Adenovirus-Mediated Gene Therapy Enhances Parainfluenza Virus 3 Infection in Neonatal Lambs

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Adenovirus-Mediated Gene Therapy Enhances Parainfluenza Virus 3 Infection in Neonatal Lambs

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Parainfluenza viruses are a common cause of seasonal respiratory disease, but in high-risk individuals (e.g., young children) these viruses can cause severe clinical manifestations that require hospitalization. Beta-defensins are a subclass of antimicrobial peptides with antiviral activity. Use of adenovirus-mediated beta-defensin gene expression has been proposed as therapy for chronic bacterial infections commonly seen in cystic fibrosis patients; however, its use during parainfluenza virus 3 (PIV3) infection has not been evaluated. The hypothesis in this experiment was that adenovirus expression of human beta-defensin 6 (HBD6) would diminish concurrent PIV3 infection in neonatal lambs. The group infected with adenovirus HBD6 and PIV3 had increased levels of pulmonary neutrophil recruitment compared to those for the group infected with PIV3 or PIV3 and adenovirus, with an increased respiration rate and body temperature late in the course of the PIV3-adenovirus HBD6 infection. Interestingly, the adenovirus-treated groups had higher levels of immunohistochemical staining for PIV3 and syncytial cell formation than the group infected with PIV3, suggesting that treatment with the adenovirus vector, regardless of whether it was carrying a target gene, exacerbated the PIV3 infection. The levels of expression of mRNA for antimicrobial surfactant proteins A and D and sheep beta-defensin 1 were increased by PIV3 and adenovirus treatment, and the increased levels of expression roughly corresponded to the degree of inflammation. While pulmonary administration of a high-dose adenovirus vector has been associated with undesirable inflammation, this is the first study to show that it can exacerbate concurrent viral infection, a concern that needs to be addressed for future studies of adenovirus in the lung. Additionally, this study showed that adenovirus-mediated HBD6 expression increases neutrophil recruitment, a recently described attribute of beta-defensins, with mild accentuation of PIV3 activity and inflammation.
cidin antimicrobial peptide (LL-37 [also known as hCAP-18]) showed success in killing bacteria, suggesting that they may be useful for antimicrobial gene therapy in patients predisposed to PIV3 infections (4); however, evaluation of antimicrobial peptide gene therapy as a way to diminish viral infection has not been applied. Human beta-defensin 6 (HBD6), a recently described beta-defensin selectively expressed in the epithelium, was chosen as the antimicrobial peptide in this model due to its lack of expression in the human lung or in sheep (38). The hypothesis of this study was that HBD6 gene expression in the neonatal lamb lung would diminish concurrent PIV3 infection in neonatal lambs.

MATERIALS AND METHODS

Animals. All animal studies were approved by the Iowa State University Animal Care and Use Committee. Neonatal lambs (n = 20) were obtained from Iowa State University’s Laboratory Animal Resources. At 3 to 5 days after birth (which allowed time for acclimation) the lambs were inoculated intratracheally with two 20-ml volumes at approximately 30-min intervals. The order and composition of the inoculum for each group (n = 4) were as follows: group A, sterile medium and sterile medium (control group); group B, sterile medium and PIV3; group C, Ad vector (no gene insertion) and PIV3 (PIV3/Ad group); and group D, Ad HBD6 and PIV3 (PIV3/Ad/HBD6 group). At day 7 of infection the lambs were euthanized by intravenous injection of sodium pentobarbital for collection of tissues. From our preliminary data this time was chosen because it is the time of significant lesion development and active viral clearance. The ovine isolate of PIV3 was grown to 10^7.8 50% tissue culture infective doses/ml (20 ml per lamb), as described previously (20). The replication-defective human Ad 5 vector had a cytomegalovirus promoter and was commercially acquired with the human HB6 gene (identification number 2602099; IMAGE Consortium), the HB6 gene with a lacZ reporter gene, or no gene insert (VirQuest, Inc., North Liberty, Iowa). Ad (no gene insert) and Ad/HBD6 had a deletion of the gene for protein E1 (nucleotides 358 to 3328), and the Ad/HBD6/lacZ insert for 5-bromo-4-chloro-3-indolylphosphate (X-Gal) staining had deletions at nucleotides 358 to 3328 and nucleotides 28592 to 30470, with the concurrent loss of protein E1 and E3 functions. Each lamb received approximately 10^12 virus particles in 20 ml of sterile saline.

