Insect Sting Reactions and Specific IgE to Venom and Major Allergens in a General Population

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Abstract

Background: Insect sting reactions are frequently reported, but population studies documenting the frequency and the relation to IgE-sensitization and serum tryptase are scarce.

Methods: Questionnaire data and results from measurements of specific IgE against venom, major allergens and cross-reacting carbohydrate determinants (CCDs) were collected from 2,090 adult participants in a cross-sectional survey.

Results: 13\% of the population reported symptoms of sting reactions and about half were systemic in nature. In all, 15\% were sensitized to venom but only 31\% of these had reacted to stings and only 38\% of those with reactions had IgE to venom. In addition, 12\% with IgE to venom were double-sensitized (DS), i.e. to both bee and wasp venom. Among DS IgE to major venom allergens, rApi m 1, rVes v 1 and rVes v 5 were negative and of no help in 31\%, but 59\% could be identified as likely sensitized to bee or wasp. IgE to CCDs occurred in only 0.7\%, but 80\% of these were DS. Finally, 36\% with IgE to CCDs had had symptoms, mostly local. Serum tryptase was not associated with a history of sting reactions.

Conclusions: In a temperate climate, self-reported insect sting reactions and sensitization to venom are frequent, but in most cases, these are not seen in the same individual. In DS individuals, measurements of IgE to major allergens can be helpful in some but not all cases and additional analyses are needed. IgE to CCDs may have some clinical relevance.

Key Words
Cross reactions · Hymenoptera venom · Insect sting allergy · Specific IgE · Tryptase

Introduction

Most people are exposed to insect stings several times during their lifetime. In the European temperate climate, wasps and honeybees from the Hymenoptera order are often the offenders. While the majority of stings just result in pain and limited swelling, some individuals get large local swellings, and in a small number, more severe systemic reactions occur. The prevalence of such reactions in the general population is only scarcely described [1, 2].

A demonstration of sensitization to insect venom is needed to make a proper allergy diagnosis, i.e. for the documentation of an allergic mechanism and to safely iden-
tify the offending insect. Such identification is crucial when allergen-specific immunotherapy is considered. Allergenic cross-reactivity with or without clinical relevance is a major challenge in this context [3]. Elevated serum tryptase has been associated with more severe insect sting reactions [4, 5], but it is unknown if it is a risk factor for such reactions in the general population.

The aim of this study was to estimate the prevalence of insect sting reactions in a general population based on questionnaires. In addition, the study aims to compare the results with levels of specific IgE to venoms, to venom major allergens and to cross-reacting carbohydrate determinants (CCDs) in the same individuals, focusing on clinical and immunological cross-reactivity and the influence of tryptase level.

**Material and Methods**

**Study Design**

The study was part of a cross-sectional population survey, including serum-specific IgE and tryptase determinations and structured questionnaires. It was based on the five-year follow-up of the Health2006 cohort. A detailed description of the baseline examination has been published elsewhere [6]. The participants in the baseline Health2006 cohort were drawn as a random sample from the background population aged 18–69 years, living in 11 municipalities in suburban Copenhagen. A total of 3,471 individuals (44.7% of the sample) entered the study and participated in the health examinations, which took place between June 2006 and June 2008. In 2011–2012, participants in the baseline Health2006 were invited for a 5-year follow-up examination, including essentially the same study protocol. A total of 3,405 were eligible for invitation (21 had emigrated and 45 had died). A total of 2,308 (45.8% men) agreed to participate and were re-examined between November 2011 and November 2012 [7]. The median age was 55.7 (range 24–76) years. For 88 participants, the serum sample left for analysis was not sufficient for all of the planned in vitro measurements and for a further 30, a valid questionnaire was not available. Thus, data from both questionnaires and the in vitro measurements were available for 2,090 individuals. In this group, the median age of 55.7 years (range 24–76 years) and gender (46.0% men) did not differ significantly from the total participating group.

**Structured Questionnaires**

The structured questionnaire was self-administered and included questions on various health, lifestyle and socioeconomic aspects. The insect-related questions were 'Did you ever have a swelling of more than 10 cm from side to side after a sting from a wasp or bee? (Yes/No)', 'Did you ever have an itching rash on larger areas of the body after a sting from a wasp or bee? (Yes/No)', 'Did you ever feel malaise with the need to lie down or with breathing problems after a sting from a wasp or bee? (Yes/No)' and 'If you had any of these symptoms, which insect(s) was/were responsible: wasp? (Yes/No), bee? (yes/no), unknown insect? (Yes/No)?'. Having had an insect sting reaction was defined as being able to confirm at least one of the symptoms mentioned above.

