INTRODUCTION

Allergic diseases are a rising global health threat and economic burden.1–3 In classical type I hypersensitivity, Immunoglobulin E (IgE) is the key molecule in the development of allergic reactions towards allergens.4,5 Specific IgE reacting with allergens triggers the release of inflammatory mediators through allergen-mediated cross-linking of the high-affinity receptor, FcεRI on allergic effector cells such as mast cells and basophils.6–8

Another IgE receptor, CD23 (FceRII), has largely been overlooked as a potentially important molecule in the field of allergy research.9 This is possibly the case because CD23 is involved in a complex variety of different immunological processes.10 Besides its role as an IgE receptor, CD23 plays a role in the development and growth of normal and leukemic B cells.11,12 Furthermore, it acts as a C-type lectin facilitating antimicrobial immunity13–17 and can even be engaged by sialylated IgG to act as a Fcγ receptor.18–20 Apart from its form as a membrane receptor, CD23 can be cleaved into soluble fragments (sCD23) which was studied as a disease marker in allergy, rheumatoid arthritis, and leukemia.21–24 Furthermore, sCD23 can activate monocytes via CD11b and CD11c integrins.25–27

Here, we focus on CD23 and discuss several mechanisms related to IgE binding, as well as the impact of the IgE/antigen-binding on different immune cells expressing CD23. One recent paper has shown that free IgE preferentially binds to FcεRI whereas IgE-ICs are preferentially captured by CD23. Binding of IgE-ICs to CD23 on B cells can, on one hand, regulate serum IgE and prevent effector cell activation and on the other hand facilitate antigen presentation by delivering the antigen to dendritic cells. These data argue for a multifunctional role of CD23 for modulating IgE serum levels and immune responses.
2 | CD23: A LECTIN AS AN FC RECEPTOR FOR IgE

The CD23 molecule is a trimeric glycoprotein member of the calcium-dependent (C-type) lectin family with a 45 kDa subunit. However, even though CD23 is a C-type lectin and IgE is heavily glycosylated, the interaction between IgE and CD23 is independent of carbohydrates and takes place for CD23 in the lectin-like head domain in a non-lectin manner.28 An early study showed that non-glycosylated recombinant IgE fragment has a high binding activity to CD23 even higher than that of myeloma IgE.29 This is also true for enzymatic deglycosylation of human hybridoma or myeloma IgE which does not show decreased binding to CD23 (not published data). CD23 presents a short cytoplasmic N-terminus followed by a single transmembrane region and a long C-terminal extracellular domain.30 The extracellular part consists of three regions: 1) an alpha-helical coiled-coil stalk region which mediates the formation of trimers, 2) a lectin head that binds IgE, and 3) a modified RGD sequence that binds to α5β1 integrins.31 There are two major CD23 isoforms, CD23a and CD23b32 which only differ in their intracellular region only 21 or 22 amino acids long for CD23a and CD23b, respectively.

CD23 is expressed initially as a membrane-bound molecule but it may be cleaved from the cell surface by metalloproteinas such as ADAM-10 resulting in soluble CD23 fragments (sCD23) of different molecular weights (37, 33, 25 and 17 kDa).33,34 Structural studies have shown that IgE interacts in different manners with its two receptors, FcεRI and CD23.35 IgE binds to FcεRI with high affinity (K_D between 0.01 nM and 0.1 nM). This occurs through the Cε3 domain in an open conformation allowing the binding of the FcεRI to a binding pocket formed by two sub-sides 1 (Cε3A) and 2 (Cε3B).36,37 In previous work, Shade et al. demonstrated the importance of glycans to IgE biology and identified a key glycan in the Cε3 domain that was required for IgE folding and IgE binding to FcεRI.38 By investigating the glycan patterns of IgE, they demonstrated that sialylation of the Fc part of IgE is associated with allergic pathogenicity and might be important for regulating atopic disease.39 In contrast, in the case of CD23, the crystal structure of the complex shows the interaction of two CD23 heads binding to Cε3 and Cε4 domains of a single IgE molecule with different affinities. One binding site has an affinity of around 1 μM whereas the other one is weaker by one order of magnitude (K_D around 14.4 μM).40-42 This interaction is characterized by bringing two Cε3 domains together in a “closed” conformation incompatible with the binding to FcεRI.

