Oral Immunotherapy (OIT): A Personalized Medicine

Francesca Mori *, Simona Barni, Giulia Liccioli and Elio Novembre

Allergy Unit, Department of Pediatrics, Anna Meyer Children’s University Hospital, 50139 Florence, Italy; s.barni@meyer.it (S.B.); giulialiccioli@gmail.com (G.L.); e.novembre@meyer.it (E.N.)
* Correspondence: f.mori@meyer.it; Tel.: +39-055-566-2034

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Abstract: Oral Immunotherapy (OIT), a promising allergen-specific approach in the management of Food Allergies (FA), is based on the administration of increasing doses of the culprit food until reaching a maintenance dose. Each step should be adapted to the patient, and OIT should be considered an individualized treatment. Recent studies focused on the standardization and identification of novel biomarkers in order to correlate endotypes with phenotypes in the field of FA.

Keywords: children; desensitization; egg allergy; food allergy; milk allergy; oral immunotherapy; peanuts allergy; tree nuts allergy; wheat allergy

1. Introduction

Oral Immunotherapy (OIT), a promising allergen-specific approach in the management of Food Allergies (FA) [1], is based on the administration of increasing doses of the culprit food until reaching a maintenance dose, after which a regular intake of the specific food allergen is mandatory for “desensitizing” the patients to prevent exposure triggering an allergic reaction. While the main goal of the research is to induce sustained unresponsiveness or tolerance, OIT serves to introduce allergenic food into the normal diet, or in case of high-risk individuals, to introduce low doses in order to prevent severe reactions after accidental exposure [2].

In this regard, some researchers believe it is clinically and psychologically significant if a small amount of tolerance exists, enabling patients to tolerate accidental ingestion of the culprit food. Normally, FA is not just about tolerance or non-tolerance of the offending food. Recently, different phenotypes have been described, for example, patients who tolerate the allergenic food heated but are not able to tolerate the whole food uncooked. For this reason, OIT introduces new concepts such as the eliciting dose, the “matrix effect” (i.e., the complex interaction with other proteins, fats, and carbohydrates in the food matrix able to change protein allergenicity), and the way of processing foods. As a result, there is huge diversity among studies on the target food and the vehicle substance employed for OIT (i.e., some foods available commercially are used in their natural forms, while others use processed foods like defatted peanuts or dehydrated egg white), so the comparison among them could be difficult. Consequently, in the US, the Food and Drug Administration (FDA) has imposed standardization in terms of safety and quantification/identification of allergenic proteins. In fact, pharmaceutical-grade peanut flour is currently being subjected to phase 3 trials (AImmune) and is expected to be widely available in the coming years [3].

Another standardized drug for peanut OIT, called AR101, was studied in a phase 3 clinical trial that enrolled 551 patients with peanut allergy aged from 4 to 55 years [4].

Nonetheless, OIT should be considered an individualized treatment; hence its standardization in a general protocol represents a challenge for investigators and clinicians alike. Each step must be adapted to the patients’ specific situation (i.e., infections, gastrointestinal disorders, drug assumption, exercise, adverse reactions during treatment). It is strongly recommended to conduct an oral food
challenge (OFC) in order to establish the lowest reaction eliciting dose. Moreover, OFC could be used to assess the efficacy of desensitization while undergoing therapy, or functional tolerance when not on therapy and on a limited diet for a period of at least four weeks [5].

The initial doses for patients undergoing OIT are sufficiently low to avoid reactions and could be individualized or established for the whole study population [6,7].

Generally, the first step of OIT includes an escalation phase starting from micrograms of allergenic proteins and reaching the range of several milligrams in one to two days. The building up phase includes an increment of the dose twice or once a week until reaching a maintenance dose, or there is the onset of dose-limiting symptoms in the hospital setting under the supervision of healthcare professionals. This phase can last for varying amounts of time: from flash protocols (one week) to slow protocols (>6 months).

Low dose food challenges should be used in high-risk patients, while the more rapid introduction of an allergenic food could be performed in low-risk patients [8].

Individualized protocols may be recommended in patients with reactions to medium-high doses of the food in the challenge tests, thus making it possible to shorten the build-up phase, with savings in healthcare resources and greater comfort for the patient. However, the build-up starting dose with respect to the threshold dose has yet to be established (e.g., a cumulative dose of half a boiled egg or 50 mL of cow’s milk could be considered as the medium and high-tolerance thresholds) [9].

