Subcutaneous Immunotherapy With High-Dose Cat and Dog Extracts: A Real-life Study

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Abstract

Background: Data on the efficacy of immunotherapy administered to patients with cat or dog allergy are scarce.

Objective: We aimed to evaluate the safety and efficacy of subcutaneous immunotherapy (SCIT) in patients with allergy to cat and dog dander.

Methods: Consecutive patients with rhinitis and/or asthma related to sensitization to cat or dog dander were included in a pragmatic, real-life, prospective, observational study. All patients had specific IgE to cat, dog, or both. SCIT was administered using an infusion pump over 3 sessions as part of a rush protocol, followed by monthly administration over 12 months. We recorded adverse events, clinical outcomes, pulmonary function, FeNO, symptoms of rhinitis and asthma, quality of life (QoL), Asthma Control Test (ACT) score, and visual analog scale (VAS) score at baseline, 6 months, and 12 months.

Results: The study population comprised 66 patients (38 females, 46 allergic to cat and 20 to dog), with ages ranging from 9 to 59 years. During the up-dosing phase, in which the infusion pump was used, 8.1% of doses elicited a systemic reaction and 5.4% caused a local reaction, while 9.3% of doses administered during the maintenance phase (ie, without an infusion pump) induced a systemic reaction. No local reactions were recorded. A significant improvement in FEV1, symptoms of rhinitis and asthma, QoL, use of medication, VAS score, and ACT score was observed at 6 months and continued at 12 months. Clinical improvement with cat extract was significantly better than with dog extract.

Conclusions: High-dose SCIT has substantial clinical value in many cat- and dog-allergic patients.

Key words: Cat. Dog. Allergy. Rhinitis. Asthma. Allergen immunotherapy.

Resumen

Antecedentes: Hay pocos estudios sobre la eficacia de la inmunoterapia administrada a pacientes con alergia a perro o gato.

Objetivo: Evaluar la seguridad y la eficacia de la inmunoterapia subcutánea (SCIT) en pacientes con alergia a estos dos animales.

Métodos: Se incluyeron pacientes consecutivos con rinitis y/o asma relacionados con la sensibilización al gato o al perro en un estudio observacional prospectivo, pragmático, en vida real. Todos los pacientes tenían IgE específica para gato y/o perro. La SCIT se administró utilizando una bomba de infusión (IP), en 3 sesiones como parte de un protocolo rápido, seguido de la administración mensual durante 12 meses. Se recopilaron datos sobre efectos adversos y resultados clínicos, función pulmonar, FeNO, síntomas de rinitis y asma, calidad de vida (QoL), prueba de control del asma (ACT) y escala analógica visual (VAS) al inicio, a los 6 y 12 meses.

Resultados: Se incluyeron 76 pacientes: 38 mujeres, 46 alérgicos a gato y 20 a perro, con edades comprendidas entre los 9 y los 59 años. Durante la fase de administración ascendente, utilizando una IP, el 8,1% de las dosis provocó una reacción sistémica (SR) y el 5,4% causó una reacción local (LR), mientras que el 9,3% de las dosis administradas durante la fase de mantenimiento (es decir, sin IP) desarrolló una SR, y no se registraron LR. Se observó una mejoría significativa en el FEV1, en los síntomas de rinitis, de asma y en los cuestionarios de la calidad de vida, uso de medicación, VAS y ACT a los 6 meses y continuó a los 12 meses. La mejoría clínica con el extracto de gato fue significativamente mayor que con el perro.

Conclusiones: Las dosis altas de SCIT tienen un valor clínico sustancial en muchos pacientes alérgicos a perros y gatos.

Introduction

Data on the efficacy of allergen immunotherapy in patients with cat allergy [1-9] or dog allergy [3,5,10] remain scarce, especially in the case of dog allergy. High doses of standardized extracts have proven effective in treating patients who are allergic to cat. Previous double-blind studies with subcutaneous immunotherapy (SCIT) based on standardized cat allergen extracts have demonstrated that maintenance doses containing 13.2 µg of Fel d 1 [6], 13.8 µg of Fel d 1 [4], and 15 µg of Fel d 1 [7,8] improved symptoms associated with exposure to cat and brought about immunologic changes [2,4,8,11-13]. SCIT with standardized dog allergen extract has proven less efficacious than SCIT with standardized cat allergen extract [3,5,10], despite having induced immunologic changes [12,14].

The purpose of the present study was to explore the use of high-dose SCIT in patients with allergic rhinitis and asthma caused by exposure to cat and dog dander in real-life clinical practice.

