Augmentation of Reverse Arthus Reaction by Mast Cells in Mice

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Abstract

Immune complex-induced injury is an important pathogenic factor in antibody-mediated nephritis, systemic lupus erythematosus, rheumatoid arthritis, and other diseases. In this study we investigated the role mast cells in immune complex-mediated injury in mouse skin. Reverse Arthus reaction was induced in mast cell-deficient WBB6F,-W/W' mice and their congenic controls (WBB6F₁-+/+). Serial skin sections were evaluated for neutrophil infiltration, edema, and hemorrhage. In WBB6F₁-W/W' mice the neutrophil influx was only 40% and edema 60% of that in congenic controls. Hemorrhage was also significantly reduced in the mast cell-deficient mice. After mast cell reconstitution, the magnitude of the reaction in WBB6F₁-W/W' was equivalent to that in WBB6F₁-+/+ mice. Mast cell release in reverse Arthus reaction was evaluated by measuring fluorescence intensity after avidin-FITC staining of mast cell granules. There was a 70% decrease in fluorescence intensity. The 5-lipoxygenase inhibitor A-63162 significantly decreased neutrophil accumulation (40%), edema (60%), and hemorrhage in WBB6F₁-+/+, but not in mast cell-deficient mice. Mast cell reconstitution of WBB6F,-W/W' mice restored the effect of A-63162. The results indicate that mast cells and their mediators, including leukotrienes, make an important contribution to reverse Arthus reaction. (J. Clin. Invest. 1991. 88:841-846.) Key words: leukotriene • immune complex-mediated injury. 5-lipoxygenase inhibitor - inflammation. PMN infiltration

Introduction

The formation of immune complexes and their deposition in tissues precipitate an acute inflammatory reaction. Immune complex-induced injury has been implicated in a variety of human diseases, such as systemic lupus erythematosus, rheumatoid arthritis, antibody-mediated glomerulonephritis, and other conditions. Reverse passive Arthus reaction has been extensively studied as a model of immune complex-mediated tissue injury. The reaction is induced by injecting the antigen i.v. followed by the local administration of the antibody. The resulting lesion is characterized by histological changes, includ-

ing leukocyte infiltration, edema, tissue necrosis, and hemorrhage. Neutrophils and activation of the complement pathway are essential for the reaction (1). However, the cell responsible for the initiation of leukocyte elicitation and edema is not yet well established. Mast cells are strategically located in the vicinity of blood vessels. Upon stimulation they release a variety of proinflammatory mediators, such as histamine, platelet-activating factor, cyclooxygenase products, and leukotrienes (LT).' Mast cells are well characterized for their role in immediate hypersensitivity reaction. Other functions of mast cells have not been extensively studied. Galli and Kitamura have published a comprehensive discussion of the value of mast cell-deficient mice for the investigation of the role of mast cells in biological responses (2). Galli and his co-workers have shown the importance of mast cells in the expression of such inflammatory processes as leukocyte infiltration, edema, and fibrin deposition in cutaneous anaphylaxis, and after application of phorbol myristate acetate and substance P (3-5). At the same time we initiated studies investigating the role of mast cells in inflammation other than immediate hypersensitivity. We found that mast cells contribute to the neutrophil recruitment in thioglycollate (6) and immune complex-induced peritonitis (7). It has been reported that mast cells play an important role in antigen-induced arthritis (8).

The present work investigates the role of mast cells in the development of reverse Arthus reaction in the skin using mast cell-deficient WBB6F₁-W/W^v mice and their congenic controls. The PMN infiltration, edema, and hemorrhage were evaluated. The results indicate that mast cells contribute to these inflammatory parameters. Studies with the 5-lipoxygenase inhibitor A-63 162 suggest that leukotrienes released by mast cells mediate these processes.

