Differences in the Behavior of Cytoplasmic Granules and Lipid Bodies during Human Lung Mast Cell Degranulation

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ABSTRACT We used a morphometric and autoradiographic approach to analyze changes in specific cytoplasmic granules and cytoplasmic lipid bodies associated with human lung mast cell degranulation. Mast cells were dissociated from lung tissue by enzymatic digestion and were then enriched to purities of up to 99% by countercurrent centrifugation elutriation and recovery from columns containing specific antigen bound to Sepharose 6 MB. Degranulation was induced by goat anti-IgE. At various intervals after stimulation, parallel aliquots of cells were recovered for determination of histamine release or were fixed for transmission electron microscopy. We found that lipid bodies, electron-dense structures that lack unit membranes, were present in both control and stimulated mast cells. Autoradiographic analysis showed that lipid bodies represented the major repository of 3H-label derived from [3H]arachidonic acid taken up from the external milieu. By contrast, the specific cytoplasmic granules contained no detectable 3H-label. In addition, lipid bodies occurred in intimate association with degranulation channels during mast cell activation, but the total volume of lipid bodies did not change during the 20 min after stimulation with anti-IgE. This result stands in striking contrast to the behavior of specific cytoplasmic granules, the great majority of which (77% according to aggregate volume) exhibited ultrastructural alterations during the first 20 min of mast cell activation.

These observations establish that mast cell cytoplasmic granules and cytoplasmic lipid bodies are distinct organelles that differ in ultrastructure, biochemistry, and behavior during mast cell activation.

A previous ultrastructural analysis of human lung mast cell degranulation (4) reported that this process differed from that observed in rat peritoneal mast cells (3, 5, 14, 16, 17, 20). These workers reported that the cytoplasmic granules of human mast cells stimulated with anti-IgE underwent a series of ultrastructural changes that included the transformation from complex granule substructural patterns to a reticular pattern and, finally, the alteration of the granule contents to an amorphous state before discharge. They concluded that the crystalline patterns exhibited by human lung mast cell granules represented storage forms of materials that are solubilized before fusion of the granule membrane with the plasma membrane and discharge.

While these constitute provocative ideas, our analysis of the published photomicrographs suggested that they were based largely upon the interpretation of cytoplasmic lipid bodies as "solubilized mast cell granules." Lipid bodies are nonmembrane-bound, roughly spherical structures that may occur in a variety of cell types, but which differ from specific secretory granules in genesis and biochemical content, as well as in ultrastructure (11). Although lipid bodies have been reported to occur in human mast cells (1, 6, 23), to our
knowledge a possible role for these structures during degranulation has not previously been studied.

To analyze changes in cytoplasmic lipid bodies associated with human mast cell histamine release, we studied purified populations of mast cells isolated from human lung and then stimulated to degranulate by exposure to anti-IgE in vitro. Changes in the volumes of cell organelles were then quantitated in transmitted electron micrographs by point counting. These studies confirmed that mast cell histamine release was associated with substantial reduction in the aggregate volume of intact secretory granules. However, degranulation was associated with little or no change in the total volume of cytoplasmic lipid bodies. Furthermore, although occasional lipid bodies appeared to discharge their contents into degranulation channels that were in communication with the extracellular space, we found no evidence of transformation of specific cytoplasmic granules into lipid bodies.

MATERIALS AND METHODS

Isolation of Mast Cells

Mast cells were purified from human lung as previously described (21). Briefly, grossly normal lung fragments were obtained from pneumonectomy or lobectomy specimens removed for pulmonary neoplasm. Single cells were dispersed from lung fragments by two 30-min incubations in solutions containing pronase (2 mg/ml) and chymopapain (0.5 mg/ml) followed by two incubations in solutions containing collagenase (1 mg/ml) and trypsin (10 U/ml). Lung cell suspensions containing 6 ± 0.7% mast cells were enriched to 29 ± 3.5% mast cells (range 13.5-45%) by countercurrent centrifugation elutriation. Countercurrent centrifugation elutriation fractions with 25% mast cells were dispersed from lung fragments by two 30-min incubations in solutions containing 1.5% potassium ferrocyanide (OPF) in 0.1 M sodium phosphate buffer, washed, and then suspended in 0.1 M sodium cacodylate buffer, pH 7.4, at associated histamine.

