Mast cells can promote the development of multiple features of chronic asthma in mice

Mang Yu, … , James Zehnder, Stephen J. Galli

*J Clin Invest.* 2006;116(6):1633-1641. [https://doi.org/10.1172/JCI25702](https://doi.org/10.1172/JCI25702).

Bronchial asthma, the most prevalent cause of significant respiratory morbidity in the developed world, typically is a chronic disorder associated with long-term changes in the airways. We developed a mouse model of chronic asthma that results in markedly increased numbers of airway mast cells, enhanced airway responses to methacholine or antigen, chronic inflammation including infiltration with eosinophils and lymphocytes, airway epithelial goblet cell hyperplasia, enhanced expression of the mucin genes *Muc5ac* and *Muc5b*, and increased levels of lung collagen. Using mast cell–deficient (*Kit*<sup>W-sh/W-sh</sup> and/or *Kit*<sup>W/W-v</sup>) mice engrafted with *FcR<sub>γ</sub>+</sub>+ or *FcR<sub>γ</sub>−/−* mast cells, we found that mast cells were required for the full development of each of these features of the model. However, some features also were expressed, although usually at less than wild-type levels, in mice whose mast cells lacked *FcR<sub>γ</sub>* and therefore could not be activated by either antigen- and *IgE*-dependent aggregation of *FcεRI* or the binding of antigen-*IgG1* immune complexes to *FcγRIII*. These findings demonstrate that mast cells can contribute to the development of multiple features of chronic asthma in mice and identify both *FcR<sub>γ</sub>*-dependent and *FcR<sub>γ</sub>*-independent pathways of mast cell activation as important for the expression of key features of this asthma model.
Mast cells can promote the development of multiple features of chronic asthma in mice

Mang Yu,† Mindy Tsai,† See-Ying Tam,† Carol Jones,† James Zehnder,† and Stephen J. Galli†1,2

†Department of Pathology and ‡Department of Microbiology and Immunology, Stanford University School of Medicine, Stanford, California, USA.

Bronchial asthma, the most prevalent cause of significant respiratory morbidity in the developed world, typically is a chronic disorder associated with long-term changes in the airways. We developed a mouse model of chronic asthma that results in markedly increased numbers of airway mast cells, enhanced airway responses to methacholine or antigen, chronic inflammation including infiltration with eosinophils and lymphocytes, airway epithelial goblet cell hyperplasia, enhanced expression of the mucin genes Muc5ac and Muc5b, and increased levels of lung collagen. Using mast cell–deficient (KitW+/-,W+/- or KitW+/+ mice engrafted with FcγRI−/− or FcγRI−/− mast cells, we found that mast cells were required for the full development of each of these features of the model. However, some features also were expressed, although usually at less than wild-type levels, in mice whose mast cells lacked FcγRI and therefore could not be activated by either antigen- and IgE-dependent aggregation of FcγRI or the binding of antigen-IgG1 immune complexes to FcγRIII. These findings demonstrate that mast cells can contribute to the development of multiple features of chronic asthma in mice and identify both FcγRI-dependent and FcγRI-independent pathways of mast cell activation as important for the expression of key features of this asthma model.

Introduction

Asthma is a complex inflammatory disorder that is becoming increasingly prevalent, particularly in the developed world (1–3). Once thought to be mainly a disorder of airway smooth muscle reactivity, asthma now is recognized to be a syndrome associated with persistent chronic inflammation of the airways (1–3). It has also become apparent that long-standing asthma can result in changes in the airways themselves, called “airway remodeling” (1–5). Thus, the spectrum of findings in this disorder is thought to include increased numbers of mucus-secreting goblet cells in the airway epithelium, associated with excessive production of mucus, and increased deposition of collagen in the lung as well as airway hyperresponsiveness (AHR) to immunologically nonspecific agonists of bronchoconstriction and chronic inflammation of the airway mucosa, with cellular infiltrates of lymphocytes, eosinophils, and other inflammatory leukocytes (1–8).

In many patients, asthma has an allergic or atopic component, characterized by allergic sensitivity to environmental allergens and increased serum levels of antigen-specific and total IgE antibodies (1–3). Accordingly, attempts to understand the pathophysiology of asthma often focus on the effector cells of allergic inflammation, including the mast cell (MC). MCs are derivatives of hematopoietic progenitors that differentiate and mature locally in most tissues, especially those that are exposed to the external environment, such as the airways (9–11). Moreover, some studies have reported that MCs are present in increased numbers in the airways of patients with asthma (8, 12, 13).

