Evaluation of Methods for the Estimation of Threshold Concentrations by the Skin Prick Test

Sten Dreborg\textsuperscript{a,b} Margareta Holgersson\textsuperscript{b}

\textsuperscript{a}Department of Women’s and Children’s Health, Department of Pediatric Allergology, Uppsala University Hospital, and \textsuperscript{b}Pharmacia Diagnostics AB, Uppsala, Sweden

Key Words
Skin prick test · Allergen · Histamine · Dose response · Parallel line bioassay · Threshold concentration · Proficiency test

Abstract
Background: The allergen dose-response curve is flat; thus, small changes in wheal size reflect large differences in skin sensitivity. The sensitivity as measured by provocation tests is given by the threshold concentration that causes symptoms and/or objective signs. The threshold concentrations differ by several magnitudes between the most and the least sensitive individuals clinically allergic to the same allergen. Variation in technique can be minimized by relating allergen responses to that to histamine. The aim here is to present and validate simple methods for estimation of the skin sensitivity given as the concentration inducing a wheal of the same size as that with the positive reference, 10 mg/ml of histamine HCl, in the same patient. Methods: Data from previously reported trials on the biological equilibration of allergen extracts were used to document a method to calculate the concentration of allergen required to induce a wheal of the same size as that with 10 mg/ml of histamine dihydrochloride in the same patient, and to validate the methods using the parallel line bioassay as the gold standard. Results: The validated methods correlated well with the results obtained using the gold standard method and provide results of skin prick testing based on threshold concentrations of allergen. Conclusions: The validated methods reduce the error of differences in testing techniques and make it possible to report skin sensitivity at threshold concentrations. A simple method to be used in clinical practice and a method suitable to describe changes in skin reactivity over time or during treatment are proposed.

Introduction

Until the skin prick test (SPT) was accepted in Europe as the method for routine diagnosis of sensitization and skin reactivity, most allergists used the endpoint concentration by intradermal skin titration as a measure of skin reactivity. However, from the 1970s, manufacturers provided extracts for the SPT in one concentration, i.e. for use in clinical practice, and even scientific reports did not include endpoint concentrations but rather wheal sizes, using one and the same concentration of allergen in all patients. Thus, the results of SPT’s
have been reported as the wheal diameter in millimeters, the wheal area in millimeters squared, or according to the semi-quantitative +++ system devised by Aas and Belin [1]. Thus, SPT results are not comparable to results from bronchial, oral, nasal or conjunctival provocation tests, where results are reported as threshold concentrations.

In a random sample, threshold concentrations differ more than 4-fold between the least and most sensitive patients. Threshold concentrations are log distributed [2–4]. The flat allergen wheal dose response means that a 10-fold increase in skin sensitivity corresponds roughly to an increase in wheal diameter from 3 to 4.65 mm, or from 4.65 to 7.2 mm in diameter, etc. [5]. Thus, wheals 3 and 11 mm in diameter represent a 1,000-fold difference in sensitivity.

Since 1973, the simple +++ method of Aas and Belin [1] comparing the allergen wheal size in relation to that with histamine has been widely used. Histamine reactivity represents the general skin reactivity that is correlated to the degree of sensitization [6]. However, the method is semiquantitative and has never been thoroughly documented.

Adjusting the allergen skin test with the histamine reactivity in the same patient eliminates the influence of differences in SPT technique [7]. Thus, it would be of value to be able to find a simple, practically applicable method to equalize the allergen wheal response between testing personnel. Furthermore, it would be of value to be able to express the response to allergen as a threshold concentration, comparable to provocation tests in other organs [2–4].

The aim of the present study is to validate simplified methods of estimating skin sensitivity to allergens in relation to that to histamine. These results are compared to those using the gold standard estimation of the concentration of allergen comparable to that of histamine, the parallel line bioassay. Finally, we wish to present some simple methods available for allergy practices and clinical research.

