Normal Pressure Hydrocephalus Presentation with a Large Pseudomeningocele

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Abstract

Background and Importance: This interesting case is about the presence of normal pressure hydrocephalus and a large pseudomeningocele at the same time after 13 months of posterior fossa surgery. Although the occurrence of a pseudomeningocele following posterior fossa surgery is not so rare, such a late large pseudomeningocele development with signs and symptoms of NPH after 13 months of surgery is rather peculiar.

Case Presentation: A 59-year-old man referred to our clinic for a cerebellopontine angle tumor. The patient presented with right hearing loss and mild facial palsy. We operated the patient with retrosigmoid approach, and we discharged him after five days. After about 13 months, the patient referred to our center again with complaint of progressive bulging of previous surgical region and gait apraxia, urinary incontinence and dementia. A large pseudomeningocele in the site of previous surgery was seen.

Conclusion: The authors presented a case report on late development (13 months) of a huge pseudomeningocele following vestibular schwannoma surgery with signs and symptoms of normal pressure hydrocephalus. Such a late big pseudomeningocele presentation with signs and symptoms of normal pressure hydrocephalus after posterior fossa surgery is peculiar.

Keywords: Hydrocephalus; Pseudomeningocele; Intracranial; Pressure

Background and Importance

We presented a case with normal pressure hydrocephalus (NPH) and a large pseudomeningocele at the same time after 13 months of posterior fossa surgery. A pseudomeningocele may develop after posterior fossa surgery, but the presence of such a late large pseudomeningocele with signs and symptoms of NPH after 13 months of surgery is rather peculiar.

Case Presentation

A 59-year-old man referred to our clinic with a cerebellopontine angle (CPA) tumor. On admission, the patient gave informed written consent to participate in the study. He presented with right-side hearing loss and mild facial palsy. Brain magnetic resonance imaging (MRI) showed a large right CPA (Figure 1). We operated the patient with retrosigmoid approach, and the pathology revealed a schwannoma. The patient was discharged with good general condition five days after operation. Thirteen months later, the patient referred again complaining of progressive bulging of previous surgical field from about two months before that, and gait apraxia, urinary incontinence and dementia from about 20 days before that.

Neurological examination revealed Mini-Mental State Examination (MMSE) score 23/30. Motor system examination showed normal tone and power in all four limbs. Deep tendon jerks were normal. Gait assessment revealed wide-based stance. His steps were slow, short, and shuffling. His feet could not be raised much above the ground as if they were glued but his visual
field and acuity were normal. There was no papilledema. A large pseudomeningocele in the site of previous surgery was seen (Figure 2).

Considering this information, the patient was investigated. His hematological and biochemical parameters were normal. Brain imaging showed fluid collection in the region of the right parietal-occipital scalp and symmetric dilatation of all the ventricles without periventricular edema. There were no significant white matter abnormalities. These findings were seen on T1, T2 weighted and T2 fluid-attenuated inversion recovery (FLAIR) images (Figure 3). Finally, NPH was diagnosed, and the cerebrospinal fluid (CSF) tap test (TT) was done; 50 milliliters (ml) of CSF was drained. The opening pressure was 180 millimeters (mm) of water, and cyto-biochemical examination of CSF showed proteins 21 mg/dl, sugar 70 mg/dl, and cells 0. Pre- and post-test assessments of gait and cognitive functions were performed which revealed remarkable improvement in patient's gait. With such a significant improvement with CSF TT, the patient underwent ventriculoperitoneal shunt surgery and afterward there was almost complete recovery of both cognitive and gait disturbance. The patient was discharged at forth postoperative day. After three weeks, he was re-examined, and the shunt was performing well. Neurological examination revealed MMSE score as 28/30, and he had stable improved cognitive and gait status. The pseudomeningocele disappeared thoroughly (Figures 4,5). He is still under follow-up to search shunt functions and its associated complications.

Discussion

A pseudomeningocele is an abnormal body of cerebrospinal fluid communicating with the CSF space in which the fluid has no membrane around, yet it exists in a cavity of soft tissues [1]. However, the pathophysiology of pseudomeningocele raises ambiguities for some researchers and scientists who consider the potential causes of pseudomeningocele in the subarachnoid space such as intradural defects. On the contrary, absorbing CSF along with other problems including subarachnoid scarring and hydrocephalus may also play a vital role in some others' opinion [2]. The time of pseudomeningocele occurrence was reported to range from 5 to 115 days after surgery (mean: 18.2 days) [3].

