p38 and OGT Sequestration into Viral Inclusion Bodies in Cells Infected with Human Respiratory Syncytial Virus Suppresses MK2 Activities and Stress Granule Assembly

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Respiratory syncytial virus (RSV) forms cytoplasmic inclusion bodies (IBs) that are thought to be sites of nucleocapsid accumulation and viral RNA synthesis. The present study found that IBs also were the sites of major sequestration of two proteins involved in cellular signaling pathways. These are phosphorylated p38 mitogen-activated protein kinase (MAPK) (p38-P), a key regulator of cellular inflammatory and stress responses, and O-linked N-acetylgalcosamine (OGN) transferase (OGT), an enzyme that catalyzes the posttranslational addition of OGN to protein targets to regulate cellular processes, including signal transduction, transcription, translation, and the stress response. The virus-induced sequestration of p38-P in IBs resulted in a substantial reduction in the accumulation of a downstream signaling substrate, MAPK-activated protein kinase 2 (MK2). Sequestration of OGT in IBs was associated with suppression of stress granule (SG) formation. Thus, while the RSV IBs are thought to play an essential role in viral replication, the present results show that they also play a role in suppressing the cellular response to viral infection. The sequestration of p38-P and OGT in IBs appeared to be reversible: oxidative stress resulting from arsenite treatment transformed large IBs into a scattering of smaller bodies, suggestive of partial disassembly, and this was associated with MK2 phosphorylation and OGT addition. Unexpectedly, the RSV M2-1 protein was found to localize in SGs that formed during oxidative stress. This protein was previously shown to be a viral transcription elongation factor, and the present findings provide the first evidence of possible involvement in SG activities during RSV infection.

Human respiratory syncytial virus (RSV) is the most important viral agent of severe respiratory tract disease in infants and young children worldwide. RSV is a cytoplasmic, enveloped virus with a nonsegmented negative-strand RNA genome that is classified in the genus Pneumovirus, family Paramyxoviridae (1). There presently is no specific anti-RSV therapy or vaccine, although infants at high risk for severe RSV disease due to underlying conditions can be substantially protected by passive antibody immunoprophylaxis (2).

RSV forms cytoplasmic inclusion bodies (IBs) during infection, as also has been reported for measles virus and human metapneumovirus, also of the family Paramyxoviridae (3–5). In the case of RSV, the IBs have been shown to contain the RSV nucleoprotein (N), phosphoprotein (P), M2-1 protein, and large polymerase (L) protein (4, 6). The expression of viral N and P proteins is sufficient for the appearance of IBs (4, 7). Viral genomic RNA also localizes in IBs (8), consistent with the presumption that these are sites of nucleocapsid assembly and RNA synthesis. Furthermore, heat shock protein Hsp70 has been shown to associate with IBs, although no functional role was determined (9). Overall, the formation and function of the IBs are not well understood.

As an obligate intracellular parasite, RSV interacts with host signaling networks and machinery both to block antiviral responses and to promote viral replication. Previous work implicated the mitogen-activated protein kinases (MAPKs), in particular the extracellular signal-regulated kinase (ERK) and p38 MAPK, in the tropism as well as entry of RSV (10–12). The p38 MAPK is a central mediator involved in regulating cellular inflammatory and stress responses, as well as cellular protein synthesis (13, 14). Thus, any alteration of p38 signaling during a viral infection has the potential for multifield impact on virus-host interactions. p38 and one of its downstream substrates, MAPK-activated protein kinase 2 (MK2), play important roles in posttranscriptional mRNA metabolism during stress conditions. In particular, activated MK2 promotes the stability of AU-rich element (ARE)-containing mRNAs, such as those encoding proinflammatory and antiviral proteins, including beta interferon (IFN-β), interleukin 1β (IL-1β), and tumor necrosis factor alpha (TNF-α) (15–17). Thus, interference with signal transduction through p38 and MK2 can reduce the stability of the mRNAs encoding these innate response proteins and thereby reduce their production.

Of the four p38 isoforms (α, β, γ, and δ), p38α appears to be responsible for MK2 activation. Thermodynamic and steady-state kinetic characterization using p38α indicated a high-affinity binding with MK2 (Kd [equilibrium dissociation constant] = 2.5 nM), and the complex is required in stress-dependent-activation of MK2 (18, 19). Furthermore, the formation of this complex seems to be critical for the stabilization of both proteins, as p38 accumulation is significantly reduced in MK2-deficient cells and, con-
versely, MK2 accumulation is reduced in p38α-knockout mouse embryonic fibroblasts (20, 21). Earlier studies could not rule out a role for the β isform because the inhibitors involved affected both p38α and p38β (22), but subsequent studies showed that MK2 stability and signaling are unaffected in knockout mice lacking the p38β isoform (23).

