Abstract. Prematurely delivered lambs were treated with radiolabeled natural surfactant by either tracheal instillation at birth and before the onset of mechanical ventilation, or after 23±1 (±SE) min of mechanical ventilation. Right ventricular blood flow distributions, left ventricular outputs, and left-to-right ductal shunts were measured with radiolabeled microspheres. After sacrifice, the lungs of lambs receiving surfactant at birth inflated uniformly with constant distending pressure while the lungs of lambs treated after a period of ventilation had aerated, partially aerated, and atelectatic areas. All lungs were divided into pieces which were weighed and catalogued as to location. The amount of radiolabeled surfactant and microsphere-associated radioactivity in each piece of lung was quantified. Surfactant was relatively homogeneously distributed to pieces of lung from lambs that were treated with surfactant at birth; 48% of lung pieces received amounts of surfactant within ±25% of the mean value. Surfactant was preferentially recovered from the aerated pieces of lungs of lambs treated after a period of mechanical ventilation, and the distribution of surfactant to these lungs was very nonhomogeneous. Right ventricular blood flow distributions to the lungs were quite homogeneous in both groups of lambs. However, in 8 of 12 lambs, pulmonary blood flow was preferentially directed away from those pieces of lung that received relatively large amounts of surfactant and toward pieces of lung that received less surfactant. This acute redirection of pulmonary blood flow distribution may result from the local changes in compliances within the lung following surfactant instillation.

Introduction

The intratracheal administration of a variety of surfactant preparations to prematurely delivered animals and infants with respiratory distress syndrome (RDS) has improved pulmonary function. In most studies, surfactant treatments were administered by injecting suspensions of surfactant into the fluid-filled airways of animals at birth and before the initiation of breathing (1-3). A surfactant suspension also can be administered to previously ventilated animals by tracheal instillation (4, 5). While the tracheal instillation of surfactant after a period of ventilation may lead to a clinical response of shorter duration than a treatment given at birth (4), such an approach has been used for most of the clinical trials (6-8). After surfactant instillation, both premature animals and infants with RDS have prompt improvements in PaO₂ values, compliance, and a temporary clearing of the chest x-ray (1-8). Surfactant when administered to both prematurely delivered and adult animals can be found by histologic techniques in the alveoli (9, 10). However, if surfactant suspensions distribute to the lung as do particles administered by either aerosol or by tracheal instillation, a nonhomogeneous distribution should occur (11). Such a distribution of a substance that will acutely change the degree of alveolar aeration and lung compliance may secondarily affect the distribution of pulmonary blood flow. Thus, we measured the distribution of both natural surfactant and pulmonary blood flow following surfactant instillations given to premature lambs either at birth and before breathing, or after the initiation of ventilation.

Methods

Delivery and instrumentation of lambs. After premedication of date-mated pregnant ewes with ketamine and atropine and spinal anesthesia,
16 lambs at 120-d gestational age (term = 150 d) were delivered by hysterotomy as previously described (4, 5). In brief, the fetal neck was exposed through a small uterine incision and an endotracheal tube was secured by tracheotomy. Approximately 10 ml of fetal lung fluid was removed through the endotracheal tube. Following sampling of cord venous blood, the lambs were delivered, dried, and ventilated with 100% oxygen by hand for ~30 s with an anesthesia bag by using a peak inspiratory pressure of 30 cm H2O and a rate of 40 breaths/min. Then, the lambs were ventilated with Sechrist IV-100 pressure-limited infant ventilators (Sechrist Industries, Inc., Anaheim, CA) with 100% humidified oxygen. Ventilator settings were an initial peak inspiratory pressure of 28 cm H2O, a positive end expiratory pressure of 2 cm H2O, a rate of 30 breaths/min, and an inspiratory time of 1 s. Subsequently, only peak inspiratory pressures were changed in an attempt to normalize Pco2 values. A 5-Fr catheter was placed in the distal aorta via an umbilical artery for continuous recording of blood pressure, heart rate, and sampling of blood for blood-gas measurements. Soon after birth and following infiltration of the superficial tissues of the neck with local anesthesia, a 3.5-Fr infant-feeding tube was passed into the right ventricle via the right external jugular vein. A second 3.5-Fr catheter was passed into the left ventricle via the right carotid artery. Catheter positions were confirmed by the contour of the pressure tracings. All lambs were paralyzed with 0.1 mg/kg pancuronium bromide (Pavulon, Organon Teknika Corp., Aurora, CO). Body temperature was maintained with radiant warmers and supplemental heat lamps. The lambs received a continuous infusion of 100 ml/kg per 24 h of 10% dextrose in water via the arterial catheter. Expiratory tidal volumes and lung compliance values were measured and reported as before (4). The lambs were sacrificed by a cisternal injection of lidocaine and exsanguination.

