FcγRIV is a mouse IgE receptor that resembles macrophage FcεRI in humans and promotes IgE-induced lung inflammation

David A. Mancardi,1,2 Bruno Iannasccoli,1,2 Sylviane Hoos,3,4 Patrick England,3,4 Marc Daéron,1,2 and Pierre Bruhns1,2

1Institut Pasteur, Département d’Immunologie, Unité d’Allergologie Moléculaire et Cellulaire, Paris, France.
2INSERM, U760, Paris, France.
3Institut Pasteur, Département de Biologie Structurale et Chimie, Plateforme de Biophysique des Macromolécules et de leurs Interactions, Paris, France.
4CNRS,URA 2185, Paris, France.

Introduction

A novel murine receptor for the Fc portion of mAbs (FcR) was recently cloned on the basis of bioinformatics database search. This receptor is among the many FcR-like (FCRL) molecules identified in mammals, and it was first named murine FCRL3 (NCBI sequence BC027310) (1). FCRLs have no known ligand except murine FCRL3. As it was found to bind IgG, mouse FCRL3 was renamed mFcRIV.

mFcRIV binds mouse IgG2a and IgG2b with an intermediate affinity (equilibrium association constant \([K_a] \approx 2.9 \times 10^7 \text{ M}^{-1}\) and \(1.7 \times 10^7 \text{ M}^{-1}\), respectively; ref. 2). Two main types of FcRs can be distinguished on the basis of their affinity for immunoglobulins. Monomeric immunoglobulins can bind to high-affinity (\(K_a \approx 10^9-10^{10} \text{ M}^{-1}\)) but not to low-affinity (\(K_a \leq 10^6 \text{ M}^{-1}\)) receptors. As a consequence, a proportion of high-affinity receptors are occupied in vivo, whereas low-affinity receptors remain free, even though they are exposed to high concentrations of circulating immunoglobulins in vivo (3). Immune complexes (ICs) bind to low-affinity receptors with a high avidity. They also bind to high-affinity receptors. Both types of receptors signal when they are aggregated at the cell surface by mAbs and multivalent antigen (Ag). Rather than on the affinity of receptors, signals generated upon FcR aggregation depend on functional motifs contained in the intracellular domains of FcR subunits engaged in receptor aggregates.

mFcRIV is an activating receptor (2). Like most activating FcRs, it associates with the common FcRγ subunit. FcRγ is a homodimer that contains 2 immunoreceptor tyrosine-based activation motifs (ITAM). The phosphorylation of FcR ITAMs by Src kinases initiates the constitution of an intracellular signaling complex, which activates an array of metabolic pathways leading to cell responses. FcRγ-dependent activation signals are amplified by FcRβ (4), another ITAM-containing subunit expressed in mast cells and basophils. mFcRIV does not associate with FcRβ.

In order for mFcRIV to be expressed at the cell membrane, it must associate with FcRγ (2). FcRγ indeed determines the membrane expression of multichain FcRs, i.e., mFcRIV, human and murine high-affinity receptors for IgE (FcεRI) and IgG (FcγRI), and human and murine low-affinity receptors for IgG (FcγRIIa) (5). FcγRII associates with multichain FcRs (6) expressed in mast cells and basophils (7). It is, however, mandatory for the expression of mFcεRI only (8, 9). FcεRI can therefore be expressed with FcRγ [hFcεRI(αγ)] in mast cells and basophils or without in monocytes, macrophages, and neutrophils [hFcεRI(γ)], especially in atopic individuals. hFcεRI may therefore be expressed in 2 forms depending on the cell type and on the species. mFcRIV is expressed in mouse monocytes, macrophages, and neutrophils.

mFcRIV was recently reported to bind mouse IgE of the b (IgEβ) but not of the a (IgEα) allotype (10). We show here that mFcRIV is a low-affinity receptor for IgE irrespective of the 2 known allotypes (11, 12). FcγRIV does not exist in humans. On the basis of sequence

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Nonstandard abbreviations used: Ag, antigen; BAL, bronchoalveolar lavage; BMMC, BM-derived mast cell; FcR, receptor for the Fc portion of mAbs; GaM, goat anti-mouse; IC, immune complex; IgE, IgE of the b allotype; IgG, IgE of the a allotype; i., intranasally(ly); ITAM, immunoreceptor tyrosine-based activation motif; \(K_a\), equilibrium association constant; quintuple-KO mice, mFcRIV(−/−) or hFcεRI(−/−) mice; mFcRIV−/−mTT23−/− deficient mice; SPR, surface plasmon resonance; TNP, 2,4,6-trinitrophenyl hapten.

Conflict of interest: The authors have declared that no conflict of interest exists.

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Figure 1
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IgE ICs, but not monomeric IgE, bind to mFcγRIIIA. (A) Schematic representation of FLAG-tagged FcγR α chains associated or not with FcγRγ expressed by transfectants. Green boxes represent ITAMs; the gray box represents an immunoreceptor tyrosine-based inhibition motif; the black stripe represents the FLAG-tag. Histograms show the binding of anti-FLAG mAb (black line) to FLAG-tagged FcγR on CHO transfectants and the binding of anti-mFcγRI mAb (black line) to mFcγRII on BMMCs or the binding of an isotype control (solid gray). (B) Histograms show the binding of mouse IgE to FcγRII CHO and to BMMCs, using 50 μg/ml ultracentrifuged or nonultracentrifuged mouse IgE and 15 μg/ml FITC-conjugated F(ab′)2 anti-mouse Ig alone. The binding of SPE-7 to FcγRII was analyzed using 2 different lots of SPE-7 and each gave similar results. (C) Histograms show the binding of IgG2b ICs, IgE ICs, or IgE ICs to FcγRII CHO and to BMMCs as revealed by neutravidin staining. ICs were made using TNP5-BSA-biotin and 15 μg/ml anti-TNP mAbs. Solid gray histograms show the binding of Ag alone as revealed by neutravidin staining. N.T., not tested. Data are representative of 2 (B) or 5 (C) independent experiments.

homology in extracellular domains, hFcγRIIIA was proposed to be the human homolog of mFcγRIII (1). We show that hFcγRIIIA has no detectable affinity for human IgE.

