Development of an antibody that neutralizes soluble IgE and eliminates IgE expressing B cells

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Immunoglobulin E (IgE) plays a key role in allergic asthma and is a clinically validated target for monoclonal antibodies. Therapeutic anti-IgE antibodies block the interaction between IgE and the Fc epsilon (F\textsubscript{\v{e}}) receptor, which eliminates or minimizes the allergic phenotype but does not typically curtail the ongoing production of IgE by B cells. We generated high-affinity anti-IgE antibodies (MEDI4212) that have the potential to both neutralize soluble IgE and eliminate IgE-expressing B-cells through antibody-dependent cell-mediated cytotoxicity. MEDI4212 variants were generated that contain mutations in the Fc region of the antibody or alterations in fucosylation in order to enhance the antibody’s affinity for Fc\textsubscript{\v{e}}RIIIa. All MEDI4212 variants bound to human IgE with affinities comparable to the wild-type (WT) antibody. Each variant was shown to inhibit the interaction between IgE and Fc\textsubscript{\v{e}}RI, which translated into potent inhibition of Fc\textsubscript{\v{e}}RI-mediated function responses. Importantly, all variants bound similarly to IgE at the surface of membrane IgE expressing cells. However, MEDI4212 variants demonstrated enhanced affinity for Fc\textsubscript{\v{e}}RIIIa including the polymorphic variants at position 158. The improvement in Fc\textsubscript{\v{e}}RIIIa binding led to increased effector function in cell based assays using both engineered cell lines and class switched human IgE B cells. Through its superior suppression of IgE, we anticipate that effector function enhanced MEDI4212 may be able to neutralize high levels of soluble IgE and provide increased long-term benefit by eliminating the IgE expressing B cells before they differentiate and become IgE secreting plasma cells.

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INTRODUCTION

Immunoglobulin E (IgE) is the least abundant immunoglobulin isotype present in blood (0.05% of Ig concentration),\textsuperscript{1} and is capable of activating potent inflammatory reactions. Asthma is a chronic inflammatory disease of the airways characterized by recurring symptoms, airways hyperresponsiveness and variable, reversible airway obstruction and bronchospasm. Asthma is thought to be caused by a combination of genetic and environmental factors. A number of different asthma phenotypes exist, of which allergic asthma, mediated by endogenous allergen-specific IgE antibodies, accounts for at least 50% of incidence rates.\textsuperscript{2} Most patients with mild-to-moderate disease are treated as symptoms arise with inhaled corticosteroids and long-acting beta\textsubscript{2} agonists combined with short-acting beta\textsubscript{2} agonists. Chronic severe asthma occurs in approximately 5% of the asthmatic population. These patients may continue to experience persistent symptoms, airflow obstruction or frequent exacerbations despite aggressive treatment, including oral corticosteroids.\textsuperscript{3} These conditions negatively affect quality of life and place undue burden on patients and healthcare providers due to adverse effects from regular systemic steroid use. There is a significant unmet medical need for patients with poorly controlled asthma.

The mechanism by which IgE elicits type I hypersensitivity in asthma is well understood. Upon release of IgE from plasma cells, IgE binds \textit{via} its Fc domain to the high- and low-affinity IgE receptors (F\textsubscript{\v{e}}RI and F\textsubscript{\v{e}}RII) present on mast cells and basophils. Crosslinking of this receptor-bound IgE by an allergen triggers cell activation and degranulation, resulting in the release of histamine and other mediators of the allergic response.
The role of the low-affinity Fcγ receptor, CD23, is complex, but consequences of CD23 interaction with IgE include regulation of IgE synthesis, allergen presentation, allergen transport and cell-mediated effector functions. The beneficial effects of targeting the IgE pathway for type I hypersensitivity (allergic) responses is well documented and, in asthma, a clinically validated target. Omalizumab binds to IgE and blocks the interaction between IgE and FcγRII and FcγRII. This block results in a rapid neutralization of free IgE and gradual down-regulation of FcγRI on basophils, dendritic cells and mast cells. Although IgE is produced and secreted by plasma cells, IgE memory B cells are rare. However, B cell-associated membrane IgE has been described by a number of groups and antibodies targeting this region are currently in clinical trials. Membrane IgE-specific monoclonal antibodies are designed to eliminate IgE-expressing B cells before they become IgE-secreting plasma cells and, overtime, curtail the amount of total free IgE. Because these antibodies target a region that is specific and proximal to the membrane, these antibodies will not bind and neutralize soluble, circulating IgE or inhibit the interaction between IgE and FcγRII and FcγRI. Hence, targeting membrane IgE is unlikely to provide an initial benefit to the patients due to the sustained existence of soluble IgE and long-lived plasma cells that secrete IgE.

