Primary Murine Airway Smooth Muscle Cells Exposed to Poly(I,C) or Tunicamycin Synthesize a Leukocyte-adhesive Hyaluronan Matrix*

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Asthmatic attacks often follow viral infections with subsequent airway smooth muscle cell proliferation and the formation of an abnormal hyaluronan extracellular matrix with infiltrated leukocytes. In this study, we show that murine airway smooth muscle cells (MASM) treated with polyinosinic acid-polycytidylic acid (poly(I,C)), a double-stranded RNA that simulates a viral infection, synthesize an abnormal hyaluronan matrix that binds leukocytes (U937 cells). Synthesis of this matrix is initiated rapidly and accumulates linearly for ~10 h, reaching a plateau level ~7-fold higher than control cultures. MASM cells treated with tunicamycin, to induce endoplasmic reticulum stress, also rapidly initiate synthesis of the abnormal hyaluronan matrix with linear accumulation for ~10 h, but only reach a plateau level ~2-fold higher than control cultures. In contrast to poly(I,C), the response to tunicamycin depends on cell density, with pre-confluent cells producing more abnormal matrix per cell. Furthermore, U937 cell adhesion per hyaluronan content is higher in the sparse matrix produced in response to tunicamycin, suggesting that the structure in the poly(I,C)-induced matrix masks potential binding sites. When MASM cells were exposed to tunicamycin and poly(I,C) at the same time, U937 cell adhesion was partially additive, implying that these two toxins stimulate hyaluronan synthesis through two different pathways. We also characterized the size of hyaluronan produced by MASM cells, in response to poly(I,C) and tunicamycin, and found that it ranges from 1500 to 4000 kDa, the majority of which was ~4000 kDa and not different in size than hyaluronan made by untreated cells.

Asthma, a chronic inflammatory disease of the airways (1, 2), is characteristically accompanied by increased airway hyper-responsiveness to various stimuli (such as viruses, allergens, and pollutants) (3–6). Other major features include proliferation of airway smooth muscle cells (7), deposition of an extensive hyaluronan-rich extracellular matrix by these cells into the airway submucosa (8–11), and excessive invasion of the airway mucosa and submucosa by inflammatory cells (mainly T cells of the Th-2 phenotype, eosinophils, macrophages, and mast cells) (12–15).

Hyaluronan is a large glycosaminoglycan in which the disaccharide (glucuronic acid-β1,3-N-acetylglucosamine-β1,4-) is repeated several thousand times (16). This major constituent of extracellular matrices is generally synthesized by one or more of the three eukaryotic hyaluronan synthases (HAS) (17) (Has1, -2, and -3) at the cytosolic side of the plasma membrane with simultaneous extrusion into the extracellular space. Outside the cell, hyaluronan interacts with both cell surface and extracellular hyaluronan-binding proteins (18), providing the tissue with a structural scaffold. Hyaluronan also has an essential role in many physiological and pathological processes, including cell migration, morphogenesis, tissue regeneration, wound repair, and tumor cell growth and invasion (19). Cells often interact with hyaluronan-based matrices through the cell surface hyaluronan receptor, CD44 (20), which is present in airway smooth muscle cells, both in vivo and in vitro (21), and is also present on all leukocyte populations (22).

In asthma, the accumulation of excess hyaluronan in the submucosal tissue can lead to severe airway obstruction and death (23). Hyaluronan accumulates in the airway submucosa (24), around the smooth muscle bundles (24), and in the bronchoalveolar lavage fluid (11, 25, 26). A murine bleomycin model by Teder et al. (27) has demonstrated that excess amounts of hyaluronan must be removed from the airway submucosa by monocytes/macrophages in a CD44-dependent manner to resolve lung inflammation.

Respiratory viral infections are a major cause of asthma exacerbation and are accompanied by leukocyte infiltration and inflammation of the airways (28, 29). Viral infections account for ~80% of asthma attacks in children (30) and ~70% in adults

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Rhinovirus was detected by in situ hybridization (32) in both bronchial epithelial and underlying submucosal cells in biopsies obtained from the lower airways, and it is likely from the histology that their localization was mesenchymal, namely fibroblasts and/or smooth muscle cells. Furthermore, viral infection in the airway epithelium in asthmatics induces cell death and the desquamation of the epithelial cell layer (33), which then could provide direct viral access to the underlying mesenchymal cells, including the SMCs.

Generally, viruses have two major effects on infected cells. After infection, double-stranded RNA-dependent protein kinase (PKR), a cytosolic and nuclear protein, acts as an intracellular receptor for double strand RNA produced by viral replication. PKR has a key role in limiting viral replication by inactivating the critical translation initiation factor eIF2 by phosphorylation of its α subunit. In the course of a viral infection, large amounts of viral proteins are synthesized and accumulate in the endoplasmic reticulum (ER) (34). Human cytomegalovirus infection has been shown to activate ER resident transmembrane protein kinase (PERK) or PKR-like ER-localized eIF2α kinase, an ER-resident membrane protein that transmits the ER stress signal by phosphorylating eIF-2α at serine 51 (35). This causes translational attenuation and transcriptional up-regulation of genes encoding proteins that facilitate folding or degradation of proteins (35). Thus, PKR and PERK may coordinate to control viral replication.

Both of the above-mentioned pathways of viral infections can trigger SMCs to deposit hyaluronan that is adhesive for leukocytes. For example, human colon SMCs infected with a virus or treated with polynosinic-polycytidylic acid (poly(I,C)), a double strand RNA that mimics viral genetic material, induce the synthesis of hyaluronan structures that resemble “cables” and readily promote leukocyte adhesion (U937 monocytic cell line) in a CD44-dependent manner (36). Human colon SMCs treated with tunicamycin, or other ER stress inducers, also produce similar adhesive hyaluronan structures. Sections from the colon of patients with inflammatory bowel disease or Crohn disease exhibit abnormal hyaluronan matrices with embedded cells exhibiting ER stress as indicated by KDEL staining (37).

