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Skin testing improves predictive value of mid-range peanut specific IgE and Ara h 2 levels in the LEAP study

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1 **Skin testing improves predictive value of mid-range peanut specific IgE and Ara h 2 levels in the**
 2 **LEAP study**

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26
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42 Clinical Implications:

43 There is a need to easily predict clinically important peanut allergy without OFC. The presence of both a
 44 positive SPT (3mm) and Ara h 2-sIgE ≥ 0.29 kU/L at 60 months may preclude the need for OFC in
 45 children meeting LEAP study inclusion criteria.

46 Peanut allergy has a prevalence of ~2% and is the leading cause of food-induced fatal and near-fatal
47 anaphylaxis. [1] Distinguishing between sensitization and clinically relevant peanut allergy remains
48 challenging, especially for patients whose peanut-specific immunoglobulin E (PN-sIgE) levels are neither
49 very low nor very high.

50 PN-sIgE is the most sensitive test for prediction of clinically important peanut allergy (sensitivity
51 (SN)=0.93), whereas Ara h 2-sIgE is the most specific (specificity (SP)=0.92). [2] Ara h 2 is the best
52 overall individual diagnostic test by receiver operative characteristic (ROC) analysis, with an area-under-
53 the-curve (AUC) of 0.84, compared to peanut-sIgE, which has an AUC of 0.69. [2]

54 Historically, PN-sIgE ≥ 0.35 kU/L was reported as a positive result, with low specificity (23-58%) and
55 positive predictive value (PPV) (44-71%).[3] Some laboratories now report PN-sIgE of ≥ 0.10 kU/L as a
56 positive test but the value of this cut-off has not been well studied.

57 Accurately predicting peanut allergy improves with higher peanut and Ara h 2-sIgE values. Previous data
58 suggest a PN-sIgE of 6 kU/L has a 90% predicted probability of clinical reactivity and at 13 kU/L this
59 probability increases to 95% with a sensitivity of 60%, specificity of 96%, PPV of 99% and negative
60 predictive value (NPV) of 35%. [4] A peanut skin prick wheal size of ≥ 8 mm and PN-sIgE ≥ 15 kU/L are
61 reported to predict peanut allergy with 95% and 92% certainty for a positive challenge, respectively. [5]
62 For Ara h 2-sIgE, a cutoff of ≥ 1.75 kU/L has a 100% positive predictive value, and ~30% of patients
63 could be diagnosed with 100% accuracy.[6] The addition of other measurements such as skin prick test
64 (SPT) wheal size or ex vivo basophil reactivity further increases the accuracy of these tests.[7, 8]

65 The Learning about Early Peanut Allergy (LEAP) trial was pivotal in demonstrating the benefit of early
66 introduction of peanuts in high-risk children to decrease the frequency of peanut allergy.[9] Santos et al.
67 used this data set to identify severity and threshold of reactions during oral peanut challenges.[7]
68 Optimal cutoffs conferring high risk for developing severe allergic reactions included BAT 48%, Ara
69 h 2-sIgE of 1.4 kU/L, PN-sIgE > 5 kU/L and a peanut SPT of 8mm. Multivariate models were
70 superior to individual biomarkers and were used to calculate the probability of serious adverse
71 events during oral food challenge (OFC). [7]

72 The LEAP study team developed a model using SPT, PN-sIgE, Ara h 1, Ara h 2, and Ara h 3-sIgE
73 values to predict peanut allergy in the absence of an OFC. [8] This prediction model, applied to 617
74 LEAP participants with a determinate OFC, had an AUC of 0.99, with an overall error rate of 2.6%.
75 However, mismatches between observed and predicted OFC results occurred when PN-sIgE ranged
76 from 0.20-2.17 kU/L. [8]

77 There remains a need to develop simple approaches using easily obtainable biomarkers to predict the
78 presence of clinically important allergies to peanuts without performing OFC in subjects with PN-
79 sIgE values in the non-extreme range of ≥ 0.1 and < 15 kU/L.

80
81 To address this need, we focused on the peanut avoidance group (n=321) in the LEAP study data set
82 because this group is a cohort of subjects at highest risk of having demonstrable peanut allergy at 5
83 years (60 months) of age when the graded OFC was performed.

