Strategies for Mast Cell Inhibition in Food Allergy

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Mast cells are tissue resident allergic effector cells that drive IgE-mediated food allergies. There are several steps leading to mast cell activation in the context of allergic disease that can be targeted to prevent mast cell activation and degranulation. These include blocking IgE-FcεRI crosslinking and type 2 cytokine receptor activation; modulating cell-surface neural chemical receptors; stabilizing mast cell membranes to prevent co-localization of activating receptors; impeding intracellular signaling; and engaging cell surface inhibitory receptors. This review highlights several ITIM-containing inhibitory mast cell surface receptors that could serve as pharmaceutical targets to prevent mast cell activation and degranulation in the context of food allergy. When activated, these ITIM-containing inhibitory receptors recruit the phosphatases SHP-1, SHP-2, and/or SHIP to dephosphorylate the tyrosine kinases responsible for activation signals downstream of the IgE-FcεRI complex. We describe several members of the Ig and Ig-like inhibitory receptor and C-type lectin inhibitory receptor superfamilies. Fundamental studies exploring the behavior of these receptors within the context of experimental food allergy models are needed. A deeper understanding of how these receptors modulate mast cell-driven food allergic responses will shape future strategies to harness these inhibitory receptors to treat food allergy.

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Abbreviations: FcεRI, Fc Epsilon Receptor I, the high affinity IgE receptor; FDA, US Food and Drug Administration; Ig, immunoglobulin; ITAM, immunoreceptor tyrosine activation motifs; ITIM, immunoreceptor tyrosine-based inhibition motifs; KIR, Killer cell inhibitory receptors; LIR, leukocyte Ig-like Receptor; MAFA, mast cell function-associated antigen; MC, mast cell; IgE, immunoglobulin E; MC\textsubscript{T}, tryptase-producing mast cell; MC\textsubscript{C}, chymase-producing mast cell; MC\textsubscript{CT}, chymase- and tryptase-producing mast cell; NK, natural killer; OIT, oral immunotherapy; PCA, passive cutaneous anaphylaxis; PI3K, phosphatidylinositol 3-kinase; PECAM-1, Platelet Endothelial Cell Adhesion Molecule-1; PSA, passive systemic anaphylaxis; RBL, rat basophil leukemia; SCF, stem cell factor; SHP, Src homology 2 domains containing protein tyrosine phosphatase; SHIP, SH2-containing inositol 5’ phosphatase; Siglec, Sialic acid-binding immunoglobulin-like lectins; TNF, tumor necrosis factor.

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INTRODUCTION

Mast cells are one of the critical, primary effector cell types that drive IgE-mediated food allergies. These tissue resident cells contain cytoplasmic secretory granules packed with pre-formed mediators that include histamine, tumor necrosis factor (TNF)-alpha, serotonin, and multiple serine proteases including tryptase and chymase. Activated mast cells can release these granules into the surrounding tissue microenvironment in addition to synthesizing several pro-inflammatory mediators including leukotrienes, prostaglandins, chemokines, and cytokines [1]. Human mast cells (MCs) are commonly separated into two subsets, tryptase-producing mucosal MC (MC₅₆) and connective tissue MC that express chymase and tryptase (MC₇₇) [2]. There is increasing recognition of MCc which express chymase, but not tryptase [3], and play a role in tumor biology [4]. Murine MCs are typically divided into connective tissue MCs or mucosal MCs depending on their anatomic location and granule contents [3]. Transcriptional profiling techniques have revealed that mast cells have unique transcriptional profiles compared to other allergic effector cells, i.e. basophils, and that the striking heterogeneity across tissue mast cell populations is driven by their tissue microenvironment [1]. Despite this, all mast cells express the high affinity IgE receptor, FcεRI, and can be involved in the development of allergic disease [5].

During sensitization to a food antigen (or allergen), B cells produce food allergen-specific IgE that later binds FcεRI on both basophils and mast cells. After subsequent exposure to that particular food, allergenic food proteins crosslink the food allergen-specific IgE bound to the FcεRI, activating sensitized basophils and mast cells, resulting in the release of pre-formed and newly synthesized mediators that drive allergic inflammation and its associated symptoms [6]. Despite decades of research, a cure for food allergies remains elusive. However, treatments for food allergy, for instance, epicutaneous, sublingual, and oral immunotherapy (OIT), are under active investigation [7,8]. Investigational OIT targeting the top eight food allergens in the US [9] and a recent, US Food and Drug Administration (FDA)-approved OIT to treat peanut allergy [10] consistently promote robust desensitization to these foods [8]. In a subset of patients with egg and peanut allergy, OIT even leads to sustained unresponsiveness, or temporary resolution of food allergy, although the durability of this limited remission is unclear [8].