Clinical parameters. The lambs were assessed daily for different clinical parameters. Body weight, temperature, and respiratory rate were taken at the same time of day, following the morning feeding. The lambs were treated daily with antibiotic (ceftiofur at 10 mg/day subcutaneously) to prevent complications associated with secondary bacterial infections (34). Tissues. Lung tissue was consistently collected from the cranial and caudal lobes on the left and right sides. The tissue samples were placed in 10% neutral buffered formalin and processed routinely for hematoxylin-eosin staining or immunohistochemistry. In addition, lung tissue was snap-frozen in liquid nitrogen for RNA isolation and real-time fluorogenic PCR.

RNA isolation. RNA isolation and cDNA production were performed as described previously (24). Briefly, 0.3 g of tissue and 3 ml of Trizol reagent (Invitrogen) were homogenized for 30 s, and the homogenate was allowed to sit for 5 min at room temperature. Chloroform (0.06 ml) was added, and the mixture was shaken vigorously for 15 s and then allowed to sit for 3 min at room temperature. This mixture was microcentrifuged for 10 min at 4°C; the resulting top aqueous layer was retrieved and mixed with isopropanol (1.5 ml), and the mixture was allowed to sit for 10 min at room temperature. The solution was again microcentrifuged for 10 min at 4°C, the aqueous-isopropanol layer was removed, and the visible pellet was resolubilized in nuclelease-free high-pressure liquid chromatography-grade water containing 0.1 mM EDTA. Spectrophotometric measurements (detection at absorbances of 260 and 280 nm with 1:40 sample dilutions) of each RNA sample were then taken to assess the isolates for purity and quality.

cDNA production. The RNA samples were treated with a commercially available DNase (RQ1 RNase-free DNase; Promega) to remove potential genomic DNA contamination. Immediately after this, reverse transcription was performed with the reagents available in a commercially available kit (TaqMan reverse transcriptase reagents; PE Applied Biosystems), according to the directions of the manufacturer. Briefly, 2 μg of DNase-treated RNA from each sample and control was added to a reaction mixture which contained final concentrations of 1X TaqMan reverse transcriptase buffer, 5.5 mM MgCl2, 2 mM deoxyribonucleoside triphosphate mixture (500 μM each deoxyribonucleoside triphosphate), 2.5 μM random hexamers, 1.25 U of murine leukemia virus reverse transcriptase per μl, 0.4 to 0.8 U of RNASense inhibitor per μl, and high-pressure liquid chromatography-grade water. The following thermocycler conditions were then used: 10 min at 25°C, 30 min at 48°C, and 5 min at 95°C. The resulting cDNA was stored in nuclelease-free microcentrifuge tubes at 20°C.

Primer and probe design. Sequence-specific oligonucleotide primers and a fluorescent probe were designed with software (ABI Prism Primer Express, version 1.5; PE Applied Biosystems), according to the suggestions of the software manufacturer. The similarities of the resultant potential primer and probe sequences to DNA sequences available in databases were determined by use of the search tool BLAST (Basic Local Alignment Search Tool, version 1.4; National Center for Biotechnology Information), and only unique sequences were used.

