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1 The Association of *S. aureus* colonization with Food Allergy Occurs Independent of 2 Eczema Severity

- 2 Eczema Severity
- 3

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37 Abbreviations

- 38 CI Confidence interval
- 39 LEAP Study Learning Early About Peanut Allergy Study
- 40 LEAP-On Study 12 month extension of LEAP Study: Persistence of Oral Tolerance to Peanut
- 41 OR Odds Ratio
- 42 SCORAD SCORing Atopic Dermatitis
- 43 S. aureus Staphylococcus aureus
- 44 SEB staphylococcal enterotoxin B

S. aureus and food allergy in LEAP/LEAP-On

- 45 SPT Skin prick test
- 46 slgE specific Immunoglobulin E
- 47
- 48

49 **Conflict of Interest disclosure statement:**

S. aureus and food allergy in LEAP/LEAP-On

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- 59 Abstract
- 60

61 **Background:** *S. aureus* has been implicated in the pathophysiology of eczema, allergic rhinitis, 62 asthma, and food allergy. *S. aureus* is a marker of more severe eczema which is a risk factor for 63 food sensitization/allergy. It may therefore be that the association between *S. aureus* and food 64 allergy in eczematous patients is related to eczema severity.

65

66 **Objective:** To investigate the association of *S. aureus* colonization with specific IgE (sIgE) 67 production to common food allergens and allergies in early childhood independent of eczema 68 severity. We additionally determined the association of *S. aureus* colonization with eczema 69 severity and persistence.

70

71 Methods: In LEAP participants, eczema severity was assessed and skin/nasal swabs cultured 72 for *S. aureus*. Sensitization was identified by sIgE. Peanut allergy was primarily determined by 73 oral food challenge and persistent egg allergy by skin prick test.

74

Results: Skin *S. aureus* colonization was significantly associated with eczema severity across LEAP while at 12 and 60 months of age it was related to subsequent eczema deterioration. Skin *S. aureus* colonization at any time-point was associated with increased levels of hen's egg white and peanut slgE, independent of eczema severity. Participants with *S. aureus* were more likely to have persistent egg allergy and peanut allergy at 60 and 72 months of age, independent of eczema severity. All but one of the 9 LEAP consumers who developed peanut allergy (9/312) were colonized at least once with *S. aureus*.

82

83 **Conclusion:** *S. aureus,* independent of eczema severity, is associated with food sensitization 84 and allergy and may impair tolerance to foods. This could be an important consideration in 85 future interventions aimed at inducing and maintaining tolerance to food allergens in 86 eczematous infants.

87

88

Clinical Implications:

91 There may be a role for *S. aureus* eradication in interventions aimed at inducing and 92 maintaining tolerance to foods in eczematous infants.

Capsule Summary:

S. aureus colonization, independent of eczema severity, is associated with hen's egg and 97 peanut sensitization and allergy. *S. aureus* colonization may impair tolerance to foods.

10 Keywords:

101 Food Sensitization. Food Allergy. Peanut Allergy. Egg allergy. Eczema. Atopic Dermatitis. S.

102 aureus. Prevention. LEAP. Microbiome

105 Main manuscript total word count: 4,325

109 **INTRODUCTION**

110 There are many studies that implicate Staphylococcus aureus (S. aureus) in the 111 pathophysiology of eczema and other atopic outcomes. Epicutaneous sensitization with 112 staphylococcal enterotoxin B (SEB) elicits local cutaneous inflammation consistent with eczema 113 in mice (1) and subjects with normal and atopic skin (2). Prospective population-based birth 114 cohorts report that skin (3) or nasal (4) colonization by S. aureus precedes the clinical diagnosis 115 of eczema in infancy. Patients with eczema are more likely to be colonized with S. aureus than 116 healthy controls and disease severity is associated with S. aureus colonization on the lesional 117 skin (5). Additionally, patients with allergic rhinitis are more frequently colonized with nasal S. 118 aureus (6, 7) or sensitized to S. aureus enterotoxins (8) than healthy controls, and those that 119 are S. aureus positive have more severe allergic rhinitis than the S. aureus negative (6, 7). 120 Furthermore, S. aureus enterotoxins trigger airway inflammation and increased airway 121 responsiveness (9) and SEB facilitates allergic sensitization in murine asthma models (10). 122 Clinically, nasal S. aureus or serum IgE to S. aureus toxins is associated with wheeze and/or 123 asthma in children and adults (11-13). Finally, the presence of S. aureus or IgE to S. aureus 124 toxins is related to asthma severity (12-14), poor asthma control (15) and higher prevalence of 125 aeroallergen sensitization (14). Therefore, there are indications that S. aureus is associated with 126 the development and/or severity of these atopic outcomes.

127

128 Interestingly, S. aureus colonization has also been associated with food sensitization and 129 allergy. Jones et al retrospectively analysed skin culture results from eczematous children, aged 130 0-18 years, and report that those with skin S. aureus had peanut, egg, and milk specific IgE 131 (slgE) levels that correlated to a greater than 95% positive predictive value of oral food 132 challenge reactions to the respective allergen (16). As eczema and eczema severity are risk 133 factors for food sensitization and allergy (17, 18) and S. aureus is a marker of more severe 134 eczema, it may be that the association between S. aureus and food allergy in patients with 135 eczema is related to eczema severity.

136

In the Learning Early About Peanut Allergy (LEAP) Study, we sequentially recorded eczema severity and tested for *S. aureus* colonization at 4 different time points in 640 children (19). This design provides a unique opportunity for the detailed investigation of the relationship between *S. aureus* and food allergy. In an exploratory secondary analysis, we aimed to investigate the association of *S. aureus* colonization with slgE production to common food allergens and food allergy in early childhood independent of eczema severity. In addition, we sought to determine the association of *S. aureus* colonization with eczema severity and persistence.

