Jugular Vein Aneurysm, When We Have to Do Surgery?

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ABSTRACT

Background: The most common lesions of jugular vein dilatation are aneurysms and ectasia. A jugular vein aneurysm is less common compared to an arterial aneurysm in adults. Because of the rare incidence, treatment guidelines primarily associated with the timing of surgery are not clearly established. Proper treatment can reduce patient complaints without excessive intervention. Case report: A 54 years old woman complained of swelling in the right neck that started three years ago and cephalgia for two years. From CT angiography, we obtained a jugular vein dilatation of 2.3 cm. During periodic evaluation from ultrasonography doppler, there is no increase in the size of the jugular vein. Conclusion: Jugular vein aneurysm presenting in adults is an infrequent phenomenon. It is a benign condition, and conservative observation is advised. It should be operated only if symptomatic or progressive enlarging. A periodic examination must be done to evaluate the size of the jugular vein before a surgical decision

1. Introduction

Jugular venous aneurysm (JVA) is a rare condition. It occurs equally between man and woman. Approximately 75% of cases are diagnosed in children. The dilatation of the jugular vein can be fusiform or saccular without torsion in the jugular vein wall. Some authors often use "ectasia" to explain fusiform dilatations and "aneurysms" for saccular ones. However, both of them are commonly used as synonyms.

A clear etiology has not yet been established. The pathology is relatively benign, depending on the localization, and can lead to severe complications such as thrombosis, pulmonary embolism, rupture, bleeding, or pressure over neighboring structures. To prevent those complications, some clinicians do surgery. However, treatment guidelines especially associated with the time of surgery, are not established.

2. Case report

A 54-year-old woman came to the emergency unit with a painless mass involving the right neck since three years ago. Initially, the swelling used to appear on coughing or loud speaking. However, the node became firm and did not reduce spontaneously later. She denied any history of trauma or other chronic diseases except hypertension. She also complained about intermittent cephalgia. There was no stridor, dysphagia, hoarseness, vomiting, dyspnea, neck pain, and chest pain.

From the physical examination, we found 1.5×2 cm, soft swelling in the right lower neck, near the clavicle,
extending to the anterior. The swelling is pulsating. There was no enlargement of the lymph node. Her blood pressure is 170/100mmHg. She has bad compliance to come to the hospital regularly, and in three consecutive visits at the hospital, she has uncontrolled hypertension. The CT scan angiography revealed that dilatation of the right internal jugular vein was approximately 23mm as high as the right thyroid level.

Meanwhile, the left internal jugular vein was normal. The patient was lost to follow up for a year. When she revisits the hospital, we do ultrasound (US) Doppler for evaluation. Figure 1 shows the result in which the right internal jugular vein was not enlarged compared to CT scan results with a diameter of 14.9x12.9mm and no thrombus formation.

Figure 1. Short axis view shows the diameter of the right internal jugular vein

From the follow-up after a year, the size of the right internal jugular vein was not increased, and no thrombus formation. The chief complaint was just intermittent cephalgia without vomiting or intracranial extension sign. She has uncontrolled hypertension and shoulder pain that can indicate neuromuscular problems as other possibilities of cephalgia’s reason. The cephalgia was not increased for a year. We consulted the neurology department and got cervical root syndrome dextra and nerve irritation C5, C6, and C7 dextra demyelinating sensory mononeuropathy. She did not bother with the cosmetic aspect of the neck swelling. Conservative therapy was decided because of the minimum symptoms and neither high-risk complications found in her jugular vein aneurysm. Periodic observation with US Doppler every three months was planned to evaluate the internal jugular vein aneurysm progression. We also optimize her antihypertensive therapy to control blood pressure.

