Safety and efficacy of venom immunotherapy: a real life study

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Abstract

Introduction: Venom immunotherapy (VIT) is recommended as the first-line treatment for patients allergic to Hymenoptera venom.
Aim: To analyze the safety and efficacy of VIT in a real life setting.
Material and methods: One hundred and eighty patients undergoing VIT were studied to evaluate the safety, efficacy, incidence and nature of symptoms after field stings and adverse reactions to VIT.
Results: Significantly more patients were allergic to wasp than bee venom (146 vs. 34, \( p < 0.0001 \)). Early and late side effects were more common during the maintenance (48 patients, 26.7%) than during the induction of VIT (32 patients, 17.8%), were more frequent in patients allergic to bees, and were not associated with angiotensin convertase inhibitors (ACEi) or \( \beta \)-adrenergic antagonists use. Systemic reactions were observed in 4 individuals on wasp VIT (2.7%) and in 6 patients allergic to bees (17.6%). The VIT was efficacious as most patients reported no reactions (50%) or reported only mild local reactions (43.75%) to field stings. The decrease in sIgE at completion of VIT correlated with the dose of vaccine received (\( r = 0.53, p = 0.004 \)). Beekeeping (RR = 29.54, \( p < 0.0001 \)) and female sex (RR = 1.27, \( p = 0.033 \)) were associated with a higher risk of venom allergy.
Conclusions: Venom immunotherapy is highly efficacious and safe as most of the adverse events during the induction and maintenance phase are mild and local. Side effects of VIT are more common in subjects on bee VIT. Beekeeping and female sex are associated with a higher risk of allergy to Hymenoptera venom.

Key words: venom immunotherapy, side effects, bee, wasp, Hymenoptera.

Introduction

Stings by Hymenoptera insects are relatively common in the population and may lead to a range of reactions from mild and local symptoms to life-threatening anaphylaxis. In Europe mainly honeybee (Apis mellifera) and wasp (US nomenclature: yellow jacket, Vespula germanica and vulgaris) are responsible for those incidents [1]. The allergy to Hymenoptera venom affects around 15–30% of the general population (as confirmed by skin prick tests or sIgE) and the frequency of systemic reactions due to Hymenoptera venom allergy varies between 0.35 to 4% [2, 3]. The incidents of fatalities due to stings are estimated at 0.03 to 0.48 deaths per 1 000 000 citizens per year [4]. Venom immunotherapy (VIT) represents an effective causative treatment for these patients preventing further sting-induced anaphylactic reactions. Immunotherapy may be associated with a risk of local and systemic side effects. Adverse reactions during venom immunotherapy are relatively common and are reported in up to 50% of patients, mostly during the build-up phase. Systemic reactions occur in 12% to 30% [5, 6] of subjects undergoing VIT, which results in a common belief that VIT is potentially dangerous. As a consequence, a limited access to this life-saving procedure may be seen. What is striking, a high level of variations (0–46%) and inconsistency in the side-effects frequency is reported in the literature [7–10]. Moreover, there are just a few comprehensive studies dedicated to this topic [11–16].

Aim

Thus, the goal of this study was to analyze safety, efficacy and adherence to treatment in a real-life setting in a group of patients who underwent immunotherapy due to bee or wasp venom allergy. The incidence and nature of symptoms after field stings and adverse reactions to...
VIT were studied. Factors associated with a higher risk of allergy to Hymenoptera venoms were also analyzed.

Material and methods

One hundred and eighty adult patients undergoing VIT at the Department of Internal Medicine, Asthma and Allergy, Medical University of Lodz, Poland, including 146 subjects allergic to the wasp venom and 34 subjects allergic to bee venom, were included into the study. Their medical histories were analyzed retrospectively. Due to the design of the study, no institutional review board approval was required. All patients included in the study were undergoing VIT against Hymenoptera venom (honey bee or wasp venom) due to systemic reactions after stings. The following criteria in the baseline patients’ characteristics were taken into consideration: sex, age, the type of the insect, levels of sIgE, Muller’s scale after stings before VIT, the criterion of VIT termination. Patients were also asked about the association with beekeeping (questions of the beekeeper in the family and in the close neighborhood) and medications taken – antagonists of β-girotensin convertase (ACE) inhibitors and β-adrenergic antagonists.

