Increased Severity of Local and Systemic Anaphylactic Reactions in gp49B1-deficient Mice

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Abstract

Gp49B1 is an immunoglobulin (Ig) superfamily member that inhibits FceRI-induced mast cell activation when the two receptors are coligated with antibodies in vitro. The critical question of in vivo function of gp49B1 is now addressed in gene-disrupted mice. gp49B1-deficient mice exhibited a significantly increased sensitivity to IgE-dependent passive cutaneous anaphylaxis as assessed by greater tissue swelling and mast cell degranulation in situ. Importantly, by the same criteria, the absence of gp49B1 also resulted in a lower threshold for antigen challenge in active cutaneous anaphylaxis, in which the antigen-specific antibody levels were comparable in gp49B1-deficient and sufficient mice. Moreover, the absence of gp49B1 resulted in a significantly greater and faster death rate in active systemic anaphylaxis. These results indicate that gp49B1 innately dampens adaptive immediate hypersensitivity responses by suppressing mast cell activation in vivo. In addition, this study provides a new concept and target for regulation of allergic disease susceptibility and severity.

Key words: mast cells • immunoreceptor tyrosine-based inhibitory motif • passive cutaneous anaphylaxis • active cutaneous anaphylaxis • active systemic anaphylaxis

Introduction

Gp49B1 is a member of the Ig superfamily with two extracellular C2-type domains (1, 2) and two cytosolic immunoreceptor tyrosine-based inhibitory motifs (ITIMs) (3). This transmembrane receptor is expressed on mast cells (4), macrophages (5, 6), and cytokine- or virally stimulated NK cells (7–9). Gp49B1 is encoded by the gp49B gene (2), whereas the highly homologous gp49A gene (10) encodes the gp49A protein that does not have an ITIM (1) and, like gp49B1, is expressed in higher amounts on immature bone marrow–derived mast cell (BMMC) progenitors than on mature peritoneal mast cells (4, 10). Ab-mediated coligation of gp49B1 to FceRI inhibits IgE-induced mast cell activation in vitro (3) by a mechanism requiring the ITIM tyrosines of gp49B1 and the src homology domain 2–containing phosphatase (SHP)-1 (11).

The ITIM was first identified in FcγRIIB (12, 13), whose extracellular domain is only distantly related to gp49B1 (3). FcγRIIB also differs because its ITIM inhibits cell activation through the recruitment of a src homology domain 2–containing inositol polyphosphate 5-phosphatase (SHP-1) (14). FcγRIIB is widely expressed in the immune system, including on B cells and mast cells, and inhibits cell activation signaled by the B cell receptor and FcγRI in vitro (12, 13). FcγRIIB-deficient mice exhibit increased vascular permeability in IgG-dependent passive cutaneous anaphylaxis (PCA) (15) and a higher death rate during IgG-or IgE-dependent passive systemic anaphylaxis compared with normal mice (16). Because both IgG- and IgE-antigen complexes bind FcγRIIB (17), the increase in sensitivity of the deficient mice to anaphylactic reactions reflects the loss of direct feedback counterregulation during adaptive immediate hypersensitivity reactions.

The gp49 family is homologous with other families of activating and inhibitory receptors of the Ig superfamily that have C-2 type domains and are expressed on various cell types in the immune system. The inhibitory members of the human killer cell Ig-like receptor (KIR), leukocyte Ig-like receptor (LIR), and mouse paired Ig-like receptor (PIR) families share with gp49B1 the expression of ITIMs that recruit SHP-1 (for reviews, see references 18–20). The ligands for KIRs and two of the LIRs are polymorphic MHC class I molecules, and the binding of autologous class I by the receptors provides self-recognition that counter-
regulates inappropriate activation signals. The ligands for the PIRs and most of the LIRs are unknown.

