Designer covalent heterobivalent inhibitors prevent IgE-dependent responses to peanut allergen

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Allergies are a result of allergen proteins cross-linking allergen-specific IgE (sIgE) on the surface of mast cells and basophils. The diversity and complexity of allergen epitopes, and high-affinity of the sIgE–allergen interaction have impaired the development of allergen-specific inhibitors of allergic responses. This study presents a design of food allergen-specific sIgE inhibitors named covalent heterobivalent inhibitors (cHBIs) that selectively form covalent bonds to only sIgEs, thereby permanently inhibiting them. Using screening reagents termed nanoallergens, we identified two immunodominant epitopes in peanuts that were common in a population of 16 allergic patients. Two cHBIs designed to inhibit only these two epitopes completely abrogated the allergic response in 14 of the 16 patients in an in vitro assay and inhibited basophil activation in an allergic patient ex vivo analysis. The efficacy of the cHBI design has valuable clinical implications for many allergen-specific responses and more broadly for any antibody-based disease.

IgE | allergy | epitope | selective inhibition | peanut

Allergy symptoms can vary from a rash to a potentially fatal anaphylactic reaction. Food allergies, especially peanut allergy, fall into the latter. These types of allergic reactions (IgE-dependent type-I hypersensitivity reactions) can lead to life-threatening systemic inflammatory responses such as anaphylactic shock (1, 2). The critical step in a peanut-allergy reaction is the binding of peanut allergens to allergen-specific IgE antibodies (sIgEs) attached to the high-affinity Fc epsilon receptors (FceRI) on mast cell surfaces that results in their cross-linking. This cross-linking triggers intracellular signaling that culminates in cellular degranulation that releases preformed mediators stored in cytoplasmic granules, including vasoactive amines, neutral proteases, proteoglycans, cytokines, and chemokines (3).

Currently, in the clinic, there are no medications that can prevent the degranulation response from mast cells, aside from a nonspecific IgE depleting therapy, wherein omalizumab, a monoclonal antibody, binds and inhibits all IgE from binding the surface receptor. Although it has shown effectiveness in clinical treatment of asthma, omalizumab has not been approved for the treatment of peanut allergies (4, 5). Furthermore, the inhibition of all IgE can lead to nonspecific immune suppression, leading to increases in parasitic infection and cancer (6, 7). A targeted and specific sIgE inhibitor would alleviate these concerns. One strategy for inhibition of peanut-allergy reactions (and any other food allergy) is to prevent the binding of allergen epitopes to sIgEs, but there are currently no functional examples of specific sIgE inhibitory compounds in the clinic or scientific literature.

The major challenge in developing targeted sIgE inhibitors is to identify the critical immunogenic epitopes from the large number of potential epitopes on allergen proteins (8, 9). Literature reports of several sIgE binding epitopes of the two most immunogenic peanut proteins, Ara h 2 and Ara h 6, exist (10, 11). These studies, however, did not assess the relative importance of those epitopes for IgE-dependent activation, instead only determining that they bind sIgEs. Here, we developed a physiologically relevant, nanoparticle-based assay system we termed nanoallergens, to identify the most critical immunogenic epitopes of Ara h 2 and Ara h 6. Next, we used this information to rationally design a class of sIgEs inhibitors, named covalent heterobivalent inhibitors (cHBIs), to irreversibly bind to and inhibit peanut allergen/sIgE interactions, thereby selectively and exhaustively inhibiting peanut-allergy reactions (SI Appendix, Fig. S1). We demonstrated, by using peanut allergic patient sera, that cHBIs specifically and irreversibly target the most crucial sIgEs for allergic responses and yield almost complete inhibition of degranulation responses to peanuts in cellular assays with patient serum. This study presents compelling preclinical data for further development and assessment of the cHBI strategy as inhibitors of peanut allergies and other food allergies to improve patient outcomes.

Results

Identification of Immunodominant Ara h 2 and Ara h 6 Epitopes Using the Nanoallergen Assay. To facilitate design of cHBIs, we first identified the most critical immunogenic epitopes from Ara h

Significance

IgE antibody interaction with allergen proteins is the main driver for life-threatening allergic responses. Using a nanoparticle-based method, we identified the most crucial binding sites for IgEs on peanut proteins by testing a small population of patient serum with a clinical history of severe allergies to peanuts. We then synthesized inhibitors, we call covalent heterobivalent inhibitors, which specifically and permanently prevent IgEs from binding to peanut proteins and triggering allergic responses. This is significant as it seems that only a few binding sites on peanut allergen proteins drive allergic responses. Also, this is a functional example of selective IgE inhibition to a relevant food allergen and establishes cHBIs as a class of allergy therapeutics.