Moreover, immunotherapy still carries a risk of allergic reactions due to allergen exposure, and to date, FDA-approved peanut OIT only provides protection against accidental peanut exposure. It does not cure peanut, or any other food allergy. Thus, there remains a critical need for the development and deployment of therapies that treat food allergies in addition to and aside from peanut allergy. Ideally, these therapies should target multiple food allergies at once, since 30% of those with food allergy are allergic to more than one food [11]. Furthermore, simultaneous treatment of multiple food allergies improves the quality of life for the caregivers of allergic individuals [12]. FDA-approved biologics like omalizumab and dupilumab used to treat allergic diseases like asthma, atopic dermatitis, and nasal polyposis, show promise in the mitigation of food allergy because they target critical components of Type 2 allergic inflammation, including IgE (omalizumab) and IL-4/IL-13 receptor signaling (dupilumab) [6]. But while studies are ongoing to examine a role for these biologics in treating food allergy [11,13,14], there is increasing interest in developing therapies that target the final common pathway leading to symptom development and anaphylaxis in food allergy – MC activation, degranulation, and allergic inflammatory mediator release. There are several components leading to MC activation in the context of allergic disease that can be targeted to prevent MC activation and degranulation. These include blocking IgE-FcεRI cross-linking as well as cell-surface neural chemical receptor activation and type 2 cytokine receptors; stabilizing MC membranes to prevent co-localization of activating receptors; impeding intracellular signaling; and engaging cell surface inhibitory receptors. This review will focus on engaging inhibitory cell surface receptors as a strategy to prevent MC activation and degranulation in the context of food allergy. Since rodent MC lines and murine models of experimental food allergy are valuable tools to study potential therapies, we will review inhibitory receptors on mouse and rat as well as human MCs.

SIGNaling downstream of the high affinity IGE receptor, FcεRI

Both the human and rodent FcεRI are tetrameric immunoreceptor family members, containing alpha (α), beta (β), and 2 gamma (γ) subunits [15-17]. The alpha chain binds IgE and the beta and gamma chains have immunoreceptor tyrosine activation motifs (ITAMs) that enable activation of the receptor [17]. Though beta chains are always present in murine FcεRI, they may be absent in human FcεRI. When the beta subunits are present in human FcεRI, they serve as amplifiers of activation signals [17,18].

During conventional IgE-FcεRI downstream signaling, cross-linking of IgE bound to the FcεRI by antigen leads to the activation of the Src tyrosine kinase, Lyn [17,18]. Lyn phosphorylates ITAMs on beta and gamma chains of the FcεRI leading to the recruitment of the cytosolic signal transduction protein Syk to the ITAMs [17].
Syk is then phosphorylated by Lyn and via autophosphorylation, starting the downstream cascade of signaling molecules that increase intracellular calcium, activate transcription factors responsible for cytokine production, induce lipid mediator production, and cause mast cell degranulation (Figure 1) [17,18]. In addition to Lyn, downstream FcεRI signaling can be initiated by other Src kinases like Fyn, which then activates the phosphatidylinositol 3-kinase (PI3K) signaling pathway.

While Lyn functions, in part, as an initiating signal proximal to FcεRI coaggregation, it also has regulatory function in response to high or supra-optimal antigen stimulation [19,20]. In addition, Lyn-deficient bone marrow derived mast cells exhibit over-exuberant degranulation when activated, and Lyn-/− mice have increased serum IgE levels, increased histamine levels, increased mast cell numbers, increased expression of FcεRI, and increased occupancy of FcεRI by IgE [21]. These are all features that predispose to allergic inflammation in murine models, suggesting that Lyn signaling can negatively regulate FcεRI signaling in the context of allergic inflammation [21].

INHIBITORY RECEPTORS ON MAST CELLS: ADRENERGIC RECEPTORS

The classic means of inhibiting MC activation and degranulation hinges on engaging cell surface alpha and beta-adrenergic receptors which bind the catecholamine neurotransmitter epinephrine (adrenaline). For example, epinephrine bound to beta-2 adrenoreceptors blocks human lung MC activation in vitro [22]. This inhibitory effect can be attenuated in the presence of the mast cell growth and survival factor stem cell factor (SCF) [22] or if human lung MC are co-cultured with airway smooth muscle cells [23]. Genetic polymorphisms in the gene encoding the beta-2 adrenoreceptor can also impact the inhibitory activity of beta-2 adrenergic agonists [24]. Studies in rat peritoneal mast cells have shown that epinephrine engaging alpha-adrenergic receptors suppresses mast cell exocytosis, a critical component needed for mast cell degranulation, in a dose-dependent fashion [25]. This is the rationale behind using epinephrine autoinjectors to treat anaphylaxis and short and long acting beta-agonists in the management of allergic asthma and anaphylaxis involving bronchospasm [26]. Although epinephrine, in particular, can attenuate mast cell degranu-
ITIM-CONTAINING INHIBITORY IMMUNE RECEPTORS ON MAST CELLS