Another aspect of the cellular response to stress is the formation of stress granules (SGs). These are complex ribonucleoprotein aggregates that contain untranslated mRNAs and form under stress conditions. SGs constitute an important intermediate step in the equilibrium between active translation and mRNA decay (24). Regulation of SG dynamics involves posttranslational modifications of a number of proteins by methylation, acetylation, phosphorylation, and the addition of O-linked N-acetylglucosamine (OGN). In particular, OGN-modification, mediated by OGN-transferase (OGT), of various ribosomal proteins is required for the aggregation of untranslated messenger ribonucleoproteins and assembly into SGs (25–27). Interestingly, p38 has been shown to interact with and recruit OGT to specific targets (28, 29).

Numerous viruses have been shown to manipulate or modulate SG pathways to impact translation and potentially affect innate antiviral mechanisms (27). The role of SGs in RSV infection remains to be elucidated. In one study, SGs were thought to promote antiviral mechanisms (27). The role of SGs in RSV infection late SG pathways to impact translation and potentially affect in-

### MATERIALS AND METHODS

**Cells.** A549 cells (ATCC CCL-185) were grown in F-12K medium (ATCC) containing 10% fetal bovine serum (FBS). Vero cells were grown in Opti-MEM I medium (Invitrogen) containing 10% FBS. The respective media containing 2% FBS were used during all experiments described below.

**Virus, infection, and inhibitor.** Recombinant RSV (RA2) was constructed from the wild-type A2 strain as previously described (31). Virus stocks were propagated and titrated on Vero cells. For titration, plaques were stained using monoclonal antibodies (MABs) specific to RSV F protein, followed by peroxidase-labeled anti-mouse IgG(H+L) (KPL) for detection. RSV was added to A549 cells at a multiplicity of infection (MOI) of 0.5 PFU per cell for all experiments unless indicated otherwise. Inhibitor CMPD-1 [4-(2’-fluorobiphenyl-4-yl)-N-(4-hydroxyphenyl)-butyramide] (EMD Biosciences or Tocris), which selectively inhibits the phosphorylation of MK2 by p38α, was prepared as a stock solution in dimethyl sulfoxide (DMSO) and used at a final concentration of 50 μM (32). Mock-treated cells were given an equivalent amount of DMSO (final concentration, 0.2%).

**RSV-specific FISH.** Fluorescence in situ hybridization (FISH) was performed as previously described (33) and adapted for the present study. Briefly, cells were fixed with 4% paraformaldehyde and hybridized overnight at 50°C with a mixture of antisense digoxigenin-UTP-labeled riboprobes representing the RSV N, P, M2-1, NS1, NS2, and F genes. These probes were 285 to 432 nucleotides in length (sequences are available upon request) and were synthesized commercially (Lofstrand Labs, Ltd., Gaithersburg, MD). Following hybridization, cells were blocked with 2% horse serum, 2% sheep serum, and 0.2% fish skin gelatin in 0.1 M Tris (pH 7.4) buffer and incubated with sheep anti-digoxigenin-alkaline phosphatase (Roche Molecular Biochemicals). Finally, for detection and visualization, Alexa 594-conjugated tyramide (Invitrogen) was applied in a tyramide signal amplification diluent (1:100) (PerkinElmer). Samples were then rinsed sequentially in 0.1 M Tris (pH 7.4) containing 0.1% Tween 20, 0.1 M Tris (pH 7.4), and phosphate-buffered saline (PBS) and were mounted in ProLong Gold antifade reagent containing the nuclear stain 4’,6-diamidino-2-phenylindole (DAPI) (Invitrogen). Samples were analyzed by confocal microscopy.

**Confocal microscopy.** Fluorescent images were captured on a Leica TCS-SP5 confocal microscope (Leica Microsystems, Germany) equipped with a white light laser using a 63×/NA1.4 oil immersion objective. The dynamic range of pixel intensities was determined so to avoid saturation of the brightest signal (such as in the dense IB aggregates) in an experiment. Each image was taken as a z-stack of 0.25-μm-thick slices. Except as noted, individual slices from a z-stack of images are shown. In some cases, as noted, 3-dimensional renderings were constructed from a z-stack of images to illustrate colocalizations. Huygens Essential (version 3.6; Scientific Volume Imaging, the Netherlands), Leica Lite (version 2.6.0; Leica Microsystems, Germany), and Imaris (version 7.1; Bitplane AG, Switzerland) were used for deconvolution and image processing.

**Antibodies.** Polyclonal antibodies (PAb) against p38α, MAB against p38β, MAB against M2-1, PAb against OGT, and PAb against β-actin were obtained from Abcam. MAB against P was obtained from Antibodies-Onlin. MAB against OGN was obtained from Covance. MAB against MK2 was obtained from Epitomics. PAb against TIA-1 was obtained from Santa Cruz Biotechnology. MAB against eukaryotic initiation factor 3 (eIF3) was obtained from Cell Signaling. Two different preparations of PAb against RSV were used: for Western blotting, we used a rabbit PAb that we used in preparing use sucrose gradient-purified RSV that had been de-natured and reduced, and for immunofluorescence, we used a goat PAb that was obtained from AbD Serotec. Secondary antibodies used for Western blotting were species specific, affinity purified, and peroxidase labeled (KPL). Secondary species-specific antibodies used for immunofluorescence were labeled with either Alexa 488 (Invitrogen or Jackson ImmunoResearch) or Alexa 594 or Alexa 647 (Invitrogen).