**Serum Specific IgE**

Sera were stored frozen until tested. All sera were tested for IgE to honeybee venom (ii), venom from wasp species (i3), the recombinant major allergens rVes v 1 (i211), rVes v 5 (i209) and rApi m 1 (i208) if IgE to venom was ≥0.10 kU A /l, the isolated glycan part from bromelain (o214), as a CCD marker, and a screening mixture of inhalant allergens (Phadiatop®) using the commercially available UniCAP 250 system (Phadia Diagnostics, Uppsala, Sweden, now Thermo Fisher Scientific). Serum tryptase was measured with an ImmunoCAP® tryptase assay, which determines total tryptase, i.e. pro-forms and mature forms of α- and β-tryptase. In a study by the manufacturer, the 95th percentile was 11.4 µg/l in 126 healthy individuals (aged 12–61 years; 48% males). This value has been adopted as the upper reference level in several insect allergy studies [4, 8].

**Ethical Issues**

All individuals gave written consent to participate in the studies, which were approved by the corresponding Institutional Review Boards (Ethics Committee of the Capital Region of Denmark, code H-3–2011–081), and conformed to the current Helsinki Declaration.

**Statistical Analyses**

The χ2 and the Fisher’s exact tests were, where appropriate, used to investigate the association between categorical variables. Logistic regression without adjustment was applied to assess the odds ratio (OR) for the outcome between exposure groups.

**Results**

**Questionnaire Data**

In all, 13% (n = 272) of the population had experienced reactions to stings (table 1). The majority reported large local reactions (51% were large local reactions only), while about one third reported rash and one third reported malaise. The pattern of reactions was similar for bee and wasp stings.

More individuals reported reactions to wasp stings (n = 203; 9%) than to stings from bees (n = 83; 4%) and unknown insects (n = 38; 2%). Finally, 2.5% (n = 54) reported reactions to both bees and wasps.

**in vitro Data**

In parallel to the reported clinical reaction pattern, more individuals had IgE ≥0.35 kU A /l to wasp than to bee venom (13.6 and 3.3%, respectively), with a range <0.35–67.5 and <0.35–15.8 kU A /l, respectively. IgE to both bee and wasp venom was seen in 1.8% and 84.9% had no IgE to the venoms.

A total of 30% of the population had specific IgE ≥0.35 kU A /l to airborne allergens (Phadiatop). This was significantly associated with being sensitized (i.e. having specific IgE) to insect venom: 67.1% of the bee-sensitized

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[OR 5.4; 95% confidence interval (CI) 3.3–8.9] and 47.2% of the wasp sensitized were sensitized to airborne allergens as well (OR 2.6; 95% CI 2.0–3.3; \( p < 0.0001 \)).

Only 14% (10/73) of the individuals sensitized to bee venom had IgE to the major allergen (rApi m 1) \( \geq 0.35 \text{kU} \text{A/l} \). In wasp venom, 35% of those sensitized to the venom (103/303) had IgE to rVes v 1, 63% (186/303) to rVes v 5 and 82% (248/303) to at least one of these. IgE to CCDs could partly explain the discrepancies since this IgE occurred in 19% (12/63), with IgE to bee venom but not to the major allergens, and in 11% (6/55) to wasp venom but not to the major allergens, compared with 10% (1/10) and 2.4% (6/248) in individuals with IgE to venom as well as to major allergens. In addition, 12% (40/336) of the individuals with positive sera were positive to both venoms, i.e. they were double-sensitized (DS). Having IgE to one venom increased the risk of having IgE to the other (OR 8.7; 95% CI 5.4–14.0). About half of the DS had IgE to major allergens from one of the species only, indicating either bee or wasp as the primary sensitizer (table 2). In a third of the DS, no IgE to the major allergens was found and the diagnosis was left open. In contrast to IgE against bee and wasp venom in the population, IgE against the major bee and wasp allergens in the DS did not correlate.

The occurrence of specific IgE to common CCDs in the venom extracts might, at least partly, explain the double sensitization, since 80% with IgE to CCDs were DS compared to only 1.3% with no IgE (table 3). Additionally, 50% (6/12) of the DS with no IgE to major allergens had IgE to CCDs compared to 22% (6/27) of the DS with IgE to at least one major allergen.

**Questionnaire and in vitro Data**

There was a correlation between sensitization and clinical symptoms; however, only 38% of the individuals reporting abnormal sting symptoms (to any insect) had IgE \( \geq 0.35 \text{kU} \text{A/l} \) to at least one venom (table 4). Excluding individuals with IgE to CCDs would change the frequency to 37%.