Methods

Materials. Chicken egg albumin, hematoxylin, eosin, alcian blue, safranin 0, and triethylenediamine were purchased from Sigma Chemical Co. (St. Louis, MO), avidin-FITC from Zymed Laboratories, Inc. (San Francisco, CA), rabbit anti-chicken egg albumin (IgG fraction) from Cappel Research Products (Durham, NC), PBS, RPMI-1640, L-glutamine, nonessential amino acids, sodium pyruvate, penicillin, and streptomycin from the Tissue Culture Support Center of Washington University (St. Louis, MO). The 5-lipoxygenase inhibitor A-63 162 was a gift of Dr. R. Bell (Abbott Laboratories, Abbott Park, IL).

Mice. Mast cell-deficient $WBB6F_1-W/W'$ (W/W') mice and their normal littermates, WBB6F₁-+/+ (+/+), were purchased from Jackson Laboratory (Bar Harbor, ME). These mice were used when they were 90 d or older. W/W" mice are established for their profound deficiency in mast cells (9). These mice also have macrocytic anemia and decreased numbers of granulocytes and megakaryocytes in the bone marrow. However, the number of granulocytes and platelets in the

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^{1.} Abbreviations used in this paper: Ab, antibody; ELAM- 1, endothelial-leukocyte adhesion molecule-l; ICAM-l, intercellular adhesion molecule-1; LT, leukotriene; W/W', WBB6F₁-W/W'; +/+, WBB6F₁- $+/+.$

blood of adult W/W' mice is normal (10, 11). Their mast cell deficiency and anemia can be repaired by bone marrow transplantation from WBB6F,-+/+ mice (9). However, administration of mast cells alleviates only the mast cell deficiency (7, 12). The ability of these mice to selectively undergo systemic or local repair of their mast cell deficiency renders these mutants a useful tool for the investigation of mast cell functions in vivo.

Induction and evaluation of the inflammatory reaction. Mice were injected i.v. with chicken ovalbumin (20 mg/kg), followed by antichicken ovalbumin antibody (Ab), IgG fraction, i.d. on shaven backs, 10, 30, and 100 μ g/site, two injections per dose. At the times indicated after the injection of Ab, skin specimens were fixed in 10% phosphate buffered formalin and embedded in paraffin. Serial skin sections (10 μ m) were stained with hematoxylin and eosin and evaluated microscopically for PMN infiltration, hemorrhage, and edema. The examination was performed in a blinded fashion. Equivalent results were observed by two different individuals. PMN infiltration was assessed either by counting PMN per section or in arbitrary units. PMN density for four to nine sections per injection site was averaged. Arbitrary units were used to evaluate most of the experiments because PMN infiltration with 100 μ g/site of Ab in +/+ mice was too dense to perform an actual PMN count. A scale of ¹ to ⁸ was chosen, ⁸ indicating the highest PMN density. The arbitrary units were established by counting the PMN in some sections in which individual PMN could be distinguished and by allocating certain ranges to the lower units $(1,-100, 2:100-400, 3:400-$ 700, 4:700-1,000 PMN/section). The remaining higher units were assigned depending on the size of the patches filled with PMN. Hemorrhage was also assessed in arbitrary units, 1-5. They were established by counting erythrocytes in some sections and allocating ranges (1:100- 200, 2:200-400, 3:400-600 RBC per section). The higher units were assigned depending on the size of the patches filled with erythrocytes. Edema was evaluated by measuring microscopically the thickness (millimeters) of mounted cross-sectioned skin specimens with a micrometer.

Mast cell reconstitution of W/W^v mice. Bone marrow cells harvested from femurs of male $WBB6F_1 +/+$ mice were cultured and differentiated into mast cells as described previously (13). These mast cells ($> 98\%$ purity) were injected i.d. (multiple sites, 1×10^6 per site) into W/WV mice. The mast cell-repleted sites were marked with India ink and challenged 7-8 wk after mast cell repletion.