146 METHODS

147 Study population, design and procedures

148 This is a secondary analysis of LEAP and LEAP-On (20) outcomes that includes all participants 149 recruited to these studies. Full study details have been previously published (19, 20). The LEAP 150 Study enrolled infants aged ≥ 4 to <11 months with severe eczema and/or egg allergy. 151 Participants were randomly assigned to avoid (LEAP avoiders) or consume peanut (LEAP 152 consumers). Assessments were undertaken at baseline (age 4-11 months) visit and at age 12, 153 30 and 60 months. They included eczema clinical evaluation, acquisition and culture of skin and 154 nasal swabs, food allergen SPT and slgE as well as total IgE. The LEAP-On Study 155 assessments were undertaken at 72 months of age, after 12 months of peanut avoidance in 156 both groups. Concurrent and past medication use was recorded at all LEAP and LEAP-On study 157 visits.

158

159 Clinical assessment of eczema severity

160 Eczema was clinically evaluated by a pediatric allergist at baseline, and at 12, 30, 60 and 72 161 months of age; eczema severity was determined according to the SCORAD (SCORing Atopic 162 Dermatitis) index. Mild, moderate and severe eczema was defined as SCORAD values <15, 163 >15-40, and >40 respectively. Persistent eczema was defined as eczema where the severity did 164 not decrease over sequential time points.

165

166 Skin and nasal swabs and S. aureus assessment

167 Skin and nasal swabs were obtained at baseline, and at 12, 30, and 60 months of age. Samples 168 were taken using sterile, cotton tipped transport swabs suitable for isolating aerobes and 169 anaerobes. A skin swab was obtained from the most severe eczema lesion or - in the absence 170 of eczema - the knee flexure. If the skin was dry, a drop of sterile water was placed on the skin 171 prior to the swab being taken. The skin swab was then placed in medium. The nasal swab was 172 inserted into one anterior nostril, and was then slowly withdrawn with a rotating motion and 173 subsequently placed in medium (Amies Medium used for both samples). Swabs were incubated 174 overnight and plated directly onto Columbia Blood Agar, CLED or MacConkey Agar (aerobic 175 incubation) and Chocolate Agar (CO2). Sensitivity was reported using BSAC (British Society for 176 Antimicrobial Chemotherapy) or via BioMerieux analyser Vitek2.

177

178 SPTs, slgE and total IgE measurement

179 SPTs and allergen slgEs were conducted at baseline, 12, 30, 60 and 72 months of age. Total 180 IgE was measured at all visits except for 12 months. Test methodologies and SPT materials 181 have been published previously (19-21).

182

183 **Definitions of peanut allergy**

184 Peanut allergy was determined by means of an oral peanut challenge at 60 and 72 months (20,

185 21). At 72 months, the allergic status of participants for whom the results of the oral peanut 186 challenge were inconclusive or not available was determined as per the diagnostic algorithm

published previously (21).

- 187
- 188

189 **Definitions of egg allergy**

S. aureus and food allergy in LEAP/LEAP-On

At baseline, egg allergy was defined as an SPT \geq 6 mm to raw hen's egg white and no history of previous egg tolerance, or an SPT \geq 3 mm to pasteurized hen's egg white and allergic symptoms related to exposure to hen's egg. At 60 and 72 months of age we defined persistent egg allergy as SPT \geq 6mm to raw or pasteurized hen's egg in the participants diagnosed as egg allergic at baseline.

195

196 Statistical analysis

197 Statistical analyses were performed on all LEAP and LEAP-On Study participants for whom an 198 outcome measurement was obtained. No imputation for missing data was conducted. Two 199 separate repeated measures longitudinal models were used to assess if Skin or Nasal S. 200 aureus (independent variable) was associated with concurrent eczema severity as assessed by 201 SCORAD (dependent variable). Analogously another two separate repeated measures 202 longitudinal models were used to assess if Skin or Nasal S. aureus at the immediately 203 preceding visit was associated with eczema persistence. Average Peanut and Egg slgE levels 204 (dependent variables) were compared between those who ever had Skin S. aureus to those 205 who never had Skin S. aureus (independent variable) via longitudinal repeated measures 206 models (one for peanut and one for egg respectively) which also included a covariate for 207 SCOARD. All repeated measures longitudinal models utilized an unstructured covariance 208 structure to model the correlation among time points within each subject, treated time as 209 categorical and also included covariates for time and the interaction between time and S. 210 aureus colonization status. Bootstrap sampling of 1,000 replicates within each time point was 211 utilized to assess where (or if) a divergence existed in the relative distribution of IgE production 212 to Egg, Peanut and Milk slgEs and Total IgE comparing those who ever had Skin S. aureus to 213 those who never had Skin S. aureus. As peanut and egg allergy (independent variables) were 214 only assessed at 60 and 72 months, four (peanut allergy at 60 and 72 months, egg allergy at 60 215 and 72 months) separate logistic regression models were constructed for each S. aureus 216 colonization location (skin, nose, and combination of skin or nose - dependent variables). 217 These logistic regression models included covariates for SCORAD (collected at 60 or 72 218 months respectively), LEAP treatment assignment, and the interaction between LEAP treatment 219 assignment and S. aureus colonization status. As there were a small number of subjects with 220 peanut allergy and complete separation occurred, the Firth penalized likelihood method was 221 used only for the peanut allergy models. These were secondary analyses on study outcomes, 222 and no adjustments have been made for multiple comparisons. All analyses were performed at 223 the 0.05 level of significance using SAS software version 9.4 or JMP version 12. Datasets for 224 the analyses are available through TrialShare, a public Web site managed by the Immune 225 Tolerance Network (https://www.itntrialshare.org/LEAP JACI 2019.url)

- 226
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- 229 **RESULTS**
- 230

231 **Participants**

The characteristics of participants screened and enrolled in the LEAP and LEAP-On Studies have been previously published (19, 20).