**Immunofluorescence.** Cells were fixed in 4% paraformaldehyde in Hanks’ balanced salt solution for 15 min at room temperature and were permeabilized with 1% Triton X-100 (Roche) in PBS for 5 min. The cells were subsequently stained with primary and appropriate fluorescent dye-labeled secondary antibodies and analyzed by confocal microscopy as described above.

**Western blot analysis.** Cells were lysed in NuPAGE LDS sample buffer (Invitrogen) supplemented with 1× phosphatase (PhosSTOP; Roche) and 1× protease inhibitor (Complete, Roche). The lysate was centrifuged through a QIAshredder Mini Spin Column (Qiagen) and subjected to heat denaturation with 2-mercaptoethanol (Sigma). Equal amounts of cell lysate were separated using 4 to 12% Novex Tris glycine gels (Invitrogen) and transferred to polyvinylidene difluoride (PVDF) membranes using an iBlot gel transfer device (Invitrogen). The membrane was incubated with primary and peroxidase-conjugated secondary antibodies. Specific bands were visualized by chemiluminescence using SuperSignal West Pico substrate (Thermo Scientific) and exposure to BioMax LIGHT films (Kodak).

### RESULTS

Effects of RSV infection on p38-P distribution. To investigate the role of the p38/MK2 pathway in RSV infection, we treated human airway epithelial A549 cells with inhibitor CMPD-1, which selectively inhibits p38/MK2 signaling by inhibiting phosphorylated p38α from phosphorylating and activating MK2 (32). Treatment with inhibitor CMPD-1 was initiated 30 min before infection to assess any inhibitory effect on viral entry or 75 min after infection, following virus adsorption. Viral infection was assessed by confocal microscopy. Western blotting using a rabbit antiserum raised against sucrose gradient-purified RSV, PAb specific to p38α, and a MAB specific to p38-P (Fig. 1). This showed that the level of p38α
inhibitor, viral RNA was readily detected in cytoplasmic aggregates from the input inoculum. In contrast, in the absence of inhibitor CMPD-1, and this small level of RNA might have been due to insufficient MK2 replacement due to the overall inhibition of protein synthesis. Remarkably, infection with RSV also resulted in a substantial decrease in the level of MK2. CMPD-1 resulted in a substantial decrease in the level of MK2. Consistent with the strong inhibition of viral protein synthesis observed in Fig. 1, only trace amounts of infectious virus were recovered from RSV-infected cells treated with inhibitor CMPD-1 (Fig. 2A). Furthermore, in situ hybridization of RSV-infected cells with a fluorescent dye-labeled viral RNA probe revealed only a small amount of viral RNA present in cells incubated in the presence of inhibitor CMPD-1, and this small level of RNA might have come from the input inoculum. In contrast, in the absence of inhibitor, viral RNA was readily detected in cytoplasmic aggregates (Fig. 2B) that were subsequently identified as viral IBs by the presence of viral P and M2-1 proteins detected by immunofluorescence (Fig. 3 and 4). The presence of RSV RNA in IBs had been documented previously (8). The accumulation of viral RNA in the IBs was maximal at 18 h postinfection, and the formation of IBs was most developed in terms of size at this time; therefore, 18 h postinfection was chosen for all subsequent experiments.

We also investigated whether RSV infection altered the cellular localization of p38-P. A549 cells were mock infected or infected with RSV, were treated or not with inhibitor CMPD-1 beginning 30 min before infection or 75 min postinfection, and at 18 h postinfection were analyzed by immunofluorescence staining with antibody specific to p38-P. The results revealed striking differences in cellular localization of p38-P under the various conditions (Fig. 3). These results showed that the inhibitor added 75 min after infection; the results with the inhibitor added 30 min before infection were indistinguishable and are not shown). In mock-infected cells, p38-P was detected as a diffuse signal throughout the cytoplasm (Fig. 3 and 4; the diffuse pattern is barely visible because the signal intensities of all of the images were reduced equally to avoid saturation of the more intense signals). In contrast, in RSV-infected cells, p38-P was found in dense cytoplasmic aggregates (Fig. 3 and 4). The appearance of this dense staining was not due to an increase in synthesis or stability, as was shown by the Western blot in Fig. 1. These aggregates were identified as viral IBs based on containing with MAbs against the viral M2-1 (Fig. 3 and 4A) and P (Fig. 4B) proteins, as well as by the presence of RSV RNA as already shown (Fig. 2B)(8). Infected cells treated with inhibitor CMPD-1 either 30 min before or 75 min after infection failed to form viral IBs, as would be expected since viral protein synthesis was strongly inhibited. These cells lacked detectable aggregation of p38-P (Fig. 3). The colocalization of p38-P with the viral M2-1 and P proteins was further illustrated by cross-sectional analysis and intensity profiling (Fig. 4). This showed that p38-P, M2-1, and P accumulated throughout the IBs, although the viral proteins tended to concentrate more heavily toward the periphery of the large IBs.

