Current views on fetal surgical treatment of myelomeningocele — the Management of Myelomeningocele Study (MOMS) trial and Polish clinical experience

Jacek Zamłyński, Ewa Horzelska, Mateusz Zamłyński, Katarzyna Olszak-Wąsik, Leszek Nowak, Piotr Bodzek, Tomasz Horzelski, Rafał Bablok, Anita Olejek

Department of Gynecology, Obstetrics and Gynecologic Oncology in Bytom, Medical University of Silesia, Bytom, Poland

ABSTRACT

Myelomeningocele (MMC) constitutes the most common congenital defect of the central nervous system, with no satisfactory alternative to the postnatal treatment. Prenatal repair of MMC is aimed at protecting from a Chiari type II malformation. The main goal of fetal MMC repair is to improve the development and quality of life in children with a Chiari type II malformation. The Management of Myelomeningocele Study (MOMS), which was published in 2011, confirmed the effectiveness of prenatal surgery. In this paper, we compared the MOMS results with our own clinical experience. Owing to high effectiveness and significant improvement in the safety of the maternal-fetal surgery, prenatal MMC repair has become a new standard of treatment.

Key words: myelomeningocele, prenatal surgery, spina bifida, Chiari malformation

INTRODUCTION

Fetal myelomeningocele (fMMC) is one of the most common neural tube defects (NTDs). The results of the current postnatal treatment are not satisfactory. Owing to significant advances in ultrasound diagnostic imaging and primary perinatal care, fMMC may be identified as early as 16–18 weeks of gestation. Early detection of different types of NTDs enables to arrange the delivery, followed by postnatal surgical treatment, in a tertiary referral center. However, only prenatal surgical treatment, known as the in utero MMC repair (IUMR), protects the fetus from developing a Chiari II malformation (CM II).

Fetal MMC, a serious congenital neural tube defect, is also the most common defect of the central nervous system [1]. The incidence of MMC in the USA has been estimated at 2/10 000 live births, whereas in Poland MMC affects approximately 6/10 000 live births [2, 3].

The risk factors for the development of fMMC include antispasmodic, antihistamine and sulfonamide drug use in pregnancy, maternal diabetes and obesity, as well as familial NTD. Insufficient folic acid supplementation and the presence of the 677C > T polymorphism in the methylenetetrahydrofolate reductase gene are the key factors for developing NTD [4].

POSTNATAL MMC REPAIR AND FOLLOW-UP IN CHILDREN WITH CM II

Since the 1960s, postnatal MMC surgery technique of tensing the surrounding layers of fascia, muscles and skin with Limberg’s modification has remained the primary treatment option, performed in the first 48 hours of life. Figure 1 shows newborns after delivery scheduled for postnatal MMC repair. In utero evolution of CM II entails serious neural defects, which are generally caused by hydrocephaly (HC) and hindbrain herniation (HH) [5]. As a consequence of spinal cord exposure to the amniotic fluid, motor function of the lower extremities is lost, and bladder and bowel dysfunction are observed [6].
Implications depend on the level of spina bifida and spinal cord impairment. Defects lower down of the spinal cord are associated with better prognosis. A defect in cerebrospinal fluid circulation, which is caused by the displacement of the hindbrain structures into the spinal canal, results in HC in 80–90% of the children with fMMC. About 80% of patients with fMMC require postnatal treatment such as endoscopic third ventriculostomy (ETV) or insertion of the ventriculoperitoneal shunt (VPS), which allows excess CSF to drain to the peritoneal cavity [7]. VPS placement is associated with numerous complications concerning its obstruction or infection [8]. Early Polish results of postnatal treatment show high rate (85%) of progressive HC when temporary Rickham reservoir placement, followed by VPS insertion, was necessary [9].

The prognosis for children with MMC is grave. Despite intensive treatment, almost 14% of all children with MMC die during the first 5 years of life, and the mortality rate increases up to 35% when cerebellar dysfunction (secondary to CM II) develops [10]. About 70% of children reach the IQ level > 80, but only approximately half of them are self-reliant in adulthood [11].

**THE REASONS FOR PRENATAL FMMC SURGERY AND EARLY CLINICAL EXPERIENCES**

In 1990, Heffez et al., introduced the ‘two-hit’ theory, demonstrating a connection between neurological defects in children with MMC and the effect of two hits: neurulation in the early embryo life occurs, and then myelodysplasia is exaggerated in the mechanism of inflammation — the first hit (activated by the components of the amniotic fluid) — and mechanical destruction of spinal cord hitting the womb — the second hit [12].