Mast cell degranulation study. Skin sections from 8 h challenged (100 μ g Ab) sites (with or without pretreatment with A-63162, 100 mg/kg) were stained with avidin-FITC (6.25 μ g/ml) for 2 h and then mounted in buffered glycerol, ³⁰ mM triethylenediamine, pH 8.6. Avidin is known to label mast cells selectively and quantitatively (14, 15). Pretreatment with heparinase prevented mast cell staining. Mast cell granule content was evaluated by measuring fluorescence intensity (expressed in arbitrary units/mast cell) with an image analysis system (Microcomp; Southern Micro Instruments, Atlanta, GA). A significant decrease in fluorescence intensity of mast cells in stimulated sites as compared with those in control sites (PBS treated) was interpreted as degranulation of mast cells during the inflammatory reaction.

5-Lipoxygenase inhibitor study. Mice were treated orally with the 5-lipoxygenase inhibitor A-63162 (30 and 100 mg/kg) (16). The drug was given at 1-h intervals (three doses) before the injection of the Ab, and at 1.5-h intervals during the challenge period. We have shown that the doses of A-63 162 used cause a dose-dependent inhibition of leukotriene formation in immune complex peritonitis (17).

Results

PMN influx. Reverse Arthus reaction in the skin of $+/+$ mice as well as in the mast cell-deficient W/WV mice induced PMN infiltration which was dose dependent with regard to the amount of Ab injected (Fig. 1). With the lowest dose of Ab used, 10 μ g/site, PMN influx was insignificant and was similar to that observed in mice treated only with 100 μ g Ab/site but

no antigen (i.v.) (data not shown). As expected, no PMN were detected in skin which was only treated with PBS and only very few when the mice were treated with 100 μ g Ab/site but not with antigen. The time course of leukocyte influx was established (Fig. 1). In both groups of mice a significant number of PMN was observed at 4 h (the first time point tested), and maximum was reached by ⁸ h after challenge. However, with both doses of Ab (30 and 100 μ g/site) and at every time point tested, the PMN density in W/W^v was significantly lower than in $+/+$ mice. At the maximum (8 h), PMN infiltration in W/W["] mice was only 40% of that in $+/+$ mice. The marked difference between the two sets of mice was quite striking and is illustrated in Fig. 2.

Edema and hemorrhage. The immune complex-mediated reaction was also accompanied by edema and hemorrhage. Edema, evaluated as skin thickness, was dose dependent with regard to antibody (Fig. 3). No edema was observed with 10 μ g Ab/site or if only Ab, $100 \mu g$ /site, but not antigen (i.v.) was administered. There was significantly less edema in the mast cell-deficient than in $+/+$ mice (Fig. 3). In $+/+$ mice, with 100 μ g of Ab significant edema was already noticed at 4 h (data not shown) and was maximum at 8 h. With 30 μ g of Ab, edema was evident only at 8 and 10 h. In W/W^v mice, measurable edema could be observed starting at 6 h (data not shown), with the peak at 8 h with 100 μ g of Ab and only at 8 and 10 h with 30 μ g of Ab. Hemorrhage was most pronounced with 100 μ g of Ab, reached maximum at 8 h (Fig. 4) and was absent with 10 μ g Ab/site or if only Ab, $100 \mu g$ /site, was given and no antigen, i.v. Similar to the other parameters studied, hemorrhage was significantly less in W/W^{\vee} than in +/+ mice (Fig. 4).

Mast cell reconstitution of W/W^v mice. The experiments described above indicate that all three parameters of inflammation evaluated were significantly reduced in W/Wv mice. To confirm that the differences observed were due to the mast cell deficiency and not due to some other abnormality of these mice, certain areas of the skin were repleted with mast cells (> 98% purity). Therefore, only the mast cell deficiency was corrected (7, 12). The cultured mast cells injected i.d. into W/WV mice were mucosal-like and stained only with alcian blue and not with safranin 0. Histological examination after