Moreover, enormous differences exist among studies in relation to the target maintenance dose range between 300 to 400 mg of allergenic proteins. In several studies, very high maintenance doses were administered (e.g., 4000 mg of peanut, equal to about 17 peanuts); however, the intake of high amounts of allergenic food is not clinically necessary and is frequently associated with a fear of reactions, and therefore, difficult to maintain because of aversion to the implicated food [10].

The target dose for milk OIT is usually 200–250 mL, even though 15 mL of milk may be considered the final milk dose in high-risk patients since it offers protection against minor accidental exposure and helps increase tolerance over time. An amount of 300 mg of powdered egg-white protein or the equivalent is considered safe for avoiding reactions deriving from traces, cross-contamination, or labeling errors [11].

In addition, the duration of the maintenance phase may be variable, as it could be prolonged for months to years and is characterized by the daily or at least biweekly administration of the offending food at home. Factors conditioning the duration of the maintenance phase are patient’s and families’ compliance and consistency, individual immunological response to an allergen, type of food and grade of allergenicity, the occurrence of an adverse reaction, etc.

Nevertheless, the extending of the maintenance phase can give rise to treatment compliance problems. In fact, drop-outs have been reported in about 60% of patients subjected to 3–5 years on cow’s milk and peanut OIT [12,13].

Moreover, it is not known what the outcome will be or how much time is required to reach permanent tolerance of the offending food. In the study on egg OIT, Jones et al. reported how tolerance is enhanced with the duration of OIT. The sustained unresponsiveness increases from 27.5% after two years of OIT, to 50% after four years of egg OIT [14].

With regard to the safety of OIT, over the last three years, major concerns have been examined in systematic reviews and meta-analyses, and it has been found that up to 91.5% of adverse reactions are observed during OIT in all patients treated and in 16% of the administered doses [15,16].

The majority of adverse reactions reported, which were mild and self-limiting, included itchiness of the mouth and lips, facial and generalized urticaria and erythema, abdominal symptoms, rhino-conjunctivitis, mild laryngeal spasms, and mild wheezing [17,18].

Severe anaphylactic reactions have been reported, and in the literature, the controlled clinical trials found that the intramuscular administration of epinephrine was necessary in 6.7% to 30.8% of all patients subjected to milk OIT, and in 20% of those subjected to egg OIT or peanut OIT [19,20].
Despite the fact that life-threatening reactions have been observed in asthmatic teenagers with poor compliance [19,20], several studies show that even though mild adverse reactions are quite common with OIT, they tend to become less frequent and less severe over time [14].

While adverse reactions normally occur with dose escalation, they are also possible during maintenance therapy with doses that were previously well-tolerated, due to being unpredictable and at times triggered by cofactors like infections, exercise, and anxiety [21,22].

In particular, adverse reactions could be the cause of patients withdrawing in 3–20% of cases of milk OIT and 0–36% of cases of egg OIT [10,23].

In addition, most withdrawals are due to the occurrence of gastrointestinal symptoms and not to anaphylaxis, with reports of eosinophilic esophagitis (EoE) in 2.7% of patients subjected to milk, peanut, egg and wheat OIT [7,12,22,24,25]. In this review, we focused, in particular, on milk, egg, and peanut because they are the most frequently implicated in food allergy in children.

2. Quality of Life (QoL)

The variable impact that OIT can have on the quality of Life (QoL) probably depends on the QoL at baseline. There was great improvement in patients with impaired QoL at the beginning, whereas deterioration was observed in those with acceptable QoL at baseline [26]. Moreover, the number of doses tolerated by patients seems to be inversely associated with QoL. The more allergic patients who had frequent reactions during the first phase of OIT, even with a few tolerated doses, failed to show any improvement in their QoL, compared to a restricted diet. Conversely, for those children who completed the OIT protocols, an improved QoL was reported [27].

3. Immunologic Changes with OIT

During OIT, the T helper (Th)2/Th1 ratio is reduced and the T regulatory effector cells increase with the production of Interleukin 10 (IL-10) by the APCs (antigen-presenting cells) and the activation of immune cells which together with Transforming growth factor-beta (TGF-β) induce production of Immunglobulin G4 (IgG4) and IgA. It is assumed that food-specific IgG4 during OIT could have an antigen-neutralizing effect and decreased basophil and mast cell responsiveness, with the suppression of IgE production. In allergen binding, both IgG4 and IgA compete with IgE, decreasing the allergen capture by basophils. This leads to a reduction in the amount of specific IgE, but also in the diversity of epitope recognition and altered affinity of IgE for antigens.