Surfactant treatments. Each lamb was treated with a surfactant suspension in 0.45% saline that contained 100 mg natural surfactant lipid, which was a mixture of natural sheep surfactant isolated following alveolar lavage of adult sheep lungs and a trace amount of [14C]palmitic acid-labeled natural sheep surfactant. The labeled surfactant was isolated from lavage fluid from young lambs ~4 h after an intravascular injection of 15 mCi [14C]palmitic acid (New England Nuclear, Boston, MA) (11). Greater than 65% of the lipid soluble radioactivity in this naturally labeled surfactant was associated with phosphatidylcholine. The labeled natural surfactant contained 0.6 μCi [3H]-labeled phosphatidylcholine. The surfactant suspension also contained 0.3 μCi [3H]dipalmitoylphosphatidylcholine (100 Ci/mole) (New England Nuclear), which had been sonicated into suspension before addition to the natural surfactant (13). The composition, surface properties, and effects on lung function after the instillation of this natural surfactant have been reported (4).

For the six lambs treated at birth, a 3.9 ml surfactant suspension was mixed with the fetal lung fluid by syringe while the lamb was still inside the uterus and before the initiation of ventilation. No attempt was made to uniformly orient these lambs within the uterus. The lambs were then delivered and ventilated with a hand bag while being rotated in an effort to optimize the distribution of the surfactant. The other 10 lambs were ventilated sternum down for 23±1 min. Then, they were disconnected from the ventilators, and the same surfactant suspension which was diluted to 15 ml was instilled over 15 s as the lambs were rotated (4, 5). The lambs were reconnected to the ventilators and were rotated for a further 30 s to optimize distribution of the surfactant suspension within the airways. The lambs were returned to a sternum-down, midline position for the duration of the experiment. This treatment procedure is similar to that used for the treatment of infants with RDS (6–8).

Radiolabeled microsphere injections. The distribution of blood flow to the lungs, the left-to-right shunt through the ductus arteriosus, and cardiac outputs were measured by using radiolabeled microspheres (14). 15±2 μm (mean±SD) diameter microspheres (New England Nuclear) labeled with 51Cr, 55Sc, or 113Sn were injected sequentially into the lambs treated with surfactant at birth. The first microsphere injection was into the right ventricle of four of six lambs at 13±1 min of age, the second injection was into the left ventricle of five of six lambs at 22±1 min of age, and the third injection was into the right ventricle of all six lambs at 23±2 min of age. For six of the lambs treated with surfactant at 23 min of age, microspheres were injected into the right ventricle at 21±1 and 44±3 min of age, and into the left ventricle in four of six lambs at 42±3 min of age. The other four lambs did not receive microspheres because the lungs were used for dry-to-wet weight ratio measurements. Starting just before the left ventricular injections, reference samples were withdrawn with a Harvard pump from the aortic and right ventricular catheters at a rate of 6 ml/min for 2 min into heparinized glass syringes (14). The blood volume of each reference sample immediately was divided into three scintillation vials and sequential rinses of the syringes were added to the vials. After sacrifice, the lungs were removed and divided into multiple pieces (see below) and the rest of the body was carbonized. Left-to-right ductal shunt expressed as percentage of left ventricular output directed to the lungs was calculated as the sum of radioactivity in the pieces of lung from microspheres injected into the left ventricle divided by the total radioactivity injected into the left ventricle. Cardiac output was calculated from the single reference sample drawn from the distal aorta. We have not detected right-to-left ductal shunts in this animal preparation (15, 16). The average number of microspheres resulting from the right ventricular injections trapped in each piece of lung was ~3,800 microspheres/piece, and the shunt fraction from the left-to-right ductal shunt resulted in >1,000 spheres/piece of lung. The radioactivity of the isotopes was determined simultaneously, and cross-channel contamination was corrected by computer using the appropriate pure isotope standards.

