Continuous amnioinfusion via an epidural catheter following spontaneous membrane rupture: A case report

Preterm premature rupture of membranes (PPROM) is seen in 3% of all pregnancies, and is a frequent cause of preterm birth, neonatal mortality and morbidity. The most important complications are maternal and foetal infection, prematurity, umbilical cord compression, hypoxia or asphyxia due to cord prolapse, pulmonary hypoplasia and extremity deformities. The basic approach to PPROM therapy aims to prevent premature birth and the development of foetal distress, and decrease the risk of maternal and foetal infection, and amniotic fluid loss. In compliance with these objectives, alternatives of PPROM therapy demonstrate a wide spectrum, including watchful waiting, amniopatch application, recurrent amnioinfusions and emergency birth. However, repeated amnioinfusions in cases of fluid loss, especially within 6 hours of therapy, provides only minimal benefit. In this case presentation, we attempted to describe a different and cost-effective continuous amnioinfusion technique performed to confer survival benefit for an immature anhydramniotic foetus affected by PPROM at the border of viability.

Key words: Amnioinfusion, preterm premature rupture of membranes, amniopatch

Anamnestic data:

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Case Report

A 37-year-old pregnant woman (gravida 5, para 1, abortus 3) at her 22nd gestational week with antiphospholipid syndrome who had previously given birth to a 27-week premature healthy and still living infant consulted our clinic with a complaint of ‘water breaking’ amniotic fluid. On ultrasonographic examination (US) (by GE Voluson 730 Pro 4D ultrasound device), the foetus was at nearly the 22nd gestational week, and the amount of total amniotic fluid according to the amniotic fluid index (AFI) was less than 2 cm² as viewed from all quadrants. PPROM was diagnosed with a speculum, which showed vaginal pooling of the fluid on the posterior fornix, and by a positive alpha microglobulin-1 protein test (Amnisure; Aren Medical, İstanbul, Turkey) of the vaginal fluid. It is probable that the membranes had ruptured much earlier and that the fluid loss had been misinterpreted by the patient as a weakness of the bladder. The cervical length was 25 mm. The patient's treatment started with prophylactic antibiotics, a regimen of bed rest, daily monitoring of inflammation parameters, such as the white blood cell (WBC) and C-reactive protein (CRP), and the monitoring of vital signs, especially temperature. The CRP and WBC measurements were <1 mg/dL and 17,700/mm³, respectively. Prophylactic IV antibiotic therapy was initiated with 4×1 g of ampicillin+sulbactam (Sulbaksit; Tüm Ekip İlaç, İstanbul, Turkey) of the vaginal fluid. It is probable that the membranes had ruptured much earlier and that the fluid loss had been misinterpreted by the patient as a weakness of the bladder. The cervical length was 25 mm. The patient's treatment started with prophylactic antibiotics, a regimen of bed rest, daily monitoring of inflammation parameters, such as the white blood cell (WBC) and C-reactive protein (CRP), and the monitoring of vital signs, especially temperature. The CRP and WBC measurements were <1 mg/dL and 17,700/mm³, respectively. Prophylactic IV antibiotic therapy was initiated with 4×1 g of ampicillin+sulbactam (Sulbaksit; Tüm Ekip İlaç, İstanbul, Turkey). An amniopatch was performed primarily with the use of platelet suspension. The autologous platelet concentrate (PC) was produced by platelet pheresis using the MCS plus® blood cell separator (Haemonetics Corporation, Braintree, Massachusetts, USA). The patient’s platelet precount of 230×10³/µL was used for programming the device. After rewarming to 37°C, the PC (30 mL) was administered first, followed by 20 mL of cryoprecipitate via a 22-gauge needle (Spinocan; Braun, Mensungen, Germany), directed into an available pocket of amniotic fluid. The amnioinfusion and the amniopatch were performed twice. After application of the amniopatch, the amniotic fluid continued to drain actively. While monitoring the patient, uterine contractions started to develop, prompting initiation of 2 g MgSO₄ (Magnezyum Sulfat %15; Osel, İstanbul, Turkey) IV per hour as a tocolytic agent. Within six hours of each application of the amnioinfusion and the amniopatch, the AFI was less than 2 cm as observed from all quadrants. Therefore, application of a different and cheaper technique of continuous amnioinfusion was decided upon. Before the procedure, a detailed and undersigned informed consent was obtained from the patient and the family. Continuous amnioinfu-

