PEARLS

- Isolated elevated intracranial pressure (ICP) commonly presents with false localizing signs.
- If elevated ICP remains unrecognized, it may lead to permanent neurologic deficits.
- Recognition of uncommon presentations, as well as radiologic signs of elevated ICP, is of paramount importance as prompt intervention minimizes morbidity.

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- Auditory neuropathy is an uncommon presentation of elevated ICP, possibly due to stretch of the vestibulocochlear nerve from the increased ICP.
- Petrous apex cephaloceles and meningoceles are uncommon radiologic features of elevated ICP.
- Ischemic central retinal vein occlusion is a rare ophthalmologic manifestation of elevated ICP that may cause irreversible vision loss if the increased ICP remains unrecognized.

Isolated elevated intracranial pressure (ICP) classically presents with headache and papilledema without focal neurologic signs.1 Cranial nerve palsies, particularly abducens nerve palsy, have been described as false localizing signs.2,3 However, if elevated ICP remains unrecognized, it may lead to permanent neurologic deficits. We report a rare presentation of chronic idiopathic ICP elevation with bilateral sensorineural hearing loss (SNHL) and central retinal vein occlusions (CRVOs).

CASE PRESENTATION

A 49-year-old Macedonian man, with a remote history of a motor vehicle accident that resulted in a leg amputation, presented to our emergency room with bilateral blindness and deafness.

Nine months earlier, he had presented to a local hospital in Macedonia with severe headache, blurry vision, and diffuse weakness. Per records, he had received a lumbar puncture (LP) with no recorded opening pressure and normal CSF findings. He was then empirically treated with plasma exchange for polyradiculoneuropathy based on an EMG/nerve conduction study.

At his presentation to our facility, he had diminished muscle strength in his bilateral upper extremities, left more than right. Deep tendon reflexes were absent on the left upper extremity and diminished elsewhere. His visual acuity was no light perception bilaterally with fixed dilated pupils. Funduscopic examination showed bilateral optic disc atrophy with mildly elevated margins consistent with atrophic papilledema and significant retinal hemorrhages consistent with CRVO. Fluorescein angiography of both eyes was also suggestive of CRVO (figure, A and B). Examination was also remarkable for hearing loss of the right ear and severely diminished hearing ability of the left ear. Audiogram showed severe bilateral low-frequency SNHL, right more than left. Auditory brainstem response was remarkable for inversion of the waveforms in response to a change in stimulus polarity, consistent with bilateral auditory neuropathies. Initial MRI of the brain was negative for any mass lesion or obstructive hydrocephalus. MRI of the internal auditory canals (IACs), however, was significant for petrous apex cephaloceles and protrusion of the meningoceles into the bilateral IACs (figure, B and C). Severe engorgement of bilateral optic veins with distention of perioptic subarachnoid space and atrophy of the optic nerves bilaterally was apparent on the MRI of the orbits (figure, E). Magnetic resonance venogram showed no evidence of venous sinus thrombosis or focal stenosis, although all venous sinuses were diffusely collapsed (figure, F). Subsequently, LP was performed with an opening CSF pressure of 45 cm H2O. The details of CSF constituents are noted in table e-1 on the Neurology® Web site at Neurology.org. CSF bacterial and fungal cultures, extensive workup for vasculitides, and HIV antibody were also negative. EMG/nerve conduction study showed multilevel cervical radiculopathy with no evidence of polineuropathy.
No significant clinical improvement was observed after a large-volume LP and administration of high-dose acetazolamide. Therefore, a ventriculoperitoneal shunt was inserted. Postoperatively, a cerebral angiogram was performed with resolution of the diffuse venous collapse on the venous phase (figure, G and H). The acuity in the left ear improved, which was also evident on repeat audiogram at 1-month follow-up. His visual acuity improved to light perception bilaterally with resolution of optic disc elevation on funduscopic examination.

**DISCUSSION** Recognition of uncommon manifestations as well as radiologic signs of elevated ICP can be of great value, as delay in diagnosis may result in permanent deficits. The classic reported otologic features of elevated ICP are tinnitus and aural fullness; however, both conductive hearing loss and SNHL have been reported. Our patient presented with bilateral severe auditory neuropathies possibly due to stretch of the vestibulocochlear nerves from the increased ICP. Protrusion of the meningoceles into the IACs was also deemed to be a contributing factor to the hearing loss in our case. Petrous apex cephaloceles and meningoceles have been described as rare radiologic signs of elevated ICP.

Our patient showed significant improvement in auditory acuity in his left ear after the ventriculoperitoneal shunt insertion. However, his vision loss only minimally improved from no light perception to light perception bilaterally. Elevated ICP classically affects the visual system through papilledema. The profound vision loss in our patient was caused by optic neuropathy and bilateral ischemic CRVO, both of which are likely secondary to papilledema. While improvement following ICP normalization was minimal, this can be attributed to irreversible damage to the retina and ganglion cells sustained before presentation to our institution.

Extensive investigation for a secondary cause of elevated ICP in our patient was unrevealing. Given his clinical improvement following the surgical management of ICP, we conclude that this was the primary pathologic process. We present this case to highlight uncommon presentations of elevated ICP, so that prompt intervention can minimize morbidity.

**AUTHOR CONTRIBUTIONS**
Mersedeh Bahr Hosseini: primary writer of the manuscript. Laura Stone McGuire: writer of the manuscript. Milena Stosic: case data collection. Heather E. Moss: manuscript revision. Michael D. Carrrithers: manuscript revision.

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REFERENCES

Pearls & Oyster: A rare presentation of chronic intracranial hypertension with concurrent deafness and blindness
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