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2 and Ara h 6 using the nanoallergen assay system developed in our laboratory (12). Nanoallergens provide multivalent display for allergen epitopes on liposomal surfaces, which are then used to evaluate their immunogenicity via in vitro mast cell degranulation responses (Fig. 1A). Currently used microarray techniques only determine if IgE binds a particular epitope but lack proper epitope display and valency to trigger cellular degranulation (13). Nanoallergens overcome these shortcomings and provide a physiologically relevant assay system to evaluate immunogenicity of epitopes by directly comparing cellular responses not just relative binding.

Nanoallergen preparation required synthesis of epitope-lipids, for which potential epitopes (synthesized as linear peptides) are conjugated to palmitic acid tails via ethylene glycol (EG). The epitope-lipid was purified, mixed at precise ratios with phospholipids [1,2-distearoyl-sn-glycero-3-phosphocholine (DSPC)] and extruded through membranes to form 100-nm nanoparticles (SI Appendix, Fig. S2). These nanoallergens multivalently present epitopes on their surfaces with high-precision and controlled stoichiometry (12, 14). Nanoallergens were used to trigger degranulation response in RBL-SX38 cells [model mast cell line which expresses human IgE receptor (FceRI)] (15) primed with a specific patient serum (Fig. 1A). This assay enables patient-specific assessment of epitope immunogenicity, rather than only assessing binding interactions, and hence accurately identifies the most antigenic epitopes.

We synthesized nanoallergens which present epitopes for 10 commonly cited peanut protein epitopes in the literature, epitopes 1–5 of Ara h 2, and epitopes 1–5 of Ara h 6 (SI Appendix, Table S1) (10, 11, 16–18). To evaluate these epitopes, we used human serum from 16 peanut-allergic patients obtained from PlasmaLabs and Massachusetts General Hospital (SI Appendix, Table S2). Serum samples were selected based on clinical history of peanut-allergic responses (as verified by commercial source or by food challenge) and a peanut sIgE >50 kU/L as determined by ImmunoCAP (SI Appendix, Table S2). Serum from each patient was used to prime RBL-SX38 cells before incubation with nanoallergens (0–5 nM) presenting each of the epitopes (SI Appendix, Fig. S3). Degranulation was monitored via beta-hexosaminidase assay and EC₅₀ values were determined (Table 1 and SI Appendix, Table S3). Our results identified Ara h 2 epitope 5, and Ara h 6 epitope 5 to be immunodominant, given their low EC₅₀ (EC₅₀ < 10 pM for Ara h 2 epitope 5 and EC₅₀ < 100 pM for Ara h 6 epitope 5) for most patient samples. Note that as demonstrated previously (12), degranulation intensity does not correlate with epitope dominance due to inhibitory cascades [such as Src homology 2 (SH2) domain containing inositol polyphosphate 5-phosphatase 1 (SHIP)] that attenuate degranulation (19). Hence, EC₅₀ values were used as the primary metric for epitope dominance.

Identification of Potentially Cross-Reactive Peanut Protein Epitopes

via Analysis of Epitope Sequence Homology. The major challenge of preventing allergic reactions with sIgE inhibitors is the large number of epitopes on allergy proteins that could bind to a diverse set of sIgEs (20). In peanuts, among the 13 known immunogenic proteins, Ara h 2 and Ara h 6 are the immunogenic proteins most closely associated with severe allergic reactions (21). Other proteins such as Ara h 1 and Ara h 3, however, are also immunogenic, albeit triggering less severe reactions (21). While it is not practical to develop inhibitors for each epitope/sIgE interaction, it is known that allergens often have homologous epitopes between two different proteins, and sIgEs can be