84
85 All LEAP study participants randomized to the peanut avoidance group with very low PN-sIgE < 0.1 or
86 very high PN-sIgE > 9 kU/L at 60 months (n= 186) had predictable outcomes of either negative or positive
87 OFC respectively (Figure E1). Further biomarkers are not needed as the PN-sIgE data gave an AUC 1.0
88 with 100% sensitivity, specificity and accuracy. Changing the upper limit to > 13 kU/L or > 15 kU/L based
89 on the published literature did not affect this finding.

90 For subjects in the peanut avoidance group with PN-sIgE in a non-extreme range ≥ 0.1 to ≤ 9 kU/L at 60
91 months, a positive SPT at 60 months, defined as ≥ 3 mm, was superior (orange line, AUC 0.8619, $P < 10^{-11}$,
92 $n=116$) to PN-sIgE (yellow line, AUC= 0.67, $P= 0.002$, $n= 116$) or Ara h 2-sIgE alone at 60 months
93 (green line, AUC= 0.86, $p < 10^{-8}$, $n=112$) (Figure 1). These subjects have “difficult to diagnose” peanut
94 allergy, and additional biomarkers are needed to make more accurate predictions. Sensitivity, specificity,
95 accuracy, PPV, and NPV are shown for optimal cutoffs of PN-sIgE =6 kU/L (selected for PPV $\geq 90\%$) and
96 for Ara h 2-sIgE =0.56 kU/L (selected for PPV=100%).

97 The presence of a positive SPT ≥ 3 mm at 60 months markedly increases diagnostic accuracy of PN-sIgE
98 (AUC 0.8653, $p=10^{-9}$, $n=116$) but, at the chosen threshold of 6 kU/L, did not change the performance
99 metrics (blue line, Figure 2). However, for Ara h 2-sIgE, the presence of a positive SPT not only increased
100 diagnostic accuracy (purple line, AUC 0.943, $p < 10^{-12}$, $n = 112$), but also allowed a lower threshold of Ara
101 h 2-sIgE (≥ 0.29 kU/L) to reach 100% SP and PPV (no false positives) (Figure 1). A flowsheet of this
102 approach is shown (Figure E2). Importantly, as the SPT wheal size increases, the cut off for Ara h 2-sIgE
103 to achieve 100% SP and 100% PPV decreases dramatically (Figure 2).

104 In conclusion, the presence of a positive SPT and an Ara h 2-sIgE of ≥ 0.29 may preclude the need for an
105 OFC in subjects who meet the LEAP study inclusion criteria [9], with the understanding that there is a
106 significant false negative rate of 52%. Compared with Sever et al (8) where analysis of multiple
107 parameters in all subjects regardless of PN-sIgE values led to an error rate of 2.8%, our overall error rate
108 using only Ara h 2-sIgE and the presence of a positive SPT was 12.5%.

109 Strengths of this report include the focus on subjects with difficult to diagnose peanut allergies and the
110 use of easy to obtain clinical data. Limitations include the makeup of the LEAP cohort, the retrospective
111 analysis, the relatively small number of subjects in the difficult to diagnose range of PN-sIgE with
112 positive SPT, and the need to repeat this study both prospectively and with a different database.