### Table 1. Primer and probe sequences for expression of mRNAs for SP-A, SP-D, SBD1, and HBD6

<table>
<thead>
<tr>
<th>Primer or probe specificity and primer or probe</th>
<th>Sequence*</th>
</tr>
</thead>
<tbody>
<tr>
<td>SP-A Forward</td>
<td>5'-TGACCTTATGCTCCTGTGAT-3'</td>
</tr>
<tr>
<td>Reverse</td>
<td>5'-GGGCCATCCAGACAAACCATCCT-3'</td>
</tr>
<tr>
<td>Probe</td>
<td>5'-6FAM-TGGTTCCTGGCCTGAGTGCC-TAMRA-3'</td>
</tr>
<tr>
<td>SP-D Forward</td>
<td>5'-ACGGTTCTGACGTGAGAATT-3'</td>
</tr>
<tr>
<td>Reverse</td>
<td>5'-TCGGTCATGTCAGGAAAGC-3'</td>
</tr>
<tr>
<td>Probe</td>
<td>5'-6FAM-TTGACTGACGTGACAGCCCAGACATGA-TAMRA-3'</td>
</tr>
<tr>
<td>SBD1 Forward</td>
<td>5'-CCATAGGAAATAAGGCTGTGTTG-3'</td>
</tr>
<tr>
<td>Reverse</td>
<td>5'-CGGACAGGTTGCAAATCT-3'</td>
</tr>
<tr>
<td>Probe</td>
<td>5'-6FAM-CCGAGCAGGTGCCCTAGACACATGA-TAMRA-3'</td>
</tr>
<tr>
<td>HBD6 Forward</td>
<td>5'-TCCAATTCCCTCCCAAATGAC-3'</td>
</tr>
<tr>
<td>Reverse</td>
<td>5'-6FAM-TTTGTATGAGAATGTCAGAAATACAT-3'</td>
</tr>
</tbody>
</table>

* 6FAM, 6-carboxyfluorescin (fluorescent reporter dye); TAMRA, 6-carboxytetramethylrhodamine (fluorescent quencher dye).
for primer and probe design (Table 1). Sequence-specific oligonucleotide primers and a fluorescent probe for detection of cDNA corresponding to the endogenous reference gene, 18S rRNA, to which target cDNA levels were normalized, were purchased commercially (TaqMan rRNA control reagents; PE Biosystems). PCR. The 96-microwell plates were designed to enable testing of two replicates of both the target gene and the endogenous 18S rRNA of a 1:5 dilution of cDNA from all lambs, a negative control (diethyl pyrocarbonate-treated water), and five progressive 1:5 dilutions of cDNA from a control sheep for generation of the standard curve. On each plate, the wells with the target gene and the endogenous reference rRNA were run simultaneously by PCR with the tissues represented on that plate. The plate run in a sequence detection system (GeneAmp 5700 version 1.3; PE Applied Biosystems) under conditions identical to those used in the optimization and validation tests. The resultant data gathered by the detection system were exported from the machine to a floppy disk, and all final data processing was performed by using a departmentally designed Excel software file to compare the target to the reference cDNA signals in order to create graphs of the relative levels of mRNA expression.

PIV3 immunohistochemistry. Formalin-fixed, paraffin-embedded sections were deparaffinized in xylene, followed by passage through a series of alcohol baths; and pronase E (0.1 g of protease XIV and 0.1 g of CaCl2 per 100 ml of Tris-buffered saline [pH 7.4]) was applied for antigen retrieval. After the sections were washed (with phosphate-buffered saline), the background blocker (20% normal swine serum), followed by primary antibody (goat polyclonal anti-bovine PIV3, 1:1000 [Veterinary Medical Research & Development, Inc.]), was applied to the tissues. These sections were washed, hydrogen peroxide (3% in methanol) was applied to block endogenous peroxidases, and the sections were washed again. Secondary antibody (1:400 biotinylated rabbit anti-goat antibody) was applied, followed by washing in supersensitive-horseradish peroxidase and chromogen (Vector Red; Vector). The slides were counterstained with hematoxylin, dehydrated in a series of alcohol baths and then with xylene, and placed under a coverslip, to which Permount was applied.

Morphometry. Measurement of microscopic morphological changes, including cellular staining, rate of mitosis, neutrophil density, and syncytial cell formation, were assessed by a pathologist blinded from the study using a grid (18). Cranial and caudal lung lobes (bilaterally) were used for the study. For each slide, the pathologist randomly chose 10 sites under low power, and then the selected morphological alteration was counted under high power. This was done for each segment of lung, and the results were totaled and averaged for the grid area (13,000 or 52,000 µm²).