Pathological and morphological studies on the spinal cord of fetuses (early and late pregnancy) and newborns with MMC reported progression of plaque defragmentation, spinal cord injury and nerve fiber destruction. These changes escalated with the duration of pregnancy [13].

The next experimental studies concerning IUMR in a sheep model disclosed that the loss of neurological function is then arrested. Especially, it became possible to maintain the motor function of the lower extremities, sustain normal cortex width and vegetative functions [14].

Finally, an experimental study of IUMR in human fetus showed reversibility and even regression of HC and HH, which are typical for CM II [15].

First IUMRs were reported in 1997–1998 at the Vanderbilt University Medical Center and at the Children’s Hospital of Philadelphia (CHOP) [16, 17]. Their early reports confirmed the theoretical and experimental assumption that in utero protection of the nerve fibers suppresses CM II evolution, resulting in HH reduction, stationary HC and maintaining motor function of the lower extremities [18, 19]. In the follow-up study, the VPS implantation rate was lower in the IUMR group as compared to the group treated postnatally (90% vs. 46%). The observed motor function of the lower extremities reflected motor functions typical for the injury of two spinal segments below the actual anatomical MMC level [19].

**THE MOMS RANDOMIZED STUDY RESULTS**

In 2011, the MOMS randomized study proved beyond any doubt the efficacy of prenatal surgery [20]. After enrollment and randomization of 183 out of the 200 recruited patients, the benefits of the prenatal surgery were confirmed and the study was completed. The rate of VPS placement in 1-year-old children who had undergone IUMR was significantly lower as compared to the postnatally treated patients (40% vs. 80%; relative risk, 0.8; 97.7% CI, 0.36–0.64; p < 0.001). Motor function of the legs (at 30 months of follow-up) was also better in the prenatally treated children: 42% of the prenatally and 21% of the postnatally treated children (relative risk 2.01; 95% CI, 1.16–3.48; p = 0.01) could walk independently. Among children in the IUMR group, the rate of patients without HH was decreased as compared to the postnatally treated group (36% vs. 4%; relative risk 0.67; 95% CI, 0.56–0.81, p < 0.001). Vegetative dysfunctions, especially concerning bladder, are the most burdensome problems in MMC patients. Unfortunately, so far it has not been proven that surgical treatment in utero could favorably influence the functioning of the urinary system [21].

The MOMS study pointed out that iatrogenic complications, which arise from the invasiveness of the procedure, belong to the main IUMR limitations. IUMR increases the risk of iatrogenic preterm premature rupture of the membranes (iPPROM) (46% vs. 4%; relative risk 6.15; CI, 1.16–3.48; p < 0.001), oligohydramnios (21% vs. 4%; relative risk 5.47; 95% CI, 1.16–18.4; p < 0.001) and inflammation-induced preterm labor (iPTL) (79% vs. 15%). As a result, higher incidence of children with low birth weight and respiratory distress syndrome is observed [20].

A five-year study of preschool children with MMC treated prenatally revealed that psychomotor development corresponded to their age in 83% of the cases. A 10-year observational study showed that behavioral functions were normal.

**Figure 1. fMMC of three newborns delivered at 36–37 weeks for a scheduled operation after birth**
in most patients and improvement of motor functions of legs was maintained in 90% of the affected children [23].

**FMMC REPAIR — POLISH CLINICAL EXPERIENCE**

In Poland, the first successful in utero FMMC repair was performed by Preis in 2004 [24]. Since 2005, 74 prenatal surgeries have been conducted at the Clinical Department of Gynecology, Obstetrics and Oncological Gynecology of the Silesian Medical University in Bytom, in cooperation with the pediatric surgeons from the Clinical Department of Pediatric Surgery in Katowice. In our first report on prenatal MMC closure in 10 patients, no increase in the lateral ventricular width was observed postoperatively [25]. Figure 2 demonstrates basic stages of IUMR. Since the surgery up to the delivery, the value remained within the qualification criterion of < 18 mm. In the next report, concerning 31 IUMR cases, control fetal MRI revealed that 13 out of 19 fetuses demonstrated HH reduction and no progression was noticed in 6 cases [26]. In our non-randomized study, we compared a group of 46 fetuses treated in utero to 47 patients with MMC postnatal closure [28]. In this observation up to 53 months of life, decreased VPS implantation rate the first group (27.8% vs. 80%; CI 0.35, 0.16–0.75; p < 0.008), lower HH (11% vs. 70%; CI 0.16, 0.04–0.06, p < 0.01), and better motor function were noticed in the first group. As in the MOMS study, we demonstrated that prenatal surgery is associated with the risk of iPTL, iPPROM and uterine wound dehiscence. As a consequence of iPPROM and iPTL, advanced neonatal procedures are required. After IUMR surgery, the cumulative mortality among newborns < 7 days of life reached 4.3% (2 newborns). In our last study of 71 fetuses with FMBC treated in utero, reduced HH and station ary HC were observed in 96.9% and 53.2% of the patients (36 cases), respectively. Only 15 newborns (21.1%) presented progressive HC (AD > 95 pctl.). 29.5% of the pregnancies were delivered before 30 weeks of gestation and 18.3% (13 cases) reached 37 weeks of gestation. Cumulative mortality was 4.2% (2 newborns), and was comparable to the MOMS result of 3% (2 out of 78 cases) and to the recently published study of VUMC — 5% (3 out of 43 cases) [30].