Venom immunotherapy

Depending on the type of allergy, specific and standardized allergen extracts of bee (Apis mellifera) or wasp (Vespu eruption spp.) venom were used (Pharmalgen®, Alutard®, or Venomenhal®). In majority of subjects but one, an ultra-rush protocol was used at the induction phase of VIT. The vaccine was administered subcutaneously with the increasing doses of: 1, 10, 20, 30 and finally 50 µg in the 30 min’ intervals of time (111 µg in total). After induction patients were receiving a maintenance dose (100 µg) of a vaccine every 4–6 weeks, depending on the tolerance and duration of treatment. All patients undergoing venom immunotherapy at our center received oral antihistamines as a pretreatment before the ultra-rush induction phase and during the maintenance treatment.

Data collected during the course of venom immunotherapy

The dose of the vaccine after each injection, all adverse reactions, wasp or bee-stings during VIT, and symptoms after stings were reported in the medical histories and classified according to the four-point Muller’s scale. Side effects during induction of VIT and during maintenance treatment of VIT were classified as early (occurring up to 30 min after the injection) and late (appearing after 30 min). The early side effects have been registered in the hospital. Patients were obliged to report all the late side effects, potential field stings during VIT and symptoms after stings on the next visit in the clinic. Specific IgE levels at the baseline, during and after VIT were analyzed.

Statistical analysis

To assess the relationship between two independent category groups of data, the χ² test or Fisher exact test were used. For the independent variables, including average VIT time and injections number, summary dose of vaccines, Muller’s median scale and average sIgE titer, the Mann-Whitney and t-test for independent means were used. To compare the decrease in sIgE after completed VIT, Wilcoxon test was used. Analysis of the correlation was made, depending on the type of data, with the Pearson’s (for linear correlation) and Spearman’s correlation (other kinds of correlation). Statistical analysis was performed using GraphPad Prism version 5. The confidence interval was set at 0.95 (95% CI) and the level of significance was set at p < 0.05.

Results

One hundred and eighty patients, aged 15–77 years, were included in the study. Demographic data and baseline characteristics of the study group are presented in Table 1. Comparison between subjects allergic to bee and wasp venom is presented in Table 2. There were significantly more patients undergoing VIT due to allergy to wasp than bee venom (146 vs. 34, p < 0.0001). The majority of patients included in both groups were women (wasps/bees; 61.6%/61.8% of women). Patients allergic to bee venom were significantly younger (44 vs. 50 y, p = 0.042). Average duration of VIT, at the time of the study was 35 months, the average number of injections was 39 and the total dose of vaccine was 2637.75 µg. Five patients allergic to bee venom were beekeepers and twelve had a relationship with beekeeping. The percentage of patients taking β-adrenergic antagonists or inhibitors of ACE was over 20% and was similar in both study cohorts. The adherence to treatment was high as all patients qualified for venom immunotherapy continued the treatment in line with the timelines suggested. Two subjects were referred to other allergy centers due to a change of the residence.

Symptoms after stings for all patients before and during VIT are presented in Figure 1 A and B. All patients qualified for the study presented systemic reactions. Median of Muller’s scale before VIT was 3 and during VIT was significantly lower (44 vs. 50 y, p = 0.042). Average duration of VIT, at the time of the study was 35 months, the average number of injections was 39 and the total dose of vaccine was 2637.75 µg. Five patients allergic to bee venom were beekeepers and twelve had a relationship with beekeeping. The percentage of patients taking β-adrenergic antagonists or inhibitors of ACE was over 20% and was similar in both study cohorts. The adherence to treatment was high as all patients qualified for venom immunotherapy continued the treatment in line with the timelines suggested. Two subjects were referred to other allergy centers due to a change of the residence.