Catecholamine neurotransmitter receptors are not the only inhibitory receptors present on MCs. Constitutively expressed cell surface inhibitory immunoreceptors can also block MC activation and degranulation by negatively regulating FceRI signaling. Inhibitory receptors have immunoreceptor tyrosine-based inhibition motifs (ITIMs) that, when activated, recruit the phosphatases SHP-1, SHP-2, and/or SHIP to dephosphorylate the tyrosine kinases responsible for downstream activation signals (Figure 1) [20,27]. Inhibitory receptors can either be engaged alone or, in some cases, coaggregated with FceRI to block signaling downstream of FceRI and inhibit degranulation [27]. The direct impact of ITIM-containing inhibitory receptors on allergen-induced FceRI signaling make these molecules attractive targets for potential therapeutics for food allergy.

Inhibitory immunoreceptors found on human and mouse mast cells are frequently divided into two groups based on protein structure, the Ig and Ig-like receptor superfamily and the C-type lectin superfamily. Most of the inhibitory immune receptors belong to the Ig and Ig-like receptor family, including FcγRIIB, Allergin-1, gp49B1, leukocyte Ig-like receptor (LIR)-5, paired Ig-like receptor (PIR)-B, signal regulatory protein (SIRP)α, Sialic acid binding Ig-like lectins (Siglecs), the CD300 family, Platelet Endothelial Cell Adhesion Molecule-1 (PECAM-1), and killer inhibitory receptors (KIR). The C-type lectin superfamily includes mast cell function-associated antigen (MAFA) and CD72 (Figure 2 and [27-29]). Selected members of these inhibitory receptor superfamilies on both rodent and human MCs are summarized in Table 1 and described in detail below.

Ig Receptor FcγRIIB

FcγRIIB is a low affinity receptor for IgG studied extensively in the context of mast cell inhibition. FcγRIIB driven inhibition of MC activation occurs through the recruitment of SHIP to the FcγRIIB ITIM [30-34]. Crosslinking of FceRI with FcγRIIB leads to inhibition of MC activation in RBL-2H3 cells transfected with murine FcγRIIB and in murine MCs [35]. Multiple studies have shown that interactions between antigen-specific IgG antibodies and FcγRIIB can inhibit MC degranulation [36,37]. Fusion proteins and bispecific antibodies have been created to coaggregate FceRI and FcγRIIB for inhibition of MCs [16,38-40]. For instance, Zhu et al. created a bifunctional human fusion protein (GE2) targeted toward Fcγ and Fcε receptors, to examine effects of simultaneously targeting and juxtaposing FcγRIIB and FcεRI in human basophils and mouse MCs from transgenic mice engineered to express the human FcεRI [16]. Their chimeric protein complexed with both FcγRIIB and FcεRI,
Allergin-1. Allergy-inhibitory receptor (Allergin)-1 is expressed in both human and mouse mast cells [44]. Unlike mice, humans express 3 isoforms of Allergin-1: Allergin-1L, Allergin-1S1, and Allergin-1S2, with Allergin-1S1 being the main form expressed on human mast cells [44]. Co-ligation of FcεRI with Allergin-1 on primary human mast cells [44] or with transfected Allergin-1 expressed in a rat basophil leukemia mast cell line (RBL-2H3) [45] inhibits IgE-mediated degranulation. Mice lacking Allergin-1 have increased allergic responses in both PCA and passive systemic anaphylaxis (PSA) models [45]. Recent studies have shown the importance of Allergin-1, not only on MCs, but also basophils, in the inhibition of IgE-mediated food allergic responses—with specific influence on anaphylaxis [46]. In fact, Allergin-1 on bone marrow derived dendritic cells has been shown to suppress a mouse model of house dust mite-induced allergic airway inflammation through inhibition of Syk via SHP-1 [47]. Extending these studies to explore the role of Allergin-1 on lung resident mast cells in the house dust mite-induced airway inflammation model and in MCs within different murine food allergy models (peanut, egg, milk, etc.) will provide important data on Allergin-1 action within MCs that could shape future therapeutics targeting this receptor.