![Fig. 1.](image-url) cHBIs are designed to selectively and irreversibly bind sIgEs and inhibit cellular degranulation. (A) Cartoon demonstration of the nanoallergen assay system for evaluating potential immunogenic epitopes. Potential epitopes of allergen proteins are synthesized as linear peptides conjugated to lipid tails (epitope-lipid). Epitope-lipids are then incorporated during liposome synthesis to produce nanoallergens. Nanoallergens are used to screen for epitope immunogenicity in cellular degranulation assays. (B) cHBI design. cHBIs were synthesized to display an immunogenic epitope that binds to the ABS of sIgEs, a NBS ligand that binds to the NBS of sIgEs, and an aminofunctional-lysine-reactive moiety (ITC) to facilitate irreversible binding to sIgEs. Representative crystal structure of an Ig with the ABS, NBS (orange), and a lysine proximal to NBS (black) is shown. The zoomed-in Fab region describes cHBIs bivalent NBS and ABS binding, which promotes covalent conjugation to the proximal lysine. (C) Chemical structure of cHBI. The immunodominant epitope, NBS ligand, and ITC moiety are linked using flexible EG linkers to facilitate bivalent binding and covalent bond formation to sIgEs. (D) Schematic illustration of allergy inhibition by cHBIs. After initial bivalent binding to ABS and NBS, cHBI forms a covalent bond with sIgEs via ITC cross-linking of the nearby lysine, thereby inhibiting cellular degranulation.
potentially designed to have cross-reactivity for several allergenic proteins simultaneously (17, 22). We noticed a high level (>50%) of homology between the most immunogenic epitopes identified in our nanoallergen analysis, Ara h 2 epitope 5 and Ara h 6 epitope 5, and epitopes from Ara h 1, Ara h 2, Ara h 3, and Ara h 6 (SI Appendix, Table S4). Hence, we hypothesized that by developing inhibitors of immunodominant epitope/sIgE interactions, namely Ara h 2 epitope 5 and Ara h 6 epitope 5, we could inhibit epitope/sIgE interactions for a broad range of immunogenic epitopes because of homology and possible cross-reactivity with sIgEs and improve the therapeutic potential of cHBI.

Rational Design and Synthesis of cHBIs to Inhibit Binding of sIgEs to Immunodominant Allergy Epitopes. Next, we engineered cHBIs that irreversibly inhibit binding of sIgEs to these two immunodominant epitopes. The cHBI design required three moieties linked by ethylene-glycol spacers: (i) the immunodominant epitopes from Ara h 2 and Ara h 6 to target the antigen binding site (ABS), (ii) a ligand to target the nucleotide binding site (NBS), and (iii) a reactive functional group (RFG) to form a covalent bond with target sIgEs (Fig. 1 B and C). NBS is a conserved binding site located proximal to the ABS between the heavy and light chains of all immunoglobulins (Fig. 1B) (23, 24). We identified a molecule, indole-3 butyric acid (IBA) that specifically binds to NBS with moderate affinity ($K_d = 2$–$10 \mu M$) (23, 25, 26). Targeting two independent sites on sIgEs simultaneously, ABS (with a peptide epitope) and NBS (with IBA), enables cHBIs to have both increased avidity and selectivity for target sIgEs. The moderate affinity of IBA minimizes its binding to off-target antibodies and requires an ABS ligand for effective inhibition, while the bivalent binding to both sites improves overall avidity and selectivity for sIgEs (23). The third critical design factor for cHBIs is the RFG which facilitates covalent bond formation after a cHBI binds bivalently to target sIgEs, thereby permanently inhibiting sIgE/allergen interaction. For this purpose, we chose an amine-reactive moiety, isothiocyanate (ITC), as the RFG in the design of cHBIs (Fig. 1C). ITC compounds are common in nature and are nontoxic (27). ITC groups form thiourea bonds with primary amines, such as lysines, in elevated pH solutions (>9) but have limited reactivity at pH 7.4 ($t_{1/2} > 50$ h) (28). Consequently, under physiological conditions, ITC has negligible reactivity with random lysines, reducing off-target conjugations with other serum proteins. Furthermore, this limited reactivity means the preassociation of cHBIs with an ABS/NBS is necessary to enhance the kinetics of covalent bond formation on sIgEs via increased ITC effective molarity, thereby providing selectivity. We previously confirmed that the NBS pocket is highly conserved in antibodies by analyzing over 260 antibody crystal structures (29) and have identified a lysine within 2 nm of the NBS pocket lip on several antibody crystal structures (SI Appendix, Fig. S4). This lysine provided the potential reactive site on sIgE $F_{ab}$ for the irreversible conjugation of cHBIs, thus enabling selective and irreversible inhibition of allergic reactions (Fig. 1D).