The humoral component of the immune system is responsible for antibody-mediated response to pathogens and toxins. This response includes the ability to engage and recruit effector cells. There are multiple antibody classes and isotypes in the human immune system, each with a palette of effector functions, presumably tailored to the nature of the invading pathogen or antigen. Much of our understanding of effector functions mediated by monoclonal antibodies comes from in vitro analyses of antibody-mediated killing. Most antibody-dependent cell-mediated cytotoxicity (ADCC) is performed by natural killer (NK) cells that express FcγRIIIa as the receptor that interacts with the Fc region of antibodies. Although a variety of techniques have been used to engage effector function and enhance ADCC by monoclonal antibodies, most rely on an improvement in the affinity of the Fc region of the antibody for FcγRIIIa. Certain mutations in the Fc region or afucosylation have been shown to enhance the Fc: FcγRIIIa binding affinity. Whether omalizumab can target membrane-bound IgE B cells and decrease the generation of new IgE-secreting plasma cells in patients is not well understood.

We have developed an antibody approach that may have the capacity to both neutralize soluble IgE and eliminate IgE-expressing memory B cells through an enhanced ADCC mechanism. This approach should have the combined clinical benefits of rapidly neutralizing soluble IgE and eliminating IgE B cells, which, overtime, should reduce the pool of IgE-secreting plasma cells.

**MATERIALS AND METHODS**

**Recombinant protein and antibody generation**

MEDI4212 wild-type (WT) monoclonal antibody (mAb) was expressed in Chinese Hamster Ovary (CHO) cells and purified as previously described. MEDI4212 aFuc was produced in FUT8-deficient CHO cells. Antibodies were purified by Protein A affinity chromatography. MEDI4212 2M2 (S239D/I332E) and MEDI4212 3M (S239D/A330L/I332E) variants were generated via site-directed mutagenesis using the parental MEDI4212 gene. These constructs were transiently expressed in CHO-CEP6 cells using lipofectimine LTX and CD CHO medium (Life Technologies, Carlsbad, CA, USA). The culture medium was collected 10 days after transfection. MEDI4212 2M2 and 3M were purified by protein A affinity chromatography. Soluble aggregate content was determined by analytical size-exclusion chromatography and removed by preparative size-exclusion chromatography. IgE was purified from U266 cells. Anti-migis antibody (an antibody specific to the membrane region of IgE) was generated internally as described previously.

**RBL-2H3 generation and calcium signaling**

FcγRI-expressing RBL-2H3 cell were generated as described previously. In short, human FcγRI was cloned from human peripheral blood lymphocytes into pcDNA3.1 vector and transfected, using a standard electroporation method, into RBL-2H3 cells. Transfected cell were cloned by limiting dilution and analyzed for surface FcγRI expression. The resulting cells were seeded at 5 × 10⁴/100 μl/well into 96-well, black-walled, flat-bottomed tissue culture plates (Costar). After 24 h, medium was replaced with test antibodies followed by addition of IgE to a final concentration of 25 ng/ml. Following a 4 h incubation at 37 °C, antibody/IgE mixture was aspirated, leaving the cell monolayer intact, and replaced with 100 μl/well of FLUO-4AM loading buffer (Dulbecco’s modified Eagle’s medium with 0.1% FBS, 20 mM HEPES, 2.5 mM probenecid and 2 μg/ml FLUO-4AM (Invitrogen, Life Technologies, Carlsbad, CA, USA)) for 1–2 h at 37 °C. Cells were washed with phosphate-buffered saline (PBS) and placed in 100 μl/well of Fluorometric Imaging Plate Reader (FLIPR) buffer (125 mM NaCl, 5 mM KC1, 1 mM MgCl₂, 1.5 mM CaCl₂, 30 mM HEPES, 2.5 mM probenecid, 5 mM glucose, 0.01% v/v fetal calf serum) for 5–45 min at 37 °C. To measure calcium mobilization following addition of crosslinking anti-IgE, the FLIPR (Molecular Devices, Sunnyvale, CA, USA) was calibrated for suitable exposure according to the manufacturer’s instructions. Anti-IgE (BioSource, Life Technologies, Carlsbad, CA, USA), diluted in FLIPR buffer, was added to the assay plates to a final concentration of 2.3 μg/ml. Fluorescence of the FLUO-4AM dye was recorded over 2 min and the peak response exported for analysis using GraphPad Prism software. For mAb crosslinking experiments, RBL-2H3 cells were cultured in the presence of 1 μg/ml IgE for 4 h before loading with FLUO4-AM and crosslinking with anti-IgEs as above.