We recently identified the integrin αvβ3 as a functional ligand for gp49B1 in vitro (21). αvβ3 is widely distributed in vivo, being expressed on endothelial cells, fibroblasts, epithelial cells, and certain hematopoietic lineage cells (for a review, see reference 22). However, before addressing the ligand in vivo, it was critical to determine whether the absence of gp49B1 provided a mast cell phenotype with increased sensitivity to FcεRI-mediated, antigen-specific activation. We therefore generated mice with a targeted disruption of the gp49B gene and focused on the effects of gp49B1 deficiency on local and systemic anaphylactic reactions that have a major mast cell component. In PCA and with defined IgE inputs, gp49B−/− mice showed markedly greater and prolonged tissue swelling concomitant with a more extensive degranulation of tissue mast cells compared with gp49B+/+ mice. The increase in tissue swelling and the accelerated and greater rate of anaphylactic death in active cutaneous and systemic anaphylaxis, respectively, in gp49B−deficient mice, establishes for the first time the capacity of an ITIM-bearing receptor to attenuate active immediate hypersensitivity reactions. Thus, our results recognize a hyperresponsive mast cell phenotype in gp49B−/− mice, which in normal mice is dynamically counterregulated by the interaction of gp49B1 with innate tissue elements.

Materials and Methods

Generation of gp49B−/− Mice. A 6.4-kb 129/Sv EcoRI→HindIII genomic fragment containing the gp49B gene was subcloned into a modified pBluescript vector that had a thymidine kinase gene on the 3′ end. The region between the NcoI sites in intron 2 and exon 3 of the gp49B gene (2) was excised and replaced with the pMC1neo vector in the forward orientation (see Fig. 1 A). The targeting construct was transfected by electroporation into 129/Sv embryonic stem (ES) cells, and cells were selected with G418 and gancyclovir. To confirm homologous recombination, genomic DNA was extracted from the drug-selected cells, digested with SphI, and hybridized with a HindIII probe common to the 3′ regions of the gp49A and gp49B genes (see Fig. 1 A). The probe was predicted to hybridize with a 4.8-kb band reflecting the disrupted gp49B gene and with an ~9.5-kb band from the wild-type gp49B gene. All the clones had a 3.6-kb fragment from the wild-type gp49A gene, which has an SphI site in intron 5 that is not present in the gp49B gene (10). When the same blots were hybridized with an XhoI→SalI pMC1Neo probe, the clones that showed disruption of the gp49B gene with the HindIII→SphI probe demonstrated ~5.8- and 4.8-kb bands consistent with the gene disruption (data not shown). Moreover, there were no bands indicative of unintentional disruption of the gp49A gene. One of the clones with a disrupted gp49B gene (no. 69) was injected into BALB/c blastocysts, and chimeric mice were generated and crossed with BALB/c mice to produce male and female gp49B−/− mice. Those mice were intercrossed to yield gp49B+/+, gp49B−/−, and gp49B+/− mice (see Fig. 1 B). The homozygous mice on the 129/Sv × BALB/c background were then bred to yield gp49B+/+ and gp49B−/− mice for study. All animal studies were approved by the Animal Care and Use Committee of the Dana-Farber Cancer Institute.

Culture of BMMC. Bone marrow cells from gp49B+/+ and gp49B−/− mice were cultured for 3–8 wk in enriched medium containing 50% WEHI-3 cell (American Type Culture Collection) conditioned medium as described (3). After 3 wk, >97% of the nonadherent cells in the cultures were mast cells as assessed by metachromatic staining with toluidine blue.

Flow Cytometry. Cells (2.5 × 106) were incubated in 25 μl of calcium- and magnesium-free HBSS supplemented with 0.1% BSA and 0.02% sodium azide (HBA) alone or with saturating amounts of either rat mAb B23.1 (3) or rabbit anti-gp49B (20,21) (10) for 30 min at 4°C. Cells were washed with HBA, incubated for 30 min at 4°C with FITC-labeled, F(ab′)2 fragments of either goat IgG anti-rat IgM or goat anti-rabbit IgG-Fc (Jackson Immunoresearch Laboratories) as appropriate for the first Ab, and washed again with HBA. Cells were resuspended in 2% paraformaldehyde in HBSS and analyzed on a Becton Dickinson FACSort™ using logarithmic fluorescence amplification.