Processing of lungs. After sacrifice, the lungs were removed intact from the animals while still attached to the endotracheal tube. The lungs of the lambs treated at birth and sacrificed at 24±1 min of age were inflated to 30 cm H2O pressure for 30 s. The pressure then was decreased to 15 cm H2O and the lobes were separated by placing small pieces of aluminum foil in the fissures. The lungs of these six lambs inflated uniformly. The lungs were frozen by immersion in liquid nitrogen while maintaining inflation to 15 cm H2O pressure and stored at −20°C. While frozen, the upper lobes were cut from the lungs with a band saw and divided into pieces. The left and right lower lobes were cut into vertical slices, and then into pieces of ~1 cm2. The location of each piece within the lung was recorded and each piece was weighed. The mean weight of the six lungs was 83±6 g and the mean number of pieces resulting from each lung was 101±9 pieces.

The lungs of six of the lambs treated with surfactant at 23 min of age and sacrificed at 46±3 min of age following microsphere injections had an average weight of 78±8 g. The lungs were similarly inflated with 30 cm H2O pressure and then, held at 15 cm H2O distending pressure. However, the lungs did not inflate uniformly. Large areas were atelectatic and other areas of the lungs appeared either partially or fully inflated. Therefore, these three different appearing areas of lung were outlined with a vital stain. The lungs were divided into 51±4 approximately equal-sized pieces, and each piece was categorized as atelectatic, partially aerated, or fully aerated. The lobar location and weight of each piece were recorded.
The microsphere-associated radioactivity was determined for all pieces and then, each piece of lung was homogenized with an Ultra-turax homogenizer (Tekmar Co., Cincinnati, OH) in water such that the homogenate contained 20% tissue (wt/vol). An aliquot of each homogenate was extracted with a 2:1 mixture of chloroform:methanol (17).

The lungs of four lambs treated with surfactant after ventilation similarly were removed and inflated. One lung did not have clearly distinguishable aerated and atelectatic volumes, and this lung was not processed further. The other three lungs were divided into 45±1 pieces weighing 1.7±0.04 g based on the degree of aeration as above; approximately one-third of the pieces from each lung were classified as aerated, partially aerated, or atelectatic. The weighed pieces were dried at 70°C for 3 d and reweighed. A dry-to-wet ratio was calculated for each piece.

Phospholipid analysis. Lipid extracts of the homogenates were concentrated under N2 and aliquots of each extract were plated in duplicate on silica gel H thin layer plates. Phosphatidylcholine was separated from the other lipids by chromatography in one dimension (18), and the phosphatidylcholine was located with iodine vapor. One phosphatidylcholine spot was used for a phosphate assay according to Bartlett (19); the duplicate spot was solubilized in Aquasol II scintillation fluid (New England Nuclear) and the radioactivity quantified. The micromoles of phosphatidylcholine and 14C and 3H radioactivity per gram wet tissue were calculated.

Data analysis and presentation. A mean value per piece of lung for each lamb was obtained for each measurement. All values, then, were divided by the mean value to normalize the numbers. These normalized values are then presented as histograms with interval widths of 10% about the mean value of 1.0. For example, if all pieces from a lung had the same amount of surfactant per gram lung tissue, then 100% of the pieces would be in the interval bracketed by 0.95–1.05 of the mean value (Fig. 1). All pieces having a normalized value <0.15 or >1.85 times the mean were grouped at the extremes of the distribution intervals.

All values are presented as means±SE, unless otherwise indicated. Slopes of curves were calculated by linear regression by the method of least squares, and were tested for significance compared with a slope of 0 by unpaired two-tailed t tests. Differences in the distributions of sur-

![Figure 1](https://via.placeholder.com/150)

Figure 1. Normalized distributions for six lambs treated with surfactant at birth. All values were calculated as described in Methods and are presented as mean±SE percentage of pieces of the lungs vs. the 10% distribution intervals. (A) Distribution of surfactant phosphatidylcholine per gram lung tissue. (B) Distribution of right ventricular output to the lung per gram tissue at 13±1 min of age. (C) Distribution of right ventricular output to the lung per gram tissue at 23±2 min of age. (D) Distribution of ratios of relative blood flow from the left ventricle via the ductus arteriosus at 22±1 min of age to relative blood flow from the right ventricle at 23±2 min of age. (E) Distribution of ratios of right ventricular blood flow per gram tissue at 23±2 min of age to surfactant per gram tissue.
Surfactant and blood flow to pieces of lung were tested for significance by ANOVA followed by the Student Newman-Keuls multiple comparison procedure.

