Bombesin-like Peptide Mediates Lung Injury in a Baboon Model of Bronchopulmonary Dysplasia

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Abstract

The etiology of bronchopulmonary dysplasia (BPD), a chronic lung disease of infants surviving respiratory distress syndrome, remains fundamentally enigmatic. BPD is decreasing in severity but continues to be a major problem in pediatric medicine, being especially prevalent among very premature infants. Increased numbers of pulmonary neuroendocrine cells containing bombesin-like peptide (BLP) have been reported to occur in human infants with BPD. We tested the hypothesis that BLP mediates BPD using the hyperoxic baboon model. Urine BLP levels increased soon after birth only in 100% O₂-treated 140-d animals which developed BPD, correlating closely with severity of subsequent chronic lung disease. Similar elevations in urine BLP were observed in the 125-d baboon “interrupted gestation” model of BPD. Postnatal administration of anti-BLP antibody attenuated clinical and pathological evidence of chronic lung disease in the hyperoxic baboon model. Urine BLP could be a biological predictor of infants at risk for BPD, and blocking BLP postnatally could be useful for BPD prevention. (J. Clin. Invest. 1998. 102:584–594.) Key words: radioimmunoassay • monocolonal antibody • oxygenation index • histopathology • proliferating cell nuclear antigen

Introduction

Bronchopulmonary dysplasia (BPD), a chronic inflammatory lung disease of premature infants (1), remains a major problem in pediatric medicine, paradoxically due to success in improving survival of preterm infants now that respiratory distress syndrome is routinely treated with prenatal glucocorticoids and surfactant replacement therapy (2). It is estimated that 20% of ventilated newborns develop BPD (3), representing at least 3,000 infants annually in the USA (4). In particular, from 1976 to 1990, the incidence of BPD increased from ~ 10 to 33% in very low birth weight infants (1,500 grams or less). As a result, BPD now accounts for the highest mean health care costs per child and also is second only to asthma in terms of total health care costs per year in a given population (5). However, this increase in BPD is not explained simply by the decrease in acute mortality in this population (2). The pathophysiology of BPD is believed to be multifactorial, with factors contributing to lung injury believed to be trauma from mechanical ventilation, oxygen toxicity, infection, inflammation, and immaturity (4). Improvements in medical management, especially high frequency ventilation, positive pressure ventilation using nasal prongs, surfactant replacement therapy, lower administered FIO₂s, and careful monitoring and prompt treatment of infections, have reduced the severity of BPD occurring in human infants over the past decade (2, 4, 6, 7). Although prenatal maturational therapy with dexamethasone, with or without postnatal surfactant therapy, has slightly lowered the overall incidence of BPD, it is controversial as to whether surfactant deficiency per se is a contributing factor (8, 9). High dose surfactant alone has been found to have a more prolonged beneficial effect on oxygenation, with subsequent reduction in the incidence of BPD (10). Beneficial effects of postnatal dexamethasone on BPD are likely to be due to generalized antiinflammatory effects of glucocorticoids. However, in spite of all of these advances, there is no biological marker for identifying which very low birth weight infants will go on to develop chronic lung disease (11).