The second major ligand of CD23 is the complement receptor, CD21. It was shown that the CD21 binding site on CD23 does not overlap with IgE binding sites and is of low affinity within micromolar range (KD ~ 8.7 x 10^-7 M).43,44 The interaction of CD23 to CD21 occurs in short repeats in CD21 and by using CD21 mutants carrying extracellular point mutations. An early study has shown that it involves both carbohydrate-dependent and independent interactions.45

3 | IgE TARGETING TO THE CD23 PATHWAY

The importance of CD23 as a target of IgE is not entirely clear. However, even though CD23 binds IgE with clearly lower affinity compared with FcεRI, it was shown that CD23 can oligomerize on the surface of B cells leading to enhanced IgE binding through an avidity effect (Figure 1A,B). The leucine zipper region in the stalk was proposed to play an important role in CD23 oligomerization.47 In turn, a more recent study has suggested a direct involvement of the CD23 stalk region in IgE binding potentially explaining why IgE binds well to B cells despite the relatively poor affinity binding to the previously described lectin domain binding site.48 A further mechanism that was reported to enhance IgE binding to CD23 is the presence of calcium, which induces structural changes in CD23.49 A less studied aspect that could be very important in regulating CD23 targeting is the valency of IgE. The binding of IgE in complex with an allergen (IgE-immune complex) was shown to impact the binding of IgE to CD23.50 Furthermore, we recently showed that IgE in complex with allergen is preferentially bound by CD23, whereas the binding to FcεRI is diminished by IgE complexation.51 However, those findings are rather new and require confirmation specifically in regards to different IgE/antigen systems. Furthermore, no direct structural evidence for such a relationship between IgE-immune complexes and FcεRI has yet been published. Moreover, the physiological relevance of IgE-allergen immune complexes (IgE-ICs) in healthy and allergic patients is not entirely clear; even though their existence has been described a long time ago.52,53 Similar to IgE-ICs, the well-documented presence of natural anti-IgE antibodies could also lead to multivalent IgE complexes that could potentially regulate CD23 versus FcεRI targeting.54-57

IgE binding to CD23 may also be enhanced by other receptors. CD21, which binds CD23, is an interesting co-receptor for IgE-ICs. Even though IgE itself does not fix complement, the inclusion of complement-fixing IgG could impact the immune complex binding to CD23. It was shown that IgE-ICs formed in allergic patients include IgE, IgG1, and IgG4.58 The binding of immune complexes to B cells via CD23 could therefore be regulated by the IgG subclass present in the complex. As IgG4 does not fix complement well, higher IgG4 content versus IgG1 could reduce CD23/CD21 interaction and thus B cell targeting of the immune complex. (Figure 1C). Surprisingly, IgG binding via FcγRI was not found to play a role in that study. Potentially, the co-ligation of FcγRs with CD23 could impact complex binding and subsequent cellular activation or inhibition of signaling via immunoreceptor tyrosine-based activation (ITAM) or inhibition (ITIM) motifs, depending on the involved FcγR.59 Overall, the role of IgG and complement factors in regulating IgE-IC binding to CD23 on B cells or other cell types requires more detailed investigations.

