Mouse and human neutrophils induce anaphylaxis

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Anaphylaxis is a life-threatening hyperacute immediate hypersensitivity reaction. Classically, it depends on IgE, FcεRI, mast cells, and histamine. However, anaphylaxis can also be induced by IgG antibodies, and an IgG1-induced passive type of systemic anaphylaxis has been reported to depend on basophils. In addition, it was found that neither mast cells nor basophils were required in mouse models of active systemic anaphylaxis. Therefore, we investigated what antibodies, receptors, and cells are involved in active systemic anaphylaxis in mice. We found that IgG antibodies, FcγRIIIA and FcγRIV, platelet-activating factor, neutrophils, and, to a lesser extent, basophils were involved. Neutrophil activation could be monitored in vivo during anaphylaxis. Neutrophil depletion inhibited active, and also passive, systemic anaphylaxis. Importantly, mouse and human neutrophils each restored anaphylaxis in anaphylaxis-resistant mice, demonstrating that neutrophils are sufficient to induce anaphylaxis in mice and suggesting that neutrophils can contribute to anaphylaxis in humans. Our results therefore reveal an unexpected role for IgG, IgG receptors, and neutrophils in anaphylaxis in mice. These molecules and cells could be potential new targets for the development of anaphylaxis therapeutics if the same mechanism is responsible for anaphylaxis in humans.

Introduction
Anaphylaxis is a systemic hyperacute allergic reaction (1) responsible for more than 1,500 deaths per year in the US (2). Anaphylaxis is associated with intense vasodilatation and bronchoconstriction, severe laryngeal edema, drop of cardiac pressure, and hypothermia. As anaphylaxis is a life-threatening medical emergency, the mechanisms thought to be responsible for anaphylaxis have been mostly investigated in animal models. Two types of models have been developed since the initial description of anaphylaxis in dogs (3): active anaphylaxis, in immunized animals, and passive anaphylaxis, in nonimmunized animals injected with antibodies. Indeed, susceptibility to anaphylaxis can be transferred by serum from immunized donors or by purified antibodies. IgE-induced passive systemic anaphylaxis (PSA) is elicited by injecting mice systemically with IgE antibodies 24–48 hours before an i.v. challenge with specific antigen. The anaphylactic shock that develops within minutes can be easily assessed by monitoring the decrease in body temperature. IgE-induced PSA observed in WT mice was abrogated in mice deficient for FcεRI, the high-affinity IgE receptor expressed by mast cells and basophils (4), and in mast cell–deficient W/W mice (5). It was also abrogated in histidine decarboxylase–deficient mice, which lack histamine (6), and in mice injected with histamine receptor antagonists (7). Anaphylactic symptoms could be induced by an i.v. injection of histamine (6). These findings together demonstrate the mandatory role of mast cells and of FcεRI in IgE-induced PSA, and they emphasize the contribution of histamine, contained in mast cell granules that are rapidly released during exocytosis. This mechanism has been widely accepted as a paradigm of the anaphylactic reaction.

IgG-induced PSA is elicited by injecting mice systemically with IgG antibodies 2–3 hours before an i.v. challenge with specific antigen. Alternatively, preformed IgG immune complexes (IC) can be injected i.v. Similar symptoms develop, with comparable kinetics, during IgE- and IgG-induced PSA. IgG1 is the dominant antibody subclass raised during humoral responses to protein antigens in mice, and passively administered IgG1-IC are sufficient to induce anaphylaxis. Because the low-affinity IgG receptor FcγRIIIA was shown to trigger mast cell activation in vitro (8) and passive cutaneous anaphylaxis in vivo (9), it has been generally accepted that these receptors account for IgG1-induced PSA. No published paper formally demonstrated this assumption, but we confirmed that, indeed, IgG1-induced PSA was abrogated in FcγRIIIA-deficient mice (P. Bruhns and M. Daëron, unpublished observations). Surprisingly, IgG1-induced PSA was not abrogated in mast cell–deficient mice (5), but it was reported to be abrogated in basophil-depleted mice (10). This suggests that mouse basophils express FcγRIIIA. FcγRIIIA are also expressed by other myeloid cells. Upon activation, mouse basophils rapidly release granular mediators, including histamine, but also lipid-derived mediators such as platelet-activating factor (PAF). Like histamine, PAF could, by itself, reproduce the clinical signs of an anaphylactic shock when injected in animals (11). PAF, but not histamine, was shown to be responsible for IgG1-induced PSA (10). These findings together indicate that IgG1-IC trigger anaphylaxis through the release of PAF, probably by aggregating FcγRIIIA on basophils.