**LAD2 β-hexosaminidase assay**

LAD2β-hexosaminidase assays were performed as described previously. Briefly, LAD2 cells were maintained in serum-free medium (StemPro-34; Life Technologies) supplemented with StemPro-34 nutrient supplement, 2 mM L-glutamine and
100 ng/ml recombinant human stem cell factor (R&D Systems, Minneapolis, MN, USA). Cells were seeded at a density of 2.5×10^4 cells/well and incubated in a 96-well polypolyethylene plate with test antibodies for 30 min at 37 °C before addition of IgE to a final concentration of 0.15 nM. Following a 4-h incubation at 37 °C, cells were washed with buffer to remove excess IgE, and IgE bound to FcεRI on the LAD2 cells was crosslinked with anti-IgE (600 μg/ml goat-261 anti-human IgE; Sigma-Aldrich, St. Louis, MO, USA) for 30 min at 37 °C. The incubation was stopped by centrifugation at 4 °C, and the cell supernatants analyzed for β-hexosaminidase content.

Membrane-associated IgE-expressing cell lines
IgE-expressing cell lines were generated internally and described previously. Briefly, nucleoporation was used for generating transfected cell lines expressing membrane IgE (mIgE) on the surface of 293T or NS0 cells. Cells were cotransfected with: (i) a linearized bicistronic mammalian expression vector coding for the mIgE heavy chain and a κ-light chain; the variable domains of the mIgE were derived from an antibody that binds the F-protein of respiratory syncytial virus; and (ii) a linearized bicistronic mammalian expression vector coding for CD79a and CD79b. Those clones with constant expression of membrane immunoglobulins to form the B-cell receptor complex were expanded and tested for expression of mIgE, CD79a and CD79b associate with membrane immunoglobulins by three-color fluorescence-activated cell sorting (FACS) into a single cell per well. Those clones with constant expression of mIgE, CD79a and CD79b were subcloned by limited dilution cloning at 0.2 cells/well to ensure monoclonality.

FACS labeling
Cell surface labeling and FACS analysis were performed as described previously. Briefly, 293T IgE-expressing cells were grown in Dulbecco’s modified Eagle’s medium with high glucose, L-glutamine, sodium pyruvate, 10% FBS, 1% genetin and 0.8% hygromycin. Medium was removed and the cells were washed with PBS. TrypLE Express was added to remove cells from the plates. Serum-containing medium was added to neutralize trypsin, and cells were washed with 10 ml PBS and resuspended at 2×10^6 cells/ml in FACS block (PBS with 3% bovine serum albumin and 0.1% sodium azide, pH 7.2). Cells were incubated in FACS block for 30 min on ice. All primary antibodies were used at 10 μg/ml and incubated on ice for 1 h. Cells were washed three times by centrifugation at 2000 r.p.m. for 3 min and the cell pellet was resuspended in 200 μl of FACS buffer. A 100 μl volume of Alexa Fluor 647-AffiniPure goat-anti-human IgG Fc fragment-specific antibody diluted at 1:400 into FACS block was added to the cells, and the cells were incubated on ice for 1 h. The primary IgG control was incubated in IgG1 control antibody and the secondary control received secondary antibody only.

FcyRIIIa enzyme-linked immunosorbent assay (ELISA) and Biacore assays
An ELISA was used to characterize the binding of MEDI4212 variants to human FcyRIIIa (158V or 158F). Individual wells of a 96-well Maxisorp Immunoplate (Nunc, Thermo Scientific, Waltham, MA, USA) were coated overnight at 4 °C with a 10 μg/ml solution of mAb and blocked with Pierce protein-free T20 block (Thermo Scientific, Waltham, MA, USA) for 1 h at 22 °C. Plates were washed three times with PBST (PBS with 0.1% Triton X-100). Incubation with four-fold serially diluted samples of Flag-tagged hFcyRIIIa (158V or 158F) at concentrations typically starting from 100 μg/ml for 1 h at 22 °C. Plates were incubated with anti-FLAG BioM2 antibody (Sigma-Aldrich, St. Louis, MO, USA) at 1:100 dilution followed by streptavidin horseradish peroxidase conjugate (Invitrogen) at 0.1 μg/ml for 30 min at 22 °C. Plates were developed by adding 50 μl KPL SureBlue TMB for 10 min in the dark and stopped with the addition of KPL TMB Stop Solution (50 μl per well). All plates were read by measuring the OD at 450 nm on a Spectra MAX plate reader.