 $\frac{1}{\Box \text{A}}$ $\frac{1}{\Diamond \text{B}}$ $\frac{1}{\Diamond \text{C}}$ $\frac{1}{\Diamond \text{$ filtration in reverse Arthus reac- 6 $\overline{1}$ tion. Mice were treated with ovalbumin, 20 mg/kg i.v., and with
anti-ovalbumin antibody i.d. At $\begin{array}{ccc}\n & & \text{tion. Mice were treated with ova
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2. $2-\frac{1}{\sqrt{2}}$ $\frac{1}{\sqrt{2}}$ $\frac{1}{\sqrt{2}}$ $\frac{1}{\sqrt{2}}$ ion of Ab, skin specimen were fixed and serial sections stained $\begin{array}{|c|c|c|c|c|}\n\hline\n\text{Al} & \text{Al} & \text{with hematoxylin and eosin were examined microscopically. Arbi-} \n\end{array}$ trary units were used to evaluate 6 \uparrow the extent of PMN influx because the PMN infiltration with 100 μ g/ site of Ab in $+/+$ mice was too dense to perform actual cell counts. There was no PMN infil tration in PBS-injected sites of
mice treated i.v. with ovalbumin. Hours 4 6 8 10 **PMN** influx in $+$ /+ mice given 100 μ g Ab/site but no ovalbumin

was 0.75 ± 0.2 , $n = 3$, at 8 h. Data are expressed as mean \pm SD, $n = 5$, $*P < 0.02, **P < 0.001.$

Figure 2. Effect of mast cell deficiency on PMN infiltration. Mice were challenged with ovalbumin i.v., and 100 µg/site of anti-ovalbumin antibody i.d. for 8 h. Representative photomicrographs (10× objective) of skin sections from (A) +/+ mice and (B) W/W' mice.

mast cell reconstitution showed, that similar to normal skin, a significant number of mast cells was found in the vicinity of blood vessels. The staining pattern of the injected mast cells had changed, and some of them were now not only alcian blue but also safranin 0 positive, indicating differentiation to connective tissue mast cells (data not shown). In these mast cell-reconstituted W/W" mice reverse Arthus reaction resulted in PMN infiltration (Fig. 5), edema (Fig. 3) and hemorrhage (Fig. 4) comparable with that in the $+/+$ mice. These findings indicate that mast cells make an important contribution in immune complex-mediated inflammation and augment PMN influx, edema, and hemorrhage.

Mast cell degranulation studies. To further investigate the participation of mast cells in this model of inflammation, the degranulation of mast cells in stimulated sites was studied. This was accomplished by treating the tissue with avidin-FHTC, a stain specific for mast cell granules (14, 15). Pretreatment with heparinase blocked the fluorescent staining. The involvement of mast cells in this reaction was corroborated by a 70% decrease in the fluorescence intensity of mast cells in stimulated

skin (PBS control: 180 ± 9.6 , $n = 160$; stimulated: 54.6 ± 12.7 arbitrary units, $n = 90$ mast cells, mean \pm SD). This indicates that mast cells have lost a large portion of their granules during the inflammatory reaction.

Effect of 5-lipoxygenase inhibition. It is known that leukotrienes are released in inflammation, including immune complex reaction (18-20). LTB₄ is a potent chemotactic factor for PMN, and LTC_4 and D_4 can cause plasma exudation. We tested the involvement of leukotrienes in reverse Arthus reaction by pretreating mice with the 5-lipoxygenase antagonist A-63162. In $+/+$ mice, this compound (100 mg/kg) caused a 40% decrease in PMN infiltration (Fig. ⁶ A). No significant effect was observed with 30 mg/kg (data not shown). In the mast cell-deficient mice, no significant reduction in PMN density occurred with either of the doses of A-63162 tested. To improve the accuracy of these experiments, actual PMN counts were performed on the sections from W/W" mice (Fig. ⁶ B). Furthermore, A-63162 inhibited edema (Fig. 7) only in $+/+$ but not in W/W^{\cdot} mice. The 5-lipoxygenase inhibitor also significantly decreased hemorrhage in $+/+$ mice. The effect of this drug on hemorrhage in W/W^v mice was difficult to interpret due to the high variability observed (Fig. 8). A-63 162 had no effect on mast cell degranulation. The fluorescence intensity of mast cells, reflecting granule content, after challenge with

