

# Impact of real-world confounders on the accuracy of an AI model to support read out of skin prick automated test results

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## Dear Editor:

Skin prick testing (SPT) is the gold standard for diagnosing allergic sensitization in individuals with a suspected airborne allergy <sup>(1)</sup>. Skin tests are used as first option in 90% of individuals suffering from respiratory allergies and almost two-third of all types of allergies <sup>(2)</sup>. However, its accuracy highly depends on the operator, causing variability during pricking and readout. S.P.A.T. or skin prick automated test standardises the SPT procedure and has been clinically validated <sup>(3)</sup>. The S.P.A.T. device serves 12 simultaneous pricks delivering a fixed amount of test solution with controlled prick force to the patient's forearm. This automation has proven to lower intra-subject variability and bring more consistent test results compared to manual SPT <sup>(4,5)</sup>.

Recently, an AI-assisted readout method was developed using a train-test approach including over 10,000 wheals following S.P.A.T. <sup>(6)</sup>. Comparing the measurements by AI with those of the physician revealed an accuracy of 95.4%. Performance evaluation in an independent cohort of 95 patients showed misclassifications that impacted the test interpretation in 0.5% of cases. Given that false positive or negative test results were mostly due to scars, hair, hyperpigmentation or darker skin tone, these real-world confounders were investigated in a post-hoc analysis.

Images of 217 patients (2,604 wheals) of the validation cohort

recruited in the previous pivotal study were analysed. The images of the forearms were analysed for skin tone, presence of hair, scars, tattoos and hyperpigmentation. Analysis of skin tone was performed using the individual typology angle (ITA) method <sup>(7)</sup>. Other possible confounders were assessed by visual inspection of the images and applying a qualitative score.

Table S1 provides an overview of the real-world confounders that were assessed. Figures S1-S2 show representative images for each confounder subgroup. Darker skin tone (ITA <0) was observed in 16.8% of patients. Presence of hair, hyperpigmentation, scar tissue or tattoos near the prick location was observed in respectively 30.4%, 3.5%, 1.0% and 1.5% of pricks. In patients with a darker skin tone (ITA < -50, ITA -50 to <-25, -25 to <0) accuracy decreased to respectively 85.4%, 90.3% and 92.0% compared to other ITA categories (96.5-96.7%) (Figure 1A). In patients with tattoo marks (category III), accuracy dropped to 88.3% (compared to 95.7-97.9%) (Figure 1B). For the other confounders, accuracy remained >90% irrespective of the presence of the confounding variable (Figure 1C-E).

The limited representation of patients with darker skin tones, or hyperpigmentation, scars and tattoos is a weakness of this study and may underestimate the extent of bias. Visuo-perceptive data and scoring are prone to inter-observer variability. It should also