Active systemic anaphylaxis (ASA) is elicited by an i.v. injection of antigen into mice immunized with that antigen. Similar symptoms develop with comparable kinetics during ASA and PSA in WT mice. More mice, however, die during ASA than during PSA. Different adjuvants can be used for immunization. It is generally

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accepted that alum favors IgG1 and IgE antibodies, whereas CFA favors IgG2 antibodies. In both cases, however, IgG1 antibodies are the most abundant and IgE the least abundant. ASA was not affected in Ct-deficient mice, which produce no IgE (12). Antibodies other than IgE are therefore sufficient to induce ASA. Supporting this conclusion, ASA was not altered in FcRII-deficient mice, but it was abrogated in FcRγ-deficient mice (5), which express no activating receptors for IgE (FcεRI) or for IgG (FcγRI, FcγRIIIA, FcγRIV). Activating FcRs are therefore mandatory for ASA. ASA was not altered in mast cell-deficient W/Wv or W/Wv mice (5, 13), in basophil-deficient mice (14), or in basophil-depleted WT mice (10). Neither mast cells nor basophils are therefore mandatory for ASA. Furthermore, ASA-induced decrease in body temperature still occurred in the absence of both cell types, but ASA-associated mortality was abolished (10). Basophils and mast cells can therefore contribute to ASA. Reduced anaphylactic shock was observed in PAF receptor-deficient (PAF-R-deficient) mice (15) and in mice injected with PAF-R antagonists (16). These findings together indicate that antibodies other than IgE, activating FcRs other than FcεRI, cell types other than basophils and mast cells, and mediators other than histamine contribute to ASA. These antibodies, FcRs, cells, and mediators are unknown.

We unravel here an unexpected role of neutrophils and an under-estimated contribution of IgG and IgG receptors to anaphylaxis. Neutrophils and basophils both contributed to ASA in WT mice. Neutrophils, however, were sufficient for ASA in genetically modified mice expressing no activating IgG or IgE receptors on mast cells and basophils. Importantly, like murine neutrophils, human neutrophils restored ASA in ASA-resistant mice, suggesting that these cells can contribute to anaphylaxis in humans. Neutrophil-dependent ASA was mediated by PAF. Neutrophils contributed also to IgG-induced PSA. Finally, 2 IgG receptors, FcγRIIIA and FcγRIV, accounted for ASA in WT mice.

Results

**FcγRIV can trigger ASA.** An i.v. antigen challenge induced a decrease of body temperature in WT or in mast cell-deficient W/Wv mice immunized with antigen in CFA/incomplete Freund adjuvant (IFA), but not in FcRγ−/− (offically referred to as FcεRγ−/−) mice, which lack all activating IgG and IgE receptors (data not shown). ASA indeed depends on antibodies. Immunizations in CFA/IFA lead to the production of comparable IgG1 and IgG2 antibody levels in WT and FcRγ−/− mice and, as expected (5), of higher levels of IgE antibodies in FcRγ−/− mice than in WT mice (Supplemental Figure 1A; supplemental material available online with this article; doi:10.1172/JCI45232DS1). Non-immunized WT, but not FcRγ−/−, mice underwent anaphylaxis upon challenge with antigen if injected with serum from mice immunized with BSA in CFA/IFA (Supplemental Figure 1B).

ASA was not altered in FcεRIα/FcεRIα−/− (double-deficient (FcεRIIIA−/−) or in FcγRIα-deficient (FcγRIIA−/−) mice compared with WT mice (Figure 1A). Unexpectedly, ASA was also unaltered in FcεRI/FcγRIIB/FcγRIIIA/FcγRI/FcγRIV−/− (5KO) mice (Figure 1B and Supplemental Table 1). 5KO mice lack all IgE and IgG receptors except the activating IgG2 receptor FcγRI (17). Immunizations did not significantly modify the expression of FcγRIV (Supplemental Figure 1C). Anti-FcγRIV blocking mAbs abolished ASA in 5KO mice (Figure 1C). FcγRIV can therefore trigger anaphylaxis.

**Neutrophils mediate FcγRIV-dependent ASA and secrete PAF.** FcγRIV is expressed by monocytes/macrophages and neutrophils (17, 18), but not by basophils, mast cells, and eosinophils (Figure 1D).

Monocytes and macrophages can be depleted by injecting toxic liposomes in vivo (19). Both control and toxic liposomes had the same effect on ASA, reducing ASA-associated mortality without altering ASA-associated temperature drop (Figure 1E). Toxic liposomes, but not control liposomes, however, depleted monocytes (example shown in Supplemental Figure 1D). Depletion of monocytes/macrophages induced by toxic liposomes, therefore, did not alter ASA (Figure 1E). Supporting this result, ASA was unaffected in 5KO mice injected with gadolinium, which inhibits monocyte/macrophage function (Supplemental Figure 1E). Monocyte/macrophages are therefore not mandatory for ASA in 5KO mice. Noticeably, ASA-associated temperature drop and mortality were abrogated by neutrophil depletion in 5KO mice (Figure 1F). Anti-Gr1 injections depleted neutrophils, but did not reduce basophil numbers (Supplemental Figure 2, A and B). ASA was restored within 7 days in neutrophil-depleted mice, and it was correlated with blood neutrophil recovery (Figure 1G). Neutrophils, but not monocytes/macrophages, are therefore required for FcγRIV-dependent ASA. Importantly, purified bone marrow neutrophils from 5KO mice, but not from FcRγ−/− mice, restored ASA-associated temperature drop in BSA-immunized FcRγ−/− recipients (Figure 2A). Neutrophils are therefore necessary and sufficient for FcγRIV-dependent ASA. Mediators released and/or secreted by activated neutrophils should therefore be responsible for the anaphylactic shock observed. Among them, PAF (20–22) was shown to mimic anaphylaxis when injected in mice (11). We detected PAF in the supernatant of neutrophils purified from 5KO, but not from FcRγ−/− mice stimulated with IgG2b-IC in vitro. Anti-FcγRIV blocking mAbs abrogated PAF production by neutrophils from 5KO mice (Figure 2B). PAF may therefore be responsible for FcγRIV-induced ASA in 5KO mice.