Material and Methods

Patients

We conducted a prospective observational study of consecutive patients with rhinitis and/or asthma due to sensitization to cat or dog dander and for whom treatment with immunotherapy was indicated [15]. All candidates for inclusion expressed a willingness to initiate SCIT after receiving information on the possible benefits of this approach. For a patient to be included, a strong association between clinical symptoms and exposure to cat or dog was required. An additional requisite was positive specific IgE to cat or dog extract as evidenced by the skin prick test (ALK, Denmark) and/or in serum and specific IgE to whole cat or dog extracts or extracts of major allergens: Fel d 1, 15 µg/mL; Can f 1, 3.21 µg/mL; Can f 5, 0.72 µg/mL [16].

Up-dosing consisted of 3 progressively increasing doses from the maintenance vials (100 000 SQ/mL; ie, 10 000, 50 000, and 100 000 SQ) administered at weekly intervals. Doses were administered using an infusion pump (Infusa T1, Medis), with infusions lasting 30 minutes, as previously described [17,18]. Patients were observed for 30 minutes after the infusion was complete. Subcutaneous injections were applied for monthly maintenance doses.

Adverse reactions (ARs) (local, systemic, immediate, or delayed) were recorded according to EAACI guidelines [15]. Delayed ARs were monitored by telephone interview 48 hours after SCIT; for this purpose, patients received instructions on how to monitor ARs if necessary. Patients were not routinely premedicated.

Clinical Outcomes

We performed spirometry and bronchodilation testing and measured the fractional exhaled nitric oxide (FeNO) concentration. We also administered a series of questionnaires validated for the Spanish population, as follows: ESPRINT-15 (health-related quality of life in allergic rhinitis), Asthma Quality of Life Questionnaire (AQLQ), Asthma Control Test (ACT), and a 10-cm visual analog scale (VAS) for assessment of nasal, ocular, and bronchial symptoms. Scores for nasal symptoms (itching, congestion, rhinorrhea, sneezing), ocular symptoms (tearing, itching, gritty sensation), and pulmonary symptoms (cough, wheezing, dyspnea, exercise-induced asthma) (0, no symptoms; 1, mild; 2, moderate; 3, severe) and use of medication were applied. Medication use was scored as follows: antihistamines, 6 points; short-acting β2-agonists, 2 points; inhaled corticosteroids, 2 points; nasal topical corticosteroids, 2 points; and oral corticosteroids, 4 points. All assessments were performed at baseline and at 6 and 12 months.

To measure the response to SCIT, the minimum clinically important differences considered were as follows: ESPRINT-15, >0.9 [19]; AQLQ, >0.5 per dimensional item, with an average change of 1.0 considered moderate and a change of at least 1.5 considered large [20]; ACT >3 [21]; and VAS >2 between visits.

Associations between the IgE molecular profile and safety outcomes and the clinical efficacy of cat and dog extract were analyzed, as was the difference in the clinical efficacy of SCIT between cat and dog extract.

Statistical Analysis

The statistical analysis was based on the Fisher exact test, Wilcoxon test, Friedman test, Kruskal-Wallis test, and a mixed-effects model. P values <.05 were considered significant.

Results

Patients

Sixty-six patients were included (38 females and 28 males, of whom 46 were allergic to cat and 20 to dog), with a mean age ranging from 9 to 59 years (34.23 [12.1]). Of these, 36.3% were sensitized to pollen, 4.1% to profilin, 6% to mites, and 6% to other allergens. Allergic rhinitis was present in 65 patients (98.5%) and asthma in 64 (97%). Most patients had persistent and moderate symptoms. Rhinitis was intermittent in 13.8%, mild persistent in 21.6%, and moderate/severe persistent in 64.6%. Asthma was intermittent in 23%, mild persistent in 36%, and moderate/severe persistent in 41%. Sixty-one patients (92%) had either a dog or cat at home (daily direct contact), while 3 (4.6%) had indirect contact with these animals, and 2 (3%) were veterinarians.

Four patients dropped out of the study at the end of the up-dosing phase to continue their treatment in another center; a further 4 patients withdrew from the study between the third
and sixth months (3 due to poor SCIT tolerance, 1 for personal reasons), and 7 patients between the sixth and twelfth month (all for personal reasons such as lack of time, distance from the hospital, travel abroad, and discontinuation of contact with pets). Fifty-one patients concluded the study (34 cat-allergic and 17 dog-allergic).

**Adverse Events**

During the up-dosing phase, 18 doses (8.1%) of SCIT (all with cat extract) triggered a systemic reaction (SR), and 12 doses (5.4%) produced a local reaction (LR); by comparison, 3 doses (9.3%) administered during the maintenance phase caused an SR; no LRs were recorded. No SRs were triggered by doses of SCIT with dog extract in the up-dosing phase or in the maintenance phase (ie, with or without the infusion pump), and 2 doses (2.1%) delivered using an infusion pump caused an LR, none of which occurred in the maintenance phase. ARs to SCIT tended to occur in patients with more severe asthma and poorer control, although this difference did not reach statistical significance.