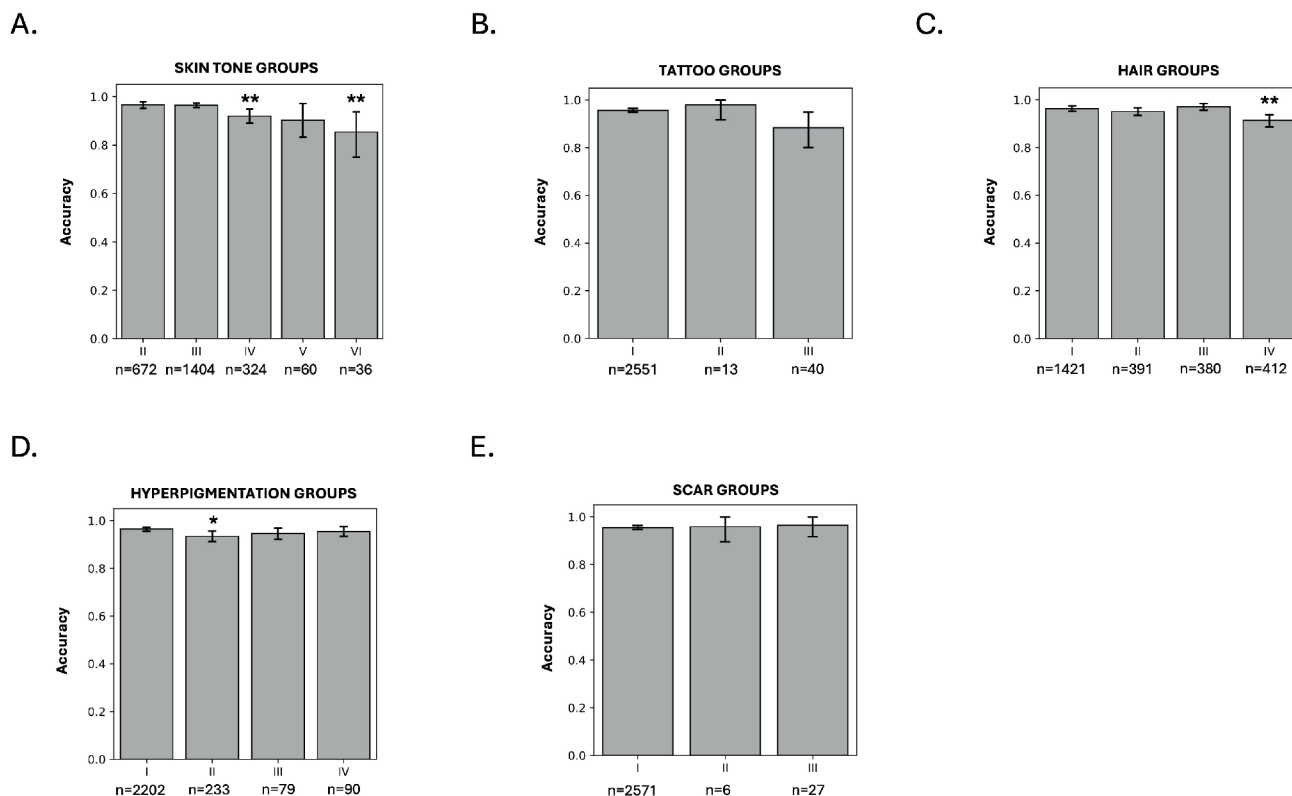


Figure 1. Analysis of accuracy of the AI model in patients stratified by real-world confounders. (A) Skin tone assessed by individual typology angle analysis. I: ITA > 50, II: ITA 25 to 50, III: ITA 0 to 25, IV: ITA -25 to 0, V: ITA -50 to -25, VI: ITA < -50. (B) Presence of tattoos assessed by visual inspection: I: no tattoos visible, II: tattoo visible at the side of the arm, III: tattoo visible in close proximity to this prick location. (C) Presence of hair assessed by visual inspection: I: no hair visible, II: minimal hair visible at the side of the arm, III: minimal hair visible in close proximity to this prick location, IV: abundant hair visible in close proximity to this prick location. (D) Presence of hyperpigmentation assessed by visual inspection: I: no hyperpigmentation visible, II: diffuse hyperpigmentation visible, III: Isolated hyperpigmentation visible at the side of the arm, IV: Isolated hyperpigmentation in close proximity to this prick location. Presence of scars assessed by visual inspection: I: no scars visible, II: scar visible at the side of the arm, III: scar visible in close proximity to this prick location. Data are presented as mean with 95% confidence intervals. Between-group comparison by generalised estimating equations model analysis. \*:  $p < 0.05$ ; \*\*:  $p < 0.01$ .

be noted that for now, the AI-assisted readout method can only process images in combination with S.P.A.T. given the need for 32 images captured under various lightning conditions for optimal accuracy of the AI model<sup>(8)</sup>. Strengths of our study relate to the large number of skin reactions analysed and the systematic evaluation of multiple confounders providing a comprehensive real-world picture.

The SPAT AI-assisted readout method showed high accuracy (>90%) in most patients irrespective of hair, hyperpigmentation or scar tissue, reinforcing the robustness of the AI model in real-world conditions. Darker skin tone and the presence of tattoo marks were observed in some patients and lowered performance. A brightness slider has recently been added in the web viewer, allowing better visualisation of images with darker skin tone and subsequent evaluation of the AI-assisted readout. Further evaluation of this method in more diverse patient po-

pulations is needed to confirm external validity. This will lead to more fair and inclusive AI models to support allergy diagnostics so more patients can benefit from these emerging technologies.

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## Authorship contribution

VH, AC, GDG, WL, ALP, and LVG were responsible for recruitment. KR, SFS, DL, RD, and LVG were responsible for data analysis. RD was responsible for the development of the AI algorithm. The study was conceived, designed, set-up, analyzed, and interpreted by KR, SFS, DL, SG and LVG. SFS, DL and LVG have verified the underlying data. All authors accept responsibility for the decision to submit for publication.