While the mechanisms responsible for the increased numbers of MCs in the airways of some asthmatic patients have not been defined, there is no doubt that MCs have the potential to influence multiple aspects of the pathology and pathophysiology of asthma. For example, MCs express the high affinity receptor for IgE (i.e., FcγRI) on their surface and therefore can bind IgE (14–16). Upon exposure to di- or multivalent antigen recognized by FcγRI-bound IgE, these FcγRI undergo aggregation, initiating a complex process of signal transduction that culminates in the secretion of a broad spectrum of proinflammatory mediators, cytokines, and chemokines (11, 14–16). In the mouse, the FcγRI chain (FcγRI) is shared by FcγRI and FcγRIII (17), and MCs can be activated to release mediators via Ag-, IgG1-, and FcγRIII-dependent mechanisms (17–22) as well as by Ag-, IgE-, and FcγRII-dependent mechanisms (14–16).

These MC-derived products can have diverse effects in the airways, including those that might, at least in part, contribute to multiple features of this disorder (1, 8, 13, 23, 24). Thus, individual MC mediators, cytokines, or chemokines can promote AHR and chronic allergic inflammation, goblet cell hyperplasia, enhanced mucin gene expression and mucus secretion, and collagen deposition (1, 8, 13, 23, 24). Moreover, patients with active asthma or asthma patients challenged via the airways with allergens to which they have sensitivities exhibit evidence of MC activation, such as increased levels of MC-derived products in bronchoalveolar lavage (BAL) fluid (13, 24). Finally, histological signs of airway MC degranulation in vivo can be identified in biopsy specimens derived from asthmatic patients (6, 13, 24).

Yet it is not at all clear to what extent MCs and IgE-dependent MC activation actually are important for the long-term pathology of asthma. First, a number of other inflammatory cells, including T lymphocytes, macrophages, eosinophils, basophils, and neutrophils, have also been implicated as potential effector cells in asthma (6, 8, 25–28). However, in humans, it can be difficult to ascertain the individual contributions of each of these potential effector cells to the pathology of the disorder. Second, while attempts to treat asthma by reducing levels of IgE with the humanized anti-IgE antibody omalizumab have produced clinical benefit in some patients, the magnitude of such benefit often is...
limited (29, 30). The extent to which the relatively modest clinical results obtained with omalizumab reflect an inability to reduce IgE levels to sufficiently low levels, as opposed to the presence of mechanisms of disease pathogenesis that operate independently of IgE, is not clear.

Studies of asthma models in mice can exploit the power of genetic approaches in analyzing how the presence or absence of potential effector elements can influence the expression of the various features of the models (23, 31). Such work has implicated MCs in the expression of certain features that develop in relatively short-term models of asthma, including AHR and eosinophil-containing inflammatory infiltrates in the lung (32–34). However, in other short-term models of asthma, which typically involve sensitization with antigen administered together with an adjuvant such as aluminum hydroxide, AHR and eosinophil infiltration can be elicited in animals that genetically lack MCs (23, 25, 33, 35).

Even though humans with allergic asthma are typically exposed to allergens repeatedly over long periods of time, only a few studies of mice have investigated protocols of allergen challenge that attempt to mimic this pattern of exposure (36–38). Moreover, of the features of chronic asthma. We then used that model to term models of asthma, including AHR and eosinophil-containing cell infiltrates. We also performed invasive measurements of lung function to assess AHR in anesthetized, tracheostomized mice (Supplemental Figure 1B). When data from individual mice were pooled, the biphasic response to OVA challenge in such mice, were largely or entirely MC dependent.

We also measured airway responses to antigen or PBS. Individual OVA-sensitized Kit WT/mice exhibited a biphasic response to the ninth OVA challenge, with peaks of responsiveness at approximately 3 or 6 hours and at approximately 9 or 12 hours; the responses were substantially reduced by 24 hours after OVA challenge (Supplemental Figure 2). When data from individual mice were pooled, the biphasic response to OVA was still evident in OVA-sensitized Kit WT/mice but not in OVA-sensitized Kit WT/mice (Figure 1B). However, peak airway responses in these 2 groups were quite similar in magnitude.

In contrast, there were no detectable responses to OVA in OVA-sensitized Kit WT/mice, nor did any group of PBS-treated mice exhibit a response to PBS challenge (Figure 1B). Very similar results were obtained when the same experiments were performed in C57BL/6-Ki TWT-/ mice and C57BL/6-Ki TWT+/ mice (Supplemental Figure 1B). Thus, airway responses induced by OVA challenge in OVA-sensitized Kit WT+/mice or Kit WT+/mice, like Penh responses to methacholine after OVA challenge in such mice, were largely or entirely MC dependent.