Results

Description of lambs. The umbilical vein blood-gas values for these 16 lambs were pH, 7.31±0.02; P02, 28±2 mmHg; and PCO2, 43±2 mmHg. The lambs weighed 1.8±0.1 kg. The six lambs treated with surfactant at birth and sacrificed at 24±1 min of age had aortic blood-gas values of pH, 7.39±0.04; P02, 204±36 mmHg; and PCO2, 34±4 mmHg just prior to sacrifice while being supported with 28 cm H2O pressure. Their respiratory tidal volumes and lung compliance values were 10.9±0.8 ml/kg and 0.42±0.03 ml/cm H2O per kg, respectively. The mean blood-gas values for the other 10 lambs were pH, 7.05±0.04; P02, 66±18 mmHg; and PCO2, 81±6 mmHg at 20 min of age just prior to surfactant instillation. At that time, expiratory tidal volume was only 2.6±0.9 ml/kg and lung compliance was 0.09±0.01 ml/cm H2O per kg. The lambs responded to the surfactant with blood-gas values at sacrifice 23 min after treatment of pH, 7.23±0.03; P02, 161±32 mmHg; and PCO2, 44±5 mmHg while the lambs were being ventilated with 28±1 cm H2O peak inspiratory pressures. Mean tidal volume increased to 7.0±0.7 ml/kg and compliance increased to a value of 0.24±0.02 ml/cm H2O per kg. Cardiac outputs were measured for six lambs and were 208±28 cm3/kg per min. The left-to-right ductal shunt measured in these lambs was 31±3% (range, 24–42%) of left ventricular output. This shunt value has been corrected for the 4±1% of spheres that were not trapped by the systemic capillary bed, as determined by the reference sample drawn from the right ventricle. Less than 4% of microspheres injected into the right ventricle were detected in the carcass, documenting relatively complete trapping of microspheres by the pulmonary vasculature and no significant right-to-left ductal shunt.

Dry-to-wet weight ratios. The dry-to-wet weight ratios were calculated for the differentially aerated pieces of lung from three lambs following treatment with surfactant after a period of ventilation. The mean±SD values were: aerated, 0.0854±0.0106; partially aerated, 0.0872±0.0154; and atelectatic, 0.0946±0.0070. The ratios for aerated and partially aerated pieces of lung were not different; however, there was less water in the atelectatic pieces than in either of the other two groups (P > 0.01). The absolute differences were very small and subsequent data are expressed per gram wet weight without a correction for the ~1% difference in lung water between atelectatic and aerated or partially aerated pieces.

Distribution of surfactant. The lungs of lambs treated at birth with surfactant were not atelectatic when inflated after sacrifice, and the surfactant was distributed relatively homogeneously throughout the pieces of lung tissue (Fig. 1 A). 48% of the pieces contained an amount of surfactant that was within ±25% of the mean, and few pieces contained either very little or large amounts of surfactant. For each lamb, the distribution of surfactant to different lobes of the lung was not uniform; however, mean values of surfactant per gram indicate no preferential lobar distribution of surfactant after treatment at birth (Table I).

In contrast, the lungs of the lambs that were treated after a period of ventilation did not inflate uniformly. Large contiguous volumes of lung were only partially aerated or entirely