**Conflict of interest**

SFS, RD, DL are employees of Hippocreates BV. SFS, RD, DL, SG and LVG hold shares of Hippocreates BV. SFS serves/ed as associate editor for Respiratory Medicine (ongoing) and Heliyon Immunology (2022-2024). AMC reports grants, speaker honoraria, consultancy or advisory fees and/or research support and other, all via Technical University of Munich from Allergopharma, ALK Abello, Astra Zeneca, Bencard/Allergen Therapeutics, GSK, Novartis, Hippo Dx, LETI, Roche, Zeller, Sanofi, Regeneron, Thermo

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## SUPPLEMENTARY MATERIAL

### Materials and Methods

#### Ethics

The study (CIV-23-04-042754) complied with all relevant ethical regulations, was approved by the institutional review boards (Belgium: a Belgian recognized Ethics Committee assigned by the Belgian competent authority; Germany: Ethikkommission der Technischen Universität München) and registered online at [www.clinicaltrials.gov](http://www.clinicaltrials.gov) (NCT05918354).

#### Accuracy evaluation of the AI algorithm

Images of 217 patients (2,604 wheals) of the validation cohort recruited in the previous pivotal study (NCT05918354) were analysed. The AI measurements of the longest wheal diameter were compared with the measurements by the treating physician. A confusion matrix (2x2 table) was created with binary values (true: wheal  $\geq$  4.5mm or false: wheal  $<$  4.5mm) of the AI and physician measurements per wheal for each of the pre-defined patient subgroups (based on the confounder categories, see Table S1). Accuracy was determined for each the confusion matrices.

#### Quantitative & qualitative scoring of the images

The images of the forearms were analysed for skin tone, presen-

ce of hair, scar tissue, tattoos and hyperpigmentation. Analysis of skin tone was performed quantitatively using the individual typology angle (ITA) method per patient <sup>(7)</sup>. Other possible confounders were assessed by visual inspection of the images per prick location and applying a qualitative score by 2 observers separately. In case of mismatch between the 2 observers, these images were evaluated together after which consensus was reached.

#### Statistical analysis

To analyze differences in accuracy between groups, we fitted a generalized estimating equations (GEE) model with a binomial distribution and logit link function. This approach accounts for the non-independence of repeated observations within clusters (i.e. the patients). Group was included as a categorical predictor, with group I (no visible signs of the confounder) serving as the reference category, except for skin tone where the largest group (group III) was selected as reference category to account for bias based on skin tone. Patients with a tattoo (group II-III) were excluded from the skin tone analysis to avoid bias because of the tattoo colour in the ITA analysis. Accuracy with 95% confidence intervals were reported in the graphs. All analyses were conducted in Python using the statsmodels package.

Table S1. Applied measurements for the confounder analysis of the AI-assisted readout method.

| Confounders              | Measurement                                                                                                     | Number of pricks | Percentage |
|--------------------------|-----------------------------------------------------------------------------------------------------------------|------------------|------------|
| <b>Skin tone</b>         | <i>ITA (Individual Typology Angle) score</i>                                                                    |                  |            |
| I                        | >50                                                                                                             | 0                | 0.0%       |
| II                       | 25 and 50                                                                                                       | 672              | 26.9%      |
| III                      | 0 and 25                                                                                                        | 1404             | 56.3%      |
| IV                       | -25 and 0                                                                                                       | 324              | 13.0%      |
| V                        | -50 and -25                                                                                                     | 60               | 2.4%       |
| VI                       | < -50                                                                                                           | 36               | 1.4%       |
| <b>Hair</b>              | <i>Qualitative score</i>                                                                                        |                  |            |
| I                        | No hair visible                                                                                                 | 1421             | 54.6%      |
| II                       | Hair visible at the side of the arm (not in close proximity to this prick location)                             | 391              | 15.0%      |
| III                      | Minimal hair visible in close proximity to this prick location                                                  | 380              | 14.6%      |
| IV                       | Abundant hair visible in close proximity to this prick location                                                 | 412              | 15.8%      |
| <b>Hyperpigmentation</b> | <i>Qualitative score</i>                                                                                        |                  |            |
| I                        | No hyperpigmentation visible                                                                                    | 2202             | 84.6%      |
| II                       | Diffuse hyperpigmentation visible (and no isolated hyperpigmentation in close proximity to this prick location) | 233              | 8.9%       |
| III                      | Isolated hyperpigmentation visible at the side of the arm                                                       | 79               | 3.0%       |
| IV                       | Isolated hyperpigmentation in close proximity to this prick location                                            | 90               | 3.5%       |
| <b>Scar</b>              | <i>Qualitative score</i>                                                                                        |                  |            |
| I                        | No scars visible                                                                                                | 2571             | 98.7%      |
| II                       | Scar visible at the side of the arm (not in close proximity to this prick location)                             | 6                | 0.2%       |
| III                      | Scar visible in close proximity to this prick location                                                          | 27               | 1.0%       |
| <b>Tattoo</b>            | <i>Qualitative score</i>                                                                                        |                  |            |
| I                        | No tattoos visible                                                                                              | 2551             | 98.0%      |
| II                       | Tattoo visible at the side of the arm (not in close proximity to this prick location)                           | 13               | 0.5%       |
| III                      | Tattoo visible in close proximity to this prick location                                                        | 40               | 1.5%       |