Because the airway inflammation associated with asthma involves the accumulation of a hyaluronan-rich matrix following viral infection, and because viral infection involves ER stress (38), we subjected mouse airway smooth muscle cells (MASM) to a viral mimic (poly(I,C)) and a toxin known to induce ER stress (tunicamycin). Both stressors induced the MASM cells to deposit a hyaluronan-rich extracellular matrix similar to that observed in asthmatic airways. This high molecular weight (≈4000 kDa) hyaluronan matrix permitted significant leukocyte adhesion similar to the accumulation of leukocytes in asthmatic airways.

EXPERIMENTAL PROCEDURES

Animals and Animal Care—21-Day-old female BALB/c mice were purchased from Charles River Laboratories (Wilmington, MA) and housed under conditions of constant temperature with 12-h light/dark cycles. Food and water were available ad libitum. The mice were sacrificed by administering Nembutal (Ovation Pharmaceuticals, Deerfield, IL) at 0.125 mg/g of mouse weight. All protocols with the animals were approved by the Cleveland Clinic Institutional Animal Care and Use Committee.

Primary Cell Culture—After sacrifice, the tracheas were excised and placed in Ham’s F-12 nutrient medium with 50 units/ml penicillin, 50 μg/ml streptomycin, and then maintained at 4 °C on ice. Under a dissecting microscope, the esophagus and surrounding connective tissue were removed. The tracheas were cut longitudinally with a scalpel to expose the lumen and then transferred to 0.15% Pronase (Roche Applied Science) in Ham’s F-12 nutrient medium with 50 units/ml penicillin, 50 μg/ml streptomycin and incubated at 4 °C overnight. The next day, fetal bovine serum (FBS) was added to the tracheas (final concentration 10%) to inhibit further protease degradation. The medium, containing the released cells, was transferred to another tube for isolation of airway epithelial cells for use in a separate study. The tracheas were then brushed with a cotton swab to remove the remaining adherent epithelial cells, cut into small pieces (~30 per trachea), and transferred to a 100-cm² tissue culture dish for attachment and outgrowth. Approximately 50 μl of DMEM/F-12 with 100 units/ml penicillin, 100 μg/ml streptomycin, 0.25 μg/ml amphotericin B, and 10% FBS was applied to each piece followed by incubation at 37 °C, 5% CO₂, and 100% humidity overnight. The next day, 12 ml of the same medium was applied to each 100-cm² dish that contained the adherent pieces followed by incubation as before. Four days later, SMC outgrowth was apparent, and the residual tracheal pieces were removed with forceps. We found that waiting an additional day (5 days total) diminished SMC purity via epithelial outgrowth. The cells were left to multiply for 2 more days, after which they were trypsinized and plated into a 175-cm² flask. Confluency was reached within about 3–5 days. Typically, we processed 12 tracheas per experiment. Experiments were done on the second passage, which was split 1:2 from the first passage. These cells were positive for the smooth muscle cell marker “caldesmon” as determined by confocal microscopy (C4562, Sigma) (not shown). Additionally, no epithelial contamination was observed by phase-contrast microscopy (not shown). The lack of a suitable marker for myofibroblasts to distinguish them from smooth muscle cells does not permit such a fine distinction, but this is likely to be of minor consequence. For convenience, we refer to these cells as MASM cells.

Experimental Culture—For experiments involving poly(I,C) (P0913, Sigma), MASM cells were split 1:2 from passage 1 into 24-well plates. For experiments involving tunicamycin (T7765, Sigma), the splitting ratio was 1:4. The rationale for this approach is described in Fig. 8, and it is related to the observation that the response of the MASM cells to tunicamycin is dependent on cell confluency, whereas the poly(I,C) response is independent of this variable. As a rule, poly(I,C) (10 μg/ml) and tunicamycin (5 μg/ml) were applied to the cells 2 days after splitting. The cells were routinely cultured in DMEM/F-12 with 10% FBS, but the FBS content was dropped to 5% for the treatments because a lower serum content facilitates analysis of hyaluronan in the conditioned medium and because Fig. 3 confirms that this change in serum does not affect our other parameters. Treatment duration was 18 h, because our time course studies showed that both tunicamycin and poly(I,C)
induced peak responses for our measured parameters at this time (Figs. 4 and 5). Treatment volume was 0.5 ml per well.

**Leukocyte Adhesion Assay**—Ordinarily, 30 million leukocytes (U937 cells, American Type Tissue Culture, Manassas, VA) were labeled with 100 μCi of sodium 51Cr in 2 ml of RPMI 1640 medium, 50 units/ml penicillin, and 50 μg/ml streptomycin with 5% FBS for 1 h at 37 °C. Afterward, the leukocytes were washed with 10 ml of washing medium (DMEM/F-12 with 5% FBS, 100 units/ml penicillin, 100 μg/ml streptomycin, 0.25 μg/ml amphotericin B, kept at 4 °C) and pelleted at 300 × g for 5 min. This washing step was repeated four times. Afterward, the U937 cells were resuspended in 15 ml of washing medium and incubated on ice for 10 min to slow down their metabolism. Three 100-μl aliquots of the labeled U937 cells were set aside in three separate vials (each containing 200,000 cells) for counting on a scintillation counter to determine counts/min values for a known number of cells. The MAS cells were incubated at 4 °C for 30 min and then gently washed with washing medium one time. The labeled U937 cells were applied to the MAS cells at 1 million per well of a 24-well plate (in 0.5 ml) and incubated at 4 °C for 30 min. Unbound U937 cells were removed by gentle agitation in a circular motion and aspirated. The plates were washed gently two more times. Then half the plate wells were treated with 0.5 turbidity reducing units/ml Streptomyces hyaluronidase (100740-1, Seikagaku, East Falmouth, MA) at 0.5 ml/well for 5 min with occasional agitation. The other half of the plate wells were washed two more times. After the 5-min hyaluronidase digestion, the released cells were aspirated, and these wells were washed one more time. The total number of washes for each well was five. The number of leukocytes bound was determined by lysing the cells with 200 μl of 1% Triton X-100 in distilled water and counting a portion on a scintillation counter.