We also show that, in spite of having an intermediate affinity, mFcγRIII binds mouse IgG2a and IgG2b as monomers and functions as a high-affinity receptor. IgE ICs can, however, displace IgG2 from mFcγRIII. When aggregated by IgE ICs of both allotypes, mFcγRIII triggered Ca2+ responses in transfected cells and induced macrophage-like transformed cells and peritoneal macrophages to secrete TNF-α.

hFcγRIII(εγ) has not only an affinity for IgE but also the same quaternary structure and a similar tissue distribution as mFcγRIII. The engagement of hFcγRIII(εγ) was reported to activate monocytes (13) and neutrophils (14). We found that, when expressed by peritoneal macrophages from transgenic mice, hFcγRIII(εγ) triggered similar TNF-α responses as endogenous mFcγRIII upon aggregation by IgE ICs. The expression of hFcγRIII(εγ) is increased by monocytes, macrophages, and neutrophils of asthmatic patients. We found that mouse bronchoalveolar lavage (BAL) macrophages express neither mFcεRI nor mCD23 but express mFcγRIII, and they secreted TNF-α upon stimulation with IgE ICs. Finally, using mice deficient for mFcεRI/IIb/IIIA−/−mFcεRI−/−mCD23−/− (quintuple-KO), that only expressed mFcγRIII, we demonstrate that the in vivo engagement of mFcγRIII by IgE ICs synergizes with mediators released by IgE-activated mast cells to induce lung inflammation.

On the basis of these results, we propose that mFcγRIII is a low-affinity IgE receptor expressed by a subset of myeloid cells, and that, rather than hFcγRIIIA, hFcγRI(εγ) expressed by monocytes, macrophages, and neutrophils of atopic donors is the human equivalent of mFcγRIII. These properties endow mFcγRIII with what we believe to be novel physiopathological roles in murine models of allergic diseases and parasite infections.

**Results**

mFcγRIII, but not hFcγRIII, is a low-affinity IgE receptor. The binding of murine IgE was investigated on CHO transfectants expressing similar levels of FLAG-tagged murine FcγRI (FcγRI CHO) (Figure 1A). High concentrations of 100,000 g ultracentrifuged monomeric IgE bound to mFcγRII-expressing mouse BM-derived mast cells (BMMCs) used as positive controls, but not detectably to FcγRI CHO (Figure 1B). Monomeric IgE of the 2 known allotypes, IgEa and IgEb, displayed the same binding properties. Nonultracentrifuged preparations of the same IgE, however, bound to mFcγRIII and mFcγRIIIA but not to mFcγRI or mFcγRIIB, except SPE-7, which bound also to mFcγRII. They bound in a manner similar to monomeric IgE binding to BMMCs (Figure 1B). Nonultracentri-
fuged IgE bound more efficiently to mFcγRIV than nonultracentrifuged IgE. These results indicate that nonultracentrifuged IgE preparations contained IgE aggregates that were removed upon ultracentrifugation, and that IgE aggregates but not monomeric IgE could bind to mFcγRIV. Untreated and ultracentrifuged IgE solutions were analyzed for homogeneity and aggregate content by dynamic light scattering (Supplemental Figure 1). Both untreated but not ultracentrifuged IgE and IgE contained aggregates, but IgE contained these in higher proportion than IgE. Noticeably, untreated IgE SPE-7 contained aggregates of bigger size than other IgE preparations. The higher proportion of aggregates among IgE than among IgE preparations may indeed explain their differential binding to mFcγRIV.

As mFcγRIV bound IgE aggregates, we investigated the binding of preformed IgE-Ag ICs and, as positive controls, of IgG2b ICs (2), to the same cells. IgG2b ICs bound comparably to mFcγRI, mFcγRIIB, mFcγRIIIA, and mFcγRIV. IgE ICs bound as efficiently as IgG2b ICs to mFcγRIV, less efficiently to mFcγRIIB and mFcγRIIIA, and not detectably to mFcγRI. mFcγRIIB and mFcγRIIIA were previously reported to behave as low-affinity IgE receptors (15). IgE ICs of the 2 allotypes bound comparably to mFcγRIV (Figure 1C). These results suggest that mFcγRIV has a low affinity for IgE.

To measure the affinity of mFcγRIV for IgE, FLAG-tagged extracellular domains of the 4 murine FcγRs and, as a positive control, of hFcεRI were produced in HEK293T cells. These molecules were N-glycosylated as demonstrated by SDS-PAGE analysis before and after peptide:N–glycosidase F treatment (Figure 2A). They were covalently immobilized onto activated dextran surfaces and used for surface plasmon resonance (SPR) analysis. IgE bound to mFcγRI, mFcγRIIB, and mFcγRIIIA but not to mFcγRI. IgE bound to mFcγRIV with a $K_A$ of approximately $2.6 \times 10^5$ M$^{-1}$ (Tables 1 and 2). IgE bound with a 10-fold lower $K_A$ to mFcγRIIB and mFcγRIIIA. It dissociated much faster from FcγRs than from hFcεRI (Figure 2B). The association rate constant ($k_{on}$) of IgE for mFcγRIV and for hFcεRI were of similar magnitudes, but the dissociation rate constant ($k_{off}$) of IgE for mFcγRIV was 250-fold higher (Table 1). As a consequence, IgE bound much more transiently to mFcγRIV ($t_{1/2} = 5$ seconds) than to hFcεRI ($t_{1/2} = 17$ minutes). Comparable kinetic parameters were obtained for 3 IgE$^a$ and 2 IgE$^b$ (Table 2).
Since hFcγRIIIA has been proposed to be the homolog of mFcγRIV, we investigated the binding of human IgE on another set of CHO transfectants expressing similar levels of FLAG-tagged hFcγRIIIA or hFcγRIIB (Supplemental Figure 2A). The 2 known polymorphic variants of hFcγRIIIA (16) and the 3 known polymorphic variants of hFcγRIIIB (17, 18) were included. BMMCs from hFcγRIIIs mice (19) were used as positive controls. High concentrations of monomeric human IgE bound to hFcγRIIIs BMMCs but not to transfectants expressing hFcγRIIIA or hFcγRIIIB (Supplemental Figure 2B). The same concentrations of the same IgE failed to bind to hFcγRIII+ CHO when in complex with F(ab′)2 anti-human F(ab′)2 (Supplemental Figure 2C). They also failed to bind to transfectants expressing hFcγRI, hFcγRII (H31a or R31a variants; ref. 20), hFcγRIIB, or hFcγRIIC (data not shown). These complexes, however, bound more avidly than monomers to hFcγRIITs BMMCs. Human IgE, therefore, do not bind to hFcγRII. Supporting this observation, SPR analysis showed no measurable affinity of hFcγRIIIa (F176γ or V176α) or hFcγRIIIB (NA1, NA2, or SH) for 3 human IgE (data not shown). These data altogether indicate that hFcγRIIIA is not the homolog of mFcγRIV as for their interactions with IgE.