We fixed cell suspensions by diluting them in a 10-fold excess of a mixture of 1% paraformaldehyde, 1.25% glutaraldehyde, 0.003% human serum albumin (Miles Laboratories, Inc., Elkhart, IN), and 0.1% glucose. Mast cell purity in these preparations was 84 ± 3% (range 65-99%).

Mast Cell Degranulation

Mast cells were stimulated to degranulate by exposure to antibody to IgE (goat anti-IgE, 2 μg/ml in PAG buffer plus 1 mM CaCl2 at 37°C [21]). In four separate experiments, mast cells were incubated with anti-IgE or buffer alone (controls) and fixed for ultrastructural studies 1, 3, 5, 10, 15, 20, and 30 min thereafter. To measure the extent of mast cell degranulation, the supernatants of parallel aliquots of anti-IgE-stimulated or control mast cells were analyzed for histamine content by the automated fluorometric technique of Sircar-Gupta (22). Total mast cell-associated histamine was determined by lysing aliquots of control cells with 0.4 N perchloric acid. We then calculated the percent of histamine release in control or IgE-stimulated mast cell populations according to the formula: % histamine release = histamine in supernatant + total cell-associated histamine.

Transmission Electron Microscopy

We fixed cell suspensions by diluting them in a 10-fold excess of a mixture of 1% paraformaldehyde, 1.25% glutaraldehyde, 0.025% CaCl2 in 0.1 M sodium cacodylate buffer, pH 7.4 (9). Cells were fixed for 1 h at room temperature, washed, and then suspended in 0.1 M sodium cacodylate buffer, pH 7.4, at 4°C. The cells then were transferred to microtubes, centrifuged at 1,500 g for 1 min, suspended in warm 2% agar, and centrifuged again. Some cells were postfixed as a pellet in agar for 2 h at 4°C in 2% aqueous osmium tetroxide and 1.5% potassium ferricyanide (OPF) in 0.1 M sodium phosphate buffer, pH 6.0 (9). Other pellets were postfixed for 2 h at room temperature in 2% collidine-buffered osmium tetroxide and stained en bloc with uranyl acetate (OCUB) for 2 h at room temperature (9). Cell pellets were then dehydrated in a graded series of alcohols and embedded in a propylene oxide–Epon sequence. Sections were cut with diamond knives, placed on copper grids, stained with lead citrate, and viewed in a Philips 400 transmission electron microscope.

We cut three randomly placed thin sections from separate blocks of cells exposed to anti-IgE or buffer (controls) for each time interval in each of the four experiments. In all, 117 thin sections were examined in the electron microscope. All mast cells in each grid (~100 cells) were examined at medium to high magnifications (2,800–30,000).