We synthesized two versions of cHBIs displaying Ara h 2 epitope 5 (cHBI$^{Arah2/ep5}$) or Ara h 6 epitope 5 (cHBI$^{Arah6/ep5}$) using Fmoc solid-phase peptide synthesis (SI Appendix, Fig. S5). ITC on cHBI was stable at pH 7.4, but became reactive at pH 10.0, indicating that cHBIs will have negligible off-target conjugation at pH 7.4 (SI Appendix, Fig. S6). We also synthesized fluorescein-tagged cHBIs which were used to detect cHBI binding to sIgEs (Table 2). Note that a 1:1 combination of cHBI$^{Arah2/ep5}$, cHBI$^{Arah6/ep5}$ (which will be referred to as cHBI$^{Arah2, Arah6}$) was used for all following experiments unless otherwise noted.

### Table 1. Identification of the immunogenic epitopes from Ara h 2 and Ara h 6 by nanoallergen analysis

<table>
<thead>
<tr>
<th>Protein Epitope</th>
<th>Serum</th>
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<tbody>
<tr>
<td>Ara h 2</td>
<td></td>
</tr>
<tr>
<td>Ara h 2</td>
<td>1</td>
</tr>
<tr>
<td>Ara h 2</td>
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</tr>
<tr>
<td>Ara h 2</td>
<td>3</td>
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<tr>
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<tr>
<td>Ara h 6</td>
<td>5</td>
</tr>
<tr>
<td>Ara h 6</td>
<td>6</td>
</tr>
</tbody>
</table>

EC$_{50}$ < 1 pM

1 pM < EC$_{50}$ < 10 pM

10 pM < EC$_{50}$ < 100 pM

100 pM < EC$_{50}$ < 1 nM

1 nM < EC$_{50}$

No response
Table 2. Chemical structures of cHBI inhibitors used in this study

<table>
<thead>
<tr>
<th>Compound</th>
<th>Structure</th>
<th>m/z (Da)</th>
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<tbody>
<tr>
<td>cHBI_Ara h 2/ep5</td>
<td></td>
<td>2915.43</td>
</tr>
<tr>
<td>cHBI_Ara h 6/ep5</td>
<td></td>
<td>2268.16</td>
</tr>
<tr>
<td>FITC-cHBI_Ara h 2/ep5</td>
<td></td>
<td>3560.66</td>
</tr>
<tr>
<td>FITC-cHBI_Ara h 6/ep5</td>
<td></td>
<td>2913.39</td>
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</tbody>
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cHBI\textsuperscript{Arah2:Arah6} inhibited >80% of degranulation in 14 of 16 sera for at least one of the peanut-extract concentrations tested. Note that due to varying levels of sensitivity of the individual subjects, some patients had lower inhibition at higher allergen concentrations (Fig. 3A and SI Appendix, Figs. S12 and S13). In control experiments performed using either cHBI\textsuperscript{Arah2/epi5} or cHBI\textsuperscript{Arah2/epi5} alone, we observed <30% inhibition, demonstrating that effective inhibition required a combination of more than one immunodominant epitope (SI Appendix, Fig. S14A). Western blot analysis revealed that cHBI\textsuperscript{Arah2:Arah6} suppressed crude peanut extract-induced phosphorylation of intracellular proteins associated with degranulation response (SI Appendix, Fig. S14B). This inhibition was also long lasting as primed RBL-SX38 cells incubated with cHBI\textsuperscript{Arah2:Arah6} retained resistance to crude peanut extract-induced degranulation even when challenged 3 d after cHBI\textsuperscript{Arah2:Arah6} was washed away (SI Appendix, Fig. S15). Additionally, inhibition via cHBI\textsuperscript{Arah2:Arah6} was concentration and time dependent; more inhibition occurred at increased concentrations and incubation times (Fig. 3B). Taken together, these results demonstrate a combination of just two engineered epitope inhibitors, cHBI\textsuperscript{Arah2:Arah6}, deliver potent inhibition of immune cell and peanut allergen interactions.