Effects of RSV infection on MK2 accumulation. Since RSV infection did not appear to affect the accumulation and phosphorylation of p38-P but drastically changed its intracellular localization, we examined the accumulation and phosphorylation status of its downstream signaling substrate MK2. Following the same treatment and infection protocol as in the previous experiments, cell lysates were prepared 18 h postinfection and analyzed by Western blotting with a MAb against MK2/MK2-P. This showed that phosphorylation of MK2 in mock-infected cells was below the limit of detection under these conditions (Fig. 5A). The results also showed that treatment of mock-infected cells with inhibitor CMPD-1 resulted in a substantial decrease in the level of MK2. This reduction in MK2 presumably was due to protein instability as a result of perturbed p38/P2K binding due to CMPD-1, as well as insufficient MK2 replacement due to the overall inhibition of protein synthesis. Remarkably, infection with RSV also resulted in a strong reduction in the level of MK2 (Fig. 5A) in this regard mimicked the outcome of the inhibitor. However, in this case, we suggest that the loss of MK2 was a result of the sequestration of p38-P in the IBs, making it unavailable to bind to and stabilize MK2. The addition of inhibitor to RSV-infected cells had no additional effect on the accumulation of MK2. As a positive control for activation of MK2, we subjected mock-infected cells to a 30-min treatment with arsenite, which is known to strongly induce...
oxidative stress. Under these conditions, essentially all of the MK2 was converted to MK2-P (Fig. 5), demonstrating the expected strong activation of the p38/MK2 pathway under stress conditions. When RSV-infected cells were similarly treated with arsenite for 30 min at 18 h postinfection, the MK2 protein, which was present at a greatly reduced level as already noted, was quantitatively converted to its activated, phosphorylated form, MK2-P (Fig. 5B).

The finding that MK2 was phosphorylated in RSV-infected cells following arsenite treatment was somewhat surprising, given that p38-P was sequestered in IBs in untreated RSV-infected cells. We therefore examined the localization of p38-P by immunofluorescence (Fig. 6). Following the brief arsenite treatment, immunostaining of RSV-infected cells with the P-specific (Fig. 6A) or M2-1-specific (Fig. 6B) MAb revealed IBs that appeared to be smaller and more numerous than in RSV-infected cells that were mock treated in parallel. This suggested that the arsenite treatment had the effect of disrupting the IBs. The observed MK2 activation in the case of the infected cells could be due, at least in part, to released p38-P.

Effects of RSV infection on OGT distribution and SG formation. As noted above (see the introduction), p38 and MK2 play important roles in posttranscriptional regulation of mRNA metabolism under stress conditions, roles that also involve SGs and OGT. p38 also binds to and recruits OGT (28, 29), which, in turn, mediates OGN modification of key target proteins that is necessary for aggregation of untranslated mRNA-containing ribonucleoproteins into SGs (26). Given the effects of RSV infection on the localization, accumulation, and activation of p38-P and MK2, we used immunofluorescence staining to investigate possible effects on OGT and SGs.

RSV-infected cells were analyzed at 18 h postinfection with antibodies specific to OGT, to the SG marker TIA-1, and to viral P (Fig. 7) or M2-1 protein (Fig. 8). Mock-infected cells exhibited a diffuse distribution of OGT (Fig. 7A and 8A; the diffuse pattern is barely visible because the signal intensities of all of the images were
reduced equally to avoid saturation of the more intense signals). In contrast, in RSV-infected cells, OGT was readily detected in dense cytoplasmic bodies. These bodies were found to contain dense accumulation of the RSV P (Fig. 7A) and M2-1 (Fig. 8A) proteins and thus were IBs. This indicated that similar to p38-P, OGT was sequestered in IBs during RSV infection.

TIA-1 also was present in a diffuse distribution in mock-infected cells, indicating the absence of SGs (Fig. 7A and 8A). When mock-infected cells were treated with arsenite for 30 min, the formation of SGs, revealed as intense speckles containing TIA-1, was evident (Fig. 7A and 8A). The distribution of TIA-1 also was diffuse in the majority of RSV-infected cells, indicating a lack of SGs in most cases. Treatment of RSV-infected cells with arsenite resulted in the appearance of aggregates of TIA-1, indicative of SGs. In addition, the arsenite treatment resulted in IBs that were smaller and more numerous than in non-arsenite-treated cells, as already noted (Fig. 7A and 8A). The appearance of SGs in RSV-infected cells following arsenite treatment indicated at least a partial recovery of OGT function.

