Observational long-term follow-up study of rapid food oral immunotherapy with omalizumab

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Abstract

Background: A number of clinical studies focused on treating a single food allergy through oral immunotherapy (OIT) with adjunctive omalizumab treatment have been published. We previously demonstrated safety and tolerability of a rapid OIT protocol using omalizumab in a phase 1 study to achieve desensitization to multiple (up to 5) food allergens in parallel, rapidly (7–36 weeks; median = 18 weeks). In the current long-term, observational study, we followed 34 food allergic participants for over 5 years, who had originally undergone the phase 1 rapid OIT protocol.

Methods: After reaching the maintenance dose of 2 g protein for each of their respective food allergens as a part of the phase 1 study, the long-term maintenance dose was reduced for some participants based on a pragmatic team-based decision. Participants were followed up to 62 months through standard oral food challenges (OFCs), skin prick tests, and blood tests.

Results: Each participant passed the 2 g OFC to each of their offending food allergens (up to 5 food allergens in total) at the end of the long-term follow-up (LTFU) study.

Conclusion: Our data demonstrate the feasibility of long-term maintenance dosing of a food allergen without compromising the desensitized status conferred through rapid-OIT.

Trial registration Registry: Clinicaltrials.gov. Registration numbers: NCT01510626 (original study), NCT03234764 (LTFU study). Date of registration: November 29, 2011 (original study); July 26, 2017 (LTFU study, retrospectively registered)

Keywords: Follow-up, Food allergy, Maintenance dosing, Omalizumab, Oral immunotherapy

Background

Food Allergy is a serious immune response that disproportionately affects children [1–3] and its prevalence among children has increased over the past decade [4–7]. Food allergy is a burden for both the individual and the family, decreasing quality of life due to anxiety of potential reactions [8]. It can be associated with increased anxiety in the child [9].

There is no FDA approved therapy for food allergies, but oral immunotherapy (OIT), a treatment in which the patient eats small but slowly increasing doses of their allergen until they can tolerate a specified dose, was shown in research settings to be successful in children and adults for single foods as well as up to five foods in parallel [10–14].

In current care, individuals who are successfully desensitized to their offending food(s) at the end of OIT continue to ingest small amounts of these foods to maintain desensitization. However, studies following the participants after OIT long-term were focusing on single food OIT trials and without omalizumab as adjunct therapy [15–17].
In this observational, retrospective study, we followed the participants from a single site, phase 1 study in which omalizumab was used as adjunctive therapy with OIT (rapid-OIT) [14]. Our results for up to 62 months maximum during long-term follow-up (LTFU) dosing (after 2 g of food protein maintenance dose was reached for each food allergen by the participant) suggest that simultaneous desensitization to food allergens is possible over the long term.

**Methods**

**Study design and participants**

Participants who were successfully desensitized in a phase 1 omalizumab OIT (rapid-OIT) trial [14] [Investigational New Drug (IND) 14831, NCT01510626)] were invited to continue in an IRB-approved LTFU study at Stanford University. In the original publication [14], only a subset of participants with peanut in their OIT were included. However, additional participants with other food allergies (as determined by a validated food challenge) underwent the same protocol and were also invited to enroll in this LTFU study. The main additional allergies included allergies to almond, cashew, egg, hazelnut, milk, peanut, pecan, and walnut (Table 1). Quality of life studies for this cohort have been published [18, 19]. At the start of the LTFU study, the participants were switched from food flours/powders to food equivalents based on exact protein amounts for each specific food allergen. Participants of the LTFU study returned to the clinic on average every 6–12 months for questionnaires, skin tests, blood tests, and oral food challenges (OFCs). Initial maintenance dose in the phase 1 study was 2 g protein per food allergen. Then, for the LTFU, high (approximately 2 g protein) or low (approximately 300 mg protein) maintenance doses for each of the participant’s food allergens were selected in conjunction with the patient and the clinician. All participants were required to carry reaction medications throughout the LTFU study. Participants were closely monitored via routine clinic visits and phone calls during the LTFU study. Self-reported adverse Events were documented as per CTCAE v4.03 [20] criteria and according to regulatory guidelines. An independent Data Safety and Monitoring Board (DSMB) met every 6–12 months to review participant safety data.