**cHBIs Prevent Cellular Degranulation Induced by Crude Peanut Extract in a Basophil Activation Test.** Finally, to test the potential for a clinical evaluation of the inhibitors, we evaluated cHBIs by using a basophil activation test (BAT). The BAT is a physiologically relevant assay in which whole blood is taken from a patient, incubated briefly with an allergen, and the allergic response is monitored via flow cytometry by measuring cellular markers of activation and degranulation, including CD63, CD107a, or CD203c (30). Whole blood drawn in heparin vacutainer tubes was taken from a single patient with a history of severe allergic reactions to peanuts, incubated with cHBI\textsuperscript{Arah2:Arah6}, challenged with 100 ng/mL crude peanut extract, and analyzed with BAT. The percent of cells expressing CD63 above a threshold value [CD63\textsuperscript{+}, threshold shown via black line at 2 × 10\textsuperscript{5} mean fluorescence intensity (MFI)] increased from 1.4% to 70% when blood was challenged with crude peanut extract in the absence of cHBI\textsuperscript{Arah2:Arah6} inhibitors (Fig. 3C). When the blood was treated with cHBI\textsuperscript{Arah2:Arah6}, however, CD63\textsuperscript{+} cells decreased from 70% to 16–23% (Fig. 3C). In parallel, the percent maximal MFI of CD63 decreased when blood was treated with 0.4 and 2 μM of cHBI\textsuperscript{Arah2:Arah6} to 44% and 40% of maximal response respectively, Fig. 3D, \( P < 0.05 \). As a control, we took the epitope sequences in cHBI\textsuperscript{Arah2:Arah6}, synthesized cHBIs presenting Scrambled epitopes of Ara h 2 epitope 5 and Ara h 6 epitope 5, and mixed them at a 1:1 ratio (cHBI\textsuperscript{Scrambled}). cHBI\textsuperscript{Scrambled} did not reduce the levels of CD63 expression on basophils after crude peanut extract challenge, demonstrating that cHBIs have high selectivity (Fig. 3C and D). In addition to CD63, cHBI\textsuperscript{Arah2:Arah6} treatment reduced the levels of other markers of degranulation, such as CD107a and CD203c (SI Appendix, Fig. S16). Among these markers, CD63 expression in particular has been shown to be a predictive marker of physiological severity of an allergic response (31). To further validate these inhibitors, we performed the BAT assay on three additional allergic patients and two nonallergic control patients (Fig. 3E and SI Appendix, Fig. S17). There was a significant drop in CD63 and CD107a expression when patient samples were treated with 1 μM cHBI\textsuperscript{Arah2:Arah6}, while the nonallergic patients had no significant...
change in CD63 or CD107a expression. In particular, the significant reduction in the number of CD63+ cells in allergen-challenged cells when treated with cHBI Arah2:Arah6 demonstrate the clinical potential for cHBIs to prevent severe allergic reactions to peanuts. Combined, these results demonstrate that cHBIs have promising clinical potential to prevent allergic reactions to peanuts and improve patient outcomes; nevertheless, studies with larger patient populations are required to further validate these claims.

Discussion

Here, we described the design and evaluation of cHBIs, a selective sIgE inhibitor, to target and permanently inhibit the antigen binding site of allergen-specific IgE antibodies that recognize specific allergen epitope sequences. The most critical component of cHBI design required the identification of the immunodominant epitopes that are critical mediators during the initiation of degranulation responses. Using the nanoallergen platform, we identified these immunodominant peanut-allergen epitopes for...
a diverse set of serum samples in a functional cellular degranulation assay. The results showed that the immunodominant epitopes were commonly shared by all of the 16 subjects. Therefore, two cHBIs were synthesized to mimic these two epitopes and were evaluated.

One of the most significant results in our current study was that two cHBIs, targeting only two of the peanut epitopes, was sufficient to inhibit degranulation responses to crude peanut extracts with over 80% efficiency in 14 of 16 patient samples, although this effect was dependent upon the concentration of allergen challenge. Given that patients generate IgE to multiple proteins and multiple epitopes within each protein, the inhibition of responses by only two epitopes highlights that we still know very little about engagement of IgE by complex allergens. In our previous work we demonstrated that low-affinity epitopes can cooperate with high-affinity epitopes in IgE engagement and degranulation responses (32). Moreover, we previously observed that certain peanut epitopes trigger degranulation at significantly lower concentrations of nanallergens, but when used in combination with an epitope of lower potency, there is a synergistic enhancement of degranulation (12). This combinatorial aspect is important and contrasts with many studies that use “allergens” with high numbers of multimerized epitopes. This work suggests several things that are important in considering allergen reactivity. First, we suggest that there are public epitopes, reactivity that is shared across a broad portion of the peanut-allergic human population, in IgE recognition of peanut proteins. Second, we suggest that these public allergen epitopes are required for allergen responses and can be therapeutically targeted to inhibit responses. This finding potentially eliminates a major obstacle in designing allergen-specific inhibitors, suggesting that inhibitors would not need to be personalized and that common reagents could be used by a broader segment of the peanut-allergic population than initially perceived. This will be a critical concept to explore in other allergens to determine if our approach to inhibitor generation can be broadly applied to allergy treatment.

