine (IL-10) and Th2 cytokine (IL-4) included the same covariates as were included for

the inflammatory cytokines, in addition to maternal history of allergy. To test the hypothesis that the effect of maternal AGEs intake on inflammatory cytokines and chemokines may be exacerbated by maternal obesity and excessive gestational weight gain, we included interaction terms between maternal AGEs intake with pre-pregnancy BMI category and with IOM gestational weight gain category. Non-significant interactions (p > 0.05) were removed from the final adjusted models. A sensitivity analysis was again performed dividing AGEs intake into quartiles to test for non-linear associations.

297

#### 298 Results

#### 299 Descriptive statistics

300 Table 1 reports descriptive statistics for maternal and offspring characteristics of the overall sample 301 of Healthy Start participants who had both maternal AGEs intake and offspring electronic medical 302 record data. Table 1 also compares maternal and offspring characteristics between the subset of 303 participants with cytokine/chemokine data and the subset without cytokine/chemokine data. There 304 were no significant demographic differences in the cohort included in analysis (N = 962, 305 Supplemental Figure 1) and those excluded (data not shown). The median [IQR] AGEs intake for the 306 overall sample was 11919 kU/day [8293, 16573]. Accounting for daily caloric intake, the median 307 [IQR] AGEs intake was 6485.0 kU/1000kcal [4669.6, 8546.6]. Birthweight (g) was statistically, but not 308 clinically, significantly higher in participants with cytokine and chemokine data compared to 309 participants without cytokine and chemokine data (3290±433 vs. 3196±541, p=0.002), as was 310 gestational age at birth (weeks) (39.48±1.24 vs. 39.21±1.95, p=0.01). Maternal age (years) at 311 delivery was also statistically, while not substantially, different at 28.25±6.03 for those with cytokine 312 and chemokine data compared to 29.17±5.84 for those without (p=0.02). AGEs intake, total caloric 313 intake, breastfeeding duration, gestational age, age of introduction of solids, nulliparity, gestational 314 smoking, ethnicity, pre-pregnancy BMI, IOM gestational weight gain, maternal history of allergy, and 315 child sex were not significantly different between those with and without cord sera cytokine and 316 chemokine data. Supplemental Table 1 shows cumulative incidence of allergy outcomes in the 317 offspring. There were no statistically significant differences between those participants with or without 318 cytokine data for any of the outcomes studied: any outcome, wheeze, asthma, atopic dermatitis, 319 allergic rhinitis, and food allergy (p >0.05, data not shown).

320 321

Associations between maternal AGEs intake and offspring diagnosis of allergic outcomes

322 Unadjusted analysis showed a positive association between a 1000kU/day increase in maternal 323 AGEs intake in pregnancy and development of allergic rhinitis up to 8 years of age (HR = 1.034; 95% 324 CI:1.012, 1.057), but the association was attenuated and no longer significant in adjusted model 1 325 (HR = 1.01 (95% CI: 0.986, 1.043). In the base adjusted analysis for any outcome, asthma, atopic 326 dermatitis, food allergy, and wheeze, there were no statistically significant associations between 327 AGEs scores and childhood allergy outcomes up to 8 years of age, with all Hazard Ratio values 328 nearing 1 (p>0.05, Table 2). For these allergic outcomes, fully adjusted regression models gave 329 similar results (Figure 1). There was also no evidence of a non-linear relationship between AGEs 330 intake and any outcome (p > 0.05, Supplemental Table 4). There were no significant interactions 331 between AGEs and BMI category (p > 0.05) for any of outcomes except for allergic rhinitis (p = 0.01) 332 and wheeze (p = 0.02), and thus interactions are not reported in the adjusted model 2 in Table 2. For 333 allergic rhinitis, women with a pre-pregnancy BMI categorized as "lean" were the only group with a 334 significant AGEs effect with a hazard ratio of 1.054 for a 1000kU/day increase in AGEs (95% CI: 335 1.014, 1.095; p = 0.008). For wheeze, only women with a pre-pregnancy BMI categorized as "obese" 336 had a significant AGEs effect with a hazard ratio of 0.934 for a 1000kU/day increase in AGEs (95% 337 CI: 0.883, 0.989; p = 0.02) (Supplemental Table 3).