PCA. The left and right ears of mice were injected intradermally with 20 μl of saline alone or containing 25 or 2.5 ng of mouse monoclonal IgE anti-DNP (clone SPE-7; Sigma-Aldrich), respectively. 20 h later, the mice were challenged with a 100 μl intravenous injection containing 100 μg of DNP-human serum albumin (HSA; Sigma-Aldrich), and ear thickness measurements were made with a caliper (Dyer Company). Net ear swelling was calculated as the difference between the thicknesses of the right and left ears of each mouse.

Active Cutaneous and Systemic Anaphylaxis. Mice were immunized intraperitonally on days 0 and 5 with 10 μg of chicken OVA adsorbed to 1 mg of alum. Approximately 3 wk after the initial injection, the left and right ears of mice were injected intradermally with 20 μl of saline alone or containing 50 ng OVA, respectively. Net ear swelling was calculated as for PCA. For systemic anaphylaxis, immunized mice were injected intravenously with 10 μg of OVA in 100 μl of PBS. The outcome was quantified by the number of animals dying and the time to death.

Histology. Ears were excised, fixed in 4% paraformaldehyde for at least 8 h, and embedded in glycolmethacrylate. Sections (2 μm) were placed on slides, air dried, and stained for 20 s in a 5% ethanolic solution of toluidine blue or in Diff-Quik (Dade Behring, Inc.). For enumeration of intact mast cells, 10 high power fields (50× objective) of each ear were examined on a Leica Di- alux 20 microscope.

Statistical Analyses. Significance was assessed with Student’s unpaired, two-tailed t test. P values < 0.05 were considered to be significant.

Results and Discussion

Generation of gp49B−/− Mice. ES cells (129/Sv strain) were transfected with a targeting construct that had a pMC1Neo construct replacing a region of the gp49B gene running from intron 2 through most of exon 3 (that encodes the NH2-terminal Ig-like domain; Fig. 1 A). Of 104 G418- and gancyclovir-resistant clones, 2 had a disrupted gp49B gene by Southern blot analyses. A clone was microinjected into blastocysts from BALB/c mice to generate chimeric mice, which were bred to BALB/c mice to produce heterozygotes that were then intercrossed to yield gp49B+/+, gp49B−/−, and gp49B+/− mice (Fig. 1 B). gp49B−/− mice were fertile, had litters of normal size, and remained healthy when living in a viral Ab-free, barrier vi-
The expression of gp49B1 and gp49A proteins was assessed on BMMCs from gp49B+/+ and gp49B−/− mice. The gp49B1-specific mAb B23.1 (3) bound to BMMCs generated from gp49B+/+ but not gp49B−/− mice (Fig. 1 C). gp49A-specific anti-gp49A201–216 (10) bound equivalently to BMMCs from gp49B+/+ and gp49B−/− mice. In addition, peritoneal mast cells from gp49B−/− mice did not bind mAb B23.1, as assessed by two-color flow cytometry using anti-c-kit to identify mast cells (data not shown). Hence, the gp49B gene product was specifically absent in cells from gp49B−/− mice. These results also confirm our previous conclusion, based on flow cytometric analyses of gp49A and gp49B transfectants, that mAb B23.1 is gp49B1-specific (3).