Side effects during venom immunotherapy for all patients

Side effects during VIT were reported by 64 patients (43.8% of all patients). Early and late side effects were more common during the maintenance phase of VIT (48 pa-
Table 1. Demographic data and baseline characteristics of the study group

| Baseline characteristics of the study group | Number of patients | % of group |
|--------------------------------------------|--------------------|------------|
| Number of patients/female/male             | 180/111/69         | 100/62/38  |
| Age [years] (min.–max.)                    | 49 (15–77)         | –          |
| Number of patients allergic to wasp venom/bee venom | 146/34             | 81/19      |
| Number of beekeepers                       | 6                  | 3.33       |
| Number of persons whose husband/wife/neighbors have a bee yard | 12                 | 6.67       |
| Median of Muller’s scale before VIT        | 3                  | –          |
| Average time of VIT [months]*              | 35                 | –          |
| Average injections number/average total dose of vaccine [µg] | 39/2637.75         | –          |
| Scheme of the initial phase of VIT – ultra-rush | 179                | 99.44      |
| Scheme of the initial phase of VIT – cluster other than ultra-rush | 1                  | 0.66       |
| Number of patients on ACE inhibitors or β-adrenergic antagonists during VIT | 37                 | 20.56      |

*Time of observation of the patient during VIT.

Table 2. Baseline characteristics of patients allergic to wasp and bee venom undergoing VIT and subgroup of patients who completed a 5-year cycle of VIT

| Variable                                                                 | Wasp     | Bee     | P-value   |
|--------------------------------------------------------------------------|----------|---------|-----------|
| Patients allergic to wasp and bee venom:                                 |          |         |           |
| Number of patients                                                       | 146      | 34      | < 0.0001  |
| Age [years] (min.–max.)                                                  | 50 (18–77)| 44 (15–71)| 0.042  |
| Females                                                                  | 90 (61.6%)| 21 (61.8%)| 1.00    |
| Average time of VIT [months]                                             | 34       | 40      | 0.449    |
| Average injections number/average total dose of vaccine [µg]             | 37/2576.38| 47/2901.31| 0.152/0.375 |
| Number of beekeepers                                                     | 1 (0.68%)| 5 (14.1%)| 0.0012   |
| Number of persons related to beekeeping                                  | 0        | 12      | < 0.0001 |
| Number of patients taking β-adrenergic antagonists or inhibitors of ACE | 30 (20.55%)| 7 (20.6%)| 0.66     |
| Number of patients stung during VIT                                      | 38 (26%) | 10 (29%)| 0.672    |
| Median of Muller’s scale after stings before VIT/during VIT              | 3/0      | 3/0     | 0.1712/0.933 |
| Median of IgE before VIT (class)/average IgE before VIT [UI/ml]          | 2.55/6.19| 2.5/5.36| 0.887    |
| Patients who completed a 5-year cycle of treatment:                      |          |         |           |
| Number of patients who completed a 5-year cycle of treatment             | 18       | 3       | 0.7692    |
| Average length of VIT in patients who completed VIT [months]             | 69.4     | 94      | 0.0392    |
| Average injections number in persons who completed VIT                   | 60       | 96      | 0.0552    |
| Average total dose of the vaccine in persons who completed VIT [µg]      | 4569.4   | 6046.4  | 0.0294    |
| Median IgE after 5-year treatment of VIT (class)/average IgE after 5-year treatment of VIT [UI/ml] | 2.05/1.36| 2.6/2.53| 0.064     |

*χ² test. **Mann-Whitney U test.
Agnieszka Kołaczek, Dawid Skorupa, Monika Antczak-Marczak, Piotr Kuna, Maciej Kupczyk

