Emerging therapies for food allergy

Corinne A. Keet, Robert A. Wood


Food allergy is a common condition for which there are currently no approved treatments except avoidance of the allergenic food and treatment of accidental reactions. There are several potential treatments that are under active investigation in animal and human studies, but it is not yet clear what the best approach may be. Here, we review approaches that are currently in clinical trials, including oral, sublingual, and epicutaneous immunotherapy, immunotherapy combined with anti-IgE, and Chinese herbal medicine as well as approaches that are in preclinical or early clinical investigation, including modified protein immunotherapy, adjuvants, DNA vaccines, and helminth administration. We discuss the importance of fully exploring the risks and benefits of any treatment before it is taken to general clinical practice and the need for clarity about the goals of treatment.

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Introduction
Food allergy is defined as an immunologically mediated response that occurs repeatedly on exposure to a food (1). Up to 12% of children report an allergy to one or more foods, making it one of the most common chronic diseases of childhood (1). Although food allergy encompasses both IgE-mediated disease, typically characterized by acute symptoms such as hives or respiratory distress, and non-IgE-mediated conditions, such as milk-protein proctocolitis and eosinophilic gastrointestinal diseases, this review focuses on IgE-mediated allergy.

There are currently no approved treatments for food allergy except for avoidance of the allergenic food and treatment of accidental reactions. While much childhood food allergy is outgrown, a substantial proportion of even milk and egg allergy persists into adulthood, and some food allergies, such as peanut, tree nuts, and shellfish, are usually persistent (1). Reported rates of reaction due to accidental exposure vary widely, from 5% per year to 58% per five years for peanut (2, 3) and up to 80% per year for milk (4). Although mortality from food-related anaphylaxis is rare, estimated to be approximately 1.8 per million person-years among food-allergic subjects (5), fear of accidental reactions and social effects of avoidance diets contribute to significantly impaired quality of life for food-allergic children and their caregivers (6). Moreover, avoidance diets can leave food-allergic children at nutritional risk (7). It is estimated that food allergy costs the US almost 25 billion dollars per year in direct and indirect expenses (8).

For all of these reasons, effective treatment for food allergy would be highly desirable. However, while potential treatments are under active investigation, it is still not at all clear what the best approach may be. Further, there is lack of consensus as to what the goals of therapy should be: while the ability to incorporate the food into the diet would be ideal, some would argue that simply minimizing the risk of reaction with accidental exposure would be of sufficient value to justify treatment. These questions are vital because we need criteria by which to evaluate novel treatments, especially if they carry risk of serious adverse reactions. In fact, whether the emerging therapies for food allergy are sufficiently beneficial to justify any significant risk remains to be determined.

Conflict of interest: Robert A. Wood receives royalties from Up-To-Date, is a consultant for the Asthma and Allergy Foundation of America, and is on the Medical Advisory Board for the Food Allergy Research and Education Foundation.

Citation for this article: J Clin Invest. 2014;124(5):1880–1886. doi:10.1172/JCI72061.

Pathophysiology of food allergy
IgE-mediated food allergy is characterized by Th2-dominant immunologic responses, with allergen-specific IgE present in circulating forms and bound to mast cells and basophils. Allergen-specific, Th2-deviated CD4+ T cells predominantly produce cytokines, such as IL-4, IL-5, IL-9, and IL-13, which promote IgE production, eosinophil proliferation, and trafficking of inflammatory cells to tissues. The predominant mechanism by which food allergy develops remains controversial, with some studies suggesting that primary sensitization through skin contact may be even more important than exposure via the gut (9). Active gastrointestinal tolerogenic mechanisms appear to be important in preventing food allergy in general, as children with genetic defects in generating regulatory T cells frequently have severe allergic disease (10, 11). APCs in the gut, particularly DCs, clearly direct these T cell responses and are themselves responsive to the context in which they receive antigen. Abnormal function of both DCs and T cells has been linked to food allergy (12–20). Contextual clues that influence DC responses include costimulatory signals through a variety of receptors, including the TLRs. These signals come from multiple sources, including those associated with tissue damage, commensal bacteria, and the allergen itself (20, 21). Responses to these signals may vary considerably due to genetic predisposition (22).