4 | CD23 AS A REGULATOR OF IgE LEVELS

A main function attributed to CD23 is the regulation of IgE synthesis. Both in vitro and in vivo studies have shown that CD23 plays a central
role in regulating IgE synthesis. However, the exact mechanism of IgE down-regulation is a matter of debate. It was shown quite some time ago that mice overexpressing CD23 display reduced IgE levels after primary immunization with antigen in alum\textsuperscript{60,61} while CD23\textsuperscript{−/−} mice show enhanced IgE production.\textsuperscript{62} Furthermore, anti-CD23 antibodies inhibit antigen-specific IgE responses in mice.\textsuperscript{63}

In human B cells, IgE synthesis can be inhibited in vitro by direct targeting of CD23.\textsuperscript{65} This supports a model of either positive or negative feedback mechanism depending on the concentration of IgE and cleavage of membrane CD23.\textsuperscript{64} Thus, high levels of IgE or antibodies against the lectin head of CD23 stabilize membrane CD23 preventing its proteolytic cleavage and thereby inhibit IgE synthesis. In turn, the cleavage of CD23 by allergens has been a proposed mechanism of enhanced IgE responses.\textsuperscript{65} Allergen-cleaved CD23 would lose the ability to suppress IgE synthesis and hence lead to elevated IgE levels. CD23 binding by antibodies recognizing the stalk region of CD23 or metalloproteinases such as ADAM10 are additional ways in which CD23 cleavage and production of soluble CD23 can occur.

It has been proposed that CD23 cleavage not only prevents negative regulation, but may even enhance IgE synthesis by acting on other cells as soluble CD23. However, the mechanisms by which sCD23 enhances IgE synthesis are unclear. Potentially, released soluble CD23 may up-regulate IgE synthesis by cross-linking membrane IgE and CD21. The activity of the soluble fragments depends on their oligomeric state namely soluble CD23 monomers inhibit whereas oligomers stimulate IgE synthesis.\textsuperscript{66} The fact that IgE and CD21 have distinct binding sites and bind CD23, simultaneously supports this hypothesis.\textsuperscript{64} In contrast, the co-ligation of membrane IgE and membrane-bound CD23 via allergen-IgE complexes has been suggested as a negative feedback mechanism of IgE synthesis but more experiments need to be performed to confirm this hypothesis.

A recent paper has shown that CD23 as well can negatively regulate BCR activation on B cells by promoting B cell contraction. This explains the down-regulation of CD23 on memory B cells that mount a higher response of memory B cells to antigenic stimulation.\textsuperscript{75} In contrast, up-regulation of CD23 on switched memory B cells correlates with antigen-specific IgE levels and may be involved in some pathologies such as allergic rhinitis.\textsuperscript{76}

A further mechanism by which CD23 may regulate IgE levels is by acting as a direct decoy receptor for FcεRI. It was shown in mice, that B cells regulate serum IgE levels directly by absorbing free IgE molecules, thus preventing FcεRI loading and allergic sensitization.\textsuperscript{51,67,68} This more novel model of IgE regulation fits well with the generally higher IgE levels in CD23 deficient mice. CD23 cleavage could then be a mechanism to suppress this serum clearance and thereby enhance IgE levels.

5 | CD23 IN THE ACTIVATION OF B CELLS AND MONOCYTES

Many functional investigations on CD23 have demonstrated mechanisms triggered by CD23 cross-linking. The role of CD23 in monocyte-related cells is generally tricky to assess, as they can also express FcεRI. For example, human monocytes express FcεRI, whereas IL-4 stimulation up-regulates CD23 on those cells.\textsuperscript{69} Therefore, anti-CD23 antibodies were often used for specific CD23 cross-linking. It has been known for quite some time that CD23 cross-linking leads to internalization. The mechanism of uptake is different for the two CD23 isoforms, CD23a facilitates endocytosis, and CD23b phagocytosis.\textsuperscript{32} The differential expression of CD23a and CD23b in B cells and monocyte-related cells, respectively, has led to several interesting comparative studies showing differential signaling. Specifically, signaling via Fyn and Syk and Akt pathways resulting in IFN-γ production was only reported for CD23 cross-linking in B cells whereas cells of the monocytic lineage have been described to signal via ikB and produce inflammatory chemokines and cytokines such as TNF, IL-1β, IL-1ra, IL-10, IL-8, MCP-1, and MIP-1α.\textsuperscript{69-74}