During the induction of treatment early side effects were reported by 9.4% of patients and late side effects were reported by 8.3% of patients. During the maintenance phase, early side effects were rare (7.2%), however the late side effects were more frequent (19.4%). Systemic reactions as side effects during the induction and maintenance phase of VIT were observed in 4 individuals undergoing VIT against the wasp (2.7%) and in 6 patients allergic to bee venom (17.65%). Only in 1 case, the administration of adrenaline was needed. Majority of systemic reactions were reported in the induction phase (1.9% of injections), not in the maintenance phase (0.19% of injections). Most of the systemic reactions (68.2%) developed within 30 min after the injection. All late systemic reactions (31.8% of events) were restricted to skin changes (urticaria, flush, reddening of the skin). The frequency and nature of adverse reactions to VIT are presented in Figure 2. Treatment with β-adrenergic antagonists or ACE inhibitors was not associated with any kind of adverse events during VIT (p = 0.773), systemic adverse events during VIT (p = 0.74) or more severe symptoms during field stings (p = 0.804).

Figure 1. Symptoms after stings for all patients before VIT (A), and symptoms after field stings during VIT (B). Severity of reactions described with Muller’s scale for all patients after stings before VIT (C), and after field stings during VIT (D)
Safety and efficacy of venom immunotherapy: a real life study

Comparison of side effects during venom immunotherapy in patients allergic to bee and wasp venom

Early and late side effects during induction of VIT were more frequent in patients allergic to bees than in patients allergic to wasps. Similarly, during maintenance treatment early and late side effects were more common in patients allergic to bees than in patients allergic to wasps (Table 3). Most of the side effects were local and included local swelling, reddening of the skin and itch at the site of injection (Figure 2). Local urticaria was more frequent as an early side effect during induction of VIT and was more common in patients allergic to bee venom ($p = 0.003$). There were no significant differences in late side effects during induction of VIT in patients allergic to bee and wasp venom. Acute local reaction, weakness, dyspnea, local itch were more frequent as early side effects in patients allergic to bee venom during the maintenance treatment ($p < 0.05$). Dyspnea, rhinitis and abdominal pain were more common as late side effects in patients allergic to bee venom during the maintenance phase ($p = 0.003$). Analyzing stings before VIT, blurred vision ($p = 0.02$) and hypotension ($p = 0.02$) were more frequent in patients allergic to wasp venom.

Side effects during venom immunotherapy depending on sex

In patients allergic to wasp venom, early and late side effects during the build-up and maintenance phase of VIT were more frequent in women. In contrast, in subjects allergic to bees both early and late side effects dur-

| Side Effects                        | Frequency |
|-------------------------------------|-----------|
| Burning face                         | 0.6%      |
| Wheeze                              | 0.6%      |
| Dysphonia                           | 0.6%      |
| Diarrhea                            | 0.6%      |
| Vomiting                            | 0.6%      |
| Sensation of obstruction in the throat | 0.6%    |
| Cough                               | 0.6%      |
| Abdominal pain                      | 0.6%      |
| Nosal blockage                      | 0.6%      |
| Syncope                             | 0.6%      |
| Hipotension                         | 0.6%      |
| Dizziness                           | 0.6%      |
| Scotoma                              | 0.6%      |
| Heat feeling                        | 0.6%      |
| Reddening of the skin               | 1.7%      |
| Itch of limbs and face              | 1.7%      |
| Dyspnoea                            | 1.7%      |
| Weakness                            | 1.7%      |
| Local swelling                      | 3.3%      |
| Itch                                | 0.6%      |
| Anxiety                             | 0.6%      |
| Somnolence                          | 0.6%      |
| Pins and needles/numbness           | 0.6%      |
| Rash                                | 0.6%      |
| Feeling cold                        | 0.6%      |
| Dyspnoea                            | 0.6%      |
| Feverishness                        | 0.6%      |
| Urticaria                           | 1.1%      |
| Local reddening                     | 5.6%      |
| Local swelling                      | 5.6%      |
| Attack of atrial fibrillation       | 0.56%     |
| Diarrhea                            | 0.56%     |
| Blurred vision                      | 0.56%     |
| Hyperhidrosis                       | 0.56%     |
| Chills                              | 0.56%     |
| Fever                               | 0.56%     |
| Rhinitis                            | 1.1%      |
| Tightness in the throat             | 1.1%      |
| Weakness                            | 1.1%      |
| Anxiety                             | 1.1%      |
| Numbness of limbs                   | 1.1%      |
| Dyspnoea                            | 1.1%      |
| Abdominal pain                      | 1.1%      |
| Headache                            | 1.7%      |
| Urticaria                           | 1.7%      |
| Drug eruption                       | 2.2%      |
| Itch                                | 1.8%      |
| Local reddening                     | 3.3%      |
| Local swelling                      | 15%       |