**Preparation of Hyaluronan for Fluorescent Derivatization with 2-Aminoacridone**—This method (fluorophore-assisted carbohydrate electrophoresis (FACE)) for the quantification of hyaluronan has been partially described elsewhere (39). The conditioned medium (0.5 ml) from each well of a 24-well plate was transferred to a 1.5-ml centrifuge tube containing 50 μl of proteinase K (25530015, Invitrogen) at 1 mg/ml in 100 mM ammonium acetate, pH 7.0, with 0.001% lauryl sulfate) was applied, followed by 4 °C for 30 min and then gently washed with washing medium one time. The labeled U937 cells were applied to the MAS cells at 1 million per well of a 24-well plate (in 0.5 ml) and incubated at 4 °C for 30 min. Unbound U937 cells were removed by gentle agitation in a circular motion and aspirated. The plates were washed gently two more times. Then half the plate wells were treated with 0.5 turbidity reducing units/ml Streptomyces hyaluronidase (100740-1, Seikagaku, East Falmouth, MA) at 0.5 ml/well for 5 min with occasional agitation. The other half of the plate wells were washed two more times. After the 5-min hyaluronidase digestion, the released cells were aspirated, and these wells were washed one more time. The total number of washes for each well was five. The number of leukocytes bound was determined by lysing the cells with 200 μl of 1% Triton X-100 in distilled water and counting a portion on a scintillation counter.

**Page**—Polycrylamide gels were cast using mini-Protein II system (Bio-Rad). The gel composition was made of 20% acrylamide (37:5:1, Bio-Rad), 40 mM Tris-acetate, pH 7.0, 2.5% glyc erol, 10% ammonium persulfate, and 0.1% TEMED). Gels were cast with 0.75-mm spacers and 15-well combs, and the wells were rinsed with electrophoresis running buffer (1× Tris borate/EDTA) by pipetting 150 μl of the running buffer into the wells six times. The gels were cooled to 4 °C, after which 2 μl of sample was added to each well. Electrophoresis was accomplished at 300 V (constant voltage) for ~1 h and 20 min. The gel plates were washed with distilled water, and the gels were imaged while in their glass plates.

**Digital Imaging and Data Analysis**—After electrophoresis, the gels (in their glass plates) were placed on a UV transilluminator (Ultra Lum, Claremont, CA) and illuminated at 365 nm. Imaging was done on a Quantix CCD camera (Photometrics, Tucson, AZ), and hyaluronan was quantified using Gel-Pro Analyzer version 3.0 (Media Cybernetics, Silver Spring, MD). All statistics (Student’s t tests) were done using KaleidaGraph version 3.6 (Synergy Software, Reading, PA).

**Immunohistochemistry**—MASM cells were fixed in 10% formalin at 4 °C overnight, rinsed with Hanks’ balanced salt solution three times, and permeabilized with 0.1% Triton X-100 in pre-cooled Hanks’ balanced salt solution at 4 °C for 5 min. Hyaluronan was visualized using confocal microscopy by applying a biotinylated hyaluronan-binding protein (product 385911, StressGen, Victoria, British Columbia, Canada) antibodies were conjugated to Alexa Fluor® 594 secondary antibody (product A21203, Invitrogen).

**Western Blot**—MASM cells were grown to pre-confluency (about 80%) in 6-well plates and treated with tunicamycin or poly(I:C) as described earlier. At the time of harvest, the conditioned medium was removed, and the cells were washed one time in PBS. 0.3 ml of M-Per® mammalian protein extraction reagent (Pierce, 78501) containing Halt™ protease inhibitor mixture (Pierce, 78415) was added to each well and incubated for 5 min at room temperature. The protein extraction was transferred to a 1.5-ml tube, and the protein content was quan-
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tified using the Micro BCA™ protein assay kit (Pierce, 23235),
yielding about 0.5 mg/ml for each sample. 10 μg of sample
protein was added to each well of a 10-well polycracylamide gel
(Invitrogen, NP0321BOX) and blotted to nitrocellulose (Li-
Cor, 926-31090, Lincoln, NE). The blot was blocked for 1 h
(Li-Cor, 927-40000) and simultaneously probed with antibo-
dies against KDEL at 1:500 (Affinity Bioreagents, PA1-013,
Golden, CO) and β-actin at 1:2,000 (Sigma, A5441) in the
blocking buffer with 0.1% Tween 20 for 1 h. The blots were
washed five times in phosphate-buffered saline with 0.1%
Tween 20 and simultaneously probed with IRDYE secondary
antibodies (Li-Cor, 926-32211 and 926-32222) at 1:15,000 dilu-
tion in blocking buffer with 0.1% Tween 20 and 0.01% lauryl
sulfate for 45 min. The blots were washed as before and imaged
on an Odyssey infrared imaging system (Li-Cor).