The second most significant result is increase in efficacy and the lack of complications or off-target effects due to incorporation of TFC in the cHBI design. From our experience with allergen model systems, we predicted that without a permanent covalent linkage, the epitope mimics would be unlikely to sufficiently inhibit IgE binding due to the short half-life of pegystatins and the eventual dissociation of the inhibitor from sIgE (33). One major concern is that cHBIs will indiscriminately conjugate proteins and trigger nonspecific inflammatory responses or side effects. While we did not observe any acute toxicity in our mouse experiments and have demonstrated that these inhibitors are very selective in vitro, further studies concerning the chronic toxicology of these inhibitors on immune responses is necessary. Based on the data from this study, we expect cHBIs to be nontoxic and the irreversible action of cHBI inhibitors could potentially yield long-lasting inhibition, since IgE bound to mast cells and basophils can remain in the body for months (34).

The cHBI design would be a significant improvement over current allergy therapeutics. Drugs that are currently available, such as epinephrine or antihistamines, only transiently attenuate such as epinephrine or antihistamines, only transiently attenuate the intensity of the degranulation response and allergic symptoms. This type of approach to immune inhibition is nonspecific and results in blocking immune components that could be important for antipathogen or antitumor immunity. Attempts at developing specific allergy inhibitors have not been successful due to the complexity of the immune response and the challenges arising from the multivalent nature of the interactions. This study describes the cHBI design, which provides a solution to these issues by using a lesser-known binding site on the FcεRI arm of antibodies with a multivalent design, and presents a successful example of a specific allergen inhibitor to be used on a clinically relevant protein-based allergen.

### Materials and Methods

#### Materials

- NovaPEG Rink amide resin, 5(6)-carboxy-fluorescein, 2-(1H-benzotriazole-1-yl)-1,1,3,3-tetramethylyuroniumhexafluorophosphat (HBTU), Fmoc-Lys(Dde)-OH, all Fmoc protected amino acids, and 10 KDa 0.5 mL entrapment filters were purchased from EMD Millipore.
- N,N-dimethylformamide (DMF) (>99.8%), dichloromethane (DCM) (>99.8%), N,N-diisopropylethylamine (DIEA), methanol, hydrazide, piperidine, trifluoro-acetic acid (TFA), triisopropylsilanes (TIS), trypsin, 2-naphthylethanesulfonyl chloride (NEDC), diisopropylazocarboxylate (DIAA), 2,4-dinitro-fluorobenzene (DNFB), acetonitrile, acetic acid, methanol, carbonate-bicarbonate buffer, Tween 20, indole-3-butycarboxylic acid (IBA), biotin, PBS, streptavidin conjugated to HRP (Step-HRP), defatted peanut meal (crude peanut extract), carboxyfluorescein (CF), and cytoschalasin D were purchased from Sigma-Aldrich.
- IBA-Boc is a Boc protected 4-methyl-4-phenyl-1,2,3- triazolinium betain salt, used as the cationic component of the Fmoc-protected amino acids.
- 10 kDa 0.5 mL ultrafiltration membranes were purchased from Amicon Ultra-15 Centrifugal Filter Units (EMD Millipore).
- 2% Fmoc-epitope lipid, 5% PEG 2000 were purchased from Invitrogen.
- 4-(dime-thylamino)pyridine (DMAP), succinic anhydride, carbon disulfide (CS2), dibenzyl-disulfonylazocarboxylate (DIAE), 2-(2-fluorobenzoyl)-1,3-indanedione (FIOB).
- Fluo-rescein isothiocyanate (FITC) was purchased from Vector Laboratories.
- Human serum was obtained from two different sources. Peanut-allergic patient serum (sera 1, 2, 3, 4) and healthy nonallergic control patient serum were purchased from Plasma Lab International. Sera 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, and 16 were taken from patients enrolled in an oral peanut immunotherapy (OIT) clinical trial administered at Massachusetts General Hospital under the direction of Wayne Shreffler (NCT01324401). Samples were obtained from patients before OIT and selected from patients having IgE levels for peanuts > 50 kU/L as determined by Immucap.
- Blood from human subjects was obtained by venipuncture from patients with a clinical history of severe allergic reactions to peanuts or from healthy volunteers. All volunteers consented and all studies were approved by either the Massachusetts General Hospital Institutional Review Board or the Indiana University Institutional Review Board.

