Case Report

Surgical management of caudal duplication syndrome: A rare entity with a centered approach on quality of life

Adilson de Oliveira, Clarissa Nascimento, Diego Ramos, Hamilton Matushita

Department of Neurosurgery, Faculty of Medicine, São Paulo University, São Paulo, Brazil.

E-mail: *Adilson de Oliveira - adilsonvalmont@gmail.com; Clarissa Nascimento - clangnnnc@gmail.com; Diego Ramos - diego.angelis@hc.fm.usp.br; Hamilton Matushita - matushita@uol.com

INTRODUCTION

The rare caudal duplication syndrome (CDS) includes a broad spectrum of anomalies involving the gastrointestinal, genitourinary, neurological, and vertebral skeletal systems. This entity was first described by Dominguez et al., in 1993; the prevalence is <1/100,000 births. Pang et al.,[5,6] it described a severe form of a type 1 split cord malformation but did not report any patients with such extensive duplication of the bony elements. The CDS is attributed to the incomplete separation of monovular twins, which helps explain, as in this case, duplication of the spine/spinal cord, hindgut (ileum/colon), and lower urinary tract.[1,2] In this similar case, the surgical approach must be tailored to the individual using a multidisciplinary team.

ABSTRACT

Background: Caudal duplication syndrome (CDS) is associated with complete/partial duplication of the spine, spinal cord, duplication/malformations of other caudal structures, and often neurological dysfunction.

Case Description: A 9-month-old with the prenatal diagnosis of multiple fetal malformations was born through cesarean section at 39 weeks. Computed tomography and magnetic resonance imaging revealed a series of abnormalities: intramedullary cyst at the L4 level with tethering of the conus with a lipoma, a duplication of the sacrum, duplication of the intestines, a ventricular septal defect, bladder duplication, four kidneys, two fully formed functional penises, two anuses (imperforate on the left), and a paramedian ossified appendage suggestive of a malformed leg. The child, at an 8 months of age, underwent a laminectomy at the L4 level for resection of the conus lipoma, release of the tethered cord, and resection of the intramedullary cyst.

Conclusion: Spine/spinal cord duplication syndromes are typically associated with spine/spinal cord abnormalities and are often associated with moderate/severe neurological deficits. Additional malformations associated with the CDS include duplication of vascular structures/organisms (e.g., bladder and distal gastrointestinal tract). Here, we focused on both the patient's quality of life and the multidisciplinary approach to managing the duplication syndrome associated with multiple system malformations (e.g., neurological/spinal, bladder, and gastrointestinal, others).

Keywords: Congenital malformation, Pediatric neurosurgery: Caudal duplication, Quality of life, Spine duplication
CASE REPORT

Documentation of anomalies at birth

A 9 months old without a history of consanguinity or familial genetic diseases had the prenatal diagnosis of multiple fetal malformations established utilizing ultrasonography. Cesarean section delivery at 39 weeks prompted the performance of repeat ultrasonography followed by computed tomography and magnetic resonance imaging. These studies documented the following anomalies; intramedullary cyst at the L4 level with a tethered cord/lipoma, duplication of the sacrum, intestine, kidneys (e.g., 4 of these), bladder, two functional penises and anuses (one imperforate on the left) with enterovesical fistulas, and several cardiac defects (e.g., ventricular septal defect, interventricular communication, atrial septal defect, and persistent ductus arteriosus). In addition, there was a paramedian ossified appendage, suggestive of a malformed leg [Figures 1-3].

Surgery

A laparotomy was performed on the 5th day of life due to an acute abdomen; an enterovesical fistula was repaired. At 8 months, the patient underwent a laminectomy at L4 for untethering and resection of the lipoma and intramedullary cyst involving the conus. In the future, a multidisciplinary team will perform corrective surgery of the bones, genitalia, and urinary tracts (e.g., when child reaches the ages of 2–3 [presently 20 months old]). This will warrant; pelvic fusion, malformed leg amputation, amputation of one of the penises with urological and plastic surgery, and correction of the abdominal wall herniation.

DISCUSSION

The rare CDS, first described by Dominguez et al., in 1993, is defined as the association between malformations and duplications involving the gastrointestinal, genitourinary, and spine/spinal cord (e.g., neural tube defects).\(^1\)

In our literature review, we did not find previous reports of patients who remained neurologically intact. Here, we used a multidisciplinary approach to ensure normal function in this 20-month-old patient with CDS. This first warranted emergently treating an enterovesical fistula at 5 days of age (acute abdomen) followed at 8 months by an L4 laminectomy for the treatment of a tethered cord/lipoma/intramedullary cyst. In other cases, those treated too aggressively sustained poor outcomes (e.g., including an adult with original normal neurological function but postoperative deterioration).\(^3\)

---

Figure 1: The child at 9 months (a) anterior, (b) posterior. Thin arrow – phallus; wide arrow – legs; discontinue thin arrow normal – anus; discontinue wide arrow – imperforate anus; asterisk – abdominal hernia; two asterisks – lipoma.

Figure 2: Computed tomography scan coronal view (a) 3D bone reconstruction, (b) soft tissue. Thin arrow – duplication of pelvis; wide arrow – absence of pubic symphysis.

Figure 3: Lumbosacral magnetic resonance imaging with fat suppression (a) sequence weight in T1B, (b) sequence weight in T2. Thin arrow – spinal cord duplication; wide arrow – intramedullary cyst; asterisk – tethered spinal cord; two asterisks – medullary cone lipoma.
CONCLUSION

Here, we report on a 9 months old with a CDS syndrome characterized by the duplication of multiple organ systems and neurological spine/spinal cord malformations. The acute abdomen due to an enterovesical fistula was treated 5 days after birth, and the patient underwent spine surgery at 9 months of age consisting of an L4 laminectomy to treat a tethered cord/lipoma and intramedullary cyst. Future operations are planned to address multiple additional duplications (e.g., at ages of 2–3).

Declaration of patient consent

The authors certify that they have obtained all appropriate patient consent forms. In the form, those responsible for the patient have given his/her/their consent for his/her/their images and other clinical information to be reported in the journal. Those responsible for the patient understand that their names and initials will not be published and due efforts will be made to conceal their identity, but anonymity cannot be guaranteed.

Financial support and sponsorship

Nil.

Conflicts of interest

There are no conflicts of interest.

REFERENCES

1. Dominguez R, Rott J, Castillo M, Pittaluga RR, Corriere JN Jr. Caudal duplication syndrome. Am J Dis Child 1993;147:1048-52.
2. Harris J, Blackwood B, Pillai S, Kanard R. Caudal duplication: Management of a rare congenital condition. Am Surg 2016;82:e227-9.
3. Hu T, Browning T, Bishop K. Caudal duplication syndrome: Imaging evaluation of a rare entity in an adult patient. Radiol Case Rep 2016;11:11-5.
4. Mei JY, Nguyen MT, Raz S. Female caudal duplication syndrome: A surgical case report with 10-year follow-up and review of the literature. Female Pelvic Med Reconstr Surg 2018;24:e16-20.
5. Pang D, Dias MS, Ahab-Barmada M. Split cord malformation: Part I: A unified theory of embryogenesis for double spinal cord malformations. Neurosurgery 1992;31:451-80.
6. Pang D. Split cord malformation: Part II: Clinical syndrome. Neurosurgery 1992;31:481-500.
7. Samuk I, Levitt M, Dlugy E, Kravarusic D, Ben-Meir D, Rajz G, et al. Caudal duplication syndrome: The vital role of a multidisciplinary approach and staged correction. Euro J Pediatr Surg Rep 2016;4:1-5.