To assess specifically airway responses distal to the trachea, we also performed invasive measurements of lung resistance (Rl) and dynamic compliance (Cdyn) in mice treated with aerosolized methacholine 24 hours after the ninth challenge with OVA or PBS. OVA-sensitized Kit WT/+ or Kit WT+/mice exhibited similar levels of AHR after OVA challenge whereas the OVA-sensitized MC-deficient Kit WT+/mice showed responses that were statistically indistinguishable from those of the PBS-treated control groups (Figure 1, C and D). Thus, our findings using invasive measurements of lung function to assess AHR in anesthetized, tracheostomized mice are basically in accord with those we obtained by using Penh to assess airway responses in conscious mice.
Antibody responses in this model are MC independent. MCs can be activated by antigen to secrete mediators by both IgE/FcεRI- and IgG/FcRIII-dependent mechanisms (14–22). Such secreted products include cytokines that can regulate the production of IgE and other antibodies (23, 39, 40). MCs can also express other functions that have the potential to influence the development of adaptive immune responses (23, 39, 40). Accordingly, it has been proposed that MCs might function in part to enhance levels of antibody production during some acquired immune responses (23, 39, 40).

We found that OVA sensitization resulted in the development of significant antigen-specific IgE and IgG1 responses, but the levels of these antibodies measured at the end of our experiments in groups of OVA-sensitized Kit−/−, KitW/W−, or Kit−/+ BMCMCs→kit+/+ mice were statistically indistinguishable (Supplemental Figure 3). Thus, the differences in airway responses to OVA or in the AHR to methacholine among OVA-sensitized Kit−/−, KitW/W−, or Kit−/+ BMCMCs→kit+/+ mice cannot be explained by differences in levels of OVA-specific IgE or IgG1 antibodies.

This asthma model is associated with increased numbers of MCs and MC-dependent lung inflammation. OVA treatment resulted in significant increases in the numbers of MCs in the lungs of the WBB6F1→Kit−/− mice, including the appearance of intraepithelial MCs (Figures 2A and 3A); similar findings were observed in the C57BL/6→Kit−/− mice (Supplemental Figure 4A and data not shown). These findings have also been reported in some patients with asthma (13). OVA treatment also induced other features of allergic inflammation, such as prominent infiltrates of leukocytes, including lymphocytes and eosinophils (Figures 2, D and G, and 3, B and D); again, similar findings were observed in C57BL/6→Kit−/− mice (Supplemental Figure 4, B and D).

In certain settings, KitW/W− mice can develop MC populations by mechanisms that do not require normal signaling via the c-Kit receptor (11). However, the lungs of OVA-treated KitW/W− mice (Figures 2B and 3A) or KitW/,W− mice (Supplemental Figure 4A) remained completely devoid of MCs. In contrast, lung MC numbers in OVA- or PBS-treated Kit−/− BMCMCs→Kit−/− mice were statistically indistinguishable from those in Kit−/− mice (Figures 2C and 3A); similar findings were observed in Kit−/− BMCMCs→KitW/W− mice (Supplemental Figure 4A).

In OVA-sensitized/challenged WBB6F1→Kit−/− mice, the ninth OVA challenge resulted in a rapid rise in the levels of histamine detectable in the serum, with levels remaining significantly elevated over baseline (i.e., pre-OVA challenge) values even 24 hours after OVA challenge (Supplemental Figure 5 and Figure 3C). In OVA-sensitized/challenged mice, serum histamine measured 24 hours after the last OVA challenge was not only significantly elevated in Kit−/− mice, but also, albeit to a lesser extent, in Kit−/− BMCMCs→Kit−/− mice; in contrast, only very low and statistically indistinguishable levels of serum histamine were detected in OVA- or PBS-treated KitW/W− mice (Figure 3C); similar findings were observed in C57BL/6→Kit−/−, C57BL/6→KitW/W−, and C57BL/6→Kit−/− BMCMCs→KitW/W− mice (Supplemental Figure 4C). The corresponding values (serum histamine 24 hours after challenge) for OVA-sensitized, PBS-challenged WBB6F1→Kit−/−, WBB6F1→KitW/W−, or WBB6F1→Kit−/− BMCMCs→KitW/W− mice were 31.4 ± 4.2, 6.0 ± 1.0, and 28.7 ± 4.6 nM, respectively. In naive WBB6F1→Kit−/−, WBB6F1→KitW/W−, or WBB6F1→Kit−/− BMCMCs→KitW/W− mice, serum histamine levels were 25.9 ± 5.1, 5.1 ± 1.2, and 23.6 ± 3.4 nM, respectively. These results indicate that, in this setting, serum histamine is derived solely or largely from MCs and/or is derived from other cellular sources in an MC-dependent manner.