In addition to the differential signaling, the processing of IgE and IgE-ICs also depends on the cell type. In monocyte-derived cells or dendritic cells only expressing CD23b, IgE, and allergen are targeted to a degradative pathway after CD23 cross-linking. In contrast, human primary B cells expressing CD23a in addition to CD23b, protect IgE and allergen from degradation and recycle IgE-ICs via CD23 allowing transfer to other immune cells\textsuperscript{77,78} (Figure 2). These findings

FIGURE 1  Mechanisms that regulate targeting of IgE to CD23. (A) Free IgE binds only weakly CD23 monomer whereas it binds much more strongly to oligomerized CD23 (B) IgE-antigen immune complexes (IgE-ICs) bind to CD23 much stronger than free IgE (C) Binding of antigen-specific IgG to IgE-ICs could enable complement fixation via CD21 and lead to co-aggregation of CD23 and CD21.
are consistent with studies in mice showing that circulating murine B cells transport IgE-ICs to the spleen.\textsuperscript{79–81}

Those findings in B cells fit well to results showing that CD23a expressing human intestinal epithelial cells\textsuperscript{82,83} as well as mouse intestinal epithelial cells can shuttle IgE and IgE-immune complexes through the epithelial layer by transcytosis.\textsuperscript{84} Like in human B cells, food allergens were also shown to be protected from degradation during epithelial transcytosis.\textsuperscript{85} Interestingly, intestinal epithelial cells (IEC) were also shown to release CCL20 in response to CD23 cross-linking, suggesting that CD23 may act as a critical receptor in initiating an allergic response by the release of chemokines capable of recruiting cells of the innate and adaptive immune system.\textsuperscript{86} Furthermore, CD23-dependent transcytosis of IgE-immune complexes was described for human airway epithelial cells (AEC), however, in contrast to B cells and IEC, CD23b was the reported isoform involved in AEC.\textsuperscript{83}
6 | CD23 MEDIATED IMMUNE RESPONSE

The consequence of CD23 mediated IgE-immune complex processing or trafficking is still not understood in detail. However, it is thought that IgE modulates immune responses to an antigen via CD23, as was shown in mice for antibody and T cell responses.86,87 The mechanism of antigen presentation mediated by CD23 has been referred to as IgE-facilitated antigen presentation (FAP) (Figure 3). As B cells are antigen-presenting cells expressing MHC class II, B cells could potentially also degrade antigen and display peptides on MHC class II for antigen presentation. This was indeed shown using EBV-transformed human B cells which directly present IgE-immune complexes to T cells.48,88-91 However, as previously mentioned, in primary B cells, IgE-immune complexes are protected from degradation. This difference in processing between normal B cells and EBV-transformed cells requires more detailed investigations, to better understand the mechanism of immunomodulation. Although primary human B cells fail to directly induce T cell proliferation, they can transfer the IgE-immune complexes to human dendritic cells to induce T cell proliferation.77 Fittingly, in mice, IgE-mediated antigen presentation was, though initiated by B cells, ultimately dependent on dendritic cells.80,92 The mechanism by which antigen could be transferred from B cells to other cell types is not entirely clear. A potential role in CD23-induced IgE or antigen shipping between immune cells could be attributed to exosomes. It was shown that B cell-derived exosomes can play a role in presenting allergen peptides to activate T cells92,93 (Figure 3). Independently, it has been described that the CD23 sheddase ADAM10 can mediate the sorting of CD23 into B cell-derived exosomes.94,95 The concept of exosome transfer between B cells and dendritic cells has also been put forth in mice.96,97 The consequence of CD23-mediated T cell proliferation and whether it is pro- or anti-inflammatory in the allergic context has not been resolved yet, and evidence is generally conflicting. In mouse models of allergic asthma, it was both postulated that CD23 could positively98 as well as negatively regulate allergic airway inflammation.99,100