Recently, BPD has been clinically defined as oxygen dependence at 28 d postnatal age or 36 wk after conception age, so we cannot draw exact parallels between the 6- to 10-d baboon model and the human disease. Nonetheless, there are close similarities in the apparent disease processes. In human hyaline membrane disease, numbers of cells containing pulmonary bombesin-like peptide (BLP) are low, due to either degranulation of NE cells and/or generalized epithelial cell necrosis. In lung sections from one of four infants with hyaline membrane disease, we previously detected marked elevation in BLP mRNA using in situ hybridization, suggesting that BLP might be implicated in the earliest stages of development of BPD (12). Pulmonary NE cell hyperplasia, which represents predominantly a cell differentiation response (13), occurs frequently in infants progressing to BPD (14, 15). This observation could be relevant to the pathophysiology of BPD because BLP are known to be potent bronchoconstrictor peptides as well as growth factors for pulmonary fibroblasts and epithelial cells. Thus, early overproduction of BLP could mediate the peribronchiolar and interstitial fibrosis and reactive Airways disease which are hallmarks of BPD. Although definitive testing of this hypothesis will require a clinical investigation, valuable insight can be gained from relevant animal models.
Although many species of rodents, nonhuman primates, and other laboratory animals develop acute hyaline membrane disease (HMD) (16–22) and even chronic interstitial lung disease similar to BPD (23–25), preterm baboons comprise the best nonhuman animal model for both HMD and its chronic sequelae, which appears clinically and pathologically most similar to human BPD (26–31). The increasing incidence of BPD as one of the major unsolved problems in pediatric medicine resulted in the initiation of a collaborative program by the National Institutes of Health whereby the baboon model of BPD established by Coalson et al. in San Antonio has been targeted as a key tool in investigating this illness. Preterm animals delivered by Caesarean section at 140 d gestation (term = 180 d) and maintained on 100% O₂ (140-d/100% O₂) for 10 d (without surfactant replacement therapy) develop clinical and pathological features typical of moderate to severe BPD (“old BPD”) similar to those described by Northway in 1967 (1). In brief, these changes include: severe distortion of lung architecture with segments of atelectasis and mucous plugging alternating with gross overexpansion of adjacent lung segments (compensatory emphysema), generalized decreased alveolarization, interstitial and peribronchiolar fibrosis, epithelial hyperplasia and squamous metaplasia, and smooth muscle hypertrophy with associated chronic reactive airways disease. In contrast, control 140-d gestation preterm baboons maintained for 10 d on oxygen PRN (to keep arterial blood hemoglobin above ~90% O₂ saturation [140-d/PRN]) do develop acute respiratory distress syndrome (RDS) with hyaline membrane disease between 1 and 48 h after delivery, but recover from the acute injury and do not develop subsequent clinical or pathological changes reminiscent of BPD. More recently, Coalson et al. have developed a model of milder BPD occurring in extremely premature baboons (125 d gestation) receiving O₂ PRN and requiring exogenous surfactant to survive, which is much more similar to the version of BPD currently seen in human infants (32). In this model of “new BPD,” >90% of 125-d/PRN animals develop characteristic clinical and pathological features of chronic lung disease found in ventilated extremely low birth weight human infants: mild interstitial and peribronchiolar fibrosis, chronic reactive airways disease, and especially arrested alveolar septation (31).

Because infants with BPD have increased pulmonary NE cells containing BLP, a growth factor and bronchoconstrictor (15), we hypothesized that BLP mediates lung injury in BPD. The kinetics of BLP mRNA and peptide levels in normal baboon lung are similar to those in humans (Sunday, M.E., and R.L. Emanuel, manuscript in preparation). The present study demonstrates that urine BLP double between 24 and 72 h after birth in BPD animals, but not in controls. The severity of chronic lung disease correlates directly with this early increment in urine BLP. Blocking anti-BLP antibody administered to five BPD baboons soon after birth protects against histopathological changes of BPD. Although two of these baboons became septic, the others were clinically near normal. These observations suggest that elevated urine BLP might predict infants at highest risk for developing BPD, and blocking BLP after birth could be a novel treatment for preventing BPD.