Figure 2. Side effects during VIT: early side effects during induction of VIT (A), late side effects during induction of VIT (B), early side effects during maintenance treatment of VIT (C), and late side effects during maintenance treatment of VIT (D)
Field stings during venom immunotherapy

Ten patients allergic to bee venom and 38 allergic to wasp venom were stung during VIT. The number of stings during VIT by wasps was 52 and by bees reached 22. Subjects allergic to wasp venom were stung usually in August, while those allergic to bee venom in July (Figure 3). Most of patients reported no reactions (50% of patients stung during VIT) or reported only mild immediate local reactions (43.7%). None of patients allergic to bee venom reported anaphylaxis, but three patients allergic to wasp venom (7.9%) had systemic reactions. All of those subjects with a systemic reaction on VIT received doubled dose (200 µg) of vaccine and did not report any further systemic reactions after field stings [1]. A weak reverse correlation between the side effect after stings by bees and wasps classified in the Muller's scale and patient’s age was found ($r = -0.36; p = 0.007$), which suggests that younger patients had stronger reactions after field stings during VIT than older subjects.

Sub-group of patients who completed venom immunotherapy

There were 18 patients allergic to wasp venom and 3 allergic to bee venom who completed at least a 5-year cycle of VIT (Table 2). The average duration of treatment, number of injections and the total dose of vaccine was higher in patients allergic to bee than to wasp venom ($p < 0.05$). A significant decrease in the sIgE levels was found in those patients who completed VIT. In detail, average sIgE in the wasp VIT group decreased from 6.19 IU/ml (before VIT) to 1.36 IU/ml (after 5-year VIT). Average sIgE in the bee VIT group decreased from 5.36 IU/ml to 2.53 IU/ml. The decrease in sIgE titer correlated with the total dose of vaccine received ($r = 0.53$, $p = 0.004$).

Factors associated with a higher risk of Hymenoptera venom allergy

Beekeeping is associated with a higher risk of bee venom systemic reactions (RR = 29.54, $p < 0.0001$). Female sex seems to be associated with a higher risk of any Hymenoptera venom systemic reactions (RR = 1.27, $p = 0.033$). Female sex seems to be associated with a higher

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**Table 3. Side effects (SE) during VIT for patients allergic to bee and wasp venom, side effects during VIT depended on sex (wasps), side effects during VIT depended on sex (bees)**

| Side effects (SE) during VIT for the whole study cohort | Wasps (%) of patients allergic to wasps | Bees (%) of patients allergic to bees |
|--------------------------------------------------------|----------------------------------------|--------------------------------------|
| SE during VIT                                           | 34                                     | 15                                   |
| Early SE during induction of VIT                         | 8.22                                   | 5.44                                 |
| Late SE during induction of VIT                          | 7.53                                   | 11.76                                |
| Early SE during maintenance treatment of VIT            | 3.42                                   | 20.59                                |
| Late SE during maintenance treatment of VIT             | 18.48                                  | 23.53                                |

| Side effects (SE) during VIT in patients allergic to wasp venom | Women | Men |
|----------------------------------------------------------|-------|-----|
| Early SE during induction of VIT                         | 11    | 5   |
| Late SE during induction of VIT                          | 9     | 5.36|
| Early SE during maintenance treatment of VIT            | 11    | 0   |
| Late SE during maintenance treatment of VIT             | 5.36  | 1   |

| Side effects (SE) during VIT in patients allergic to bee venom | Women | Men |
|---------------------------------------------------------------|-------|-----|
| Early SE during induction of VIT                             | 11    | 0   |
| Late SE during induction of VIT                              | 9     | 5.36|
| Early SE during maintenance treatment of VIT                | 3     | 2   |
| Late SE during maintenance treatment of VIT                 | 3     | 7.7 |

