Vertebral Artery Dissection: 
a Contemporary Perspective

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ABSTRACT
Arterial dissections are among the most frequent causes of stroke in young adults. Usually they are 
associated with trauma, but as the modern imaging tools are evolving, more dissections are being di-
agnosed and more etiologies are being described. Vertebral artery dissections (VADs) have the distinct 
particularity that they can cause ischemic stroke (in the brainstem, cerebellum or even the spinal cord), 
but also subarachnoid hemorrhage, when the dissection occurs in the intracranial segment of the ver-
tebral artery. We present a review of the literature, going over etiology, clinical aspects, diagnosis and 
treatment of VADs and we illustrate the theory with three different types of VAD from our clinical 
experience.

Keywords: vertebral artery dissection, ischemic stroke, subarachnoid hemorrhage, doppler ultrasound

The vertebral artery has its origin in 
the subclavian artery and is divided 
into four segments: V0 (the origin), 
V1 (short segment before taking the 
laterocervical pathway), V2 (the 
passage through the transverse apophyses 
of the cervical spine), V3 (the retromastoid seg-
ment), and V4 (the intracranial portion, up to 
the point where it joins the other vertebral ar-
tery to form the basilar artery). Dissections can 
occur in any segment of the vertebral artery, 
but opinion differs among authors regarding 
the susceptibility to dissection of the four seg-
ments. The distal V1- and the proximal V2-seg-
ment (at the level of C6 vertebral body) were 
the most frequent locations of dissections (43% 
of cases) in a study published by Bartels which 
included 28 patients (1), while other authors 
found that the location of the dissection was 
more often in the pars transversaria (V2; 35%) 
or atlas loop (V3; 34%) than in the prevertebral 
(V1; 20%) or intracranial (V4; 11%) segment 
(2). The V4 segment has a different histological 
structure, compared with the other segments of 
the vertebral artery, as the thickness of the tu-
ника media and the adventitia tapers when the 
vessel pierces the dura. Also, the vessel is sub-
ject to shearing forces associated with head 
motion (3). In a study performed on 983 cases 
of cervical artery dissections (CeAD), 149 
(15.2%) presented with multiple CeADs (4). 

In the majority of cases, dissection occurs at 
the endothelial level, with rupture of the arte-
rial intima, and the development of a false lu-
men. Blood will enter between the intima and 
the media, will quickly coagulate, and will lead 
to different degrees of stenosis or even an oc-
cclusion. The hematoma is distributed around 
the circumference of the vessel, and has a cres-
cent-shape, which is clearly visible on axial
MRI sections (at cervical or cranial level). In the next weeks, up to 60% can recanalize, by resorption of the hematoma, 39% reaching complete recanalization (5). The other histological site of dissection is much more rare, consists of the cleavage of the arterial wall between adventitia and media, with leakage of the blood in the subarachnoid space, and occurs in V4 segment. Clinical presentation with subarachnoid hemorrhage (SAH) is more rare, but it is associated with worse outcome compared with non-SAH VAD, one of the causes being the high incidence of rebleeding (6).

There are several instances which favor the occurrence of a dissection: trauma, infection and inflammation, smoking, a particular genetic background, or certain diseases such as Ehlers-Danlos or Marfan disease.

A person who has a dissection of the vertebral artery can display an entire range of symptoms, from completely asymptomatic to a severe stroke in the vertebro-basilar territory. The consequences of a dissection depend on several factors: the location of the dissection, the degree of obstruction, the functionality of the collateral circulation. Vertebral artery dissection is one of the most frequent causes of stroke in people aged between 18-45 years, the estimated frequency being between 1-2.6 per 100000 (7). Strokes occur in the medulla or in the cerebellum (territory of the postero-inferior cerebellar artery, PICA), but infarction can appear also in the distal part of the vertebro-basilar system or in the spinal cord (since the anterior and posterior spinal arteries have their origin in the vertebral arteries).

Case 1. A 58-year–old male smoker, with no other vascular risk factors, presented to our Emergency Department because of the abrupt occurrence of postural imbalance and numbness of the left face and lower right limb. The patient denied any neck pain or any acute traumatic injury at the cervical level. Upon admission, the neurologic examination revealed a left sided Horner syndrome, decreased pinprick and thermal sensation in the entire territory of the left trigeminal nerve and in the right lower limb. He had a wide-based, unsteady gait. No other neurological abnormalities were noticed.