In arsenite-treated RSV-infected cells, the IBs and SGs were scattered in the cytoplasm, with no apparent colocalization (Fig. 7A and 8A). The RSV P protein showed clear spatial segregation with the IBs but did not colocalize with the SGs (Fig. 7A). The RSV M2-1 protein was similarly present in IBs but, surprisingly, was also found colocalized with TIA-1 in the SGs that formed in response to arsenite treatment (Fig. 8A). The lack of colocalization of IBs and SGs, and the difference between the P and M2-1 proteins with regard to colocalization with SGs, was further illustrated by cross-sectional analysis and three-dimensional modeling of the fluorescence confocal images, shown both as selected images (Fig. 7B and 8B, respectively) and as animations (see Videos SA and SB, respectively, in the supplemental material). Thus, when SGs formed in RSV-infected cells following arsenite treatment, the M2-1 protein translocated to the newly forming SGs. Whether the SG-associated M2-1 protein was derived from IBs or had been free in the cytoplasm is unknown.

To further investigate the effect of OGT sequestration within the viral IBs, we investigated the intracellular distribu-
tion of OGN, marking the locations of proteins that had been modified by OGT. A549 cells were infected or mock infected in the same way as in previous experiments, and at 18 h postinfection immunofluorescence was performed with antibodies against RSV, OGT, and OGN (Fig. 9) or with RSV, OGN, and the SG marker eIF3 (Fig. 10). In these experiments, limitations in the available primary and secondary antibodies precluded the use of murine antibodies that were monospecific to RSV P.

**Fig 4** Immunofluorescence imaging and intensity profiles of viral IBs. A549 cells were infected with wt-rRSV at an MOI of 0.5 or were mock infected. At 18 h postinfection, the cells were fixed and stained with antibodies for either the viral M2-1 (A, green) or P (B, green) protein and p38-P (red). Nuclei were stained with DAPI (blue). The samples were analyzed by confocal microscopy. In panels A and B, the bottom row of images shows enlargements from the middle row as well as cross-sections in the x and y dimensions. The panels on the right show a single mid-cell layer from a z-stack of confocal images (top images). The white line with arrowheads below each end indicates the track of a line intensity profile across one (A) or two (B) selected IBs (bottom images).
or M2-1, and instead we used polyspecific goat antibodies against RSV virions; as a result, RSV-specific staining was not limited to IBs.

In mock-infected cells, OGT was distributed diffusely (Fig. 9A; the diffuse pattern is not visible because the signal intensities of all of the images were reduced equally to avoid saturation of the more intense signals). In the same cells, OGN was faintly detected, with small punctate spots and diffuse patches in the cytoplasm. When mock-infected cells were treated with arsenite, the distribution of OGT was not detectably changed, but OGN was evident in a large number of cytoplasmic aggregates (Fig. 9A), indicating the enzymatic activity of OGT. Immunostaining for eIF3 identified these aggregates as SGs (Fig. 10).

In RSV-infected cells, OGT was concentrated in IBs, as already noted (Fig. 7 and 8), whereas the accumulation of OGN was diffuse and similar to that in mock-infected cells (Fig. 9A). Arsenite treatment of RSV-infected cells resulted in the accumulation of OGN in SGs, identified by colocalization with eIF3 (Fig. 10). The confocal images of arsenite-treated, RSV-infected cells from Fig. 9A and 10A were illustrated by cross-sectional analysis and reconstructed as three-dimensional images, shown in Fig. 9B and 10B, respectively. This further illustrated that OGN showed clear colocalization with eIF3 in SGs (Fig. 10B), whereas OGT did not colocalize with SGs (Fig. 9B), as already shown.

These results showed that RSV infection resulted in the sequestration of OGT in viral IBs but that the size of the IBs was greatly

**FIG 5** Western blot analysis of the intracellular expression of MK2 and phosphorylated MK2 (MK2-P) in response to RSV infection and treatment with inhibitor CMPD-1. (A) Monolayer cultures of A549 cells were infected with wt-rRSV at an MOI of 0.5 or were mock infected. Replicate cultures were treated with the inhibitor CMPD-1 or mock treated, beginning 30 min before infection (~30 min) or 75 min postinfection (~75 min). Mock-infected cells were treated with arsenite for 30 min at 18 h postinfection as a control. The cultures were harvested and analyzed by Western blotting with antibodies against MK2/MK2-P and β-actin as a loading control. (B) A549 cell cultures were infected or mock infected as for panel A and were treated with arsenite or mock treated for 30 min at 18 h postinfection. The cells were harvested and analyzed by Western blotting with antibodies against MK2/MK2-P, and β-actin was used as a loading control.