**Neutrophils are immediately and systemically activated during active anaphylaxis.** Upon activation, blood neutrophils release granules that contain mediators and enzymes, including myeloperoxidase (MPO). MPO can be released within minutes by activated neutrophils (data not shown and ref. 20) and, under inflammatory conditions, by monocytes/macrophages (23). MPO can oxidize luminol, and oxidized luminol emits luminescence. Luminescence could, indeed, be detected in vivo in WT, but not in Mpo−/− mice injected with luminol and challenged with PMA or LPS (24). Immunized, but not naive, 5KO mice exhibited a massive, systemic luminescence emission following antigen challenge and luminol injection (Figure 2C and Supplemental Figure 2C). Luminescence was detectable within minutes after antigen challenge and lasted for at least 20 minutes (Figure 2D and Supplemental Figure 2D). These results suggest that neutrophils are systemically activated within minutes in ASA.

**FcγRIV, neutrophils, and PAF also contribute to PSA.** We investigated whether neutrophil depletion or FcγRIV blocking, which both affect ASA, may also affect an IgG-induced PSA in immunodeficient Rag2−/− 5KO mice. These mice, indeed, lack endogenous IgG that might compete with IC for FcγRIV binding. FcγRIV is indeed a high-affinity IgG receptor (17). An i.v. injection of monoclonal IgG2b-IC induced a significant, although nonlethal, temperature drop in Rag2−/− 5KO mice, but not in Rag2−/−FcγRγ−/− mice (Figure 3A). FcγRIV blockade or neutrophil depletion abolished monoclonal IgG2b-induced PSA in Rag2−/− 5KO mice (Figure 3B). FcγRIV can therefore induce IgG2b-induced PSA that depends on neutrophils. Immunocompetent 5KO mice also developed IgG2b-induced PSA (data not shown). Because neutrophils from 5KO mice secrete PAF
Figure 1
Neutrophils mediate FcγRIV-dependent active anaphylaxis. Indicated mice were immunized and challenged with BSA. Central temperatures and survival rates were monitored. (A) ASA in WT (n = 4), FcγRIIIA−/− (n = 5), and FcεRI/II−/− (n = 4) mice. (B) ASA in WT (n = 9) and 5KO mice (n = 7). (C) ASA in 5KO mice injected with anti-FcγRIV mAbs (n = 10) or isotype control (n = 11) before BSA challenge. (D) Representative expression of FcγRIV on mouse blood and peritoneal cells: B cells (CD19+), T cells (CD4+), monocytes (mono.) (CD11b+Gr1−), neutrophils (neutro.) (CD11b+Gr1+), basophils (baso.) (IgE+DX5+), eosinophils (eosin.) (Gr1+SiglecF+), macrophages (macro.) (CD11b+Gr1+), and mast cells (IgE+CD117+). (E and F) ASA in 5KO mice injected with (E) PBS liposomes (lipo.) (n = 7) or clodronate liposomes (n = 8), (F) anti-Gr1 mAbs (n = 8) or isotype (iso.) control (n = 9) before BSA challenge. Data are a compilation of 2 experiments. Note that both PBS and clodronate liposome injections inhibited ASA-associated mortality. (G) Immunized 5KO mice were injected with anti-Gr1 mAbs or isotype control on day 0, challenged with BSA on day +1, retroorbitally bled on days +2 and +6, but otherwise left untouched until rechallenged with BSA on day +7. Upper panel: representative density plots of blood leukocytes stained as indicated. Percentages of Gr1hiCD11bhi cells (neutrophils) are indicated. Lower panel: ASA in immunized 5KO mice at day +7 after depletion (Iso, n = 3; anti-Gr1, n = 4). (A–C and E–G) Data are represented as mean ± SEM. (A–F) Data are representative from at least 2 independent experiments (A, 2; B, 5; C, 4; D, 3; E, 2; and F, 4 experiments). ***P < 0.001. X’s represent mortality in the 100% experimental group.
when stimulated in vitro with IgG2b-IC (Figure 2B), we measured PAF in the plasma of 5KO mice undergoing neutrophil-dependent IgG2b-induced PSA. We detected elevated PAF levels in the plasma of challenged, but not in nonchallenged (no detectable PAF; not shown), 5KO mice (Figure 3C). FcγRIV blockade abolished the increase in circulating PAF levels. Elevated PAF levels in plasma therefore correlate with FcγRIV-triggered, neutrophil-dependent PSA. Surprisingly, FcγRIV blockade or neutrophil depletion abolished monoclonal IgG2b-induced PSA in WT mice also (Figure 3D). FcγRIV is therefore responsible for IgG2b-induced neutrophil-dependent PSA in WT mice.