Surface plasmon resonance was used to measure kinetic rates and binding constants as described previously. The equilibrium dissociation constant (Kd) for the binding of MEDI4212 IgG Fc variants to hFcyRIIIA-158V and hFcyRIIIA-158F was measured on a Biacore 3000 instrument (Biacore, Piscataway, NJ, USA). The MEDI4212 IgGs were immobilized at high density on a CM5 sensor chip using a standard amino coupling chemistry as outlined by the instrument manufacturer. The final surface density of MEDI4212 measured approximately 2000 resonance units. A reference flow cell was also prepared on this sensor chip using the identical immobilization protocol minus IgG. The stock solution and final concentration series of hFcyRIIIA-158V and hFcyRIIIA-158F were prepared in instrument buffer as 0.978–16 000 nM and 1.95–32 000, respectively. The instrument buffer was composed of PBS/Tween/ethylenediaminetetraacetic acid buffer containing 50 mM phosphate, pH 7.4, 150 mM NaCl, 3 mM ethylenediaminetetraacetic acid and 0.005% Tween-20. Receptor solutions were injected over IgG surfaces for 50 min at a flow rate of 5 μl/min, and the binding data were collected. Following each injection, bound receptor was removed with a brief pulse (injection) of 5 mM HCl. Once binding data had been collected for the entire concentration series, the steady-state data were fit to a one-site (1:1) binding model to determine the Kd for each interaction.

Primary peripheral blood mononuclear cell (PBMC) IgE class switch, IgE quantitation and IgE ELISpot
IgE class switching in PBMCs was described previously. Human whole blood was collected from healthy donors in accordance with IRB approval and corporate bioethics policies in sodium citrate cell preparation tubes and centrifuged at 1700g for 25 min at room temperature. The red blood cells were predominantly beneath the density gradient, and the
supernatant, containing PBMCs, was decanted. PBMCs were centrifuged at 524g for 7 min and the supernatant was aspirated. Pellets were resuspended in 10 ml of complete medium (RPMI minus phenol red, 2% penicillin-streptomycin, 1% L-glutamine, 0.1% β-mercaptoethanol, 10% FBS and 0.5% HEPES). Cells were centrifuged for 10 min at 200g with the brake off. Supernatants were aspirated and residual red blood cells were lysed in ACK lysis buffer according to the manufacturer’s instructions. Lysis was quenched in less than 5 min with the addition of 10 ml of complete medium. Cells were centrifuged for 5 min at 524g. Cell pellets were resuspended in 10 ml of complete medium and passed through a 50 μm filter. Freshly isolated PBMCs (1×10^6 cells/well) were IgE class-switched by incubation with interleukin-4 (IL-4) (10 ng/ml) and anti-CD40 (1 μg/ml) and co-incubated with 10 μg/ml of either IgG1 isotype control, MEDI4212 aFuc or anti-migis F4 aFuc (positive control) at 37 °C for 7 days in a 5% CO₂ incubator. Following the 7 day incubation, cells were washed twice in complete medium and transferred to Mabtech IgE ELISpot plates and incubated for 24 h at 37 °C in a 5% CO₂ incubator. The Mabtech IgE ELISpot protocol was followed for IgE ELISpot development. ELISpot images and spot quantitation were performed using a Bioreader 5000-Fx from BIO-SYS GmbH, Karben, Germany.

RNA was isolated from class-switched PBMCs with the RNeasy Plus Micro kit (Qiagen, Venlo, Netherlands), treated with DNasel and reverse transcribed with oligo(dT)20 primers using the SuperScript III first-strand synthesis kit as suggested by the manufacturer (Invitrogen). Ten percent of the cDNA reaction was analyzed using TaqMan Fast Universal PCR Master Mix (2×) andNo AmpErase UNG TaqMan chemistry (Catalog #4352042; Life Technologies), and quantitative RT-PCR was performed in a Prism 7900HT (Applied Biosystems, Foster City, CA, USA). IgE expression was calculated after amplification with a custom-designed IgE-specific TaqMan assay (forward primer: CCCTGGTAGCAACACTC; Reverse primer: GTCCCGAGCACTGTTAAG; TaqMan probe: FAM-TCTGTT- CCAAGACTGGTGGACAAC-TAMRA) and comparison to an 18S internal control (Catalog # Hs99999901-s1; Life Technologies) using the comparative C_ method. The fold-change in IgE expression among class-switch samples incubated with antibody versus control class-switched samples was determined.