**Oral food challenges, skin prick tests, and serological analysis**

We conducted separate, standardized OFCs using validated and published techniques [21] with each food allergen that had been included in the OIT. For each allergen, OFCs were done in a staged approach with approximately 150, 300, 600, 1200, and 2000 mg of each food allergen on separate days. Skin prick tests (SPTs) were performed as per published methods [14]. Allergen-specific IgE and IgG4 levels were measured at Johns Hopkins Allergy and Clinical Immunology Reference Laboratory by ImmunoCAP fluorescence enzyme immunoassay (Thermo Fisher scientific/Phadia, Waltham, MA).

**Statistical analysis**

Statistical analyses and preparation of graphics were performed in R. Kaplan–Meier methods were utilized to graphically depict time to reduction of long-term maintenance dose to for the separate foods. Differences between the Kaplan–Meier curves were assessed using a log-rank test (R package survival [22, 23]). The association of allergen specific IgG4 to IgE ratios (log 10) as well as SPT data with the two different dose levels (low vs. high maintenance dose) were evaluated for participants with available data using a linear mixed effects regression model controlling for the months of time point after maintenance was reached with random effects for participant and allergen type. Similarly, we fitted a linear mixed

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**Table 1 Demographics at the beginning of the phase I studies of participants who were enrolled in the LTFU study**

| Characteristics                        | LTFU participants |
|------------------------------------------|-------------------|
| Number of ITT participants               | 34                |
| Sex, male (n), female (n)                | M (23), F (11)    |
| Age at beginning of original phase I trial in years, median (range) | 8.7 (4.6–16.9)  |
| With other atopic conditions             |                   |
| Atopic dermatitis (%)                    | 57%               |
| Atopic rhinitis (%)                      | 73%               |
| Asthma (%)                               | 63%               |
| With number food(s) in OIT               |                   |
| 1 food in OIT, n (%)                     | 6 (17.6%)         |
| 2 foods in OIT, n (%)                    | 7 (20.6%)         |
| 3 foods in OIT, n (%)                    | 5 (14.7%)         |
| 4 foods in OIT, n (%)                    | 9 (26.5%)         |
| 5 foods in OIT, n (%)                    | 7 (20.6%)         |
| With food in OIT*                        |                   |
| Almond, n (%)                            | 6 (17.6%)         |
| Cashew, n (%)                            | 18 (52.9%)        |
| Egg, n (%)                               | 11 (32.4%)        |
| Hazelnut, n (%)                          | 7 (20.6%)         |
| Milk, n (%)                              | 10 (29.4%)        |
| Peanut, n (%)                            | 26 (76.5%)        |
| Pecan, n (%)                             | 8 (23.5%)         |
| Walnut, n (%)                            | 10 (29.4%)        |

*OIT oral immunotherapy
a Other foods with n = 1 not listed: e.g. Barely, Soy, Rye
effects regression model to assess the association of the outcomes of the OFCs to 2 g throughout the LTFU study with the two different dose levels (low vs. high maintenance dose), controlling for the months of time point after maintenance was reached, the allergen type and the participant.

Results
Study participants and characterization of their food allergies
In this LTFU study, 34 participants previously desensitized to 1 or up to 5 foods in parallel during a rapid-OIT trial (with 16 weeks of adjunct omalizumab), were followed. Methods for the rapid desensitization are published elsewhere [14]. The participants were followed for up to 62 months after the respective participant reached the 2 g maintenance dose for the first food (median 53 months, range 32–62 months, Fig. 1). The participants entered the LTFU study at different dates and they were followed until May 2017, which resulted in the variation of their time in the LTFU study. A summary of the clinical characteristics and demographics of the participants who enrolled in the LTFU study is displayed in Table 1. The largest proportion of participants (n = 9, 26.5%) was dispensed four food allergens in their OIT. The most common food allergen was peanut (n = 26, 76.5%) and the second most common food allergen in the OIT was cashew with 18 (52.9%) participants. Overall, we followed the long-term maintenance dosing of eight different food allergens (Table 1). The participants were desensitized to various combinations of these food allergens. It was previously shown in a study of DBPCFCs in 60 multi-allergic children that several combinations of food allergies often occur together [24]. The number of participants that were sensitized, underwent OIT and then followed for each pairwise combination of two foods is shown in Fig. 2. Of the 11 participants that were desensitized to egg and the 10 that were desensitized to milk, 8 were desensitized to both of these foods. Of the 8 participants with pecan in their OIT and 10 with walnut, 7 were desensitized to both foods, corresponding with previous reports of cosensitization between pecan and walnut [25, 26].