Gp49B1. Gp49B1 is a transmembrane mouse receptor within the gp49 family that contains two ITIMs [48]. Co-ligation of gp49B1 with FcεRI inhibited degranulation and time-dependent inhibition of antigen-specific IgE-driven histamine production with modified Syk signaling in human basophils and muted antigen-specific IgE-mediated passive cutaneous anaphylaxis (PCA) in human FcεRI transgenic mice [16]. They later showed that even when the fusion protein was deployed after human blood basophils and human lung mast cells had been sensitized, it could inhibit IgE-mediated responses in these cells [41]. Studies using murine bone marrow-derived MCs have shown that supra-optimal antigen stimulation of MCs, with associated supra-optimal FcεRI cross-linking, actually triggers antigen/IgE-dependent and independent co-localization of FcγRIIB and FcεRI, dampening MC activation [30]. In addition, a candidate vaccine engineered to induce protective immune responses against peanut allergy in a mouse model of peanut allergy was found to protect against peanut challenge in sensitized mice by generating peanut-allergen-specific IgG that engaged FcγRIIB [42]. Notably, expression of FcγRIIB is tissue specific; primary human skin MCs do not express FcγRIIB while gastrointestinal MCs do [43]. This may mean that therapeutic targeting of FcγRIIB could have differential impact on symptom development and resolution in food allergy, depending on the organ most significantly affected.

**Table 1. Summary of human and mouse mast cell ITIM-containing receptors.**

| Receptors  | Species | ITIM or ITIM-like domains | Phosphatases critical for inhibitory signaling | Ligands | References |
|------------|---------|---------------------------|-----------------------------------------------|---------|------------|
| FcγRIIB    | H, M    | 1                         | SHIP                                         | IgG     | [30-34]    |
| Allergin-1 | H, M    | 2                         | SHP-1                                         | Unknown | [47]       |
| Gp49B1     | H (LIR-5), M | 3, 2*                     | SHP-1                                         | Integrin avβ3 | [52] |
| PIR-B      | M       | 4*                        | SHP/SHIP independent                         | MHC-I   | [58]       |
| SIRPα      | H, M    | 4                         | SHP-2                                         | CD47    | [60,61]    |
| Siglec-2   | H, M    | 4                         | SHP-1                                         | Sialic acid | [64] |
| Siglec-3   | H, M    | 2, 1*                     | SHP-1                                         | Sialic acid | [63] |
| Siglec-8   | H, M (Siglec-F) | 2                     | SHP-1                                         | Sialic acid | [63] |
| CD300a     | H, M    | 4, 2*                     | SHP-1, SHIP                                   | Phosphatidylethanolamine and Phosphatidylinerine | [76,77] |
| CD300f     | H, M    | 3                         | SHP-1, SHP-2                                  | Ceramide and sphingomyelin | [83,84] |
| PECAM-1    | H, M    | 2                         | SHP-2                                         | PECAM-1, Integrin avβ3, CD38, and CD177 | [89,92] |
| KIR2DL4    | H       | 1                         | SHP-2                                         | HLA-G   | [33]       |
| CD72       | H, M    | 2                         | SHP-1, Cbl-b**                                | CD100/Semaphorin 4D | [96,97] |
| MAFA       | H       | 1                         | SHP-2, SHIP                                   | Unknown | [98]       |

*Indicative of number of ITIM or ITIM-like domains for mouse receptor if different from human. **Cbl-b is a ubiquitin ligase also important for inhibitory signaling of CD72. H=Human, M=Mouse

Ig-like Superfamily Inhibitory Receptors

**Allergin-1.** Allergy-inhibitory receptor (Allergin)-1 is expressed in both human and mouse mast cells [44]. Unlike mice, humans express 3 isoforms of Allergin-1: Allergin-1L, Allergin-1S1, and Allergin-1S2, with Allergin-1S1 being the main form expressed on human mast cells [44]. Co-ligation of FcεRI with Allergin-1 on primary human mast cells [44] or with transfected Allergin-1 expressed in a rat basophil leukemia mast cell line (RBL-2H3) [45] inhibits IgE-mediated degranulation. Mice lacking Allergin-1 have increased allergic responses in both PCA and passive systemic anaphylaxis (PSA) models [45]. Recent studies have shown the importance of Allergin-1, not only on MCs, but also basophils, in the inhibition of IgE-mediated food allergic responses—with specific influence on anaphylaxis [46]. In fact, Allergin-1 on bone marrow derived dendritic cells has been shown to suppress a mouse model of house dust mite-induced allergic airway inflammation through inhibition of Syk via SHP-1 [47]. Extending these studies to explore the role of Allergin-1 on lung resident mast cells in the house dust mite-induced airway inflammation model and in MCs within different murine food allergy models (peanut, egg, milk, etc.) will provide important data on Allergin-1 action within MCs that could shape future therapeutics targeting this receptor.

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tion of mouse MCs [49]. Castells et al. later identified αβ, as a ligand for gp49B1 and demonstrated that binding of αβ to gp49B1 also led to inhibition of antigen-induced IgE-mediated MC degranulation [50]. Gp49B1 also suppressed SCF-induced MC degranulation and associated tissue swelling. Intradermal injection of SCF into gp49B1-deficient mice resulted in four times more degranulating MCs and twice the amount of tissue swelling than in wildtype mice [51]. Gp49B1 recruits SHP-1 phosphatase, modulating downstream calcium release to induce inhibitory effects [52]. The human homologue of gp49B1 appears to be Leukocyte Ig-like receptor (LIR)-5 (also known as HM18, ILT3, LILRB4, CD85k, and LILB1-5) [53]. LIR-5 is expressed intracellularly in human MC, migrating to the cell surface after FcεRI cross-linking [54]. LIR-5 expression in dendritic cells marks them as tolerogenic in healthy individuals [55]. Its expression in myeloid-derived suppressor cells helps drive immune tolerance in cancer [55,56]. To date, however, the impact of MC-expressed LIR-5 in the severity of symptoms and development of anaphylaxis in food allergy remains unknown.