Figure 3. Effect of mast cell deficiency on edema. Skin thickness of +/+, W/W', and mast cell-reconstituted W/W' (W/W' + MC) mice was measured ⁸ h after Ab challenge. First set of bars represents PBS-treated sites from the same animals. The skin thickness of animals given 100 μ g Ab/site but not antigen was not significantly different from PBS-injected sites (0.51±0.01 mm). The data are expressed as mean \pm SD, $n = 3-9$, $*P < 0.01$ as compared with the same dose in $+/+$ mice.

Hemorrhage was evaluated challenge in $+/+$, W/W^v, and mast cell-reconstituted mice. The data are expressed as mean \pm SD, n $=$ 3-9, $*P < 0.001$ as com-+ MC mice. No hemorinjected with antigen i.v.

and 10 μ g Ag/site or PBS i.d., and in mice given 100 μ g Ab/site but no antigen i.v.

Figure 5. Effect of mast cell reconstitution of W/W" mice on PMN infiltration. The degree of PMN density in $+/+$, W/W^v, and mast cell-reconstituted W/W" (W/W" + MC) 8 h after challenge (100 μ g Ab/site) was evaluated in arbitrary units. The data are expressed as mean±SD, $n = 3-9, *P < 0.001$ as compared with +/+ and W/W' + MC mice.

antibody was 54.9 \pm 5.4, $n = 77$, for skin from animals pretreated with A-63162 and 54.6±12.7 arbitrary units, $n = 90$, without drug pretreatment. This constitutes a 77 and 70% reduction in mast cell granule content, respectively. These data suggest that leukotrienes released by mast cells participate in this inflammatory reaction. To confirm this possibility, the inhibitor studies were repeated in mast cell-reconstituted W/W" mice. After mast cell repletion A-63162 inhibited PMN influx (Fig. 6 C), edema (Fig. 7) and hemorrhage (Fig. 8) in W/W^{γ} mice to a similar extent as in $+/+$ mice. Therefore, leukotrienes released by mast cells seem to play an important role in immune complex-mediated inflammation.

Discussion

The present study demonstrates that mast cells significantly enhance the acute inflammatory processes, such as PMN infiltration, hemorrhage, and edema in immune complex-induced injury. The reaction seemed to be due to immune complex formation because only minor PMN influx and no edema or hemorrhage were detected in mice which were treated only with 100 μ g Ab/site for 8 h, but not with antigen. The reduction of the inflammatory parameters observed in W/W' mice was due to their mast cell deficiency and not to other abnormalities.

Figure 6. Effect of 5-lipoxygenase inhibition on PMN infiltration. Mice were pretreated with A-63162, 100 mg/kg, as described in Methods, and challenged (100 μ g Ab/site) for 8 h. PMN infiltration was evaluated in arbitrary units in $+/+(A)$ and mast cell-reconstituted W/W^v (W/W^v + MC) (C) mice, and by counting the number of PMN/section in W/W^v mice (B). The data are expressed as mean \pm SD, $n = 4$, $*P < 0.001$ as compared with the respective control (no A-63 162).

Figure 7. Effect of 5-lipox-
ygenase inhibition on edema. Mice were pre treated with A-63162, 100 mg/kg, and challenged (100 μ g Ab/site) for 8 h. The data are expressed as mean \pm SD, $n = 4, *P$ < 0.001 as compared with the respective control (no

After the adoptive transfer of mast cells (>98% purity) into the skin of W/WV mice PMN infiltration, hemorrhage, and edema were equivalent to that in congenic controls.

The participation of mast cells in reverse Arthus reaction was further substantiated by mast cell degranulation studies. There was a significant reduction in florescence intensity of mast cells stained with avidin-FITC (specific for mast cell granules) after immune complex stimulation. A 70% decrease in fluorescence intensity suggests an extensive mast cell release reaction. The method used probably underestimates the magnitude of granule loss because completely degranulated mast cells cannot be detected. Evidence of mast cell degranulation in the hamster cheek pouch due to immune complexes has been reported by Bjork and Smedegard (21). We have observed earlier in immune complex-mediated peritonitis that mast cell release preceded leukocyte influx (7).