Quantitative Analysis of Mast Cell Degranulation

Two experiments were analyzed quantitatively, using cells from two different donors that gave similar levels of histamine release (~40%). Thin sections from each time point were examined in the same order, beginning at the top left corner of the grid and proceeding to the lower right systematically, until at least 50 recognizable mast cells (nondegranulated or anti-IgE activated) in control or stimulated populations had been photographed. All photomicrographs were printed at ×15,000, and at least 10 prints from each group were selected at random for quantitative analysis by point counting (25). Most mast cell images (e.g., Fig. 1) included part of the nucleus, and one or more cytoplasmic lipid bodies, and multiple specific cytoplasmic granules. For analysis of cell or organelle volumes by point counting (23), multipurpose screens with 1 point/cm2 were drawn onto the photomicrographs using a Hewlett-Packard HP86 computer interfaced with a Hewlett-Packard HP 7470 plotter (Hewlett-Packard Co., Palo Alto, CA). We determined the volumes of the entire cell, nucleus, cytoplasmic granules, cytoplasmic lipid bodies, degranulation channels, cytoplasm, and “other” organelles (mitochondria, rough and smooth endoplasmic reticulum, and Golgi structures and cytoplasmic vesicles). The demarcation between a degranulation channel and the plasma membrane was taken as the narrowest constriction of the channel near its apparent point of fusion with the cell membrane. The membranes of swollen cytoplasmic granules and chains of interconnected granules containing altered granule matrix material were counted as part of degranulation channels. Mean cell volume (Vc) was calculated from the measured mean profile area (A) using the Weibel and Gomez formula: $Vc = A/2.66π$ (24). Nuclear volume (Vn) was estimated as the number of points falling on the nucleus (Pn) and the total number of points falling on the cell profile (Pc), and by taking Vn = (Pn/Pc) × Vc. Cytoplasmic volume (Vcyt) was then calculated as Vcyt = Vc - Vn and was used to estimate organelle volume (Vo = [Pn/Pc] × Vcyt). The changes in cell or organelle volume during exocytosis were examined for statistical significance using Student’s t-test.

Autoradiographic Analysis of Mast Cells Labeled with [3H]Arachidonic Acid

We recently reported that several kinds of inflammatory cells, including macrophages, granulocytes, lymphocytes, and mast cells, incorporate 3H-label derived from [3H]arachidonic acid (3H-AA) predominantly into cytoplasmic lipid bodies (6). Because human lung mast cell degranulation is associated with the generation of oxidative products of AA metabolism (19), we used electron microscopic autoradiography to determine whether mast cell activation resulted in alterations of the distribution of 3H-AA-derived label in mast cells allowed to incorporate this precursor over an 18 h period of labeling in vitro.

**Labeling with [3H]-AA:** Mast cells (6–20 x 106, 92 ± 2% mast cells) were washed with PAG buffer and then cultured for 18 h in RPMI 1640 medium with 25 mM Hepes, 1 mM glutamine, 10% fetal calf serum, and 100 μg/ml Kanamycin in a volume of 1.5–2.5 ml in 16 mm flat-bottomed tissue culture wells in humidified 5% air/5% CO2. Immediately before addition of cells, 50 or 100 μCi [3H]-AA (NET-398, New England Nuclear, Boston, MA) was placed in each culture well and solvent evaporated under a stream of nitrogen.

**Stimulation of Mast Cell Degranulation:** After 18 h labeling with [3H]-AA, mast cells were washed 3 x with PAG, placed in 0.1 ml of PAG plus 1 mM CaCl2 at 37°C, and then were stimulated with ionophore A23187 (1 μg/ml), goat anti-IgE (2 μg/ml) or, as a control, normal goat serum. After incubation for 15–30 min, cells were diluted in PAG, centrifuged in a microfuge, and the supernatant removed in order to quantitate release of 3H species and histamine. Radioactivity was quantitated by liquid scintillation spectroscopy in a Triton X-100–toluene scintillation cocktail. Histamine was quantitated by an automated fluorometric method (22). Parallel aliquots of 3H-labeled cells were fixed and processed for EM or analyzed for radioactive lipid content by thin layer chromatography (19).

**Abbreviations used in this paper:** AA, arachidonic acid; BPO-FLYS, N-α-benzylpenicilloyl-N-α-formyl-L-lysine; EM, electron microscopy; OCUB, osmium collidine uranyl en bloc; OPF, osmium tetroxide-potassium ferricyanide; PAG buffer, 25 mM piperonazine-N,N′-bis-2-ethanesulfonic acid, 140 mM NaCl, 6 mM KCl, 0.003% human serum albumin, and 0.1% glucose.
**FIGURE 1** Control mast cell illustrating cytoplasmic granules containing scroll-like inclusions and several large cytoplasmic lipid bodies (arrows). OCUB × 6,000.