338

339 Associations between maternal AGEs intake and cord sera cytokine/chemokine levels

Supplemental Table 2 reports the frequency and percentage of values outside of the detection range
for each cytokine/chemokine, and records whether the out-of-range values were low (below the lower
limit of detection) or high (above the upper limit of detection). The median and IQR for levels of each
cytokine/chemokine are also reported.

344

The hypothesized interaction between AGEs intake and pre-pregnancy BMI category for each of the cytokines/chemokines examined was non-significant (all p>0.05). After removing all non-significant interaction terms, the associations between continuous maternal AGEs scores and cord sera levels of the cytokines/chemokines remained non-significant in the final adjusted models (all p>0.05; Tables 3 and 4). There was also no evidence of a non-linear relationship between AGEs intake and any cord sera levels of the cytokines/chemokines (p > 0.05, Supplemental Table 5)

- 351
- 352 Discussion

353 Previous studies have indicated that AGEs intake may be implicated in non-communicable diseases 354 such as diabetes,<sup>31,32</sup> chronic kidney disease,<sup>33</sup> non-alcoholic fatty liver disease,<sup>34</sup> coronary heart 355 disease,32 obstructive airway diseases,<sup>35</sup> cancer,<sup>36</sup> dementia and aging<sup>37</sup>, arthritis.38 356 asthma/wheeze,<sup>39,40</sup> chronic bronchitis. <sup>41 37,42,43</sup> Studies focusing on allergy outcomes, showed an 357 association between the childhood diet, studied at a similar time as allergic outcomes.<sup>12,44,45</sup> In this 358 study, we examined the association between maternal AGEs intake during pregnancy and offspring 359 allergy outcomes. The unadjusted analysis showed a positive association between maternal AGEs 360 intake in pregnancy and offspring allergic rhinitis up to 8 years of age, but the association was no 361 longer significant in adjusted models. For all other offspring allergy outcomes, we did not find a 362 significant association with maternal AGEs intake, before or after adjusting for potential confounders. 363 For allergic rhinitis, a significant interaction between maternal pre-pregnancy BMI and AGEs intake 364 was observed, showing that the association between increased AGEs intake and "lean" based on 365 their pre-pregnancy BMI. A significant interaction between AGEs intake and pre-pregnancy BMI was 366 also observed for wheeze, indicating increased intake of AGEs was associated with reduced risk of 367 offspring wheeze only among women categorized as obese based on their pre-pregnancy BMI. 368 Immunological data from previous studies<sup>12,15,16</sup> indicates that intake of AGEs may be associated with 369 development of allergic disease via its effect on RAGE receptors, leading to a cascade of 370 inflammatory processes. In the study presented here, no significant associations between maternal 371 AGEs intake and cord sera levels any of the cytokines or chemokines studied were observed.

372

373 Studies examining the relationship between AGEs intake and allergy outcomes, have been 374 conducted in children, focusing on the child's diet and the child's concurrent allergic symptoms. The 375 International Study of Asthma and Allergies in Children (ISAAC) study reported that eating fast foods 376 2 3 times a week by adolescents was associated with asthma, rhino-conjunctivitis and eczema in 377 these study participants.<sup>45</sup> The authors hypothesized this this effect may be due to the AGEs content 378 of the food. Another prospective study reported on intake of AGEs in children and found that higher 379 fast food consumption rates in urban children in South Africa was associated with atopic dermatitis 380 than those with lower consumption rates. Urban children with high fried/microwaved meat 381 consumption also had higher rates of any allergy compared to those with lower intakes.<sup>44</sup> These 382 foods were once again used as a proxy for AGEs intake, rather than using a robust measurement of 383 AGEs content of the whole diet, as was done for the Healthy Start cohort in the study presented 384 here. More recently, a study from the US indicated that increased AGEs intake was significantly

associated with increased odds of wheezing, wheeze-disrupted sleep, and wheezing requiring
 prescription medication.<sup>40</sup>