Kit−/− and Kit−/+ BMCMCs→KitW/W− mice were also very similar in measurements of the inflammation associated with responses to OVA sensitization and challenge, with markedly increased numbers of inflammatory cells in the lungs versus only minimal responses in KitW/W− mice (Figures 2, D–I, and 3, B and D). OVA-treated Kit−/− or Kit−/+ BMCMCs→KitW/W− mice also exhibited significant elevations in BAL fluid monocytes, macrophages, neutrophils, eosinophils, and lymphocytes (Figure 3D). While there were some differences in the number of individual types of leukocytes in the BAL fluid of antigen-challenged Kit−/− versus Kit−/+ BMCMCs→KitW/W− mice, taken together, the data in Figures 2, D–I, and 3, B and D, demonstrate that MCs
are largely responsible for the antigen-induced leukocyte recruitment and chronic inflammation in this model. The same conclusion was supported when the same experiments were repeated using C57BL/6-Kit-/+; C57BL/6-KitW/w-/-; and C57BL/6-KitW/w-/- BMCMCs→KitW/w-/- mice (Supplemental Figure 4, B and D).

Activation of MCs via IgE or, in the mouse, IgG1 antibodies requires signaling through the FcRγi (14, 17). We found that levels of FcRγi mRNA in lungs obtained 24 hours after the last challenge with OVA or PBS were highly upregulated in OVA-treated Kit+/+ or Kit-/- BMCMCs→KitW/w-/- mice but not in MC-deficient KitW/w-/- mice (Figure 4A); similar results were obtained in C57BL/6-Kit-/-; C57BL/6-KitW/w-/-; and C57BL/6-KitW/w-/- BMCMCs→KitW/w-/- mice (Supplemental Figure 6). Although FcRγi is not restricted to MCs (41), these results are in accord with our findings that levels of tissue MCs and other hematopoietic cells are much more markedly increased in the lungs of OVA-challenged as opposed to PBS-challenged wild-type, Kit+/+ BMCMCs→KitW/w-/-, or Kit-/- BMCMCs→KitW/w-/- mice than in OVA-challenged MC-deficient KitW/w-/- or KitW/w-/- BMCMCs→KitW/w-/- mice (Figures 2, A–I, and 3, A, B, and D; Supplemental Figure 4, A, B, and D). We also quantified levels of a transcript which is thought to be restricted to T cells, i.e., T cell–associated GTPase (42), and found that levels of this mRNA were also highly upregulated in the lungs of OVA-treated Kit+/+ or Kit-/- BMCMCs→KitW/w-/- mice but not in MC-deficient and KitW/w-/- mice (Figure 4B).

Taken together, these quantitative RT-PCR results are consistent with our histological findings, which showed that OVA treatment was associated with increased numbers of MCs (Figures 2, A and C, and 3A; Supplemental Figure 4A) and striking lung infiltrates of lymphocytes, as well as of other leukocytes (Figure 2, D, F, G, and I, and data not shown).

MC-dependent enhancement of airway goblet cell hyperplasia, mucin gene expression, and collagen deposition. Quantitative RT-PCR analysis demonstrated a highly MC-dependent upregulation of expression of genes encoding mucins SAC and 5B, which are the major components of mucus secreted by goblet cells and submucosal glands, respectively (7, 43) (Figure 5, A and B; Supplemental Figure 7, A and B). These findings are consistent with our histological observations, which revealed that the striking increases in numbers of mucus-secreting goblet cells in the airway epithelium of OVA-treated mice were largely or fully MC-dependent (Figures 2, J–L, and 5C; Supplemental Figure 7C).

**Figure 3**

Features of allergic inflammation in this chronic asthma model 24 hours after the ninth OVA or PBS challenge. (A and B) Numbers of lung MCs (A) and eosinophils (B). (C) Serum histamine concentration. (A–C) White bars, PBS-treated group; black bars, OVA-sensitized/challenged group. **P < 0.01 or ***P < 0.001 versus corresponding PBS controls (n = 6); **P < 0.01 or ***P < 0.001 versus group indicated (n = 6 per group). ND, not detected. (D) Numbers of leukocytes in BAL fluid from the right lungs of WBB6F-Kit+/+ (black bars), WBB6F-KitW/w-/- (white bars), and KitW/w-/- BMCMCs→KitW/w-/- (gray bars) mice following OVA sensitization/challenge (O) or PBS treatment (P). **P < 0.01 or ***P < 0.001 versus corresponding PBS controls (n = 8 per group); **P < 0.01, or ***P < 0.001 versus group indicated (n = 8 per group). MON, monocytes; MAC, macrophages; PMN, neutrophils; EOS, eosinophils; LYM, lymphocytes.