IgE ICs displace monomeric IgG from mFcγRIV. mFcγRIV was described as having an intermediate affinity for mouse IgG2a and IgG2b. Although this affinity is at least 10-fold higher than that of murine low-affinity FcγR2 (2), IgG ICs bound similarly to mFcγRIV, to mFcγRII, and to mFcγRIIIA (Figure 1C). This affinity is 5-times lower than that of high-affinity mFcγR (3), and we wondered whether monomeric IgG would bind to mFcγRIV. The same CHO transfectants that were used in Figure 1 were incubated with 100,000 g ultracentrifuged monomeric IgG1, IgG2a, or IgG2b. As expected, neither IgG1 nor IgG2b bound to mFcγRI, mFcγRII, or mFcγRIIIA, whereas IgG2a bound to mFcγRI but not to mFcγRII or mFcγRIIIA. Both IgG2a and IgG2b, but not IgG1, bound to mFcγRIV (Figure 3A). mFcγRIV, therefore, binds monomeric IgG2a and IgG2b, even though it has an intermediate affinity.

To investigate whether, when bound onto mFcγRIV and cross-linked, monomeric IgG2 can activate cells, we generated quintuple-KO mice. Such mice express mFcγRIV as their sole FcR. Thio-glycollate-elicited peritoneal macrophages from these mice secreted comparable amounts of TNF-α upon challenge with F(ab′)2 goat anti-mouse (GaM) when preincubated with monomeric IgG2a or IgG2b but not when preincubated with monomeric IgEα or IgEβ (Figure 3B). mFcγRIV, therefore, functions as an activating high-affinity receptor for IgG2a and IgG2b.

Table 1  Kinetic parameters of soluble FcR ectodomains for IgEα (C48-2)R

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<th>(K_a(M^{-1}))</th>
<th>(K_o(M))</th>
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<td>5.0 × 10^{-5}</td>
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<td>n.m.</td>
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</tr>
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<td>1.000</td>
<td>5.9 × 10^{-10}</td>
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\( ^a \)Data generated by injecting the concentrations of soluble monomeric IgG1 (C48-2) indicated in Figure 2B onto immobilized FcR ectodomains; \( ^b \)Dissociation constant; \( k_{on} \), association rate constant; \( k_{off} \), dissociation rate constant; n.m., not measurable.

mFcγRIV engagement by IgE ICs activates macrophages. The aggregation of mFcγRIV by IgG2a ICs was reported to trigger a Ca\(^{2+}\) response in DT40 cells (2). Likewise, IgG ICs, but not monomeric IgE, induced an increase in the intracellular Ca\(^{2+}\) concentration of mFcγRIV+ DT40 transfectants (Figure 4A). IgE of the 2 allotypes behaved similarly, except SPE-7, which triggered Ca\(^{2+}\) fluxes in the absence of Ag.

To determine whether mFcγRIV-dependent IgE-induced intracellular signals can induce biological responses, we first used the MH-S cell line. Because these cells express mFcγRI, mFcγRIIIB, and mFcγRIIV but not mFcγRI or mCD23 (data not shown), they were preincubated with 2.4G2 F(ab′)2, to block all IgE-binding receptors but mFcγRIV. Indeed, although intact 2.4G2 IgG bound to mFcγRIV+ CHO, 2.4G2 F(ab′)2 neither bound to mFcγRIV+ CHO nor blocked the binding of IgE ICs. mAb 9G8 specifically recognized mFcγRIV and blocked IgE IC binding (Supplemental Figure 4, A and B). When crosslinked with F(ab′)2, anti-Ig, 9G8 induced 2.4G2 F(ab′)2–blocked MH-S cells to secrete TNF-α (Figure 4B). As previously reported, IFN-γ upregulated the expression of mFcγR2 (2), but not that of mCD23, on MH-S cells (Figure 4C). IFN-γ–treated, 2.4G2 F(ab′)2–blocked MH-S cells secreted TNF-α when challenged with IgE ICs. They produced comparable amounts of TNF-α in response to IgE ICs or to IgG2b ICs (Figure 4D) and in

mFcγRIV being also a low-affinity receptor for IgE, we wondered whether IgE ICs could bind to mFcγRIV in the presence of IgG. The binding of IgE ICs was reduced when mixed with a saturating concentration of IgG2a (Supplemental Figure 3A). IgG2a bound to mFcγRIV under these conditions, albeit slightly less than in the absence of IgE ICs (Supplemental Figure 3B). Likewise, the binding of IgE ICs was reduced but remained detectable when mFcγRIV+ CHO were incubated with IgE ICs diluted 1:2 in normal mouse serum (Figure 3C). IgEα and IgEβ behaved similarly in every condition. Therefore, IgE ICs and noncomplexed IgG2a compete with each other for binding to mFcγRIV.

We next investigated whether IgE ICs could bind to mFcγRIV when saturated by IgG. We found that IgE ICs could displace previously bound IgG2a (Supplemental Figure 3C) and bind to mFcγRIV (Supplemental Figure 3D). Likewise, we found that IgE ICs bound comparably to mFcγRIV+ CHO, whether these were preincubated with normal mouse serum diluted 1:2 or not (Figure 3D). IgE of both allotypes behaved similarly. These data imply that IgG2a can dissociate from mFcγRIV. IgG2a and IgG2b indeed dissociated rapidly from mFcγRIV at 37°C (3 ≤ 10 minutes) (Figure 3E). This observation is in keeping with the fast dissociation rates measured from SPR analysis (\( k_{off} \) [IgG2a] = 0.0631 ± 0.0145 s\(^{-1}\) and \( k_{off} \) [IgG2b] = 0.117 ± 0.011 s\(^{-1}\); Figure 3F).

Table 2  Kinetic parameters of mFcγRIV ectodomains for mouse IgEα and IgEβ

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<th>(t_{1/2}(s))</th>
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<td>IgEα (27-74)</td>
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<td>1.7</td>
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<tr>
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<td>1.5</td>
<td>1.4 × 10^{-5}</td>
<td>7.1 × 10^{-6}</td>
</tr>
<tr>
<td>IgEβ (C48-2)</td>
<td>1.0 × 10^4</td>
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<td>6.0</td>
<td>2.0 × 10^{-5}</td>
<td>4.9 × 10^{-6}</td>
</tr>
<tr>
<td>IgEβ (SPE-7)</td>
<td>3.3 × 10^4</td>
<td>0.044</td>
<td>6.8</td>
<td>7.5 × 10^{-6}</td>
<td>1.3 × 10^{-4}</td>
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\( ^a \)Data generated by injecting soluble monomeric IgEα and IgEβ onto immobilized mFcγRIV ectodomains.
response to IgE ICs or to IgE ICs (Figure 4E). IgE IC–induced TNF-α secretion was abrogated by preincubating cells with 9G8 or mFcγRII-blocking mAb 9E9 (Figure 4F and confirmed by anti–TNF-α ELISA in Supplemental Figure 4C).