**A NEW STANDARD — FETAL MMC CLOSURE. POST-MOMS ERA**

The main goal of prenatal MMC closure is to improve psychomotor development and the quality of life in patients with Chiari malformation. The latest studies, known as ‘post-MOMS era’ and published after 2011, showed high safety of the maternal-fetal surgery [13]. Bennet et al. compared the results of a cohort study of 43 pregnancies after IUMR with the results of the randomized MOMS study results, and demonstrated that improved technical skills in the IUMR procedure lowered the rate of iPPROM (22% vs. 46%, p = 0.011) and chorioamniotic separation (0% vs. 26%, p < 0.001). Mean time of pregnancy after IUMR in the cohort study was extended up to 34.4 ± 6.6 vs. 34.1 ± 3.1 weeks of gestation. Also, the number of preterm births < 30 weeks was reduced to 2 (4%) vs. 10 (13%) (p = 0.08).

Satisfactory results of perinatal care in patients who underwent IUMR depend not only on the fetal-surgery team, but also on their understanding of the need to cooperate and achieve common goals [31]. Figure 3 presents an MMC site after prenatal surgery. The presented results have demonstrated high efficacy of IUMR. The studies from the last three decades have been finally accepted by the interna-
tional societies of obstetricians and gynecologists (ACOG, SOGC) for selected inclusion and exclusion criteria (Table 1) [32–34]. The Polish Gynecological Society published their recommendations about eligibility and the IUMR surgical procedure in 2006 [35].

EDUCATION OF THE PARENTS AND THE APPROACH AFTER MMC DIAGNOSIS

In cases when a NTD such as non-lethal fMMC is diagnosed, the parents should be informed about three possible treatment options:

- postnatal treatment performed within 24 hours of life,
- prenatal fMMC closure for patients who meet the eligibility criteria,
- pregnancy termination.

Medical consultation in cases when fMMC is diagnosed should not be directive. The decision of the prospective parents should be autonomous and based on the information presented during the consultation. Informed consent should be obtained.

If the parents decide to choose IUMR or if the consultation about fMMC is problematic, all necessary help and quick pre-qualification is provided also via e-mail: spinabifida@o2.pl, phone number: 32 7861540, or personal consultation at the Department of Gynecology, Obstetrics and Gynecologic Oncology in Bytom, Medical University of Silesia, Stefana Batorego 15, 41–902 Bytom, Poland. Patients with diagnosed fMMC should not be referred for pregnancy termination without being fully informed about all possible treatment options.

REFERENCES


Table 1. Criteria for fetal surgery

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
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<tbody>
<tr>
<td>Gestational age: 20–25 weeks</td>
<td>Fetal anomaly unrelated to myelomeningocele</td>
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<td>Maternal age ≥ 18 years</td>
<td>Pregestational diabetes</td>
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<td>Width of lateral ventricle of the brain AD &lt; 18 mm</td>
<td>Severe kyphosis</td>
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<td>Symptoms CM II &gt; 0° HH</td>
<td>Isthmo-cervical insufficiency</td>
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<tr>
<td>Myelomeningocele with the upper boundary located between ≤ Th12 — ≥ S1</td>
<td>Placenta previa</td>
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<td>Bilateral leg movement</td>
<td>BMI ≥ 35</td>
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<tr>
<td>Singleton pregnancy</td>
<td>TORCH, HIV, HCV, HBS infection</td>
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<tr>
<td>Normal karyotype</td>
<td>Anatomical uterine anomaly</td>
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Figure 3. Result of prenatal MMC surgery of three newborns at 36–37 weeks of gestation, operated at the Department of Gynecology, Obstetrics and Gynecologic Oncology in Bytom, Medical University of Silesia (healed wound, no cerebrospinal fluid leak)