Although basic science has been helpful in understanding the mechanisms by which potential treatments for food allergy might work, to date, the most promising therapies have not come as a result of these discoveries but instead from clinical observation and the modification of therapies previously developed for other allergic diseases. For example, immunotherapy, discussed at length below, was first described as a treatment for IgE-mediated allergic disease in 1911 (23), more than 50 years before the discovery of IgE (24). Some therapies, such as the recombinant peanut vaccine described below, emerged from basic science and appeared effective in animal models, but failed when tested in humans. Still other approaches, such as DNA vaccines, which appear promising in animal models, may be difficult or impossible to safely translate to humans. In the following sections, we will first review approaches that are under active clinical investigation, after which we will review potential approaches that are in preclinical or early clinical investigation.

Current clinical investigations
Current clinical investigations are summarized in Table 1. For venom and Aero allergies, immunotherapy using intact, and often

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Review
The proliferate of small and medium-sized studies of these methods was not published until 2005 (30). In recent years, there has been an increase in the number of studies examining the efficacy and safety of oral immunotherapy. A Cochrane review of trials of oral immunotherapy in children found that 70% of treated subjects increased their food challenge threshold (6). The results of these studies are promising, with 60% of treated subjects tolerating a 5-gram challenge at one year compared with none of the placebo subjects (44). Most of the studies have been done in school-aged children, but a randomized trial of 60 children found that milk oral immunotherapy was also effective in young children, with 90% of treated subjects tolerating a 5-gram challenge at one year compared with none of the placebo subjects (44). Fewer studies have been done with sublingual immunotherapy, but thus far it appears that efficacy is much less than with oral immunotherapy. A CoFAR placebo-controlled study of peanut sublingual immunotherapy in 40 adolescents and adults found that 70% of treated subjects increased their food challenge threshold.

These findings are similar to other controlled studies of oral immunotherapy. A multi-center, randomized, placebo-controlled trial of egg oral immunotherapy conducted by the Consortium for Food Allergy Research (CoFAR) in 55 children found that 55% were able to consume a 5-gram serving of egg after 10 months of therapy compared with 0% of placebo-treated patients. After an additional 12 months of therapy, 75% of those on active treatment were able to consume a full 10-gram serving (53). For peanut, a randomized, placebo-controlled trial of oral immunotherapy in 28 children found that 84% were able to tolerate a 5-gram challenge at one year compared with none of the placebo subjects (44). Most of the studies have been done in school-aged children, but a randomized trial of 60 children that milk oral immunotherapy was also effective in young children, with 90% of treated subjects tolerating a 5-gram challenge at one year compared with none of the placebo subjects (44). Fewer studies have been done with sublingual immunotherapy, but thus far it appears that efficacy is much less than with oral immunotherapy. A CoFAR placebo-controlled study of peanut sublingual immunotherapy in 40 adolescents and adults found that 70% of treated subjects increased their food challenge threshold.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Selected recent clinical studies</th>
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<tr>
<td><strong>Allergen-specific immunotherapies</strong></td>
<td>Drug name</td>
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<tr>
<td>Oral immunotherapy</td>
<td>N/A</td>
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<tr>
<td>Sublingual immunotherapy</td>
<td>NA</td>
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<tr>
<td>Oral immunotherapy plus omalizumab (anti-IgE)</td>
<td>Omalizumab</td>
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<tr>
<td>Recombinant protein</td>
<td>EMP-123</td>
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<tr>
<td>Epicutaneous patch</td>
<td>ViaSkin, others</td>
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<tr>
<td><strong>Nonspecific</strong></td>
<td>Chinese herbal medicine</td>
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<td>Anti-IgE</td>
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AE, adverse event.
old by at least 10-fold at 44 weeks, compared with 15% of placebo-treated subjects, but none of the treated subjects were able to pass a full food challenge at that time (47). Our group compared milk oral immunotherapy to sublingual immunotherapy in 30 children and found that after approximately 18 months, 70% of oral immunotherapy subjects could tolerate a full serving of milk compared with 10% of sublingual immunotherapy subjects (76). However, sublingual immunotherapy appears to be safer than oral immunotherapy, with significantly fewer multi-system, gastrointestinal, and lower respiratory reactions (76).