Table I. Relative Distribution of Surfactant and Blood Flows by Lobes

<table>
<thead>
<tr>
<th></th>
<th>Left upper lobe</th>
<th>Left middle lobe</th>
<th>Left lower lobe</th>
<th>Right upper lobe</th>
<th>Superior right middle lobe</th>
<th>Inferior right middle lobe</th>
<th>Right lower lobe</th>
<th>Total left lung</th>
<th>Total right lung</th>
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<tr>
<td>Lambs treated at birth, NS* per gram lung</td>
<td>0.83 ±0.50</td>
<td>1.07 ±0.51</td>
<td>0.95 ±0.42</td>
<td>1.08 ±0.47</td>
<td>1.05 ±0.43</td>
<td>0.98 ±0.33</td>
<td>1.05 ±0.37</td>
<td>0.98 ±0.45</td>
<td>1.02 ±0.40</td>
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<td>Lambs treated after ventilation, NS per gram lung</td>
<td>1.61±0.88</td>
<td>1.94±0.75</td>
<td>1.04 ±0.75</td>
<td>1.03±0.75</td>
<td>1.02±0.94</td>
<td>1.92±0.57</td>
<td>0.58 ±0.41</td>
<td>1.26±0.84</td>
<td>0.77 ±0.65</td>
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<tr>
<td>RV* blood flow in lung before NS treatment</td>
<td>0.95 ±0.16</td>
<td>0.93 ±0.17</td>
<td>1.06 ±0.21</td>
<td>0.94 ±0.56</td>
<td>0.79 ±0.19</td>
<td>0.96 ±0.19</td>
<td>1.07 ±0.21</td>
<td>1.03 ±0.21</td>
<td>0.98 ±0.31</td>
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Values are given as mean±SD. * NS, natural surfactant; RV, right ventricle. † Total left lung > total right lung (P < 0.01). § Upper lobes > lower lobes (P < 0.05).
atelectatic to visual inspection. Sequentially, more surfactant was found in aerated than partially aerated pieces of lungs, and more was found in partially aerated than in atelectatic pieces of lung (Table II). The aerated pieces of lung contained 5.4 times more surfactant per gram than atelectatic pieces and 2.5 times more surfactant per gram than partially aerated pieces. Thus, visual inspection of the lungs held at 15 cm H2O static airway pressure accurately identified the location and distribution of the surfactant in lambs treated with surfactant after a period of ventilation. The pieces of lung for these lambs distributed such that only 24% of pieces received amounts of surfactant per gram tissue within ±25% of the mean (Fig. 2A). On the average, more surfactant was found in pieces from the left than right lungs ($P < 0.01$), and the upper lobes of each lung received

<table>
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<th>Table II. Relative Amount of $^{3}$H-labeled Natural Surfactant per Gram Tissue (No. of pieces assayed)</th>
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<tr>
<td><strong>Aerated</strong></td>
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<tr>
<td>Lamb 1</td>
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<tr>
<td>2</td>
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<td>3</td>
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<td>4</td>
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<tr>
<td>5</td>
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<tr>
<td>6</td>
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<tr>
<td>Average</td>
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</table>

Figure 2. Normalized distributions for six lambs treated with surfactant at 23±1 min of age. All values were calculated as described in Methods and are presented as mean±SE percentage of pieces of the lungs vs. the 10% distribution intervals. (A) Distribution of surfactant phosphatidylcholine per gram lung tissue. (B) Distribution of right ventricular output to the lung per gram tissue at 21±1 min of age. (C) Distribution of right ventricular output to the lung per gram tissue at 44±3 min of age. (D) Distribution of ratios of relative blood flow from the left ventricle via the ductus arteriosus at 42±3 min of age to relative blood flow from the right ventricle at 44±3 min of age. (E) Distribution of ratios of right ventricular blood flow per gram tissue at 44±3 min of age to surfactant per gram tissue.
more surfactant than the lower lobes (Table I). The synthetic dipalmitoylphosphatidylcholine was distributed to the lungs of both groups of lambs in the same pattern as was the natural surfactant (data not shown).

**Distribution of pulmonary blood flow.** Radiolabeled microspheres were injected into the right ventricle to measure the distribution of blood flow to the lungs. In lambs treated with surfactant at birth, the pulmonary blood flow at 13 and 23 min of age was relatively homogenous to the pieces of lung. At the early and late injection times, 69 and 63% of the pieces received blood flows within ±25% of the mean (Fig. 1 B and C). In the five lambs receiving injections of microspheres into the left ventricle, the distribution of the normalized shunt flow from the aorta via the patent ductus arteriosus to the lungs had the same distribution as the blood flow from the right ventricle (Fig. 1 D). The results indicated very good mixing of blood from the two ventricles.

Pulmonary blood flow distribution was measured in lambs treated with surfactant after a period of ventilation by injecting microspheres immediately before and 22±2 min after surfactant treatment. The histograms of normalized blood flow per gram tissue before surfactant treatment indicated a relatively homogenous distribution of blood flow throughout the lung with 77% of the pieces receiving flows within ±25% of the mean (Fig. 2 B). There was no consistent pattern of preferential blood flow from the right ventricle to different lobes of the lung per gram tissue (Table I). After surfactant therapy, the overall distribution of flow was somewhat less homogenous with 67% of the pieces receiving flows within ±25% of the mean flow (Fig. 2 C). The histogram of the ratios of normalized blood flow from the right ventricle and from the left-to-right ductal shunt in these lambs documented good mixing of the pulmonary blood flow from the two sources (Fig. 2 D).

