In Utero Meconium Exposure Increases Spinal Cord Necrosis in a Rat Model of Myelomeningocele

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Background/Purpose: The rationale for in utero repair of myelomeningocele has been supported experimentally by the observation of preserved neural function after prenatal closure of surgically created defects compared with nonrepaired controls. The mechanism of injury to the exposed neural elements is unknown. Postulated mechanisms include trauma to the herniated neural elements or progressive injury from amniotic fluid exposure as gestation proceeds. A component of amniotic fluid that may contribute to neural injury is meconium. In the current study the effect of human meconium on the exposed spinal cord in a fetal rat model of myelomeningocele was examined.

Methods: Twenty time-dated pregnant rats underwent laparotomy at 18½ days of gestation. The exposed uterus was bathed in ritrodrine for tocolysis. The amniotic cavity was opened over the dorsal midline of the fetal rat, and, under a dissecting microscope (×25), a 2- to 3-level laminectomy was performed. Under magnification (×40), the translucent dura was opened using a 25-gauge needle as a knife. Two fetuses per dam were operated on. In the control group, the amniotic fluid was restored with saline solution, whereas in the experimental group a solution of Human meconium diluted (10%) in saline was used to restore the amniotic fluid. Fetuses were harvested by cesarean section at 21½ days’ gestational age. The liveborn pups were then killed and fixed in 10% formalin. Sections 10 μm thick were stained with H&E and studied by light microscopy for evidence of spinal cord injury.

Results: Seven of 20 (35%) experimental rat pups and 6 of 20 (30%) control rat pups were liveborn. All liveborn pups had severe paralysis of the hindlimbs and tail, so that functional differences between the 2 groups could not be detected. Histologic examination of 13 spinal cords at the site of surgical exposure showed that necrosis of neural tissue in 5 of 7 meconium-exposed rat pups was increased when compared with that observed in the 6 fetuses exposed to amniotic fluid without meconium. In general, inflammation was greater and repair processes appeared delayed in meconium-exposed rat pups.

Conclusions: Exposure of the spinal cord of fetal rats to amniotic fluid by surgically created myelomeningocele leads to severe functional impairment. Histologically recognizable necrosis of neural elements was increased in those animals that were exposed to diluted human meconium in the amniotic fluid. The results support the hypothesis that meconium may contribute to the pathophysiology of spinal cord injury observed in myelomeningocele.

INDEX WORDS: Myelomeningocele, rat, fetal surgery, meconium

Myelomeningocele is a congenital anomaly associated with significant lifelong disabilities that include paraplegia, hydrocephalus, sexual dysfunction, skeletal deformation, and bowel and bladder incontinence. These disabilities limit the average longevity of patients born with myelomeningocele to less than 40 years and are, in part, secondary to the severe spinal cord damage sustained before birth. According to the “two-hit hypothesis” the exposed neural elements in myelomeningocele primarily are the result of defective spinal cord development (myelodysplasia), whereas a secondary event resulting in erosion and necrosis of the exposed spinal cord results in progressive damage with gestational age. This hypothesis is supported by several observations suggesting that neural damage acquired during fetal life is increased by mechanical trauma or chemical toxicity of the amniotic fluid.

These findings have led some investigators to propose repair of myelomeningocele in utero as an alternative approach to minimize neural damage before birth. However, the specific mechanism of neural injury remains unknown. There is now evidence that in utero defecation may normally occur and that physiological passage of meconium increases late in gestation. This led
us to hypothesize that spinal cord damage in myelomeningocele may be related to contamination of the amniotic fluid by meconium. To test this hypothesis, we carried out an experimental study in the fetal rat model of myelomeningocele. Our results support the conclusion that prenatal exposure to meconium may contribute to damage of the neural elements in myelomeningocele.

MATERIALS AND METHODS

All experimental protocols were reviewed and approved by the Institutional Animal Care and Use Committee and followed guidelines set forth in the National Institutes of Health Guide for the Care and Use of Laboratory Animals.

Experimental Model

Female rats were obtained from a commercial breeder (225 g, Criffa, SA, Barcelona, Spain). Animals were mated, and females were checked daily for introtial plugging. The day of plugging was defined as gestational day 0 for time dating. The pregnant rats were kept in individual cages and maintained under 12 hours light/12 hours dark cycling conditions. On day 18½ (term, 22 days) anesthesia was induced by inhalation of sevoflorane (Sevorane, Abbot Laboratories) supplemented by intraperitoneal sodium pentobarbital (15 mg/kg intraperitoneal).

Open spinal dysraphism was created as described by Heffez et al. Briefly, a hysterotomy was performed by placement of a 6-0 nylon purse-string suture incorporating the amniotic membranes followed by a 3-mm incision placed within the pursestring overlying the back of the fetal rat. For fetal immobilization, the dorsal skin of the fetus was sutured to the border of the opened uterus with a 6-0 nylon. Under 25X magnification (Leica, Wild M651.MS-D, Herbrugg, Switzerland) the paraspinal muscles were dissected, and then a 2- to 3-level laminectomy was performed. The translucent dura was opened carefully under 40X magnification using a 26-gauge needle. The fetus then was returned to the uterus, the amniotic fluid volume was restored, and the exposed uterus, after closure, was bathed with ritrodrine for tocolysis (10 mg/mL; Pre-par, Solvay). The maternal laparotomy was closed in 2 layers with a continuous 4-0 silk suture. Fetuses were harvested by cesarean section at 21.5 days’ gestational age.