The most frequent symptoms of SRs with SCIT based on cat extract were rhinitis (71.4%) and asthma (71.4%), followed by conjunctivitis (43%) and urticaria (24%). Most SRs were immediate (90%) and grade 1 (62%), and the rest were grade 2 (8 SRs). All SRs were controlled and resolved after treatment with antihistamines (100%), inhaled β₂-agonists (71%), systemic corticosteroids (47%), and epinephrine (42.8%). No cases of anaphylactic shock or hypotension were reported. The onset, grade, and treatment of SRs were similar in both the up-dosing phase and the maintenance phase. Onset was late and intensity mild in all LRs, and none required treatment. In total, 3 patients were withdrawn from the study owing to adverse events (all with cat extract).

**Specific IgE**

All patients had a positive IgE level (>0.35 kU/L) and prick test result with cat and dog extracts. Among the cat-allergic patients, the results were positive for Fel d 1 in 83% of cases, Fel d 2 in 26%, and Fel d 4 in 50%. Among the dog-allergic patients, the results were positive for Can f 1 in 79%, Can f 2 in 47%, Can f 3 in 26%, and Can f 5 in 63%. In the case of dog-allergic patients, 21.1% were monosensitized to Can f 5; none were monosensitized to lipocalins or serum albumins. All dog-allergic patients recognized at least 1 commercially available allergen, while 12% of cat-allergic patients did not recognize any.

**Other Outcomes**

The results of the pulmonary function test, FeNO values, symptom scores, and the ACT, VAS, and medication scores are shown in Figure 1.

Baseline spirometry was normal in most patients with asthma, and the bronchodilation test did not reveal significant differences and sixth months (3 due to poor SCIT tolerance, 1 for personal reasons), and 7 patients between the sixth and twelfth month (all for personal reasons such as lack of time, distance from the hospital, travel abroad, and discontinuation of contact with pets). Fifty-one patients concluded the study (34 cat-allergic and 17 dog-allergic).

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At the beginning of the study, FeNO was elevated (>50 ppb) in 77% of patients (47) and remained elevated at 6 months in 49% of patients (27) and at 12 months in 54.6% [30]. Mean FeNO values decreased by around 10% at the end of the study (Figure 1).

The greatest decrease in medication use was seen with antihistamines and short-acting β-agonists. Doses of inhaled corticosteroids were approximately halved. No patient required systemic corticosteroids during the study (Figure 1).

The results of quality of life questionnaires for rhinitis and asthma are presented in Figure 2. The ESPRINT-15 questionnaire showed that 87.3% of patients improved at month 6 and that 80% maintained this improvement at month 12. The decrease in total score and all dimensions was greater than the minimal important difference (0.9).

At months 6 and 12, all dimensions of the AQLQ (activity limitation, symptoms, emotional function, environmental stimuli, and total) increased by more than 0.5 points, the minimal important difference; in many patients, this increase was more than 1, which is considered a moderately important change. The AQLQ showed an overall improvement in 87% of patients in the first 6 months, with 83% maintaining the improvement at 12 months. The response to SCIT with cat and dog extract was good or very good in 66.7% of patients. Patients reported a subjective improvement with the third maintenance dose (month 2 of treatment).

The molecular profile (number of allergens to which the patients were sensitized or the quantitative value of specific IgE to allergens) was not related to the response to SCIT with cat or dog extracts \((P=.646)\) or with safety profile \((P=.109)\), FeNO values \((P=.592)\), direct contact with cats or dogs \((P=.39)\), age \((P=.218)\), or gender \((P=.697)\).

At the baseline visit, patients with dog allergy experienced a more deleterious impact on their quality of life in terms of asthma symptoms and emotional parameters \((P=.041\) and \(P=.009\), respectively), required greater use of medication in general (especially inhaled corticosteroids \([P=.005]\) and antihistamines \([P=.006]\)), and had higher FeNO values \((P=.001)\) and higher perceived symptom intensity (VAS). At the 12-month visit, the perceived intensity of symptoms remained unchanged \((P=.008)\). In addition, dog-allergic patients had an increased need for medication \((P=.02)\), poorer rhinitis-related quality of life in relation to their daily activities \((P=.042)\), and a lower FEV\(_1\)/FVC ratio \((P=.049)\) than patients with cat allergy. The ACT scores at 6 and 12 months of treatment revealed a tendency toward improved disease course among patients receiving SCIT with dog extract than in those receiving SCIT with dog extract, although the differences were not statistically significant.