**PIR-B.** Paired immunoglobulin (Ig)-like receptor (PIR)-B is a glycoprotein with 4 ITIMs expressed on the cell surface of mouse mast cells, and closely related to gp49B1 and the human LIR family of receptors [57]. PIR-B is constitutively tyrosine phosphorylated and predominantly expressed on murine mast cells compared to its activating isoform, PIR-A [57,58]. Co-ligation of FcεRI and PIR-B leads to inhibition of IgE-mediated serotonin release [58], while co-ligation of c-kit and PIR-B inhibits calcium mobilization [57]. Interestingly, mouse bone marrow-derived MC PIR-B did not require SHP-1 for inhibitory function, setting it apart from PIR-B inhibitory receptors observed in other cell types [57,58]. Uehara et al. found that one of the 4 ITIMs responsible for PIR-B inhibitory function did not bind to the SHP-1, SHP-2, or SHIP phosphatases traditionally associated with the inhibitory effects of ITIMs. Thus, MC PIR-B has the potential to inhibit MC activation in a SHP/SHIP-independent fashion [58]. Similar inhibition through PIR-B and FcεRI co-ligation was also observed in RBL-2H3 mast cells with chimeric receptors, where the 3rd and 4th ITIMs were critical for inhibitory function of IgE-mediated calcium mobilization and degranulation [59].

**SIRPα.** Signal regulatory protein α (SIRPα), a Src homology-containing receptor phosphotyrosine phosphatase found on human and rodent MCs and other innate immune cells, possesses two conventional ITIMs and two ITIM-like domains [60]. Coaggregation of SIRPα with FcεRI on RBL-2H3 MCs leads to inhibition of IgE-mediated responses, through a reduction in calcium, decreased phosphorylation of FcεRI ITAMs, and loss of MAPK activation through recruitment of SHP-1 and SHP-2 [60]. Pan and colleagues found that murine MC activation reduced expression of SIRPα. Knocking down SIRPα in these cells in vitro enhanced IgE-FcεRI-mediated allergic (IL-4, IL-13) and inflammatory (IL-6, TNF-alpha) cytokine production. Intraderal ear injection of SIRPα-deficient MCs sensitized with dinitrophenyl (DN) specific IgE into MC-deficient mice followed by intravenous challenge with DNP heightened the allergic response in this PCA murine model compared to intradermal injection of sensitized SIRPα-wildtype MCs [61]. SIRPα negatively regulated FcεRI signaling by sequestering SHP-2 phosphatase from a PI3K regulatory subunit, resulting in impaired PI3K activation and blunted IgE-FcεRI-mediated responses [61].

**Siglecs.** Sialic acid-binding immunoglobulin-like lectins (Siglecs) are type 1 transmembrane proteins expressed on multiple immune cells that contain an N-terminal sialic acid binding domain [62,63]. There are 15 known Siglecs in human and nine in mice [62]. In humans, Siglecs-2, 3 (also known as CD22 and CD33, respectively) and 5-12 are considered inhibitory as they contain ITIMs. When Siglecs bind to sialic acid moieties on the ends of glycoproteins and glycolipids of the same cell or different cells, this alters the spatial location of Siglecs relative to transmembrane immune receptors, facilitating the immunomodulatory effects of these receptors [64]. CD22 (Siglec-2), CD33 (Siglec-3), and Siglecs-5 through 10 have been observed at varying levels of expression on human mast cells [64-68], with Siglec-7 and 8 most consistently associated with human mast cells [64].

When inhibitory Siglecs are brought in close proximity to the IgE-FcεRI signaling complex on MCs, they can also reduce MC activation. When Siglec-7 and Siglec-9 stably transfected into the MC RBL-2H3 cell line were crosslinked to the FcεRI, this inhibited serotonin release by RBL-2H3 cells [69]. Recently, Duan et al. highlighted the role of CD33 in inhibiting IgE-mediated activation of human and murine MCs. Using liposomes containing the ligand for CD33 and the model allergen trinitrophenyl (TNP), Duan and colleagues targeted TNP-specific IgE-FcεRI complexes on MCs, forcing proximity between the inhibitory CD33 and FcεRI to promote inhibition [68].