**ANALYSIS OF ULTRASTRUCTURAL AUTORADIOGRAPHY:** Electron microscopic autoradiographs were prepared by looping 100-nm thick sections on copper grids with Ilford 1-4 emulsion (Ilford Ltd., Ilford, Essex, England). These were then exposed at 4°C, developed in Microdol-X and stained in lead (6). Chi-square analysis was performed on electron microscope prints magnified to ×16,200 to determine whether an association existed between cell organelles and silver grains (26). A standard grid composed of circles (r = 260 nm, approximately the size of specific mast cell granules and smaller than all other structures analyzed) with a center-to-center distance of 2 cm was employed for morphometric analysis (grain counting).

**RESULTS**

**Control Mast Cells**

Unlike rat peritoneal mast cells, whose cytoplasmic granules are homogeneously electron dense and therefore present a uniform ultrastructural appearance (5, 17, 20), human mast cell granules may exhibit a variety of different ultrastructural patterns (1, 2, 4, 13, 15, 18, 23). In our preparations, the most common of these consisted of granules with one to many cylindrical scroll-like inclusions (Fig. 1, Fig. 2, A and B). The scrolls themselves varied in overall electron density (Fig. 2A), and some scrolls contained central deposits of homogeneously electron dense matrix (Fig. 2, A and B). Because scrolls with central densities appeared in both control and in anti-IgE stimulated mast cells, we do not consider this finding evidence of an early stage of granule dissolution. Some granules had crystalline (Fig. 2C), particulate (Fig. 2D) or reticular (Fig. 3, A–C) patterns. Although the reticular pattern has been attributed to dissolution of highly ordered granule contents (4), we observed a small number (<5%) of granules with this ultrastructure in both control and stimulated mast cell preparations. A few individual granules of occasional (<5%) control mast cells were swollen and contained separated or distended scrolls and/or scant particulate electron dense material (cf. in Fig. 3, D and E). These changes were similar to those characteristic of the majority of granules at early intervals after mast cell stimulation with anti-IgE; their presence in a small minority of control mast cells may have reflected prior stimulation or secretory activity, occurring either in vivo or during the cell isolation procedure.

In addition to specific cytoplasmic granules, control mast cells also contained cytoplasmic lipid bodies (Fig. 1 and 3F). Lipid bodies are roughly spherical, variably dense structures that are often enmeshed in cytoplasmic filaments and that, on average, are larger and less numerous than the specific cytoplasmic granules. They also differ from granules in that they lack a unit membrane (Fig. 3F). They have been identified in the mast cells of human adenoids (1) and in human nasal and bronchial mucosal mast cells (23), and are seen regularly as a component of the cytoplasm of human lung mast cells (6). We have never observed them in human skin mast cells (A. M. Dvorak, unpublished data).
Mast Cells Stimulated by Anti-IgE