Ad distribution. OCT-embedded frozen sections were cut to a thickness of 6 µm and placed in a bath of 4% formaldehyde for 5 min, and the slides were then washed twice in Hanks balanced salt solution (HBSS). X-Gal staining solution (20 mg of 5-bromo-4-chloro-3-indolyl-b-D-galactopyranoside per ml of N,N-dimethyl formamide) was added (20 µl/ml) to a fresh solution of 5 mM potassium ferrocyanide–5 mM potassium ferricyanide–2 mM magnesium chloride in HBSS for the final X-Gal working solution. The slides were then exposed to the final X-Gal working solution for 10 h at 37°C. The slides were washed in HBSS counterstained with nuclear fast red and placed under a coverslip. Sections were then examined for cells that stained blue, which is indicative of LacZ expression (23).

Statistics. For assessment of clinical and fluorescent real-time PCR data, significance was determined to be a P value of <0.05. Nonparametric Wilcoxon tests were used to assess samples of unequal sizes for statistical differences. t tests that assumed unequal variances were used to assess the means for groups of equal sizes for differences.

RESULTS

The daily body temperatures showed early and late increases through the course of infection. On day 2, the body temperatures of all animals in the PIV3-infected groups were generally increased, with the animals in the PIV3/Ad/HBD6 group having a significantly higher temperature than those in the control group (P < 0.01) (Fig. 1). Starting on day 5 the animals in the PIV3/Ad/HBD6 group had significantly increased body temperatures compared with those of the animals in the control group, and these continued throughout the duration of the experiment (P < 0.05). The animals in the PIV3 and PIV3/Ad groups had moderately increased body temperatures that were not statistically significant different except for those for the animals in the PIV3 group on day 6 (P < 0.01).

The respiration rate on day 2 was significantly higher in the animals in the PIV3/Ad group (P < 0.05) and moderately elevated in the animals in the PIV3/Ad/HBD6 group (P = 0.08) compared to that in the animals in the PIV3 group (Fig. 2). By day 4 the respiration rates in the animals in the PIV3-infected groups were generally elevated (P < 0.05) compared with those in the animals in the control group, except for an increase that lacked statistical significance for the PIV3/Ad/HBD6 group on day 6. Interestingly, on days 6 and 7 the animals in the PIV3/Ad/HBD6 group had a slightly higher respiration rate than those in the PIV3 or PIV3/Ad group.

Daily milk consumption (in liters per kilogram of body weight) was evaluated to determine if there were any clinical differences in appetite. For the duration of the experiment, the animals in the PIV3, PIV3/Ad, and PIV3/Ad/HBD6 groups had decreased (P < 0.05) average milk consumption compared with the controls on day 6 (P < 0.05).
to that for the animals in the control group, with no significant difference among the PIV3-treated groups (Fig. 3).

The growth of the lambs (average daily gain, in kilograms per day) during the experiment corresponded to the milk consumption, with significantly decreased growth of the animals in the PIV3 ($P < 0.05$) and PIV3/Ad/HBD6 ($P < 0.05$) groups (Fig. 4). The animals in the PIV3/Ad group showed a similar decrease in growth, but the decrease was not statistically significant.

Grossly, the control lambs had no visible lesions, while the lambs in the PIV3, PIV3/Ad, and PIV3/Ad/HBD6 groups had a cranioventral to hilar distribution of multifocal plumb-red consolidation. There was a slightly wider lesion distribution in the lambs in the PIV3/Ad and PIV3/Ad/HBD6 groups compared with that in the lambs in the PIV3 group. Microscopically, the PIV3-induced lesions consisted of a multifocal thickening of the alveolar septa by macrophages and a small number of lymphocytes, neutrophils, and plasma cells. The bronchiolar lumens contained small aggregates of neutrophils and sloughed cellular debris. Within these areas, the bronchiolar epithelium was thickened due to hyperplasia, with folds of piled cells having pale cytoplasms, increased mitotic figures, and infrequent syncytial cells. The animals in the PIV3/Ad/HBD6 group appeared to have relatively higher numbers of neutrophils within the lungs compared to the numbers in the lungs of the animals in the group inoculated with PIV3 alone.

To evaluate neutrophil recruitment to the lungs following treatment, pulmonary neutrophil density was determined by morphometry. The lungs of each of the PIV3-infected groups had significantly increased ($P < 0.05$) neutrophilic infiltrates compared with the lungs of the controls (Fig. 5). In addition, the PIV3/Ad/HBD6 group had a higher neutrophil count than either the PIV3 or the PIV3/Ad group ($P < 0.05$).