The symptoms constellation of NPH include gait apraxia, urinary disturbance and memory impairment. According to reports, gait disturbance is the most common symptom occurring earlier than other complications [4]. Much more research needs to be performed to discover the aspects of pathophysiology of NPH which is assumed to be related to decreased absorption of cerebrospinal fluid rather than over-production [5]. There are two types of NPH: Idiopathic and secondary. The latter may result from brain trauma and tumors, subarachnoid hemorrhage, infection of the central nervous system, or occur after cranial surgery [6]. Relkin and his colleagues in their study suggested that NPH had a slow onset in someone over 40 years old with worsening symptoms occurring for three to six months without other neurological presentations. Brain imaging should show
ventriculomegaly without periventricular white matter change and evidence of parenchymal cerebral atrophy. For a strong diagnosis, a patient should have the gait disturbance and at least two of nine following symptoms: decreased step height and length, a wide-based gait, slowed walking, truncal swaying during walking, toes deviating outward, spontaneous retropulsion, difficulty turning, and difficulty keeping balance while walking [7-10].

According to references, there are very important factors related to intracranial pressure (ICP) namely the volume of intracranial fluid, the elastance, the contribution of the atmosphere, and the orientation of the craniospinal axis relative to the gravitational vector [11-13].

According to Tanna and colleagues' study, cranial CSF volume is about 164.5 mL, with a range of 62.2 to 267 mL. Ventricular volume also varies considerably from 7.49 to 70.5 mL, with a mean of 31.9 mL [14]. Elastance is defined by the pressure change per unit of volume change. We cannot consider this relationship here absolutely linear in all volumes with the same status in all physiologic conditions. Compliance is considered as the opposite of elastance. Two main ways of compensation include expansion or loss of volume when volume is added to the system. In a physiologic sense, this can occur either by distention of the spinal dura mater or by displacement of CSF and blood [13,15,16].

According to prior studies and references, there was linear relationship between intracranial volume and ICP which can be better understood by showing the response of ICP to volume added into the neural axis (Figure 6.A). Ryder et al.
explained that this relationship described hyperbolic curve in normal adults. According to prior studies, in spatial compensation as volume augments the intracranial pressure is minimally influenced. Still, the increased pressure changes per unit volume occurred with declining compliance; this portion is named the spatial decompensation [13,15-17]. As explained in the exemplified curve with compliance diminishing, the slope had a swift growth during spatial decompensation [13,17-19]. According to references, placing intracranial pressure logarithmically against volume is an alternative method, which gives a straight line. Its slope is the pressure-volume index (PVI), or the calculated volume in milliliters needed to raise ICP by a factor of 10 (Figure 6.B). In normal adults, the index ranges from 25 to 30 mL. Then compliance decreases through pathologic processes (decreased index). Finally, a few changes volume would cause significant alterations in pressure [13,15,17].

According to above, we suggest a hypothesis that could explain this case. The pseudomeningocele and the hydrocephalus result from the same hydrodynamic mechanisms. Progressive hydrocephalus can induce both ventricular and extradural collection dilatation. It seems that the surgical procedure to remove the vestibular schwannoma weakens the dural layer creating a "locus minori resistentiae" through which the CSF finds its way to the soft tissues at the base of the neck, forming the pseudomeningocele. We assume that the initial problem was a chronic hydrocephalus but the important point in this patient was the absence of intracranial hypertension signs and symptoms including papilledema, periventricular edema, and high opening pressure in tap test. It seems that the large pseudomeningocele acted as a reservoir to increase the system compliance, therefore prolongation of spatial compensation phase, and as a result, manifestation of a chronic hydrocephalus in delayed mode as NPH after 13 months of surgery without signs and symptoms of elevated ICP occurred. Unfortunately the patient did not refer to us for follow-up visits. More images in follow-up period could be helpful in understanding how the hydrocephalus developed over time.

Conclusion
The authors of this study presented a case report on late development (13 months) of a huge pseudomeningocele following vestibular schwannoma surgery with symptoms of NPH. Such a late big pseudomeningocele presentation with signs and symptoms of NPH after posterior fossa surgery can be considered peculiar.

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Conflicts of interest
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References