Decreased allergen-induced skin prick test (SPT) and basophil activation during the first few months of immunotherapy have been observed in OIT studies. However, a typical early increase in food-specific IgE levels has been demonstrated during the initial months of OIT. After 6 to 12 months of OIT, a transition has been observed from a Th2 predominant cytokine signature to a Th1-associated pattern, while immune suppression by T regulatory (T reg) cells and clonal anergy occur later during OIT. Syed et al. reported an increase in the function of antigen-specific CD4+CD25+Foxp3+T Treg cells following OIT, corroborating the theory of active suppression of the immune response by food allergens [24,25,28–33].

4. Sustained Unresponsiveness

Immunologic changes occurring during OIT-treatment seem to be temporary, revealing interindividual variability in immune suppression and clinical response. In egg OIT, from 71% to 90% of maintenance-phase patients retain desensitization after 1–6 years of follow-up [34–38]. In the literature, the length of follow-up reported in milk OIT ranges from 3 to 5.8 years. Desensitization to a full serving dose of cow milk (CM) equal to 200 mL is maintained in a range of between 31 and 100% of subjects [12,39–41]. The avoidance period before retesting tolerance has been described as being as long as 1–4 months. One study on peanut allergy reported that 50% of patients (3/6 individuals) who passed an initial challenge test after a three-month avoidance diet following successful completion of OIT, had a positive second challenge test after the avoidance period was prolonged for a further three
months [32]. Today, milk and peanut OIT are the most widely studied (Tables 1 and 2). There are very few publications on the egg (Table 3), other tree nuts (Table 4), wheat (Table 5), or fish OIT.

| Reference, Year | Design | Sample Size (n) | Subject | Maintenance Dose | Duration | Conclusions |
|-----------------|--------|----------------|---------|-----------------|----------|-------------|
| Meglio P. et al., 2004 [42] | open-label | 21 | 6–10 | 200 mL | 6 mon | 72% achieved desensitization to 200 mL of cow’s milk daily |
| Longo G. et al., 2008 [43] | randomized open-label | 30 | 5–17 | 150 mL | 10-day rush escalation, 1 yr maintenance | 36% completely tolerant (≥150 mL) and 54% partially tolerant (5–150 mL) |
| Skripak JM. et al., 2008 [11] | randomized, placebo-controlled | 13 | 6–17 | 500 mg milk protein | 23 wk | Median milk challenge threshold increased from 40 mg at baseline to 5140 mg after OIT |
| Narisety SD. et al., 2009 [44] | open-label (follow-up) | 13 | 6–16 | 500–4000 mg milk protein | 3–17 mo | Ongoing milk intake demonstrated tolerance from 1000 to 16,000 mg (median, 7000) with 33% tolerating 16,000 mg on OFC. |
| Pajno GB. et al., 2010 [45] | randomized, placebo-controlled | 15 | 4–10 | 200 mL | 18 wk | 67% tolerant to 200 mL cow’s milk |
| Martorell A. et al., 2011 [46] | randomized, placebo-controlled | 30 | 2–3 | 200 mL | 1 yrs | 90% showing complete desensitization |
| Keet CA. et al., 2012 [47] | randomized, placebo-controlled | 20 for OIT | 6–17 | 1000–2000 mg | 60 wk | 70% of patients receiving OIT passed an 8 g OFC; only 40% passed OFC when treatment was discontinued for 6 wk. |
| Goldberg M. et al., 2015 [48] | open | 14 | 6.5–12.7 | 1.3 g of BM protein | 12 mo | Only 3 (21%) of 14 patients tolerated the 1.3 g/d BM dose. Patients who successfully reached maintenance had decreased milk-specific IgE reactivity. |
| Takahashi M. et al., 2016 [49] | open | 31 (48 tot, 31 OIT, 17 controls) | 5–17 | 200 mL of microwave heated cow’s milk every day (fresh cow milk was warmed in a microwave oven at 550 W for 100 s) | 12 mo | No children in the untreated group did not pass an open food challenge to CM. Of the 31 children in the OIT group, 14 (p = 0.002) achieved desensitization, and eight (p = 0.036) achieved two-weeks-SU to CM at 1 year from the start of OIT. Two years after the start of OIT, both the rate of desensitization and the rate of the two-week-SU in the OIT group significantly increased compared with the rates at one year (p = 0.025 and p = 0.008, respectively). |