PIV3 immunohistochemistry produced no staining in the control lungs, while cellular staining was present in the lungs of all lambs in the PIV3-infected groups. This was characterized by small granular red staining within the cytoplasm (Fig. 6). Affected cells most often were epithelial cells of the distal bronchi to terminal bronchioles. Morphometric quantification of the cellular PIV3 immunostaining showed increased numbers of immunoreactive cells in the PIV3/Ad ($P < 0.05$) and PIV3/Ad/HBD6 ($P < 0.05$) groups compared to the number in the PIV3 group (Fig. 7).

The mitotic rate was significantly increased in all of the PIV3-infected groups compared with that in the control group (Fig. 8). No significant changes in the PIV3-infected groups were evident. PIV3-induced syncytial cell formation was not present in the control group but was present in all of the PIV3-infected groups. The combined Ad gene therapy groups had higher syncytial cell densities than the PIV3-infected groups ($P < 0.05$) (Fig. 9).

The Ad vector was localized by X-Gal staining to a small number ($<5\%$) of individual epithelial cells of the bronchi and terminal bronchioles (data not shown). This staining was not seen in samples from two control lambs inoculated with sterile medium.

Real-time fluorogenic PCR was used to assess the lambs for transfection of HBD6 and for alterations in host SBD1, SP-A, and SP-D mRNA. No amplification of HBD6 was detected in the control groups (inoculated with PIV3/Ad) by the fluorogenic PCR, while all four of the HBD6-inoculated lambs had late-cycle amplification, indicating weak HBD6 expression (data not shown). Expression of SP-D in each PIV3-infected group was significantly elevated compared with that in the control group, and the levels of expression tended to increase from the PIV3/Ad group to the PIV3/Ad/HBD6 group (Fig. 10). Similar trends in SP-A and SBD1 mRNA expression were present, with statistically significant differences seen for the levels of expression in the PIV3/Ad and PIV3/Ad/HBD6 groups combined compared with that in the PIV3 group for both SP-A and SBD1 ($P < 0.05$) (Fig. 11 and 12).
DISCUSSION

The hypothesis of this study was that beta-defensin (HBD6) expression during concurrent PIV3 infection would diminish disease progression. Clinical parameters were monitored during the course of infection, and day 7 of infection was chosen for necropsy, as the lesions were usually marked and active viral clearance was seen in the lambs (7, 20). All PIV3-infected groups had clinical evidence of infection (e.g., elevated temperature and tachypnea and depressed milk consumption and growth), gross and microscopic lesions consistent with PIV3 infection, and immunohistochemical staining in PIV3-infected animals only. Ad transfection was confirmed by X-Gal staining of the Ad/HBD6/lacZ-infected airway epithelium on day 4 and by HBD6 mRNA expression on day 7 of infection.

The animals in the Ad-treated groups (the PIV3/Ad and PIV3/Ad/HBD6 groups) had increased temperatures and respiration rates on day 2 of infection. This was consistent with previous work in which BALB/c mice showed increased levels of the proinflammatory cytokines tumor necrosis factor-alpha (TNF-α) and interleukin-6 (IL-6) within even hours of intrapulmonary inoculation of human Ad 5 (40). TNF-α and IL-6 are endogenous pyrogens that account for the increased body temperatures, and TNF-α mediates bronchial and vascular smooth muscle contractility, which, along with pyrexia, could have increased the respiratory rate (2, 35). Other clinical alterations compared to the clinical conditions of the controls were generally attributable to PIV3 infection. Of particular interest was the fact that the animals in the Ad-treated groups had increased staining of PIV3 and syncytial cell formation.
compared with that in the animals in the PIV3 groups not treated with Ad. Antigen staining occurs at sites of PIV3 protein synthesis, and syncytial cell formation is mediated by the fusion protein and is regulated by the hemagglutinin-neuraminidase protein; both indicators are by-products of PIV3 replication (28).