A nonlethal PSA could also be observed in Rag2−/− 5KO mice injected with IC made of GPI and K/BxN serum (GPI/anti-GPI-induced PSA) (25, 26). Neutrophil depletion abolished this GPI/anti-GPI-induced PSA in Rag2−/− 5KO mice (Figure 3E). As expected, FcγRIV blockade only slightly reduced GPI/anti-GPI-induced PSA in WT mice. Neutrophil depletion, which express FcγRIIIA and FcγRIV, was sufficient to abolish GPI/anti-GPI-induced PSA in these mice (Figure 3F). Similar results were observed in 5KO mice (data not shown).

Whereas basophils have been reported to mediate monoclonal IgG1-induced PSA (10), neutrophil depletion, but not basophil depletion, abolished GPI/anti-GPI-induced PSA in the same experiment (Figure 3H). Similarly, whereas mast cells have been reported to contribute to monoclonal IgG1-induced PSA (5), mast cell-deficient W6/32 mice showed unaltered GPI/anti-GPI-induced PSA. Neutrophil depletion abolished GPI/anti-GPI-induced PSA in W6/32 mice as efficiently as in WT mice, whereas basophil depletion had no effect (Figure 3I). Together, these results demonstrate that neutrophils, but not mast cells or basophils, are mandatory for polyclonal IgG-induced PSA in nonimmunized WT mice.

FcγRIIIA and FcγRIV account for ASA in WT mice. We next investigated the contribution of activating Fc receptors to ASA in WT mice. WT mice express 1 activating IgE receptor, FcεRI, and 3 activating IgG receptors, FcγRI, FcγRIIIA, and FcγRIV. No anti-FcεRI- or anti-FcγRI-blocking antibodies are available. The recently described (27) mAb 275003, which is specific for FcγRIIIA (Figure 4A and Supplemental Figure 3, A and B), and which can block the binding of IgG-IC to FcγRIIIA in vitro (Supplemental Figure 3C), abolished FcγRIIIA-dependent (9) IgG-induced PSA (Figure 4B). FcγRIIIA blockade also significantly reduced ASA-associated temperature drop and mortality in WT mice (Figure 4, C and D). FcγRIV blockade reduced ASA-associated temperature drop similarly to FcγRIIIA blockade, but reduced mortality less efficiently (Figure 4D). Concomitant...
FcγRIIIA and FcγRIV blockade, however, were necessary to abolish ASA in WT mice (Figure 4, D and E). FcγRIIIA and FcγRIV therefore account for ASA in WT mice.

Neutrophils and basophils contribute to ASA in WT mice. FcγRIIIA and FcγRIV are both expressed by neutrophils, monocytes, and macrophages. FcγRIIIA, but not FcγRIV, is expressed also on basophils, eosinophils, and mast cells (Figure 5A and Figure 1D). All these cell types can potentially contribute to ASA. Inhibition of monocyte/macrophages (Figure 5B) or depletion of eosinophils (Supplemental Figure 3D) did not affect ASA in WT mice. Importantly, neutrophil depletion reduced temperature drop and prevented ASA-associated death in WT mice (Figure 5C). Basophil depletion (example shown in Supplemental Figure 3E) had no effect unless neutrophils were also depleted, suggesting a dominant role for neutrophils in ASA (Figure 5C). Similar results were obtained in mast cell–deficient Wsh mice (Figure 5D) and in FcγRIIIA−/− mice that express no activating FcγRs on mast cells and basophils (Figure 5E). ASA in WT mice is therefore unaffected by a deficiency in a single cell population, except that of neutrophils.

We observed that the proportion of blood neutrophils significantly increased following immunizations with antigen in CFA/IFA. Neutrophilia is not specific to this adjuvant, as it was also observed following immunization with alum or with alum plus Pertussis toxin (Figure 6A). This observation could possibly explain the dominant contribution of neutrophils to ASA following CFA/IFA immunizations. To test this hypothesis, we delayed the antigenic challenge for several weeks until neutrophil numbers dropped back to baseline. Six weeks after immunization, neutrophil numbers in 5KO mice immunized with BSA in CFA/IFA were comparable to neutrophil numbers in naive mice (Figure 6B), while the levels of anti-BSA antibodies in the serum remained high (Figure 6C). When challenged with BSA, these 5KO mice developed ASA, as expected, and ASA-associated temperature drop and mortality were both dependent on neutrophils.
The dominant role for neutrophils in ASA is therefore not due to higher neutrophil numbers in immunized mice. 