Methods

Animals. The National Research Council Guide for the Care and Use of Laboratory Animals was strictly adhered to throughout all phases of this study. The Animal Care Committee of the Southwest Foundation for Biomedical Research reviewed and approved the protocols used in this study. Details of the neonatal intensive care unit (NICU) animal care have been described previously (29). Values from clinical monitoring at 1–2 h intervals were used for the calculation of mean oxygenation indices (\(\Delta_{E}SE\) [OI = (F\(_{O2}\) × mean airway pressure) × 100/P\(_{O2}\)] for three groups of experimental animals between birth and 10 d of age: 140-d/PRN (O\(_2\) PRN, non-BPD); 140-d/100% O\(_2\) without antibody treatment (100% O\(_2\)/Neg, severe BPD); and 140-d/100% O\(_2\) receiving intravenous anti-BLP blocking antibody treatment (100% O\(_2\)/2A11). 2A11 murine monoclonal antibody (1 mg/ml, sterile, pyrogen-free, and azide-free) was administered intravenously as an infusion starting 2 h after birth (5 mg/kg, given over 2 h), with two boosts given similarly on days 3 and 6 (4 mg/kg, starting at ~72 and 144 hours).

Urine BLP RIA. Urine was collected from newborn baboons as 24-h pooled specimens (5–50 ml total volume) which was stored at -70°C, shipped frozen on dry ice, and thawed only once at the time of assay, using an aliquot (~1 ml) for RIA. BLP levels were quantitated by RIA (Inc Star Corp., Stillwater, MN) using the clear supernatant of urine samples, containing 25 mM acetic acid and stored at -70°C. The assay uses a rabbit antibody to amphibian bombesin, has a detection limit of 100 pg/ml, and detects bombesin and its mammalian homologue gastrin-releasing peptide (GRP) which is the major known pulmonary BLP (12), but not related neuropeptides neurenom B, substance P, nor neurotensin. Each sample was measured in at least two separate radioimmunoassays. Urine BLP values were normalized to their respective creatinine content (mg/ml) which was measured using a kit (Sigma Chemical Co., St. Louis, MO) to correct for variations in urinary concentration or dilution. Each value shown on the plots represents a 24-h urine collection ending at the indicated time points; the normalized value for the first day urine collection is defined as 100%, with subsequent values expressed as the mean percent change compared to the 24-h urine BLP level.

Histopathological and immunohistochemical analyses. Tissues were fixed in 4% paraformaldehyde for 18 to 24 h before routine processing into paraffin, as described previously (33). Histochemical stains included hematoxylin and cosin for general tissue architecture, Masson’s trichrome for fibrosis, and elastin stain for demonstrating pulmonary alveolar architecture (34).

Routine immunostaining was carried out on 3-μm sections as described previously using PCNA monoclonal antibody clone PC10 (1:20 dilution) (Dako Corp., Carpinteria, CA) (33), our own rabbit antibombesin antisera (1:500 dilution) (35), or rabbit polyclonal anti-NEP 9.5 (1:500 dilution) (Ultrade Inc., Isle of Wight, UK) with the avidin–biotin complex (ABC) technique (Vector Laboratories, Burlingame, CA) (33). The peroxidase substrate was diaminobenzidine and methyl green was used as the counterstain. Negative controls run in parallel consisted of the following: for PCNA, the isotype-matched MOPC-21 (IgG1) (Sigma Chemical Co.) was used; and for antibombesin antisera, bombesin antisera preabsorbed overnight with bombesin antigen (10 μg/ml) (36).

Localization of 2A11 mouse monoclonal antibody in baboon tissues was carried out using a 1:10 dilution of biotinylated horse anti-mouse IgG (Vector Laboratories) as the primary antibody or biotinylated normal horse IgG as the negative control incubated overnight at 4°C. Subsequent incubations with methanol/H\(_2\)O, the ABC reagent, and biotinylated tyramide (Tyramide-System Amplification [TSA] from DuPont-New England Nuclear, Boston, MA) were the same as for the standard immunoperoxidase method.

Statistical analyses. For group comparisons, statistical analyses used the Student’s t test (unpaired), one-way ANOVA, or two-way ANOVA and results are given as mean values ± SE.

