Case Report

Decompressive craniectomy and early cranioplasty in a 15-year-old boy with *N. meningitidis* meningitis

Julius Hoehne, Monika Friedrich, Alexander Brawanski, Michael Melter, Karl-Michael Schebesch

Departments of Neurosurgery, Pediatrics, University Medical Center Regensburg, Franz-Josef-Strauss Allee 11, 93053 Regensburg, Germany

E-mail: *Julius Hoehne - julius.hoehne@ukr.de; Monika Friedrich - monika.friedrich@ukr.de; Alexander Brawanski - alexander.brawanski@ukr.de; Michael Melter - michael.melter@ukr.de; Karl-Michael Schebesch - karl-michael.schebesch@ukr.de

*Corresponding author

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Abstract

**Background:** Intracranial hypertension is a well-known life-threatening complication of bacterial meningitis. Investigations on decompressive craniectomy after failure of conservative management are scarce, but this surgical treatment should be considered and performed expeditiously, as it lowers the intracranial pressure and improves brain tissue oxygenation. Early cranioplasty can further aid the rehabilitation.

**Case Description:** A 15-year-old boy was admitted to our emergency department because of sudden onset of neurologic decline and consecutive loss of consciousness. Clinical examination and imaging showed elevated intracranial pressure, leading to the suspected diagnosis of meningitis. Intracranial pressure monitoring was installed, but the initiated conservative management failed. Finally, the patient underwent bilateral decompressive craniectomy. The microbiological test showed growth of *Neisseria meningitidis*. After full neurologic recovery, cranioplasty with two CAD/CAM titanium implants was conducted successfully.

**Conclusions:** This unique report shows that decompressive craniotomy with duroplasty may be a crucial therapeutic approach in bacterial meningitis with refractory increased intracranial pressure and brainstem compression. Early cranioplasty with a patient-specific implant allowed the early and full reintegration of the patient.

**Key Words:** Cranioplasty, computer-aided design/computer-aided manufacturing, encephalitis, intracranial pressure, meningococcal disease, meningitis, *Neisseria meningitidis*, W/Y-135

CASE REPORT

A 15-year-old Turkish boy was admitted to the emergency department after loss of consciousness. 3 days prior to admission, he had experienced fever and ague with a gradual decline of his general condition. When presenting to his primary care physician, he was treated with nonsteroidal antiinflammatory drugs (NSAIDs) and antitussives for a suspected viral infection. Because his condition had not improved and he had developed nausea and vomiting, the patient presented to an emergency physician the night prior to hospital admission. This time, paracetamol and antiemetics were administered. During the day he was tired, disoriented, and somnolent. The next morning, he was found unconscious and was subsequently admitted to the emergency department. No clinical signs of epileptic seizures could be detected.
On admission, the patient was not adequately responsive, showing a Glasgow Coma Score of 9 including neck stiffness. The body core temperature was 36.5°C. Pupils were equal, round, as well as reactive to light and accommodation. No additional neurological abnormality was present.

The initial computed tomography (CT)-scan showed generalized brain edema [Figure 1]. Consequently, no lumbar puncture was conducted.

At the neurological department, the patient was initially treated with corticosteroids (10 mg dexamethasone), antibiotics (2 g ceftriaxone and 400 mg ampicillin), and antiviral medication (750 mg aciclovir) because of suspected meningitis. Relatives and people in close contact with the patient were treated accordingly. An intracranial pressure (ICP) probe was implanted for ICP monitoring. Despite maximal osmotic management, the patient developed intractable ICP with values up to 36 mmHg. Because the patient failed to respond to maximal conservative treatment, the team of pediatrics, neurologists, and neurosurgeons decided to conduct bilateral craniectomy and durotomy to control the extensively elevated ICP.

Immediately after decompressive surgery, the ICP decreased significantly but increased again few hours after the first surgery, while the right pupil became dilated consecutively. The immediate CT-scan showed an epidural hematoma and subgaleal blood collection. The hemorrhage was evacuated instantly [Figures 2 and 3]. During the following days, the ICP remained stable within normal limits, and the ICP-probe was removed. Radiologic follow-up by means of magnetic resonance imaging (MRI) on day 7 after ictus showed diffusion impairment in parts of the insular region and around the temporal horn. On day 13 after surgery, further radiological follow-up by means of a CT scan confirmed bilateral hygroma that were drained by bilateral subdural drainages. A lumbar drainage was implanted for 5 days. The electroencephalogram showed no signs of epilepsy; brainstem auditory evoked potentials could not be obtained on the right, whereas somatosensory evoked potentials were positive.