**Human FcεRI(αγ) is a functional equivalent of mouse FcγRII.** In spite of having a high affinity and being expressed on mast cells and basophils, hFcεRI shares with mFcγRII similar structures and tissue distributions: both can be expressed by monocytes, macrophages, and neutrophils, and they associate with FcγRγ, but not FcRβ, in these cells (3). To investigate whether hFcεRI and mFcγRI could respond similarly to IgE IC, we used hFcεRI-Tg mice. As described previously (21), hFcεRI were expressed on the same cells in these mice as in atopic patients. They were detected on blood GR1+Mac1+ polynuclear cells and GR1+Mac1+ cells (Figure 5A). They were also detected and with a higher expression on thioglycollate-elicited peritoneal macrophages from transgenic but not from WT mice as described. mFcγRI was expressed similarly in WT and in hFcεRI-Tg macrophages. IgE ICs induced WT macrophages to

Figure 3
mFcγRI is a high-affinity receptor for IgG2a/b, but IgE ICs can bind to mFcγRI in the presence of high IgG concentrations. (A) Histograms show the binding of monomeric mouse IgG1, IgG2a, or IgG2b (27–35) (10 µg/ml 100,000 g ultracentrifuged) to FcγRI CHO, revealed by FITC-conjugated F(ab′)2 GaM. (B) Thioglycollate-elicited peritoneal macrophages from quintuple-KO mice were incubated with indicated concentrations of monomeric mouse Ig (µg/ml) and assayed for TNF-α secretion following incubation with F(ab′)2 GaM. Curves represent the percentage of cytotoxicity as a function of supernatant dilution. (C and D) Histograms show the binding of IgE ICs (C38-2a) or IgG ICs (C48-2b), (C) when diluted 1:2 in normal mouse serum or in PBS to mFcγRI CHO or (D) when mFcγRI CHO were preincubated with normal mouse serum diluted 1:2. Solid gray histograms show binding of Ag alone. IC binding was revealed by neutravidin staining. (E) mFcγRI CHO were preincubated for 1 hour at 4°C with saturating concentrations of indicated IgG. Curves represent the percentage of IgG2a (open diamonds) or IgG2b (27–35) (filled diamonds) bound to these cells after an incubation at 37°C for increasing time periods. Insets show corresponding histograms at 0 (bold black line), 10 (dark gray line), 30 (black line), and 60 (light gray line) minutes. Solid gray histograms show binding of secondary Abs alone. a-, anti.-

(F) SPR sensorgrams resulting from the injection of IgG2a or IgG2b (C48-4) onto immobilized mFcγRI ectodomains. Data are representative of 2 (A–D and F) or 3 (E) experiments that gave similar results.
secrete TNF-α. IgE ICs induced hFcεRIgG macrophages to secrete higher amounts of TNF-α than WT macrophages (Supplemental Figure 5A). To evaluate the contribution of hFcεRI only, we intravenously injected 9E9 into WT or hFcεRIgG mice and recovered thioglycollate-elicited macrophages. Treatment with 9E9 inhibited 25% of TNF-α secretion by macrophages from hFcεRIgG mice, while abolishing TNF-α secretion of macrophages from WT mice (Figure 5B and Supplemental Figure 5B). The remaining TNF-α secretion observed in 9E9-treated macrophages from hFcεRIgG mice corresponds, therefore, to the sole contribution of hFcεRI.

Both human FcεRI and murine FcγRIV enable macrophages to secrete cytokines in response to IgE ICs. The engagement of alveolar macrophage mFcγRIIB by IgE ICs promotes lung inflammation. FcεRI(αγ) are expressed by human alveolar macrophage in asthmatic patients (22). We therefore examined FcγRI on mouse alveolar macrophages. CD11c+ Mac-1+ macrophages represent 95% of BAL cells (Figure 6A). BAL cells harvested from mFcεRIIB+/mFcγRIIIA+ mice expressed mFcγRIV, but neither

Figure 4
mFcγRIV engagement by IgE ICs induces cell activation in transformed cells. (A) The histogram represents the binding of 9G8 to mFcγRIV+ DT40. Transfectants were loaded with Fluo-3, and the intracellular Ca2+ concentration was monitored, following triggering (arrows) by Ag alone (black line), IgE alone (thin lines), or IgE ICs (thick lines). Curves represent the relative intracellular Ca2+ concentration as a function of time. Two different lots of SPE-7 were assayed and gave similar results. (B) TNF-α secreted by MH-S cells, induced by LPS or by 1 μg/ml 9G8 and the indicated concentrations (μg/ml) of goat anti-hamster F(ab’)2 (GaH), was titrated in supernatants. (C) Histograms represent the binding of indicated mAbs on IFNγ-treated or untreated MH-S cells. Solid gray histograms represent the binding of isotype controls. (D and E) TNF-α secreted by IFNγ-treated 2.4G2 F(ab’)2–saturated MH-S cells, induced by the following reagents, was titrated in supernatants: (D) IgG2b ICs (Gork) or IgE ICs (C48-2β) and (E) IgE ICs (C38-2α) or IgE ICs (C48-2α). (F) IFNγ-treated MH-S cells were saturated with both 2.4G2 F(ab’)2 and Polymyxin B–treated 9G8, 9E9, or irrelevant hamster IgG (Iso.). TNF-α secreted by these cells induced by IgE ICs (C48-2β) was titrated in supernatants. Curves represent the percentage of cytotoxicity as a function of MH-S supernatant dilution. All Ig concentrations are indicated in μg/ml. Data are representative of 3 (A–C) or 2 (D–F) experiments. Mean ± SD of triplicates in TNF-α bioassays are represented (B, D, and E).
mFcγRIIB/IIIA, as expected, nor mCD23, and they bound IgE ICs of either allotype. The binding of IgE ICs was abrogated by 9G8 or by 9E9. BAL cells secreted TNF-α when challenged with IgE ICs but not when challenged with monomeric IgE (Figure 6B). WT mouse 2.4G2 F(ab′)2–blocked BAL cells also secreted TNF-α in response to polymyxin-treated IgE ICs, excluding a possible LPS contamination as being responsible for TNF-α secretion (Figure 6C). IgE of the 2 allotypes behaved similarly (Figure 6D).