Results

Changes in urine BLP levels predict BPD. Results of urine bombesin-like peptide (BLP) radioimmunoassay, normalized
A

\[ \Delta \text{uBLP (from 24h)} \]

\[ \text{Postnatal Time (Hours)} \]

B

\[ \text{Urine BLP (pg/mg Cr)} \]

\[ \text{Postnatal Time (Hours)} \]

C

\[ \Delta \text{uBLP (from 24h)} \]

\[ \text{Postnatal Time (Hours)} \]

D

\[ \text{Urine BLP (pg/mg Cr)} \]

\[ \text{Postnatal Time (Hours)} \]

E

\[ \text{SF05} \]

\[ \text{S096} \]

\[ \text{Postnatal Time (Hours)} \]

F

\[ y = 22.099 + 19.858x \]

\[ R^2 = 0.866 \]

\[ P = 0.022 \]

\[ \text{Oxygenation Index at 240 Hours} \]

G

\[ \Delta \text{uBLP (from 24h)} \]

\[ \text{Postnatal Time (Hours)} \]

Sunday et al.
for creatinine to correct for variations in urinary concentration or dilution, are given in Fig. 1. Control animals (140-d/PRN) had no significant change in mean urine BLP levels between 24 and 72 h after delivery (n = 9). Mild elevation in urine BLP levels occurred after ≥ 96 h, consistent with increased numbers of pulmonary NE cells after prolonged mechanical ventilation (14). Individual values from all nine animals are shown in Fig. 1 B, with all animals manifesting either a decline or minimal increase between 24 and 72 h after birth.

In contrast, the group of 140-d/100% O₂ baboons (n = 8) developing chronic lung disease similar to the original severe form of BPD had a significant doubling in mean urine BLP levels between 24 and 48 h, termed ΔurBLP (24–48) (Fig. 1 C). All of these animals manifested moderate to severe oxygenation impairment (oxygenation index [OI] > 2.5) and chest x-ray changes at 6 to 10 d of age, where OI is defined as (F/O₂ × mean airway pressure) × 100/P/O₂. Although peaking at 48 h, urine BLP levels were still significantly elevated above the 24-h values at 72 and 96 h. Individual results from all eight of these animals are given in Fig. 1 D. A small percentage (~ 15–20%) of 140-d animals treated with 100% O₂ do not develop significant clinical evidence of chronic lung disease during their course. Urine BLP levels from two such animals are shown in Fig. 1 E: both of these baboons had a decline rather than an increase in urine BLP levels (with 48 h levels representing 66 and 58% of the corresponding 24-h values).

In the five 140-d animals treated with 100% O₂ for which serial urine collections were available, there was a direct correlation (r = 0.93, P = 0.022) between the increment in urine BLP between 24 and 48 h and the OI at 10 d (Fig. 1 F). There was no correlation between early changes in urine BLP (or absolute values of urine BLP at 24, 48, 72, or 96 h) and OI between 2 and 216 h of age (data not shown). In the 140-d/100% O₂ model, we did not observe any correlations between absolute urine BLP levels and OI, apparently due to variability in baseline urine BLP levels.

The above observations are based on the hypoxic baboon model, and we cannot be absolutely certain that increased urinary BLP does not reflect increased BLP gene expression in response to hyperoxia rather than as a result of BPD per se. To provide assessment of changes in urine BLP and BPD independently of therapy with 100% O₂, we also measured urine BLP levels in the second baboon model of “new BPD” (125-d gestation baboons treated with O₂ PRN and exogenous surfactant therapy) (30). In this group (n = 12), there was a significant (~ 30%) increase in urine BLP between 24 and 48 h after birth (Fig. 1 G), with doubling occurring at 72 h; levels remained elevated at 96 h. Most of these 125-d animals were maintained for only 6 d (n = 11), so that we could not analyze correlations with 10-d OI values. These observations suggest that elevated urine BLP levels might be associated with chronic lung disease occurring in diverse clinical settings.