Activation of mast cells by anti-IgE resulted in a characteristic set of ultrastructural alterations whose final effect was to expose the cytoplasmic granule interiors to the external milieu. Although these changes constituted a continuum, and individual mast cells occasionally demonstrated several of them simultaneously, they could be ordered into the following sequence: (a) Cytoplasmic granule swelling and fusion. As early as 1 min (Fig. 4A) and prominently by 3 min after exposure to anti-IgE, cytoplasmic granules appeared swollen and, in favorable planes of section, clearly demonstrated areas of membrane fusion with adjacent granules. Initially, presimulation matrix configurations such as scrolls, particles, crystals, and occasional reticular patterns were still identifiable in these granules, although their substructural elements progressively separated from one another. (b) Formation of chains of interconnected granules, which enlarged to form tortuous cytoplasmic channels containing altered granule matrix material (Fig. 4B). This change was observed rarely as early as 1 min after administration of anti-IgE, and was marked by 3 min. (c) Fusion of channels with the plasma membrane. Although open channels sometimes contained altered granule matrix material, neither membrane-free granules nor identifiable granule constituents (such as scrolls, crystals, particles, or reticular structures) was ever observed either free in the extracellular medium or attached to the surface of degranulating mast cells. This is in contrast to the degranulation patterns of rat peritoneal mast cells (16, 20), and human (8) or guinea pig basophils (7), where recognizable granule structures are extruded to the exterior. As noted by many others (1, 4, 15, 16, 23), mast cell degranulation was associated with striking increases in cytoplasmic filaments, particularly in the paranuclear region. In our material, many filaments were intimately associated with cytoplasmic lipid bodies. Despite marked reductions in the number of cytoplasmic granules during anti-IgE induced secretion, the numbers of cytoplasmic lipid bodies changed little.
or not at all (see below). At late stages of degranulation, some lipid bodies appeared close to degranulation channels (Fig. 4B) or, rarely, the plasma membrane. In a few instances we observed apparent discharge of osmiophilic content from lipid bodies into degranulation channels (Fig. 5), but we never observed discharge of lipid bodies from the cell surface. We also found no evidence of transformation of membrane-bound cytoplasmic granules into lipid bodies, such as organelles of intermediate ultrastructural appearance.

Quantitative Changes in Mast Cell Organelles during Degranulation

The two separate experiments analyzed gave very similar results that were pooled for presentation in Fig. 6. Mast cells responding to stimulation with anti-IgE exhibited a 77% diminution in the volume of unaltered cytoplasmic granules (Fig. 6). Part of this volume was shifted into that of the degranulation channels, which we classified as “intracellular”
(although in strict topological terms, channels that have fused with the plasma membrane may be considered to be outside the cell). The remainder of the loss in granule volume probably accounted for the small (but statistically insignificant) reduction in total cellular volume associated with degranulation (Fig. 6). Degranulation did not result in a significant change in the volume of cytoplasmic lipid bodies (Fig. 6). Quantitative analysis of unstimulated mast cells used as control preparations in these two experiments detected no significant changes in cell or organelle volumes 5 or 20 min after incubation in buffer (data not shown).

Distribution of $^{3}$H-AA Label in Control and Degranulated Mast Cells

Morphometric analysis of EM autoradiographs confirmed the results of our previous experiments (6), showing that nearly all $^{3}$H-label derived from $^{3}$H-AA incorporated by cultured human mast cells during an 18 h incubation in vitro was localized to cytoplasmic lipid bodies (Table I, Fig. 7A). Lipid bodies were the only cellular structure exhibiting statistically significant labeling; little or no label was associated with cell membranes. In this experiment, anti-IgE–stimulated mast cells induced ultrastructural changes similar to those described above, including the close approximation of mast cell lipid bodies with degranulation channels (Fig. 7C). Although lipid body contents and autoradiographic grains (Fig. 7B and C) were sometimes observed within degranulation channels, mast cell degranulation appeared to be associated with relatively little net loss of lipid body content. Thus, stimulation with anti-IgE provoked the discharge of only a small fraction (mean of five experiments, 3%; range 2–12%) of total cell-associated $^{3}$H-lipids (19). In addition, EM autoradiography and morphometric analysis failed to document either statistically significant autoradiographic labeling of degranulation channels (Table I) or significant changes in the overall volume of lipid bodies in mast cells exhibiting degranulation (Fig. 6). In contrast to the small amount of total $^{3}$H-lipid products released by stimulated mast cells in these experiments, there was substantial release of the granule-associated mediator histamine (~75% release in three experiments with anti-IgE, with spontaneous release in the absence of anti-IgE of <5%).