Table 1. Milk OIT studies.
| Reference, Year | Design | Sample Size (n) | Subject Age (yrs) | Maintenance Dose | Duration | Conclusions |
|----------------|--------|----------------|------------------|------------------|----------|-------------|
| Ebrahimi M. et al., 2017 [50] | open | 14 | 3.5–7 | 200 to 250 mL of cow’s milk each day for 90 days. | 90 days | The median of the difference of the wheel diameter with the control, decreased from 10 to 6 mm. After the OIT, the sIgE level of cow’s milk proteins and casein decreased from 39.30 to 10.40 and 7.72 to 2.83 (KU/L), respectively. The study doesn’t show data of sustained responsiveness in the follow-up. |
| Amat F. et al., 2017 [51] | randomized | 43 (18 high-risk arm, 23 low-risk arm) | 3–10 | “low-risk arm”: from extensively heated baked milk to the half-heated baked milk and then raw milk until 2720 mg of milk protein per day. “high-risk arm”: immediately raw milk | 9 mo | Fifteen children (36.6%) were classified as responders, 11 (26.8%) were partial responders, with an average gain in threshold of tolerance of 697 mg [27.2–2550], and 15 children (36.6%) remained non-responders. The study doesn’t evaluate sustained unresponsiveness to milk proteins. |
| Efron A. et al., 2018 [52] | retrospective, case-control | 43 (110 tot, 43 OIT, 67 controls) | 1–4 | First OFC—cookie containing ~1 g milk protein heated in frying and baking. Second OFC—pancake containing ~1 g milk protein heated in frying. Third OFC—toast containing ~4 g cheese proteins (mostly casein). Fourth OFC—yogurt containing ~4 gr of unheated cheese proteins. | 12–18 mo (3 mo each product) | At last follow-up, 88% of treated children were tolerant to unheated milk proteins vs. 52% of controls (p = 0.003). |
| Inuo C. et al., 2018 [53] | randomized, double-blind, controlled | 25 (13 pHF-pHF, 12 eHF-pHF) | 1–9 | two double-blind groups: a partially hydrolyzed cow’s milk protein-based formula (pHF)-pHF group and an extensively hydrolyzed cow’s milk protein-based formula (eHF)-pHF group | 16 wk | There was a significant increase in the threshold in the pHF-pHF group (p = 0.048), but not in the eHFpHF group (p = 0.23). Among the participants with a severe allergy, whose baseline thresholds were <4 mL, there was a significant change in thresholds between baseline and at the end of the trial in the pHF-pHF group (p = 0.023). |
| Mota I. et al., 2018 [54] | prospective | 42 | 2–18 | 200 mL | 36 mo | During the maintenance phase, 92% maintained diet without restrictions including daily ingestion of 200 mL of CM (36 of 39 adherent patients). Overall, 93% were adherent patients (39 of 42), since they keep daily ingestion of 200-mL CM. |
Table 1. Cont.

| Reference, Year | Design | Sample Size (n) | Subject Age (yrs) | Maintenance Dose | Duration | Conclusions |
|-----------------|--------|----------------|------------------|-----------------|----------|-------------|
| Kauppila T. et al., 2019 [55] | open | 180 (296 OIT, 64 controls) | 5–17 | 200 mL | 11 yrs of follow-up | Out of the initial study group, 244/296 (83%) patients participated in the long-term follow-up. Among these patients, 136/244 (56%) consumed ≥2 dL of milk daily. The median follow-up time was 6.5 years. Of the recorded markers and clinical factors, the baseline milk sIgE level was most associated with maintaining milk OIT (p < 0.001). |
| De Schryver S. et al., 2019 [56] | open | 26 (52 tot, 26 OIT and 26 controls) | 6–18 | 200 mL | 1 mo | Among the 26 children randomized to OIT, 18 were defined as desensitized to milk. The difference in the percentage of milk-desensitized children between the groups attributed to the OIT is 69.2% |
| Berti I. et al., 2019 [57] | open | 68 | 3–11 mo | up dosing until 150 mL | 3.5–16 mo | Sixty-six infants (97%) reached the target of the protocol |

Legend: y: years; mo: months; n: number; OFC: oral food challenge; OIT: oral immunotherapy; tot: total; wk: weeks; BM: baked milk; SU: sustained unresponsiveness.