#### Methods

**Nanallergen synthesis and characterization.** We utilized the epitope-epitope synthesis strategy as seen in previous publications (12). We synthesized epitope-lipid nanoparticles of all five Ara h 2 epitopes and five Ara h 6 epitopes using standard Fmoc solid-phase peptide synthesis (SPPS) chemistry on NovaPEG Rink amide resin. Residues were activated with HBTU and DIEA in DMF for 5 min and allowed to couple to resin for 30–45 min. Coupling efficiency was monitored using a Kaiser test. The Fmoc-protected residues were deprotected with a 3:1:1 mixture of pyridine, thioanisole, and 10% TFA in CH2Cl2 for 15 minutes. The purified product was characterized using a semipreparative Zorbax C18 column or Zorbax C3 column with either acetonitrile or 2-propanol gradients in the mobile phase. We monitored the column eluent with a diode array detector allowing a spectrum from 200 to 400 nm to be analyzed. The purified product was characterized using a Bruker Autoflex III Smartbeam matrix-assisted laser desorption ionization time-of-flight mass spectrometer. Liposomal nanallergens were prepared by preparing a liposome dispersion as previously described (35). Epitope lipids were then incorporated into liposomes using dry film hydration and extrusion. The liposomes were composed of 2% peptide lipid, 5% polyethylene glycol 2000 (mPEG2000)–DSPE, 5% cholesterol, and 93% DSPC. The lipid mixture was used for liposome encapsulation.
dissolved in chloroform, dried with nitrogen lyophilized for 30 min, rehydrated in PBS at 60 °C, and then extruded through a 100-nm polycarbonate filter (Avanti).

The liposomes were characterized by DLS analysis via a 90Plus nanoparticle size analyzer (Brookhaven Instruments Corp.), using 658 nm light observed at a fixed angle of 90° at 20 °C. All nanolipidic particles had a particle size between 115 and 125 nm, depending on the epitope size (Fig. 2C and SI Appendix).

chBI synthesis. All inhibitors (chBI and FITC-chBI) were synthesized using SPPS (23), with several modifications. First, the epitope was synthesized as a linear peptide using Fmoc-amino acids activated with a 3.6-fold excess of HBTU with 20-fold DIEA for 5 min before addition to the resin. Note that for FITC-chBI, before epitope synthesis, Fmoc-Lys-IvDdE-OH was added, then deprotected and Fmoc-Eg-OH was added. The Fmoc was deprotected and a HBTU-activated CF was added. Then the IvDdE group was deprotected with 2% hydrazine in DMF and epitope synthesis was performed.

Following epitope synthesis for all chBIs, a Fmoc-Eg-OH spacer was added, followed by conjugation of Fmoc-Lys(IvDdE)-OH. To the Fmoc-protected lysine, IBA-BOC was added. Following IBA-Boc addition, the IvDdE group of lysine was deprotected using 2% hydrazine in DMF in the same fashion. Fmoc-Eg-OH spacer was added to the ε-amine of this lysine.

After deprotection of the Fmoc group of the Eg spacer, ITC domain was added, just before cleavage from the resin. The deprotected amines were chemically modified into ITC moieties using a modified procedure from Munch et al. (36). Briefly, resins with deprotected primary amines were washed in anhydrous DMF three times. A 10-fold excess CS2 with a 20-fold excess of DMAP was added to DMF and allowed to react for 20 min at room temperature. The mixture was filtered and washed with anhydrous DMF. One milliliter of DMAP with a 20-fold excess of DIAE was added to resin and cooled to −20 °C in a −20 °C freezer. Then, a 2-fold excess of BocO and 0.2-fold of DMAP was added to the vessel and allowed to react for 20 min at −20 °C. The vessel was removed, allowed to warm to room temperature for 30 min, and then washed with DMF, DCM, and diethyl ether and allowed to dry in a vacuum chamber. See SI Appendix, Fig. 55 for further details.

