Respiratory syncytial virus (RSV) is a significant cause of lower respiratory tract disease in children and individuals with cell-mediated immunodeficiencies. Airway epithelial cells may be infected with RSV, but it is unknown whether other cells within the lung permit viral replication. We studied whether human alveolar macrophages supported RSV replication in vitro. Alveolar macrophages exposed to RSV demonstrated expression of RSV fusion gene, which increased in a time-dependent manner and correlated with RSV protein expression. RSV-exposed alveolar macrophages produced and released infectious virus into supernatants for at least 25 d after infection. Viral production per alveolar macrophage declined from 0.053 plaque-forming units (pfu)/cell at 24 h after infection to 0.003 pfu/cell by 10 d after infection and then gradually increased. The capability of alveolar macrophages to support prolonged RSV replication may have a role in the pulmonary response to RSV infection.
Productive Infection of Isolated Human Alveolar Macrophages by Respiratory Syncytial Virus

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Abstract
Respiratory syncytial virus (RSV) is a significant cause of lower respiratory tract disease in children and individuals with cell-mediated immunodeficiencies. Airway epithelial cells may be infected with RSV, but it is unknown whether other cells within the lung permit viral replication. We studied whether human alveolar macrophages supported RSV replication in vitro. Alveolar macrophages exposed to RSV demonstrated expression of RSV fusion gene, which increased in a time-dependent manner and correlated with RSV protein expression. RSV-exposed alveolar macrophages produced and released infectious virus into supernatants for at least 25 d after infection. Viral production per alveolar macrophage declined from 0.053 plaque-forming units (pfu)/cell at 24 h after infection to 0.003 pfu/cell by 10 d after infection and then gradually increased. The capability of alveolar macrophages to support prolonged RSV replication may have a role in the pulmonary response to RSV infection. (J. Clin. Invest. 1990.86:113–119.) Key words: mononuclear phagocyte • viral replication • fusion gene • respiratory virus • long-term cultures

Introduction
Respiratory syncytial virus (RSV) is a major cause of childhood bronchiolitis and pneumonia and demonstrates peculiar tropism for the lower respiratory tract of young children (1). RSV may induce an airway inflammatory response (2) as seen in pathological studies, which have demonstrated RSV antigen-expressing cells in alveolar lumina (3, 4) and increases in lymphocytes and mononuclear phagocytes (MP) in lung specimens from children with RSV-induced bronchiolitis (5, 6). Although tracheal epithelial cells are permissive to RSV infec-

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1. Abbreviations used in this paper: F, fusion; MP, mononuclear phagocytes; pfu, plaque-forming units; RSV, respiratory syncytial virus.

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tion in vitro (7), little is known about the replication of RSV in other pulmonary cells, particularly those involved in lung defense.

The alveolar macrophage is thought to have a major role in defense of the respiratory tract against invading pathogens, including virus (8–11). Alveolar MP may be exposed to RSV via the airstream or by virus replication within the respiratory tract (3, 9). In a previous study, we demonstrated that alveolar and blood MP exposed to RSV in vitro can express viral proteins, indicating that these cells are permissive to RSV infection (12). However, infection may result in distinct outcomes, including latent, abortive, or persistent infections characterized by absent or low levels of release of new virus (13). Viral infection may also result in cytoplytic and/or productive infection generally resulting in the release of new progeny virus (14). These separate outcomes of viral infection may have unique consequences for the host. Since the nature of the viral replicative cycle in alveolar MP has potential effects on the pulmonary immune response to RSV, we felt that it was important to determine which of these possible outcomes follows RSV infection of alveolar MP.

RSV infection of alveolar macrophages was studied initially by monitoring the expression of RSV fusion (F) RNA and the expression of viral proteins. The RSV F gene of 1.9 kb in length is transcribed eight genes downstream from the RSV promoter (15), and F protein is thought to promote intercellular viral spread (16–19). It was further determined whether RSV-infected alveolar MP produced and released infectious virus. The data obtained indicate that human alveolar MP support all stages of the RSV replicative cycle and produce infectious virus for up to 25 d after infection.