In addition to its presence in lung, GRP mRNA can be detected in baboon brain, thyroid, stomach, and pancreas at 140 d gestation and after 100% O₂ treatment for 10 d (two animals screened, data not shown). It was also detected in thymus and/or adrenal in the two animals given 100% O₂ treatment for 10 d. Using reverse transcriptase-PCR (RT-PCR), we did not detect GRP mRNA in kidney or bladder. In previous studies of human fetal tissues, the major source of GRP mRNA and BLP immunoreactivity has been demonstrated to be lung (12). We have obtained total RNA samples from lung of one 140-d animal given O₂ PRN for 24 h and another animal given 100% O₂ treatment for 24 h (thanks to Dr. Carl White, National Jewish Medical and Research Center, Denver, CO). The animal treated with 100% O₂ had about a twofold increase in lung levels of GRP mRNA as compared to the PRN animal (data not shown). When we calculate the amount of BLP excreted in a 24-h urine collection (~ 5 ng), it is less than half of the BLP content of lung tissue harvested from animals following 100% O₂ treatment for 10 d (~ 14 ng). Thus, although we cannot be certain that the lung is the only source of urine BLP, the cumulative data are consistent with lung as the major contributor to urine BLP levels.

Postnatal treatment with anti-BLP–blocking antibody. To test whether BLP might play a causative role in the pathophysiology of BPD, we administered the well-characterized murine monoclonal anti-BLP–blocking antibody 2A11 to five 140-d/...
100% O₂ baboons beginning 2 h after birth. The dosage and frequency of administration were based on results from previous preclinical pharmacokinetics studies in mice and dogs (37): we chose a dose comparable to that required for > 95% blocking of BLP-triggered gastrin release in dogs. We sought objective evidence that 2A11 was indeed functioning as a BLP blocking antibody in the preterm baboons. First, the five 140-d/100% O₂ animals receiving 2A11 demonstrated only slightly