234

Characteristics of *S. aureus* colonization in the LEAP Study with no differences noted in *S. aureus* colonization between intervention groups.

- 237 Approximately half (48.8%) of the participants had some form of S. aureus colonization (32.2% 238 skin and 32.3% nasal) on at least one LEAP study visit (Table 1), and the majority of these 239 participants tested positive only once (Online Repository Table E1). The highest rates of 240 colonization were recorded at 4-11 months of age (18% for skin and 15% for nose); these 241 decreased up to 30 months of age with a small increase observed at 60 months of age (Table 242 1). With the exception of the results at 60 months, the skin was more commonly the sole 243 colonized location compared to the nose (Table 1). No significant differences in terms of 244 frequency and persistence in all forms of S. aureus colonization were noted between the LEAP 245 avoiders and consumers (Online Repository Table E1). There was a small but significant 246 association between S. aureus colonization in the nose and on the skin, but concordance at any 247 particular time was slight (Online Repository Table E2).
- Very few of the total *S. aureus* positive swab samples were identified as methicillin resistant [skin 7/263 (2.7%); nose 2/257 (0.8%)].
- We additionally performed an exploratory analysis to investigate the relationship between skin *S. aureus* colonization at baseline and oral or topical antibiotic/steroid medication use at baseline. We did not find a statistically significant difference (p=0.695) in terms of skin *S. aureus* colonization when comparing subjects that were reported at baseline to have received these
- 254 medications versus those that did not (data not shown).
- 255

256 S. aureus colonization affected eczema severity and resolution

257 I). Eczema Severity

258 S. aureus colonization was significantly associated with concurrent eczema severity (measured 259 by SCORAD mean (SD) and SCORAD severity classification) across all study time points. 260 Participants with skin S. aureus had higher SCORAD values compared to those who did not 261 have skin S. aureus (Table 2). The majority of the subjects that were skin S. aureus colonized 262 had concurrent moderate and severe eczema at all time points (Online Repository Figure E1). 263 Those with nasal S. aureus colonization also had higher SCORAD values compared to those 264 who did not have nasal S. aureus; however the association was less strong than that observed 265 between skin S. aureus and eczema severity (Table 2).

266

267 II). Eczema persistence and deterioration

As previously published, eczema severity decreased over time, and there was no significant

269 difference in eczema severity between the two LEAP intervention groups (21). Although

270 SCORAD generally decreased over time, this was not the case for participants who were skin

colonized with S. aureus at certain visits (Figure 1). Indeed, considering the 12-30 and 60-72

- month time intervals, eczema significantly worsened in participants with immediately preceding
 skin *S. aureus* colonization relative to those without.
- 274 Preceding nasal *S. aureus* colonization was not associated with eczema persistence or 275 deterioration (Online Repository Figure E2).
- 276 277

278 S. aureus colonization was associated with food slgE and total lgE production

Hen's egg white and peanut slgE production at each LEAP and LEAP-On study visit was significantly associated with skin *S. aureus* positivity at any time point in the interval from baseline to 60 months (Online Repository Figure E3 and Figure 2 respectively). Importantly, these associations were corrected for eczema severity at each time point.

283

284 Notably, high levels of hen's egg white and peanut slgE production at each visit were also 285 associated with skin S. aureus positivity at any time point in the interval from baseline to 60 286 months (p<0.05) (Figure 3). In Figure 3, the divergence in the distribution at each time point 287 demonstrates that high level hen's egg white and peanut slgE values were disproportionately 288 represented in those participants who were skin colonized with S. aureus compared to those 289 who were not. For peanut slgE, this association was most apparent at 30 months but remained 290 subsequently. In contrast, the association for hen's egg white slgE became stronger over time 291 with S. aureus positive participants comprising over half of the upper tail of the relative 292 distribution of sIgE despite only representing a third of the overall sample. Furthermore, we 293 investigated the relationship between skin S. aureus and high level sIgE production to cow's 294 milk, and found a similar relationship with that observed for egg white and peanut. Indeed, at 295 30, 60, and 72 months, high levels of cow's milk slgE were associated with skin S. aureus 296 colonization at any time point in the interval from baseline to 60 months (Online Repository 297 Figure E4). Finally, high levels of total IgE at all assessments, were associated with any skin S. 298 aureus positivity (Online Repository Figure E4).

299

300 In order to assess if the observed associations between S. aureus colonization and high slgE 301 production to foods were food specific or confounded by total IgE, we examined the correlation 302 between total IgE and each of the three food sIgEs (cow's milk, egg white, and peanut). The 303 three pairwise correlations between each food and total IgE were moderate and consistent over 304 the 4 study visits (Online Repository Figure E5). Using multivariate logistic regression models, 305 egg white and peanut sIgE levels at 60 months were significantly associated with skin S. aureus 306 positivity after adjusting for total IgE at 60 months (Online Repository Figure E6). This 307 association was less strong for cow's milk slgE. In contrast, after adjustment with each food sIgE, total IgE levels were no longer significantly associated with skin S. aureus positivity 308 309 (Online Repository Figure E6).

310

311 S. aureus colonization was related to persistence and development of food allergy

312 I). Persistence of egg allergy

313 Of the 408 subjects with protocol defined egg allergy at baseline, 42.7% and 38.1% had

314 persistent egg allergy at 60 and 72 months respectively.