Doppler ultrasound of the cervical arteries showed a high-grade left vertebral artery stenosis in the V1/V2 segment; in the V3-V4 segments the flow velocities were lower, with a poststenotic shape. The presence of a stenosis associated with the absence of other atherosclerotic lesions raised the suspicion of a dissection in the V1 segment of the left vertebral artery. A brain MRI was performed on the day of admission. Diffusion-weighted (DWI) MRI showed a small lesion in the dorsolateral portion of the left lower medulla, behind the retro-olivary sulcus, not visible on T2 sequences (Figure 1). On the contrast enhanced T1-weighted images of the neck, the lumen of the left vertebral artery was thinned out throughout the entire course. A thickening of the wall with nonspecific signal transduction was observed in the distal V2 segment. MRI findings were consistent with a dissection of the left V2 and V3 vertebral artery, in the subacute stage.

Resting 12 lead ECG revealed a left bundle branch block. Transthoracic echocardiography showed left ventricle concentric hypertrophy and grade I diastolic dysfunction. Routine blood analysis highlighted a slight disturbance of the lipid profile (total serum cholesterol was 239 mg/dl and HDL cholesterol was 40 mg/dl).

The patient was given unfractionated heparin for 5 days and then switched to dual antiplatelet therapy (aspirin 75 mg daily and clopidogrel 75 mg daily) and statin (rosuvastatin 20 mg daily). His clinical condition improved within a few days. Upon discharge, 9 days after admission, he no longer had numbness of the left face or postural imbalance, and he was able to walk unassisted. At discharge: left

FIGURE 1. Case 1. DWI-MRI shows an acute ischemic stroke (the white arrow: hyperintense signal) in the left part of the lower medulla, behind the retro-olivary sulcus.
Horner syndrome and a mild impairment of temperature and pinprick sensation in the right lower limb. At the 30 days follow-up, the Doppler ultrasound showed partial recanalization of the left vertebral artery, and a minimal hyposthesia in the right lower limb.

Case 2. A 61-year old male, with a history of alcohol abuse disorder, presented to the Emergency Department for the sudden onset of slurred speech, dysphagia and dysphonia, followed by drooping of the left eyelid, numbness on the left side of his face, unsteady gait, difficulty when using his left limbs and vertigo. The symptoms began several hours before arriving at the hospital, he mentions he felt slight pain on the left side of his neck prior to symptom onset.

On clinical examination he had moderate dysarthria and dysphonia, inward deviation of the left eye, limitation of conjugated eye movements towards the left, peripheral facial palsy, rotational nystagmus on looking towards the right and upwards, and difficulty swallowing, with an impaired pharyngeal reflex on the left side. He also had ataxia of all limbs, more severe on the left side, left inferior limb paresis, brisk reflexes and clonus of the foot bilaterally, a wide-based gait, could not stand upright unaided, and thermal and pain sensation were decreased on the right side of his body. Given the sudden onset and the presenting symptoms, a posterior circulation stroke was the most likely cause, and given the associated left cervical pain we suspected a vertebral artery dissection. Initial cerebral CT scan was normal.

Doppler ultrasound examination of the cervical and cerebral arteries revealed a systolic flow speed of 180 cm/sec in the left vertebral artery (V2 segment) and absence of the flow in V4 segment, supportive of our clinical suspicion.

Brain MRI (performed 9 days from onset) showed a subacute ischemic stroke in the territory of the postero-inferior cerebellar artery. The left vertebral artery had a thick wall, isointense on T1 and T2 weighted sequences, and, on MRA, a discontinuous flow in the second and third segment and slow flow in the fourth segment, consistent with a left vertebral artery dissection in the second and third segment.