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Agnieszka Kołaczek, Dawid Skorupa, Monika Antczak-Marczak, Piotr Kuna, Maciej Kupczyk
risk of development of wasp venom allergy (RR = 1.27, \( p = 0.045 \)). A similar trend, however not reaching statistical significance, was found in women allergic to bee venom.

Discussion

Venom immunotherapy is the only causal treatment for patients allergic to Hymenoptera venom which is a potentially life-threatening condition. Patients allergic to bee or wasp venom, who have experienced a systemic reaction after sting and are not referred to receive VIT, are at a significant (40–75%) risk of the same or more severe reaction after another sting [6, 17]. The effectiveness of VIT ranges from 97.3% [16] to 93.8% [8] and the risk of systemic reactions after field stings in subjects treated with immunotherapy decreases to 6%. In a multicenter study conducted in 2013, the VIT failure rate (defined as a systemic reaction after a field sting by the culprit insect) reached 6.2%, as 22 out of 357 patients had systemic reactions [8]. In another study, the VIT failure rate was estimated at 6.5% of all treated cases [9]. This is confirmed by results of our study as in our cohort 3 (6.25%) patients experienced systemic reactions after field stings during VIT. All 3 reactions were graded 3 in Muller’s scale and were reported in patients allergic to wasp venom. Patients with a systemic reaction to field sting on VIT received doubled dose (200 µg) of vaccine, as suggested by the literature [1] and did not report any further systemic reactions after field stings. In the most comprehensive Cochrane Database systematic review of venom immunotherapy trials published till 2012, 7 studies enrolling 392 patients were analyzed [16]. It has been confirmed that VIT is effective in preventing allergic reactions as only 2.7% of participants treated had a subsequent systemic reaction to a sting, compared with 39.8% of untreated participants. Venom immunotherapy has been found to be effective in preventing large local reactions and improving quality of life of allergic patients.

Unfortunately, VIT may be associated with a risk of adverse events [7]. We have found that side effects during VIT are quite frequent (reported by 43.8% of patients in our cohort). The range of side effects reported in the literature varies from 17.9% to 45% of VIT applications [10–15]. Patients allergic to bee venom were found to be at a higher risk of a systemic allergic reaction during VIT (17.65% of events vs. 2.7% in wasp allergic patients). This is in line with several previous reports published to date showing an increased risk of side effects of VIT in bee allergic subjects [10, 13, 18–20]. In the Cochrane Database meta-analysis mentioned above, systemic adverse reactions occurred in 14.2% of participants treated with bee venom and only 2.8% in those treated with wasp venom [16]. The described phenomenon may be explained by higher allergenic potency of bee venom itself or the properties of the bee venom extract used for immunotherapy, as it has been proposed previously [18–20].