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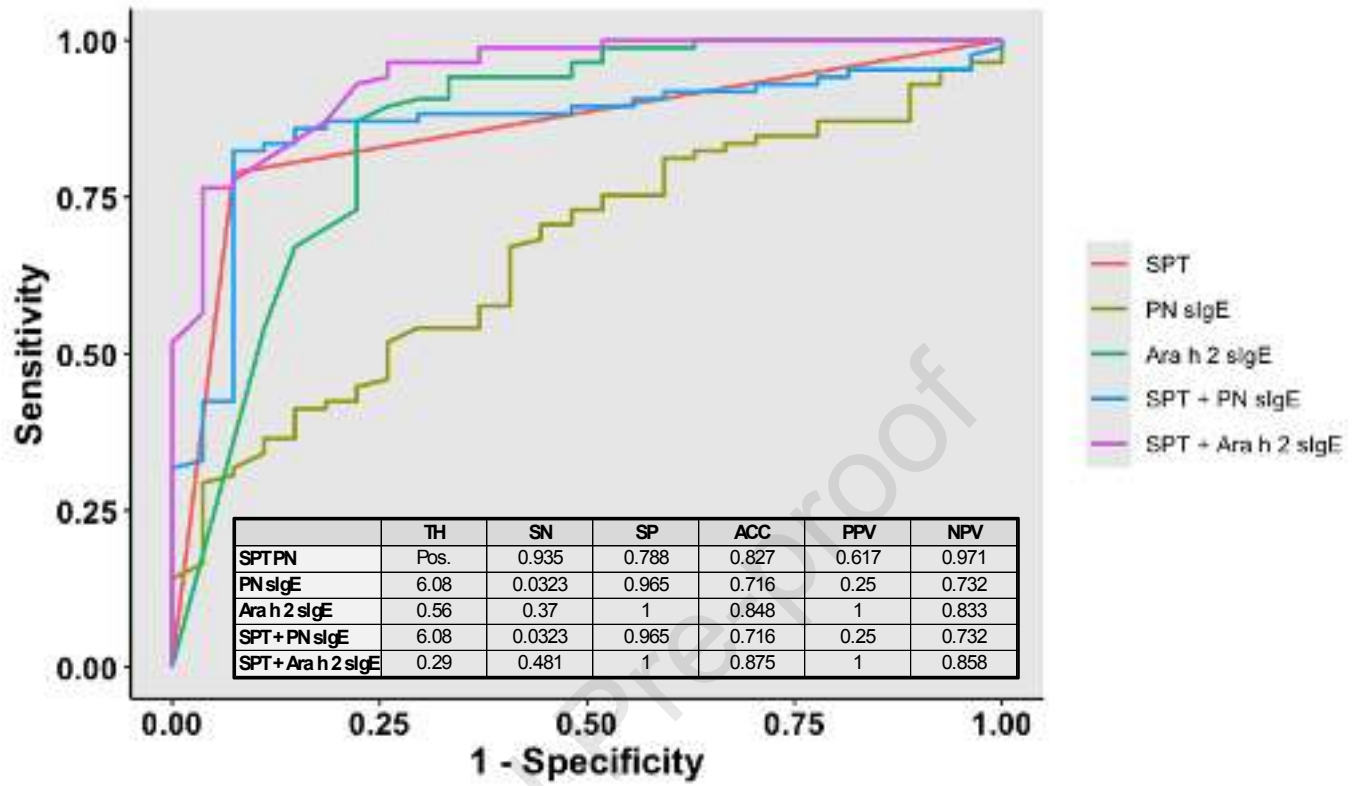
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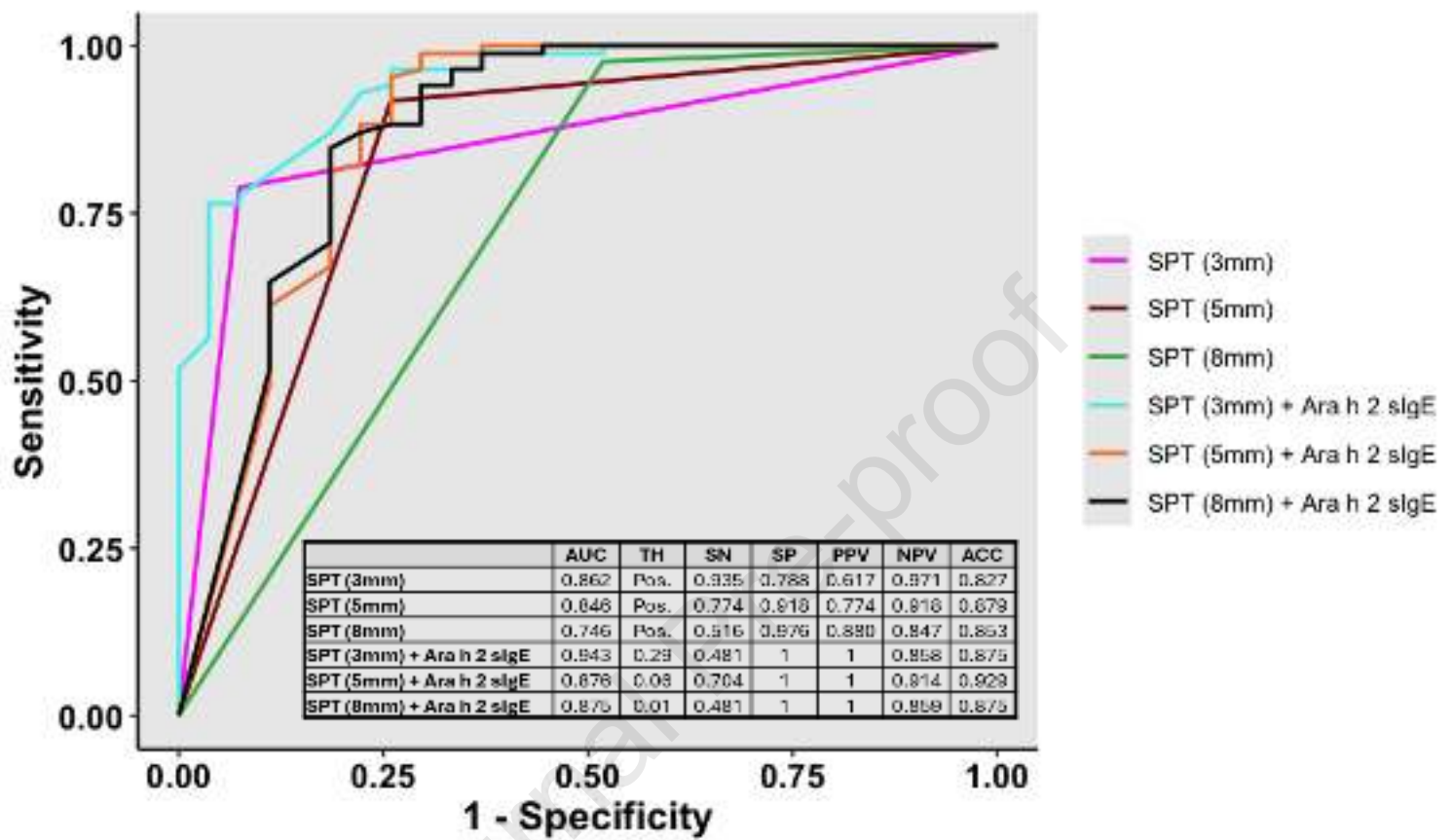
154 **Figure legends**