Materials and Methods

Patients

The patient samples investigated have been presented previously [4, 8]. Briefly, 36 wall pellitory-sensitive patients [8] and 26 mite-sensitive patients [4] from previously published papers on the biological equilibration of allergen extracts were included consecutively, i.e. without any selection. The mean age of the patients sensitive to wall pellitory (Parietaria officinalis) was 31.5 years (range 15–49), and that of the mite-sensitive patients was 32 years (range 17–45). The inclusion criteria were a clinical history of wall pellitory or possible mite respiratory allergy; a positive SPT with wall pellitory extract (P. officinalis and P. judaica) or Dermatophagoides pteronyssinus and D. farinae allergen extracts, respectively; a histamine wheal (1 mg/ml) at least ≥4 mm in diameter; never having received allergen-specific immunotherapy; the allergen investigated being an adequate problem in the region; being aged 15 years or older, and not having taken antiallergic drugs for a period adequate to avoid an influence on the skin test results [9]. The local ethics committees approved the original trials [4, 8].

Test Solutions

Freeze-dried, standardized allergen preparations of wall pellitory (P. officinalis; batch D 821216/2, 18% protein) [8] and house dust mite (D. pteronyssinus; batch D 820421/2, 44% protein) [4] were used (in-house references; Pharmacia Diagnostics AB, Uppsala, Sweden). The extracts were purified by removal of low-molecular-weight substances by ultrafiltration. The freeze-dried extracts were reconstituted on the day of testing with Albumin Diluent® (0.03% human serum albumin and 0.4% phenol in saline; Pharmacia Diagnostics AB). Histamine dihydrochloride 1 mg/ml (5.43 mmol/l or 0.63 mg/ml histamine base; Pharmacia Diagnostics AB) and histamine dihydrochloride 10 mg/ml (54.3 mmol/l or 6.3 mg/ml histamine base, Pharmacia Diagnostics AB) were used. The potency of the extracts is given in dry weight per millilitre, which is the most exact method for dispensing allergen extracts for final production.

SPT Method

The method described by Østerballe and Weeke [10] was employed, using a steel lancet with a 1-mm tip and shoulders to prevent the lancet from further penetration of the skin. The tip was pressed at a 90-degree angle against the skin surface through a drop of allergen or histamine, applying the same pressure each time. One lancet was used per test. After completion of the test, the test solution remaining on the skin surface was removed by pressing a soft tissue against the skin. In principle, the criteria set up in the EAACI position paper on skin testing [11] were followed: after 15 min, wheals were encircled on the surrounding erythema, the drawing was transferred by means of a translucent tape to a record sheet, and the area was estimated by a digitizer.

Original Test for Biological Equilibration

The original trials were performed as part of a pan-European effort to document the equilibration of the total allergenic potency of common inhalant allergen extracts. To obtain similar samples of patients, the above inclusion criteria were followed in all such published trials [3, 4, 8, 12–14].

Design

First, a preliminary test was performed on the volar aspect of the forearm to decide upon the three concentrations to be used in the final test on the back. Due to the poor precision of the SPT [4], several parallel tests with the same concentration were needed to define the response to each concentration and to estimate the dose-response relationship by parallel line bioassay within each patient. The dose response was best fitted to the model: log D = A + a + b log C, within the normal range of wheal sizes obtained by the SPT, where A is area, D is diameter and C is concentration. The allergen
concentrations causing wheals similar in size to that caused by 10 mg/ml of histamine HCl (C_{H10}) in the same patient, and concentrations 10 times lower and higher than that, were selected for the final test on the back [4, 8, 15].

**Definite Test on the Back**

The concentration of allergen inducing a wheal reaction similar to that with C_{H10}, the allergen dilutions 10 and 100 times stronger as well as C_{H1} and C_{H10} were all tested in quadruplicate, allocated on the back in a mirrored pattern (fig. 1c).

**Methods for Evaluation in This Study**

The Published Method for Biological Equilibration

The geometric mean of the four tests with C_{H10} and each of the three individually selected concentrations of allergen were used for further calculations in the published reports [4].

In preliminary trials [12, 14, 16], the dose response of allergen skin test reactivity within the response level used for the SPT had been found to be best fitted to the log/log model: \( \log A = a + b \log C \).

The four tests with C_{H10} and the four allergen tests were performed with each concentration of allergen allocated over both sides of the back, in a mirrored, upside-down manner. The allergen concentration eliciting a wheal of the same size as that with C_{H10} in the individual patient was estimated by parallel line bioassay (fig. 1c).

