Pearls & Oy-sters: Angioplasty and Stenting as New Treatment Method for Cough Headache With Stenotic Internal Jugular Vein

Case Report With 12-Month Follow-up

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Pearls

- Drainage dysfunction of stenotic internal jugular veins (IJVs) is considered an etiology of cough headache as a result of fluctuation and imbalance of intracranial pressure.
- Angioplasty and stenting procedure has been shown to be a new effective method for treating cough headache caused by severe stenosis of IJVs.

Oy-sters

- It is inaccurate to determine the etiology of cough headache by a singular diagnostic imaging modality, emphasizing the need to use multimodal imaging modalities.

A 39-year-old man presented with symptoms including recurrent headaches lasting for several hours over the past 3 years, moderate to severe in intensity, localized on bilateral occipital and temporal areas, and sometimes accompanied by nausea, photophobia, and phonophobia, with no cranial autonomic symptoms. The headache was primarily provoked by sudden body postural changes and partially alleviated by resting, and gradually appeared during coughing, sneezing, and bending down, but was not triggered by sustained physical exercise. The neurologic examination was unremarkable and the lumbar opening pressure was 200 mm H2O at rest but >350 mm H2O after Valsalva maneuver (VM). MRI showed Chiari malformation type I with cerebellar tonsil hernia, with normal enhancement imaging (figure). Magnetic resonance angiography and magnetic resonance venography (MRV) revealed no abnormalities. The patient was identified as having cough headache.

Color Doppler ultrasound showed normal bilateral IJVs at normal breathing, with bilateral retrograde venous flow at VM (768 ms on the right vs 312 ms on the left). Cerebral vascular angiography located stenoses of both IJVs. The patient was recommended indomethacin (75 mg twice a day) to relieve pain and then dabigatran (150 mg twice a day) to prevent thrombosis, but symptoms were not relieved during the 4-month follow-up. Following the third edition of the International Classification of Headache Disorders (ICHD-3), the patient was diagnosed with secondary cough headache rather than primary cough headache based on both his structural cranial abnormalities and clinical features, including age <50 years, primarily provoked by sudden body postural changes rather than coughing, lasting several hours, which is much longer than 1 minute, and no response to indomethacin.1

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For this patient, surgical treatment of Chiari malformation type I was not suggested by neurosurgeons owing to the absence of indication. Therefore, we targeted IJV stenoses–related cough headache by performing angioplasty and stenting as a new treatment approach.

The patient received dual antiplatelet therapy (aspirin 100 mg daily and clopidogrel 75 mg daily) for 4 days before the procedure. The thromboelastography mapping test for antiplatelet drugs was recommended, and the inhibition rates of arachidonic acid and adenosine diphosphate were higher than 50% and 40%, respectively.

Angiogram revealed severe stenoses of the bilateral IJVs in the venous phase and slow outflow of the bilateral transverse sinus (figure, B–D). Following this, an 8F guiding catheter was inserted into the IJV. A Renegade microcatheter was manually inserted across the stenotic segment. A pressure transducer connected with the Renegade microcatheter was used for functional assessment of IJV stenoses. The pressure gradient of bilateral IJVs was measured to be 9 mm Hg (distal and proximal pressure of stenosis of the left and right IJVs were 14 and 5 mm Hg, 13 and 4 mm Hg, respectively).

A 0.014-inch (300 cm) Transend microwire was manipulated in the IJV. Following balloon inflation, 2 Precise stents were deployed in the bilateral stenotic IJVs. Angiography showed mild residual postprocedural stenoses of bilateral IJVs, and the VasoCT scan revealed full expansion of the stents (figure, E and F). Clopidogrel and aspirin were maintained for 6 months after the operation. During a 12-month follow-up, the headache was mostly relieved from coughing, sneezing, and bending down, but could appear when bowing the head.

Discussion

Chiari malformation type I is the most common cause of secondary cough headache, which manifests pressure difference between the ventricles and the lumbar subarachnoidal space after a VM. IJV stenosis is also a factor to alter intracranial pressure caused by drainage dysfunction, so it is speculated as another underlying etiology of cough headache. Our study reports a patient with cough headache diagnosed with pressure gradient across the IJV stenosis. This case led us to investigate the association between cough headache and IJV stenosis. We performed angioplasty and stenting as a new approach to treat IJVs stenosis–induced cough headache with satisfactory outcomes.

This study reveals the importance and necessity of multimodal imaging to diagnose cough headache, especially for those with additional headache triggers, higher pain intensities, and diverse headache duration (as observed in the present case). For this case, we observed normal MRV and right internal jugular reflux, with opposite results from angiography. Because the intracranial and extracranial venous system is complex with variability between individuals and is often asymmetric, Dolic et al. regarded it almost impossible to determine the relevance of a single structural or hemodynamic venous abnormality and emphasized the need to use multimodal imaging modalities.

The diagnostic reasoning of cough headache was based on ICHD-3. Although the patient had been diagnosed with secondary cough headache, other possible causes of headache were systematically assessed but were eventually precluded. ICHD-3 diagnosis 6.6, “Headache attributed to cranial venous disorder,” could be ruled out because the patient did not...
present with cerebral venous thrombosis. Moreover, this patient did not fulfill the diagnostic criteria of ICHD-3 diagnosis 7.7, “Headache attributed to Chiari malformation type I,” for several reasons. First, there is no evidence of a clear relation between headache development and Chiari malformation progression, or no surgical treatment for the diagnosis. Second, headache episodes are not associated with any symptoms or clinical signs of brainstem, cerebellar, lower cranial nerve, or cervical spinal cord dysfunction, which do not satisfy the criteria in the ICHD-3.7 However, we could not exclude this diagnosis because the headache did not disappear after treating the stenotic IJV.

Angioplasty and stenting have been used to treat stenotic IJV since Ryu et al.8 used IJV stent for treating traumatic occlusion in 1997. Afterwards, several IJV stents were used to treat chronic cerebrospinal venous insufficiency that might induce multiple sclerosis, tinnitus, visual impairment, dizziness, sleep disturbance, and neck discomfort or pain.9 In recent years, angioplasty and stenting for IJV have been rarely used because of the complex JVI variability among different individuals. However, a relatively low rate of restenosis and complications have proved stenting a safe operation.10 Donnet et al.11 performed cranio cervical MRV and found stenosis of venous sinuses or IJV in 5/7 patients in the cough headache group; they did not perform any surgical intervention for the stenosis. Our previous study has confirmed that stent implantation or balloon dilation of cerebral venous sinus can decrease intracranial pressure and improve symptoms immediately.10 We further extended the treatment to IJVs and investigated the efficacy of using angioplasty and stenting to treat cough headache with severe stenotic IJVs.


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Disclosure
The authors report no disclosures relevant to the manuscript. Go to Neurology.org/N for full disclosures.

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References


