Clinical Study
Safety and Efficacy of Tree Pollen Specific Immunotherapy on the Ultrarush Administration Schedule Method Using Purethal Trees

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1. Background

Specific immunotherapy (SIT) is the common therapy method used in contemporary allergology. The efficacy and safety of SIT are discussed. Allergen-specific immunotherapy has been found to be effective in reducing symptoms of allergic rhinitis, especially in patients with house dust mite and pollen allergies. However, there is a difference in opinions as to whether SIT significantly improves asthma symptoms and about the long-term effects of SIT [1–5]. A new specimen for immunotherapy and an easier administration method (sublingual immunotherapy) aim to increase the efficacy and widespread use of this type of treatment [2, 5]. The accelerated administration schedule of SIT using cluster therapy or ultrarush is available mainly for venom allergies and sometimes for pollen allergies. However, there is insufficient information about the possibility of using these methods for other allergies [6, 7]. This study evaluates SIT with Purethal Tree (birch, hazel, and alder) chemically modified allergens at a concentration of 20,000 BAU (Bioequivalent Allergy Unit)/mL for tree pollen to assess the treatment’s efficacy and safety.

2. Materials and Methods

Twenty-two patients with tree pollen (birch, alder, and hazel) allergies, which manifest as seasonal rhinoconjunctivitis without bronchial asthma, were included in the study. Their allergy was confirmed using the skin prick test (SPT; HAL Allergy, Holland) and a concentration of pollen-specific IgE (sIgE) against birch, alder, and hazel. A positive SPT result was defined by the presence of a wheal with a diameter greater than 3 mm and with a histamine wheal greater than 5 mm. Tests of serum-specific IgE concentrations were performed using the Pharmacia CAP System FEIA (ThermoFisher, Sweden) immunoenzymatic method. The results of these assays
Table 1: Comparison of two different administration schedules of Purethal Tree SIT.

|                          | Group A                                                                 | Group B                                                                 |
|--------------------------|-------------------------------------------------------------------------|-------------------------------------------------------------------------|
| **Dose administration—period of time** | 0.1 mL, 0.2 mL, 0.3 mL, 0.4 mL—every week                               | 0.1 mL, 0.2 mL, 0.3 mL, 0.4 mL—every 15 min                              |
|                          | 0.5 mL—every two weeks, four times                                       | 0.5 mL—every 15 min, four times                                         |
| **Total BAU dose per year of treatment (10 000 BAU/mL)** | 100 000 BAU                                                             | 100 000 BAU                                                             |
| **Total BAU dose after entire therapy (3 years)**     | 300 000 BAU                                                             | 300 000 BAU                                                             |

BAU: Bioequivalent Allergy Unit.

were evaluated according to the manufacturer’s instructions [8]. A serum concentration of IgE greater than 0.75 IU/mL was confirmed as a positive result. Patients with concomitant bronchial asthma, allergy to grass pollens, house dust mites, Alternaria, or Cladosporium and those with contraindications to SIT were excluded from the study. Asthma was excluded based on a negative history, an examination, and a negative bronchial reversibility test according to Global Initiative for Asthma (GINA) [9].

All patients were randomized and divided into two groups:

(i) Group A received a conventional administration schedule of SIT with Purethal Trees as preseasonal therapy (October–January): the first dose of 0.1 mL, a second dose of 0.2 mL, and a third dose of 0.4 mL every week, and doses four, five, six, and seven of 0.5 mL every two weeks;

(ii) Group B received an ultrarush administration schedule of SIT with Purethal Trees as follows: the first dose of 0.1 mL, the second dose of 0.2 mL, the third dose of 0.4 mL, and doses four, five, six, and seven of 0.5 mL every 15 minutes (December or January). A comparison of the groups is shown in Table 1.

All patients were randomized and divided into two groups:

After three years of SIT, comparable effects were observed in both groups. A statistically significant reduction (P < 0.05) in nasal symptom scores during tree pollen season was found in Group A (3.991 ± 0.804 to 1.634 ± 0.540 points) and Group B (3.845 ± 0.265 to 1.501 ± 0.418). Nasal score values during all tree pollen seasons are shown in Table 3.

A statistically significant reduction was found in the scores of other allergic symptoms (P < 0.05). Scores in
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Table 2: Study groups characteristics.

|                          | Group A       | Group B       | P     |
|--------------------------|---------------|---------------|-------|
| Age                      | 20.9 ± 4.2    | 21.9 ± 3.2    | NS    |
| Men/women                | 6/4           | 6/6           | NS    |
| Disease duration          | 4.7 ± 1.5     | 4.9 ± 1.3     | NS    |
| Mean score of nasal symptoms before SIT | 3.991 ± 0.804 | 3.845 ± 0.265 | NS    |
| Mean score of other symptoms before SIT | 4.127 ± 0.652 | 4.011 ± 0.407 | NS    |
| Serum concentration of total IgE | 402.78 ± 54.8 | 387.66 ± 74.61 | NS     |
| Serum concentration of sIgE to birch | 32.72 ± 8.77  | 35.33 ± 6.87  | NS    |
| Serum concentration of sIgE to alder | 15.92 ± 3.91  | 18.73 ± 4.61  | NS    |
| Serum concentration of sIgE to hazel | 28.55 ± 6.12  | 30.34 ± 5.88  | NS    |

sIgE: specific IgE; SIT: specific immunotherapy; NS: not statistically significant.