**FIG 6** Immunofluorescence analysis of the intracellular distribution of p38-P and viral P (A) or M2-1 (B) protein in response to RSV infection and treatment with arsenite. Monolayer cultures of A549 cells were infected with wt-rRSV at an MOI of 0.5 or were mock infected. Cultures were treated with arsenite or mock treated for 30 min at 18 h postinfection, fixed, and immunostained using antibodies to the P (A, green) or M2-1 (B, green) protein and p38-P (red). Nuclei were stained with DAPI (blue). Samples were analyzed by confocal microscopy.
FIG 7 Immunofluorescence analysis of the intracellular distribution of OGT, the viral P protein, and the SG marker TIA-1 in response to RSV infection and treatment with arsenite. Monolayer cultures of A549 cells were infected with wt-rRSV at an MOI of 0.5 or were mock infected. Cultures were treated with arsenite or mock treated for 30 min at 18 h postinfection, fixed, and immunostained using antibodies specific to the P protein (green), OGT (red), and TIA-1 (violet), and nuclei were stained with DAPI (blue). Panel B shows enlargements from the bottom row of panel A, with further enlargements rendered as three-dimensional images. White arrowheads in panel B indicate representative IBs, illustrating that the IBs contain P and OGT but do not localize with TIA-1. An animation of panel B is shown in Video SA in the supplemental material.
Immunofluorescence analysis of the intracellular distribution of OGT, the viral M2-1 protein, and the SG marker TIA-1 in response to RSV infection and treatment with arsenite. Monolayer cultures of A549 cells were infected with wt-rRSV at an MOI of 0.5 or were mock infected. Cultures were treated with arsenite or mock treated for 30 min at 18 h postinfection, fixed, and immunostained using antibodies specific to M2-1 (green), OGT (red), and TIA-1 (violet), and nuclei were stained with DAPI (blue). Panel B shows enlargements from the bottom row of panel A, with further enlargements rendered as three-dimensional images. The white arrowhead in panel B indicates a representative IB containing M2-1 and OGT, but not TIA-1, and the white arrows in panel B indicate representative SGs containing M2-1 and TIA-1. An animation of panel B is shown in Video SB in the supplemental material.
FIG 9 Immunofluorescence analysis of the intracellular distribution of OGT and OGN in response to RSV infection and treatment with arsenite. Monolayer cultures of A549 cells were infected with wt-rRSV at an MOI of 0.5 or were mock infected. Cultures were treated with arsenite or mock treated for 30 min at 18 h postinfection, fixed, and immunostained using antibodies against RSV (green), OGT (red), and OGN (violet). Nuclei were stained with DAPI (blue). Panel B shows enlargements from the bottom row of panel A, with further enlargements rendered as three-dimensional images. Incidentally, the anti-RSV antibodies used in this particular experiment had been raised against whole virus and RSV staining was not limited to IBs; thus, colocalization of viral protein with OGT in IBs (one example is indicated by a white arrowhead in panel B) was much less evident than in other experiments.
diminished by the induction of severe oxidative stress by a brief arsenite treatment. Thus, the sequestration appeared to be reversible. Our data did not reveal any discernible differences in the spatial localization of proteins modified by the addition of OGN in mock- versus RSV-infected cells, although the species of proteins that are OGN modified may differ under these two conditions and merit further investigation.

Negative association between IBs and SGs. We noted that...
RSV-infected cells that contained well-developed, large IBs did not have detectable SGs, as is illustrated in Fig. 11 with the cell marked “a.” In contrast, a small fraction of RSV-infected cells were found to contain small IBs, and these cells also contained SGs, as is illustrated in Fig. 11 with the cell marked “b.” We quantified the relative abundance of these two phenotypes in A549 cells that were infected with RSV at an MOI of 0.5 PFU per cell and analyzed 18 h postinfection (as in all of the experiments in this study). Two replicate experiments were performed: in each, 25 microscopic fields were counted, with a total of approximately 300 RSV-infected cells. The percentages of infected cells with SGs in the two experiments were 5.8% and 4.3%. Thus, most RSV-infected cells formed large, prominent IBs that apparently precluded SG formation, whereas a small fraction of infected cells formed smaller IBs that permitted SG formation. The pattern of small, numerous IBs present in this small subpopulation of RSV-infected cells was similar in appearance to the pattern that was induced in the majority of RSV-infected cells by brief asenite treatment (Fig. 7 and 8): in each case, the presence of small IBs was associated with the presence of SGs, and there was a negative association between large, prominent IBs and SGs.