**Figure 4**

Lung mRNA levels of genes encoding (A) FcRγi chain and (B) T cell–specific GTPase 24 hours after the ninth OVA or PBS challenge. White bars, PBS-treated group; black bars, OVA-sensitized/challenged group. **P < 0.01 versus corresponding PBS controls (n = 6 per group); ***P < 0.001 versus group indicated (n = 6 per group). Eq, equivalents.
Masson trichrome staining revealed that OVA treatment resulted in enhanced deposition of collagen in the airways of Kit\(^{-/-}\) or Kit\(^{+/+}\) BMCMCs→Kit\(^{-/-}\) mice but had little or no effect in the airways of Kit\(^{+/+}\) mice (Figure 2, J–L); similar results were observed in C57BL/6-Kit\(^{+/+}\), C57BL/6-Kit\(^{-/-}\), and C57BL/6-Kit\(^{+/+}\) BMCMCs→Kit\(^{-/-}\) mice (data not shown). Much of this enhanced collagen deposition was localized in the airways immediately below the airway epithelium, a location that also exhibited what appeared to be an increased amount of airway smooth muscle (Figure 2, J and L).

Quantification of lung hydroxyproline in WBB6F\(^1\) mice confirmed that the OVA-induced increase in lung collagen in this model was predominantly MC-dependent (Figure 5D).

MC expression of FcRγ is required for optimal expression of enhanced airway responses, numbers of airway MCs, airway inflammation, and mucin gene expression. MCs can be activated to release mediators in response to ligand-dependent engagement of multiple distinct cell surface receptors and signaling pathways (10, 11, 39). However, antigen-dependent activation of MCs via either IgE/FcεRI or IgG1/FcγRIII requires signaling mediated by the γ chain (FcRγ) common to both of these receptors (14, 17). To examine the extent to which FcεRI/FcγRIII-mediated MC activation is required for expression of our chronic asthma model, we compared the expression of various features of our model in Kit\(^{+/+}\) or Kit\(^{-/-}\) mice that had been engrafted with C57BL/6-FcRγ\(^{-/-}\) versus C57BL/6-FcRγ\(^{+/+}\) BMCMCs.

MC expression of FcRγ in BMCMCs→Kit\(^{+/+}\) mice had no detectable effects on the levels of total or antigen-specific IgE or IgG1 antibodies (Supplemental Figure 8). However, OVA treatment induced little or no enhancement of Penh responses to methacholine in FcRγ\(^{-/-}\) BMCMCs→Kit\(^{+/+}\) mice, but substantial enhancement of the responses was observed in FcRγ\(^{+/+}\) BMCMCs→Kit\(^{+/+}\) mice (Figure 6A). Similarly, OVA challenge of OVA-sensitized mice induced a much stronger airway response in FcRγ\(^{+/+}\) BMCMCs→Kit\(^{+/+}\) mice than in FcRγ\(^{-/-}\) BMCMCs→Kit\(^{+/+}\) mice (Figure 6B). Very similar results were obtained in C57BL/6-FcRγ\(^{-/-}\) versus C57BL/6-FcRγ\(^{+/+}\) BMCMCs→Kit\(^{-/-}\) mice (Supplemental Figure 9), except that the FcRγ\(^{-/-}\) BMCMCs→Kit\(^{-/-}\) mice, in contrast to the FcRγ\(^{+/+}\) BMCMCs→Kit\(^{+/+}\) mice, exhibited no detectable enhancement of Penh in response to OVA challenge (compare with Supplemental Figure 9B and Figure 6B).

In accord with these results, we found that expression of FcRγ by adoptively transferred MCs significantly enhanced the ability of such MCs to orchestrate most of the other features of this chronic asthma model in the recipient Kit\(^{+/+}\) mice (Figures 7 and 8, Supplemental Figures 10 and 11). Indeed, OVA-induced increases in numbers of lung MCs appeared to be entirely dependent on the expression of FcRγ by the adoptively transferred MCs (Figure 7A and Supplemental Figure 10A). These results strongly suggest that an effect mediated through MC-FcRγ, perhaps including the ability of IgE antibodies to promote MC survival or proliferation even in the absence of known antigen (44–48), contributed to the increased numbers of MCs observed in this setting.