Methods
Preparation and RSV infection of monocytes and alveolar macrophages. 12 healthy, nonsmoking adult volunteers each underwent venipuncture and bronchoalveolar lavage using standard techniques. Informed consent was obtained from each participant and studies were approved by the institutional review board, University Hospitals of Cleveland. PBMC adherent to plastic petri dishes precoated with AB sera (Whittaker M. A. Bioproducts, Walkersville, MD) were prepared after Ficol-Hypaque centrifugation as previously described (20).
Alveolar macrophages were isolated from bronchoalveolar lavage fluid by minor modifications of methods previously described (21). After bronchoscopy, segments of the right middle and lower lobes were lavaged with sterile saline (0.9%, 240 ml, 37°C) in 60-ml aliquots followed by immediate aspiration. Lavage fluid was centrifuged at 500 g for 10 min, and the cell pellet was washed twice with RPMI-1640 (Gibco Laboratories, Grand Island, NY). Alveolar macrophages were isolated by adherence to plastic precoated with AB sera for 1 h at 37°C. Nonadherent cells were removed by washing three times with cold HBSS (Gibco Laboratories). Alveolar and blood MP preparations were >95% viable and >90% nonspecific esterase positive.
Blood and alveolar MP (5 × 10⁶ cells/0.5 ml) were allowed to
adhere to 12-well tissue culture plates (Costar, Cambridge, MA) and then exposed for 2 h at 37°C to Eagle's MEM (Whitaker M. A. Bioproducts) containing 2% (vol/vol) FCS (Hyclone, Logan, UT) or to this medium containing RSV at a multiplicity of infection of 3 plaque-forming units (pfu)/cell. Cells were then washed once in RPMI without serum and cultured in RPMI containing 10% (vol/vol) FCS and 2 mM glutamine, 1 mM nonessential amino acids, 100 U/ml penicillin, 100 μg/ml streptomycin, and 0.25 μg/ml amphotericin B (culture medium) at 37°C in 5% CO₂. Culture medium from individual, parallel wells was recovered at 2-, 6-, 24-, and 120-h intervals, and after centrifugation at 500 g cell-free supernatants were frozen at −70°C for determination of viral titers. Adherent alveolar MP were washed once in culture medium at each time interval and harvested by gentle scraping with a rubber policeman for determination of viability, immunofluorescent staining, and viral titer. Parallel uninfected and infected cultures were washed once and incubated in fresh culture medium at each 5-d interval for the 25 d of study.

Viability of MP cultures was determined after resuspension of the cells in 0.05% (wt/vol) trypsin blue in PBS and enumeration of the percentage of cells that excluded the dye by light microscopy. In selected experiments, viability of adherent MP cultures was assessed by fluorescent microscopy after staining with acridine orange and ethidium bromide, each dye at 1 μg/ml in PBS, for 1 min. Acridine orange stains the nuclei of viable cells green, while ethidium bromide stains the nuclei of nonviable cells orange (22). The results obtained by these separate assays for viability agreed within ±15%. Latex particle ingestion by MP cultures was performed as previously described (20).

RSV preparation and viral titer. Respiratory syncytial A2 strain virus was propagated in CV-1 cells (African green monkey kidney cell line; American Type Culture Collection, Rockville, MD) in Eagle’s MEM as previously described (12). Virus titer was determined by incubation of serial dilutions of virus stocks, cell-free supernatants, or sonicated alveolar MP (15 s twice, sonic dismembrator; Artex Systems Corp., Farmingdale, NY) on CV-1 monolayers, overlaid with 0.5% (wt/vol) agarose and stained with 2% (wt/vol) crystal violet in 10% (vol/vol) formalin after 5 d of incubation. Inactivated virus was prepared by exposure to ultraviolet (UV) light (15 W) at a distance of 10 cm for 1 h or by storage at room temperature for 7 d. Viral titer per cell was determined from total virus in supernatant and disrupted cells divided by the number of viable cells times the percentage of infected cells for each donor at each time point.