DISCUSSION

We found that mast cells derived from human lung, like those identified in other human tissues (1, 23), contained both specific granules and lipid bodies. As in other cells (11), mast cell lipid bodies appear in electron micrographs as round, osmiophilic, nonmembrane–bound structures. In both control and degranulating mast cells, lipid bodies were distributed predominantly near the nucleus, generally in intimate association with cytoplasmic filaments. In addition to differing from cytoplasmic granules in ultrastructure, mast cell lipid bodies also differ from specific granules in size, number, and apparent growth mechanism.

During degranulation, cytoplasmic granules exhibited a

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FIGURE 5 Mast cells 20 min after stimulation (42% histamine release) showing discharge of lipid bodies into degranulation channels. In (A and B), multiple lipid bodies (arrows) appear to be discharging into degranulation channels that also contain agar, indicating communication with the extracellular space. (C and D) Higher power electron photomicrographs of lipid bodies discharging into degranulation channels. The arrow in D indicates an unaltered scroll-filled granule. (A–D) OPF. (A and B) × 9,500; (C) × 53,000; (D) × 51,000.

A series of characteristic changes including swelling, intergranule fusion to form degranulation channels, and fusion of channels with the plasma membrane. A similar sequence has been proposed by Trotter and Orr (18, 23), who examined mast cells in biopsies of human nasal and bronchial mucosa. These qualitative changes were accompanied by a precipitous reduction in the number of intact cytoplasmic granules and by substantial histamine release. Throughout the early stages
and 5 or 20 min after stimulation with anti-lgE in vitro. Electron micrographs were printed at ×15,000 and point counting was used to calculate the cell volume and the volumes of the nucleus, the specific granules, the degranulation channels, the lipid bodies, the other organelles (mitochondria, rough and smooth endoplasmic reticulum, Golgi structures, and vesicles), and the residual cytoplasm. Cells from two experiments that gave similar levels of mast cell stimulation, as judged by histamine release (see inset), were analyzed. At least 10 prints were analyzed at each time point in each experiment. The two experiments gave similar results which were pooled and are presented as M ± SE. Significant changes in organelle volume associated with degranulation are indicated (Student’s t test). Morphometric analysis of control mast cells exposed to goat serum instead of anti-lgE disclosed no change in this volume of the cell or any of the organelles over the duration of the experiment (data not shown).

**Mast cells were labeled with 3H-AA for ~18 h in vitro, then fixed for electron microscopy**. Electron micrographs were printed at ×15,000 and point counting was used to calculate the cell volume and the volumes of the nucleus, the specific granules, the degranulation channels, the lipid bodies, the other organelles (mitochondria, rough and smooth endoplasmic reticulum, Golgi structures, and vesicles), and the residual cytoplasm. Cells from two experiments that gave similar levels of mast cell stimulation, as judged by histamine release (see inset), were analyzed. At least 10 prints were analyzed at each time point in each experiment. The two experiments gave similar results which were pooled and are presented as M ± SE. Significant changes in organelle volume associated with degranulation are indicated (Student’s t test). Morphometric analysis of control mast cells exposed to goat serum instead of anti-lgE disclosed no change in this volume of the cell or any of the organelles over the duration of the experiment (data not shown).

**FIGURE 6 Quantitative analysis of human lung mast cell cellular and organelle volumes before and 5 or 20 min after stimulation with anti-lgE in vitro.** Electron micrographs were printed at ×15,000 and point counting was used to calculate the cell volume and the volumes of the nucleus, the specific granules, the degranulation channels, the lipid bodies, the other organelles (mitochondria, rough and smooth endoplasmic reticulum, Golgi structures, and vesicles), and the residual cytoplasm. Cells from two experiments that gave similar levels of mast cell stimulation, as judged by histamine release (see inset), were analyzed. At least 10 prints were analyzed at each time point in each experiment. The two experiments gave similar results which were pooled and are presented as M ± SE. Significant changes in organelle volume associated with degranulation are indicated (Student’s t test). Morphometric analysis of control mast cells exposed to goat serum instead of anti-lgE disclosed no change in this volume of the cell or any of the organelles over the duration of the experiment (data not shown).