Only 31% of the individuals with IgE \( \geq 0.35 \text{kU} \text{A/l} \) to at least one venom reported sting reactions/symptoms (to any insect; table 4). Excluding individuals with IgE to CCDs would change the frequency to 30%.

In addition, 11 of the 14 individuals with specific IgE to CCDs \( \geq 0.35 \text{kU} \text{A/l} \) and clinical data available were DS. Four of these (36%) reported sting reactions. This reaction rate did not differ from that of individuals sensitized to at least one insect venom. All 4 had large local swellings and 1 had malaise and generalized rash in addition to the swelling. Two people with IgE to CCDs, but not to venom, had no symptoms related to the stings.

Among individuals with low IgE to venom (0.20–0.35 kU A/l) and reporting either generalized rash and/or malaise, 3 of 71 (4%) had IgE \( \geq 0.35 \text{kU} \text{A/l} \) to the major wasp allergens. A similar ability to identify individuals sensi-

### Table 1. Questionnaire data based on answers from 272 of the 2,173 individuals (13%) who reported abnormal reactions to insect stings

<table>
<thead>
<tr>
<th>Stinging insect</th>
<th>Reaction, n (%)</th>
<th>large local reaction (only)</th>
<th>generalized rash (without malaise)</th>
<th>malaise</th>
<th>systemic reaction (rash or/and malaise)</th>
<th>at least 1 reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wasp</td>
<td>104 (51)</td>
<td>58 (29)</td>
<td>68 (34)</td>
<td>99 (49)</td>
<td>203 (100)</td>
<td></td>
</tr>
<tr>
<td>Bee</td>
<td>43 (52)</td>
<td>23 (28)</td>
<td>28 (34)</td>
<td>40 (48)</td>
<td>83 (100)</td>
<td></td>
</tr>
<tr>
<td>Unknown</td>
<td>19 (50)</td>
<td>11 (29)</td>
<td>11 (29)</td>
<td>19 (50)</td>
<td>38 (100)</td>
<td></td>
</tr>
</tbody>
</table>

1 More than one insect could be recorded for each individual.

### Table 2. Specific IgE to venom major allergens from bee (rApi m 1) and wasp (rVes v 1 and rVes v 5) in individuals with IgE to both bee and wasp venom, i.e. showing double sensitization

<table>
<thead>
<tr>
<th>rVes v 1 and/or rVes v 5</th>
<th>rApi m 1</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Positive1</td>
<td>4 (10%)</td>
<td>21 (54%)</td>
</tr>
<tr>
<td>Negative</td>
<td>2 (5%)</td>
<td>12 (31%)</td>
</tr>
<tr>
<td>Total</td>
<td>6 (15%)</td>
<td>33 (85%)</td>
</tr>
</tbody>
</table>

1 Specific IgE \( \geq 0.35 \text{kU} \text{A/l} \). \( p = 1.000 \), Fisher’s exact test.
tized but with IgE to venom below the detection limit was

not seen for the major bee allergen IgE assay.

Median serum tryptase was 4.8 μg/l (95% CI 2.1–11.7
μg/l). In total, 5.4% of the population had serum tryptase
above the reference limit of 11.4 μg/l. Elevated levels did
not occur more frequently in individuals with sting reac-
tions (11/246; 4.5%), not even malaise (4/82; 4.9%) com-
pared to nonreactors (102/1,844; 5.5%; n.s). For each type
of reaction and for no reactions, serum tryptase concen-
trations did not differ when comparing individuals with
or without specific IgE to venom.

Discussion

A prevalence of insect sting allergies of up to 43% has
been reported in selected populations and probably varies
with exposure, as determined by climate and activities [3, 9]. Few general population studies have been performed.
An Austrian study [2] found large local reactions in 4.6%
and systemic reactions in 3.3%, i.e. only about half the
prevalence in our larger study. The methodology using
telephone interviews versus administered questionnaires
with only a few questions might explain some of the dif-
fferences. Systemic reactions in our study were either
‘malaise’ and/or ‘itching rash covering large parts of the
body’ and, most likely, ‘malaise’ would include some va-
sovagal and severe pain reactions or hyperventilation in
addition to anaphylactic reactions. A large study on an
adolescent subtropical population reported figures of the
same magnitude as our study [10]. The explanation might
be that a lower exposure level in our colder climate com-
pensated for the longer exposure time in our adult popu-
lation with an older median age. A smaller rural area
study on adults [11] and a larger study among industrial
workers [12] in another subtropical area only found sys-
temic reactions in 2.3%. However, in the rural study, the
proportion sensitized in the symptomatic group was only
slightly higher than in our study.