To determine whether bronchoalveolar macrophage mFcγRIIV-dependent IgE-induced signals can induce an inflammatory reaction in vivo, we first used mice lacking mFcεRI and mCD23, the 2 classical IgE receptors. When instilled intranasally (i.n.), IgE ICs induced no significant infiltration of polymorphonuclear cells in the bronchoalveolar space. We hypothesized that mast cell activation is required for initiating the inflammatory response in this model, as it has been described in an autoantibody-dependent arthritis model (23). As the mast cells in the mFcεRI−/− mCD23−/− mice lack the high-affinity receptor for IgE, they cannot be activated by IgE ICs. Therefore, we chose to substitute in vivo airway mast cell activation by i.n. instillation of supernatant of WT BMMCs activated in vitro. Supernatant from high numbers of IgE-sensitized BMMCs challenged with Ag induced a dose-dependent alveolar infiltration, starting at a dose equivalent to 5 × 10^6 stimulated BMMCs (Figure 7A, 2 left panels). IgE ICs, which failed to induce a significant inflammation when administered alone, induced a robust influx of Mac1+ Gr1+ polymuclear cells in bronchial alveoli when administered to mFcεRI−/− mCD23−/− mice 1 day after an i.n. instillation of a dose of IgE-stimulated BMMC supernatant, which did not induce a detectable inflammation (2.5 × 10^6 stimulated BMMCs, inducing 1% infiltration, equivalent to the infiltration induced by unstimulated BMMC controls) (Figure 7A, 2 right panels). Because mFcεRI−/− mCD23−/− mice express mFcγRIIB and mFcγRIIIA, which could possibly bind IgE ICs, we repeated the experiments in quintuple-KO mice, which express no activating FcR. IgE ICs induced a similar weak infiltration in quintuple-KO and FcRγ−/− mice. They induced a marked Mac1+ Gr1+ polymuclear infiltration in quin-

**Figure 5**

Both hFcεRI and mFcγRIIV engagement by IgE ICs induces TNF-α secretion by peritoneal macrophages. (A) Density plots show the binding of indicated mAbs to white blood cells from WT and hFcεRI Tg transgenic mice. Histograms show the binding of anti-hFcεRI mAb to cell populations defined by 5 gates from the density plots of blood cells from hFcεRI Tg (open black) or WT (solid gray) mice. (B) Indicated mice were injected intravenously with 9E9 or irrelevant hamster IgG 1 day before recovery of thioglycolate-elicited macrophages. The cells were assayed for TNF-α secretion by ELISA following incubation on IgE-, Ag-, or IgE IC (C48-2b)–coated wells. All reagents were treated with Polymyxin B. Mean ± SD of triplicates are represented. Significant differences between cells triggered by IgE ICs are indicated (**P < 0.001; Student’s t test). The horizontal gray line represents background TNF-α levels. Data are representative of 3 (A) and 2 (B) independent experiments.
tuple-KO (Figure 7B) but not in FcRγ−/− mice (Figure 7C) when administered after a dose of IgE-stimulated BMMC supernatant, which induced no detectable infiltration. No significant increase in lymphocyte populations in the BALs from quintuple-KO mice could be observed, either by flow cytometry (BMMC supernatant, 0.10% ± 0.03%; IgE-ICs, 0.09% ± 0.06%; BMMC supernatant plus IgE-ICs, 0.17% ± 0.10%; variations not significant, Student’s t test) or by cytospin analysis (data not shown). Mediators released by IgE-sensitized mast cells upon engagement of FcεRI by Ag can, therefore, enable bronchoalveolar macrophages to induce lung infiltration, upon engagement of FcγRIV by IgE ICs.

Discussion

mFcγRIV shares several properties with hFcγRIIIA (hCD16A) and, for this reason, it was first named CD16-2 (1). mFcγRIV was found to have an intermediate affinity for mouse IgG2a (Kₘ = 2.9 × 10⁷ M⁻¹) and IgG2b (Kₘ = 1.7 × 10⁷ M⁻¹) (2). hFcγRIIIA was also found to have an intermediate affinity for human IgG (Kₘ = 2.5 × 10⁷ M⁻¹) (24, 25). The higher affinity of hFcγRIIA than that of hFcγRIIB was proposed to result from the association of hFcγRIIIA with the FcRγ subunit (26). This association confers on mFcγRIV and hFcγRIIIA their cell-activating properties. The high amino acid sequence homology (64.9%) of the extracellular domains of mFcγRIV and hFcγRIIIA may also account for their similar intermediate affinities for IgG. These common features, altogether, support the current view that hFcγRIIIA is a human equivalent of murine FcγRIV. Our data challenge this view. We show here that mFcγRIV is an IgE receptor. IgE ICs, but not monomeric IgE, bound to mFcγRIV. This was observed on mFcγRIV−/− CHO, transformed macrophagic cells, BM-derived cultured macrophages (data not shown), and freshly isolated ex vivo BAL macrophages. IgEα and IgEβ bound to all 4 cell types similarly. When measured by SPR analysis, the mean affinity of mFcγRIV for 5 different IgE was Kₘ ≈ 4 × 10⁷ M⁻¹. IgEα and IgEβ bound with the same affinities. These data, altogether, identify mFcγRIV as a low-affinity receptor for IgE, irrespective of IgE allotypes. Inter-
Interestingly, 2 amino acids (K$_{117}$ and E$_{132}$) (27), which are involved in the formation of 2 salt bridges between the extracellular domains of hFc$_{\varepsilon}$RI and the Fc portion of human IgE and which are critical for the binding of IgE, are conserved in mFc$_{\gamma}$RIIV. mFc$_{\gamma}$RIIV was recently reported to bind mouse IgE but not IgE$_a$ (10). We found the same preference for IgE$_b$ when using the same commercially available IgE under the same conditions, i.e., when not ultracentrifuged and at the same high concentrations. Binding of IgE of both allotypes, however, was abrogated when aggregates were removed by ultracentrifugation. The 2 IgE$_b$ anti-DNP, SPE-7 and C48-2, were reported to spontaneously aggregate and to activate mast cells in the absence of Ag (28). Aggregates present in higher amounts in IgE$_b$ preparations could account for the preference of mFc$_{\gamma}$RIIV for nonultracentrifuged IgE$_b$ observed by Hirano et al. (10) and by us. In contrast with mFc$_{\gamma}$RIIV, hFc$_{\gamma}$RIIIA is not an IgE receptor. hFc$_{\gamma}$RIIIA had no measurable affinity for human IgE when studied by SPR, and human IgE failed to bind to hFc$_{\gamma}$RIIIA CHO, whether as monomers or as F(ab')$_2$-anti-human F(ab')$_2$ complexes. hFc$_{\gamma}$RIIIA and hFc$_{\gamma}$RIIB gave similar results, irrespective of known polymorphisms.