Figure 2. 140-d baboons treated with 100% O₂ and anti-BLP antibody (2A11). (A) The change in urine BLP from 24 to 48 h in non-BPD 140-d/PRN baboons (PRN, n = 9), 140-d/100% O₂ baboons with severe BPD that did not receive antibody treatment (100% O₂/Neg, n = 8), and the five 140-d/100% O₂ baboons that received 2A11 shortly after birth (100% O₂/2A11). Only the non-2A11–treated animals exposed to 100% O₂ had a significant increase in urine BLP at 48 h. Results are given as mean values ± SE (*P < 0.001 compared to 140-d/PRN controls using the Student’s t test, unpaired). (B–E) Lung sections from representative 2A11-treated animals demonstrate BLP-positive cells. (B) BLP immunostaining of a section of lung tissue from a 2A11-treated animal (4N96) demonstrates a small cluster of pulmonary NE cells in a bronchiole. (C) A thin section immediately serial to (B) run in parallel using antibombesin antiserum preabsorbed with bombesin antigen (10 μg/ml) demonstrates loss of immunostaining of the same NE cells. (B–E: magnification of 100, methyl green counterstain). (D and E) There are frequent BLP-positive cells present in the small bronchioles (D) and alveolar ducts (E, arrows); these cells also contain the NE-specific marker PGP 9.5 (data not shown). All immunostaining was carried out as detailed in Methods using 3-μm thick lung sections and our own rabbit antibombesin antiserum (1:500) with the ABC-complex immunoperoxidase technique. The substrate was diaminobenzidine (33). (Magnification of 80, methyl green counterstain). (F) Horse anti–mouse IgG immunostaining of lung tissue sections from a representative 2A11-treated animal demonstrates staining for 2A11 (murine IgG) in the undifferentiated mesenchymal tissue around developing blood vessels and airways (between arrows), but not in differentiated smooth muscle around blood vessels (v) or bronchioles (b). (G) A section immediately serial to (F) incubated in parallel with normal horse serum lacks this mesenchymal immunostaining.
Figure 3. Kinetics of oxygenation in 140-d preterm baboons treated with O2 PRN versus 100% O2±anti-BLP antibody 2A11. (A) Mean oxygenation indices±SE $[\text{OI} = (\text{FiO}_2 \times \text{mean airway pressure}) \times 100/\text{P}_{\text{O}_2}]$ are given at hourly intervals for three groups of experimental animals between birth and 12 h of age: 140-d/PRN (O2 PRN, non-BPD, $n = 13, 46\%$ male); 140-d/100% O2 without antibody treatment (100% O2/Neg, severe BPD, $n = 13, 38\%$ male); and 140-d/100% O2 receiving intravenous anti-BLP blocking antibody treatment (100% O2/2A11, $n = 5, 20\%$ male). Antibody (1 mg/ml) was administered intravenously as an infusion starting 2 h after birth (5 mg/kg, given over 2 h). † $P < 0.002$, and * $P < 0.05$ using the Student’s $t$ test. (B) Mean oxygenation indices±SE are given at 12- to 24-h intervals for the same three groups of experimental animals between birth and 10 d of age as given in A: 140-d/PRN; 140-d/100% O2/Neg; and 140-d/100% O2 receiving anti-BLP blocking antibody treatment. Antibody (1 mg/ml) was administered as an infusion 2 h after birth (details given in A), with two boosts given on days 3 and 6 (4 mg/kg, infusions starting at $\sim 72$ and 144 h). † $P < 0.001$, and * $P < 0.04$ using the Student’s $t$ test. (C) Kinetics of individual OI values for each of the five 140-d/100% O2 animals treated with antibody 2A11. (D) Mean OI values between birth and 6 d of age are given only for females from the same three groups of experimental animals as shown in A: 140-d/PRN, $n = 13$; 140-d/100% O2/Neg, $n = 13$; and 140-d/100% O2/2A11, $n = 5$. Comparison of 140-d/100% O2/Neg and 140-d/100% O2/2A11 groups by one-way ANOVA yields * $P < 0.001$. Neg, $n = 8$; and 140-d/100% O2/2A11, $n = 4$. † $P < 0.001$, and * $P < 0.02$ using the Student’s $t$ test. (E) Mean OI values between 5 and 10 d of age are given only for animals without clinical evidence of sepsis: 140-d/PRN, $n = 7$; 140-d/100% O2/
Clinical parameters relevant to the medical status of preterm infants are given.  *The number of animals in each treatment group is given in parentheses.  ‡Whether or not animals were given 2A11 intravenously beginning 2 h after birth is indicated.  §Birthweight is given in grams

Table I. Clinical Profiles of 140-d Baboons Treated for 10 d with PRN O2 versus 100% O2 with or without 2A11 Treatment

<table>
<thead>
<tr>
<th>Treatment (n)*</th>
<th>2A11†</th>
<th>Birthweight (grams)</th>
<th>% Male§</th>
<th>% HFOV</th>
<th>% Air leak**</th>
<th>OI\textsubscript{i}i</th>
<th>urBLP (24–48 h)ii</th>
</tr>
</thead>
<tbody>
<tr>
<td>O2, PRN (13)</td>
<td>No</td>
<td>534±18</td>
<td>38 (5)</td>
<td>54</td>
<td>31</td>
<td>3.0±0.6</td>
<td>105±17 (9)</td>
</tr>
<tr>
<td>100% O2 (13)</td>
<td>No</td>
<td>527±14</td>
<td>46 (6)</td>
<td>46</td>
<td>31</td>
<td>6.9±1.6</td>
<td>199±29(i) (9)</td>
</tr>
<tr>
<td>100% O2 (5)</td>
<td>Yes</td>
<td>498±6</td>
<td>20 (1)</td>
<td>0</td>
<td>0</td>
<td>2.5±0.4(i)</td>
<td>132±13 (5)</td>
</tr>
</tbody>
</table>

Clinical indices (100% O2 (5) Yes 498 ± 6

Clinical profiles of 140-d baboons treated for 10 d with PRN O2 versus 100% O2 with or without 2A11 treatment.

