Complete intraventricular migration of the ventriculoperitoneal shunt

Elif Basaran Gundogdu, Ufuk Ozsoy, Yusuf Tuzun

Department of Neurosurgery, University of Health Sciences, Bursa Yuksek Ihtisas Training and Research Hospital, Bursa, Turkey

ABSTRACT

Intracranial migration of the ventriculoperitoneal shunt is an extremely rare situation. A 2-month-old infant who had undergone ventriculoperitoneal shunt for hydrocephalus in the neonatal period presented with feeding problems and increasing head circumference. There was intracranial migration on cranial computed tomography and shunt survey images. Endoscopically the shunt was removed and a new shunt was applied. The patient was discharged with cure. This extremely rare shunt complication and its surgery both have serious complications. Therefore, the surgical technique, proper fixation of the shunt and the use of the right shunting material are significantly important.

Keywords: Shunt complication, intraventricular migration, endoscopic procedures

Introduction

The frequency of intracranial migration of the ventriculoperitoneal shunt is reported to be 0.1-0.4% in the literature [1]. Direction of the migration is determined by the pressure gradient between the cranial and the peritoneal cavities. The case being reported is an example for the intracranial migration of the ventriculoperitoneal shunt which is extremely rare.

Case Presentation

A 2-month-old infant who had undergone ventriculoperitoneal shunt for hydrocephalus in the neonatal period presented with feeding difficulty and increasing head circumference. She was hospitalized. The shunt catheter was not palpable on the cranium, the cervical region or the anterior chest wall. On cranial computed tomography (CT), the shunt was completely in the ventricle and acute hydrocephalus was seen. Also the parenchyma was extremely thinned. On shunt survey images, there was no continuity of the shunt catheter, so an external ventricular drain (EVD) catheter was introduced immediately and a clear cerebrospinal fluid (CSF) leak was observed (Figure 1 a, b, c). The patient was reoperated since there was no bacterial growth in the CSF culture and there was no fever. The EVD catheter
was removed, and the new shunt catheter was inserted endoscopically. After inserting the new shunt catheter, the old catheter was removed from the right frontal region (Figure 1d, e, f). The patient was discharged with cure.

Discussion

Complications of the ventriculoperitoneal shunt are frequently reported for patients with hydrocephalus in the literature. Although migration of the distal edge of the ventriculoperitoneal shunt is common, intracranial migration is a rare complication and it has been reported in 0.1-0.4% of the cases [1]. Despite continuously developing shunting technology, 33% of the patients who had undergone shunt experience dysfunction of the shunt at the end of the first year, 50% after 2 years and 70% after 10 years [2].

In migration of cranial edge of the shunt, lower pressure and mechanical factor providing continuation of the shunt catheter within cranium have been considered as responsible [3, 4]. In other words, if the cranial pressure is high, the catheter will move towards the abdomen, and if the abdominal pressure is high, the catheter will move towards the cranium [3, 5]. As Ceran-Rous et al. [6] have reported, underlying diseases (porencephaly), dynamic causes such as abdominal peristaltic movements, dynamic translocation factors such as cervical movements, dynamic attraction factors such as increased CSF reabsorption may cause migration of the shunt. Flexion and extension movements of the neck in childhood can play a role in upward migration of the shunt catheter. In children, since the distance between the ventricular and the peritoneal edges of the catheter is shorter than that in adults, proximal migration more readily occurs. Having seizures and constipation play roles in proximal migration of the shunt catheter [4, 7]. Yılmaz et al. [8] suggested that, detachment of the shunt from its attachment sites due to development of

Figure 1. Preoperative (a, b, c) and postoperative (d, e, f) x-rays images of ventriculoperitoneal shunt.
the child, as well as abdominal peristaltic movements, cervical flexion and extension movements, and the history of seizures might be responsible for migration of the ventricular catheter.

The second most common cause of shunt migration in pediatric patients is interruption and detachment of the shunt. Predisposing factors are shunt design, the material used and the surgical technique. Any fixation point can cause interruption by causing tension of the shunt. In addition, immune reaction can cause calcification and dissolution of the shunt material [9, 10]. During surgical procedure, a loose binding or use of absorbable suture material on connection part can lead to detachment in the shunt system by causing tension. Metal instruments can cause small erosions or even full-thickness tears leading to detachments. Risk for shunt detachment occurs only if a connection part is present in the shunt system. For one-piece shunts there is no such problem [11, 12]. Within shunt material, a catheter whose distal tube is stuck with valve and other wall is coated with pure silicone (to prevent calcification) is the best choice when ventricular edges are bound appropriately [13].

When complete intraventricular shunt migration was searched in the PubMed, 10 cases were encountered. Although this situation is extremely uncommon, both the case itself and the surgical procedure are quite risky. In cases in which the shunt is migrated towards the ventricle; anemia, sepsis and hydrocephalus cause cortical atrophy [14]. Situations such as presence of adhesions to the choroid plexus can lead to intraventricular hemorrhage while the shunt is being extracted [15]. To prevent this, avoiding burr holes and wide dural incisions, and attention to proper fixation of the shunt are critical [16, 17].

**Conclusion**

Complete intraventricular shunt migration is an extremely rare case. The situation itself and its surgery both have extremely severe complications. To prevent this, the surgical technique, proper fixation of the shunt and the use of the right shunting material are significantly important.

**Informed consent**

Written informed consent and photography release forms were obtained from the parents of the patient for the publication of this case report.

**Conflict of interest**

The authors declared that there are no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

**References**