Almost all but one patient in our cohort were treated with the ultra-rush protocol. Several studies to date proved that cluster protocols, including the rush or ultra-rush scheme are significantly safer and much better tolerated by patients in contrast to a traditional protocol [12, 14, 15, 19, 20]. Early and late side effects are more common during the maintenance than during the induction of VIT with the ultra-rush protocol, whereas systemic side effects are significantly more frequent at the induction of VIT. Most of the side effects reported in our study were mild and localized, and included local swelling, reddening of the skin and itch at the site of the injection. Systemic
reactions as side effects during the induction and maintenance phase of VIT were observed in 4 individuals in our study group undergoing VIT against the wasp (2.7%) and in 6 patients allergic to bee venom (17.65%). Only in one case, the administration of adrenaline was needed. Thus, the incidence of systemic adverse reactions during VIT in our cohort may be estimated to reach 1 event per 706 injections (1.4% of injections), and potentially life-threatening reactions (defined as a need of adrenaline treatment) as 1 per 7060 injections (0.14% of injections). These findings are in line with a study by Ruëff et al. [18] where systemic adverse reactions were observed in 6–7% of patients undergoing VIT, but only in a few cases adrenaline was necessary [19, 20]. No fatalities due to VIT have been described in the literature to date [16]. In our study majority of systemic reactions were observed in the induction phase (1.9% of injections), not in the maintenance phase (0.19% of injections). This confirms findings of Mosbech et al. [13] who in a cohort of 840 patients undergoing VIT reported majority of systemic side effects (1.9%) during dose increase, and not in the maintenance phase (0.5% of injections). We have seen most of the systemic reactions (68.2%) within 30 min after the injection. All late systemic reactions (31.8% of events) were restricted to skin changes (urticaria, flush, reddening of the skin).

More than 20% of our study cohort patients were on ACEi or β-adrenergic antagonists. Interestingly, treatment with β-adrenergic antagonists or ACE inhibitors did not increase the risk of any kind of adverse events during VIT, systemic adverse events during VIT and was not associated with more severe symptoms during sting stings. This is in contrast to the findings of Ruëff et al. [9] who observed a cohort of 1532 patients and reported that ACE inhibitor medications were the strongest factors associated with more severe symptoms during sting challenge, followed by honeybee venom allergy and systemic allergic reaction during VIT, whereas double VIT and longer duration of therapy reduced the failure rate. It needs to be noted that the number of patients on anti-hypertensive medications in our study was probably too small to make any meaningful observations, as that was an observational, real-life analysis, not a hypothesis-driven and not powered based on a pre-defined hypothesis.

There are several limitations inherited in the design of our study. The study group consisted of only 180 patients however, most of the studies performed in one center analyze cohorts of around 100 patients [11, 12]. What is more important only 34 patients were allergic to bee venom, which may bias results of observations in this sub-group and may result in underestimation of statistical significance of our findings. On the other hand, all real life and epidemiological studies confirm that wasp venom allergy is more frequent, thus we believe that our results mimic the situation found in the every-day clinical practice. All patients undergoing venom immunotherapy at our center received oral antihistamines as a pretreatment before the ultra-rush induction phase and during the maintenance treatment. This is a well-accepted clinical practice, however it needs to be noted that this may result in underestimation of the number and/or severity of adverse events during VIT in our cohort. Reimers et al. [11] in a cohort of 57 patients on VIT due to honeybee venom allergy found that fexofenadine (in a dose of 180 mg) significantly reduced local allergic reactions and generalized symptoms of the urticaria and angioedema type, but did not mask the anaphylaxis. A higher frequency of sting allergy in females was found in our study. In most published studies there is a 60 : 40 male predominance. This raises questions of relative exposure to stings in males and females in this population, referral patterns, access to care, cultural biases or other, unknown reasons that might affect the gender ratio. Another limitation of our work is that we have not studied any other than sIgE possible biomarkers (e.g. serum tryptase), and thus we were not able to include those parameters in our analysis. However, current guidelines do not recommend any other biomarkers in a longitudinal follow-up of venom allergic subjects.

Conclusions

In the population studied, we have found that wasp is more frequent than bee venom allergy. Venom immunotherapy is highly efficacious and safe, as most of the adverse events during the induction and maintenance phase are mild and local. Side effects of VIT are more common in subjects on bee VIT. Majority of systemic side effects are reported during the induction phase, within 30 min after the injection, and the risk of adverse effects is not increased in patients receiving ACEi or β-adrenergic antagonists. Beekeeping and female sex are associated with a higher risk of allergy to Hymenoptera venom.

Conflict of interest

The authors declare no conflict of interest.

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