Antibody-dependent cellular cytotoxicity assay
The ADCC activity of the antibodies was assessed as follows. Transfected IgE target cells, 293T cells and 293T expressing membrane IgE or N50 and N50 cells expressing membrane IgE, CD79a and CD79b were harvested using cell dissociation buffer (Invitrogen) and resuspended in ADCC assay buffer (RPMI 1640 supplemented with 5% FBS) at a density of 2×10^5 cells/ml. The cells were added to a 96-well round bottom tissue culture plate (BD Biosciences, Franklin Lakes, New Jersey, USA) at 50 μl/well, along with various concentrations of antibody at 50 μl/well in ADCC assay buffer, and pre-incubated at 37 °C for 30 min. As a positive control, ADCC activity of anti-CcmX.migis antibodies with mIgE-expressing 293T and N50 cells as targets was tested at different doses. A transfected NK cell line was used as effector cell at an E/T ratio of 2.5:1. Cells were incubated with serial dilutions of antibody for 5 h. Target cell lysis was measured by detecting the release of lactate dehydrogenase (LDH). All assays were performed in triplicate. For maximum cell lysis, 25 μl/well of 9% Triton X-100 (BD Biosciences, Franklin Lakes, New Jersey, USA) was added to the control wells. The plates were centrifuged at 300g for 3 min and incubated at 37 °C for 4 h. Plates were then centrifuged at 300g for 10 min, and 50 μl of supernatant from each well containing LDH released from lysed cells was transferred to MaxiSorp 96-well plates (BD Biosciences). A 50 μl volume of reconstituted substrate mix (CytoTox 96 Non-Radioactive Cytotoxicity Assay kit; Promega) was added to the wells, and plates were incubated in the dark at room temperature for 30 min. Reactions were terminated with 50 μl stop solution (Promega). LDH activity was quantified by measuring the absorbance at 490 nm. Percent cytotoxicity was calculated as follows:

Percent cytotoxicity = (experimental − effectors spontaneous − target spontaneous)/(target maximum − target spontaneous) × 100

where experimental corresponds to the signal measured in experimental wells; effector spontaneous corresponds to the signal measured in the presence of PBMCs alone; target spontaneous corresponds to the signal measured in the presence of 293T (transfected or untransfected) target cells alone; and target maximum corresponds to the signal measured in the presence of detergent-lysed 293T cells (transfected or untransfected).

NK/FcγRIIIα-NFAT cells and assay description
Classic ADCC is triggered when the Fc domain of a mAb associates with FcγRIIIα receptors on the surface of effector cells. This antibody-FcγRIIIα binding event triggers signaling events that induce the release of granzyme B and perforin, causing perforation of the target cell membrane and entry of granzyme B, which results in cell killing. A surrogate reporter bioassay for ADCC effector function was generated by transducing (lentiviral vectors) FcγRIIIα (158V) in NK-92 cells. Infected populations were sorted for high levels of cell surface FcγRIIIα expression. Surrogate monitoring of ADCC was achieved by co-expression of NFAT-luciferase reporter genes. Activation of FcγRIIIα induces NFAT signaling. A ratio of 2:1 (T/E) was found to achieve the most consistent results with the NK/FcγRIIIα-NFAT reporter cells.

RESULTS
MEDI4212 variants have increased affinity for FcγRIIIα
We previously reported a high-affinity neutralizing anti-IgE, MEDI4212. Fc region variants of this antibody were generated to enhance ADCC effector function. Four separate constructs were generated: (i) MEDI4212WT; (ii) MEDI4212 2M2, which
contains mutations S239D and I332E in the Fc region that have been shown to enhance ADCC; (iii) MEDI4212 3M, which contains mutations S239D, A330L and I332E in the Fc region that have been shown to enhance ADCC; and (iv) MEDI4212 aFuc, which is WTMEDI4212 produced in FUT8-deficient CHO cells and elicits an afucosylated IgG1 with the Fc region and its receptor, FcγRIIIa. In vitro assays were established to evaluate the affinity of the Fc region of the MEDI4212 variants to FcγRIIIa. Two commonly occurring genetic polymorphisms of the FcγRIIIa receptors (158V and 158F) have been described and are known to influence binding to IgG1. 158V mediates high-affinity binding and 158F mediates lower-affinity binding to Fc receptors. Binding of MEDI4212 to FcγRIIIa was evaluated by ELISA (Figure 1 for 158V or 158F, respectively). As anticipated, the binding of MEDI4212 WT was considerably lower (approximately 100-fold) for 158F than for 158V (Table 1). All MEDI4212 variants showed significant improvement in FcγRIIIa binding (Figure 1), and this improvement was particularly dramatic for 158F (Figure 1b). Surface plasmon resonance was used to determine the $K_D$ for MEDI4212 variants and FcγRIIIa. The relative improvement in affinity was similar to the improvements observed by ELISA (Table 1).