Table 2. Peanut OIT studies.

| Reference, Year | Design | Sample Size (n) | Subject Age (yrs) | Maintenance Dose (mg) | Duration | Conclusions |
|-----------------|--------|----------------|------------------|-----------------|----------|-------------|
| Jones SM. et al., 2009 [24] | open-label | 29 | 1–16 | 1800 | 36 mo | 93% passed 3.9 g peanut OFC |
| Blumchen K. et al., 2010 [58] | randomized, open-label | 23 | 3–14 | 500 | 7-day rush escalation, 8 wk maintenance | 64% reached their maintenance dose of 500 mg peanut |
| Varshney P. et al., 2011 [25] | randomized, placebo-controlled | 19 | 3–11 | 2000 | 48 wk | 84% passed 5000 mg peanut OFC |
| Anagnostou K. et al., 2011 [59] | open-label | 22 | 4–18 | 800 | 32 wk | 64% tolerated 6.6 g OFC |
| Anagnostou K. et al., 2014 [60] | randomized, placebo-controlled | 39 | 7–16 | 800 | 26 wk | 62% tolerated 1400 mg challenge |
| Vickery BP. et al., 2014 [10] | open-label | 24 | 1–16 | ≤ 4000 | ≤ 5 y | 1 mo after OIT stopped, 50% achieved sustained unresponsiveness to 5000 mg OFC |
| Narisety SD. et al., 2015 [6] | randomized, placebo-controlled | 16 | 7–13 | 2000 | 12 mo | Significantly greater increase in OFC threshold in OIT vs. SLIT; low rate of sustained unresponsiveness 85% of patients passed the build-up phase, and 67% tolerated 5 g of peanuts during the post-treatment challenge |
| Kukkonen K. et al., 2017 [61] | double-blind, placebo-controlled | 39 (60 tot, 39 OIT and 21 controls) | 6–18 | 100-2000 | 8 mo | |
| Reference, Year | Design | Sample Size (n) | Subject Age (yrs) | Maintenance Dose (mg) | Duration | Conclusions |
|-----------------|--------|----------------|------------------|----------------------|----------|-------------|
| Vickery B. et al., 2017 [62] | double-blind, placebo-controlled | 40 (40 OIT and 154 controls) | 9–36 mo | 300-3000 | 29 mo | overall 78% of subjects receiving E-OIT demonstrated sustained unresponsiveness to peanut four weeks after stopping E-OIT and reintroduced peanut into the diet. 79% and 62% AR101 subjects tolerated >443 mg and 1043 mg respectively, versus 5 of 26 (19%) and 0 of 26 (0%) placebo subjects (both p < 0.0001) |
| Bird JA. et al., 2018 [63] | double-blind, placebo-controlled | 29 (tot 55, 29 OIT and 26 controls) | 4–26 | 300 | 20-34 wk | 250 of 372 participants (67.2%) who received active treatment, as compared with 5 of 124 participants (4.0%) who received placebo, were able to ingest a dose of 600 mg or more of peanut protein, without dose-limiting symptoms, at the exit food challenge |
| PALISADE group, 2018 [4] | double-blind, placebo-controlled | 372 (496 tot, 372 OIT and 124 controls) | 4–17 | 300 | 24 wk | Of the 145 patients treated, 113 (77.9%) were fully desensitized to 3000 mg of peanut protein, 20 (13.8%) patients were partially desensitized to 300-2400 mg, and 12 patients (8.3%) failed. 63/64 patients (98.4%) consuming 1200 mg maintenance dose were successfully re-challenged to 3000 mg. All patients in the high dose group (3000 mg) who continued regular consumption and arrived for follow-up (n = 22) passed a challenge to 3000 mg. |
| Nachshon L. et al., 2018 [64] | prospective | 139 (145 tot, 139 < 18 y) | 4–18 | 1200 or 3000 | 6 mo | 16 children (67%) passed the 133-mg OFC, and 14 (58%) passed the 795-mg OFC. Only 1 child (10%) in the historical control group passed the 133-mg OFC (p = 0.006). Ultimately, eight children (33%) in the OIT group achieved sustained unresponsiveness |
| Nagakura K. et al., 2018 PAI [65] | prospective, open-label | 24 (24 OIT, 10 controls) | 5–18 | 133 | 12 mo | 15/22 patients (68.1%) in the OIT group achieved sustained unresponsiveness, whereas only 2 (18.1%) in the control group passed the second OFC |
| Nagakura K. et al., 2018 [66] | double-blind, placebo-controlled | 22 (22 OIT, 11 controls) | 5–18 | 795 | 2 y | OIT participants who underwent dose variations on the unexpired lots of peanut flour were able to successfully tolerate the 100% dose increase, following a two-week tolerance of a 50% dose reduction on an unexpired lot of peanut flour |
| Anvari S. et al., 2018 [67] | double-blind, placebo-controlled | 15 | 5–16 | 3900 | 3 mo | |
Table 2. Cont.