**Fig. 1.** Original test procedure [4, 15]. The pretest on the forearm is performed using duplicate tests with the three lowest of six 10-fold concentrations. The horizontal line denotes the mean wheal diameter obtained with 1 mg/ml of histamine in the same patient (D_{H1}) or the mean wheal area obtained with 1 mg/ml of histamine in the same patient (A_{H1}). The slope represents the allergen dose response, and the rings the mean wheal responses to the three lowest concentrations of the test solution, respectively. The filled circles indicate the concentration (C) eliciting a mean wheal diameter (D_A) or mean wheal area obtained with the same allergen concentration in the same patient (A_A) of similar size as that with D_{H1} or A_{H1}, and the rectangles indicate the three concentrations chosen for the final test on the back. a The D_A or A_A elicited by the middle C_A tested was of similar size as that with D_{H1} or A_{H1} and, therefore, was chosen as the lowest C_A tested in the final test. b The D_A or A_A elicited by the highest C_A tested was closest to the size of that with D_{H1} or A_{H1} and, therefore, this and two higher allergen concentrations were tested in the final test on the back. c The C_A chosen in the pretest and two higher concentrations were tested in quadruplicate on the back, i.e. 12 SPTs with allergen 4 with C_{H10} in total. The concentration eliciting a wheal of the same size as that with C_{H10} in the same patient at the same time is calculated according to the model \( 10 \log A = a + 10 \log C \).
In this study, we used either four tests obtained with the three selected allergen concentrations or one test per concentration, selected at random, to estimate $C_{H10}$ in the individual patient, using the model: $\log A = a + b \log C$. The results are given in micrograms dry weight per milliliter, the geometric mean, the 95% CI and the range.

The method using four parallel tests that was originally published in 1987 [4] was later adopted by the Nordic Council on Medicines and included in an appendix of the second edition of the Nordic guidelines on allergen standardization [15].

The Original Method of Björkstén et al.

Originally, Björkstén et al. [16] used a single test with one concentration of allergen and histamine of 1 mg/ml, respectively. In a preliminary study, they found the best-fitted model was a log-log model: $\log D = a + b \log C$.

In this study, we used the same model but chose the concentration with which most patients had been tested in the original trial [4]. Since we had calculated the areas, the model used was: $\log A = a + b \log C$.

The method of Björkstén et al. [16] and its principles are shown in figure 2a. The proposed method for clinical use is available in online supplementary Appendix 1 (for all online suppl. material, see www.karger.com/doi/10.1159/000366203).

The Modified Method of Björkstén et al.

Either one or four tests with one concentration of three tested, with a (mean) wheal size most similar to that with $C_{H10}$, were used for estimating the $C_{H10}$ using the area of the wheal(s) and the log-log model (see fig. 2b and online supplementary Appendix 2 for further details):

$$C_{H10} = [Ah/Aa]^{2.5} \times C_A,$$

where $C_A$ is the concentration of allergen. In both these simplified methods, when using a single test concentration, the test utilized was chosen at random by noninvolved personnel among the four allergen tests.

Proposed Single-Concentration Methods

The proposed methods presented are as follows: (1) in allergy practices, use the diagnostic allergen extract normally used in the practice, i.e. a single fixed concentration of allergen and $C_{H10}$ (online suppl. Appendix 1), and (2) in scientific work, a limited titration with 10-fold concentrations of allergen and $C_{H10}$ (online suppl. Appendix 2).

Results

All patients fulfilling the inclusion criteria used in this paper who participated in the original wall pellitory and mite trials, which aimed at determining the biological activity of partly purified wall pellitory (P. officinalis) and mite (D. pteronyssinus) allergen extracts, were included in the evaluations. The results of the wall pellitory and mite individual $C_{H10}$ and median $C_{H10}$ concentrations using the three different methods are shown in figure 3a and b. The 95% CI of the reference parallel line bioassay method overlapped that of the unmodified and modified methods of Björkstén et al. [16], using one or four tests. Thus, all showed similar $C_{H10}$ and geometric means as the parallel line bioassay, i.e. the gold standard. The concentration of allergen eliciting a wheal of the same size as that with $C_{H10}$ can therefore be estimated by using one concentration of histamine and one concentration of allergen.