387

388 Following on from the results of these studies,<sup>40,44,45</sup> we sought to test whether maternal AGEs intake 389 during pregnancy was associated with offspring allergic outcomes. In both the unadjusted and 390 adjusted models for any outcome, asthma, atopic dermatitis, and food allergy up to 8 years, we 391 found no statistically significant findings for the association between maternal AGEs intake and 392 offspring allergy outcomes. The primary reason we suspect may explain the discrepancy between 393 our study findings and the findings of the previous studies listed, is that previous studies examined 394 concurrent AGEs intake of the child in which allergic disease was also studied, whereas we 395 prospectively studied maternal intake of AGEs during pregnancy and examined allergy in offspring. 396 This may indicate that AGEs intake in pregnancy may not affect child allergy outcomes. The results 397 of our study indicated that among women with a pre-pregnancy BMI categorized as lean weight 398 women, increased AGEs intake during pregnancy was associated with increased risk of offspring 399 allergic rhinitis; and among women with a pre-pregnancy BMI categorized as obese, increased AGEs 400 intake was associated reduced risk of offspring wheeze. These findings are difficult to interpret and 401 further studies may be able to clarify these results.

402

403 The discrepancy in findings between our study and previous studies may also be explained by 404 different measures and methods used to determine AGEs intake, or different methods for 405 assessment of allergy outcomes. Previous studies investigating the association between AGEs and disease outcomes, used serum levels of AGEs,<sup>31,46</sup> or activity of the age receptor (RAGE)<sup>37,42,43</sup> 406 407 rather than dietary intake of AGEs as in our study. Some studies measured reported high-fructose 408 corn syrup sweetened soft drinks, fruit drinks<sup>38</sup> or apple juice, apple juice, fruit drinks and soda 409 intake.<sup>39,41</sup> It is postulated that intake high fructose corn syrup, leads to *in situ* formation of AGEs, 410 hence making these foods a suitable proxy for studying AGEs intake.<sup>47</sup>

411

412 Only three studies used dietary intake of AGEs measured by food frequency questionnaires to define 413 AGEs intake; one study defined AGEs intake by food group<sup>35</sup> and two studies quantified AGEs intake 414 based on FFQs measuring AGEs intake of commonly eaten foods.<sup>3337</sup> Our study is the first study to 415 our knowledge that investigated AGEs intake in pregnancy and its association with child health 416 outcomes. We have used detailed 24 hour recalls completed 2-8 times over the course of pregnancy to calculate average AGEs intake throughout pregnancy. The median AGEs intake of the women in the Healthy Start study was 6485.0 kU/1000 kcal. In our cohort, the median [IQR] AGEs intake for the overall sample was 11919 kU/day [8293, 16573]. There are no studies reporting AGEs intake in pregnant women, but a review paper by Nowotny et al.<sup>48</sup> indicates that AGEs intakes in adults can range from 6 000 – 27 000 kU/day, which indicates that our calculated AGES intake falls within this range.

423

424 No previous studies have explored the relationship between maternal AGEs intake during pregnancy 425 and cord blood cytokines and chemokines. We had expected to see that higher maternal AGEs 426 intake would be positively associated with proinflammatory cytokines and chemokines (IL-1β, IL-6, TNF- $\alpha$ , MCP-1, IL-8)<sup>49</sup> and negatively associated with anti-inflammatory cytokines such as IL-10.<sup>4</sup> In 427 428 particular, we expected to see some association between AGEs intake and TNF- $\alpha$  and MCP-1 levels. 429 TNF- $\alpha$  is a known pro-inflammatory cytokine and higher levels of TNF- $\alpha$  has been associated with 430 increased expression of RAGE.<sup>50</sup> MCP-1 is thought to play a role in cell-surface expression of 431 adhesion molecules<sup>51</sup> and RAGE is closely related to other genes coding for cell adhesion 432 molecules. However, we saw no significant associations between maternal AGEs intake and cord 433 sera cytokine or chemokine levels.