**MEDI4212 variants bind cell surface IgE**

293T and N50 cells were engineered to express IgE-containing the membrane receptor. To verify that the cells expressed IgE at the surface and that it could be bound by MEDI4212, flow cytometry was performed. MEDI4212 did not bind to the surface of control 293T (Figure 2b). However, MEDI4212 and variants all showed significant binding to the membrane IgE-transfected 293T cells (Figure 2a) and N50 cells. No binding was observed with an IgG1 control (Figure 2a, red line) or anti-human Alexa 647 secondary control (Figure 2a, gray line). These data suggest that the vast majority of IgE-engineered cells express IgE at the surface, and that the antibodies bind to the surface IgE.

**MEDI4212 variants inhibit IgE–FcγRI interaction**

The primary mechanism of action by which anti-IgE (omalizumab) inhibits the IgE-mediated phenotype is by inhibition of IgE interacting with FcγRI on the surface of basophils and mast cells. MEDI4212 variants were evaluated for their ability to prevent IgE–FcγRI interaction. RBL-2H3 (rat basophilic cell line) cells were stably transfected with human FcγRI. Activation of FcγRI receptor by crosslinking receptor-bound IgE leads to calcium mobilization that was detected using an FLIPR. All MEDI4212 variants were potent inhibitors of IgE-induced signaling (Figure 2c). Changes in the MEDI4212 Fc region had no effect on the antibody's ability to potently inhibit IgE–FcγRI interaction (Calcium Flux). Additionally, human mast cells (LAD2 cells) that naturally express human FcγRI were used to evaluate activation of FcγRI signaling. Addition of IgE to these cells leads to crosslinking the receptor bound IgE, activation of cells and secretion of mediators such as histamine and beta-hexosaminidase. All MEDI4212 variants were potent inhibitors of IgE-induced beta-hexosaminidase release (Figure 2d).

**MEDI4212 variants have enhanced ADCC activity**

An increase in antibody affinity for FcγRIIIa has been shown to enhance effector function and ADCC activity. A variety of assays were performed to demonstrate that MEDI4212 variants enhance ADCC. 293T or N50 cells that overexpress surface IgE (membrane-bound) were used as target cells for these assays. First, a surrogate reporter assay for ADCC was established using NK/FcγRIIIa-NFAT cells that overexpress FcγRIIIa (158V) and NFAT-luciferase and provide a relative measure for activation of the FcγRIIIa signaling pathway. Consistent with its weak binding to FcγRIIa (Figure 1), MEDI4212 WT had minimal activity in this assay (Figure 3a). MEDI4212 aFuc, 3M and 2M2 had average EC50 values of 28.6±11.2 ng/ml, 28.9±9.2 ng/ml and 38.9±13.5 ng/ml, respectively, $n=5$ (Figure 3a). Second, as a more direct measure of cell killing, KC133 natural killer cells were added to 293T-IgE

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**Figure 1** MEDI4212 variants have increased affinity for FcγRIIIa (ELISA). MEDI4212 variants were coated on plates and assessed for binding to: (a) FcγRIIIa-FLAG (158V) MEDI4212 aFuc, 3M 2M2 and WT had EC50 values of 64.1±23.6 ng/ml, 19.0±7.3 ng/ml, 20.5±11.8 ng/ml and 272.7±78.3 ng/ml, respectively ($n=3$); (b) FcγRIIIa-FLAG (158F) MEDI4212 aFuc, 3M 2M2, and WT had EC50 values of 72.0±25.8 ng/ml, 56.0±6.4 ng/ml, 112.0±31.2 ng/ml and 27696±4977 ng/ml, respectively ($n=3$). MEDI4212 variants demonstrated enhanced binding to both FcγRIIIa types compared with MEDI4212 WT.
target cells at a 2.5:1 (E/T) ratio. Cytotoxicity was assessed by quantitatively measuring the release of lactate dehydrogenase. MEDI4212 variants had enhanced ADCC-mediated killing when compared to MEDI4212 WT. MEDI4212aFuc, 3M and 2M2 had average EC50 values of 18.9 ± 11.5 ng/ml, 13.3 ± 9.7 ng/ml and 21.7 ± 6.0, respectively, n = 6 (Figure 3b). Although nearly 100% of the cells appeared to be labeled by MEDI4212 in the flow cytometry analyses (IgE expression at the cell surface), only 20%–30% cytotoxicity was observed in the ADCC assay. Longer incubation times and a higher E/T ratio did not improve the percent cytotoxicity observed. Similar studies were performed using IgE-NS0 target cells (Figure 3c).