**PAF mediates neutrophil-dependent ASA.** Because neutrophils play a dominant role in ASA in SKO mice and in WT mice, because neutrophils secrete PAF in vitro (Figure 2B), and because PAF concentration is elevated in plasma during neutrophil-dependent PSA (Figure 3C), we analyzed the contribution of PAF to ASA in 5KO and in WT mice. Two PAF-R antagonists (Figure 7A), but not a histamine receptor-1 antagonist (Figure 7B), markedly reduced ASA in 5KO mice. PAF-R antagonists also abolished ASA-associated death and inhibited temperature drop in WT mice, whereas histamine receptor-1 antagonist had a much milder effect (Figure 7C).
PAF production requires cytosolic phospholipase A2 (cPLA2) (28–30). ASA was strongly inhibited in cPLA2-deficient mice (Figure 7D). PAF may therefore account for neutrophil-dependent ASA in 5KO mice and contributes to ASA in WT mice.

Human neutrophils restore ASA in FcγRII–/– mice. Like murine neutrophils (Supplemental Figure 4A), human neutrophils express FcRs. The only activating FcRs expressed by neutrophils from normal donors are FcγRIIA (Figure 8A). Human neutrophils,
indeed, do not express FcγRI, FcεRI, or FcγRIIIA. They, however, express the GPI-anchored FcγRIIIB (31, 32). FcγRIIA, but not FcγRIIIB, could bind IC made with mouse IgG1, IgG2a, or IgG2b (Figure 8B). Human neutrophils could be activated in vitro by IC made with mouse polyclonal IgG (Figure 8C) or mouse monoclonal IgG1 or IgG2b (Supplemental Figure 4B), as could murine neutrophils from WT and 5KO, but not from FeRγ−/−, mice (Supplemental Figure 4C). Neutrophils purified from normal human donors could restore ASA when injected i.v. into immunized FeRγ−/− mice prior to antigen challenge (Figure 8, D–F). Importantly, the same neutrophils purified from individual donors induced no anaphylaxis upon antigen challenge in nonimmunized mice (Figure 8, E and F). The intensity of the shock was proportional to the number of human neutrophils transferred into FeRγ−/− mice. Human neutrophils can therefore substitute for murine neutrophils in ASA.

Discussion

We show here that IgG antibodies, activating FcγRs, neutrophils, and PAF are the main players in ASA, whereas IgE antibodies (12), FcεRI (5), mast cells (5, 13), and histamine do not play a major role. Several cell types have been involved in anaphylaxis; among them are mast cells and basophils. We demonstrate that neutrophils are not only sufficient to induce ASA, but play a dominant role. Indeed, neutrophil depletion, but not basophil depletion, eosinophil depletion, monocyte/macrophage inhibition or mast cell deficiency, abrogated ASA-associated death, and reduced temperature drop in WT mice. Importantly, transfer of mouse neu-
trophils expressing FcγRIIV, but not of neutrophils expressing no activating FcγRs, restored ASA in FcγR−/− mice. Neutrophil activation could be visualized in vivo within minutes in mice undergoing ASA. Although monocytes/macrophages were reported to be involved in a model of anaphylaxis induced by an i.v. injection of goat IgG in mice immunized with goat IgG anti-mouse IgD (33), we could not detect any role for monocytes/macrophages in ASA in mice immunized with antigen in CFA/IFA. The depletion of eosinophils did not impair ASA either. Basophil depletion, which did not affect ASA by itself, further reduced ASA when combined with neutrophil depletion. Basophils therefore contribute to ASA together with neutrophils, but neutrophils play a dominant role. One possible explanation is that neutrophils are much more numerous than basophils in blood.

Noticeably, we observed neutrophilia following immunization of mice with antigen in CFA/IFA. We excluded that neutrophilia accounted for the dominant contribution of neutrophils to ASA by delaying antigen challenge until neutrophil numbers in immunized mice were comparable to those in naive mice. Under these conditions, neutrophils were also mandatory for ASA in SKO mice. We found that neutrophils also predominantly contributed to IgG-induced PSA (whether induced by monoclonal IgG2b-IC or polyclonal-IC) in nonimmunized WT mice that have normal numbers of neutrophils. We also observed neutrophilia following immunizations with alum or with alum plus Pertussis toxin. In preliminary experiments, neither neutrophil depletion nor basophil depletion significantly reduced temperature drop, but a depletion of both cell types abolished ASA in mice immunized with antigen in alum (our unpublished observations). Taken together, our results demonstrate that neutrophils contribute to anaphylactic shock in 2 models of ASA, i.e., following immunization in CFA/IFA or in alum, and in 2 models of PSA, i.e., induced by polyclonal IgG-IC or monoclonal IgG2b-IC.