A strength of this study is that we present information from a well-characterized cohort with a large sample size, which enabled us to take various relevant covariates into account. We were able to study the association between a comprehensive and detailed measure of dietary AGEs intake during pregnancy, physician diagnosed recorded outcomes, and a range of cytokines and chemokines in cord sera. We are also the first research group to study the association between maternal AGEs and cord sera cytokine levels as a possible underlying mechanism for the development of offspring allergies.

We have used detailed 24 hour recalls completed 2-8 times over the course of pregnancy to calculate average AGEs intake throughout pregnancy. Approximately 76% of the participants completed  $\geq$ 2 dietary recalls over the pregnancy, with a median of 3 recalls. Two recalls can be representative of the entire pregnancy, given that dietary intake is relatively stable across pregnancy<sup>52</sup> and two or more recalls are sufficient to estimate usual dietary intake, per the National Cancer Institute Dietary Assessment Primer.<sup>53</sup> In addition, we also compared the estimated daily AGEs values for those with below the median number of intakes (<3) to those with 4 or more entries. The distributions were not statistically different with those with 3 or less entries (N = 556) having a median estimated daily AGEs intake of 12104 (IQR: 7400-17148) and those with 4+ entries (N = 406) with a median of 11824 (IQR: 8978-15577) (p = 0.49). We have used a robust method to calculate AGEs intake, but the AGEs content of many foods is unknown and calculation of the AGEs scores of composite foods is difficult to standardize. One particular limitation of the dietary AGEs calculation may be that discretionary fats and sugars were excluded due to difficulties with calculating accurate AGEs scores. These issues may partially explain the null associations.

455

Future studies may benefit from using an AGEs food frequency questionnaire, validated against reliable biomarkers, such as AGEs serum levels, and other forms of dietary intake to measure AGEs instead. This information will enable us to assess if current levels of foods are indeed correct, especially composite foods and provide us with validated measures of dietary intake. There are currently no studies reporting on validating measurement of AGEs intake by using biological samples such as blood<sup>12</sup> or skin.<sup>33,54</sup> Some covariate information was collected through self-report rather than objective measures, which might have influenced data quality.

463

464 Our results are limited as only one Th2 and one T-regulatory cytokine were measured. Measuring 465 cytokines in cord blood mononuclear cells, after sufficient stimulation with antigen, is preferable to 466 using frozen cord sera. Future studies should focus on measuring a wider range of cord sera 467 cytokines and chemokines, using antigen stimulation of cord blood mononuclear cells.

468

469 Other limitations included the potential bias associated with the self-report of dietary intake.55,56 In 470 addition, the sensitivity and specificity of using electronic medical record data for allergy outcomes is 471 not known, but this approach has been used in European cohort studies to report and validate on 472 asthma outcomes.<sup>57</sup> We also acknowledge that the mechanistic consequences of the AGE/RAGE 473 interaction could either impact on immune responses increasing a tendency to allergic sensitization 474 and/or enhancing susceptibility to inflammation independent of allergy. The latter may have been 475 impacted by pregnancy intake of AGEs and future studies should focus on separating "allergic" from 476 "inflammatory" outcomes. In particular, wheezing in infancy may not indicate an allergic phenotype, 477 though its inflammatory characteristics<sup>58</sup> have been well described. Most of the findings have wide 478 confidence intervals, highlighting the fact that further studies are required. 479

#### 480 Conclusion

481 In this study, we did not observe significant associations between maternal AGEs intake during 482 pregnancy, and offspring allergy outcomes or cord blood cytokines and chemokines. These findings 483 suggest that maternal AGEs intake during pregnancy may not impact development of allergic 484 diseases in offspring. Exposure to AGEs during pregnancy may not have the same impact on child 485 development as postnatal AGEs exposure.

486

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#### 651 Figure Legends/Text:

Figure 1. Hazard ratios with confidence intervals for the associations between continuous maternal
AGE (kU/day) intake and Offspring Allergy Outcomes up to 8 years; adjusted for maternal
race/ethnicity, child sex, nulliparity, maternal smoking prior to pregnancy, maternal history of allergy,
mode of delivery, breastfeeding duration, age of introduction of solid foods, maternal pre-pregnancy
BMI category, and energy intake during pregnancy (all models exclude interactions).