Skin tone was assessed by individual typology angle (ITA) analysis per patient. The presence of hair, hyperpigmentation, scars and tattoos was assessed qualitatively per prick location.

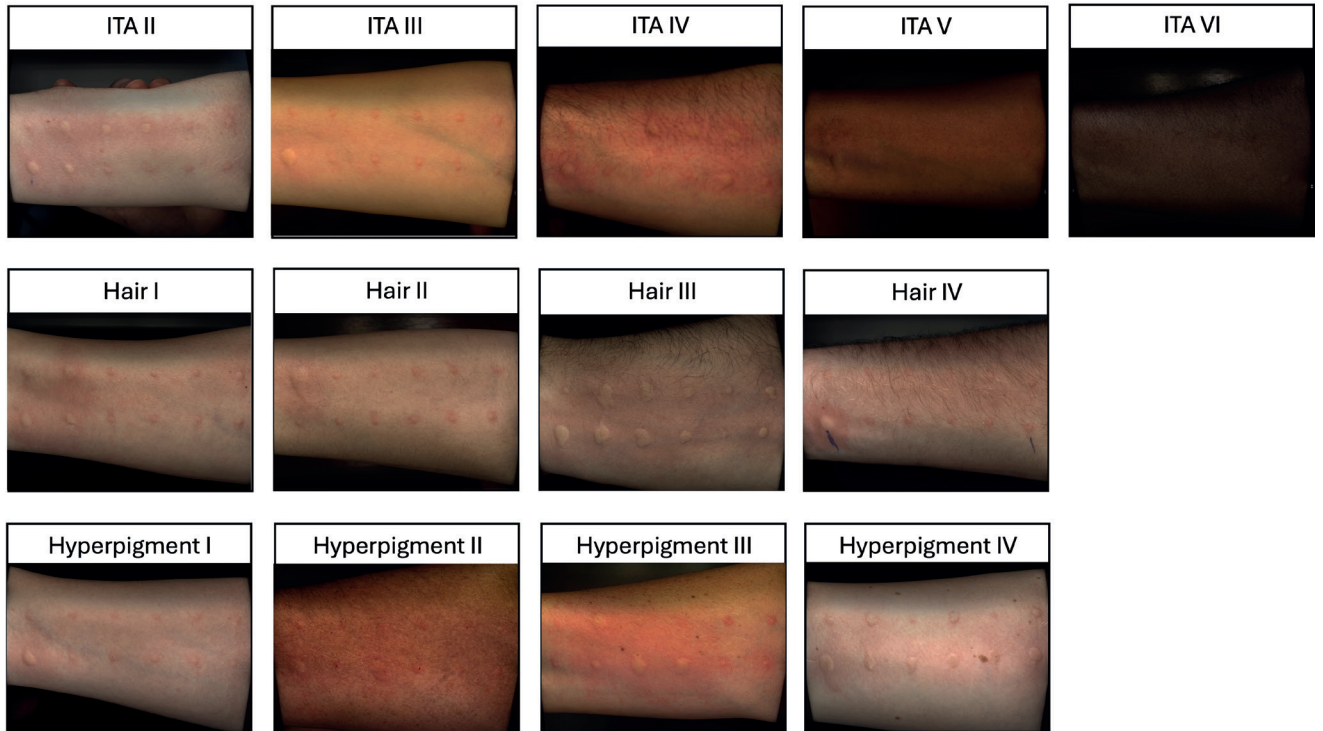


Figure S1. Representative images of patient's forearms per skin tone, hair and hyperpigmentation class. Skin tone was assessed by individual typology angles (ITA) analysis per patient. The presence of hair or hyperpigmentation was assessed qualitatively per prick location.



Figure S2. Representative images of patient's forearms per scar and tattoo class. The presence of scars or tattoos was assessed qualitatively per prick location.