We also found that IgG2a and IgG2b monomers bind to mFc$_{\gamma}$RIV and that mFc$_{\gamma}$RIV-bound IgG2 triggers cytokine secretion when crosslinked on ex vivo macrophages. mFc$_{\gamma}$RIV therefore fulfills the criteria defining high-affinity receptors. If so, one expects mFc$_{\gamma}$RIIV to be occupied by IgG in vivo. mFc$_{\gamma}$RIIV may therefore not be available for IgE ICs, especially as the concentration of IgG is much higher than that of IgE in vivo. However, we found that IgG2a/2b rapidly dissociated from mFc$_{\gamma}$RIIV at 37°C and that IgE ICs replaced IgG2a when added to IgG-saturated mFc$_{\gamma}$RIIV CHO. This finding is of critical importance as it provides a possible biological significance to mFc$_{\gamma}$RIV as a low-affinity receptor for IgE.
The ITAM-containing FcR\(\gamma\) subunit provides mFc\(\gamma\)RIV with cell-activating properties. IgE ICs triggered an increase in the intracellular Ca\(^{2+}\) concentration in mFc\(\gamma\)RIV- DT40 cells. IgE\(\alpha\) and IgE\(\beta\) induced similar Ca\(^{2+}\) responses. As expected, the anti-DNP IgE\(\alpha\) SPE-7 also triggered Ca\(^{2+}\) responses in the absence of Ag, supporting the interpretation that it spontaneously aggregates. Noticeably, SPE-7 contained qualitatively bigger aggregates than other IgEs tested by dynamic light scattering. We also observed that, differing from all other IgE tested, SPE-7 bound to mFcRI. SPE-7 has been reported previously to exhibit Ag multiplicity, binding other molecules than DNP and IgE-binding molecules, to preexist in 2 different conformations, and to exist as 4 different conformations in the presence of Ag (29). It has also been reported that SPE-7 activates mast cells in the absence of Ag (28, 30), but that monovalent hapten abolishes these properties (31), suggesting that SPE-7 auto-aggregates due to a low affinity for its own structure and/or recognizes other cell surface components (32) (reviewed in ref. 33). Altogether, these data obtained by others strongly support our data that SPE-7 contains high–molecular weight aggregates, induces activation in the absence of Ag, and binds other molecules than DNP and IgE-binding molecules, including mFcRI. Data obtained with this mAb could be questionable, especially when used at high local concentrations in vitro and in vivo: a local injection may considerably affect the probability of high local concentrations of SPE-7 aggregates compared with a systemic (intravenous) injection and therefore induce or not induce cross binding to other molecules.

In addition to triggering calcium responses, IgE ICs triggered TNF-\(\alpha\) secretion by a macrophage cell line, MH-S, and by ex vivo macrophages. IgE\(\alpha\) and IgE\(\beta\) ICs induced the secretion of similar amounts of TNF-\(\alpha\). The TNF-\(\alpha\) secretion was due to mFc\(\gamma\)RIV, as responses of both cell types were not affected by 2.4G2 F(ab\(^{\prime}\))\(_{2}\), while responses of MH-S were abolished by 9G8. ICs made of IgG2 ICs and IgE ICs induced TNF-\(\alpha\) secretions of similar intensities, indicating that they bound to mFc\(\gamma\)RIV with similar avidities, although mFc\(\gamma\)RIV has markedly different affinities for IgG2 and IgE. IgE ICs can therefore activate macrophages via mFc\(\gamma\)RIV. IgE ICs may form when IgE is synthesized locally. A local IgE production was described in the upper respiratory tract of atopic patients and in the bronchial mucosa of asthmatic patients (34). Antibody class-switching to IgE, and therefore local IgE production, is enhanced locally in the nasal mucosa in grass pollen-allergic subjects suffering from rhinitis during the pollinic season (35). They can also form in helminth infections, in which specific IgE and parasite Ag are present in high concentrations.

Human macrophages do not express FcR\(\gamma\)RI. They can, however, express another IgE receptor, hFcRRI. Although hFcRRI has a high affinity whereas mFc\(\gamma\)RIV has a low affinity for IgE, both can be engaged by IgE ICs. Moreover, hFcRRI is expressed without FcR\(\beta\) in macrophages and neutrophils and especially in atopic patients (13, 22). Mouse macrophage/neutrophil FcR\(\gamma\)RIV and human macrophage/neutrophil FcR\(\gamma\)RII(\(\alpha\)) have therefore the same quaternary structure: an IgE-binding subunit and the same ITAM-containing FcR\(\gamma\) subunit. hFcRRI expressed in macrophages from transgenic mice triggered TNF-\(\alpha\) to a similar extent as mFc\(\gamma\)RIV when engaged by IgE ICs. mFc\(\gamma\)RIV and hFcRRI have therefore similar functional properties. hFcRRI aggregation was described to trigger Ca\(^{2+}\) responses and cytokine secretion in monocytes from atopic patients (13) and in neutrophils from asthmatic patients (14). As hFcRRI(\(\alpha\)) is overexpressed on monocytes, neutrophils, and eosinophils in atopic patients and particularly on alveolar macrophages (22), we examined freshly isolated BAL cells for mFc\(\gamma\)RIV expression and function. BAL macrophages from WT, mFc\(\gamma\)RII/III/IIA\(^{-}\), and mFc\(\gamma\)RII/III/IIA\(^{-}\) mice expressed mFc\(\gamma\)RIV, but neither mFc\(\gamma\)RII (mCD23) nor mFc\(\gamma\)RI. Binding of IgE ICs to these cells was mFc\(\gamma\)RIV dependent, as binding was abolished by 9G8 or 9E9. IgE ICs, but not monomeric IgE, induced these BAL cells to secrete TNF-\(\alpha\). Comparable results were obtained with BAL cells from mFcRRI/III/IIA\(^{-}\) mice. IgE\(\alpha\) ICs and IgE\(\beta\) ICs gave similar results. These IgE-induced, mFc\(\gamma\)RIV-dependent biological responses of alveolar macrophages are of potentially high physiological relevance. TNF-\(\alpha\), which induces bronchial hyperresponsiveness (36), airway infiltration by neutrophils and eosinophils (37), and activates airway smooth muscle (38), was indeed recognized as playing a major role in asthma-associated remodeling and pulmonary inflammation, especially in asthma refractory to corticosteroid therapy (39).