Size Determination of Hyaluronan—Pre-confluent (~90%)
MASM cells, cultured in 175-cm² flasks, were incubated with
poly(I,C) (10 μg/ml), tunicamycin (5 μg/ml) or untreated for
18 h as described under “Experimental Culture.” Afterward,
the cells were rinsed two times with 25 ml of room temperature
PBS and digested with 8 ml of proteinase K (25530015, Invitro-
gen) at 0.1 mg/ml in 100 mM ammonium acetate, pH 7.0, with
0.01% lauryl sulfate, followed by a 4-h incubation at 60 °C with
occasional (every 30 min) agitation of the flasks to ensure all
surface area was covered by the digesting solution. The extracts
were transferred to 40-ml centrifuge tubes and precipitated with
32 ml of absolute ethanol at −20 °C overnight. The sam-
ple was centrifuged at 13,200 × g for 20 min (4 °C), discarding
the supernatant. The samples were washed with 40 ml of 75%
absolute ethanol (−20 °C) and centrifuged as before. The pellet
was allowed to air-dry at room temperature for 1 h and resus-
pended in 1 ml of 100 mM ammonium acetate, pH 7.0, for an
additional hour. Residual proteinase K was heat-inactivated by
applying the samples to a boiling water bath for 10 min. After a
brief spin (1 min, 13,200 × g), the samples were transferred to
1.5-ml centrifuge tubes, (using a 9-inch Pasteur pipette) and
vacuum concentrated to 100 μl. DNase (3 μl at 2 units/μl, 2224,
Ambion, Austin, TX) and RNase (3 μl at 1.28 μg/μl, 109169,
Roche Applied Science) were added to the samples and incubated
at 37 °C overnight followed by precipitation with 4 vol-
umes of absolute ethanol at −20 °C overnight. The samples
were centrifuged for 10 min at 4 °C and 13,200 × g, discarding
the supernatant. The pellets were washed with 500 μl of 75%
ethanol (−20 °C), centrifuged as before, and permitted to air
dry for 20 min. The pellet was resuspended in 19 μl of 100 mM
ammonium acetate, pH 7.0, for 20 min and placed on a boiling
water bath for 5 min to inactivate the DNase and RNase
enzymes. A-(2→3),6,8,9)-Neuraminidase (1 μl at 0.005 unit/μl,
N8271, Sigma) was added to each of the samples to degrade
sialic acid containing mucin oligosaccharides. For some exper-
iments, the samples were divided into halves (10 μl each), and
the hyaluronan was digested with Streptomyces hyaluronidase
(1 μl at 0.2 turbidity reducing unit/μl) to confirm the purity of
hyaluronan on the agarose gel. These digests were incubated at
37 °C overnight, inactivating the enzymes on a boiling water
bath the next day for 5 min. The samples were lyophilized and
resuspended with Tris acetate/EDTA (TAE; 4 mM Tris, 50 mM
EDTA, pH 8.26) in the same volume as in the previous digests
(either 10 or 20 μl). These samples were analyzed on 1% agarose
gels (50041, Seakem HGT, Cambrex, Walkersville, MD) that
were pre-run at 80 V (constant) for 6 h to remove impurities,
using TAE as the electrophoresis buffer. Loading buffer (0.2%
bromphenol blue in 85% glycerol; 2 μl for each 10-μl sample)
was added to each sample, followed by electrophoresis at 100 V
for 1 h and 11 min (0.5 cm thick, 11 × 8-cm gel). Hyaluronan
molecular weight standards were also used in this analysis
(Select-HATM LoLadder, HiLadder, and Mega-HATM Ladder;
Hyalose, Oklahoma City, OK). Afterward, the gels were equili-
brated in 200 ml of 30% ethanol with gentle shaking (~60 rpm)
for 1 h and stained with Stains-All (200 ml at 6.25 μg/ml in 30%
ethanol; E-9379, Sigma) overnight in darkness. The next day,
the stain was discarded, and the gels were rinsed with distilled
water, exposing them to light for ~25 min until background
was minimal. The gels were imaged on a light box with a stand-
ard, color, digital camera.

RESULTS

Leukocytes Bind Hyaluronan Cables Produced by MASM
Cells Stimulated with the Viral Mimic Poly(I,C)—Under rou-
tine conditions (Fig. 1, A–C), cultured MASM cells are coated
with small amounts of hyaluronan (B, green) on the apical sur-
face that is not adhesive for leukocytes. MASM cells treated
with poly(I,C) (Fig. 1, D–F) synthesize an abnormal hyaluronan
matrix (E, green), which has been described previously as
resembling cables (36). U937 leukocytes, stained with an anti-
sperm for human CD44 that does not interact with murine
CD44 (Fig. 1D, red), adhere to this matrix. The following exper-
iments define processes by which MASM cells initiate synthesis
of the hyaluronan matrices with structural information that
promotes leukocyte adhesion and subsequent degradation of
the abnormal matrix.

MASM cells were cultured for 18 h in 5% FBS with different
centration of poly(I,C) (0–50 μg/ml). Fig. 2A shows
increased accumulation of hyaluronan in the cell layer at 1
μg/ml poly(I,C) with a further increase to a plateau level
worth 4-fold above the control (without poly(I,C)) in 10 μg/ml
poly(I,C) (p < 0.0001). Fig. 2B shows that the adhesion of U937
leukocytes (4 °C, 30 min) increased to a plateau level at 1 μg/ml
poly(I,C) (p < 0.002), and that the majority of U937 leukocytes,
which are not adhere to HA (4 °C, 30 min) in the presence of
poly(I,C), adhesion to HA (in the presence of poly(I,C)) is
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Poly(I,C), the hyaluronan content increased in the cell layer with increasing concentrations of FBS (Fig. 3A, white bars) but remained constant in the medium fractions (Fig. 3A, gray bars). In the absence of poly(I,C), hyaluronan secreted into the medium fractions remained nearly constant (15.2 ± 1.6 μg/culture) with increasing FBS (Fig. 3B, gray bars) and close to the amounts secreted into the medium fractions in poly(I,C)-treated cultures (21.4 ± 2.3 μg/culture). In contrast, for each FBS concentration, the amounts of hyaluronan in the cell layers (Fig. 3B, white bars) were much lower when untreated than for the poly(I,C)-treated cultures.

Adhesion of leukocytes (4 °C, 30 min) was measured in parallel cultures treated with (Fig. 3C, white bars) or without (Fig. 3D, white bars) poly(I,C). Replicate cultures in each case were treated with Streptomyces hyaluronidase to determine hyaluronan-mediated leukocyte adhesion (Fig. 3, C and D, gray bars). Poly(I,C) treatment in the absence of FBS showed a significant (p < 0.002) hyaluronan-mediated leukocyte adhesion that increased further at 0.5% FBS (Fig. 3C, white bars compared with gray bars). Residual Streptomyces hyaluronidase independent leukocyte adhesion was highest in the culture without FBS and decreased with increasing FBS, reaching a minimum at 5% FBS (Fig. 3C, gray bars). The cultures without poly(I,C) treatment showed a small proportion of hyaluronan-mediated leukocyte binding, which was essentially independent of FBS concentration (Fig. 3D, white bars compared with gray bars). Hyaluronan-mediated adhesion was determined from the data in Fig. 3, C and D, by subtracting the values for the hyaluronidase residual bound leukocytes from the total bound (white bars minus gray bars), and the results are plotted in Fig. 3E. The values for poly(I,C)-treated cultures in the presence of 0.5% and higher FBS were up to 10-fold higher (p < 0.003) than the values in the untreated cultures.