The increased immunostaining and syncytial cell formation were both indicative of enhanced PIV3 activity (21, 28). This novel finding suggests that pulmonary Ad-mediated gene therapy can exacerbate concurrent PIV3 infection. Proteins of Ads, including proteins E1A, E1B, and E3, have been shown to have immunomodulating functions, most of which are immunosuppressing (17, 29). The vector used in this study had an E1 deletion, but the E3 gene was intact. E3 gene products inhibit peptide processing for major histocompatibility complex (MHC) I presentation; activation of nuclear factor-kappa beta (NF-κβ); and apoptosis through the fatty acid synthase, TNF-related apoptosis-inducing ligand, and TNF-α pathways (5, 8, 17, 29, 32). Processing for MHC I presentation and apoptosis of virus-infected cells are important mechanisms for viral clearance, and the synthesis of antiviral type I interferon along with other inflammatory mediators are mediated in part through NF-κβ activation. Early protein E3 inhibition of antiviral pathways could have allowed more enhanced PIV3 replication and infection. While the immunosuppressive effects of the Ad E3 gene may participate in enhancing PIV3 infection, the low level of cellular distribution of Ad suggests that this may not be a significant component. Another mechanism for enhanced PIV3 infection may be the general inflammatory and cellular changes caused by the addition of the Ad vector. In previous work, pretreatment with 4-ipomeanol similarly enhanced PIV3 activity in a calf model (21). The compound 4-ipomeanol is biotransformed in Clara cells into a free-radical metabolite which causes necrotizing bronchiolitis and interstitial pneumonia. The investigators that used the PIV3-infected calf model hypothesized that 4-impomeanol may (i) suppress pulmonary innate defenses, (ii) increase the number of susceptible cells, or (iii) increase the levels of factors (e.g., proteases) that enhance viral replication. Early immunosuppression by Ad proteins and/or altered cellular susceptibility to PIV3 infection and replication may contribute to this increased PIV3 activity in Ad-treated lambs. Further work needs to be done to clarify and define the mechanism.

We further evaluated the levels of mRNA for innate antimicrobial peptides and proteins to determine if Ad gene therapy altered mRNA expression. SP-D levels were increased in each PIV3-infected group compared with those in the control group (P < 0.05), with moderate increases in the PIV3/Ad/HBD6 group compared to that for the PIV3 group. SP-A and SBD1 mRNA expression was significantly increased compared to that in the PIV3 group. SP-D, SP-A, and SBD1 mRNA expression is partially regulated by the status of pulmonary inflammation.
and lung injury (22, 30). The increases seen here were likely a coordinated response to the exacerbated inflammation and injury associated with inoculation of the Ad vector and exacerbated PIV3 infection.

The effect of HBD6 expression in vivo on PIV3 infection was unanticipated, as it tended to be proinflammatory and aggravated the overall clinical situation. Microscopically, neutrophil infiltration was increased in the PIV3/Ad/HBD6 group, and this was confirmed by morphometry, in which the neutrophil density was increased \( P < 0.05 \) compared with those in either the PIV3 or the PIV3/Ad group. The clinical scores for body temperature and respiration rates were increased for the PIV3/Ad/HBD6 group compared with those for the control group and with slight increases compared with those for the PIV3/Ad group. While beta-defensins are increasingly known for their antimicrobial properties, they have additional functions, such as leukocyte chemotaxis and immunomodulation (30). Recently, HBD2 was shown to function in vitro as a chemoattractant for TNF-α-primed neutrophils, and the HBD2 ligand is not for TNF-α. Contrarily, HBD2 was shown to function in vitro as a chemoattractant and with slight increases compared with those for the PIV3/Ad group. Temperature and respiration rates were increased for the PIV3/Ad/HBD6 group compared with those for the control group and with slight increases compared with those for the PIV3/Ad group. While beta-defensins are increasingly known for their antimicrobial properties, they have additional functions, such as leukocyte chemotaxis and immunomodulation (30). The effect of HBD2 expression in vivo on PIV3 infection was unanticipated, as it tended to be proinflammatory and aggravated the overall clinical situation. Microscopically, neutrophil infiltration was increased in the PIV3/Ad/HBD6 group, and this was confirmed by morphometry, in which the neutrophil density was increased \( P < 0.05 \) compared with those in either the PIV3 or the PIV3/Ad group.

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