315 Overall, participants that had skin and/or nasal S. aureus colonization in the interval from 316 baseline to 60 months were 1.57 (95% CI, 1.02-2.42; p=0.042) times as likely to have persistent 317 egg allergy at 60 months of age as opposed to those that did not (Table 3). This association 318 was slightly stronger for nasal (OR 1.61; 95% CI, 1.03-2.52; p=0.036) as opposed to skin (OR 319 1.39; 95% CI, 0.88-2.19; p=0.160) S. aureus colonization. Skin S. aureus colonization prior to 320 72 months of age was the only colonization pattern significantly associated with the likelihood 321 (OR 1.77; 95% CI, 1.09-2.89; p=0.022) of egg allergy persisting until that age. There was a non-322 significant trend for preceding nasal (OR 1.54; 95% CI, 0.95-2.49; p=0.079) as well as skin 323 and/or nasal (OR 1.59; 95% CI, 0.99-2.55; p=0.055) colonization and egg allergy persisting at 324 72 months. When comparing the LEAP intervention groups, no association was noted between 325 persistent egg allergy and S. aureus colonization. All odds ratios were corrected for eczema 326 severity at 60 or 72 months accordingly (Table 3).

- 327
- 328 II). Development of peanut allergy

Overall, participants that had skin and those that had nasal *S. aureus* colonization in the interval from baseline to 60 months were 2.94 (95% Cl, 1.11, 7.76; p=0.029) and 2.41 (95% Cl, 1.04, 5.59; p=0.04) times as likely to have a diagnosis of peanut allergy at 60 months respectively as opposed to those that were not colonized. In addition, any preceding form of *S. aureus* colonization was significantly associated with peanut allergy at 72 months of age. All odds ratios were corrected for eczema severity at 60 or 72 months accordingly (Table 4).

335

336 Within the peanut consumption group, subjects that were skin S. aureus colonized at any study 337 point through LEAP were 7.13 (95% CI, 1.14, 44.47; p=0.035) and 3.87 (95% CI, 1.02, 14.65; 338 p=0.047) times as likely to be diagnosed with peanut allergy at 60 and 72 months of age 339 respectively compared with participants that were never skin S. aureus colonized (Table 4 and 340 Figure 4). With regards to nasal or 'skin and/or nasal' colonization at both time points, this 341 association was statistically significant only when it concerned nasal S. aureus and peanut 342 allergy at 60 months of age (Table 4 and Online Repository Figures E7 & E8). These odds 343 ratios are based on a small number of subjects who developed peanut allergy within the LEAP 344 consumers group. Specifically, there were only 9 (6 by 60 months and an additional 3 by 72 345 months) LEAP consumers who did not have peanut allergy at baseline and were diagnosed with 346 peanut allergy at 60 and/or 72 months. All but one of these 9 LEAP consumers (9/312) had S. 347 aureus colonization at one or more time points (Online Repository Fig E9). The 6 LEAP 348 consumers who were diagnosed with peanut allergy at both 60 and 72 months had all stopped 349 consumption well before 60 months of age due to suspected allergic reactions following peanut 350 consumption. In addition, there were 7 individuals in the consumption group who were allergic at 351 baseline. Of these, 6 had some form of S. aureus colonization at some point during the study 352 (data not shown). Within the avoidance group, there was no higher risk for peanut allergy at 60 353 or at 72 months in the subjects with any S. aureus colonization (Table 4).

354

The increased risk of peanut allergy at 60 or 72 months of age among the peanut avoiders compared to the consumers was less marked in those who had any *S. aureus* compared to those without *S. aureus* (Table 4, Panel B in Fig 4 and Online Repository Fig E7 & E8).

359 **DISCUSSION**

Previous findings that *S. aureus* colonization in eczema is associated with food sensitization and allergy (17, 18) may be confounded by eczema severity. In the LEAP and LEAP-On Studies we aimed to elucidate the relationship between *S. aureus* and food sensitization/allergy by correcting our analyses for eczema severity.

364

In the LEAP Study cohort, approximately half of the participants were found to be colonized by
 S. aureus. (Table 1 and Discussion in Online Repository). We demonstrate that skin
 colonization with *S. aureus* was related to eczema severity, persistence and deterioration.
 (Table 2, Fig 2 and Discussion in Online Repository).

369

370 In addition, we demonstrate that - even after correcting for eczema severity - hen's egg white 371 and peanut slgE values at each visit in LEAP and LEAP-On were significantly associated with 372 skin S. aureus positivity at any LEAP study time point (Online Repository Fig E3 and Fig 2). This 373 relationship was even stronger when we looked into high-level hen's egg white and peanut slgE 374 production (Fig 3). Similar findings are noted for cow's milk, where high level slgE production to 375 milk at 30, 60 and 72 months of age was related with any skin S. aureus colonization (Online 376 Repository Figure E4). Together these data suggest that S. aureus is associated with hen's egg, 377 peanut and cow's milk allergy.

378

379 Moreover, high levels of total IgE production were significantly associated with any skin S. 380 aureus colonization (Online Repository Figure E4) which is consistent with literature reporting 381 that S. aureus can promote a polyclonal IgE response [12]. In order to investigate whether sIgE 382 to foods in subjects with S. aureus colonization is explained by total IgE production, we explored 383 the relationship between total IgE levels and food sIgE levels to cow's milk, hen's egg white, 384 and peanut and found a significant but moderate correlation (Online Repository Figure E5). 385 Furthermore, we found that the association between egg white or peanut slgE at 60 months and 386 S. aureus colonization was not explained by total IgE (Online Repository Figure E6). However, 387 the association between total IgE levels and skin S. aureus was not significant when we 388 adjusted our analysis for each food sIgE (milk, egg white, peanut) (Online Repository Figure 389 E6). Overall these results indicate that in our study population high polyclonal IgE production in 390 the subjects with S. aureus colonization could only partly account for the association between 391 skin S. aureus colonization and high levels of egg white and peanut sIgE.