However, in some other features of the response that were strongly expressed in FcRγ\(^{+/+}\) BMCMCs→Kit\(^{+/+}\) or FcRγ\(^{-/-}\) BMCMCs→Kit\(^{-/-}\) mice, OVA treatment also induced significant, albeit relatively modest, responses in the FcRγ\(^{-/-}\) BMCMCs→Kit\(^{+/+}\) or FcRγ\(^{-/-}\) BMCMCs→Kit\(^{-/-}\) mice. For numbers of lung eosinophils (Figure 7B and Supplemental Figure 10B), numbers of most types of leukocytes present in the BAL fluid 24 hours after OVA challenge (Figure 7D and Supplemental Figure 10D), and levels of mRNA of genes encoding mucin 5AC and mucin 5B (Figure 8, B and C; Supplemental Figure 11, B and C), FcRγ, and T cell GTPase (Supplemental Figure 12, A and B), the responses in the FcRγ\(^{-/-}\) versus FcRγ\(^{+/+}\) BMCMCs→Kit\(^{+/+}\) mice.
BMCMCs→Kitw+W−/− or FcRγ−/− BMCMCs→Kitw+W−/− mice, although significant compared with responses in the PBS-challenged mice, were significantly lower, and in many cases at least 50% lower, than those in the OVA-challenged FcRγ−/− BMCMCs→Kitw+W−/− or FcRγ−/− BMCMCs→Kitw+W−/− mice. These results indicate that FcRγ-independent mechanisms of MC activation can contribute to the development of these features of this chronic asthma model but that optimal responses require MC expression of FcRγ.

FcRγ-independent mechanisms of MC activation can significantly contribute to elevations of serum histamine and increased numbers of airway goblet cells. Serum histamine levels were significantly lower in OVA-treated FcRγ−/− versus FcRγ−/− BMCMCs→Kitw+W−/− mice, but the reduction was only approximately 19% (Figure 7C). Given that virtually all of the elevation of serum histamine in this model is MC dependent (Figure 3B), the most straightforward interpretation of this finding is that in WBB6F1 mice, OVA-treatment either induced histamine release from other sources. Notably, in mice on the C57BL/6 background, FcRγ-independent mechanisms appear to be less important than FcRγ-dependent mechanisms in regulating OVA-induced histamine release (Supplemental Figure 10C).

The increased numbers of goblet cells induced in OVA-treated versus PBS-treated FcRγ−/− BMCMCs→Kitw+W−/− mice reached levels that were nearly as high as, and were statistically indistinguishable from, the levels induced in the corresponding FcRγ−/− BMCMCs→Kitw+W−/− mouse (Figure 8A). In contrast, the enhanced levels of Mac5ac and Mac5b mRNA in OVA-challenged mice were significantly reduced in FcRγ−/− versus FcRγ−/− BMCMCs→Kitw+W−/− mice (Figure 8, B and C). FcRγ-independent mechanisms appeared to contribute significantly both to the increases in goblet cell numbers and to the enhanced levels of Mac5ac and Mac5b mRNA in OVA-challenged C57BL/6 mice as well (Supplemental Figure 11).

**Discussion**

We have developed a mouse model in which repetitive i.n. challenge of sensitized mice with antigen induces changes in the airways and lungs that mimic many of the features of chronic asthma in humans. Our studies in MC-engrafted Kitw+W−/− mice indicate that MCs are not required for the generation of antigen-specific IgE or IgG1 antibody responses in this model but are required for the development of every other feature of the model that we analyzed. Moreover, in many instances, the optimal development of...
individual features of the pathology and pathophysiology in this model require that MCs express FcRγ, and MC expression of FcRγ appears to be required for virtually all of the increase in lung MC numbers observed in OVA-treated mice (Supplemental Table 2). Notably, using the more recently described MC-engrafted C57BL/6-Kit/J-W-sh mouse model (49, 50), in which both the adoptively transferred FcRγ−− or FcRγ++ BMCMCs and the recipient genetically MC-deficient Kit/J-W-sh mice are all on the C57BL/6 background, we obtained results very similar to those obtained in experiments employing the transfer of C57BL/6-FcRγ−− versus C57BL/6-FcRγ++ BMCMCs into WBB6F1-kit/J-W-sh mice.

The simplest interpretation of these findings is that these MC-FcRγ−− dependent features of the response reflect consequences of the activation of MCs by antigen and antigen-specific IgE and/or IgG1 antibodies (11, 14, 18). However, some features of the response, including the marked increase in lung MCs, may also (or alternatively) reflect actions of IgE on MCs that are mediated through FcRγ but can be observed in the absence of known specific antigen (47, 48); these effects include the promotion of MC survival, proliferation, and cytokine production (44–48). Nevertheless, our findings clearly identify the activation of MCs via FcRγ-dependent mechanisms as a key promoter of many of the features of this chronic asthma model. Such findings support the view that IgE-dependent MC activation represents an attractive therapeutic target in asthma (14, 30, 51).