Unlike its companion Siglecs described above, Siglec-8 does not require co-ligation or other means of forced ligation with FcεRI to exert its inhibitory effects on MC activation [63]. Yokoi et al. showed that while Siglec-8 engagement on eosinophils resulted in eosinophil apoptosis, Siglec-8 engagement on human mast cells did not induce apoptosis. Rather, it blocked FcεRI-dependent histamine and prostaglandin release as well as calcium flux in MCs, and prevented anti-IgE-induced human bronchial ring contraction in a model of allergic asthma [63]. Experiments employing RBL-2H3 cells transfected...
with human Siglec-8 confirmed that Siglec-8-induced inhibition of calcium flux and MC degranulation depended on a functional membrane-proximal ITIM domain [63]. Kerr and colleagues used gene expression profiling and flow cytometry to demonstrate that AK002, a humanized monoclonal antibody specifically targeting Siglec-8, could decrease sputum eosinophil counts and inhibit FcεRI-activated lung mast cells [70]. This antibody also blocked MC signaling pathways and MC activation in a model of non-allergic airway inflammation [71]. Anti-Siglec-8 monoclonal antibody also effectively blocked MC activation during passive systemic anaphylaxis in a humanized mouse strain that produces mature Siglec-8-expressing MC populations [72].

Engaging and activating Siglecs expressed on immune cells involved in food allergy, aside from MCs, has shown promise as a potential means of treating type-2 allergic inflammation driven by food. In a mouse model of oral egg allergen-induced eosinophilic intestinal inflammation, Dong et al. showed that activating Siglec-F (the mouse equivalent of human Siglec-8) with an anti-Siglec-F antibody blunted intestinal eosinophilic inflammation, thus improving the integrity of the intestinal barrier [73]. It also reduced Th2 cytokine and IgE levels in mice with egg ovalbumin-induced eosinophilic inflammation, reducing their concomitant diarrhea and weight loss [73]. Siglec-engaging Tolerance-inducing Antigenic Lipo-somes (STALs) created with a high affinity inhibitory Siglec ligand juxtaposed to an antigen or immune-modulating therapeutic of choice can be used to induce tolerance in murine models of allergy and autoimmune diseases by enhancing the co-localization of the inhibitory Siglec with the specific antigen-receptor or drug-binding protein [74,75]. Orgel and colleagues deployed STALs that incorporated a ligand for B-cell expressed CD22 (Siglec-2) adjacent to Ara h2, the predominant allergic epitope in peanut allergy. In a murine model of peanut allergy, Orgel et al. showed that targeting both a Siglec and a food-antigen specific receptor prevented sensitization to Ara h2 and significantly blunted the severity of allergic reactions to whole peanut extract [74].

Targeting Siglecs on B cells and eosinophils has shown promise as a therapeutic approach in mouse models of food-induced allergic inflammation, while targeting Siglecs on MCs has proven effective in reducing MC activation in both allergic and non-allergic airway inflammation models. Given the variety of tools under active investigation, from liposomes to monoclonal antibodies, engineered to engage Siglecs, this particular class of ITIM-containing inhibitory receptors appears to have the greatest promise as a potential target in treating food allergy. Future studies should target Siglecs highly expressed on MCs, especially Siglec-8, in the context of in vivo food allergy models, taking advantage of both STALs technology and the Siglec-8-specific monoclonal antibody.

**CD300 family.** CD300 multigene family members are type 1 transmembrane proteins belonging to the Ig-superfamily inhibitory receptors. Genes encoding for these receptors are located on chromosome 17 in humans and chromosome 11 in mice. CD300a and CD300f possess long cytoplasmic tails containing ITIMs making them the only inhibitory receptors within the family [76]. As with other ITIM-containing inhibitory receptors, the suppressive functions of CD300a and CD300f are linked to the recruitment of SHP-1 and SHIP to their ITIMs. Both receptors are expressed on several immune cells, including allergic effector cells like MCs, basophils, and eosinophils [76]. Bachelet et al. showed that CD300a is constitutively expressed on human MCs; its expression could be decreased in vitro with exposure to eosinophil-derived major basic protein and eosinophil-derived neurotoxin [77]. Immune complex-mediated crosslinking of CD300a on human MCs inhibited IgE-mediated activation. Neutralization of the murine homolog of CD300a enhanced allergic inflammatory mediator release in a model of allergic peritonitis [77]. Co-ligation of CD300a and FceRI through simultaneous engagement of CD300a and IgE also inhibited MC activation and associated kinase phosphorylation in human and mouse MCs [78]. Since the identification of CD300a natural ligands, phosphatidylserine (PS) and phosphatidylethanolamine (PE), studies have examined the impact of these natural CD300a ligands on mast cell regulation. Both PS and PE are present on the outer portion of cell plasma membranes of infected, transformed, apoptotic, or activated cells [76]. Using mouse bone marrow derived mast cell cultures, Wang et al. showed that cis binding of PS exposed on the MC outer membrane during degranulation with CD300a is important for self-regulation of murine MC degranulation [79]. Moreover, mice deficient in CD300a took longer to recover from anaphylaxis in a PSA model [79].