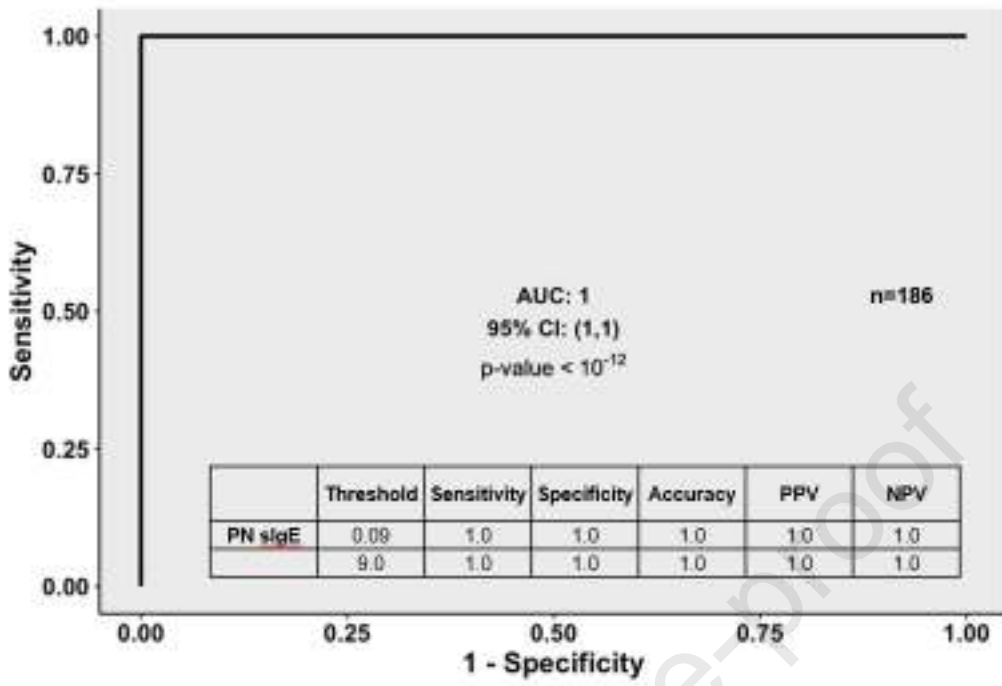
155 **Figure 1.** A positive SPT (3mm) augments prediction of peanut allergy when PN-sIgE ≥ 0.1 and ≤ 9 kU/L
 156 at 60 months. TH=threshold, SN=sensitivity, SP=specificity, ACC=accuracy, PPV=positive predictive
 157 value, NPV=negative predictive value.