To evaluate the chBI inhibitors, RBL-SX38 cells were similarly plated and primed with patient serum. However, before incubation with allergen, cells were treated with DABBS (15 min–24 h), and similarly challenged with allergens as described above. Note that concentrations of crude peanut extract used for inhibition assays shown in Fig. 3A varied, given the sensitization of the patient to crude peanut extract. A total of 1 ng/mL was used for serum 6 and sera 8–16, 100 ng/mL was used for sera 3 and 7, and the remaining sera 1, 2, and 4 were challenged with 1,000 ng/mL.

For DNP-BSA degranulation assays, a similar procedure was used with the following exceptions: RBL-2H3 cells were used instead of RBL-SX38 cells, and cells were incubated with a 25%/75% mixture of anti-DNP IgE and anti-cyclin A for 24 h instead of patient serum, and DNP-BSA was used as the allergen. Percent degranulation was calculated as previously described (32). Briefly, experimental signals were calculated as a percentage between positive control (100%) and negative control (0%). Degranulation maximum (Dmax) and degranulation areas were calculated using Origin 7 software, using Hill's curve fit for sigmoidal curves using percent degranulation values. Note that if nanoallergen did not induce significant degranulation above error (determined as SD of triplicate experiments, P < 0.05), it was deemed to have “no response.” If a significant signal was present but there was insufficient data to determine an EC50 value, the EC50 value was estimated to be greater than an EC95 value calculated assuming that the highest signal reached was Dmax.

ELISA assay. Binding interactions between chBI and antibodies were detected using a sandwich ELISA. Before the ELISA test, patient serum was incubated at various concentrations with FITC-chBIs at 37 °C for 5 h. Then, a high-binding 96-well ELISA plate was coated with 100 μL of 2 nM of anti-IgE antibody for 2 h in bicarbonate buffer (pH = 10). Plates were washed to remove unbound antibody. Wells were blocked with a blocking buffer (5% BSA, 0.2% Tween-20 in PBS) for 1 h. Inhibition was performed with various concentrations with FITC-chBIs at 37 °C for 5 h. A 1:1 mixture was used for all other samples. After blocking, 100 μL of patient serum was incubated with 100 μL of FITC-chBI (diluted 1:5,000 in blocking buffer) for 1 h. The plate was washed again and incubated with an anti-FITC mouse IgG-HRP conjugate (diluted 1:5,000 in blocking buffer) for 1 h. The plate was washed again and an Amplex Red Kit was used to quantify the ELISA signal using a SpectraMax M5 spectrophotometer according to the manufacturer’s instructions.

For the Ara h 2 competitive binding inhibition, a similar ELISA was performed with some modifications. Before the ELISA test, patient serum was incubated in various concentrations with unlabeled chBIs at 37 °C for 5 h. Then, a high-binding 96-well ELISA plate was coated with 100 μL of 2 nM of anti-IgE antibody for 2 h in bicarbonate buffer. Plates were washed to remove unbound antibody. Wells were blocked with the blocking buffer for 1 h, washed, and incubated with serum-chBI mixture 1:1:1 with the blocking buffer for 16 h. The plate was washed again and incubated with biotin-tagged HRP (diluted 1:5,000 in blocking buffer) for 1 h. The plate was washed again, incubated with streptavidin-HRP conjugate, washed, and an Amplex Red Kit was used to quantify the ELISA signal using a SpectraMax M5 spectrophotometer according to the manufacturer’s instructions.

Animal studies. C57BL/6 female mice (7- to 8-wk old) were obtained from Harlan Bioball Biosciences. Mice were maintained in pathogen-free conditions, and studies were approved by the Institutional Animal Care and Use Committee. Toxicology studies were carried out in a similar manner as previously published in the University of Indiana in Vivo Therapeutics Core (32).

Basophil activation test. The BAT assay protocol was used as previously described (38). Briefly, blood collected in vacutainer tubes containing heparin was processed within 24 h of collection. For basophil activation: 100 μL of whole blood was diluted 1:1 in serum-free RPMI containing chBIs for 4 h at 37 °C. Cells were activated with 100 ng/mL of crude peanut extract or media (blank) for 30 min at 37 °C. RBS were lysed using ALC lysis buffer and remaining cells were stained and fixed for flow cytometry. Basophils were selected as viability dye negative, SSC−, HLA-DR−, and CD123+. Each sample was analyzed in triplicate with a cutoff of 1,000 basophils per sample. Basophil activation was assessed by measuring fluorescence of CD63, CD107a, and CD203c. Percentage maximal CD63 MFI was found by calculating the percentage between the blank (negative control) and the no inhibitor (positive control) MFI for CD63.

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