Despite the initial antibiotic and antiviral treatment without prior lumbar puncture, microbial testing of cerebrospinal fluid at a later stage showed Neisseria meningitidis type W/Y-135; the polymerase chain reaction (PCR) test for herpes simplex (HS) virus type 1/2 was negative. Uncomplicated pneumonia was treated successfully with antibiotics but prolonged the intubation period. Two weeks after admission, inflammatory hematological parameters had improved significantly, and the boy was extubated. Rapidly, the patient’s general condition improved, hence he was discharged on day 45 without any neurologic deficit and soon afterwards returned to school.

After 4 weeks, patient specific CAD/CAM titanium cranioplasties (CL instruments, Attendorn, Germany) were implanted bilaterally. Wound healing was uneventful and postoperative imaging showed accurate fitting without relevant artifacts [Figure 4].

**DISCUSSION**

Basically, permanent neurological deterioration due to meningitis is reported for up to 16% of affected children with neurologic sequelae; this rate refers to survivors in developed countries, irrespective of maximal conservative treatment. Neurological and neuropsychological sequelae have been mainly reported in cases of bacterial meningitis. In patients with bacterial meningitis, brain edema has been observed in up to 24% of patients. Monitoring of ICP has been proposed by Barnett et al. as a useful adjunct for therapy and provided indication for prognosis in an outcome analysis of 10 patients with encephalitis. Direct intraparenchymal ICP monitoring is accurate, low risk, and routine procedure in pediatric...
Surgical Neurology International 2015, 6:58

neurosurgery that is superior to radiological evaluation alone. Only anecdotal evidence is available for decompressive craniectomy as a treatment option for bacterial meningitis with intractable ICP. A 28-year-old female with bacterial N. meningitidis infection underwent decompressive craniectomy, after diagnostic lumbar puncture lead to temporal herniation and raised ICP could not be managed conservatively. A 17-year-old boy with meningitis after exacerbated pansinusitis developed brain edema with consecutive unilateral anisocoria and underwent bilateral decompressive craniectomy. Seven weeks later, the autologous bone was successfully reimplanted without any neurological deficit. A 34-year-old female suffered from Streptococcus pneumonia meningitis and developed intractable intracranial hypertension exceeding 30 mmHg with a GCS of 3. Bilateral decompression was performed and the patient was at discharge, apart from bilateral hearing loss, fully independent. In a further report two patients with S. pneumonia meningitis underwent surgical intervention. First, a 35-year-old male with chronic pansinusitis neurologically deteriorated. Subsequently late left hemispheric craniectomy was performed but the patient died due to the sequelae of territorial ischemia. The other patient, a 19-year-old male, suffering from meningitis as a result of prolonged and exacerbated maxillary sinusitis, was treated successfully with bifrontal craniectomy and recovered without any deficit. Finally, in a 56-year-old female with bacterial meningitis, treatment with bilateral decompressive craniectomy and guidance by cerebral microdialysis and brain tissue oxygen pressure measurement resulted in a favorable outcome.

Most other patients described in the literature had viral infections. The treatment of elevated ICP – refractory to conservative and osmotic management – is not new. Supporting evidence mainly stems from studies on ischemic stroke and traumatic brain injuries, in which early decompressive craniectomy was beneficial for survival and neurological outcome. Contrary to ischemic brain injury, decompressive surgery in traumatic brain injury is still controversial. The titanium implant offers complete restoration of cosmesis and good defect closure to prevent neurologic sequelae. Early cranioplasty in children by means of the CAD/CAM technique has not been reported before. In our patient, the interval between craniectomy and cranioplasty was 4 weeks, whereas longer waiting periods have been proposed. The recommended time span before implantation ranges from as early as 2 weeks to more than 2 months.

CONCLUSION

Decompressive craniectomy has been shown beneficial in certain conditions other than severe traumatic brain injury resulting in severe refractory ICP. In this unique case of a 15-year-old boy suffering from bacterial meningitis with N. meningitidis, bilateral decompressive craniectomy was conducted as last resort because of generalized brain edema and massively increased ICP that was refractory to any conservative and osmotic treatment. Bilateral preformed titanium cranioplasty implants were used at an early stage, and neurological recovery was excellent.

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