| Table 1 158V and 158F affinity measurements |
|-------------------------------------------|
|                                          |
| **MEDI4212 3M**                          |
| Average of three studies                  |
| EC50 ng/ml (s.d.)                         |
| 158V ELISA                                |
| 19.0 (7.3)                                |
| 158F ELISA                                |
| 56.0 (6.4)                                |
| 158V Biacore                              |
| 7                                            |
| 158F Biacore                              |
| 36                                           |
| **MEDI4212 2M2**                          |
| Average of three studies                  |
| EC50 ng/ml (s.d.)                         |
| 20.5 (11.8)                               |
| 112.0 (31.2)                             |
| 7                                          |
| 67                                          |
| **MEDI4212 aFuc**                         |
| Average of three studies                  |
| EC50 ng/ml (s.d.)                         |
| 64.1 (23.6)                               |
| 72.0 (25.8)                               |
| 50                                         |
| 389                                        |
| **MEDI4212 WT**                           |
| Average of three studies                  |
| EC50 ng/ml (s.d.)                         |
| 272.7 (78.3)                              |
| 27696 (4977.5)                            |
| 502                                        |
| 4963                                       |

Abbreviation: WT, wild-type.

Figure 2 MEDI4212 variants inhibit Calcium Flux and degranulation, and bind cell surface-associated IgE. 293T and NS0 cells were engineered to express IgE at the surface. (a) MEDI4212 was used to detect the IgE-expressing cells by flow cytometry. MEDI4212 variants bind to the surface of IgE-expressing cells as efficiently as MEDI4212 WT. (b) MEDI4212 variants do not bind to the surface of WT293T cells. (c) RBL-2H3 (rat basophilic cell line) cells were stably transfected with human FcεR1. Activation of FcεR1 receptor by crosslinking receptor-bound IgE leads to calcium mobilization that can be detected using FLIPR (n = 3, representative shown). (d) Human mast cells (LAD2 cells) naturally express human FcεR1 (n = 3, representative shown). Activation of FcεR1 receptor by crosslinking receptor-bound IgE leads to activation of cells and secretion of mediators such as beta-hexosaminidase. MEDI4212 WT and ADCC-enhanced variants all inhibit calcium mobilization (c) and beta-hexosaminidase release (d) in a dose-dependent manner. ADCC, antibody-dependent cell-mediated cytotoxicity; FLIPR, Fluorometric Imaging Plate Reader; IgE, immunoglobulin E; WT, wild-type.
MEDI4212 variants consistently enhanced cell killing. MEDI4212 aFuc, 3M and 2M2 had average EC50 values of 25.6 ± 7.2 ng/ml, 20.0 ± 9.0 ng/ml and 41.7 ± 12.6 ng/ml, respectively (n = 6). A maximum cytotoxic effect of 30–50% was observed using IgE NS0 cells as the target cells. Finally, granzyme B and downstream caspase activity are fundamental early biochemical signatures of cytotoxicity. PanToxiLux is a commercially available kit designed to measure the cytotoxicity by lymphocytes to a cell-permeable fluorogenic substrate that is measured by flow cytometry. It is a cell-based assay using a cell-permeable fluorogenic substrate that is measured by flow cytometry. Granzyme B signaling was detected by flow cytometry after 15 min. MEDI4212 aFuc, 3M and 2M2 had average EC50 of 18.9 ± 11.5 ng/ml, 13.3 ± 9.7 ng/ml, and 21.7 ± 6.0, respectively (n = 6). (c) KC133 and NS0-IgE target cells at a 2.5 : 1 (E/T) ratio. LDH release was quantified as a measure of cell cytotoxicity after 14 h. MEDI4212 aFuc, 3M and 2M2 had an EC50 of 25.6 ± 7.2 ng/ml, 20.0 ± 9.0 ng/ml and 41.7 ± 12.6 ng/ml, respectively (n = 6). (d) KC133 and NS0-IgE target cells at a 5 : 1 (E/T) ratio. Granzyme B signaling was detected by flow cytometry after 15 min. MEDI4212 aFuc, 3M and 2M2 had an average EC50 of 160 ± 30.2 ng/ml, 64.8 ± 23.0 ng/ml and 55.4 ± 21.6 ng/ml (n = 3). MEDI4212 ADCC variants, but not MEDI4212 WT, mediated effective target cell killing in all assay formats tested. ADCC, antibody-dependent cell-mediated cytotoxicity; IgE, immunoglobulin E; LDH, lactate dehydrogenase; NK, natural killer; WT, wild-type.