Figure 4. Histopathological analyses and PCNA immunostaining in lung tissues from 140-d baboons treated with O2, PRN or 100% O2±2A11.  (A–D) Lung lobes were inflated with 4% paraformoldehdy (A–C: elastic tissue stain, magnification of 8; D: hematoxylin and eosin, magnification of 80). Representative sections of lung tissue are shown from one animal for each group: (A) A typical non-BPD 140-d/PRN control animal maintained for 10 d. There is normal alveolarization and no evidence of fibrosis.  (B) A 140-d/100% O2 baboon with severe chronic lung disease after 10 d of treatment. There is extensive interstitial and peribronchial fibrosis with adjacent areas of compensatory emphysema and airspace dilatation.  (C and D) A 140-d/100% O2 baboon receiving three doses of 2A11 with 10 d of 100% O2. There is minimal evidence of interstitial fibrosis (best appreciated with elastin stain in C), most of which occurs in areas with intraalveolar hemorrhage and/or resolving hyaline membrane disease (C, arrows). Although there is some evidence of airspace dilatation, the magnitude of these changes is much less than that occurring in most animals not receiving 2A11.  Pockets of alveolar hemorrhage and sparse neutrophil infiltrates occur frequently in the vicinity of residual hyaline membranes (long arrows in D indicate intraalveolar hyaline membranes; short arrow indicates hyaline membrane phagocy-tosed by an alveolar macrophage). Although there is some perivascular fibrosis (V, arrow in D), there is minimal to no evidence of peribronchial fibrosis (aw, airway lumen) (D: hematoxylin and eosin).  (E–H) Immunostaining of noninflated lung tissue slices fixed for 18–24 h with 4% paraformaldehyde.  (E–G) PCNA immunostaining, carried out as described previously (33).  (H) MOPC IgG1 negative control: all magnification of 80. Representative sections of lung tissue are shown from one animal for each group, with representative positive cells indicated by arrows (short arrows, epithelial cells; arrowheads, mesenchymal cells).  (E) A typical non-BPD 140-d/PRN control animal maintained for 10 d. There is PCNA immunostaining localized predominantly to epithelial cells in airways and airspaces.  (F) A 140-d/100% O2 baboon with severe chronic lung disease after 10 d of treatment demonstrates extensive PCNA labeling of mesenchymal cells in the interstitium and peribronchial regions, coinciding with areas of fibrosis, and also intense PCNA labeling of epithelial cells in airways and airspaces.  (G and H) A 140-d/100% O2 baboon receiving three doses of 2A11 with 10 d of 100% O2.  PCNA immunostaining is predominantly localized to the epithelial cells in airways and alveoli. There is much reduced PCNA immunostaining compared to either the 140-d/PRN animals (as in E) or the 140-d/100% O2 animals (as in F).
Immunohistochemical analyses for proliferating cell nuclear antigen (PCNA) as a marker for proliferating cells revealed moderate labeling of predominantly the epithelial compartment in 140-d/PRN animals (Fig. 4E) and intense labeling of both epithelium and distal lung mesenchyme in 140-d/100% O2 baboons, consistent with ongoing interstitial fibrosis (Fig. 4F). In contrast, there was marked reduction in PCNA labeling of lung parenchyma in 2A11-treated animals (Fig. 4G), in which most of the residual proliferating cells were epithelial in location.