However, our study also shows that certain features of this model are significantly influenced by effects of MCs that are independent of MC-FcRγ. Such antigen- and MC-dependent but MC-FcRγ−− independent effects contribute importantly in this model to the striking increases in the numbers of epithelial goblet cells (especially in mice on the WBB6F1 background) and also contribute significantly to the elevated levels of mRNA for mucins SAC and SB in the lungs (in both WBB6F1 and C57BL/6 mice). These findings have interesting implications for our understanding of the factors that contribute to enhanced mucin production in human asthma. Among the 18 Muc genes that have been identified in humans, MucSac and MucSb gene products represent the major gel-forming mucins in airway secretion (7, 43). And it has been reported that increased levels of mucin SAC and mucin SB can be detected in patients with asthma (6, 7). Increased mucin production contributes significantly to airway obstruction and AHR in asthma; accordingly, several potential therapies targeting mucus hypersecretion are currently under development (7, 52). Our data indicate that, at least in mice, MCs are major contributors to goblet cell hyperplasia and increased expression of MucSac and MucSb that are observed in this model of chronic asthma.

FcRγ−− independent mechanisms of MC engagement also contribute significantly to elevations of levels of serum histamine (especially in mice on the WBB6F1 background), lung infiltration with eosinophils and lymphocytes, and recruitment of leukocytes to the airways in response to antigen challenge, as reflected in counts of cells in BAL fluid. Moreover, some of the entirely MC-dependent airway responsiveness to antigen challenge in WBB6F1 mice (Figure 1B) and perhaps even some of the MC-dependent AHR to methacholine (Figure 1A) in this model also appear to reflect contributions by MC-dependent but MC-FcRγ−− independent processes (Figure 6, A and B). In contrast, in C57BL/6 mice, there was no detectable contribution of MC-FcRγ−− independent processes to the MC-dependent enhancement of Penh responses to either methacholine or antigen (Supplemental Figure 9).

In summary, our findings in this mouse model show that MCs can represent key drivers of many important inflammatory, structural, and functional changes of the lungs that are also observed in chronic human asthma. These observations are thus in accord with the many reports implicating MCs and their products in the pathology of human asthma (13, 24). However, while some of the features of our chronic asthma model appear to be strongly dependent on MC expression of FcRγ, other effects are clearly antigen- and MC-dependent but also significantly MC-FcRγ−− independent. Our findings show that this MC-FcRγ−− independent pathway can elicit important contributions of MCs to several critical features of chronic asthma, including epithelial goblet cell hyperplasia and markedly enhanced expression of genes encoding mucins SAC and SB.

On the other hand, the importance of such MC-dependent but MC-FcRγ−− independent pathways in this model may be influenced by the genetic background of the mice, in that such pathways appeared to be more important in the development of goblet cell hyperplasia and in the elevations of serum histamine after OVA challenge observed in WBB6F1 as opposed to C57BL/6 mice.

Adding even more complexity, a very large number of candidate mechanisms might be proposed to account for the MC-dependent but MC-FcRγ−− independent pathways that appear to contribute to the development of some of the features of this model. MCs can be activated independently of FcRγ by multiple factors that may be present in tissues affected by asthma in humans or by models of asthma in mice, including products of complement activation (C5a and C3a) as well as certain neuropeptides, neurotrophins, chemokines, products of recruited leukocytes, and other types of products that can induce or enhance MC activation (10, 39, 40, 53–55). Complicating further any analysis of the potential roles of such factors in our model are observations indicating that many (if not all) of these factors may be able to influence features of asthma independently of any effects they may have on MCs.

For example, we previously reported that C5a and C3a can promote contractile responses in intestinal smooth muscle independently of any contribution of MCs (56). Notably, aerosolized C5a can also induce elevations of Penh and increased numbers of BAL leukocytes independently of MCs in mice on either the WBB6F1 or C57BL/6 background (Supplemental Figures 13 and 14). Accordingly, we expect that extensive work (and probably some good fortune) may be required to identify the important MC-dependent but MC-FcRγ−− independent pathways that can contribute significantly to the development of some of the features of this model of chronic asthma.

The extent to which our observations can be extended to human asthma remains to be determined. However, the findings in this mouse model raise the possibility that therapies that target the IgE-dependent activation of MCs and other effector cells may not be fully effective in the treatment of chronic asthma even if it were possible fully to eliminate IgE-dependent effector cell activation by such approaches. Indeed, our data strongly suggest that MCs themselves and FcRγ-dependent pathways of MC activation in addition to IgE represent promising therapeutic targets in patients with chronic asthma.