It is likely that the differences between oral immunotherapy and sublingual immunotherapy, with regard to both efficacy and safety, primarily represent differences in the antigen doses that are used, given that typical sublingual immunotherapy maintenance doses are under 10 mg compared with 1 to 4 grams for oral immunotherapy. Improved sublingual immunotherapy efficacy might therefore be possible if higher doses were used, and in theory, an ideal sublingual immunotherapy dose might achieve efficacy similar to oral immunotherapy at far lower — and therefore safer — doses, given the high density of tolerogenic APCs in the sublingual space (77). However, significantly higher sublingual immunotherapy doses will not be possible unless more concentrated extracts or alternative delivery systems, such as the tablets that have been formulated for grass pollen sublingual immunotherapy (78), are developed. Until then, maximum sublingual immunotherapy dosing will remain limited by the concentration of the available aqueous extracts and the volume of liquid that can be safely administered sublingually.

The mechanisms of action underlying oral immunotherapy and sublingual immunotherapy are not entirely clear, although it is likely that a combination of suppressive mechanisms, anergy, and deletion of reactive T cells is important (refs. 21, 35, and Figure 1). With treatment, allergen-specific IgE tends to rise initially and then fall marginally by the completion of treatment. Specific IgG4 levels increase, and markers of basophil activation and mast cell reactivity (as evidenced by skin test responses) typically decrease (47, 53, 76). Although these changes occur in most subjects, especially with oral immunotherapy, and may be associated with more positive outcomes, the data thus far do not allow for any of these measures to be used as reliable biomarkers of clinical response.

Although short-term desensitization appears to be common with oral immunotherapy, the prospect for complete resolution of allergy — or even a sustained period of unresponsiveness (also sometimes called "clinical tolerance") — appears to be limited with current methods. Although the persistence of desensitization after avoidance of the allergen has only been assessed in a few studies, the results are sobering overall. In our study of milk allergy, 40% of subjects desensitized with oral immunotherapy remained unresponsive at six weeks, and some regained reactivity within a week (76). Similarly, in the egg oral immunotherapy study, only 36% of desensitized subjects had sustained unresponsiveness four to six weeks after the discontinuation of therapy (39). This concept is very important because if sustained protection is not the norm, the long-term safety of food immunotherapy after treatment may be far more problematic than the known short-term risks during treatment. Although data on long-term outcomes are limited, we recently reported in a follow-up of two milk oral immunotherapy studies three to five years after study completion that only 25% of subjects were consuming normal servings of milk without any symptoms, that almost 30% were having regular or predictable symptoms, and that, most concerning, almost 20% had anaphylaxis during the follow-up period, including some who appeared to have had an excellent response to treatment (79). While ongoing consumption of the problem food may afford protection for most, this does not appear to always be the case and, even if it were, this is something that can be difficult to achieve, even in the research setting (80).

**Anti-IgE therapy.** Anti-IgE agents such as TNX-901 and omalizumab, which is licensed for the treatment of asthma, have been shown to increase the food challenge threshold to peanut. These treatments have the advantage over immunotherapy that they are non-allergen
CD63 expression (104, 106). A controlled efficacy study is currently being conducted in adolescents and adults (ClinicalTrials.gov identifier: NCT00602160), and studies using FAHF-2 along with oral immunotherapy will hopefully begin in the near future.

**Novel therapeutic strategies in preclinical or early clinical investigation**

**Modified protein immunotherapy.** Immunotherapy with proteins that are modified so that IgE-binding epitopes are removed or significantly altered, while maintaining relevant T cell binding, could provide efficacy similar to that of the unmodified protein, with an improved safety profile. This approach could theoretically make it possible to induce tolerance with far shorter courses of therapy by safely providing higher doses of the tolerogenic epitopes with little or no need for gradual dose escalation. Two general approaches are under investigation, one relying on the modification of the IgE-binding sites to reduce reactivity and the second based upon the identification of specific tolerogenic epitopes that are spliced from the larger molecule and provided as “peptide immunotherapy.”