**Matching of surfactant and blood flow distributions.** In lambs treated with surfactant at birth, the normal distribution of the ratio of surfactant per gram lung to blood flow per gram lung resulted in a more dispersed distribution than was the distribution of surfactant per gram lung. Only 35% of pieces of lung received within ±25% of the mean value (Fig. 1 E). 15±4% of the pieces of lung from the six lambs had a ratio of surfactant to blood flow of >1.85. This dispersion of the distribution histogram indicated that blood flow distribution did not match surfactant distribution. The correlation of surfactant distribution with blood flow for the 133 pieces of lung from a single lamb indicated a significant negative effect of the presence of surfactant on the distribution of pulmonary blood flow (Fig. 3). The decrease in perfusion to pieces of lung receiving relatively large amounts of surfactant was detected by linear regression analysis in four of six lambs (P < 0.01). No correlation between distributions of perfusion and surfactant was detected in the other two lambs.

An effect of surfactant distribution on pulmonary blood flow distribution also was detected in the lambs treated with surfactant after a period of ventilation. The ratio of surfactant distribution to blood flow distribution was bimodal (Fig. 2 E) and similar to that of the histogram for the surfactant distribution alone (Fig. 2 A). The effect of surfactant on pulmonary blood flow distributions was tested in these six lambs by comparing by linear regression analysis changes in blood flow distributions before and after surfactant treatments. In four of six lambs, there was a significant negative correlation between surfactant per gram tissue and pulmonary blood flow per gram tissue (P < 0.04). The presence of surfactant in the tissue resulted in a relative fall in blood flow to those pieces of lung and an increase in relative blood flow to pieces of lung that received very little of the exogenously administered surfactant (Fig. 4). In two of six lambs, pulmonary blood flow distribution did not change significantly.

**Discussion**

We treated lambs with surfactant at birth and before the initiation of ventilation because most experimental demonstrations of the efficacy of surfactant administrations have involved treatment of prematurely delivered animals before the onset of ventilation (1–4, 20, 21). Surfactant treatments of premature rabbit lungs improve lung compliance with a “dose-response curve” that demonstrates no further improvements in compliance measurements at doses of surfactant >1.1 mg surfactant lipid/g lung (22). The compliance changes were insensitive to the volume used to instill the surfactant. Lungs of premature rabbits and lambs treated with surfactant at birth were generally well aerated visually; however, patchy atelectatic areas were described and confirmed microscopically (1, 9). This patchy atelectasis could

Figure 3. Correlation of normalized (norm.) values of right ventricular blood flow per gram tissue with surfactant per gram tissue for the 133 pieces of lung from a lamb. The slope of the regression line fit by the method of least squares is significantly different from zero (P < 10⁻³).
represents volumes of lung not receiving the exogenously administered surfactant; however, such atelectasis also is characteristic of the lungs of the term newborn in the early hours of life. The assumption has been that the normal physiologic clearance of fetal lung fluid after birth via the distal airways and alveoli primarily to the vascular space (23) will result in the exogenously administered surfactant being well-distributed throughout the lung. The surfactant was relatively homogeneously distributed throughout the uniformly inflated lungs of lambs treated with surfactant at birth. However, this technique of surfactant administration is not easily adapted to clinical practice.

In all clinical studies to date, surfactant has been administered from shortly after the onset of breathing to up to 33 h after ventilatory support (6, 7, 24). Except for the use of dry surfactant (24), the infants were treated by tracheal injection with a suspension of surfactant in a volume of ~3.5–7 ml/kg containing ~50–125 mg surfactant lipid/kg (6, 7). The dose was based on a dose-response curve for the treatment of prematurely delivered lambs with natural sheep surfactant (25). The volume of the suspension has been an empirical compromise designed to optimize surfactant distribution while minimizing potential problems resulting from flooding the airways. While the efficacy of varying volumes of the suspension has not been studied systematically, 7 ml/kg surfactant suspensions do not significantly change gas exchange or cardiovascular status in premature lambs (26). Thus, we chose a dose and volume of surfactant suspension consistent with previous clinical and experimental usage. Surfactant was localized to alveoli following administration to fetal or adult animals; however, the distribution seemed to be non-uniform (1, 9). Radiolabeled particles given to rats and hamsters by instillation in 1.5 ml/kg suspensions were nonuniformly distributed with preferential deposition to dependent lung locations, while aerosolized particles were more evenly distributed with preferential deposition to the apical lobes (11).