| Reference, Year | Design | Sample Size (n) | Subject Age (yrs) | Maintenance Dose (mg) | Duration | Conclusions |
|-----------------|--------|----------------|------------------|----------------------|----------|-------------|
| Zhong Y. et al., 2018 [68] | open-label | 7 (9 total, 7 completed protocol) | 8–14 | 3000 | 12 mo | Of the seven who completed OIT, six tolerated 6000 mg of peanut protein at the first OFC at six months of maintenance phase; the last patient was afraid of consuming more than 3000 mg of peanut protein but passed the challenge with 3000 mg. After 12 months of maintenance therapy, only 3 of the 7 subjects consented to 4 weeks of abstinence. Of these, only 1 passed the challenge with 6000 mg of peanut protein. |
| Fauquert JL. et al., 2018 [69] | double-blind, placebo-controlled | 21 (30 tot, 21 OIT and nine controls) | 12–18 | 400 IN CAPSULES | 24 wk | Unresponsiveness to 400 mg of peanut protein was achieved in 17/21 peanut group patients (two patients withdrew) and 1/9 in the placebo group. |
| Blumchen K. et al., 2019 [70] | double-blind, placebo-controlled | 31 (62 tot, 31 OIT and 31 controls) | 3–17 | 125–250 | 16 mo | Twenty-three of 31 (74.2%) children of the active group tolerated at least 300 mg peanut protein at final food challenge compared with 5 of 31 (16.1%) in the placebo group (p < 0.001). Thirteen of 31 (41.9%) children of the active versus 1 of 31 (3.2%) of the placebo group tolerated the highest dose of 4.5 g peanut protein at final OFC (p < 0.001) |
| Wasserman RL. et al., 2019 [71] | retrospective record | 270 | 4–18 | 3000 | 36 mo | All patients who reached the 3000 mg target dose (214/262 81%) were challenged with 6000 mg of peanut protein and all but 1 patient passed the challenge. 14 had demonstrated sustained unresponsiveness with 6000 mg |

Legend: Y: years; mo: months; n: number; OFC: oral food challenge; OIT: oral immunotherapy; tot: total; wk: weeks; SLIT sublingual immunotherapy.

Table 3. Egg OIT studies.

| Reference, Year | Design | Sample Size (n) | Subject Age (yrs) | Maintenance Dose (mg) | Duration | Conclusions |
|-----------------|--------|----------------|------------------|----------------------|----------|-------------|
| Buchanan AD. et al., 2007 [72] | open-label | 7 | 1–16 | 300 mg | 24 mo | 57% passed 8 g OFC. 29% passed OFC after 3–4 mo period of egg avoidance |
| Vickery BP. et al., 2010 [73] | open-label | 8 | 3–13 | 300–3600 mg | 18–50 mo | 75% passed a 10 g OFC 1 mo after stopping OIT |
| Burks AW. et al., 2012 [28] | randomized, placebo controlled | 40 | 5–11 | 1600 mg | 22 mo | 75% passed 10 g OFC, but only 28% demonstrated SU on re-challenge 6–8 wk later |
| Escudero C. et al., 2015 [74] | double-blind, placebo-controlled | 30 (61 tot, 30 OIT, 31 controls) | 5–17 | 1 undercooked egg every 48 h | 3 mo | At 4 months, 1/31 (3%) in CG passed DBPCFC and 11/30 (37%) of OITG (95% CI, 14 to 51%; p = 0.003) |
Table 3. Cont.