The patient received unfractionated heparin and was afterwards switched to aspirin, clopidogrel and a statin. The patient had a satisfactory evolution during his stay in our clinic, with improvement of the eye movement, ataxia and dysarthria. The swallowing difficulty persisted on discharge, albeit slightly improved, and we decided to keep the nasogastric tube we inserted. The patient returned for follow-up one month later, with significant improvement in gait and coordination, dysphonia and nystagmus, but also being able to swallow food of moderate consistency and not requiring the nasogastric tube. Ultrasound examination showed the same appearance.

Case 3. A 68 year-old female, was admitted to our clinic for dizziness, a balance disorder, vertigo, vomiting and transient neck pain which had begun 4 days prior to her admittance. Her medical history included diabetes mellitus and arterial hypertension. On neurologic examination, the patient had a positive Romberg sign with a tendency to retropulsion and no other neurological signs. Her CT scan showed no recent intracranial vascular lesions, but revealed a spontaneously hyperdense left vertebral artery with a larger diameter; ultrasound examination showed high velocities on the vertebral arteries, suggesting a 50% left vertebral artery stenosis and a 60-70% right vertebral artery stenosis. A few days after her admittance to our clinic...
department she developed intense headache, photophobia, phonophobia, vomit and neck stiffness. An emergency CT scan excluded new lesions. Spinal tap revealed a hemorrhagic cerebrospinal fluid (traumatic spinal tap was excluded). A subarachnoid hemorrhage was diagnosed, due to a vertebral artery dissection. MRI confirmed the vertebral artery dissection, with occlusion of the left vertebral artery, and a stenosis >70% of the right vertebral artery, without any image of ischemic stroke (Figure 3).

The three cases illustrate the clinical versatility of the vertebral artery dissection (with two different location of ischemic stroke for the same site of the VAD, and one subarachnoid hemorrhage), and the importance of choosing the right method for diagnosing a stroke (including the etiological diagnosis).

Ultrasound examination can be a very useful tool for diagnosing blood flow disturbances in cervical and cerebral arteries. Typical configurations for dissection include narrow, bi-directional complexes (with positive and negative flow), caused by an increased resistance to the blood flow, when dissection is in the upper segment of the vertebral arteries and double lumen or an intimal arterial flap in the lumen, but this is rarely seen; stenosis or occlusion can be frequently observed, but ultrasound cannot point out exactly the site of the stenosis, and cannot differentiate between stenosis due to atherosclerosis or dissection. When the patient has no other atherosclerotic lesions, and is young, the scales can easily sway towards dissection, but in older people, other methods must be used to prove it. Ultrasound criteria for diagnosing a vertebral artery stenosis are less precise compared to those for internal carotid artery.

Other options to show a vertebral artery stenosis or occlusion are CTA, MRA or DSA, but the hallmark of the dissection is the presence of the arterial wall hematoma, identifiable on axial MRI slices. For occluded arteries the sensitivity of duplex without colour, duplex with colour, time of flight (TOF) MRA and contrast enhanced (CE)-MRA was over 98%, with a specificity ranging from 90.8% (95% CI 89.4 to 100) for duplex to 100% (95% CI 97.5% to 100) for both non contrast and CE-MRA (8).

MRA (magnetic resonance angiography) has a tendency to overrate the degree of stenosis, and data regarding the sensitivity and specificity of the method given by different studies ranges between 53.8% to 100%, for both non-contrast and contrast enhanced MRA (9-12).

CTA (computed tomography angiography) has few studies but the results suggest that the method can be superior to TOF MRA, when slow flow is present, or for the detection of intracranial vertebral artery stenosis or occlusion (13). DSA (digital subtraction angiography) remains the gold standard for exploring the residual arterial lumen. Exploring the internal carotid artery, the flame shape of the occlusion can be a hint for dissection, but in vertebral arteries this is less obvious. Other changes (thrombembolism or atherosclerosis) can produce a stenosis or occlusion, so catheter angiographic findings are nonspecific. For patients who presented with SAH, the most frequently reported angiographic findings were fusiform

![FIGURE 3. MRA (left panel, white arrow): occlusion of the left vertebral artery; MRI (Right panel, white arrow): Enlargement of the diameter of the VA with thrombus in the left vertebral artery.](image)
aneurysms (70.8%), and pearl-and-string lesions (24.5%), meaning a succession of aneurysmal dilatations and vessel constrictions (6).

