Fetal pulmonary injury following single high-dose intra-amniotic betamethasone treatment in preterm goat kids

Preterm keçi yavrusunda tek ve yüksek doz intra-amniyotik betametazon tedavisi sonrasında fetal pulmoner zedelenme

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Abstract

Objective: Fetal lung maturation is an extremely important process that is necessary for the survival of the neonates. Conventionally, corticosteroids are administered maternally for inducing fetal lung maturation in preterm fetuses. Alternatively, single-dose intra-amniotic (IA) treatment might be speculated to improve lung maturity. In the goat model, we recently showed that high-dose IA betamethasone (BM) was associated with an increased number of stillbirths and emphysematous changes. The aim of the present study is to expand our previous findings and evaluate the histopathological effects of IA injection of a single high-dose of BM 48 h before induced preterm delivery, using our previously collected specimens.

Material and Methods: Five hair goat fetal lungs that had received 8 mg/kg IA BM at gestational day 118 (term, 150 days) and scheduled for preterm delivery by cesarean section at day 120 of gestation were examined pathologically. Specimens were stained with hematoxylin and eosin (HE) and were interpreted by light microscopy.

Results: The histopathological examination of the fetal lungs revealed edema, hemorrhage, slight inflammatory reaction, marked emphysema, and desquamation of the pneumocytes and bronchiolar or bronchial epithelial cells.

Conclusion: High-dose IA BM administrations to induce lung maturation can paradoxically cause severe pathological lesions in the fetal lungs. These might explain the toxic effects we encountered with this mode of treatment. (J Turkish-German Gynecol Assoc 2012; 13: 242-6)

Key words: Animal models, betamethasone, corticosteroids, intra-amniotic, lung maturation

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Introduction

The pulmonary system is among the last of the fetal organ systems to mature, both functionally and structurally. Because the immature pulmonary system may not oxygenate the neonate adequately, preterm birth can lead to significant neonatal morbidity or mortality (1). Reduced lung function during infancy has been shown in association with preterm birth, probably persisting through adulthood (2-4). Immature lungs may increase the risk of respiratory distress and death among both term and preterm infants during the neonatal period (2). Moreover, low birth weight secondary to preterm
delivery has been associated with reduced lung function and increased death rates from chronic obstructive airway disease in adult life (4). Animal studies and clinical human data have revealed that fetal lung maturation takes place after maternal corticosteroid administration (5). In line with this, elevated cortisol levels have been shown to enhance lung maturation (6). However, the efficacies of various corticosteroid analogs, total dose, dosing intervals, and the means of administration for fetal lung maturation have not been exactly defined. Alternative routes of administration, such as the IA route might be hypothesized to be more effective than the standard maternal method of administration. Our previous animal study showed that the dose and the application route of betamethasone (BM) are particularly important for fetal lung maturation (7). In the mentioned investigation (7), we proposed that higher (than that of previously used) doses of IA can be an attractive treatment option to induce improved antenatal lung maturation. However, we found that a high (8 mg/kg) dose of intra-amniotic (IA) and fetal i.m. (4 mg/kg) BM was not superior to the standard dose and the maternal route of administration in the goat model. Moreover, IA BM was associated with an increased number of stillbirths and macroscopic emphysematous changes, compared to controls. Despite these toxic effects, detailed histopathological examination of the fetal lungs was not included in this previous paper (7), and we were not able to identify the exact underlying mechanisms of pulmonary injury following the IA route.

Here, we report the detailed histopathological findings in the fetal lungs of kids that we gathered from our previous experimental design. These were antenatally treated with high-dose (8 mg/kg of estimated fetal weight) IA BM. Our aim was to delineate the causes for the unexpected toxic effect we encountered following high-dose IA therapy.

**Material and Methods**

All the experiments were approved by the institutional animal use and care committee at Süleyman Demirel University and performed in accordance with the National Institute of Health Guidelines for the Care and Handling of Animals. The number of animals was restricted (n=5), as suggested by the animal ethics committee. Details of the procedures were described thoroughly in our previous study (7). Briefly, five female hair goats (Capra hircus), 2 to 4 years old and 40±5 kg in weight were used for the study. Goats were fed on standard feed and tap water ad libitum. A singleton structurally normal ongoing pregnancy was confirmed by ultrasonography (Echo Camera SSD-500, Aloka, Tokyo, Japan) at gestational days 60, 75, and 90 (term pregnancy, 150-155 days). At day 118, amniocentesis with a 21 G needle under ultrasound guidance was performed (Figure 1). Betamethasone disodium phosphate plus betamethasone acetate (Celestone chronodose amp, Schering-Plough Inc, Istanbul, Turkey) equivalent to 8 mg/kg of estimated fetal weight was injected into the amniotic sac. Subsequently, at day 120, excluding the stillborn kids, cesarean section (CS) with paralumbral skin and dorsal curvature uterine incision was performed under sterile conditions at gestational day 120 (corresponding to 31-32 weeks of gestation in human pregnancy). Details of the operation are provided elsewhere (8). After preterm delivery, the kids were euthanized with high-dose (50 mg/kg) sodium thiopental (Pental Sodyum, IE Ulugay, Istanbul, Turkey) administered via the umbilical catheter, the trachea was clamped for 3 min to maintain airway pressure and necropsy performed. At macroscopic examination, any visible lung rupture or pulmonary interstitial emphysema was recorded. The lungs were excised en bloc, the trachea removed to the bifurcation and their wet weight was measured (Figure 2). The lungs were then fixed by 10% formalin inflation into the airways, and fixation continued in formalin for three days. Five different samples were taken from all of the lungs (left apical, left diaphragmatic, right apical, right cardiac and right diaphragmatic lobes). Then, tissue samples were routinely processed, paraffin embedded and cut into 5 mm sections, and slides were stained with hematoxylin and eosin (HE) to be interpreted by
light microscopy. Histopathological changes were examined in a blinded manner under the 40x objective of a Nikon E-600 trinocular microscope and microphotography apparatus.