The results in Fig. 3 indicate that poly(I,C) increases hyaluronan contents of the cell layer and hyaluronan-mediated leukocyte adhesion even in the absence of FBS, but that the presence of FBS increases the response. For comparison, in cultures without FBS, the ratio of hyaluronan in the cell layer to that in the medium was 2.8, whereas the ratio was 8.5 in cultures with 5% FBS (Fig. 3A, arrows). In contrast, the amount of hyaluronan secreted into the medium compartment was independent of FBS concentration. Thus, the increase in hyaluronan in the cell layer in response to poly(I,C) is independent of the pathway that is involved in hyaluronan secretion. Based on the results showing that the hyaluronan response in the poly(I,C)-treated cultures relative to the untreated cultures was at a plateau level in

FIGURE 1. MASM cells produce hyaluronan “cable” structures that promote leukocyte adhesion in response to poly(I,C). MASM cells were treated with (D–F) or without (A–C), poly(I,C), followed by the application of U937 leukocytes. The nontreated MASM cells do not promote leukocyte adhesion via the small amount of hyaluronan on their apical surface (B, green). In contrast, poly(I,C) induced synthesis of hyaluronan (E, green). U937 leukocytes bind to the hyaluronan produced in response to poly(I,C) (D, red) but are absent in the untreated culture (A). The leukocytes are stained with a human-specific anti-CD44 antibody that binds to the human-derived U937 cells but not the underlying MASM cells. 4',6-Diamidino-2-phenylindole-stained nuclei are blue. The merged views are in C and F, and their white bars indicate 50-μm magnification. This experiment was repeated >3 times with different experiments.

FIGURE 2. Dose relationship of poly(I,C) to the hyaluronan production and hyaluronan-mediated leukocyte adhesion of MASM cells. A shows the FACE analyses of hyaluronan for the cell matrix of MASM cells treated with different doses of poly(I,C) in 5% FBS (n = 6). B shows the number of U937 leukocytes bound (white bars) to the MASM cells for this same experiment compared with the number of leukocytes remaining after release by treatment with Streptomyces hyaluronidase (5 min, 4 °C) (gray bars) (n = 3). Error bars represent standard deviation.
The cell layer at 5% FBS with minimal non-hyaluronan-mediated leukocyte adhesion, this concentration of FBS was used in the following experiments.

**Differential Hyaluronan Kinetics of Poly(I,C) and Tunicamycin Stimulation of MASM Cells**—Fig. 4 shows the results of an experiment to determine the time course of the poly(I,C) response. All cultures were started at time 0 in the presence of 5% FBS, and a small aliquot of poly(I,C) was added at different times during the 18-h incubation to yield a final concentration of 10\(\mu\text{g/ml}\) poly(I,C). Controls (Fig. 4, A and B, bars at time 0) were incubated without poly(I,C) for the entire 18 h. Exposure of the MASM cells to poly(I,C) for 2 h already showed an increase in the amount of hyaluronan in the cell layer, with a nearly linear increase to a plateau value (\(p < 0.0001\)) at about 10 h, similar to the increase in hyaluronan in the cell layer at 5% FBS with minimal non-hyaluronan-mediated leukocyte adhesion, this concentration of FBS was used in the following experiments.

**FIGURE 4. Poly(I,C) kinetics of hyaluronan production and hyaluronan-mediated leukocyte adhesion by MASM cells.** A shows the FACE analyses of hyaluronan for MASM cells treated with 10\(\mu\text{g/ml}\) poly(I,C) and 5% FBS for different times. Hyaluronan present in the cell layer is shown as white bars, and the amount secreted into the medium is shown as gray bars (\(n = 4\)). B shows the number of leukocytes bound (white bars) to poly(I,C)-treated MASM cells for this same experiment compared with the number of leukocytes remaining after release by hyaluronidase (gray bars) (\(n = 4\)). Error bars represent standard deviation.

**FIGURE 3. Relation of serum to poly(I,C)-induced hyaluronan production and hyaluronan-mediated leukocyte adhesion of MASM cells.** A shows the FACE analyses of hyaluronan for MASM cells treated with 10\(\mu\text{g/ml}\) poly(I,C) and different doses of FBS in the cell layer (white bars) and secreted into the medium (gray bars) (\(n = 2\)). B shows the effect of FBS dosage on MASM cells that were not treated with poly(I,C) (\(n = 2\)). C and D show the number of U937 leukocytes bound (white bars) to poly(I,C)-treated (C) or nontreated (D) MASM cells for this same experiment compared with the number of leukocytes remaining after release by hyaluronan degradation (gray bars) (\(n = 3\)). E shows data from C and D expressed as the number of leukocytes bound directly to hyaluronan on poly(I,C)-treated (gray bars) and nontreated (white bars) MASM cells (\(n = 3\)). Error bars represent standard deviation for \(n = 3\) or range for \(n = 2\).
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FIGURE 5. Tunicamycin kinetics of hyaluronan production and hyaluronan-mediated leukocyte adhesion by MASM cells. A shows the FACE analyses of hyaluronan for MASM cells treated with 5 μg/ml tunicamycin for the same times (and using the same original cell preparation) presented in Fig. 4. Hyaluronan derived from the cell layer is shown as white bars, and the amounts of hyaluronan secreted into the medium are shown as gray bars (n = 4; error bars represent standard deviation). B shows the number of leukocytes bound (white bars) to tunicamycin-treated MASM cells for this same experiment compared with the number of leukocytes remaining after release by hyaluronan degradation (gray bars) (n = 2; error bars represent range).

the cell layer. The results in Fig. 4 show that the synthesis of the hyaluronan matrix that is adhesive for leukocytes is initiated rapidly upon exposure to poly(I,C) and that it continues to increase with linear kinetics for 10–12 h before stopping.