**Effects of gp49B1 Deficiency on PCA.** To determine whether gp49B1 regulates FcεRI-mediated mast cell activation in vivo, we tested the effects of gp49B1 deficiency on the IgE-mediated PCA reaction in the ear. Passive sensitization with a defined input of IgE assures that subsequent antigen challenge is directed to the mast cells (24) and avoids the variation in titers of Ab classes that can occur with active sensitization. The left and right ears of mice were injected intradermally with saline or monoclonal IgE anti-DNP, respectively. 20 h later, the mice were challenged with an intravenous injection of DNP-HSA (100 μg), and initial measurements were made 0.5 h later, a time that falls within the 15 to 45 min plateau of the peak response (25). The net swelling in the ears of gp49B+/+ mice declined progressively from the 0.5 h time point and was essentially undetectable by 4 h (Fig. 2 A). The sensitized ears of gp49B−/− mice showed two- to threefold greater net swelling at 0.5 to 2 h after challenge compared with gp49B+/+ mice, and also declined progressively from the 0.5 h peak response with barely detectable swelling at 4 and 8 h. Enhanced swelling in the ears of the deficient mice was also evident histologically 0.5 h after challenge and was due to separation of the structural elements by edema, which had largely subsided in both genotypes by 4 h (Fig. 2 B). There was no influx of neutrophils, eosinophils, or other cell types evident within the time frame for the net tissue swelling. Importantly, the number of morphologically intact mast cells, defined as cells with metachromatic granules associated with a nucleus, was a significant twofold lower at both 0.5 and 4 h after challenge of gp49B−/− mice (Fig. 2 C), indicative of an enhanced susceptibility to mast cell degranulation in the deficient animals.

To determine whether this augmented mast cell response reflected a reduced requirement for the concentration of sensitizing IgE, the PCA reaction was conducted with an

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**Figure 1.** Generation of gp49B−/− mice. (A) Genomic organization of the mouse gp49A and gp49B genes (top), structure of the targeting vector (middle), and organization of the disrupted allele. The expected fragment sizes of 3.6, ~9.5, and 4.8 kb for the wild-type (WT) gp49A, WT gp49B, and disrupted gp49B genes, respectively, in the blot analysis in B are indicated by brackets. E, EcoRI; N, NcoI; Sp, SphI; H, HindIII; Δ3, truncated exon 3. (B) DNA blot analysis of SphI-digested genomic DNA identifying gp49B+/+, gp49B−/−, and gp49B−/− mice by hybridization with a HindIII→SphI gp49A/gp49B1 genomic fragment to provide the fragment sizes defined in panel A. (C) Flow cytometric analysis of BMMCs from gp49B+/+ and gp49B−/− mice stained with mAb B23.1 (anti-gp49B1; top panels, thick lines) or polyclonal anti-gp49B1 (anti-gp49A; bottom panels, thick lines) and the respective negative controls (thin lines).
intradermal injection of 2.5 ng of IgE, a dose that is 10- to 100-fold lower than is customarily used (24, 26). In gp49B1−/− mice, maximal swelling at 0.5 h was one-half that obtained with 25 ng of IgE and was undetectable beyond 1 h (Fig. 2 D). The ear swelling in gp49B1−/− mice was also lower than that observed with the higher IgE dose, but was dramatically greater than that of gp49B1+/+ mice at the low IgE dose (Fig. 2 D). Furthermore, the magnitude of the ear swelling in gp49B1−/− mice with low dose IgE sensitzation was equivalent to that of gp49B1+/+ mice sensitized with 10-fold more IgE. Thus, the gp49B1−/− mouse has an elicited phenotype of enhanced susceptibility to IgE-mediated mast cell activation over a 10-fold range of IgE sensitizing concentrations in situ. Inasmuch as the rapid and edematous tissue swelling in PCA is mast cell dependent (24), our findings establish conclusively that gp49B1 is an endogenous inhibitor of IgE-mediated mast cell activation in vivo. The observation of others that IgE-initiated degranulation of BMMC from gp49B1−/− and gp49B1+/+ mice is comparable in vitro (23) could be explained by the absence of a tissue ligand for gp49B1 in those cell suspensions.

Recently, mice deficient in leukotriene (LT)C4 synthase were found to exhibit attenuated ear swelling in IgE-dependent PCA (25), revealing a previously unappreciated contribution of mast cell–derived LTC4 and its receptor-active metabolites in immediate tissue swelling. PCA with 50 ng IgE was attenuated in LT C4 synthase-deficient mice by 50% compared with normal littermates (25), indicating that the cysteinyl leukotrienes are as significant as the amine mediators exocytosed from mast cell secretory granules in terms of the induction of early edema. Because coligation of gp49B1 with cross-linked FcεRI inhibits the generation of LTC4 by mast cells in vitro (3), it seems likely that the augmented production of cysteinyl leukotrienes and/or their synergistic action with the released amines contributes to the increase in PCA-associated tissue swelling in gp49B1−/− mice.