Immunofluorescent staining. MP were fixed while plastic adherent with 100% methanol at 4°C for 10 min, or harvested with a rubber policeman, resuspended in 0.075 ml of PBS, and applied to microscope slides (5 × 10⁵ cells/slide), air dried, and fixed with cold acetone for 10 min at 4°C. The cells were reacted for 30 min with either a mixture of mouse MAB to RSV proteins conjugated to FITC (Bartels Immunodiagnostic Supplies Inc., Sacramento, CA) or a monospecific rabbit antibody to F (a kind gift of Samuel Levine, Wayne State University, Detroit, MI) followed by a goat anti-rabbit F(ab)₂ antibody (rhodamine labeled; Southern Biotechnology Associates, Inc., Birmingham, AL). The cells were washed three times in PBS, dried, and mounted with PBS/glycerol (1:1; vol/vol), and analyzed by phase contrast and epi-fluorescent microscopy.

RNA isolation, Northern blot analysis, and whole cell dot blot analysis. RNA was extracted from cellular pellets by published methods (23) using 4 M guanidine isothiocyanate (IBI, New Haven, CT) and centrifugation on a solution of 5.7 M cesium chloride (Sigma Chemical Co., St. Louis, MO) in a rotor (SW 50.1; Beckman Instruments, Inc., Palo Alto, CA) at 36,000 rpm for 16 h. Total RNA, quantitated spectrophotometrically, was electrophoresed through 0.9% agarose formaldehyde gels, stained with ethidium bromide (Sigma Chemical Co.), and transferred to Genescreen membranes (New England Nuclear, Boston, MA) (23). Size estimation of the RNA transcripts was measured by comparison with a formamide denatured Bst EII lambda digest DNA ladder (New England Biolabs, Beverly, MA). The membranes were baked at 80°C for 2 h.

Whole cell RNA dot blots were performed with minor modifications of published techniques (24). Briefly, twofold serial dilutions of cells in PBS were applied by vacuum to Genescreen membranes pre-wet in 2× SSC, 10% (vol/vol) formaldehyde (EM Science, Gibbstown, NJ) on a dot blot minifold (Schleicher & Schuell, Inc., Keene, NH). The following solutions were sequentially passed through the filter: 100 ml of 2× SSC, 0.2% SDS; 100 ml of 2× SSC, 15% (vol/vol) of formamide (Bethesda Research Laboratories, Gaithersburg, MD) at 60°C; and 100 ml of 2× SSC. The membranes were then incubated in 10% (vol/vol) formaldehyde, 20× SSC at 60°C for 15 min, followed by two washes (100 ml each) in 2× SSC, and then baked at 80°C for 2 h.

Prehybridization was performed at 42°C for 16 h in 35% formamide, 1× Denhardt's, 2× SSC, 0.5% SDS, and 50 μg/ml heat-denatured salmon sperm DNA (Sigma Chemical Co.). Hybridization was performed with a 32P-labeled cDNA probe for a fragment of RSV F cloned in pUC 9 (16) in the above solution without salmon sperm DNA containing 10 mM EDTA. The probes were labeled by random primer extension (Boehringer Mannheim Biochemicals, Indianapolis, IN) and specific activities exceeded 10⁶ cpm/μg (25). After hybridization, the membranes were washed sequentially in 2× SSC, 0.5% SDS; 1× SSC, 0.5% SDS; and finally, 0.1× SSC, 0.5% SDS at 60°C, dried, and exposed to Kodak XAR-5 film with two intensifying screens at 70°C.

Statistical analysis. All values are represented as mean±SEM. Comparisons between means were analyzed for statistical significance by paired t test or the Wilcoxon matched-pairs signed rank test, and were considered significant at P < 0.05.