Tunicamycin at a plateau concentration of 5 μg/ml induced synthesis of an adhesive hyaluronan matrix by MASM cells cultured for 18 h in the presence of 5% FBS (data not shown). Fig. 5 shows hyaluronan production and leukocyte adhesion for cultures treated with 5 μg/ml tunicamycin and 5% FBS for different times up to 18 h. Similar to the poly(I,C) time course, synthesis of the hyaluronan in the cell layer was initiated rapidly and increased linearly with time before reaching a plateau (p < 0.0002) between 10 and 12 h (Fig. 5A, white bars). In contrast to the poly(I,C)-treated cultures, the hyaluronan content in the cell layer at plateau was much less (25.2 ± 1.7 μg/culture (Fig. 5A) versus 111.3 ± 8.1 μg/culture (Fig. 4A); p < 0.0001). The amounts secreted into the medium compartment showed a tendency to decrease with time, particularly after 8 h of treatment (Fig. 5A, gray bars). As for the poly(I,C) treatment, tunicamycin treatment initiates an independent mechanism for formation of the hyaluronan matrix in the cell layer compared with that for secretion into the medium. Fig. 5B shows the adhesion of U937 leukocytes for parallel cultures. In this case, the adhesion showed a lag of 2–4 h before increasing to a maximum plateau value (p < 0.0001) at 8–12 h of tunicamycin treatment. Interestingly, the number of leukocytes bound per increase in cell layer hyaluronan content is much higher (p < 0.0001) for the maximum plateau values for tunicamycin treatment than for the poly(I,C) treatment (~120 versus 55 leukocytes/μg of hyaluronan, respectively) as is discussed below.

Selective Induction of MASM Cell Adhesion Molecules by Poly(I,C), but Not Tunicamycin, Despite Similar Hyaluronan-mediated Leukocyte Adhesion—Fig. 6 shows an example in which the hyaluronan synthesis and leukocyte adhesion parameters are compared for identical MASM cell cultures treated with poly(I,C) (10 μg/ml) and tunicamycin (5 μg/ml) under standard conditions (5% FBS, 18 h). In this case, at the end of the 18-h incubation, some cultures were treated with Streptomyces hyaluronidase before testing them for leukocyte adhesion (Pre-Hyaluronidase, gray bars) and others after leukocyte adhesion (Post-Hyaluronidase, black bars) (Fig. 6A). The control cultures...
Fig. 6A, No Treatment) showed some hyaluronan-mediated adhesion of the leukocytes (white bar compared with black bar). The poly(I,C) and tunicamycin cultures showed 3.6- and 2.9-fold (p < 0.0001 and 0.0008, respectively) increases in hyaluronan-mediated leukocyte adhesion compared with no treatment (Fig. 6A, white bars), respectively. Interestingly, pretreatment of poly(I,C) cultures with hyaluronidase showed a significant (p < 0.0004) increase in the number of leukocytes bound compared with the number that remained bound when the hyaluronidase treatment was done after leukocyte adhesion (Fig. 6A, gray bar compared with black bar), and both were significantly (p < 0.0001 and 0.0004, respectively) higher than the respective control cultures. This is likely because of up-regulation of VCAM on the MASM cells, which permits adhesion through interaction with VLA4 integrin on the leukocytes (36). We observed that poly(I,C), but not tunicamycin, induces VCAM-1 synthesis by the MASM cells (Fig. 6C). Additionally, we found that poly(I,C)-treated MASM cells pretreated with Streptomyces hyaluronidase promote U937 cell adhesion to cells expressing high levels of VCAM-1 (Fig. 6D). The presence of the hyaluronan matrix during the leukocyte adhesion step masks about half of the sites that are exposed when the hyaluronan matrix is removed prior to leukocyte binding. Notably, the tunicamycin-treated cultures do not show a significant difference between hyaluronidase treatments before and after leukocyte adhesion and only a modest increase relative to the control cultures. This is consistent with ER stress, which drastically inhibits new protein synthesis except for select proteins, such as chaperones, in an attempt to facilitate proper folding of proteins in the endoplasmic reticulum.

In contrast to the similarity of the leukocyte adhesion response, the increase in hyaluronan deposition in the matrix of the tunicamycin-treated MASM cells was much less (p < 0.003) than for the poly(I,C)-treated cells (Fig. 6B). Typically, tunicamycin increases hyaluronan deposition less than 2-fold, whereas poly(I,C) yields 7-fold or greater increases. The observation that similar numbers of leukocytes bind to the hyaluronan matrix in both treatments (Fig. 6A) suggests that the organization of the newly synthesized matrix may be much more dispersed in the tunicamycin-treated cultures, with less masking of potential binding sites, than for poly(I,C)-treated cultures.

Despite Similar Hyaluronan-mediated Leukocyte Adhesion by MASM Cells Stimulated with Poly(I,C) and Tunicamycin, Only the Tunicamycin Response Is Associated with the Up-regulation of ER Chaperones—Fig. 7, A–C, shows confocal micrographs of control, poly(I,C), and tunicamycin cultures stained for hyaluronan (green) and KDEL (red), a peptide sequence marker for chaperones synthesized in response to ER stress. (40) D shows a Western blot in which MASM cell protein extracts were probed for KDEL (green) and β-actin as a loading control (red) to confirm our confocal observations (A–C). Lanes 1–3 represent MASM cells untreated (NT; lane 1), or treated with poly(I,C) (PIC; lane 2) or tunicamycin (TUN; lane 3). Molecular weight standards are shown as red/yellow. This blot was repeated three times with three different samples with essentially identical results.
Hydrolysis of blunt-ended DNA molecules

The hydrolysis of blunt-ended DNA molecules was studied to understand the dynamics of DNA repair. The experiment was conducted using a series of buffer conditions to investigate the role of specific ions and pH levels. The data collected indicated that the hydrolysis rate was significantly affected by the presence of certain ions, particularly calcium ions.

The results showed a clear correlation between the hydrolysis rate and the concentration of calcium ions. At lower calcium ion concentrations, the hydrolysis rate was significantly slower, suggesting a role for calcium ions in promoting or stabilizing the DNA structure.

The study also revealed that the pH level of the buffer significantly influenced the hydrolysis rate. Optimal conditions were found to be in the pH range of 7.5 to 8.0, where the rate of hydrolysis was highest.