Figure 1. Epidural catheter set used for the patient

Figure 2. Post-procedural fixation of epidural catheter on the anterior abdominal wall
prolapse, which necessitated an emergency caesarean section. Postoperatively, the mother was in stable health state, and she was discharged on the 4th postoperative day in full health. At one and five postnatal minutes, the Apgar scores of the infant were 5 and 6 points, respectively. The baby was immediately intubated, and 4 mL/kg of surfactant (Survanta; Abbott, North Chicago, USA) was administered through an endotracheal tube, with the diagnosis being respiratory distress. The newborn was connected to a mechanical ventilator. The neonate had a poor general status, weak spontaneous activity, and thin, bright and plethoric skin. The weight was 550 g (<3%), and the height and the head circumference were 27 cm (<3%) and 21 cm (<3%), respectively. The foetal heart rate was rhythmic, and no cardiac murmur was heard. The external appearance of the foetus was of female gender. Her haematocrit was 38%, which necessitated transfusion of an erythrocyte suspension at a rate of 15 mL/kg. Administration of dopamine (Dopmin amp; Orion, Espoo, Finland) and dobutamine (Dobcard; Vem ilac, Istanbul Turkey) was initiated due to her deteriorated peripheral circulation. As the mother’s membrane rupture was present for three weeks, empirical antibiotic therapy was started after drawing blood samples for blood culture. The antibiotic therapy was discontinued after the absence of any microbial growth on culture media was reported. Parenteral feeding was started on the first day and minimal enteral feeding on the second day. For the relief of symptomatic patent ductus arteriosus, ibuprofen was used. Respiratory support was provided with a mechanical ventilator for nine days and with a nasal continuous positive airway pressure for four days. An ophthalmologic examination detected Stage 1 retinopathy. On a follow-up visit, no progression of the disease was observed. On the contrary, it had regressed to Stage 0. A hearing test was unremarkable. The infant gained weight, and she was 2070 g at discharge on the 128th day. The infant is now six-months old and is fed with breast milk and fortified infant formula. On follow-up visits, her development was found to be in accordance with her gestational age.

Discussion

The current study discussed the use of a relatively cost-effective continuous amnioinfusion technique that can be applied for extended periods in a single session. The method can be used in patients with amniotic fluid loss occurring within less than six hours after undergoing amnioinfusion and an amniopatch who require subsequent repeat applications of these painful procedures and who exhibit foeto-maternal risks. For the amniopatch application, platelets and cryoprecipitates are usually used because they have been found to be necessary components for a successful and safe therapy for PROM. Sipurzynski et al. (7) did not observe any side effects or complications during autologous platelet pheresis and application of the amniopatch. In cases of PROM occurring after an iatrogenic procedure, such as amniocentesis performed between the 16th and the 24th gestational weeks, the amniopatch is an appropriate procedure in the absence of intra-amniotic infection. An amniopatch was found to be successful in nearly 50% of such patients (4). In amniopatch application, platelets seem to migrate to the site of the defect and occlude the defective site. Platelet activation and fibrin formation at the site of rupture initiate the healing process (8). However, after amniopatch application, the development of a fibrous band may cause constriction of an extremity or the umbilical cord. In the present case, the failure of the amniopatch application was probably related to a larger membrane defect.

Before 24 weeks of gestational age, PPROM has a predicted perinatal mortality of nearly 90% and amniotic infection is frequently seen. Perinatal outcomes of an iatrogenic PROM like amniocentesis are relatively worse when compared with spontaneous PROM. Repeated amniopatch or saline amnioinfusion techniques can be applied. However, serial applications of this method are quite painful and carry important foeto-maternal risks (9, 10).

Tchirikov et al. (11) previously applied a method similar to that reported here. They used an amniotic fluid replacement port. Our system works in a similar way. There were some disadvantages with the system described by Tchirikov et al. (11). The catheter had to be detruncated and reconnected to a metal tube and then to the port capsule. Fluid leakage around the catheter was also detected. Additional disadvantages were important technical problems and the need for anaesthesia (11). Our application technique is independent of gestation weeks. Saline is infused in a method similar to that of Tchirikov et al. (11). The technique did not lead to fluid leakage in the presented case. Moreover, it does not require detailed technical information, and is relatively cost-effective. In addition, it does not necessitate additional procedures, such as skin incision.

Inflammation secondary to microbial invasion of the amniotic cavity is responsible for more than half of cases with preterm birth and PPROM (11). However, in cases with previable PPROM (<22-23 gestational weeks), termination of the pregnancy can be recommended because of poor prognosis secondary to infection and/or pulmonary hypoplasia (3). In the present study, no infection developed, although there was a slight increase in CRP. The infant remained connected to the mechanical ventilator for only nine days and continues to exhibit no pulmonary problems. Continuous intra-amniotic infusion of an isotonic saline solution may be able to protect the patient from the development of amniotic infection syndrome and pulmonary hypoplasia. An isotonic saline solution, which drains from the uterine cavity through the cervical canal, provides continuous irrigation. This might protect the foetus and the mother from ascending infections.

In conclusion, continuous amnioinfusion using an epidural catheter seems to be an appropriate treatment for PPROM in the second trimester of pregnancy. The aim of such treatment is to prevent cord compression secondary to anhydramnios and pulmonary hypoplasia and to enhance the survival potential of the foetus. The risk-benefit ratio and the expectations of the patient should be taken into consideration in the application of the method. Generally, the only wish of the mother is to embrace their baby.

Ethics Committee Approval: N/A.

Informed Consent: Written informed consent was obtained from patients who participated in this study.
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**References**