**DISCUSSION**

The MAPKs, including p38 MAPK, ERK, and Jun N-terminal protein kinase (JNK), represent interrelated pathways involved in cellular homeostasis and responses to external stimuli, including viral infection. Previous evidence indicated roles for p38 MAPK and ERK, in particular, in RSV infection (11, 12, 34). Our study demonstrated that RSV interferes with the p38 signaling pathway by spatially sequestering p38-P into viral IBs. p38 has been shown to be indispensable in the stress- and cytokine-dependent activation of MK2. In addition, the tight binding between MK2 and p38 is required for the stable accumulation of both proteins (20, 21). Consistent with this, one of the outcomes of the sequestration of p38-P in RSV IBs in the present study was the extensive depletion of MK2, effectively achieving an MK2 knockdown in RSV-infected cells. MK2 has been shown to be an essential component in the inflammatory response. For example, inactivation of the mouse MK2 gene by a targeted mutation resulted in severely reduced production of TNF-α and IFN-γ and significantly reduced production of IL-1β, IL-6, and IL-10 in response to lipopolysaccharide stimulation (16). Many of these proinflammatory proteins are encoded by mRNAs marked for rapid degradation by AREs found in their 3' untranslated regions (UTRs), and the activation of p38/MK2 pathway results in the stabilization of these ARE-containing mRNAs (15, 17, 35). Specifically, activated MK2 mediates phosphorylation of the cellular zinc finger protein tristetrapolin (TTP). While unphosphorylated TTP binds to and promotes the degradation of ARE-containing mRNAs, phosphorylation of TTP by MK2 promotes complex formation between TTP-P and the chaperone protein 14-3-3, which excludes TTP from SGs and thereby inhibits TTP-dependent degradation of ARE-containing mRNA (13, 36). Thus, viral interference with signal transduction through p38 leading to MK2 depletion would reduce the production of anti-inflammatory and antiviral factors.

We also showed that RSV infection resulted in the sequestration into viral IBs of another key player in intracellular signaling, OGT. OGT catalyzes posttranslational attachment of single OGN molecules to serine and threonine residues of cellular proteins in a dynamic interplay with their phosphorylation status, thereby altering protein stability, interactions, spatial distribution, and activity. In particular, OGT is known to be involved in modifying the translation machinery during stress induction by supporting the assembly of SGs and processing bodies (25, 26). SGs are not constitutively present in cells but can form under conditions of environmental stress or infection. They are complex cytoplasmic foci that contain stalled translation complexes that subsequently can be directed for translation or degradation. Thus, SGs play an important role in posttranscriptional control of gene expression and have particular importance for ARE-containing mRNAs, including those of proinflammatory and antiviral cytokines and other proteins. Sequestration of OGT in IBs would presumably preclude
its role in cellular regulation and, in particular, would prevent OGT from triggering the formation of SGs.

The sequestration of both p38-P and OGT into RSV IBs appeared to be partly reversed by oxidative stress induced by a brief treatment with arsenite. In both cases, this led to an apparent shrinkage and fragmentation of the IBs. Furthermore, arsenite treatment was associated with the activation of the remaining MK2, as well as the formation of SGs and the addition of OGN to target proteins within SGs. This suggested that the apparent fragmentation of IBs in response to arsenite treatment may have resulted in the release of p38-P and OGT; if so, this indicates that these proteins had been transported to IBs and sequestered but had not been damaged. Arsenite disrupts ATP production, among other things, and its ability to reduce and scatter IBs and release p38-P and OGT suggests that the maintenance of IBs and the sequestration of p38-P and OGT are highly dynamic processes that require energy.

Examination of RSV-infected A549 cells showed that ~95% of infected cells contained prominent IBs with no detectable SGs, whereas the remaining ~5% of infected cells contained smaller IBs as well as SGs. This suggests that the formation of large, prominent IBs precludes the formation of SGs. This could be explained by the observed efficient sequestration of OGT by large IBs, making OGT unavailable to induce SGs. In contrast, in the few infected cells in which IB formation was less developed, sequestration of OGT was less prominent, leaving free OGT available to induce SG formation. This suggests that RSV infection is a stimulus for SG formation, but this normally is blocked by virus-mediated sequestration of OGT. In the small fraction of RSV-infected cells that contained smaller IBs as well as SGs, the formation of large IBs may have been delayed or reduced, perhaps due to delayed or reduced viral replication in those particular cells. This might result from variability in the cell population, including factors such as host defense or permissiveness to infection, variability in the virus population, including the presence of defective interfering particles in those cells, or other stochastic factors.

Several recent studies have noted SGs in the context of RSV infection (8, 30, 37). Lindquist et al. reported SG formation in RSV-infected cells and suggested that SGs enhanced RSV replication (8). However, they observed that SGs occurred in only a small percentage of RSV-infected cells: for example, at an MOI of 1, only approximately 10% of the infected cells contained SGs at 16 h postinfection (8), which is similar to the findings in the present report (i.e., ~5% of cells at 18 h postinfection with an MOI of 0.5). Thus, the formation of SGs does not appear to be a regular feature of RSV-infected cells, and the association of SGs with enhanced RSV replication was not confirmed in a subsequent study (38). Lindquist et al. reported that the size of the IBs was larger in the small fraction of cells containing SGs than in the large fraction of cells lacking SGs. However, this size difference was small, with an average IB volume of ~0.4 μm³ in infected cells without SGs to ~0.6 μm³ in infected cells with SGs, which translates to an average radius of ~0.4 μm versus ~0.5 μm. In the present work, we observed that the large majority of RSV-infected cells contained large, prominent IBs and did not contain SGs. These large IBs ranged in size between 2 and 3 μm in radius and thus were far bigger than the ones described by Lindquist et al. We observed dense accumulation of OGT in these large IBs, suggesting that they were very efficient in sequestering OGT and thereby inhibiting SG formation. The reason for the difference in IB size between the report of Lindquist et al. and the present study is not clear. It may be relevant that the measurements were taken at different times during infection: our study focused on 18 h postinfection, which is at or slightly following the peak of viral RNA expression, when the effects of IBs on RSV replication might be the most relevant, whereas the measurements of Lindquist et al. were made at a later time, 24 h postinfection. Lindquist et al. documented a substantial increase in SG formation at this late time point, and it may be that the cells were experiencing increased stress due to the well-advanced viral infection, which may have affected IB size, among other things.