These results prompted us to examine the role of mFc\(\gamma\)RIV in a murine model of passive lung inflammation. Surprisingly, while IgE ICs did induce a weak influx of polymuclear cells in BAL (8%–10%) when instilled i.n. in mFc\(\gamma\)RI/III/IIA\(^{-}\) mice, they did induce an almost undetectable influx (2.5%) when instilled i.n. in quintuple-KO mice. Therefore, IgE ICs may induce some infiltration in BAL by triggering mFcRRII\(\alpha\) on mast cells. Lung inflammation involves multiple cell types. Among these, mast cells are well known as the initiators of IgE-induced allergic reactions. We therefore “primed” mice by instilling i.n. supernatants of IgE-sensitized mast cells challenged with Ag in vitro. Mast cell supernatants, collected 30 minutes after challenge, contained granular and lipid mediators but low levels of cytokines (e.g., 92 ± 6 pg/ml TNF-\(\alpha\)). These supernatants induced a dose-dependent lung infiltration of polymuclear cells in mFc\(\gamma\)RI/III/IIA\(^{-}\) mice. Supernatants from high numbers of activated mast cells were required for inducing a significant infiltration. When administered i.n. at a dose which induced no detectable inflammation, supernatants from activated mast cells enabled IgE ICs to induce a marked lung infiltration not only in mFc\(\gamma\)RI and mCD23-deficient mice, but also in quintuple-KO mice, which express mFc\(\gamma\)RI only, but not in FcR\(\gamma\)-deficient mice, which express no activating FcR. The absence of inflammation observed in FcR\(\gamma\)-deficient mice rules out the participation of IgE-binding molecules that are not associated with the FcR\(\gamma\) chain, e.g., galectin-3 (40). The use of quintuple-KO mice excluded the contributions not only of the 2 main IgE receptors (mFcRRI and mCD23) but also those of the 2 “minor” IgE receptors (mFc\(\gamma\)RIIB and mFcRRII\(\alpha\)) to the inflammatory process we observed. It leaves mFc\(\gamma\)RIV as the sole candidate among all known FcRs. mFc\(\gamma\)RIV-expressing cells may therefore synergize with mFcRRI-expressing mast cells to generate an IgE-induced lung inflammation. Whether unidentified products in mast cell supernatant enhanced the responsiveness of mFc\(\gamma\)RIV-expressing macrophages to IgE ICs, whether vasoactive mediators initiated the extravasation of polymuclear cells into tissues, or whether secreted proteases facilitated their migration by degrading the extracellular matrix is unknown. The responsible mediators in supernatants from activated BMMMC\(\alpha\)s may not be mast cell specific; other cells could, however, possibly cooperate with macrophages in vivo.

In conclusion, we demonstrate here that mFc\(\gamma\)RIV is a low-affinity receptor for IgE, irrespective of allotypes. It can therefore operate in all mouse strains. mFc\(\gamma\)RIV has the same quaternary structure, the same tissue distribution, the same ligands, and the same functional properties as human FcRRI(\(\alpha\)). Mice may therefore
Research article

be a better model for IgE-dependent inflammation, allergies, and parasite infections than previously thought, macrophage FceRI
playing, in murine models, the role played by macrophage FceRI
in patients. The cooperation between mast cells and hFceRI-
expressing lung cells proposed here in a mouse model of IgE-
induced lung inflammation, indeed suggests that a similar coop-
eration may occur between mast cells and hFceRI-expressing lung
cells in allergic asthma.

Methods
Cells and cDNAs. CHO-K1, HEK293T, DT40, and MH-S cells were from
the ATCC and cultured as recommended by the manufacturer. BMMCs
were obtained from mouse BM cells cultured in OptiMEM (Invitrogen)
in 10% FCS, supplemented with 5% X63-IL3-conditioned medium for 4
weeks (≥ 90% mast cells).
cDNAs coding for mouse FceRI, FcεRIIB1 (Ly17.2 haplotype), FcεRIIA
(H haplotype; ref. 41), and FcRy were previously cloned in the laboratory.
cDNA from C57BL/6 and 129/Sv spleen cells were used to clone mFceRI
and mFcεRIIB1 (Ly17.1), respectively. Both mFcεRIIB1 haplotypes gave
identical results throughout this study; only results obtained with Ly17.2
haplotype are presented. Human FcεRI and FcεRIIA(R113) cDNAs were from
J. Van de Winkel and J. Leuen (University Medical Center Utrecht, Utrecht,
The Netherlands). hFcεRIIB (N1, N2, or SH) cDNAs were provided by S.
Santoso and U. Sachs (Institute for Clinical Immunology and Transfu-
sion Medicine, Giessen, Germany). cDNA from human blood cells were used to
close hFcεRIIA(N16), and hFcεRIIA(F105) by site directed mutagenesis.
A cDNA sequence coding for a FLAG tag was inserted immediately 3′ of
the signal sequence cleavage site in all FcR cDNAs. Resulting constructs were
cloned into pNT (neomycin29), cDNAs corresponding to EC domains of all
FcRs were cloned into p3×FLAG-CMV-14 (Sigma-Aldrich). Stable transfect-
ants were obtained and sorted to equivalent surface expression by flow
cytometry on a MoFlo (Dako) or FACSAria (Becton Dickinson).

Mice. mFcεRIIB/IIIα-/- and mFcεRI/IIIA-/- C57BL/6 (N6 B6) mice (42)
were provided by S. Verbeek (Leiden University Medical Center, Leiden,
The Netherlands) and backcrossed to generation N8. hFcεRIIγ (N6 B6) and
mFcεRII-/- (N6 B6) mice were provided by J.-P. Kinet (Harvard
Institutes of Medicine, Boston, Massachusetts, USA) and backcrossed to N12
B6. mFcεRII-/- (N12 B6) mice were provided by M. Lamers (Max-
Planck-Institute for Immunology, Freiburg, Germany). Quintuple-KO
(N6 B6) mice were obtained by intercrosses. WT C57BL/6J and 129/Sv
were purchased from Charles River Laboratories, and FcRy-/- mice (N12 B6)
were purchased from The Jackson Laboratories. BAls were performed on
ketamin-xylazin anesthetized mice, using 6 washes of 1 ml normal saline
solution, and cells were purified by adherence. Peritoneal macrophages
were harvested from mice injected i.p. with 2 ml thioglycollate (Bio-Rad)
4–5 days prior to harvest and purified by adherence. All mouse protocols
were approved by the Animal Care and Use Committees of Comité régional
d’éthique en matiére d’expérimentation animale, Ile-de-France,
France.

Abs and reagents. Mouse IgEs anti-2,4,6-trinitrophenyl hapten (anti-
TNP) mIgEs (C38-2, C48-2, 15.3.2) and anti-dansyl mIgE2 (27–74), were
purchased from BD Biosciences – Pharmingen; anti-DNP mIgE (SPE-74)
was purchased from Sigma-Aldrich; mouse IgE anti-OVA (2C6) were
provided by L. Kobzik (Harvard School of Public Health, Boston, Massa-
chusetts, USA) (45); and the mouse IgE anti-DNP mAb 2682-I was used as
culture supernatant. All these IgEs were purified from culture superna-
tants of hybridomas by the manufacturers or in house and cannot there-
fore contain another allotype of IgE. Several batches of each commercial
IgE were used and gave identical results.