We measured only IgE levels to wasp and honeybee
venoms in this study. In northern Europe, the majority
of stings encountered by humans would be from wasps
(\textit{Vespula} sp.) and honeybees (\textit{Apis mellifera}). Other bees,
European hornets (\textit{Vespa crabo}) and paper wasps
(\textit{Polistes} sp.) are responsible for only a relatively small
number of stings in humans in this area, and it is un-
likely that stings with venom not tested for would be a
major explanation for reactions in individuals with no
IgE to venom. The reporting of lifetime prevalence in an
adult population implies recall bias, both with respect to
the type of stinging insect and to the symptoms, e.g. a
painful reaction a long time ago could be remembered as
larger than it actually was. Another major explanation

<table>
<thead>
<tr>
<th>Table 3. Specific IgE to CCDs in individuals with and without specific IgE to both bee and wasp venom ≥0.35 kUA/l</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific IgE to CCDs ≥0.35 kUA/l</td>
</tr>
<tr>
<td>---------------------------------</td>
</tr>
<tr>
<td>Positive (n = 15)</td>
</tr>
<tr>
<td>Negative (n = 2,205)</td>
</tr>
<tr>
<td>Total</td>
</tr>
<tr>
<td>Fisher’s test</td>
</tr>
<tr>
<td>OR (95% CI)</td>
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<table>
<thead>
<tr>
<th>Table 4. Specific IgE to venom in individuals with or without possible allergic reactions to stinging insects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific IgE to venom and/or bee total</td>
</tr>
<tr>
<td>Positive1 must have IgE ≥0.35 kUA/l</td>
</tr>
<tr>
<td>Yes</td>
</tr>
<tr>
<td>No</td>
</tr>
<tr>
<td>Total</td>
</tr>
</tbody>
</table>

p < 0.0001, \( \chi^2 \) test; OR 4.5 (95% CI 3.4–6.0).

1 Specific IgE ≥0.35 kUA/l.
2 Large local reaction, generalized rash or malaise.
IgE to major wasp allergens could document sensitization in (a few) individuals with systemic symptoms, but with IgE to wasp venom below the normal detection value. This was not the case for IgE to the major bee allergen, supporting the finding that IgE to rApi m 1 has a lower level of sensitivity [23–25].

It has been shown that insect allergic patients with IgE to CCDs have an increased risk of being misdiagnosed as having pollen sensitization [26], but in general, atopy is not regarded as a major risk factor for insect allergy [1, 12]. However, our population study with an OR of 2.6–5.4 for sensitization to wasp and bee stings could potentially be used to modify the concept. It is unlikely that sensitivity to CCDs is responsible for this finding since the prevalence was rather low in our population. An OR of >8 for sensitization to wasp venom if sensitized to bee venom is, however, in favor of common allergenic component(s), i.e. cross-reactivity rather than coincidental sensitization to different allergens. With an OR of >300 for sensitivity to both bee and wasp venom when having IgE to CCDs, CCD structure seems to be a very important candidate for such a common component. The additional observation that several of the DS individuals with IgE to CCDs in our unselected population had, in fact, had symptoms when stung, indicates some clinical relevance of this sensitization, although in many cases it occurs with cosensitization to other venom and pollen (especially grass) allergens [27, 28].

That there was no increased reaction rate to stings in individuals with serum tryptase above the normal range may be surprising, since elevated serum tryptase has been identified as a risk factor for more severe sting reactions in case control studies [5]. However, serum tryptase seems not to be associated with atopy and allergic respiratory disease [29], and patients with indolent mastocytosis without skin lesions have been found to have a high reaction rate to insect stings but a lower baseline serum tryptase than other groups of mastocytosis patients [30].

In conclusion of this first large general population study on insect allergies in northern Europe, the majority of those sensitized did not report symptoms and the majority who reported having had symptoms were not sensitized. IgE to CCDs occurred less frequently when compared to other regions, but the occurrence was highly linked to double sensitization with and without sting reactions. Our study supported that measurement of IgE to major allergens/components in the venom has an important place in the investigation of DS patients and IgE- (and skin test) negative patients with a convincing clinical history. However, ‘specificity’ and problems with differ-
entiation have been reported [31] and, in a high proportion of the population, IgE to venom was neither accompanied by IgE to CCDs nor to the major allergens tested for, so it is likely that other components (still not identified) of the many allergens in the venom [32] are responsible for the sensitization. There is a need for commercially available tests for IgE to additional venom allergens to better diagnose patients with double sensitization and also to ensure that relevant allergens are present in extracts used for immunotherapy, which has not always been the case [33].

References

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Disclosure Statement
There were no conflicts of interest.