Two mediators, PAF and histamine, were found to play a critical role in experimental anaphylaxis. Mast cells, basophils and, apparently, neutrophils (34) can release histamine. Histamine accounts for IgE-induced PSA but not for IgG1-induced PSA, whereas PAF accounts for IgG1-induced PSA but not IgE-induced PSA (10). We found that PAF has a dominant role in ASA. Indeed, PAF-R antagonists, but not histamine receptor H1 antagonists, markedly reduced temperature drop and prevented death in ASA in mice immunized with antigen in CFA/IFA. In agreement with these results, ASA-associated heart rate and arterial pressure reduction were strongly impaired in PAF-R-deficient mice (15), suggesting that ASA-associated temperature drop and mortality may also be inhibited in these mice. Indirectly supporting this assumption,
ASA was virtually abrogated in cPLA2-deficient mice (our results and ref. 29), which cannot generate several lipid-derived mediators, including PAF. Activated neutrophils (20, 22), monocytes/macrophages (35), and eosinophils (36) all produce PAF, but neutrophils were reported to be major producers (21). We found, indeed, that neutrophils secrete PAF when stimulated by IgG2b-IC, but we also found elevated PAF levels in plasma of mice undergoing neutrophil-dependent PSA. Together our results suggest that PAF released upon neutrophil activation during ASA and PSA is responsible for anaphylactic shock.

ASA depends on activating receptors (5) that associate with and whose expression and signaling depend on FcRγ. These are FcγRI, FcγRIIa, FcγRIIb, and FcγRI (37). ASA was reported in IgE-deficient mice (12), and we observed that it was comparable in WT and FcγRIIa-deficient mice. Similarly, the deletion of FcγRI did not affect ASA in mice immunized with antigen in aluim (5, 38). While blocking either FcγRIIa or FcγRI reduced ASA, blocking both receptors abrogated ASA in WT mice. IgG antibodies seem therefore more important than IgE antibodies in ASA following immunization in CFA/IFA. FcγRII contributes to ASA in WT mice, and it was necessary and sufficient to induce ASA in 5KO mice. A single activating FcγR on a single cell population is therefore sufficient to induce ASA. Our results demonstrate that FcγRIIa or FcγRI is responsible for ASA following CFA/IFA immunizations, and that each could substitute for the other to induce ASA, provided that IgG2 antibodies are produced.

That IgG1 can induce anaphylaxis was demonstrated by PSA. The only activating FcγR having an affinity for IgG1 is FcγRIIa (17, 18). Depletion of basophils using specific mAbs abrogated IgG1-induced PSA (10), although mast cells and neutrophils also express FcγRIIa. A model of basophil-deficient mice, however, challenges this result (14). Mast cells are necessary for IgE-induced ASA, provided that IgG2 antibodies are produced.

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We found, however, that not only IgG1, but also IgG2, antibo-
dies could induce PSA. IgG2-induced PSA could develop in SKO
mice, and neutrophils contributed to the shock. IgG2 can there-
fore contribute to anaphylaxis. IgG2-IC that may form in vivo
upon antigen challenge following immunization in CFA/IFA may
be responsible for the predominant role of neutrophils as IgG2
IC can engage FcyRIV (but also FcyRIIIA) on these cells. In the
absence of IgG2 antibodies, as in ASA following immunization
with antigen in alum, IgG1-IC may trigger FcyRIIIA-expressing
basophils and neutrophils. It follows that basophils contribute to
ASA and to PSA when IgG1-IC are present (this report and ref.
10), but also other cells (14), among which are neutrophils. Our
results indicating that neutrophils are mandatory for polyclonal
IgG-induced PSA differ from previous reports implicating mast
cells (5) and basophils (10) in IgG1-induced PSA. They are, how-
ever, not contradictory. Indeed, monoclonal IgG1-IC can selectively
engage FcyRIIIA, while IgG1-IC made of IgG1 and IgG2 isotypes
will engage both FcyRIIIA and FcyRIV, to induce PSA. FcR-express-
ing cells involved in each type of PSA may therefore be different.
Taken together, these data suggest that IgE, IgG1, and IgG2 can all
induce anaphylaxis when engaging FcεRI, FcγRIIIA, and FcγRIV on
mast cells, basophils, and neutrophils, respectively. Although like-
ably, it was not formally demonstrated that FcεRI or FcγRIIIA alone
could induce ASA. Using the SKO model, we provide evidence that
FcγRIV alone can. Because FcγRIV is a receptor for IgG2, but not
for IgG1, and is expressed by neutrophils, but not by basophils
or mast cells, we are able to demonstrate here a role for the most
unexpected antibody, receptor, and cell type in ASA in mice.