| Reference, Year | Design, Placebo-Controlled | Sample Size | Subject Age (yrs) | Maintenance Dose | Duration | Conclusions |
|-----------------|----------------------------|-------------|-------------------|------------------|----------|-------------|
| Giavi S. et al., 2016 [75] | Double-blind, Placebo-Controlled | 29 | 1–5.5 | 9000 mg of low allergenic hydrolyzed egg (HydE) preparation | 6 mo | No statistically significant difference was observed on the final OFC (36% and 21% had a negative OFC in the treatment and placebo groups, respectively) |
| Yanagida N. et al., 2016 [76] | Open-label | 21 (33 tot, 21 OIT and 12 controls) | 5–18 | 62 to 194 mg (=1/32 of a heated whole egg) of egg protein in a scrambled form once daily | 12 mo | Respectively, 71% (15/21) and 0% (0/12) of the patients in the OIT and control groups exhibited sustained unresponsiveness to 1/32 of a whole egg 2 weeks after stopping OIT after 12 months ($p < 0.001$); 33% (7/21) and 0% (0/12; $p = 0.032$), respectively, showed sustained unresponsiveness to 1/2 of a whole egg. |
| Jones SM. et al., 2016 (Follow-up of Burks et al., 2012) [14] | Randomized, Placebo-Controlled | 40 (55 tot, 40 OIT and 15 controls) | 5–18 | 1600 mg | 22 mo | Of 40 E-OIT-treated subjects, 20 (50.0%) of 40 demonstrated SU by year 4. SU after E-OIT is enhanced with a longer duration of therapy and increases the likelihood of tolerating unbaked egg in the diet. |
| Pérez-Rangel I. et al., 2017 [77] | Double-blind, Placebo-Controlled | 15 (33 tot, 15 OIT and 14 controls) | 5–18 | 1 undercooked egg every 48 h | 5 mo | A total of 32 patients underwent the egg ROIT protocol (ROIT2). Thirty-one children (96.9%) completed the build-up phase, and 30 completed the maintenance phase, with a 93.8% rate of treatment success at five months |
| Akashi M. et al., 2017 [78] | Double-blind, Placebo-Controlled | 18 (36 tot, 18 OIT, 18 controls) | 3–15 | 4000 mg of dry egg powder | 6 mo | Eight of the 14 (57%) patients in the OIT group passed 4 g of dry egg powder whereas none of the 16 patients in the “eliminate egg” group |
| Maeta A. et al., 2018 [79] | Open-label | 13 | 3–8 | 10 LAC, each containing 79–110 mg of egg white protein | 4 mo | After the OIT, 2 participants tolerated 2 g of hard-boiled EW. Four participants did not show any improvement in response to OIT. |
| Itoh-Nagato N. et al., 2018 [80] | Double-blind, Placebo-Controlled | 45 | 5–15 | 60 g of cooked egg and 1 g of EWP | | The early start group received rush OIT for three months, while the late-start group continued the egg elimination diet (control). In the next stage, both groups received OIT until all participants had finished 12 months of maintenance OIT. The ratio of the participants in whom an increase of the TD was achieved in the first stage was significantly higher in the early-start group (87.0%), than in the late-start group (22.7%). |
Table 3. Cont.

| Reference, Year | Design          | Sample Size | Subject Age (yrs) | Maintenance Dose | Duration | Conclusions |
|-----------------|-----------------|-------------|-------------------|------------------|----------|-------------|
| Bird JA et al., 2019 [81] | open-label      | 13          | 1–18              | 3800 mg of BE    | 2 y      | Eight subjects completed 12 months of BE OIT, and seven subjects passed the 3.8 g BE OFC. After an additional year of daily 3.8 g BE ingestion, six subjects were challenged and 5 passed a 6 g LCE OFC. The study suggests that egg-allergic children reactive to BE may be able to undergo BE OIT to accelerate desensitization to LCE. |
| Martin-Muñoz MF et al., 2019 [82] | Double-blind, placebo-controlled | 76 (101 tot, 76 OIT and 25 controls) | 6–9 | 3300 g protein (30 mL of PEW) in 38 patients daily, in 38 patients every two days. | 12 mo | At T12, 4/25 (16%) of the total control patients passed the PEW DBPCFC vs. 64/76 (84.21%) OIT patients who had reached the target dose or total desensitization. (p = 0.000). At T24, 97.43% OIT patients passed the challenge. Daily OIT maintenance achieves better adherence, effectiveness, and safety |

Legend: BE: baked egg; EWP: egg white powder; mo: months; LAC: low egg-allergen cookies; OFC: oral food challenge; OIT: oral immunotherapy; tot: total; yrs: years; LCE: light cooked egg; ROIT: rush OIT; SU: sustained unresponsiveness; E-OIT: Egg-OIT; PEW: pasteurized egg white; CG: control group; OITG: OITgroup.