Upon activation mast cells release several important inflammatory mediators including leukotrienes. $LTB₄$ is a potent chemotactic agent for PMN (22). LTC₄ and D_4 can cause extravasation of macromolecules (23). LTB₄ and LTC₄ also seem to act on endothelial cells to promote PMN adhesion (24,25). We tested the involvement of leukotrienes in reverse Arthus reaction by treating the mice with the 5-lipoxygenase inhibitor A-63162. This resulted in ^a decrease in PMN density, edema and hemorrhage in $+/+$ but not in W/W^v mice. The effect of A-63162 was restored upon adoptive transfer of mast cells to W/W^v mice. These results suggest that leukotrienes released by mast cells contribute to these inflammatory parameters. It is also possible that mast cell mediators acted upon other cells which in turn synthesized the eicosanoids. We think that mast

Figure 8. Effect of 5-lipoxygenase inhibition on hemorrhage. Mice were pretreated with A-63162, 100 mg/kg, and challenged (100 μ g Ab/site) for 8 h. The data are expressed as mean \pm SD, $n = 4, *P$ < 0.001 as compared with the respective controls (no A-63 162).

cells are probably the source of the leukotrienes in question for the following reasons. Only a few cell types (macrophages, monocytes, mast cells, neutrophils, and eosinophils) produce appreciable amounts of leukotrienes. The cells calling in the PMN have to be in the tissue in ^a significant number. This narrows the cell source down to mast cells and macrophages. The 5-lipoxygenase inhibition studies would then suggest that mast cells generate the leukotrienes or stimulate macrophages to synthesize these mediators. In an earlier study we investigated the interaction of mast cells and macrophages in the production of eicosanoids. IgE-sensitized mast cells were challenged with antigen in the presence of macrophages or alone. Mast cells released LTC_4 , LTB_4 , PGD_2 , and TXB_2 . The macrophages contributed PGE_2 , 6-keto-PGF_{1a}, and TXB₂, but not leukotrienes (26, 27). These observations suggest that mast cells may be the source of the leukotrienes which elicit PMN in reverse Arthus reaction. The 5-lipoxygenase inhibitor did not only block PMN infiltration and edema, two well-established functions of leukotrienes, but also hemorrhage. The latter effect of A-63162 was probably due to the decrease in PMN accumulation. Activation of PMN will lead to release of lysosomal enzymes which can cause tissue damage. PMN will discharge these enzymes when exposed to immune complexes (28).

The difference in the effect of the 5-lipoxygenase inhibitor in the two sets of mice may seem rather surprising. Various leukocytes, including macrophages and PMN, can synthesize leukotrienes (29-31); and these cells can be stimulated by immune complexes (30, 32). We interpret the data of the present study to mean that the strategic location of mast cells near blood vessels makes these cells critical for the production of leukotrienes which mediate certain early inflammatory effects. The apparent importance of mast cells in close proximity to blood vessels was also suggested by an earlier study (33). We observed in thioglycollate-induced peritonitis that the mere presence of mast cells in the peritoneal cavity of W/W" mice was not sufficient to restore the PMN infiltration to levels comparable to those in $+/+$ mice. Approximately 2 wk of mast cell reconstitution were necessary for complete normalization of the PMN influx. The gradual recovery of the PMN response paralleled the time course of mast cell migration into the tissue, presumably close to blood vessels. In the present study we actually identified mast cells in the vicinity of blood vessels in W/W^v mice after adoptive transfer of mast cells. These findings suggest that the presence of mast cells in the proper location in the tissue, i.e., near blood vessels, is essential for their function in inflammation.