In conclusion, this study highlights the importance of environmental factors such as calcium ion concentration and pH in the hydrolysis of blunt-ended DNA molecules, providing valuable insights for further research in DNA repair mechanisms and related biological processes.

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**Figure 1:**

A: A schematic representation of the DNA double helix with blunt-ended cuts highlighted in green.
B: A graph showing the hydrolysis rate as a function of calcium ion concentration.
C: A graph illustrating the effect of pH on the hydrolysis rate.

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**Figures and Tables:**

1. Figure 1: Illustrates the molecular structure of DNA with specific focus on blunt-ended cuts.
2. Table 1: Summary of hydrolysis rates under different buffer conditions.
3. Additional figures and tables available in the supplementary material.
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FIGURE 9. Poly(I,C) and tunicamycin induce hyaluronan synthesis and hyaluronan-mediated leukocyte adhesion by complementary, yet independent, pathways. MASM cells were either untreated (NT) or treated with poly(I,C) (PIC), tunicamycin (TUN), or a combination of both (TUN + PIC). These treatments were made at doses that were known to induce a maximum plateau response (white bars) for the leukocyte adhesion assay (10 and 5 μg/ml for poly(I,C) and tunicamycin, respectively) and a mid-plateau dose (gray bars) that gave leukocyte adhesion values that were about 50% of the maximum value (0.1 and 0.5 μg/ml for poly(I,C) and tunicamycin, respectively). A shows the number of leukocytes bound to the MASM cells under these conditions. B shows the number of leukocytes remaining after treatment with Streptomyces hyaluronidase. C presents the hyaluronan mass (by FACE analyses) of hyaluronan from the cell matrices for the same experiment. Error bars represent standard deviation.

least (27 and 42%, respectively) increase relative to the untreated control. In contrast, FACE analysis showed a similar fold increase with respect to the HABP extraction (p > 0.03) for the tunicamycin-treated cells while showing a 3.4-fold increase (p < 0.0001) for the poly(I,C)-treated cells. This confirms that the amount of freely available hyaluronan, as a result of poly(I,C) treatment, is only a fraction (29%) of the total produced by the cell. Thus, 71% of the hyaluronan produced by poly(I,C)-treated MASM cells is masked. These data also confirm that essentially all of the hyaluronan, produced by the MASM cells in response to tunicamycin, is freely available for leukocyte adhesion.

Independent and Additive Effects of Poly(I,C) and Tunicamycin—Induction of the poly(I,C)-mediated interferon pathways and ER stress, in the same cell, more closely resembles viral infection than either of these alone. Fig. 9 shows the results of an experiment to determine the effect of combining tunicamycin and poly(I,C) treatments. Pre-confluent MASM cell cultures were treated under standard conditions (18 h, 5% FBS) with tunicamycin and poly(I,C), at a dose known to induce maximum hyaluronan-mediated leukocyte adhesion (plateau doses; white bars of Fig. 9; 5 and 10 μg/ml, respectively) and a dose inducing leukocyte adhesion at about half of the maximum response (mid-plateau dose; gray bars of Fig. 9; 0.1 and 0.5 μg/ml, respectively). Hyaluronan-mediated leukocyte adhesion for the combination was partially, but not completely, additive (a + b ≠ c; 28 and 21% increase from TUN and PIC, respectively; p < 0.002–0.0002) at the plateau doses and slightly more than additive at the mid-plateau doses (a’ + b’ = c’; 50 and 63% increase from TUN and PIC, respectively; p < 0.02–0.008) when comparing the combined treatments with the individual treatments at the mid-plateau doses (Fig. 9A). Similarly, the amount of hyaluronan in the cell layer, for the combination treatments, was almost completely additive (a’ + b’ = c’; 50 and 43% increase from TUN and PIC, respectively; p < 0.02–0.008) when comparing the combined treatments with the individual treatments at the plateau doses (Fig. 9C). In contrast, the amount of hyaluronan in the cell layer at the plateau doses was not additive (a + b ≠ c) for the combination treatments but was actually lower (45%; p < 0.002) than poly(I,C) alone, although higher (48%; p < 0.0001) than tunicamycin alone. The observation that poly(I,C) and tunicamycin induce independent, yet additive, hyaluronan production and hyaluronan mediated leukocyte adhesion implies that these two toxins function in different, but related, cellular pathways.

From the confluency data (Fig. 8), we learned that leukocyte binding does not show linear proportionality to the amount of hyaluronan on the cell surface. This was probably caused by masking the available surface hyaluronan for binding. For example, a thicker coat of hyaluronan could bind the same number of leukocytes as a thinner coat if they both had the same amount of surface area hyaluronan available for binding. Thus, in Fig. 9, greater leukocyte binding from less hyaluronan (as observed for the combined treatments at plateau doses) could be explained if a thinner coat of surface hyaluronan was more evenly distributed between a greater number of cells. This could happen if a portion of the MASM cells responded to one of the two treatments better than the other.

Hyaluronan Size Determination from MASM Cells Treated with Poly(I,C) or Tunicamycin—We treated MASM cells with poly(I,C) (10 μg/ml) or tunicamycin (5 μg/ml) under standard conditions (18 h, 5% FBS). We purified the hyaluronan (and other glycosaminoglycans) through a series of enzymatic digestions (proteinase, nuclease, and glycosidase) and precipitations (as fully described in “Experimental Procedures”), analyzing the extracts by agarose gel electrophoresis (Fig. 10). This resulted in a blue, high molecular weight (HMW) smear (1500–4000 kDa) that was confirmed as hyaluronan by pre-digestion with Streptomyces hyaluronidase (Fig. 10A, lanes 4, 6, and 8). The molecular weight range was similar in each of the treatment conditions (untreated (NT), tunicamycin, and poly(I,C); lanes 3, 5, and 7, respectively). The smears below 27 kDa are likely heparan sulfate (Fig. 10A, yellow/brown) and chondroitin sulfate (purple).4 We were surprised that although the hyaluronan is present as a 1500–4000-kDa smear, a significant amount was present in a fairly defined band at about 4000 kDa. Because this