We found more surfactant in the left rather than right lungs and in upper vs. lower lobes of the lungs of lambs treated after a period of ventilation. Since these lungs were cut in pieces based on a visual assessment of degree of inflation at a static pressure of 15 cm H2O, the distribution of surfactant was not strictly comparable with that measured in the lungs of lambs treated at birth with surfactant. However, the atelectatic vs. aerated areas had predominantly lobar distributions and were large compared with the size of pieces of lung actually sampled (~1 cm3). Thus, random sampling of pieces of lung would not have changed the overall distribution pattern very much. While all pieces of lung received some surfactant, the pieces of lung that were aerated after static inflation contained relatively large amounts of surfactant, and atelectatic pieces received much less surfactant. In contrast, the lungs of lambs treated at birth were uniformly inflated despite some pieces of lung receiving relatively little surfactant. For example, ~10% of the aerated pieces of lung of lambs treated at birth contained amounts of surfactant per gram that did not result in inflation in lambs treated after ventilation. More than 40% of the aerated pieces from lungs of lambs treated at birth contained amounts of surfactant per gram lung that resulted in only partial aeration in the previously ventilated lambs. Possibly following ventilation, larger amounts of surfactant are needed per gram lung to achieve aeration because of bronchiolar epithelial damage and the resultant pulmonary edema that occurs with ventilation of the surfactant-deficient lung (20, 21). Such damage may result in the entrance into the airway of proteins that interfere with the surface tension-lowering properties of surfactant (12). Thus, it is likely that one must accept a less effective and relatively nonhomogeneous distribution of surfactant to the immature lung following airway inflation after the initiation of ventilation.

We measured relative pulmonary blood flow by measuring the trapping of radiolabeled microspheres by the pulmonary microvasculature (27). In adult animals, such measurements are made following central venous rather than right ventricular injections to ensure complete mixing of the blood and microspheres. While we have not detected large right-to-left shunts through the foramen ovale in similarly studied lambs (15, 16), we injected the microspheres into the right ventricle to avoid such shunts. The completeness of mixing can be assessed only indirectly in this model. Very few pieces of lung received large numbers or small numbers of microspheres, and the relative number of microspheres within a piece of lung with consecutive injections was similar. These results indicate good mixing. A large or variable ductal shunt also could perturb the distribution
data if that shunt flow did not mix well with the pulmonary artery blood flow. While the lambs had large left-to-right ductal shunts that accounted for 24–42% of the left ventricular output, this shunted blood was distributed equivalently to both lungs, implying good mixing.

Surfactant treatment after a period of ventilation resulted in a prompt increase in PO₂ to mean values over 200 mmHg, and a mean PO₂ of 161 mmHg at sacrifice. The percent functional right-to-left pulmonary shunt can thus be assumed to be ~40% of the cardiac output (28). We hypothesized that pulmonary blood flow would be directed toward regions of lung receiving the most surfactant, and thus, those areas should be better ventilated. While surfactant distribution and static inflation correlated very well in those lambs treated after a period of ventilation, we do not know which areas of lung actually were ventilated, and thus, where gas exchange actually occurred in the mechanically ventilated lung. The PO₂ values suggest that significant gas exchange occurred in regions of lung that appeared aerated and only partially aerated with static inflation because only 39% of lung pieces were assessed as being aerated, while 32 and 27% of lung pieces, respectively, were partially aerated or atelectatic. The decrease in blood flow to regions of lung receiving relatively large amounts of surfactant with a relative increase in blood flow to those regions receiving small amounts of surfactant in 8 of 12 lambs might be explained by pressure relationships within the lungs. The nonhomogeneous distribution of surfactant will result in a wide range of compliance changes throughout the lung. In fully aerated areas, the peak inspiratory pressures of 28–31 cm H₂O may be transmitted to the pulmonary microvasculature and inhibit blood flow (29). In fact, significant overdistension may occur in areas receiving large amounts of surfactant. In areas of low compliance, there may be very little pressure transmitted from the airways to the pulmonary microvasculature. These maladaptive changes in pulmonary perfusion following surfactant instillation are acute changes and parallel the acute changes noted in the adult dog following segmental atelectasis (30). The chronic effect of surfactant instillation on pulmonary perfusion remains to be studied.

Acknowledgments

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References