Discussion

It has been shown that relating the allergen SPT wheal response to that to histamine in the same patient reduces the main drawback of the SPT method, i.e. the difference in technique between testing personnel. Thus, the significance of the technique used by the technician is largely eliminated [7].
This paper shows that a formula can be derived by using the common slope of the log-log allergen dose-response relationship: \[ 10 \log D or A = a + b 10 \log(C) \]. The concentration eliciting a wheal the same size as that with 10 mg/ml of histamine dihydrochloride can be calculated by: 
\[
\frac{Dh}{Da} \times \frac{1}{b} \times C \text{ used}, \quad \text{where } b = 0.20 \text{ for the diameter or } 0.4 \text{ for the area. Therefore, the index (1/0.2) is equal to } 5, \text{ or simply } \frac{Dh}{Da} \times 5 \times C \text{ used, or for the area (1/0.4), it can alternatively be presented as } \frac{Dh}{Da}^{2.5} \times C. \]
Using the mean b found \[4\], 0.387 (95% CI 0.371–0.403) for the allergen and 0.334 (95% CI 0.332–0.357) for histamine, respectively, the index when calculating with the area should be 2.58 and 2.99, respectively. Strictly, the formulas shown are not valid, since the dose-response curves are not totally parallel. However, at least when using allergen wheals of a similar size as that with histamine, the above approximation can be accepted, as has been shown in this paper.

Of the two simple methods, the original method proposed by Björkstén et al. \[16\] (online suppl. Appendix 1) is recommended for use in clinical practice. In scientific work, the alternative method (online suppl. Appendix 2) may be preferred, despite the fact that the results do not differ when the two methods are applied to groups.

The proposed methods express skin reactivity in the form of threshold concentrations. The threshold concentration is a better indicator of reactivity than the diameter or area when trying to ascertain the skin response. It is also more suitable when comparing skin response to that of other organs, i.e. the conjunctiva, the nasal mucosa, bronchi and the gastrointestinal tract, the reactivities of which are always reported as threshold concentrations.

The benefit from using the methods described in the appendices is that the skin response is expressed as the threshold concentration. Threshold concentrations make it possible to express the skin sensitivity of patients in a concentration of allergen. Differences in threshold concentration are better estimates of the skin sensitivity between patients and within patients over time than are changes in wheal size.

At least duplicate tests should be used to document the coefficient of variation of the SPT in the hands of the testing personnel. In small children, regular proficiency testing and single tests can be accepted. It is essential to use a proficiency test program, thus assuring constant, reproducible results \[17\].

The Methods Proposed

The principles behind the methods that have been proposed in this paper can be summarized as follows:
• differences in technique should be compensated for by referring the allergen response to that to histamine;
• skin reactivity should be expressed as a threshold concentration, making it possible to express differences in skin reactivity between centers and within patients over time in terms of allergen concentration;
• consequently, comparison of skin test reactivity between centers in, for example, epidemiological stud-
ies and multicenter therapeutic studies becomes possible.
These methods will be validated in a forthcoming paper in relation to other methods in their ability to evaluate changes in skin sensitivity during treatment [Dreborg et al., submitted].

Acknowledgements

My sincere thanks to Dr. Antonio Basomba (Valencia), to Profs. Barry Kay, Tak Lee and Steven Durham (London) as well as to their nurses, who did the original skin testing. Thanks are also due to Margareta Holgersson, PhD, who gave statistical advice, and the statistician Anders Hansson, MSc, who did the computer work.

This work has not been published as an original paper but is a nonpublished part of paper IV in the medical dissertation by Dreborg [3].

Disclosure Statement

No financial support was provided for this publication. The biological standardization/equilibrating trials and the statistical work done 25 years ago was supported by Pharmacia Diagnostics AB, Uppsala, Sweden.

References

3 Dreborg S: The skin prick test: methodological studies and clinical applications; Linköping University Medical Diss No 239, 1987.
7 Dreborg S: Allergen skin prick test should be adjusted by the histamine reactivity. Int Arch Allergy Immunol 2015;166:77–80.