4 M. Lauer, A. Wang, and V. C. Hascall, personal observations.
phenomenon could be caused by the aggregation of multiple hyaluronan strands, we made serial dilutions from a separate aliquot of the poly(I,C)-induced hyaluronan to determine whether this unusual HMW band might be related to the concentration of hyaluronan in solution (Fig. 10B). Within the range of detection, diluted hyaluronan did not show a dissolution of the HMW band. Another possible cause for hyaluronan aggregation might be that the ionic conditions (TAE buffer and any other salts that might not have been removed by ethanol precipitation) of the samples induce aggregation. But when a hyaluronan molecular weight standard was added to a hyaluronan extract from poly(I,C)-treated MASM cells (Fig. 10C, lane 4), no differences in band migration occurred, indicating that the ionic conditions of the sample were not responsible for the high molecular weight band. Finally, we compared hyaluronan from poly(I,C)-induced MASM cells, purified under routine conditions (Fig. 10C, lane 5), with identical hyaluronan that had been exposed to 4 M guanidine, 2% CHAPS, and 0.05 M Tris-HCl, pH 7.0, as a means to disassociate any potential hyaluronan aggregates (lane 6). The failure of any of these approaches to disrupt the HMW (~4000 kDa) hyaluronan band confirms that this band represents the actual molecular weight of unassociated hyaluronan strands.

**DISCUSSION**

In this paper, we described the hyaluronan-mediated response of MASM cells to poly(I,C) and tunicamycin, which are models that mimic viral infection and ER stress relevant to airway inflammation associated with asthma (23, 27, 38, 40, 41).

Our results demonstrate that MASM cells respond to both poly(I,C) and tunicamycin to produce a leukocyte-adhesive hyaluronan matrix, but accomplish this by different cellular pathways. At all cell densities studied, MASM cells initiate synthesis of the hyaluronan matrix immediately after the addition of poly(I,C). This matrix accumulates linearly for

![FIGURE 10. Hyaluronan size determination from MASM cells treated with poly(I,C) or tunicamycin.](image-url)
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10–12 h, resulting in a 7–9-fold increase in the amount of cell-associated hyaluronan matrix. In contrast, the response of the MASM cells to tunicamycin depends on cell density, with pre-confluent densities responding most effectively. Tunicamycin-induced hyaluronan accumulation is nearly linear for 10–12 h, although the matrix accumulation is much less (~2-fold at most). In both treatments, the normal pathway for synthesis and secretion of hyaluronan into the medium is not significantly altered, which indicates that the hyaluronan response to poly(I,C) and tunicamycin is completely associated with the cell matrix.

Although U937 leukocytes adhere to the hyaluronan matrix produced by both reagents, the number bound per hyaluronan in the matrix is much higher for tunicamycin-treated MASM cell cultures. This implies that the more robust cable structures produced by poly(I,C)-treated MASM cells mask many sites that remain accessible in the more disperse hyaluronan matrix initiated by tunicamycin. These results are consistent with the observation that there is no difference between the numbers of leukocytes that bind to tunicamycin-treated MASM cells after hyaluronidase treatment compared with the number that remained bound when the hyaluronidase treatment followed the initial adhesion. In contrast, hyaluronidase digestion before leukocyte adhesion in poly(I,C)-treated MASM cultures increased the number of bound leukocytes (probably via VCAM), indicating that many of the sites on the cell surfaces were masked by the more copious hyaluronan matrix synthesized in response to poly(I,C).

Additionally, poly(I,C) treatment of MASM cells, in addition to inducing hyaluronan-mediated leukocyte adhesion, also induces the expression of various proteins, activates the ER stress response, and is therefore more sensitive to ER stress. If this is the case, the poly(I,C) response appears to be independent of the number of leukocytes.

The abnormal hyaluronan matrix increases in proportion to cell number with poly(I,C) treatment, although it decreases with increased cell number with tunicamycin treatment. This may indicate that pre-confluent MASM cells are synthesizing more protein in their proliferative phase and are therefore more sensitive to ER stress. If this is the case, the poly(I,C) response appears to be independent of the load of protein synthesis.

There are at least three cellular pathways that can initiate the synthesis of the abnormal hyaluronan matrix. In addition to the viral and ER stress responses, mesangial cells stimulated to divide in hyperglycemic medium produce a leukocyte-adhesive matrix (48). Furthermore, recent studies have shown that 3T3-L1 cells that are differentiated to adipocytes in hyperglycemic medium also produce this matrix (49). ER stress may also link obesity and insulin resistance in type 2 diabetes (50). This report showed that high fat feeding and obesity induce ER stress in liver, which suppresses insulin signaling via c-Jun N-terminal kinase activation.

Several studies indicate that leukocytes adhere to certain hyaluronan matrices synthesized by cells subjected to various stress responses, but not others, such as hyaluronan coats made by cells normally (51). For example, lymphocytes interact with the abnormal hyaluronan matrix produced by venules and endothelial cells produced in graft versus host disease, but do not do so to the adjacent normal hyaluronan matrix in the papillary dermis (52). Kidney renal tubule endothelial cell cultures stimulated with either BMP-7 or interleukin-1β increase hyaluronan synthesis (53). However, lymphocytes only adhere to
the hyaluronan matrix produced in response to BMP-7. This suggests that the increase in hyaluronan synthesis by interleukin-1β reflects stimulation of the normal hyaluronan synthesis pathway. Lymphocytes adhere much more avidly to hyaluronan with covalently bound heavy chains of inter-α-inhibitor than to unsubstituted hyaluronan (54). All of these results indicate that hyaluronan matrices produced in response to stress contain structural information that inflammatory cells can recognize.

Under normal physiological conditions, hyaluronan is cleared from connective tissues through the lymphatics (55) or by the resident cells (56, 57). In contrast, under some pathological conditions, as in asthmatic airways, the abnormal hyaluronan matrix is removed by the leukocytes/macrophages that enter the tissue. Degradation of the abnormal matrix by these cells could release hyaluronan fragments, which are now considered “danger signals” in initiating host responses to the inflammatory process and likely participate in determining the extent of the response (58). Overproduction of the matrix or over-reaction by inflammatory cells to this matrix may contribute to the inflammatory flares that are typical of asthmatic patients.

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