FITC-conjugated rat anti-mIgG2a, FITC-conjugated anti-Mac1, FITC-con-
jugated anti-hFcεRI, PE-conjugated anti-h-IIIβ, PE-conjugated anti-mCD23,
PE-conjugated anti-Gr1, and hamster IgG1 were purchased from PharMin-
gen; unlabeled-, HRP- or FITC-labeled anti-FLAG (M2) and LPS were pur-
chased from Sigma-Aldrich; unlabeled and FITC-labeled F(ab′)2 GaM were
purchased from Jackson Immunoresearch Laboratories Inc.; PE-conjugated
neutravidin was purchased from Molecular Probes; and unlabeled and FITC-
labeled F(ab′)2 goat anti-hamster (GaH) were purchased from Serotec.
Hamster anti-mFcεRI mAb 9G8 and 9E9 were provided by J.V. Ravetch
(The Rockefeller University, New York, New York, USA); mIgG1 anti-TNP (D10),
mIgG2b anti-TNP (GORK), and mIgG2a anti-SRBC (12-6-22) mAbs were
provided by B. Heyman (Uppsala Universitet, Uppsala, Sweden); and human
myeloma IgE PSA was provided by T.F. Huff (Virginia Commonwealth
University, Richmond, VA, USA). Recombinant murine IFN-γ was purchased from
Peprotech, and low-endotoxin lysozyme-free OVA was purchased from MP
Biomedicals. BSA was trinitrophenylated using trinitrobenzene sulfonic acid
to 5 or 16 moles of TNP per mole of BSA.

Monomeric Ig binding assays. Aggregates in stock solutions were removed by
an 18-hour ultracentrifugation at 100,000 g, and 2 × 106 cells were incubated
with monomeric Ig at indicated concentrations for 1 hour at 4°C. Cell-
bound Ig was detected using 15 μg/ml F(ab′)2 FITC-conjugated GaM or PE-
conjugated F(ab′)2 anti-human F(ab′)2-specific Abs for 30 minutes at 4°C.

IC Ig binding assays. Unless otherwise specified, mouse IgCs were preformed
by incubating 10 μg/ml TNP, pBSA-biotin with 30 μg/ml anti-TNP mAbs
for 1 hour at 37°C, and 2 × 106 cells were incubated with ICs for 2 hours at
4°C. ICs bound to cells were detected using PE-conjugated neutravidin at
2 μg/ml for 30 minutes at 4°C.

Serum or monomeric IgG2a versus IgE IC competition assay. A total of 2 × 106
cells were incubated successively with serum diluted 1:2 or with 30 μg/ml
mIgG2a (12-6-22), washed, and incubated with IgE ICs for 2 hours at 4°C;
or incubated simultaneously with IgE ICs diluted 1:2 in normal mouse
serum; or incubated with both mIgG2a and IgE ICs for 2 hours at 4°C.

Bound mIgG2a was revealed using 5 μg/ml FITC-conjugated F(ab′)2 anti-
mIgG2a, and bound IgE ICs were revealed using PE-conjugated neutravidin
for 30 minutes at 4°C. Saturation of mFcεRI by mIgG2a or mIgG2b was
obtained from the concentration of 30 μg/ml and up (data not shown).

Dinamic light scattering. Unreated or ultracentrifuged IgE solutions at
300 μg/ml in PBS were analyzed immediately after ultracentrifugation
for homogeneity and presence of aggregates of high molecular weight,
using a DynaPro MS800 dynamic light scattering instrument (Wyatt).
Triplicates of 20 measurements at 25°C were averaged with acquisition
periods of 10 seconds. During the illumination, the photons scattered by
proteins were collected at a 90-degree angle on a 10-second acquisition
period and were fit with the analysis software Dynamics (Wyatt). Inten-
sity fluctuations of the scattered light, resulting from Brownian motion
of particles, were analyzed with an autocorrelator to fit an exponential
decay function, then measuring a translational diffusion coefficient D.

For polydisperse particles, the autocorrelation function was fit as the
sum of contributions from the various size particles using the regular-
ization analysis algorithm. D is converted to a hydrodynamics radius Rh
through the Stokes-Einstein equation (Rh = kBT/6πηD, where η is the
solvent viscosity, kT is the Boltzmann’s constant, and T is the temperature).

Apparent molecular weights for a spherical particle were deduced from
histograms of distribution of percent intensity versus radius to identify
the peak containing monomeric IgE.

Measurement of intracellular free calcium concentration. Mobilization of
intracellular free calcium concentration was determined as described
previously (46).

Secretion of TNF-α in MH-S and BAL macrophages. For 4 hours, 2.4 × 104
MH-S cells or 1.2 × 105 alveolar macrophages were incubated with indicated
Reagins at 37°C. If indicated, preincubations were performed for 1 hour at 4°C: mFcRIIB/mFcRIIA were blocked by 20 μg/ml 2.4G2 F(ab')2, or mFcRIV was blocked by 10 μg/ml 9G8 or 9E9. If indicated, reagents were preincubated 1 hour at 37°C in the presence of 2 μg/ml Polyoxymyxin B.

Secretion of TNF-α in peritoneal macrophages. All plates were coated in sodium carbonate, with or without the presence of 100 μg/ml TNPα-BSA containing 2 μg/ml Polyoxymyxin B, saturated with 1 mg/ml BSA, and incubated with 30 μg/ml IgEα anti-TNP (C48-2) or without for 1 hour at 37°C. Cells were preincubated in the presence of 20 μg/ml 2.4G2 F(ab')2, added to wells, and incubated for 3 hours at 37°C. For peritoneal macrophages from quintuple-KO mice, cells were incubated for 1 hour with monomeric Ig as indicated, washed 3 times, and incubated with 15 μg/ml F(ab')2; GaM for 3 hours at 37°C. If indicated, mice were injected intravenously with 20 μg Hamster anti-mFcRIV mAb 9E9 or irrelevant Hamster IgG 1 day before macrophage harvest.

Serial dilutions of cell-free supernatants were harvested and titrated for TNF-α by a cytotoxic assay on L929 cells that could be abrogated by anti-TNF-α mAbs as described previously (47). Additionally, TNF-α was titrated using an anti–TNF-α ELISA kit (R&D Systems).

Production of soluble FcR ectodomains tagged with 3xFLAG fusion protein. cDNA constructs coding for soluble FcR ectodomains tagged with a 3xFLAG peptide were transfected by a standard calcium chloride technique into HEK293T cells. Fusion proteins from 96-hour supernatants were purified from HEK293T cells. Expression of high-affinity binding of human immunoglobulin E by transfected cells. Cell. 1981. Genetic polymorphism (Igh-7 allotype) for immunoglobulin E (Fc epsilon RI): role in asthma. J. Exp. Med. 337:187–189.

Results are expressed as the mean ± SD. P values of less than 0.05 were considered significant.

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Address correspondence to: Pierre Bruhns, Unité d’Allergologie Moléculaire et Cellulaire, Département d’Immunologie, Institut Pasteur, 25 rue du Docteur Roux, 75015 Paris, France. Phone: 33-1-4568-8629; Fax: 33-1-4061-3160; E-mail: bruhns@pasteur.fr.