MEDI4212 aFuc and 3M eliminate IgE class-switched human PBMCs

We sought to establish a human ex vivo assay using class-switched IgE cells. A variety of class switching conditions were tested. The most common conditions used include the addition of IL-4 and anti-CD40 or CD40L. Other studies have also included IL-17A or IL-21. We tested a variety of combinations and ratios of these conditions and found that IL-4 plus anti-CD40 gave the most consistent IgE class switching results. In purified human B-cell preparations, only a small number of the total B-cells were found to class switch to IgE as analyzed by FACS, ELISA, qPCR and ELISpot. The B cells from some donors consistently class-switched better than others. In order to understand the PBMCs from our donor population, we genotyped for FcγRIIIa variants, 158V and 158F. Interestingly, as observed by others, we found mostly heterozygous 158 V/F (48 V/F, 3 V/V and 0 F/F) in our limited donor population.37–39 Once conditions for IgE class switching had been optimized in purified B-cell preparations, PBMCs were harvested from the same donors and class-switched using identical conditions. IgEElispot, IgE ELISA and IgE qPCR were used to verify IgE class switching and expression. The addition of MEDI4212 aFuc, 3M or the anti-migis antibody, an antibody specific to the membrane region of IgE, consistently reduced the number of IgE-expressing cells to background levels noted in unstimulated cells (Figure 4a). Additionally, MEDI4212aFuc, 3M or anti-migis antibody elicited a substantial drop in the level of IgE expression as observed by qPCR (Figure 4b).
We hypothesized that targeting primary IgE B cells would provide long-term benefit through the elimination of the IgE B-cell pool. Combining this mechanisms of action with omalizumab (soluble IgE neutralization) would provide both short- and long-term benefits. Enhancing ADCC activity of the MEDI4212 antibody should provide the increased long-term benefits associated with IgE B-cell elimination. In addition, higher affinity anti-IgE (MEDI4212) should provide clinical benefit to patients with a broader range of IgE expression levels and body weights.

Experimental systems available for testing the in vivo efficacy of ADCC-enhanced antibodies are limited. Within the oncology field, NOD/SCID/γcnull mice have been used in conjunction with tumor cells, human NK cells and ADCC-enhanced anti-CD20 antibodies. These studies require a large number of K562 leukemia target cells (1×10^6 per mouse). Within the total human B-cell population, IgE-expressing B cells are exceptionally rare. Obtaining enough IgE-expressing primary target cells to facilitate in vivo studies is not feasible. In the absence of an in vivo system to target IgE-expressing B cells, we found that a limited number of isolated human B cells could be class-switched to IgE expressing cells and that these cells were susceptible to killing with an ADCC-enhanced antibody targeting IgE. What remains largely unknown is whether the ADCC-enhanced MEDI4212 would enhance IgE memory B-cell elimination in humans. IgG1 antibodies are known to stimulate modest levels of ADCC effector function, and since the number of target IgE B cells is very low and the antibody dose is likely to be high (omalizumab dose is 150–375 mg), it is possible that MEDI4212 WT (IgG1 without enhanced effector function) may facilitate ADCC-mediated killing of the very rare and limited IgE B-cell population.

In follicular non-Hodgkin lymphoma patients treated with rituximab (anti-CD20, IgG1), 158V homozygous carriers had better outcomes than those carrying the 158F genotype. This outcome is presumably due the FcγRIIIa polymorphic variant158V, which has a much higher affinity for IgG1 Fc than 158F. Increasing the affinity of the antibody for of FcγRIIIa provides a dramatic increase in bindingof158F compared to 158V. The increase in FcγRIIIa affinity may broaden the efficacy of the antibody to include those patients that are homozygous 158F carriers. However, as reported by others in small clinical trials, in our limited donor population we found only 3 V/V and 0 F/F out of 51 donors. In addition to NK cells, FcγRIIIa is also expressed on activated monocytes and macrophage subsets. Immune responses by monoocyte- or macrophage-expressing FcγRIIIa could also potentially be heightened upon of engagement of immune complexes containing enhanced FcγRIIIa affinity antibodies as has been shown with NK cells. The effects that these particular cell types have on the activity of the growing number of clinically validated FcγRIIIa affinity-enhanced antibodies remain an open question.

Omalizumab is approved for use in atopic asthma, and has been effective as an add-on therapy. However, because of dosing restrictions dictated by the regulatory agencies, about
one-third of severe asthmatics are not eligible for omalizumab due to high IgE levels and/or body weight. There are a number of reports that demonstrate an improvement in asthma in patients treated with omalizumab whose IgE levels exceeded the treatment guidelines. Modeling suggests that a high-affinity anti-IgE (MEDI4212) may expand the treatable population to include patients with higher IgE levels or body weights. ADCC-enhanced MEDI4212 has the potential to both neutralize soluble IgE and target IgE-expressing B cells through ADCC. This approach should have the combined benefits of soluble IgE neutralization and IgE B-cell elimination that, over time, should decrease the pool of IgE-secreting plasma cells.

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