The latter approach is now in phase 3 trials for cat allergy (107, 108) and is being actively pursued in preclinical trials for other allergens, including egg and fish (109, 110).

The former approach has been applied to peanut allergy by developing a recombinant vaccine in which the IgE-binding epitopes on the three major peanut allergens, Ara h1, h2, and h3, were modified by single amino acid substitutions, then encapsulated in inactivated *E. coli* (EMP-123). In mouse models, the modified allergens did not bind IgE or induce basophil reactivity (111); however, they reduced anaphylactic symptoms on rechallenge with allergen in sensitized animals (112). In spite of the encouraging results in the animal model, a recent phase I trial of EMP-123 administered per rectum was disappointing (113). In fact, acute allergic reactions were so common that five of ten peanut-allergic subjects were unable to complete dosing, indicating that IgE binding was not adequately reduced by this modification.

**Adjuvants.** Another promising approach is to combine the immunotherapeutic protein — either intact or modified — with other components to enhance tolerogenic responses. For example, the TLR9 agonist CpG oligodeoxynucleotide leads to a Th1-skewed DC response. Further, linking CpG with allergen has reduced anaphylactic reactions to allergen when used either as a preventative therapy or as treatment in the form of s.c. injection or e.c. immunotherapy (114–118) in murine models. In humans, a similar TLR9 agonist, phosphorothioate oligodeoxynucleotide DNA, was coupled to ragweed antigen and showed some efficacy in a pilot s.c. injection study (119). Another potential adjuvant is chitosan, a polymer commonly found in the cell walls of fungi and many invertebrates, which activates macrophages and modulates Th2 inflammation (119–121). Mice fed chitosan did not exhibit orally induced peanut sensitization and had reduced IL-4, IL-5, and IL-10 production and increased IFN-γ (119–121). A more direct approach is to fuse inhibitory human TLR9 ligands to Ara h2 causes aggregation of the inhibitory receptor FcγRIIB and the high-affinity IgE receptor FcεRI and thus inhibits degranulation of mast cells and basophils. In a murine model, s.c. treatment with FcγRI-Ara h2 inhibited anaphylaxis to peanut (122). However, the likelihood of efficacy in humans and the long-term disease-modifying potential of these therapies are unclear. Although we are not aware of any current human trials using these or other adjuvant strategies for the treatment of food allergy, it is critical that

**Adjuvants**

**Food Allergy Herbal Formulas (FAHF)**

- **FAHF-2** is a novel formulation of nine herbs that are commonly used in Chinese traditional medicine for the treatment of gastroenteritis, asthma, and allergic rhinitis (102).
- One component, *Rubia cordifolia*, inhibited peanut-triggered IgE production, while another, *Dianthus superbus*, inhibited peanut-induced anaphylactic reactions. The mechanism appears to be partially related to suppression of Th2 responses and enhancement of Th1 responses, as production of IL-4, IL-5, and IL-13 was significantly reduced with treatment, while IFN-γ production increased (103).
- In the murine model, suppression of anaphylactic reactions lasted at least six months after treatment (104, 105).

Preliminary data from human pilot studies suggest that FAHF-2 is safe and well tolerated and showed immunologic effects, including decreased IL-5 levels and basophil activation, as measured by
DNA vaccines. A distinct immunotherapeutic approach is to eschew protein administration altogether and instead provide exposure to allergen in the form of DNA. Using a variety of vectors, DNA-encoding allergen is administered and then incorporated into APCs, where it is ultimately translated into protein, potentially leading to Th1-biased responses (123). In a mouse model, oral gene delivery with chitosan-DNA nanoparticles protected against the development of peanut allergy (123). In humans, DNA vaccines for infectious diseases exhibit only modest immunological effects and have not yet shown efficacy when used alone (124). For allergy, a DNA-plasmid vaccine for treatment of Japanese red cedar allergy is currently in phase I trials (ClinicalTrials.gov identifiers: NCT01707069 and NCT01966224). Although the limited human trials conducted thus far have shown good tolerability, the possibility that DNA may be integrated into the host genome remains a concern and may limit application to food allergy. Further, this approach shows more promise for prevention than it does for treatment, and while preventative strategies are certainly desirable, more general approaches would be preferable to food-specific approaches such as these.