658 659

	Overall	Subset with	Subset without	
	sample	cytokine/chemokine	cytokine/chemokine	
		data	data	
Sample size (N)	962	462	500	
Continuous variables	Mean* ±	Mean* ± SD <sup>a</sup>	Mean* ± SDª	p-val
	SDª			
Maternal				
characteristics				
Advanced Glycation	11919	11921.2 [7875.5,	11918.7 [8648.2,	0.2
End Product Intake	[8293,	15998.7]	16890.9]	
(AGEs)	16573]			
(kU/day)(Median				
[IQR])				
AGEs (kU/1000kcal)	6485.0	6434.0 [4510.2,	6434.0 [4510.2,	0.1
(Median [IQR])	[4669.6,	8448.9]	8448.9]	
	8546.6]			
Age at delivery	28.7 ± 5.9	28.25 ± 6.03	29.17 ± 5.84	0.0
(years)				
Total caloric intake	2070 ±	2055.58 ± 378.81	2083.66 ± 399.48	0.2
(kcal/day)	390			
Breastfeeding	8.7 ± 6.7	8.28 ± 6.64	9.11 ± 6.71	0.0
duration (breastmilk				
months) <sup>34</sup>				
Nulliparous	475 (49%)	239 (52%)	236 (47%)	0.1
Smoking in	64 (7%)	32 (7%)	32 (6%)	3.0
pregnancy				
Pre-pregnancy body				0.2
mass index <sup>31</sup>				

Underweight	24 (2%)	14 (3%)	10 (2%)	
(<18.5 kg/m²)				
Lean (25 kg/m²)	510 (53%)	232 (50%)	278 (56%)	
Overweight (25-	245 (25%)	120 (26%)	125 (25%)	
29.99 kg/m²)				
Obese (≥30 kg/m²)	183 (19%)	96 (21%)	87 (17%)	
IOM <sup>d</sup> gestational				0.25
weight gain <sup>33</sup>				
Less than	213 (22%)	92 (20%)	121 (24%)	
recommended				
Within	286 (30%)	143 (31%)	143 (29%)	
recommended range				
More than	460 (48%)	227 (49%)	233 (47%)	
recommended				
Maternal history of	334 (35%)	160 (35%)	174 (35%)	1
allergy				
Offspring				
characteristics				
Race/ethnicity				0.94
Non-Hispanic	548 (57%)	262 (57%)	285 (57%)	
white				
Non-Hispanic	98 (10%)	50 (11%)	48 (10%)	
black				
Hispanic	213 (22%)	101 (22%)	112 (22%)	
Other <sup>c</sup>	104	49 (11%)	55 (11%)	
	(11%)			
Sex – female	503 (52%)	247 (53%)	256 (51%)	0.52
Birthweight (grams)	3242 ±	3290.4 ± 432.8	3196.3 ± 541.12	0.002
	595			
Gestational age at	39.3 ± 1.7	39.48 ± 1.24	39.21 ± 1.95	0.01
Hinth (weaks)				

i.	Age solid foods	6.1 ± 2.0	6.1 ± 1.9	$6.2 \pm 2.0$	0.12
	introduced (months)				

a) SD: Standard deviation

b) p-value for hypothesis test comparing demographic variables between those with or without any cytokine data. Statistical tests included t-tests for normally distributed continuous variables, Wilcoxon rank sum tests for non-normally distributed continuous variables, and chi-square tests for categorical variables.

c) Other race/ethnicity includes non-Hispanic Asian, American Indian/Alaska Native, Hawaiian/Pacific Islander, and multi-racial

d) IOM: Institute of Medicine

\*AGE intake was expressed as median [IQR] as a non-normally distributed variable.