Table 3: Nasal symptom scores during tree pollen seasons before and during SIT therapy.

|                          | II–V 2009 | II–V 2010 | II–V 2011 | II–V 2012 |
|--------------------------|-----------|-----------|-----------|-----------|
| Group A                  |           |           |           |           |
| N = 1209                 | 3.991 ± 0.804 | 2.441 ± 0.322 | 1.760 ± 0.454 | 1.452 ± 0.207 |
| Group B                  | 3.845 ± 0.265 | 2.219 ± 0.422 | 1.834 ± 0.231 | 1.588 ± 0.367 |
| P                        | NS        | NS        | NS        | NS        |

NS: not statistically significant.

Table 4: Use of relief drugs during tree pollen season.

|                          | II–V 2009 | II–V 2010 | II–V 2011 | II–V 2012 |
|--------------------------|-----------|-----------|-----------|-----------|
| Group A                  |           |           |           |           |
| N = 776                  | 1.661 ± 0.445 | 1.018 ± 0.578 | 0.880 ± 0.250 | 0.498 ± 0.213 |
| Group B                  | 1.788 ± 0.652 | 1.109 ± 0.299 | 0.745 ± 0.321 | 0.532 ± 0.244 |
| P                        | NS        | NS        | NS        | NS        |

NS: not statistically significant; N: number of days with score assessment.

Groups A and B fell from 4.127 ± 0.652 to 2.752 ± 0.136 and from 3.845 ± 0.265 to 2.331 ± 0.109, respectively.

After three years of SIT, use of relief drugs was significantly lower in both groups, as shown in Table 4.

3.1. Concentration of Birch, Hazel, and Alder IgG4 during SIT. Before SIT, the mean value of birch IgG4 was 1577 AU/mL (range: 24–1786 AU/mL). The value was significantly higher after three years of SIT, with values of 3566 AU/mL (range: 45–5609 AU/mL) in Group A and 3678 AU/mL (range: 78–6781 AU/mL) in Group B (P < 0.05).

Similar trends were observed in alder IgG4. At the start of the study, the mean alder IgG4 value was 790 AU/mL (range: 45–3821 AU/mL). After three years, mean IgG4 levels were 21786 AU/mL (range: 89–16780 AU/mL) in Group A and 10402 AU/mL (range: 88–15090 AU/mL) in Group B. No statistically significant differences in serum IgG4 levels for any analyzed pollens were found between Groups A and B after three years of SIT (P > 0.05).

3.2. Safety Results. No systemic anaphylactic reactions (I, II, III, or IV degree reactions) occurred in either group during the course of SIT therapy [12]. Of all 211 Purethal Trees injections performed on patients in Group A, erythema or wheals smaller than 5 cm were observed after 23 injections (10.9%) and erythema or wheals greater than 5 cm after seven injections (3.3%). Similar results were observed in Group B: erythema or wheals smaller than 5 cm were observed after 18 injections (7.8%) and erythema or wheals greater than 5 cm after eight injections (3.5%) out of 231 total injections.
There were no statistically significant differences between the groups.

4. Discussion

SIT has become a widespread and continuously improving treatment. The ultrarush administration schedule method has been safe and effective in treating allergy to hymenoptera venom [2, 6]. Although many studies emphasize the safety and efficacy of SIT on wasp or bee allergies, SIT administration of vaccines has not been widely used in other studies of allergy desensitization [13–19].

Purethal Tree, which was used in this study, has been confirmed as an effective and safe means to treat a cluster [14]. This type of treatment is an indirect form of an administration schedule between conventional SIT and the ultrarush method. Our work confirmed that Purethal therapy using an ultrarush administration schedule before pollen season is similarly safe.

The results of this study quell doubts about the safety of this treatment. A statistically significant reduction in allergy symptoms during pollen season was correlated with the use of the conventional method. However, it was a preliminary study based on a small group of patients.

The study also showed a statistically significant reduction of symptomatic drug use. In the in vitro control treatment, we used IgG4 measurements. Unfortunately, there is no superior method of assessing the effectiveness of SIT. The observed increase in IgG4 levels after three years of SIT may be further evidence that an ultrarush administration schedule of pollen SIT is effective. The available literature includes studies that primarily attempt to accelerate desensitization to allergens using sublingual immunotherapy, although there are a few case studies of SIT injection. Notably, all of these case studies showed positive efficacy and safety profiles [15–18].

The present study was not double-blinded or placebo-controlled. However, the main aim of this study was to compare two methods of administering a vaccine. An important result from this study is the demonstrated safety associated with the ultrarush treatment compared to the conventional method. No patients had systemic reactions during ultrarush treatment, unlike the group with conventional SIT, which supports currently available data [13, 18–21]. However, as this is a new method of treatment and research is still insufficient, ultrarush SIT should be conducted within 24 hours of hospitalization.

In this study, most patients receiving vaccinations using the ultrarush administration schedule method experienced more minor subsequent allergic reactions. This outcome should be confirmed in a larger group of patients. An important observation regarding this study is that none of the patients treated with an ultrarush administration experienced late reactions. Late reactions cannot be ruled out, and further studies on their likelihood are needed.

5. Conclusion

An ultrarush administration schedule of SIT with Purethal Trees is a safe treatment in preliminary observations. This therapy is comparable with conventional administration of SIT in terms of efficacy and safety. However, further investigations on larger groups of patients are required.

Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

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