### 3. Discussion

A jugular vein aneurysm is a rare condition. However, it is the most often venous deformity affecting the neck veins\(^5\). JVA is commonly from the internal jugular vein. It can also involve the external jugular vein, and the rarest affects the anterior one\(^1\). Paleri et al. stated that up to 10% percent of the jugular aneurysms could be bilateral. The cause of this disorder is still unclear, but several factors are suspected as predisposing factors\(^6\). The lack of elasticity of the vein wall is mentioned as the possible cause and commonly the dilatation in fusiform type or "ectasia" \(^5\). Meanwhile, saccular dilatation or aneurysms are caused secondary to tumors, inflammation, or trauma, although they can also appear spontaneously\(^1\). Other predisposing factors of JVA are elevated pressure in the internal jugular vein, compression between the cervical pleura and head of the clavicle, mechanical obstruction of the lower neck and upper mediastinum, thoracic outlet syndrome, and anterior scale compression. JVA is generally seen in the right jugular vein in children because the right side lung dome is higher than the left one. So, it makes the right jugular vein more susceptible to increased intrathoracic pressure\(^7,\&\). Another possible cause is the right internal jugular vein valves which are placed higher than the left one. On the contrary, in adults, JVA is mainly acquired and usually on the left side. It has
been related to elderly hypertensive patients because of compression to the left innominate vein by the atherosclerotic aorta. The most clinical manifestation of the venous aneurysm is soft compressible swelling, commonly appearing along the vein axis. It is usually asymptomatic, but the main problem that patients often complain about is the cosmetic aspect. Other rare presentations reported were atypical chest pain. In cases of JVA with malignancy, symptoms of dysphonia are usually found. JVA complications can be hoarseness of voice or Horner’s syndrome and can be complicated by thrombosis. When an asymptomatic JVA starts causing symptoms such as pain, we should suspect thrombosis. According to the literature, the risk of thrombosis in an internal jugular vein aneurysm is less than 1%. JVA usually does not grow progressively and rapidly. Until now, there has never been a spontaneous rupture of the swelling reported. Another less frequent manifestation but ever reported is the intracranial extension. It can be accompanied by CNS involvement or not. Other diagnoses of neck swelling include a laryngocele, brachial cyst, cavernous hemangioma, superior mediastinal cyst, or cystic hygroma. In this case, the swelling is painless and pulsating. In the beginning, we think about the carotid artery aneurysm, but after we do CT angiography, there is dilatation of the internal jugular vein confirmed by ultrasound doppler. To distinguish between both structures, we performed a Doppler examination. We get dilatation from the venous structure from the US doppler, which is an internal jugular vein.

Computed tomography, MRI, and ultrasound can also exclude other diseases such as dermoid cyst, cervical adenitis, bronchogenic cyst, metastatic adenopathy, or thyroglossal duct cyst. US Doppler imaging is the cheapest non-invasive diagnostic examination to differentiate between cystic and solid or vascular and non-vascular lesions. Sometimes, JVA cannot be diagnosed by a CT scan, moreover, in the case of mild internal JVA. US doppler is preferred because dilatation of the internal jugular vein can be interpreted dynamically by comparing the size of the vein at rest and the Valsalva maneuver. However, the CT scan is usually used to exclude any causative structural lesion. So, both imaging tests can be used to complement each other. Color doppler of both lower limbs and abdominal ultrasonography to see the great veins should be done to rule out any other venous malformations. A good clinical examination and US Doppler can diagnose this condition accurately and help differentiate it from other neck swellings.

Surgery indications are still debated because most JVAs are benign and asymptomatic lesions that can be conservative with regular follow-up and reassurance. However, if it has been decided on a patient not to undergo surgery, the clinician must periodically observe the lesion for changes and progressions that maybe occur. Besides, patients should be protected from all trauma to prevent bleeding and infection. The cosmetic aspect is often used as the reason for surgery in the case of JVA. It is also indicated if an aneurysm becomes painful due to thrombosis or phlebitis from the jugular venous system. However, such complication was infrequent, so prophylactic anticoagulation therapy was not routinely given. Another indication of surgery is Horner’s syndrome, intractable cough, congestive cardiac failure, massive hemorrhage secondary to trauma, and spontaneous rupture. Surgical resection eliminates the theoretical risk of those complications. Embolism or rupture of a venous aneurysm may appear in several locations resulting in emergency surgery or death. This circumstance occurs in some cases of deep venous aneurysms in the abdomen and lower extremities. Although there were no reports of life-threatening complications for JVA, most clinicians choose surgical treatment in saccular aneurysm cases. On the contrary, surgical treatment is only done in fusiform dilatation if the lesions are symptomatic or progressive enlarging. It is usually managed by a conservative and rarely needs surgery.

In the setting of our case, the patient had fusiform dilatation of the jugular vein without such complications. Follow-up evaluation after a year did not show any progression of the disease. The only symptom that complained is cephalgia. Most literature did not
mention cephalgia as the clinical manifestation of JVA. Even intracranial extension of JVA can be excluded because we did not get CNS involvement. By contrast, cephalgia is often described in jugular vein compression as the manifestation of mechanical obstruction of intracranial vein drainage\(^8,11\).

4. Conclusion

Jugular vein aneurysm presenting in adults is a very rare phenomenon. It is a benign condition, and conservative observation is advised. The surgery indicates a cosmetic or physiological concern, painful aneurysm secondary to thrombosis or phlebitis, other complications such as Horner’s syndrome, intractable cough, congestive cardiac failure, massive hemorrhage secondary to spontaneous trauma rupture. Surgery is also preferred if the lesion is symptomatic, enlarging, or disfiguring. Therefore, if conservative management is decided, it needs periodic follow-up to observing the lesion for any changes and recording the natural course.

5. Conflicts of interest

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7. Authors’ contributions

RA has given substantial contributions to the conception of the design of the manuscript. MP and JN were major contributors in writing the manuscript. IPD was editing the manuscript for publications. All authors have participated in drafting the manuscript, JN revised it critically. All authors read and approved the final version of the manuscript.

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9. Consent for publication

Written informed consent was taken from the patient to use medical data for academic and research purposes, including publication.

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