### Discussion

Pragmatic trials provide information that may be considered complementary to data from randomized clinical trials [22]. This study, which was carried out in a real-life setting, confirms the efficacy of SCIT using extracts with high doses of dog and cat allergens. We administered immunotherapy using a rush up-dosing phase with an infusion pump followed by a monthly maintenance dose over a 12-month period. The safety profile of the rush up-dosing and maintenance phases was generally good, as reported in previous studies using an infusion pump [17,18], conventional schedules [23], and other rush schedules [24], and the safety profile of the up-dosing and maintenance phases was even better than some cluster schedules described in the literature [5].

Systemic reactions to the SCIT were only observed with cat extract but not with dog extract, as reported elsewhere [5,10]. This extract-specific difference could be due to the high concentration of Fel d 1 (15 µg), the major cat allergen, in comparison with the major dog allergens, Can f 1 (3.21 µg) and Can f 2 (3.52 µg).
Can f 5 (0.72 µg) [16]. Most systemic reactions produced were grade 1 or 2 and resolved with appropriate treatment, although 3 patients withdrew from the study during the maintenance phase owing to adverse effects.

In this study and others published by our group [16,25], a high percentage of patients were monosensitized to Can f 5. Can f 1 and Can f 5 were the most frequently recognized allergens in dog-allergic patients, followed by Can f 2. In the case of cat-allergic patients, Fel d 1 was the most frequent, followed by Fel d 4 and Fel d 2. Of note, 12% of cat-allergic patients had positive IgE to whole cat extract, although they did not recognize the 3 cat allergens tested, thus emphasizing the need for more allergens in clinical practice and the need to control the presence of whole allergens in diagnostic and treatment extracts. The pattern of allergen recognition was not associated with the onset of adverse events or efficacy of treatment.

SCIT with cat or dog extract showed clear clinical efficacy at 6 months, and this efficacy was maintained at 12 months of treatment. Throughout the study, significant improvements were observed in FEV₁, rhinitis, AQLQ score, ACT score, VAS score, symptom score, and use of medication, even in cases where the patient maintained direct contact with the pet(s) at home. In addition, although the increase in FEV₁ was not clinically relevant, the improvement in rhinitis and the test scores (AQLQ and ACT) exceeded the minimal important difference. Taylor et al [1], performed a double-blind placebo-controlled study with 10 cat-sensitized patients in which they recorded decreased bronchial and cutaneous sensitivity to cat extract in the group that received SCIT with cat allergen. Similar results have been reported elsewhere [2-4]. We recorded a less substantial improvement in ocular symptoms than in airway symptoms, thus contrasting with the studies by Alvarez-Cuesta et al [6] and Varney et al [7] (both with cat extract) and Valovirta et al [10] (with dog extract), who observed decreased conjunctival sensitivity throughout treatment.

Clinical efficacy differed based on the type of extract (cat or dog). The clinical improvement with cat extract was significantly more marked in terms of symptoms, use of medication, and lower FEV₁/FVC ratio than with dog extract. Previous studies reported that SCIT with dog extract was less effective than SCIT with cat extract [3,5,12,19,26], likely owing to the higher concentration of Fel d 1 than of Can f 1 or Can f 5 [16], even using concentrations recommended in the USA (ie, 15 µg/mL of Can f 1 per dose) [26]. Specific IgG and IgG4 to cat and dog allergens were not measured in this study. However, a significant immune response in the form of increased IgG and/or IgG4 titers to cat [4,8,12,13,19] and dog [10,12,14] dander has been reported with extracts that were similar to those used in our study (Alutard SQ). The good clinical and immune response to SCIT is clearly associated with the high doses of major allergens contained in the extracts, mainly in the case of cat [3-8,13,18,19].

Our study is limited by the modest number of patients and the lack of a placebo-control group. However, it was designed as a pragmatic trial. In addition, it is difficult to confirm our results, since we did not perform nasal or bronchial challenge or assess changes in skin reactivity.

Conclusions

This real-life study reinforces the clinical efficacy of SCIT with high-dose extracts in cat- and dog-allergic patients. The safety and efficacy profile of this SCIT regimen with cat or dog extract was not related to the molecular profile of IgE.

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Conflicts of Interest

JS reports having served as a consultant to Thermo Fisher Scientific, MSD, Novartis, Genentech, Sanofi, Leti, Roche, ALK, FAES FARMA, Mundipharma, and GSK. He has also been paid lecture fees by Novartis, GSK, Stallergenes, LETI, and FAES FARMA and has received grant support for research from Thermo Fisher Scientific, ALK, and Sanofi.

SU and MJR declare that they have no conflicts of interest.

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