Long-term maintenance dosing and OFC
We followed the 34 participants for a maximum of 62 months after they reached their 2 g protein maintenance dose for the first food allergen in their OIT, total of up to 65 months of dosing. Throughout the follow-up phase for each participant for each food allergen the decision to stay on a high LTFU maintenance dose per food allergen or reduce to a low LTFU maintenance dose per food allergen was made between the clinician and the parent in a collaborative approach based on adherence and taste. No change of dose occurred because of safety reasons and each dose change was monitored closely by the trained clinical team.

The long-term maintenance doses at the end of our follow-up phase in May 2017 for each participant for
the small sample size and the small percentage of failed results have to be interpreted with care, considering..

During the dose escalation phase (Additional file 2: Figure S2). During the LTFU phase, the wheal diameter varied across a large range for the participants per allergen, however, it was not significantly dependent on a low or high maintenance dose (P = 0.1).

Safety The number of allergic reactions during the LTFU study decreased over time (Table 2). In total 1126 reactions were recorded (3.5% of maintenance doses). Of those, 1076 reactions (95.6%) were mild, 40 (3.6%) were moderate and 5 (0.4%) were classified as severe. There were no fatal events. There were no serious adverse events. There were no anaphylactic reactions. Four of the severe adverse events were skin reactions and one was nasal congestion. All five severe reactions occurred within the first 19 months after participants reached the maintenance dose and while each participant was on a 2 g dose..
maintenance dose for their food allergens. No severe respiratory, gastrointestinal or hypotension adverse events occurred. Epinephrine was used by participants for mild or moderate reactions associated with minor wheezing. There were no events involving shortness of breath, dyspnea, hypotension, dizziness, or difficulty breathing. Safety results did not differ based on low vs high LTFU maintenance dose.

**Discussion**

The efficacy of OIT for food allergy to desensitize to one or several foods has been demonstrated in several trials [10–14, 27, 28]. It is important to study the follow-up of participants after they finished the desensitization. Published follow-up studies [15–17, 29, 30] mostly focus on participants who continue on or increase their achieved maintenance dose and who were desensitized to only a
single food mostly without omalizumab therapy. In Meglio et al. [17], families who participated in a OIT trial for cow’s milk were evaluated by a structured interview after an average 4 years and 8 months after start of their OIT. The parents were advised at the end of the original trial not to discontinue the daily intake of cow’s milk (with the goal of maintaining the effects of the oral desensitization) and the follow-up results indicated a persistence of the desensitization effect [17]. Jones et al. [15] followed participants of an egg OIT trial for up to 4 years in which the participants continued egg intake or (for a shorter period) placebo. At the end of the study, the participants were tested for sustained unresponsiveness, which 50% of the egg-consuming group achieved. In a recent blinded, phase 2 clinical trial, it was shown that OIT in combination with omalizumab enables rapid and safe desensitization in multifood allergic participants. Desensitization was achieved more successfully with omalizumab than placebo as an adjunctive therapy with OIT over 36 weeks in this trial [31]. In this current observational study, we addressed a different question. Can the maintenance dose be reduced after maintenance is achieved in OIT and still maintain the effect of desensitization? Many children have aversions to the high maintenance dose of each food allergen due to taste, convenience, sense of fullness, etc. A lower maintenance dose might increase not only an individual’s quality of life but also an individual’s adherence with continued ingestion of allergens long-term.