Hanley et al. also described SGs in the context of RSV infection (30). This study similarly showed that SGs formed in only a small percentage (~5%) of cells infected with wild-type RSV, whereas SGs were much more frequent in cells infected with an RSV mutant that lacked most of the trailer region and exhibited reduced replication. These results are consistent with the present study: specifically, both studies suggest that RSV infection has the potential to induce SG formation but that the wild-type virus suppresses this induction. It may be that the trailer sequence that is missing in the mutant RSV of Hanley et al. is important for suppressing SG formation, or it may be that the mutant failed to restrict SG formation because of reduced replication and hence reduced expression of viral proteins. In any event, our work differs from these previous studies in providing a mechanism for how RSV suppresses SG formation, namely, by the sequestration of OGT, an upstream regulatory molecule in SG assembly, into the viral IBs.

Several other recent studies had noted an association between host cell molecules and RSV IBs. In the first of these reports, Brown et al. noted the appearance of heat shock protein Hsp70 within virus-induced IBs (9). Lindquist et al. also made the interesting observation that HuR, a component of SGs, localized to IBs during RSV infection, although this did not appear to contribute to RSV infection, since knockdown of HuR had no effect on viral replication (8). Another recent study that was published while the manuscript for this article was in preparation provided evidence of the sequestration of additional cellular proteins in IBs, specifically, melanoma differentiation-associated gene 5 (MDA5) protein and mitochondrial antiviral signaling (MAVS) protein (39). This was associated with suppression of IFN-β production. Our present results provide evidence that the sequestration of components of innate immune response systems and signal transduction pathways into viral IBs is a prominent feature of RSV infection (and, by analogy, possibly for other viruses, such as human metapneumovirus and measles virus, that also form viral IBs).

The observation that RSV targets and sequesters p38 and OGT is particularly noteworthy because these proteins may have a broad impact on host cell signaling and affect a wide range of cellular activities beyond the stability of ARE-containing mRNAs and the formation of SGs, including a myriad of other targets that are necessary for viral translation (40) and regulation of the adaptive immune response (41). For example, p38 activates many other protein kinases in addition to MK2, including MK3, MK5, MSK1/2, and MNK1. Moreover, transcription factors such as ATF-2, SAP1, CHOP, STAT1, NFAT, GADD153, and numerous others are downstream substrates of p38 phosphorylation (42). Similarly, OGN-modified proteins affect a variety of cellular functions, including transcription, translation, protein processing, growth, metabolism, stress, and the immune response (40, 41, 43–45). As a specific example, under IL-1 induction or osmotic...
stress, OGN modification of TAB1 has been shown to modulate TAK1 activation, leading to IκB kinase (IKK) phosphorylation and NF-κB translocation (46). By confining OGT, RSV could potentially prevent the activation of TAB1/TAK1 pathway and therefore influence NF-κB-induced cytokine production.

We were surprised to find that following arsenite treatment, the viral M2-1 protein appeared in SGs. M2-1 was previously shown to function as a viral elongation/anti-termination factor that is necessary for the synthesis of full-length viral mRNAs and is essential for RSV replication (47). M2-1 contains a Cys2-His1 (CCCH) motif that is essential for its function in viral RNA synthesis (48, 49). Interestingly, it was previously noted that this CCCH motif is a feature that M2-1 shares with TTP (48, 49). The present study takes this analogy further by showing that like TTP, M2-1 can be recruited to SGs. This raises the possibility that M2-1 may have TTP-related roles in affecting SG formation and activity and RNA metabolism. This is under investigation.

In summary, we found that p38 and OGT, two key cellular molecules that are centrally involved in posttranslational protein modification and signal transduction, are sequestered inside RSV IBs during infection. Given the wide-ranging roles of these cellular proteins, their sequestration has the potential to substantially affect cellular activities during RSV infection. In the present study, we showed that this resulted in suppression of SG formation and was an active, reversible process. Future work is needed to study the full impact of p38/OGT sequestration and to identify additional host molecules that may be sequestered during RSV infection. Furthermore, this work identified an interaction between the viral M2-1 protein and SGs that potentially suggests a TTP-related role for this viral protein.

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REFERENCES