Results

Presence of viral F RNA in alveolar macrophages, monocytes, and CV-1 cells after RSV exposure. Using a cloned cDNA probe for RSV F RNA, expression of this gene in total RNA extracted from autologous blood and alveolar MP pooled from four donors, and CV-1 cells after exposure of each cell type to RSV for 36 h at 0 or 3 pfu/cell was determined. Equivalent amounts of total RNA were hybridized with a [32P]cDNA probe for F RNA (Fig. 1). F RNA was readily detected as a 1.9-kb band in RSV-exposed monocytes (lane 2), alveolar MP (lane 4), or CV-1 cells (lane 6), whereas no detectable F RNA was present in the corresponding cultures not exposed to RSV (lanes 1, 3, and 5) or in CV-1 cells exposed to UV light-inactivated RSV at the same viral dose (lane 7). RNA from monocytes demonstrated a nonspecific hybridization signal that comigrated with the 28 S RNA species and was present at equal intensities in both uninfected and infected cells. Longer exposures of Northern blots also demonstrated hybridization signals of ~ 17 kb in size only in lanes exposed to RSV, probably indicating genomic RNA (not shown).

Figure 1. Northern blot analysis of monocyte, alveolar macrophage, and CV-1 RNA. Northern analysis of total RNA, evaluated using a [32P]cDNA probe for RSV F RNA. Lane 1, monocytes; lane 2, monocytes exposed to RSV; lane 3, alveolar MP; lane 4, alveolar MP exposed to RSV; lane 5, CV-1 cells; lane 6, CV-1 exposed to RSV; lane 7, CV-1 exposed to UV-inactivated RSV. The size of the F RNA was determined using RNA markers (see Methods). Exposure times of the autoradiograms were 12 h for monocytes, 4 h for alveolar MP, and 1 h for CV-1 cells.
F RNA expression in alveolar and blood MP and RSV protein expression. Expression of RSV F RNA was directly compared at 12 and 24 h after infection from equal numbers of autologous alveolar and blood MP (each at 5 × 10^5 cells/donor, four separate donors). Although expression of F RNA by alveolar and blood MP differed within a given donor (for example, in Fig. 2), no significant differences were seen when the results from four donors were combined. These results suggest that both alveolar and blood MP support RSV transcription and/or replication.

Separate aliquots of RSV exposed cells were analyzed for F protein expression (indirect immunofluorescence) and demonstrated undetectable F protein after 2 h exposure to RSV. However, F protein was readily detectable in cells exposed for 24 h (not shown). Alveolar MP uninfected or infected with RSV and fixed while adherent after 15 d of culture demonstrated similar morphology containing both rounded and elongated, spindle-shaped cells by phase contrast microscopy (Fig. 3 A). Approximately 30% of the RSV-infected cells continued to express RSV proteins as determined by epi-fluorescent microscopy of the same field, as shown in an example from one donor (Fig. 3 B), while uninfected cell cultures did not demonstrate fluorescent staining. Greater than 90% of the viable cells in culture for 15 d were capable of ingesting latex particles consistent with an identification as phagocytes.

Productive infection of alveolar macrophages by RSV. Release of new progeny virus from alveolar MP (five donors) after RSV exposure was determined at 2.6, 24, 48, 72, 96, and 120 h after exposure to RSV at 0 or 3 pfu/cell. Viral titer in cell-free supernatants and in washed and sonicated adherent cells was determined at each time point. Viral titer in supernatants or disrupted cells declined by >2 logs by 6 h after infection, but then increased by 24 h after infection and remained elevated through 120 h (Fig. 4).