It has also been postulated that CD23-mediated FAP can lead to IgG responses to unrelated allergens in a possible scenario of epitope spreading.8,101 Thus, CD23-expressing B cells will behave as presenting cells binding to antigens, regardless of the cell’s specificity, just as is the case with dendritic cells which can bind to different unrelated antigens via Fcγ receptors and induce antibody response to unrelated allergens. By this, CD23-FAP might explain the development of allergen polysensitivity to multiple allergens. CD23-mediated FAP is indeed as efficient as Fcγ receptors to induce antigen presentation even higher than BCR on the surface of B cells.102 This mechanism might play a role in IgE autoreactivity where low cross-reactive IgE autoantibodies can develop via FAP into high-affinity IgE autoantibodies.103 Even though this mechanism has been postulated as a potential cause of different IgE-mediated auto-immune diseases such as atopic dermatitis where the presence of IgE auto-antibodies has been associated with allergen sensitization it still requires a lot of further evidence.104

7 | CD23 IN CURRENT ALLERGY THERAPY APPROACHES

As long as the biology of CD23 is not completely understood, the potential use for CD23 as a therapeutic target is limited. However, several recent studies have begun to shed light on how current allergy therapies affect CD23.

The only disease-modifying therapy for allergies is allergen-specific immunotherapy (AIT).105,106 Multiple injections of increasing allergen doses induce the generation of tolerance via regulatory T cells and the induction of protective IgG4 antibodies.107,108 The role of CD23 in the generation of such IgG responses is unclear. However, the tolerogenic IgG induced by AIT was shown to inhibit IgE binding to CD23 and hence antigen presentation by EBV-transformed B cells.109-111

A different approach to treat allergic diseases is by anti-IgE therapy.112 Omalizumab, a monoclonal anti-IgE antibody is used for severe allergic asthma and chronic spontaneous urticaria.113,114 Mechanistically, Omalizumab inhibits both FcεRI: IgE and CD23: IgE interactions.115 A more recent anti-IgE antibody, Ligelizumab, was more efficacious in the treatment of allergic asthma.116 Functionally, Ligelizumab displayed reduced IgE:CD23 inhibition compared to Omalizumab but enhanced inhibition of IgE: FcεRI binding.117 A different anti-IgE antibody referred to as MEDI4212 is mimicking CD23 binding to IgE and was shown to inhibit allergic responses in mice and inhibit the FcεRI pathway.68 A further interesting anti-IgE termed (BD6), an anti-IgE Fab bound to IgE-Fc through a mixed protein-carbohydrate epitope, was shown to inhibit FcεRI while retaining CD23 binding.118,119 The monoclonal anti-CD23 antibody Lumiliximab, which specifically targets CD23 was shown to inhibit allergen-induced response in allergen-presenting cells and reduced Th2 response.120 However, anti-CD23 never led to particularly significant clinical outcomes in patients with asthma suggesting that blocking CD23 does not reduce allergic symptoms. Hence, studying the type of immune response elicited by CD23 is essential to understanding its role in allergy and immunotherapy as it could very well be of benefit to target IgE towards the CD23 pathway instead of blocking this interaction.

8 | CONCLUSION

The general goal of disease-modifying allergy immunotherapy is to reduce IgE responses, while enhancing IgG and regulatory T cell responses. While evidence from experimental disease models as well as allergic patient studies on CD23 is still lacking, evidence shows that i) CD23 can absorb and clear IgE from the serum in a non-inflammatory fashion, ii) CD23 reduces the synthesis of IgE from B cells, iii) CD23 regulates antigen-specific IgG and T cell responses. Together, those factors lead us to believe that CD23 deserves a closer look as a therapeutic target in allergies.
CONFLICT OF INTEREST
The authors have no conflicts of interest to disclose.

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