Here we present an observational, retrospective study of the LTFU of children after OIT with omalizumab as adjunctive therapy (omalizumab used for the first 16 weeks in the phase 1 study). The 34 participants were followed for a maximum of 62 months after they reached the 2 g maintenance dose in the original phase I trial [14], total up to 65 months after starting OIT. Throughout this time, the participants returned to the clinic on average every 6–12 months and OFCs up to 2 g for all offending foods were performed by a board-certified allergy and immunology specialist. All participants started the LTFU study at a maintenance dose of at least 2 g protein per food allergen and they either continued on this dose or reduced their dose to a low (median 300 mg) dose. This was not a randomized, placebo-controlled study but the long-term maintenance doses were decided throughout the study by the clinicians in collaboration with the parents of the participating children. Our reason for patient-based participation with a clinician was that this reflected more of a ‘real-life scenario’ and the decision to reduce the dose was not because of safety reasons but because of preference and ease of dosing compliance.

At the end of the LTFU study, of all 96 food allergens across the 34 participants, 74 (77%) were taken at a low, long-term maintenance dose (Fig. 3). Every participant passed the 2 g OFC to each of their offending foods at the end of the follow-up phase in May 2017. No severe respiratory, gastrointestinal or hypotension reactions
occurred after dosing and the overall rate of adverse events decreased over the course of the study.

This study has several strengths. The follow-up was done by in-person visits, including questionnaires but also standardized OFCs were performed to quantify the desensitization status for each food allergen. The study also has several limitations. The number of participants with each individual food and food allergen combination is too low to give adequate power for statistical analyses of these groups. Furthermore, the IgG4/IgE ratio or SPT wheal diameter data was limited in the study by some of the participants during LTFU (i.e. a convenience sample based on those individuals who agreed to blood and/or skin tests).

Conclusion
Our LTFU cohort of food OIT individuals who received omalizumab for the first 16 weeks during the original phase 1 trial supports the feasibility of long-term dosing in food-allergic individuals.

### Additional files

**Additional file 1: Figure S1.** Allergen-specific IgG4/IgE ratios for various time points during the dose escalation and after 2 g maintenance dose was reached. Each line represents one participant. The dots are colored by the dose at the specific time point.

**Additional file 2: Figure S2.** Wheal diameter of SPTs for various time points during the dose escalation and after 2 g maintenance dose was reached. Each line represents one participant. The dots are colored by the dose at the specific time point.

### Abbreviations
LTFU: long-term follow-up; OIT: oral immunotherapy; OFC: oral food challenge; SPT: skin prick test.

### Authors’ contributions
SA performed data curation, formal analysis, and writing of the manuscript. MM was involved in investigation, methodology, and writing of the manuscript. TD, WB, and RK were involved in care of participants and documentation of clinical data. DT performed data curation and data management. VS was involved in writing, editing, and reviewing the manuscript. SC was involved in care of participants, data management, and writing of protocol for IRB. KN was involved in funding acquisition, conceptualization of study,
analysis of data, and writing, reviewing, and editing of manuscript. All authors read and approved the final manuscript.

Acknowledgements
We thank Fran Meneses and Sally Mackey at the Sean N. Parker Center for Allergy and Asthma Research at Stanford University for their assistance.

Competing interests
The authors declare that they have no competing interests.

Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Consent for publications
Not applicable.

Ethics approval and consent to participate
Written consent was obtained from all participants in the study. The study was approved by Stanford IRB.

Funding
Monali Manohar, Sandra Andorf, Tina Dominguez, Dana Tupa, Rohun Kshirsagar, Vantha Sampath, Sharon Chinthrajah, and Kari Nadeau were supported by the NIAID grants R21AI095838, U19AI104209, Myra Reinhard family Foundation, the Monali Manohar, Sandra Andorf, Tina Dominguez, Dana Tupa, Rohun Kshirsagar, and the National Institute of Cancer. Common terminology criteria for adverse events (CTCAE)v. 4. 2009. https://evs.nci.nih.gov/ftp1/CTCAE/CTCAE_4.03_2010-06-14_QuickReference_Sx7.pdf. Accessed 2 Aug 2017.

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Received: 3 August 2017   Accepted: 7 December 2017
Published online: 21 December 2017

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