Human anaphylaxis is essentially active. IgG1 is the most abun-
dant IgG subclass in human plasma, and the majority of antibo-
dies raised by vaccinations (generally in alum) belong to this sub-
class. Human IgG1 binds to all activating human FcγRs (39). Both
specific IgE and IgG antibodies are found in the serum of aller-
genic patients, but the relative concentration of the various classes
is poorly known and rarely investigated. FcγRIV has no human
ortholog, and FcγRIIIA are not expressed by human neutrophils,
basophils, and mast cells (40). These 3 cell types, nevertheless,
express activating Fc receptors. All of them express another activate-
ing IgG receptor, FcγRIIA. Mast cells and basophils also express the
high-affinity IgE receptor FcεRI in normal individuals as well as
neutrophils in atopic patients (41). These 3 cell types can produce
PAF upon activation, especially neutrophils (21). PAF could play
an important role in human anaphylaxis. Indeed, plasma PAF concen-
tration has been correlated with the severity of shocks in patients
(42). Although the cellular source of PAF in human anaphylaxis
was not identified, the above-mentioned results endow neu-
rophils with a critical role. Supporting this assumption, IgG-IC
could activate human neutrophils in vitro, and a transfer of human
neutrophils restored ASA in Fcer1-deficient mice.

In conclusion, we demonstrate here a previously unexpected role
of neutrophils in anaphylaxis. An IgG2-induced, FcγRIV-depen-
dent, PAF-mediated active anaphylaxis, contingent on neutro-
phils, could be unraveled using immunized multiple Fcε-deficient
mice. IgG-induced, FcγR-dependent, PAF-mediated ASA was also
observed in WT mice, to which neutrophils contributed. Anaphy-
laxis induction is therefore a property of neutrophils, which is not
trivial considering that neutrophils are the most numerous cells
among blood leukocytes in humans. This may have important
therapeutic consequences if indeed neutrophils can induce IgG-
dependent anaphylactic reactions in humans.

Methods
Mice. SKO (N6 B6) mice have been described (17). cPla2γ−/− mice (129/B6)
were provided by J. Bonventre (Brigham and Women’s Hospital, Harvard
Institutes of Medicine, Boston, Massachusetts, USA); KRN transgenic
mice were provided by D. Mathis and C. Benoist (Harvard Medical
School, Boston, Massachusetts, USA) and the Institut de Génétique et de
Biologie Moléculaire et Cellulaire (Illkirch, France). WT C57BL/6J and
NOD mice were purchased from Charles River, and with Wbb/Wbb, FcγRIIIA−/−
and FcyRγ−/− C57BL/6J mice from Jackson Laboratories. Rag2−/− mice were used to
generate Rag2−/− SKO and Rag2−/− FcγR−/− mice by intercrosses. All
mouse protocols were approved by the Animal Care and Use Committees
of Paris, Île de France, France.

Antibodies, reagents, and cells. PBS- and clodronate-liposomes were prepared
as previously described (19). Alum hydroxide gel, pertussis toxin, BSA,
DNP-HSA, rabbit GPI, gadolinium(III) chloride, CFA and IFA, CV-3988,
ABT-491, cyproheptadine, and luminol were obtained from Sigma-Aldrich;
anti-mouse CD11b, CD4, CD19, Gr1, SiglecF, CD117, IgE, and anti-
human FcεRI, FcγRII, CD62L were from BD Biosciences; anti-mouse DX5,
mFcer1, and anti-hFcer1 were from eBioscience; anti-mFcRII (290322)
and mFcgRIIIA (275003) were from R&D Systems; anti-hFcgRIIIA mAbs
(IV.3) were from StemCell Technologies; mouse IgG3 anti-DNP were from
Serotec; and the MPO ELISA kit was from HyCult Biotech. The hybrid-
omas producing mAbs anti-mFcγRIV (9E9) were provided by J.V. Ravetch
(Rockefeller University, New York, New York, USA); anti-Gr1 (RB6-8C5)
was provided by R. Coffman (DNAX Research Institute, Palo Alto, Califor-
nia, USA); mlgG2a anti-platelet (6A6) was provided by R. Good (University
of South Florida College of Medicine, Tampa, Florida, USA); and mlgG1
and mlgG2b anti-DNP was provided by B. Heyman (Uppsala Universitet,
Uppsala, Sweden). Purified mAbs anti-hFcγRIIB/C (GB3) were provided
by U. Jacob (SuppreMol GmbH); anti-CCR3 was provided by J.J. Lee (Mayo
Clinic, Scottsdale, Arizona, USA). Anti-CD200R3 (Ba103) was produced as
described (10). CHO K1 cells stably transfected with FLAG-tagged mouse
FcγRs (17) or human FLAG-tagged FcγRs (39) were cultured as described.

Flow cytometry analysis. Cells were stained with indicated fluorescently
labeled mAbs for 30 minutes at 0°C.