Helminth administration. In an extension of the hygiene hypothesis, which states that the relative lack of infectious exposures of modern life has contributed to the epidemic of allergic diseases, some researchers have tried administration of helminths for control of allergic diseases, including allergic rhinitis and atopic dermatitis. Helminths secrete a variety of factors known to inhibit B cell and mast cell activity (125). In small studies of allergic and autoimmune diseases done thus far, administration of *Trichuris suis* (pig whipworm) was relatively well tolerated, with some gastrointestinal side effects and eosinophilia, but efficacy for allergic diseases has not yet been shown (125).

**Effects on clinical practice**

Although the FDA has not licensed any of these treatments for food allergy and most experts strongly oppose their use in current clinical practice, both oral immunotherapy and sublingual immunotherapy are being used increasingly in practice settings (126). Is this a justifiable practice? On the one hand, there is accumulating evidence that desensitization can be widely achieved with current protocols. On the other, the safety data we do have indicate that oral immunotherapy is a more risky treatment than we normally tolerate for diseases of this severity, with a relatively high per-patient rate of systemic reactions. Reactions can be unpredictable, with factors such as illness and menses that can lead to reactions with previously tolerated doses (34). Unlike s.c. injection, oral immunotherapy and sublingual immunotherapy are administered at home, without medical supervision. Most concerning, however, is our lack of understanding of the long-term trajectory of patients treated with oral immunotherapy or sublingual immunotherapy and whether they are durably protected from serious reactions. Patients and their families who choose these therapies may not fully understand the risk/benefit ratio, especially if they are being treated in clinical settings without informed consent. Although some patients and providers may believe that these therapies, especially oral immunotherapy, will limit the risk of serious or fatal reactions from food, there are currently no data that show a decreased rate of serious life-threatening reactions with oral immunotherapy, much less cost-effectiveness. In fact, all placebo-controlled trials show a higher rate of serious reactions overall in treated subjects, and it is our clear impression that the risk of significant adverse reactions is far higher in those being treated than in those practicing strict avoidance (47, 53). These risks may be justified depending on the long-term treatment outcome, but long-term results are currently unavailable, and studies to date are certainly not reassuring. More research is clearly needed to understand both sides of the risk/benefit equation and to optimize therapy to reduce risks. In addition, research should address patient preferences and quality of life with these therapies.

**Future directions**

Future research will likely maintain some form of immunotherapy at its core and focus on increasing efficacy and/or safety. It is possible that some therapies may provide a level of desensitization that protects from accidental reactions but does not eliminate all reactivity. It may also be possible that long-term, even lifetime, treatment with all of these therapies may be needed to sustain protection. Adjuvants, recombinant or pepplexins, DNA vaccines, and/or cotreatment with anti-IgE or Chinese herbs could all allow for safer and more effective therapy, but additional research is clearly needed. In the next decade, despite concerns about safety and efficacy, we do anticipate the wide use of food immunotherapy in general practice, but our hope for the next two to three decades is that therapies can be developed that are both safer and more effective and include the induction of sustained protection, or even true immunologic tolerance, for the vast majority of patients with persistent, severe food allergy.

**Acknowledgments**

Corinne A. Keet receives research support from the National Institute of Allergy and Infectious Diseases (NIAID), the National Institute of Child Health and Human Development (NICHD), and the Gerber Family Foundation. Robert A. Wood receives research support from the NIAID, the Initial Training Network (ITN), and the Food Allergy Research and Education Foundation. This work was funded in part by grant 1K23AI103187-01 (NIAID).

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