Table 4. Tree nut OIT study.

| Reference, Year | Design                             | Samples Size (n) | Subject Age (yrs) | Maintenance Dose (mg) | Duration | Conclusions |
|-----------------|------------------------------------|------------------|-------------------|-----------------------|----------|-------------|
| Elizur A et al., 2019 [83] | randomized, elimination diet controlled | 73               | 4–20 yrs          | 1200                  | 18 mo    | 89% desensitized (passed the OFC with 4000 mg of walnut) |

Legend: mo: months; n: numbers; yrs: years; OFC: oral food challenge.

Table 5. Wheat OIT studies.

| Reference, Year | Design            | Samples Size (n) | Subject Age (yrs) | Maintenance Dose | Duration | Conclusions |
|-----------------|-------------------|------------------|-------------------|------------------|----------|-------------|
| Rodriguez del Río et al., 2014 [84] | prospective, no control | 6                | 5–11              | 13 g           | 6 mo     | 85% desensitized |
| Sato S et al., 2015 [85] | prospective, historical control | 29               | Median age: 9     | 1300 mg starting dose | 24 mo    | 88.9% desensitized, 61.1% sustained unresponsiveness (passed the OFC with 4000 mg of wheat) |
| Okada et al., 2016 [86] | retrospective     | 57               | 1–11.8            | 400 mg          | 1 yrs    | 32 patients (86%) tolerated very low dose OFC (53 g of wheat protein) |
| Khayatzadeh A et al., 2016 [87] | case-control      | 13               | 5.5–19            | 5.2 g of wheat protein | Build-up phase: 3–6 days; maintenance phase: 3 months | 12 out of 13 completed maintenance phase: 12 out of 12 were desensitized |
| Rekabi M et al., 2017 [88] | prospective, no control | 12               | 2–10              | 30–70 g         | 18 months | 12 out of 12 patients tolerated 50 g of pasta |

Legend: mo: months; n: numbers; yrs: years; OFC: oral food challenge; OIT: oral immunotherapy; Baked Egg: BE.
5. Multiple Food OIT

A limit of OIT is represented by the fact that it is an allergen-specific approach, and most studies have been carried out on a single allergen, even though a great number of patients are sensitive to multiple allergens. For this reason, recent studies involving multiple foods have been described [89]. Moreover, in the case of tree nut allergy, cross-desensitization has been reported with the ingestion of only one type of nut thanks to the relevant cross-reactivity among tree nuts [83].

6. Desensitization Efficacy

A recent systematic meta-analysis of 31 clinical trials on food allergy [90] demonstrated the efficacy of desensitization, which entails an increase in the reaction threshold calculated as the food dose tolerated by the patient.

7. Personalized Medicine

It might be helpful to identify biomarkers associated with safe and successful OIT in order to select suitable subjects who are not expected to have reactions to OIT, screening out subjects in whom OIT could give rise to unnecessary risks. While it is likely that the outcome of OIT depends on numerous factors, several individual characteristics could have a predictive value.

There is evidence that the following patients are at high risk for failing OIT “desensitization”:

1. with IgE binding to a broader diversity of peptides;
2. with high IgE-binding intensity to allergens;
3. with the highest level of serum-specific IgE or the largest skin test response;
4. with more severe reactions at low doses;
5. with more severe asthma;
6. with persistent allergy (desensitization could prove to be more effective in small children, suggesting that it is easier to achieve immune modulation when started at an early age) [10,43,58,59,91,92].

There is evidence indicating that the following patients are more likely to successfully complete OIT “desensitization”, namely, those:

1. who are able to tolerate some form of allergen, e.g., patients eating cooked milk or cooked egg, may outgrow the overall allergy sooner [93–95];
2. who show a reduced skin prick test wheal size and an increase in specific IgG4-blocking antibodies after OIT to cow’s milk, egg, and peanut [14,24], with the latter possibly being a biomarker for sustained unresponsiveness [91];
3. who show a tendency towards a decrease in the specific IgE levels.

8. Conclusions

Clinical and immunopathological studies on FA-OIT focus on novel biomarkers and therapies in order to correlate FA endotypes with clinical phenotypes and propose the best-personalized treatment for each patient with FA.

Moreover, more research is necessary for understanding whether a longer course of OIT could increase tolerance rates and whether OIT only accelerates desensitization in subjects who would, in any case, progress towards natural tolerance without any intervention.

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