MRI is the only reliable method for exploring the arterial wall. A hyperintense, crescent-shaped intramural hematoma is specific for arterial dissection, on axial cuts at cervical or cranial level (depending on the site of the dissection). If there is a residual patent lumen this will appear as an eccentric flow void. The aspect of the intramural hematoma is time dependent. In the first 3 days it has a high signal intensity on T2-weighted images with intermediate signal intensity on T1-weighted images followed in the next days in most cases by a slightly or definitively increased signal intensity on T1- and T2-weighted images. The increased signal will remain high for approximately 2 months (14).

Apart from T1- and T2-weighted images, other MRI techniques were studied in order to improve the accuracy of the diagnosis. The intramural hematoma sign is considered positive if the patient has an eccentric or concentric hypointense signal lesion in the vertebral artery on susceptibility weighted imaging (SWI), a corresponding hyperintense signal on phase map and no evidence of calcification on the brain CT (15). Another technique, black blood T1-weighted imaging (3D-BB-T1WI) revealed the characteristic crescent shape of the intramural haematoma in 14 cases (87.5%), being considered a promising technique (16). Two other techniques are mentioned for detection of VAD. One is BPAS (BasiParallel Anatomic Scanning) which was designed to visualize the surface appearance of the vertebrobasilar artery within the cistern and the other is VISTA (Volumetric isotropic TSE acquisition) based on black blood imaging method, designed to evaluate the arterial wall and lumen. Although the sensitivity is still low, these are considered promising techniques for diagnosing VAD. A recent study published by Natori (17) highlights the advantages of 3D-T1-weighted imaging, upon BPAS + MRA, MRA alone or MRI alone, the technique being able to identify luminal stenosis, aneurysmal dilatation, intramural hematoma or intimal flap in 100% of cases.

Treatment of the VAD is still a subject of debate. Should we use anticoagulants or antiplatelet therapy? Is it safe to administer fibrinolitic treatment in the setting of an ischemic stroke due to VAD? When should we use endovascular treatment? These are just several questions found in currently daily practice, answers being based more on the personal experience or local habits, than on evidence based data.

MEDICAL TREATMENT

Current guidelines for the treatment of acute ischemic stroke (18) do not list arterial dissection among the contraindications of thrombolysis. Thrombolysis was independently associated with neither an unfavourable outcome, nor with an excess of symptomatic bleedings in an analysis performed on 616 cervical artery dissections from a multicentric stroke data base – CADISP (19). However, the lack of any trend towards a benefit of thrombolysis may sustain the search for more efficient treatment options including mechanical revascularization strategies.

The choice between anticoagulant and antiplatelet therapy has not been settled yet. There are no reliable data from randomised trials to decide whether anticoagulants or antiplatelet agents are better to prevent further thromboembolic events after cervical arterial dissection.

Most neurologist favor anticoagulants, especially in the VAD, but this is sustained more by personal experience and not evidence-based data. A meta-analysis (20) performed on 34 non-randomised studies, which encompassed 762 patients, showed no significant difference with regard to the risk of death (antiplatelet 5/268 (1.8%), anticoagulation 9/494 (1.8%), p = 0.88); stroke (antiplatelet 5/268 (1.9%), anti-coagulant 10/494 (2.0%), p = 0.66), or stroke and death.

The endovascular approach is an attractive alternative for the treatment of vertebral artery dissection in the acute setting. However, while various techniques have been used (although a large part of the published data refers to the treatment of local pseudoaneurysms following the dissection), these haven’t been studied in a manner which allows for the issuing of recommendations and therefore, at least for the time being, angioplasty and stenting of the vertebral arteries in the setting of an acute stroke due to VAD remain an option in selected cases.

CONCLUSION

Arterial dissections are more frequently diagnosed during the last years, as the access to performant imagistic tools has increased. VAD
has the particularity to cause both ischemic stroke and subarachnoid hemorrhage and should be taken into account as a possible etiology for these diseases, especially in young adults. Medical treatment (either antiplatelet or anticoagulant) is the first option for secondary prevention of stroke, but endovascular treatment with stenting is emerging.

Conflict of interests: none declared.

Financial support: none declared.

REFERENCES