 Table 2. Results of Cox proportional hazard models examining the association between

 maternal daily AGE intake and child allergy diagnosis

	Unadjusted Mode	el ((N = 9	Adjusted Model 1	± ((N = 9	S Adjusted Model 2 <sup>≠</sup> (N = 782)		
Outcome	HR*(95% CI)	p-value	HR*(95% CI)	p-value	HR*(95% CI)	p-value	
Any Allergy	1.007 (0.993, 1.02	0.32	0.995 (0.978, 1.01	0.56	0.997 (0.978, 1.0	0.74	
Asthma	1.014 (0.993, 1.03	0.19	1.009 (0.984, 1.03	0.48	1.008 (0.98, 1.03	0.57	
Atopic Dermatiti	1.003 (0.986, 1.02	0.70	0.993 (0.972, 1.01	0.49	0.999 (0.976, 1.02	0.93	
Food Allergy	1.012 (0.971, 1.05	0.56	0.978 (0.928, 1.03	0.41	0.975 (0.915, 1.03	0.44	
Rhinitis	1.034 (1.012,	0.002	1.014 (0.986, 1.04	0.34	1.012 (0.98, 1.04	0.58#	
	1.057)						
Wheeze	0.99 (0.969, 1.011	0.35	0.975 (0.951, 1.00	0.05	0.981 (0.955, 1.00	0.17#	

\* Hazard ratio's (HR) given for 1000kU increase in AGE value.

<sup>±</sup>adjusted for maternal race/ethnicity, child sex, nulliparity, maternal smoking prior to pregnancy and energy intake during pregnancy.

<sup>≠</sup>adjusted for maternal race/ethnicity, child sex, nulliparity, maternal smoking prior to pregnancy, maternal history of allergy, mode of

delivery, breastfeeding duration, age of introduction of solid foods, maternal pre-pregnancy BMI category, and energy intake during

pregnancy

#Significant interaction between AGE and BMI category observed. Results from models including interactions shown in

Supplemental Table 3.

662

663

Table 3. Associations between maternal AGE intake (kU) and natural log levels of cordsera cytokines and chemokines

	Unadjusted	(N= 462)		Adjusted <sup>+</sup> (N= 462)			
Outcome	β*	SE	p-value	β*	SE	p-value	
IL-6	-0.001	0.01	0.95	0.009	0.011	0.43	
IL-8	-0.01	0.008	0.18	-0.005	0.009	0.57	
TNF-α	-0.0003	0.003	0.91	0.004	0.004	0.27	
MCP-1	0.004	0.004	0.23	0.007	0.004	0.09	
IL-10	-0.007	0.006	0.22	-0.001	0.013	0.92	

\*The beta estimate represents the change in log-pg/mL of each outcome per each 1000kU increase in maternal AGE intake.

+ Adjusted for child race/ethnicity, child sex, nulliparity, energy intake during pregnancy, IOM gestational weight gain category, BMI Category. The model for IL-10 additionally adjusts for maternal history of allergy.

## Table 4. Associations between maternal AGE intake (kU) and the odds of the cord sera cytokine and chemokine value being detectable

	Unadjus	sted Model (N= 462	2)	Adjusted Model* (N= 462)			
Outcome	OR*	95% CI	p-value	OR*	95% CI	p-value	
IL-1β	0.992	(0.966, 1.019)	0.57	0.981	(0.949, 1.014)	0.26	
IL-4	0.998	(0.97, 1.026)	0.87	1.015	(0.98, 1.051)	0.41	

\* The odds ratio (OR) represents the change in the odds that the outcome is detectable per each 1000kU increase in maternal AGE intake.

+ Adjusted for child race/ethnicity, child sex, nulliparity, energy intake during pregnancy, IOM gestational weight gain category, BMI Category. The model for IL-4 additionally adjusts for maternal history of allergy.

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					-p ai		HR	95% CI	р
Any Allergy							0.99	(0.98, 1.02)	0.74
Asthma			<b></b>		•		1.01	(0.98, 1.04)	0.57
Atopic dermatitis			<b> </b>	•			1.00	(0.98, 1.02)	0.93
Food allergy			•				0.98	(0.92, 1.04)	0.44
Allergic rhinitis			<b> </b>		•		1.01	(0.98, 1.05)	0.58
Wheeze		<b> </b>	•		4		0.98	(0.96, 1.01)	0.17
0	92 0.94	4 0.96	0.98	1	1 02	1 04			
0.			4000111			-			
	Hazar	d ratio (per	1000kU	Increa	se in AG	E)			