392

Allergy to hen's egg typically resolves during early childhood (22). However, in LEAP and LEAP-On, 42.7% and 38.1% of the baseline egg allergic participants had persistent egg allergy at 60 and 72 months of age respectively. Our results demonstrate that any *S. aureus* positivity increased the odds of hen's egg allergy persisting at 60 (OR 1.57, p=0.042) or 72 (OR 1.59, p=0.055) months of age independent of eczema severity (Table 3) suggesting that *S. aureus* may prevent the acquisition of natural tolerance to hen's egg.

399

In the LEAP Study, peanut consumption was successful in preventing peanut allergy at 60
 months of age. Interestingly, LEAP consumers with *S. aureus* skin colonization were 7.13
 (p=0.035) and 3.87 (p=0.047) times more likely to develop peanut allergy primarily confirmed by

403 peanut challenge at 60 or 72 months of age respectively (Table 4, Fig 4). Whilst these 404 associations are based on only 9 (6 by 60 months and an additional 3 by 72 months) LEAP 405 consumers who did not have peanut allergy at baseline and were diagnosed with peanut allergy 406 at 60 and/or 72 months, it is worth noting that all but one of these participants were colonized 407 with S. aureus at one or more LEAP visits (Online Repository Fig E9). The 6 subjects that 408 developed peanut allergy by 60 months of age had all stopped consuming peanut well before 60 409 months of age. It could therefore be argued that the reason for failing to acquire oral tolerance 410 was inadequate consumption rather than the immunological effect of S. aureus. However, these 411 6 subjects stopped eating peanut during the course of the study because of symptoms during 412 consumption that strongly suggested peanut allergy. This indicates that the reduced duration of 413 peanut consumption was the consequence of an accelerated development of peanut allergy. 414 More specifically, there are two possible explanations for the development of peanut allergy 415 despite previous peanut consumption in these subjects: A) they developed an accelerated form 416 of peanut allergy potentiated by S. aureus, and/or B) S. aureus may have inhibited tolerance 417 mechanisms related to peanut consumption. The fact that S. aureus was associated with a 418 higher risk of peanut allergy among peanut consumers but not avoiders (Table 4, Panel B in Fig 419 4 and Online Repository Fig E7 & E8) further suggests that peanut consumption was less 420 effective in the prevention of peanut allergy among participants with S. aureus compared to 421 those with no S. aureus.

422

423 S. aureus has been implicated in the development and severity of atopic diseases such as 424 eczema, allergic rhinitis and asthma. With regards to food allergy, an epidemiological clinical 425 study indicates an association between skin S. aureus and milk, egg or peanut allergy in 426 children with eczema (16). There are murine studies that support a biological explanation 427 between S. aureus and food allergy. Indeed, SEB co-applied on the skin with ovalbumin or 428 peanut extract increases the systemic production of ovalbumin slgE (23) and enhances peanut 429 specific CD4⁺ Th2 responses on subsequent exposure to peanut extract alone (24) respectively. 430 Additionally, SEB administered orally with antigen (ovalbumin or peanut) results in highly Th2 431 polarized immune responses to the antigen, while subsequent oral challenge with the respective 432 antigen triggers anaphylaxis (25). In all three studies, the antigen specific immune responses 433 were not observed with SEB or the antigen alone suggesting that S. aureus might be acting as 434 adjuvant. Our results show an association between skin S. aureus and high slgE production to 435 hen's egg white, peanut and cow's milk as well as to high total IgE levels. However, we 436 demonstrated that the relationship between S. aureus and sIgE production to egg white and 437 peanut was primarily explained by the corresponding food allergen slgE and not total IgE levels. 438 S. aureus has been associated with more severe forms of atopic diseases, and our data extend 439 these observations in food allergy.

440

441 Study strengths include the longitudinal design of the LEAP Study with detailed clinical 442 assessments and colonization results obtained at four scheduled study intervals. As our results 443 are corrected for eczema severity, we are able to confirm that the association between *S.* 444 *aureus* carriage and egg/peanut slgE production or allergy occurred independent of eczema 445 severity.

447 There are limitations to the colonization results reported as use was made of less sensitive 448 bacteriological culture techniques and not DNA-based testing. Nevertheless, cultures allow for 449 the detection of live microorganisms and not remnant, nonviable genetic material from prior 450 infection. As we did not genotype the isolated strains, it is not possible to match organisms over 451 time and between skin and nasal swabs. Swabs were collected on only 4 occasions in LEAP 452 and were not collected in LEAP-On. Diagnostic food challenges were undertaken to peanut but 453 not hen's egg. A major limitation is related to the interpretation of the association between S. 454 aureus and peanut allergy in the consumers, which, although significant, is based on the very 455 small numbers of LEAP consumers who became peanut allergic as it is reflected in the wide 456 confidence intervals around the odds ratios. Larger numbers of participants who become peanut 457 allergic - despite being fed peanut in infancy/early childhood - would be required to assess if 458 these findings do indeed demonstrate that S. aureus colonization interferes with oral tolerance 459 induction. Finally, even after adjusting for eczema severity, we cannot rule out that the observed 460 association between colonization and food allergy could be due to other confounding factors.

461

462 S. aureus has been implicated in the development and severity of atopic diseases namely 463 eczema, allergic rhinitis and asthma; our findings extend these observations to the development of food allergy, independent of eczema severity. The role of S. aureus as a potential 464 465 environmental factor should be considered in future interventions aimed at inducing and 466 maintaining tolerance to food allergens in eczematous infants. Further prospective longitudinal 467 studies measuring S. aureus with more advanced techniques and interventional studies eradicating S. aureus in early infancy will help elucidate its role in the development of eczema or 468 469 food allergy.