Experimental Groups

Twenty dams were operated on with 2 fetuses per dam manipulated. In each dam, the amniotic fluid volume was restored with saline (37°C) in one operated fetus (control group), whereas the other received a sterile 10% (by weight) meconium in saline suspension (experimental group). Meconium was obtained from human newborns less than 24 hours of postgestational age. Sterility of meconium was confirmed by parallel aerobic and anaerobic cultures.

Data Analysis

Liveborn pups were examined for neurologic deficits assessing postural reactions and pain perception by pricking with a needle (superficial pain) and by pinching with a hemostat (deep pain). The pups were photographed and then killed and fixed in 10% formalin. Cross sections of spine from both groups were taken at the level of the surgical site and at the levels proximal and distal to the operative site. Sections (10-μm thick) were stained with H&E and studied by light microscopy for evidence of spinal cord injury.

RESULTS

Operative Results

Seven of 20 (35%) experimental rat pups and 6 of 20 (30%) control rat pups were liveborn. All liveborn pups showed severe paralysis of the hindlimbs and tail, so that functional differences between the 2 groups could not be detected.

Histologic Study

In contrast to nonexposed segments of spinal cord (Fig 1), histologic examination of the 13 spinal cords at the site of the defect showed necrosis of neural tissue that was increased in the meconium-exposed rat pups (Fig 2) relative to the rats exposed to amniotic fluid without meconium (Fig 3). In general, inflammation was greater and repair processes appeared delayed in the meconium-exposed rat pups.

![Fig 1. Normal spinal cord. Sections of the spinal cord above (A) and below (B) the level at which the operative intervention was performed. A is at the thoracic level and B at the lumbar level. The dorsal aspect is above and the ventral below. Note the cellularity of the dorsal horns. (Both H&E, original magnification ×50.)](image-url)
DISCUSSION

Spinal cord damage is the primary cause of morbidity associated with myelomeningocele. The goal of perinatal management is to avoid or prevent the neural damage commonly observed in myelomeningocele. Improved understanding of the pathogenesis of neural injury is, therefore, required for rational perinatal strategies to evolve.

In this study, the observed fetal mortality rate was higher than reported in fetal rat models of other surgically created anomalies.\textsuperscript{12} The mortality rate was the same as reported by Heffez et al\textsuperscript{13,15} and is related to the duration (typically greater than 20 minutes) and extent of the surgical procedure required to expose the fetal rat spinal cord.

In our study, we found that spinal cord exposure induced severe functional impairment in both control and meconium-exposed groups. On histologic study, however, necrosis was increased in the meconium-exposed group suggesting that exposure to human meconium can increase the neurotoxicity of the amniotic fluid. Traditionally, prenatal meconium passage was thought to occur only during fetal distress. Recently, Ciftci et al\textsuperscript{11} challenged this idea showing that meconium-stained amniotic fluid reflects impaired clearance of amniotic fluid, which normally contains meconium from physiologic defecation.\textsuperscript{11} The presence of intestinal enzymes in the amniotic fluid of healthy fetuses and its absence in fetuses with intestinal obstruction also supports the idea that meconium passage during fetal life is a normal physiological event.\textsuperscript{13-15} Proximal intestinal dilatation in fetuses with colonic atresia or anorectal malformation also favors this theory.

Previous studies have shown that meconium increases amniotic fluid toxicity for other fetal tissues, such as intestine in experimental models of gastroschisis.\textsuperscript{16} In-
flammation and induction of necrosis by meconium also has been shown in lung with meconium aspiration syndrome. It therefore is not surprising that meconium increases damage of the exposed neural elements in this model.

According to our results, fetal spinal cord exposure leads to severe neural damage, which appears increased when human meconium is added to amniotic fluid. Thus, in contrast to other studies, we believe that chemical toxicity secondary to amniotic fluid exposure is relevant to the pathophysiology of neural damage observed in myelomeningocele. Our hypothesis that meconium might be involved on the pathophysiology of spinal cord damage in myelomeningocele is in agreement with those of previous studies suggesting that most of neural damage occurs late in gestation. As mentioned above, in utero defecation is gestational age dependent and becomes more prevalent the last weeks of gestation. Therefore, gastrointestinal waste products should be added to urea, ammonia and osmotic pressure gradients as potential causes of damage to the exposed spinal cord. However, our experimental protocol cannot exclude trauma as a contributing factor to spinal cord damage.

Although the results of in utero repair of myelomeningocele remain to be determined, our findings reinforce the theory that injury of the spinal cord is, at least in part, acquired during fetal life. Thus, the rationale of prenatal treatment remains promising for fetuses with myelomeningocele. There also may be a role for other prenatal strategies directed toward prevention of spinal cord damage and disability associated with myelomeningocele. For instance, the benefit of amnioinfusion, recently documented in experimental settings, may be attributable to withdrawal and dilution of the harmful chemical substances in amniotic fluid. The identification of meconium as a source of those substances provides new evidence supporting the potential benefit of prenatal treatment.

Exposure of the spinal cord of fetal rats to amniotic fluid by surgically created myelomeningocele leads to severe functional impairment. Histologically recognizable necrosis of neural elements was increased in those animals that were exposed to human meconium diluted in the amniotic fluid. The results suggest that meconium may contribute to the pathophysiology of spinal cord injury observed in myelomeningocele.

REFERENCES