Discussion

The present study demonstrates that elevated urine BLP between 24 and 72 h after birth is a reliable predictor of which preterm baboons will develop chronic lung disease similar to BPD in two different baboon models of BPD. This effect is apparent even when gender is eliminated as a variable, and all experimental and control animals were treated during the same time frame under the direction of a single neonatologist to control for variations in intensive care unit practices. The precise mechanism for increased urinary BLP at the two different gestational ages with differing O2 exposures is not known. Similarly, the precise pathophysiology of BPD is unknown: the role of immature antioxidant defenses in the 125-d PRN animals could undoubtedly play a role in the pathobiology of BPD in these extremely premature animals (32). Currently, there is no reliable epidemiological or biological marker for identifying which preterm human infants will go on to develop chronic lung disease (11). Furthermore, we demonstrate that postnatal administration of blocking anti-BLP antibody is protective against the development of clinical and pathological evidence of BPD: none of the five animals receiving anti-BLP blocking antibody manifested significant peribronchiolar fibrosis or widespread alveolar interstitial fibrosis. At least part of this effect is probably due to blockade of BLP function as a fibroblast growth factor, as evidenced by marked diminution in PCNA immunostaining in mesenchymal cells. Multiple molecular mechanisms underlying the anti-BLP antibody effect are likely, however, including decreased macrophage activation and blockade of direct effects of BLP as a bronchoconstrictor. It is unlikely that the late beneficial effect on the course of BPD is simply due to the early improvement in OI of anti-BLP antibody (2A11)-treated animals as compared to both the untreated 100% O2 and O2 PRN animals: the male 2A11-treated animal (4N96) did resolve severe RDS, but had minimal evidence of chronic lung disease after 10 d of 100% O2.

Only a few potential adverse side effects of 2A11 are apparent. Histological analyses demonstrate scattered intraalveolar hemorrhage and sparse neutrophilic infiltrates, consistent with resolving hyaline membrane disease (Fig. 4, C and D, arrows). Pathological evidence of intrapulmonary hemorrhage was present in a majority of both the 140-d/100% O2 and the 140-d/PRN animals (all of which were ventilated and none of which received exogenous surfactant), and generally correlated with severity of acute RDS. Only the two animals which developed septicemia had more widespread, moderate to severe neutrophilic infiltrates, diagnostic of clinically significant acute pneumonia. Clinical and pathological evidence of septicemia typically occurs in >5% of either 140-d/100% O2 or 140-d/PRN animals (all of which had indwelling arterial lines).
the two baboons became septic by chance alone, especially considering the presence of indwelling arterial lines and endotracheal tubes in these preterm animals. Of additional note, cultures of the antibody stocks remaining were negative for bacterial and fungal organisms. Nonetheless, this is an unexpected, interesting scientific observation that may shed light on the role of bombesin-like peptides during acute and chronic lung injury processes.

One further point of interest concerns the significant improvement in OI in all five 2A11-treated animals during the first 36 h of postnatal life. In our previous work, we demonstrated that BLP is important for type II cell differentiation and surfactant synthesis as well as generalized cell proliferation in both murine and human fetal lung in organ culture and in murine lung in utero (40–43). Subsequent work in other laboratories confirmed and extended this result to isolated rat primary type II cell cultures (44, 45), in which BLP stimulated surfactant secretion as well as synthesis (44). The highest endogenous BLP levels occur at mid-gestation in primate lung, but often peak effects of BLP occur at subnanomolar concentrations (44, 45). We postulate that 2A11 might promote more efficient secretion of endogenous surfactant from type II cells by reducing local BLP concentrations to a more optimal concentration. High doses of BLP often downregulate biological effects, regardless of whether the cellular response is proliferative, synthetic, or secretory. Finally, we cannot exclude the possibility that the 2A11 effect might be mediated via IgG1 Fc receptor function, with nonspecific modulation of macrophage function leading to altered cytokine production. Ideally, a double-blinded, randomized baboon trial should be carried out using 2A11 versus MOPC, an IgG1 of undefined specificity, to test for care of children with chronic illnesses enrolled in the Washington state medicaid program, fiscal year 1993. Pediatrics. 100:197–204.


23. Han, R.N., S. Buch, I. Tseu, J. Young, N.A. Christie, H. Frindova, S.J.