470

471

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487

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493

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- 586

587 **Display Legends**

588

589 Table 1. Skin and Nasal *S. aureus* Colonization Prevalence Over Time in LEAP

590

	4-11 (mo)	12 (mo)	30 (mo)	60 (mo)	Ever Colonized 4-11(mo) – 60(mo)	
Skin S. aureus						
N	640	626	618	630	640	
S. aureus	115 (18.0%)	63 (10.1%)	40 (6.5%)	45 (7.1%)	206 (32.2%)	
No S. aureus	525 (82.0%)	563 (89.9%)	578 (93.5%)	585 (92.9%)	434 (67.8%)	
Nasal S. aureus						
Ν	640	626	618	630	640	
S. aureus	96 (15.0%)	35 (5.6%)	32 (5.2%)	94 (14.9%)	207 (32.3%)	
No S. aureus	544 (85.0%)	591 (94.4%)	586 (94.8%)	536 (85.1%)	433 (67.7%)	
Skin and/or Nasal S. aureus						
Ν	640	626	618	630	640	
S. aureus	166 (25.9%)	87 (13.9%)	66 (10.7%)	125 (19.8%)	312 (48.8%)	
No S. aureus	474 (74.1%)	539 (86.1%)	552 (89.3%)	505 (80.2%)	328 (51.3%)	
Skin and Nasal S. aureus Combination						
Ν	640	626	618	630		
Nasal Only	51 (8.0%)	24 (3.8%)	26 (4.2%)	80 (12.7%)		
Skin Only	70 (10.9%)	52 (8.3%)	34 (5.5%)	31 (4.9%)		
Skin and Nasal	45 (7.0%)	11 (1.8%)	6 (1.0%)	14 (2.2%)		
Neither	474 (74.1%)	539 (86.1%)	552 (89.3%)	505 (80.2%)		

591

592

593 The prevalence of skin, nasal, skin or nasal, and the combination of skin and nasal S. aureus colonization for all subjects enrolled in LEAP at

baseline (4-11 months), 12 months, 30 months, and 60 months are shown. If a subject has at least one instance of *S. aureus* colonization at any of

the 4 LEAP visits (4-11 mo to 60 mo) then that subject is summarized as 'S. aureus' in the 'Ever Colonized' column. Analogously, if a subject

S. aureus and food allergy in LEAP/LEAP-On

never has *S. aureus* at any of the 4 LEAP visits (4-11 mo to 60 mo) then that subject is summarized as 'No *S. aureus*' in the 'Ever Colonized'
 column. This definition of 'Ever Colonized' is utilized in subsequent analyses.

599 Table 2. Concurrent Skin and Nasal S. aureus Colonization and Eczema Severity

600

				SI	kin <i>S. aureu</i>	ıs						
		4-11 (mo)			12 (mo)			30 (mo)		. <u></u>	60 (mo)	
	No S. aureus	S. aureus	p-value	No S. aureus	S. aureus	p-value	No S. aureus	S. aureus	p-value	No S. aureus	S. aureus	p-value
SCORAD N Mean (SD) LS Means (SE) Diff LS Means (S. aureus - No S. aureus)	525 32.6 (18.5) 33.1 (0.8)	115 42.3 (18.6) 40.1 (1.6)	<.001 6.9 (3.6, 10.2)	563 20.5 (14.1) 21.0 (0.6)	63 31.6 (16.5) 27.5 (1.5)	<.001 6.5 (3.3, 9.6)	576 15.1 (12.9) 15.4 (0.5)	40 33.1 (16.8) 28.4 (1.8)	<.001 13.0 (9.4, 16.6)	583 5.9 (9.9) 6.3 (0.4)	45 22.1 (15.3) 17.1 (1.3)	<.001 10.8 (8.1, 13.5)
				Na	asal <i>S. aure</i>	us						
		4-11 (mo)			12 (mo)			30 (mo)			60 (mo)	
	No S. aureus	S. aureus	p-value	No S. aureus	S. aureus	p-value	No S. aureus	S. aureus	p-value	No S. aureus	S. aureus	p-value
SCORAD N Mean (SD) LS Means (SE)	544 33.5 (18.9) 33.7 (0.8)	96 39.6 (17.8) 38.5 (1.7)	0.009	591 21.4 (14.7) 21.4 (0.6)	35 26.5 (14.1) 26.4 (2.0)	0.015	584 16.0 (13.6) 16.0 (0.6)	32 21.7 (17.5) 20.6 (2.0)	0.024	534 6.5 (10.3) 6.6 (0.5)	94 10.6 (14.6) 9.5 (1.0)	0.005
Diff LS Means (S. aureus - No S. aureus)			4.8 (1.2, 8.3)			5.0 (1.0, 9.1)			4.6 (0.6, 8.7)			2.9 (0.9, 4.9)

601

602 Data is presented for Eczema severity defined by SCORAD for all participants who were in LEAP with available data for each time point divided

603 into groups based on whether a subject had S. aureus at the concurrent visit or did not have S. aureus at the concurrent visit. P-values are from a

604 longitudinal repeated measures model comparing the difference in least squares means in SCORAD between subjects without S. aureus

605 colonization to those with *S. aureus* colonization.