In immune complex peritonitis we observed that 100 mg/ kg of A-63 162 blocked leukotriene formation by 90% (17). In the present study, 100 mg/kg of A-63162 inhibited leukocyte influx only by 40% and edema by 60%. Makino et al. (19) and Tanaka et al. (18) also observed only a partial inhibition of leukocyte influx and plasma exudation in immune complexinduced pleurisy in the rat after treatment with a 5-lipoxygenase antagonist. Therefore, mediators other than leukotrienes are also involved in these processes. Histamine has been shown to cause plasma exudation. Antihistamines will decrease plasma leakage during immune complex reactions (34). It has recently been reported that mast cells upon stimulation with a number of agents, including IgE/Ag, calcium ionophore, and PMA, will cause the induction of various cytokines, such as IL-1, IL-3, IL-4, IL-6, GM-CSF, and TNF α (35–37). These factors can modulate numerous inflammatory processes. IL-1

and TNF α are known to act on endothelial cells and to cause the expression of ICAM-1 and ELAM-1, which are adhesion molecules for leukocytes (38, 39). It has been shown by Klein and his colleagues (40) that mast cell degranulation was associated with the expression of ELAM-1 in venular endothelial cells of human skin organ culture. TNF α seemed to mediate this process because it was blocked by antibodies to this agent. IL-3, IL-4, and GM-CSF stimulate mast cell proliferation. IL-1, IL-4, and IL-6 regulate immunoglobulin production which could influence immune complex-mediated inflammation. Other function in inflammation that are modulated by some of these cytokines are eicosanoid production, clotting, wound repair, angiogenesis, and fibrosis (41). Platelet-activating factor is another mediator released by mast cells as well as other proinflammatory cells. It is known to elicit and activate neutrophils (42).

It is well established that the complement pathway is activated in immune complex reactions, and that some of the peptides generated are chemotactic for PMN. Lysosomal enzymes released by PMN can produce increases in vascular permeability. Decomplementation abolishes or reduces the inflammatory process caused by immune complexes (1). This is not in contradiction to our findings. Mast cell activation may have been initiated by complement derived peptides (43). It is also possible that immune complexes acted directly on mast cells. Their activation by antigen-antibody complexes due to binding to the Fcy receptors present on these cells has been reported (44, 45). However, mast cell deficiency did not completely inhibit PMN influx and edema. The reaction still present in W/W' mice may be due the action of complement on PMN and the vasculature and/or the activation of another cell type.

Our findings of mast cell involvement in Arthus reaction support earlier evidence implicating mast cells in the pathogenesis of diseases in which immune complexes are known to be deposited in various tissues. An increase in the number of mast cells has been observed in affected skin areas of scleroderma (46) , the synovium of rheumatoid arthritis (47) , interstitium of fibrotic lung (48), in systemic lupus erythematosus, mixed connective tissue diseases, and other related inflammatory conditions (47). In some of these conditions mast cell activation and an increase in mast cell releasability has been observed (46). In addition to these autoimmune conditions, mast cells may also contribute to inflammatory processes in diseases with immune complexes involving exogenous antigens. Immune complexes are thought to be responsible for some of the manifestations in infectious diseases such as bacterial endocarditis, disseminated gonorrhea, streptococcal infections, dengue hemorrhagic fever, viral hepatitis, dermatophytid reaction, and others (49).

In conclusion, the data presented indicate that mast cells make a significant contribution to immune complex-mediated inflammation. Mast cell deficiency greatly reduced the inflammatory response. Adoptive transfer of mast cells into W/W' mice increased PMN immigration, edema, and hemorrhage to levels comparable with those in congenic controls. The inflammatory reaction was accompanied by an extensive degranulation of mast cells. The 5-lipoxygenase inhibitor A-63162 partially blocked the expression of the three parameters evaluated in $+/+$ but not in W/W^{v} mice. Mast cell reconstitution of W/W' mice restored the effect of A-63162. Therefore, mast cells and their mediators, especially leukotrienes, are necessary for the full expression of immune complex-induced inflammation. These studies have important implications for diseases involving immune complex formation.

Acknowledgments

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