**TABLE I**

| Organelle       | Control mast cells | Degranulated mast cells |
|-----------------|-------------------|-------------------------|
|                 | percent observed  | percent expected | X² | percent observed | percent expected | X² |
| Lipid bodies    | 33.2              | 41.1                | 51.2 | 26.4           | 2.7              | 75.0 |
| Granules        | 23.1              | 28.3                | 0.04 | 10.8           | 11.6             | 0.05 |
| Degranulation channels | 21.0 | 23.8 | 0.14 |
| Cytoplasm       | 36.1              | 43.2                | 0.03 | 33.4           | 39.0             | 0.02 |
| Nucleus         | 5.8               | 16.2                | 0.41 | 5.7            | 13.9             | 0.35 |
| Extracellular   | 1.9               | 8.3                 | 0.72 | 2.7            | 8.9              | 0.48 |

* Mast cells were labeled with 3H-AA for ~18 h in vitro, then fixed for preparation of EM autoradiographs (see Materials and Methods) 15 min after exposure to 2 μg/ml anti-lgE (degranulated mast cells) or equivalent normal goat serum (control mast cells). The percent of actual (i.e., observed) organelle-associated silver grains was compared to the percent of grains expected to occur over these structures by chance, assuming a uniform distribution of grains over the prints. The data were derived from 14 control mast cells (with 386 grains) and 12 degranulated mast cells (with 205 grains). Too few grains were associated with cellular membranes to permit formal statistical analysis. The labeling of lipid bodies in control or degranulated mast cells was highly significant (P < 0.01) by the X² test. No other structure was significantly labeled (all P values >0.05).

sent the major repository of 3H-label derived from 3H-AA and other 3H-fatty acids (6) incorporated by mature human lung mast cells in vitro. By contrast, the cytoplasmic granules of these mast cells contained no detectable 3H-label. Together with our other findings, these data strongly support the view (1, 23) that lipid bodies and specific mast cell granules represent distinct organelles, not different functional stages of the same entity.

However, we agree with Caulfield et al. (4) that initiation of mast cell granule matrix solubilization may be among the earliest morphologic events in anti-lgE induced mast cell degranulation, perhaps even preceding fusion of granule membranes with the plasma membrane. In this paper, the earliest ultrastructural evidence of impending degranulation was granule swelling with widening and separation of scroll-like or other granule matrix substructural patterns. It is not clear, however, exactly how this initial solubilization is effected and whether it requires exposure of granule contents to extracellular fluid. In our experience (A. M. Dvorak, unpublished data), the great majority of granules exhibiting swelling and matrix dissolution at early intervals after stimulation do not admit the extracellular tracer, cationized ferritin, when it is incubated with fixed cells. However, exposure of the cytoplasmic granule matrix to the external medium theoretically may be effected by channels too narrow or transient to be identified readily with cationized ferritin. Alternatively, the granule interiors may come into contact with solvent transported from the extracellular via traffic of cytoplasmic vesicles (10, 12). In view of these uncertainties, we feel that it is premature to conclude precisely when, in the sequence of human lung mast cell degranulation, the cytoplasmic granule interiors become exposed to the extracellular medium.

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FIGURE 7 (A–C) Electron microscopic autoradiographs of purified human lung mast cells labeled with 3H-AA in vitro and then exposed to buffer (A) or to anti-lgE (B and C) for 15 min before fixation. The control cell (A) contains heavily labeled lipid bodies, which appear larger and much darker than the more numerous specific cytoplasmic granules. The cells stimulated with anti-lgE (B and C) exhibit labeled degranulation channels that contain altered granule material. N, nucleus. (A–C) OCUB. (A) × 17,000; (B) × 21,500; (C) × 17,000.

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