158 **Figure 2.** Increasing SPT size from 3mm (n=43) to 5mm (n=27) to 8mm (n=15) decreases the threshold
 159 of Ara h 2-sIgE needed to achieve 100% SP and PPV.

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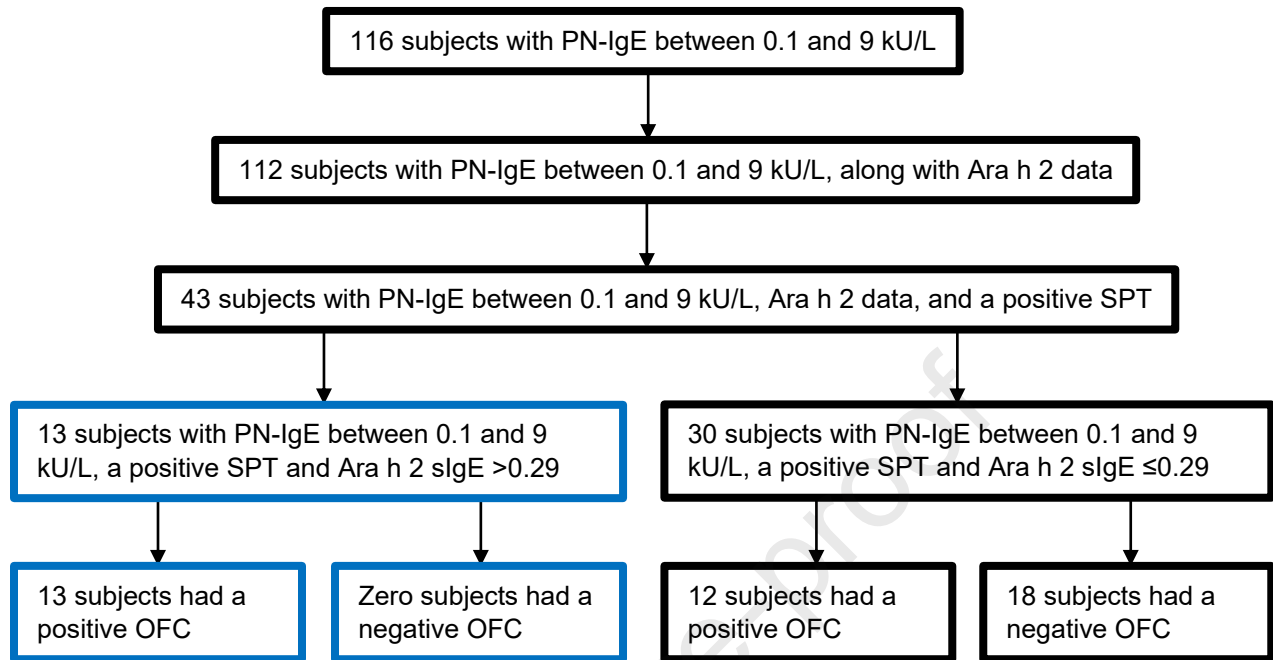


Figure E1. Peanut allergy is easy to predict in peanut avoidance group with PN-sIgE <0.1 or >9 kU/L at 60 months.

Figure E2. Flowchart. Key findings are highlighted in blue.

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