**Results**

There were 3 stillbirths out of 5 fetuses that were given IA BM. The fetal lungs from one of those kids could not be examined because of a severe autolytic reaction, probably secondary to significant trauma by other goats nearby following stillbirth. The histopathological observation of the remaining fetal lungs (a total of 8 gross specimens from 4 fetuses) revealed edema, slight inflammatory reaction, hemorrhage, marked emphysema, and desquamation of the pneumocytes (Figures 3-5). Edema was typically localized around the vessels at the interstitial tissue. Protein-rich eosinophilic edema fluid was also present in some alveolar spaces. Slight neutrophilic and mononuclear inflammatory cells were evident around the pulmonary vasculature and around bronchioles. Marked increase in alveolar macrophages was a prominent finding in the alveolar lumens. Severe desquamation was observed in alveolar cells. In addition to alveolar cells, desquamation was also prominent in bronchiolar and bronchial epithelial cells.

Pulmonary lesions were more severe in specimens from stillbirths compared to live births. Marked areas of hemorrhage and edema were common findings in stillborn kids. Thickening at the septal areas due to edema and inflammatory cell infiltrations were also characteristic in these kids. Some of the alveolar spaces were filled with desquamated epithelial cells and inflammatory cells. Atelectatic areas were also commonly encountered in fetuses with stillbirth (Figures 6, 7).

![Figure 3](image3.png) **Figure 3.** Histopathological appearance of the lungs of a kid showing severe hemorrhage in septal tissue around the bronchi (arrow heads), HE, Bar=50 µm

![Figure 4](image4.png) **Figure 4.** Severe emphysemic areas (arrow heads) and edematous interstitial tissue (arrow) in a kid, HE, Bar=100 µm

![Figure 5](image5.png) **Figure 5.** Severe alveolar emphysema (arrows) in the lungs of a kid, HE, Bar=100 µm

![Figure 6](image6.png) **Figure 6.** Inflammatory reaction (arrows) in a lung of a stillborn kid, HE, Bar=50 µm
BM given intra-amniotically at high doses (>2 mg/kg) can be speculated to be absorbed by the chorioamniotic membranes and perhaps by the fetal surface of the placenta, leading to reabsorption by the fetus. Following reabsorption in a cyclic manner, BM would be expected to accumulate toxic levels in the fetus. This toxic accumulation probably initiates lung injury and finally causes fetal death. Some fetuses tend to demonstrate an exaggerated response with prominent edema and inflammation, as supported by our findings from stillbirths. Another possibility, however, is the emergence of such histopathological characteristics following fetal loss. This may be unlikely, as the specimens were available in a short time after fetal loss except in one case, which was excluded from analysis. Septal thickening was an important histopathological feature that we encountered. In contrast, low (2 mg/kg) doses of IA BM were reported to be associated with thinning of alveolar walls, which is a favorable finding for lung maturation. Therefore, IA BM at very high doses (8 mg/kg) not only causes lung injury but also disturbs normal fetal lung maturation. It must, however, be noted that even standard doses of BM administered maternally for fetal lung maturation can induce certain adverse histopathological changes. For example, the results of a pregnant sheep model indicated that 70% of lambs delivered at 128 days of gestation, 24 h after a single injection of 0.5 mg/kg maternal i.m. BM developed pulmonary interstitial emphysema, compared with fewer than 5% of control animals (12).

Our data from the present design would be insufficient to explain the exact mechanisms and causes of increased fetal death rate (60%) following high-dose IA BM, as we do not have specimens from other organs. However, toxicity caused particularly by BM in other organ systems, including the fetal central nervous system, can be expected. At this stage, we do not know whether these unfavorable effects on the fetus are specific for BM or whether they can be generalized to other glucocorticoids.

Budesonide is a steroid derivative with minimal systemic absorption and almost no placental transfer. Budesonide is conventionally used as an oral inhalant in the treatment of childhood asthma. Limited data from animal (sheep) experimentation reveals no perinatal loss with relatively high doses of IA budesonide administration; moreover, its pulmonary maturational effects were reported to be comparable to the standard maternal BM therapy (9, 11). Hence, budesonide and other corticosteroids with minimal absorption and limited toxicity can be candidates for fetal therapy, including IA administrations. Further experimental studies will be needed on IA budesonide therapy.

Conclusion

High-dose IA BM causes severe pathological lesions, including edema and hemorrhage in the fetal lungs. These changes may elucidate the toxic effects and increased fetal losses we encountered with this mode of treatment.

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Conflict of interest
No conflict of interest was declared by the authors.

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