607 **Table 3.** Persistent Egg Allergy in Relation to *S. aureus* Colonization and Treatment Assignment

608

S. aureus Colonization	LEAP N=363			LEAP-On N=318			
(Baseline to 60 Months)	Odds Ratio	95% CI	p-value	Odds Ratio	95% CI	p-value	
×	Overall	(S. aureus vs N	lo S. aureus)			•	
Skin S. aureus	1.39	{0.88, 2.19}	0.160	1.77	{1.09, 2.89}	0.022	
Nasal S. aureus	1.61	{1.03, 2.52}	0.036	1.54	{0.95, 2.49}	0.079	
Skin and/or Nasal S. aureus	1.57	{1.02, 2.42}	0.042	1.59	{0.99, 2.55}	0.055	
Within	n Peanut Consu	mption Group (S. aureus vs I	No S. aureus)			
Skin S. aureus	1.37	{0.73, 2.58}	0.326	1.68	{0.85, 3.35}	0.139	
Nasal S. aureus	1.42	{0.76, 2.67}	0.276	1.65	{0.83, 3.26}	0.154	
Skin and/or Nasal S. aureus	1.65	{0.89, 3.03}	0.108	1.88	{0.96, 3.70}	0.066	
XX7'.(1	· D (A ·	1 0 (0	N	C)			
		dance Group (S			(0.05.2.(7))	0.072	
Skin S. aureus	1.39	$\{0.74, 2.64\}$	0.300	1.86	$\{0.95, 3.67\}$	0.072	
Nasal S. aureus	1.83	$\{0.98, 3.43\}$	0.059	1.44	$\{0.73, 2.86\}$	0.295	
Skin and/or Nasal S. aureus	1.49	{0.81, 2.73}	0.196	1.34	{0.69, 2.58}	0.385	
With	nin Those With	S. aureus (Avo	idance vs. Co	nsumption)			
Skin S. aureus	0.88	{0.44, 1.77}	0.717	0.94	{0.44, 1.99}	0.869	
Nasal S. aureus	1.02	{0.49, 2.09}	0.955	0.81	{0.37, 1.77}	0.600	
Skin and/or Nasal S. aureus	0.85	{0.47, 1.52}	0.573	0.78	$\{0.41, 1.47\}$	0.440	
	T1 XX7.41		.1				
		at S. aureus (Av			(0.47.1.54)	0.505	
No Skin S. aureus	0.86	{0.51, 1.47}	0.583	0.85	{0.47, 1.54}	0.587	
No Nasal S. aureus	0.79	$\{0.47, 1.34\}$	0.386	0.93	{0.52, 1.66}	0.799	
No Skin and/or Nasal S. aureus	0.93	{0.50, 1.74}	0.829	1.09	{0.55, 2.18}	0.797	

609

This table displays the odds ratios, 95% confidence intervals, and p-values from multiple multivariate logistic regression models. One set of models was fit for the 60 month data (outcome of interest being persistent egg allergy as assessed by raw and pasteurized egg skin prick test wheal cut-offs at 60 months), and another set of models was fit for the 72 month data (outcome of interest being persistent egg allergy as assessed by raw and pasteurized egg skin prick test wheal cut-offs at 72 months) with *S. aureus* colonization status (one model each for skin, nasal, and skin and/or nasal) adjusted for SCORAD (at 60 and 72 months respectively), LEAP treatment assignment, and the interaction between *S. aureus* status and treatment assignment. Those

615 who do not have protocol-defined egg allergy at baseline are not included in this analysis.

616

S. aureus and food allergy in LEAP/LEAP-On

617 **Table 4.** Peanut Allergy in Relation to *S. aureus* Colonization and Treatment Assignment

618

S. aureus Colonization		LEAP N=619	LEAP-On N=538			
(Baseline to 60 Months)	Odds Ratio	95% CI	p-value	Odds Ratio	95% CI	p-value
(Dasenne to oo Wonths)		dll (S. aureus vs N	1	Ouus Katio	75 /0 CI	p-value
Skin S. aureus	2.94	{1.11, 7.76}	0.029	2.19	{1.04, 4.61}	0.039
Nasal S. aureus	2.41	{1.04, 5.59}	0.040	2.18	{1.05, 4.56}	0.037
Skin and/or Nasal S. aureus	4.24	{0.97, 18.59}	0.055	2.78	{1.09, 7.07}	0.031
Wi	thin Peanut Cons	sumption Group (S	5. aureus vs	No S. aureus)		
Skin S. aureus	7.13	{1.14, 44.47}	0.035	3.87	{1.02, 14.65}	0.047
Nasal S. aureus	3.78	{0.79, 18.11}	0.096	3.88	{1.03, 14.61}	0.045
Skin and/or Nasal S. aureus	12.26	{0.68, 220.56}	0.089	5.57	{0.96, 32.26}	0.055
II.		·1 0 (6				
		bidance Group (S.			(0.65.0.05)	0.500
Skin S. aureus	1.21	{0.65, 2.25}	0.545	1.24	{0.65, 2.37}	0.508
Nasal S. aureus	1.54	{0.84, 2.82}	0.162	1.23	{0.65, 2.32}	0.519
Skin and/or Nasal S. aureus	1.47	{0.81, 2.67}	0.208	1.39	{0.75, 2.58}	0.293
W	Vithin Those Wit	h S. aureus (Avoi	dance vs. Co	onsumption)		
Skin S. aureus	4.29	{1.60, 11.51}	0.004	3.27	{1.27, 8.43}	0.014
Nasal S. aureus	5.78	{2.01, 16.65}	0.001	3.23	{1.25, 8.34}	0.015
Skin and/or Nasal S. aureus	5.86	{2.43, 14.14}	< 0.001	3.97	{1.77, 8.95}	0.001
Wi	thin Those With	out S. aureus (Ave	oidance vs. (Consumption)		
No Skin S. aureus	25.26	{4.86, 131.35}	<0.001	10.18	{3.31, 31.35}	< 0.001
No Nasal S. aureus	14.19	{3.86, 52.21}	< 0.001	10.19	{3.31, 31.33}	< 0.001
No Skin and/or Nasal S. aureus		{2.93, 815.20}	0.007	15.90	{2.98, 84.66}	0.001