**Effects of gp49B1-deficiency on Acute Active Cutaneous Anaphylaxis.** To more fully assess the biologic relevance of gp49B1 in the negative regulation of immediate hypersensitivity, we used a model of active cutaneous anaphylaxis. OVA-immunized gp49B1+/+ mice that were challenged intradermally in the ear with 50 ng of antigen demonstrated little tissue swelling as measured by micrometry (Fig. 3 A) or histology (Fig. 3 B), despite an antigen-specific Ab response (Table I). In striking contrast, gp49B1−/− mice with a
comparable Ab response showed robust swelling at 0.5 h that was somewhat greater than that obtained by PCA with 25 ng of hapten-specific IgE (Fig. 2 A), and which consisted of edema without a cellular infiltrate (Fig. 3 B). The challenged ears of $gp49B^{-/-}$ mice showed fourfold fewer intact mast cells 0.5 h after challenge compared with $gp49B^{+/+}$ mice as noted histologically (Fig. 3 C) and quantified by cell counting (Fig. 3 D).

Whereas the contribution of mast cells to early tissue swelling in active anaphylaxis in mice has not been formally established, the decrease in intact mast cells 0.5 h after intradermal antigen challenge of immunized gp49B1-deficient mice (Fig. 3 C) suggests that a concomitant, augmented release of vasoactive mediators from mast cells accounts for the tissue edema (Fig. 3 B). Moreover, the findings indicate that the absence of gp49B1 augments the effector phase of a polyclonal, acquired immune response that likely involves mast cell activation through both IgE/FcεRI and IgG/FcγRIII.

**Effect of gp49B1 Deficiency on Active Systemic Anaphylaxis.** Mice immunized with OVA were injected intravenously with 10 mg OVA. Initial observations suggested that $gp49B^{-/-}$ mice became moribund more frequently and rapidly than normal animals, so the outcome was quantified in additional experiments. Whereas 8 out of 29 (27%) $gp49B^{+/+}$ mice died in response to systemic antigen challenge, 17 out of 25 (68%) $gp49B^{-/-}$ mice died. Moreover, the mean times to death were 45 ± 9 and 26 ± 8 min in $gp49B^{+/+}$ and $gp49B^{-/-}$ mice, respectively, which were significantly different ($P < 0.0001$). In two out of three $gp49B^{-/-}$ mice that died, the lungs showed focal leukose-
questration of neutrophils and mast cells had degranulated in the trachea. In the heart, myocyte swelling with distorted myofibrils was associated with mast cell degranulation. Mast cell degranulation was not seen in three gp49B<sup>++</sup> mice at the interval by which the gp49B<sup>−/−</sup> mice died.

That the mast cell can be responsible for a fatal outcome has been established in passive models using anti-IgE or IgE sensitization followed by antigen, in which death occurs within 40 to 60 min in WBB6F1<sup>+/−</sup> mice, but mast cell–deficient WBB6F1<sup>−/−</sup> mice survive (27–29). In active systemic anaphylaxis with these two strains of mice, the mast cell contribution was most evident for the cardiac tachycardia, and death was not dependent on the mast cell, thus reflecting other pathways (27, 28). Hence, while the mast cell may not be the absolute cause of death in gp49B<sup>−/−</sup> mice, the susceptibility of the gp49B<sup>−/−</sup> mice to fatal active systemic anaphylaxis in a relatively short interval compared with gp49B<sup>+/−</sup> mice is compatible with an augmented burst of mediator release from mast cells.

In summary, our results establish that gp49B1 is an inhibitor of FcεRI–directed mast cell activation and attendant immediate hypersensitivity reactions in vivo. This constitutive effect of gp49B1 is sufficiently robust to attenuate the local and systemic anaphylactic effects of antigen challenge during an ongoing immune response. Thus, tissue mast cells through their expression of gp49B1 possess an innate system that counterregulates acquired allergic inflammation.

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