IC binding. Mouse IC were preformed by incubating 10 pg/ml DNP-DNP-
BSA-biotin with 15 μg/ml anti-DNP mAbs for 1 hour at 37°C. 2 x 106 cells,
preincubated or not with indicated blocking mAbs, were incubated with IC
for 2 hours at 4°C, which were detected using neutravidin-PE at 2 μg/ml
for 30 minutes at 4°C.

ASA. 6- to 9-week-old male mice were used for ASA. They were injected
i.p. at days 0, 14, and 28 with 200 μg BSA, either once in CFA and twice in
IFA, or 3 times in alum, or 3 times in alum plus Pertussis toxin. BSA-specific
IgG1 and IgG2a/b/c antibodies in serum were titered by ELISA at day 30.
Briefly, BSA-coated plates were sequentially incubated with dilutions of
serum, HRP-labeled anti-Ig isotype antibodies, and SIGMAFAST OPD
solution. Internal negative (serum from naïve C57BL/6J mice) and positive
(pool of serum of BSA-immunized C57BL/6J mice) controls were added to
all test plates to define a background value and a “value 1” in arbitrary units,
respectively. High titers of IgG1 antibodies were found in mice immunized
using either CFA/IFA or alum immunization protocol. IgG2 antibodies were
detected in mice immunized using CFA and IFA only. Absolute counts and
proportions of granulocytes in peripheral mouse blood were determined
using an ABC Vet automatic blood analyzer (Horiba ABX).

Mice with comparable antibody titers were challenged i.v. with 500 μg BSA
(unless otherwise specified) 10 days after the last immunization. Note that
the amount of antigen that induced ASA (Supplemental Figure 5A) was sim-
ilar to that reported to be required for IgG-induced PSA (43). Central tem-
perature was monitored using a digital thermometer (YSI), and time of death
was recorded. Except in Figure 1, E and F, and in Figure 2A, data presented in
each figure correspond to 1 representative experiment. Supplemental Table 1 lists the total number of mice that were used for indicated experimental conditions, their breeds of anaphylactic shock, and the mortality rate.

**PSA.** Mice were injected i.v. with preformed IC made of 20 μl K/BxN serum and 50 μg GPI (Figure 4B); 500 μg IgG2b mAb anti-DNP and 200 μg DNP-HSA (Figure 3, A, B, and D); and IC made of 100 μl K/BxN serum and 40 μg GPI (Figure 3, E–I). Alternatively (Supplemental Figure 1B), 500 μl of serum from BSA-immunized mice collected on day 30 was injected i.v. into naive mice and mice were challenged 3 hours later with an i.v. injection of 500 μg BSA. Central body temperature was recorded using a digital thermometer (YSI).

**Experimental immune thrombocytopenia.** Blood samples were taken retroorbitally before and at indicated time points after the i.v. injection of 5 μg of anti-platelet mAb 6A6. Platelet counts were determined using an ABC Vet automatic blood analyzer (Horiba ABX).

**In vivo blocking and depletion.** 200 μg/mouse of anti-FcγRIV mAbs were injected i.v. 30 minutes before challenge. 50 μg/mouse of anti-FcγRIIA mAbs, 2.1 mg/mouse of PBS- or clodronate-liposomes, 300 μg/mouse of anti-Gr1 or anti-CCR3 mAbs, and 30 μg/mouse of anti-CD200R3 mAbs or 1 mg/mouse of gadolinium were injected 24 hours before challenge. 66 μg/mouse CV-3988, 25 μg/mouse ABT-49, or 30 μg/mouse cyprophetamine was injected i.v. 10, 20, or i.p. 30 minutes before challenge, respectively.

Depletion of cell populations was ascertained for specificity of the depletion using flow cytometry on blood samples taken during or after the experiment (as represented in the figures, or in examples shown in Supplemental Figure 1D, Supplemental Figure 2, A and B, and Supplemental Figure 3, D and E). Depletion of specific cell populations or efficiency of blocking antibodies was also controlled using the macrophage-dependent experimental immune thrombocytopenia (EITP) model: ITP in 5K0 mice was inhibited by FcγRIV blockade (Supplemental Figure 5B), as expected (18), was inhibited by monocye/macrophage depletion using toxic liposomes (Supplemental Figure 5C), as expected (44), and was unaltered by neutrophil depletion (Supplemental Figure 5D).

**Bioluminescence imaging.** Bioluminescence from depleted and anesthetized mice injected i.p. with 15 mg/mouse luminol was acquired on an IVIS fluorescence imaging system. Mice were injected i.v. with preformed IC made of 200 μg DNP-HSA (Figure 3, A, B, and D); and IC made of 100 μg/ml of anti-FcγRIV and 1/100 dilution of K/xB/N serum) or neutrophils with plate-bound IC (DNP-HSA and indicated mouse anti-DNP antibodies).

**Statistics.** Data were analyzed using 1-way ANOVA with Bonferroni’s post-test (Figure 6A, Figure 8C, and Supplemental Figure 4B) or 2-tailed Student’s t test (all other data). P < 0.05 was considered significant.

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