CD300a shares 80% homology extracellularly with family member CD300c, an activating receptor. Both receptors also bind PS and PE [76]. Thus, future investigations should explore whether or not cross-reactive binding occurs between the two receptors. Some studies have linked upregulation of CD300a expression with clinical allergy. Sabato et al. showed that CD300a expression is upregulated after crosslinking of the FcεRI and is increased in basophils of birch pollen allergic individuals compared to healthy controls [80]. Interestingly, investigations of the ratios of CD300a compared to CD300c expressed on human basophils also highlight potential use of these receptors as biomarkers for allergy. Zenarruzabeitia et al. found that lower ratios of CD300a to CD300c correlated with increased hypersensitivity in cow’s milk allergy in children [81]. However, Larsen and colleagues...
Found no difference in CD300a expression in peripheral blood derived mast cells from peanut allergic individuals compared to controls. These authors did not report on CD300c expression [82]. Thus, whether similar expression patterns in the ratio of CD300a to CD300c exist in mast cells from food allergic patients remains unknown. Future studies should harness the existing murine models of food allergy to characterize CD300a and CD300c expression patterns on MCs across different tissue sites and across different food allergens.

CD300f has two ITIMs and one immunoreceptor tyrosine-based switch motif that can recruit SHP-1 to induce inhibitory effects in mouse [83] and human [84] MCs. The natural ligand for mouse CD300f is ceramide while both ceramide and sphingomyelin serve as ligands for human CD300f [85]. These lipid ligands can inhibit IgE-mediated activation of MCs upon binding to the receptor in the presence of FceRI engagement [83,84]. Mice deficient in CD300f show increased allergic responses in PSA and PCA models [83]. Moreover, when a mouse model of ovalbumin-induced IgE- and MC-dependent food allergy was established in CD300f-deficient mice and compared to wildtype mice, these CD300f-deficient mice developed higher levels of total and ovalbumin-specific IgE, and higher levels of mucosal mast cell protease-1, a marker of mucosal MC activation [86]. Using a different food allergy model in the CD300f-deficient mice, the authors were able to show enhanced clinical food-allergic responses, represented by larger drops in core body temperature upon allergen challenge [86]. When ceramide-CD300f binding was disrupted in wildtype mice used in the ovalbumin-food allergy model, the frequency of ovalbumin-induced allergic diarrhea increased [86]. In sum, CD300f appears to be critical in the regulation of MC-driven allergic responses in this mouse model. This suggests that pharmacologic engagement of CD300f could be used to blunt allergic symptoms in food allergy. Manipulation of the CD300f inhibitory pathway must proceed with caution, however, since CD300f has been shown to transmit activating signals via PI3K-binding motifs and the growth factor receptor-bound protein 2 (Grb2) [76,87].

**PECAM-1.** Platelet Endothelial Cell Adhesion Molecule-1 (PECAM-1) is a cell adhesion molecule linked to the transmigration of cultured bone marrow derived MCs across skin endothelial cells lining skin vessel walls [88]. Its natural ligands include PECAM-1 itself, integrin αβ3, CD38, and CD177 [89]. The ability of PECAM-1 to modulate cell motility depends on the interaction of its ITIM domain with SHP-2 [90]. The presence of this ITIM also explains its status as an inhibitory receptor. Polymorphisms in the gene encoding PECAM-1 have been associated with increased risk of asthma [91], although it is not clear whether MCs or other leukocyte populations are differentially activated. The absence of PECAM-1 in mice increased IgE-mediated allergic responses in PSA and PCA models. MCs from PECAM-1-deficient mice also showed increased MC mediator release in comparison to MCs from wild type mice after FcεRI activation [92]. However, since PECAM-1 is used to facilitate the trafficking of multiple leukocyte populations [93], it may prove a challenging target to inhibit MC activation in food allergy.

**KIRs.** Killer cell inhibitory receptors (KIR) for MHC Class I are type I transmembrane proteins expressed primarily by innate natural killer (NK) and adaptive T lymphocytes. However, when Blery and colleagues reconstituted KIR function in the non-lymphoid MC RBL-2H3 cell line through transfection, they demonstrated that KIR required co-ligation with the ITAM-dependent activating FcεRI receptor in order to exert its inhibitory effect. Moreover, it used an inhibitory pathway distinct from FcγRIIB, blocking not only extracellular calcium entry, but also extracellular calcium release from the endoplasmic reticulum [34]. Recently, others have shown endogenous expression of KIR2DL4 receptor in the LAD2 human MC line, on cultured human MCs derived from the peripheral blood of healthy individuals, and on human tissue MCs [33,94]. Ueshima et al. found that KIR2DL4 inhibits IgE-mediated responses in human mast cells through the recruitment of SHP-2 [33]. While the interaction of KIR2DL4 with its natural ligand human leukocyte antigen (HLA)-G has been explored in the context of anti-tumor immunity and pregnancy [95], a possible role for the HLA-G-KIR2DL4 interaction in mitigating MC activation in food allergy remains to be examined.