The persistence of viral replication was assessed in alveolar MP (5 × 10^5 cells/time point, five additional donors) after exposure to RSV at 0 or 3 pfu/cell at 6, 24, 120, 240, 360, 480, and 600 h after infection. Medium was changed at each 120-h interval and viral titer determined in cell-free supernatants or disrupted alveolar MP. Alveolar MP contained infectious virus and released virus to the extracellular milieu throughout this period, 600 h (25 d) (Fig. 4). Viral titer released from sonicated cells at 6 h after infection was 198±115 pfu/ml, which was significantly less than that released at 24 h after infection, 6,218±2,211 pfu/ml (P < 0.01). Total virus titer, that present in supernatant plus disrupted cells, at 24 h after infection was 7,335±2,416 pfu/ml, which was increased significantly compared with viral titer present in the cultures at 6 h after infection, 565±246 pfu/ml (P < 0.05). Taken together, these results indicated that RSV-infected alveolar MP produced new progeny virus.

Alveolar MP from all 10 donors used in these experiments permitted RSV replication. Uninfected alveolar MP cultured in parallel from each donor demonstrated undetectable virus, indicating that the alveolar MP did not harbor virus at the time of bronchoalveolar lavage (not shown). Additionally, RSV exposure of medium-control cultures in the absence of alveolar MP had undetectable viral titers after 24 h at 37°C.

Next we examined whether RSV exposure of alveolar MP resulted in cytopathic infection. There were no differences in mean viable cell number of alveolar MP between uninfected and infected cultures at any time point from these donors (Table I). Total virus (supernatant + sonicated cells) produced per infected (immunofluorescent positive), viable alveolar MP was significantly higher at 24 h after infection than that produced at 120–360 h after infection (P < 0.05). By 480 and 600 h after infection viral production per cell did not significantly differ from that seen at 24 h after infection.

Discussion

Although human monocytes can be productively infected by RSV (26), little is known about the RSV replicative cycle in these cells, nor whether alveolar macrophages differ in their ability to support productive infection. The results presented herein indicate that alveolar macrophages support each stage of RSV replication, resulting in production of infectious virus for prolonged periods.

Northern analysis of RSV-infected blood and alveolar MP and the CV-1 cell line demonstrate the presence of RSV F RNA whose size is consistent with the known size of the transcript for F gene (16). Using this probe, total RSV F RNA accumulation in alveolar MP exceeded that present in blood MP at 36 h after infection; however, comparable levels of F RNA were expressed by alveolar and blood MP at 12 and 24 h after infection. Either differences in the amount or stability of RSV RNA in MP by 36 h after infection, or differences in the viability of RSV-infected MP could account for these results. However, RSV F expression correlated with the time-dependent RSV protein production, indicating that alveolar MP support both transcription and protein expression of RSV genes.

In an earlier study we found that alveolar and blood MP were equally permissive to RSV infection, while cord blood MP were significantly more susceptible to infection with this virus (12). The data presented here on adult blood and alveolar MP confirm and extend these previous observations and demonstrate that alveolar MP cultured in vitro produce infectious virus for prolonged periods after RSV exposure. These combined results indicate that RSV infection and replication in MP may have a role in the extrapulmonary dissemination of RSV and prolonged virus shedding seen in some immunodeficient individuals (27).

Replication of RSV in alveolar MP, a tissue-derived component of the human MP series, extends previous results that demonstrated RSV replication in blood MP and the U937
Figure 3. RSV protein expression in alveolar macrophages exposed to RSV. A. Phase contrast photograph of RSV-exposed alveolar MP after 15 d of culture. B. Photograph of the same field of cells after immunofluorescent staining with mouse MAb to RSV proteins (FITC-labeled).
monocytic cell line (26, 28). The present results contrast with previous work that demonstrated resistance of bovine alveolar MP to RSV replication, perhaps suggesting species-specific differences in productive RSV infection of alveolar MP (29, 30).

Unexpectedly, RSV-infected alveolar MP showed no alteration in viability compared with uninfected controls. This contrasts with results generally seen in cells infected with RSV, where extensive cytolysis usually occurs by 5–7 d after infection (1, 3). MP infected with herpes simplex (31) or Sendai virus (32) (also a member of the paramyxoviruses, as is RSV) have been shown to produce virus for prolonged periods without an apparent effect on MP viability. We find that alveolar MP cultures contain subpopulations of cells capable of replicating or resisting RSV infection for periods up to 600 h (25 d).