619

This table displays the odds ratios, 95% confidence intervals, and p-values from multiple multivariate logistic regression models using the Firth penalized likelihood method. One set of models was fit for the 60 month data (outcome of interest being peanut allergy as assessed by oral food challenge at 60 months), and another set of models was fit for the 72 month data (outcome of interest being peanut allergy as assessed by oral food challenge at 72 months). Predictors of interest included *S. aureus* colonization status (one model each for skin, nasal, and skin and/or nasal) adjusted for SCORAD (at 60 and 72 months respectively), LEAP treatment assignment, and the interaction between *S. aureus* status and treatment assignment. Infants randomly assigned to consumption underwent a baseline, open-label food challenge; the 7 subjects who reacted to that challenge are not included in this analysis. Interpret results with caution as a small number of which with pagent allergy (agencielly in the Pagent Consumption arm) contribute to these analysis.

626 of subjects with peanut allergy (especially in the Peanut Consumption arm) contribute to these analyses.

- 627 **Figure 1.** Eczema Severity by Skin *S. aureus* Colonization at the Preceding Visit
- 628 Data is presented for all participants who were in LEAP and LEAP-On with available SCORAD 629 data for each study assessment time point divided into groups based on whether subjects had skin S. aureus at the previous visit (in red) or did not have skin S. aureus at the previous visit (in 630 blue). Black diamonds represent model predicted means, boxes represent 25th and 75th centiles, 631 632 error bars represent 2.5th and 97.5th centiles, and the middle line of the box represents the 633 median. The total number of subjects contributing to the analysis at each time point, p-values, 634 mean differences and 95% confidence intervals around that difference directly above each 635 assessment time point refer to the least squares mean difference (S. aureus - no S. aureus) and 636 p-value comparison between those who had skin S. aureus at the previous visit and those who 637 did not have skin S. aureus at the previous visit using a longitudinal repeated measures model 638 adjusted for SCORAD at the previous visit, time, S. aureus status at the previous visit, and the 639 interaction between S. aureus status at the previous visit and time.
- 640

641 **Figure 2.** Peanut sIgE Over Time by Skin S. aureus Colonization Status

- 642 Data is presented for all participants who were in LEAP and LEAP-On with available Peanut 643 Specific IgE data for each study assessment time point divided into groups based on whether 644 subjects ever had skin S. aureus from baseline to 60 months (in red) or never had skin S. aureus 645 from baseline to 60 months (in blue). Black diamonds represent model predicted means, boxes represent 25th and 75th centiles, error bars represent 2.5th and 97.5th centiles, and the middle line 646 of the box represented the median. The total number of subjects contributing to the analysis at 647 648 each time point, p-values, mean differences and 95% confidence intervals around that mean 649 difference directly above each assessment time point refer to the comparison between those who 650 never have S. aureus and those who ever have S. aureus groups using a longitudinal repeated 651 measures model adjusted for SCORAD, time, S. aureus status, and the interaction between S. 652 *aureus* status and time. Average SCORAD values at each time point are annotated directly 653 below the box plots for those who ever had skin S. aureus (red) and those who never had skin S. 654 aureus (blue).
- 655

Figure 3. Relative Distribution of Hen's Egg White and Peanut sIgE Over Time by Skin S.
 aureus Colonization Status

These figures show the relative distribution of hen's egg white-specific IgE and peanut-specific IgE between those who ever have skin *S. aureus* (shown in red) from 4-11 months to 60 months and those who never have skin *S. aureus* (shown in blue). The vertical reference lines indicate where the distribution begins to significantly differ (p < 0.05) between the two groups using bootstrap sampling of 1000 replicates of the upper percentiles indicating that those with *S. aureus* colonization are over represented in the higher end of the distribution of sIgE (which is more indicative of allergy). A reference panel is included to illustrate the 67.8% of the trial participants who never had skin *S. aureus* and the 32.2% who ever had skin *S. aureus* and what a pattern with no association of skin *S. aureus* with sIgE levels would look like.

668

669 Figure 4. Peanut Allergy in Relation to Skin S. aureus Colonization and Treatment Assignment Percents (from raw data), odds ratios and 95% confidence intervals from multiple multivariate 670 671 logistic regression models using the Firth penalized likelihood method are displayed. One model 672 was fit for the 60 month data (outcome of interest being peanut allergy as assessed by oral food 673 challenge at 60 months), and another model was fit for the 72 month data (outcome of interest 674 being peanut allergy as assessed by oral food challenge or the relevant diagnostic algorithm at 72 months). Predictors of interest included skin S. aureus colonization status adjusted for SCORAD 675 676 (at 60 and 72 months respectively), LEAP treatment assignment, and the interaction between 677 skin S. aureus status and treatment assignment. Panel A for the plot summarize the relationship 678 between peanut allergy and skin S. aureus colonization status (overall, within consumers, and 679 within avoiders). In the 'Percent' panel, the numerators refer to the number of subjects with 680 peanut allergy while the denominator refers to the number of subjects with skin S. aureus (in red) 681 and those without skin S. aureus (blue). Panel B of the plot summarize the relationship between 682 peanut allergy and peanut consumption (overall, within those with skin S. aureus, within those 683 without skin S. aureus). In the 'Percent' panel, the numerators refer to the number of subjects 684 with peanut allergy while the denominator refers to the number of subjects in the avoidance group (in grey) and those in the consumption group (green). Interpret results with caution as a 685 small number of subjects with peanut allergy (especially in the Peanut Consumption arm) 686 687 contribute to these analyses.