**C-type Lectins**

**CD72.** A 45 kilo Dalton type II transmembrane protein, CD72 (Lyb-2) is an ITIM-containing inhibitory receptor whose natural ligand is CD100 or Semaphorin 4D (Sema4D) [96]. Kataoka and colleagues showed that CD72 is expressed on human MCs generated from CD34+ peripheral blood cells of healthy human volunteers. The authors concurrently stimulated CD72 with CD100 or an agonistic anti-CD72 antibody and the canonical MC receptor KIT (whose signals are critical for MC growth and survival), with stem cell factor (SCF). This resulted in inhibition of KIT-triggered phosphorylation of Src family kinases and extracellular-regulated kinases. KIT-driven mast cell motility, proliferation, and chemokine generation were also significantly impaired [96]. Engaging CD72 on mouse bone marrow derived-MCs suppressed IgE-FcεRI-mediated degranulation in addition to cell surface KIT and FcεRI-expression [97]. By contrast, activation of CD72 on human mast cells did not impair IgE-FcεRI-driven MC degranulation, only KIT-mediated signaling [97]. This species difference suggest that even
if future murine food allergy models explore CD72 activation as a strategy to inhibit MC activation and improve symptoms, this may not be directly applicable to IgE-mediated food allergy in humans.

**MAFA.** Mast cell function-associated antigen (MAFA) is a type II membrane glycoprotein originally discovered on the rat mucosal-type mast cell line RBL-2H3 [98,99]. Like CD72, it has an extracellular C-type lectin domain that, when engaged, can inhibit FcεRI activation via Lyn-mediated phosphorylation of its ITIM domain with subsequent SHIP and SHP phosphatase recruitment to the plasma membrane to disperse the inhibitory signals [98]. Time-resolved phosphorescence anisotropy and fluorescence resonance energy transfer studies suggest that MAFA can associate with isolated or clustered FcεRIIs [100]. Interestingly, MAFA does not require co-ligation with FcεRI to prevent activation [98]. Nevertheless, MAFA-FcεRI co-clusters are more effective at inhibiting MC degranulation than MAFA clusters alone [101]. Murine (KLRG1) and human (MAFA-L) homologs of MAFA have been described. However, the murine homolog is actually expressed on NK cells, while the human MAFA-L is expressed on MCs, but is not exclusive to this immune cell type [98]. MAFA is a far more challenging inhibitory receptor target in the development of therapeutics for food allergy. The complete absence of MAFA homolog expression on murine MCs hampers the ability to study the utility of targeting this receptor in the context of a murine food allergy model. Moreover, human MAFA-L expression across different immune cell types could increase off-target effects of any pharmaceutical directed at this receptor.

**CONCLUSIONS AND OUTLOOK**

Inhibitory receptors play a role in the natural regulation of FcεRI activation on mast cells and represent attractive targets for the development of therapeutics used to treat pathologic, mast cell-driven, food allergic responses. Indeed, a recent clinical trial demonstrating the safety and efficacy of an anti-Siglec-8 antibody for eosinophilic gastritis and duodenitis, gastrointestinal conditions characterized by elevated mast cell numbers in addition to eosinophilia [102], suggest that engaging this particular ITIM-containing inhibitory receptor may hold the most promise in blocking mast cell activation that promotes food allergic symptoms. Reliable biomarkers for diagnosing and monitoring the evolution of a food allergy may arise from studying changes in the surface expression levels of various inhibitory receptors in the context of food allergy. Engaging inhibitory receptors to treat food allergy could eliminate the need for allergen specific therapy, providing protection for those with multiple allergies. Activating inhibitory receptors in conjunction with oral immunotherapy could enhance the efficacy of this treatment while reducing side effects. For many of these inhibitory receptors, fundamental studies exploring the behavior of these receptors within the context of experimental food allergy models are clearly needed. Clinical and translational studies that characterize shifts in inhibitory receptor expression over the course of allergen-specific immunotherapy could also inform our understanding of the role for these receptors in desensitization or the induction of sustained unresponsiveness in food-allergic individuals. Clarifying how these receptors modulate food allergic responses will shape the strategies used to target them (natural ligands vs. agonistic monoclonal antibodies vs. small molecule agonists). Pharmaceuticals that engage and activate mast cell inhibitory receptors may serve as a novel approach to food allergy management that brings us closer to a cure.

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