The factors determining the ability of alveolar MP to resist RSV infection or support replication will require further investigation.

The mechanism(s) by which MP support replication of some virus for prolonged periods (> 2 wk), including both DNA and RNA virus (31–37), are only partially understood. Proposed mechanisms include production of noninfectious virions that require homologous infectious virus for replication (defective interfering particles) (38), production of temperature-sensitive viral mutants (39), host cell viral resistance factors, a steady-state situation maintained by equilibrium between infected and uninfected cells (38), failure of the host cell to efficiently cleave viral glycoproteins resulting in reduced viral production, and production of antiviral proteins such as interferon, thus reducing viral production (reviewed in reference 40). Any of these mechanisms could potentially account for the findings of prolonged replication of RSV in alveolar MP cultures; however, in this, and in a previous study (12), syncytial formation has not been observed, suggesting inefficient cleavage of fusion protein. This may have a role in the prolonged replication of RSV in these cultures. It is also possible that cytokines produced by alveolar MP, including tumor factor, which directly restricts RSV infection of human blood MP (12), have a role in modulating viral replication in alveolar MP (41). Whether these latter observations have a central role in the mechanism of prolonged RSV replication in alveolar MP cultures will require studies at the individual cell level.

Viral production per cell is initially higher in RSV-infected alveolar MP at 24 h after infection than at 120–360 h after infection. This is probably not due to cell-associated virus, since repetitive washing of RSV-infected alveolar MP does not reduce virus released from disrupted cells. It is possible that the initial RSV replicative cycle in alveolar MP is more efficient than subsequent cycles, which could account for the observed decrease in plaque-forming units per cell. Alternatively, alveolar MP from bronchoalveolar lavage are a heterogeneous population differing in cell density and functional characteristics such as interleukin 1 and PGE2 production (42, 43). Viral replication may proceed at varying rates within these subpopulations, but with continued differentiation induced by plastic adherence, an eventual steady-state level of viral production may be seen. At 400–600 h after infection we observed an increase in plaque-forming units per cell, probably due to release of virus from nonviable cells.

Alveolar macrophages are felt to have a major role in pulmonary defense (8, 44, 45). Previous authors have demonstrated that RSV infection of monocytes alters the production of interleukin 1 and interferon (46, 47). Blood and alveolar MP exhibit important functional differences, including secretion of immunomodulatory and inflammatory mediators (21, 48–50), which may have a role in the pathobiology of RSV infections. Each stage of viral replication within MP could potentially alter important biological functions. Precise understanding of the alveolar MP response to RSV infection may require definition of alveolar MP function at these separate stages of RSV replication.

Acknowledgments

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**Table I. Viability of Uninfected and RSV-infected Alveolar Macrophages and Production of Virus/Viable Cell with Time after RSV Infection**

<table>
<thead>
<tr>
<th>Time in culture (h)</th>
<th>Viable cell number (×10⁶)</th>
<th>% infected cells</th>
<th>Virus production×10⁴ pfu/cell</th>
</tr>
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<tbody>
<tr>
<td>Uninfected</td>
<td>Infected</td>
<td>Infected</td>
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</tr>
<tr>
<td>24</td>
<td>35.6±3.1</td>
<td>38.0±7.0</td>
<td>41.3±3.2</td>
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<tr>
<td>120</td>
<td>36.6±4.3</td>
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<td>240</td>
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<td>35.8±8.0</td>
<td>38.4±0.8</td>
</tr>
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<td>360</td>
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<td>600</td>
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<td>9.4±1.8</td>
<td>31.2±4.6</td>
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The number of viable cells is the product of the total cells recovered times the percentage that excluded trypan blue. Plaque-forming units per cell was determined from the total viral titer (cells + supernatant) divided by the number of viable, infected cells (%) immunofluorescent) determined at the indicated times. Results are the mean±SEM for five donors.

* P < 0.01 vs. 24 h; † P < 0.05 vs. 24 h.
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References