Methods

Mice. Genetically MC-deficient (WB/Rej-W−− × C57BL/6J-W−−)F1-Kit/J-W−− mice, WBB6F1-kit/J-W−− mice, congenic normal WBB6F1-Kit/J-W−− mice, and C57BL/6J mice were purchased from Jackson Laboratory. C57BL/6-kit/J-W−− mice were generously provided by Peter Besmer (Molecular Biology Program,
Memorial Sloan-Kettering Cancer Center, New York, New York, USA); these mice were then backcrossed to C57BL/6J for more than 6 generations. FcRγ/− mice (57) on the C57BL/6J background were purchased from Taconic. Age-matched female mice were used for all experiments. All animal care and experimentation reported herein were conducted in compliance with the guidelines of the NIH and with the specific approval of Institutional Animal Care and Use Committee of Stanford University.

**MC engraftment.** Selective engraftment of MCs in MC-deficient WBB6F-Kit−/− or C57BL/6-Kit−/− mice was performed as described (33), with minor modifications. Bone marrow cells derived from 4-week-old female WBB6F-Kit−/−, C57BL/6J-FcRγ/−, or C57BL/6J-FcRγ/− mice were cultured in WEHI-3-conditioned medium (ATCC number TIB-68), as a source of IL-3, for 4–5 weeks to generate cell populations that contained more than 95% immature MCs. Via the tail vein, 5 × 10^6 BMCMCs were injected into each mouse, and the recipients (e.g., WBB6F-Kit−/− BMCMCs→Kit−/− or C57BL/6-Kit−/− BMCMCs→Kit−/− mice) were used for experiments 18 weeks later. Engraftment of Kit−/− mice with BMCMCs did not repair their anemia (as examined by hematocrit), confirming the selectivity of the MC engraftment (11, 33).

**Immunization and airway challenge with antigen.** Mice were immunized by 3 i.p. injections of 50 μg OVA (Sigma-Aldrich) in 0.1 ml PBS on days 1, 4, and 7. Starting on day 12, mice were challenged i.n. with 20 μg OVA in 30 μl PBS weekly for 9 weeks; control mice received i.p. injections and i.n. challenges with PBS on the same schedule. As shown in Supplemental Figures 15 and 16, repeated i.n. administrations of OVA also appeared to firm the selectivity of the MC engraftment (11, 33).

**Measurement of airway reactivity to methacholine.** Twenty-four hours after the eighth OVA or PBS challenge, responses to aerosolized methacholine were measured using whole-body plethysmography (Buxco) (33). Respons to antigen (OVA) or PBS were assessed by recording Penh over 10-minute time periods at intervals before and after the ninth (final) OVA or PBS challenge (in the absence of methacholine). In some studies, we also performed invasive measurements of airway reactivity in anesthetized, tracheostomized, mechanically ventilated mice (59). Aerosolized methacholine was administered in increasing concentrations (0, 6.25, 12.5, 25, 50, and 100 mg/ml), with individual doses separated by 4 minutes. Rl and Cdyn were continuously computed by fitting flow, volume, and pressure to an equation of motion for each aerosol challenge period, which consisted of a 2-minute aerosol exposure and a 4-minute period after exposure.

**Hydroxyproline assay.** Tissue hydroxyproline concentration was used to measure the collagen content of the lung. Sections of the left lung 1-mm thick were dried with acetone, approximately 10 mg of the dried samples were hydrolyzed with 0.5 ml 6 N HCl at 110°C overnight, and hydroxyproline concentration was determined as in ref. 61.

**Statistics.** Unless otherwise specified, differences in airway response between different groups were tested for statistical significance using ANOVA. The unpaired Student’s t test (2 tailed) was used for all other analyses. P < 0.05 was considered statistically significant. Unless otherwise specified, all data are presented as mean ± SEM or mean ± SEM.

**Acknowledgments**

We thank Zhen-sheng Wang and Alian Xu for technical assistance. Peter Besmer for his gift of C57BL/6-Kit−/− mice, Erwin W. Gelfand for the gift of an OVA-specific IgE standard, and members of the Galli laboratory for discussions. This work was supported by United States Public Health Service grants HL67674, AI23990, and CA72074 (to S.J. Galli) and a Ruth L. Kirschstein National Research Service Award (to M. Yu).

Received for publication May 20, 2005, and accepted in revised form April 11, 2006.

**Address correspondence to:** Stephen J. Galli, Department of Pathology, L-235, Stanford University School of Medicine, 300 Pasteur Drive, Stanford, California 94305-5324, USA. Phone: (650) 723-7975; Fax: (650) 723-6902; E-mail: sgalli@stanford.edu.


