CASE REPORT

Isolated oculomotor nerve palsy: a rare manifestation of internal carotid artery dissection

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SUMMARY

A 58-year-old man reported sudden-onset binocular double vision that appeared 3 days earlier. He denied history of headache/cervical pain or trauma. He had a medical history of well-controlled diabetes, hypertension and dyslipidaemia. Neurological examination revealed a left-sided ptosis and binocular horizontal diplopia in dextroversion without apparent extracranial-muscle paresis or pupillary involvement. Other cranial nerves were spared as well as motor, sensory and coordination systems. There were no signs of ocular erythema, proptosis or palpable orbital mass. Brain MR angiography revealed a crescent-shaped mural hyperintensity in left internal carotid artery (ICA) at skull base, extending to intrapetrous segment, with reduced calibre and flow, suggesting a left ICA dissection. The patient was started on antiaggregation therapy. A year later he was asymptomatic and CT angiography confirmed ICA recanalisation.

BACKGROUND

Internal carotid artery dissection (ICAD) is an important cause of ischaemic stroke among young and middle-aged patients, accounting for 14–20% of all cases.1 Trauma and connective tissue disorders are well-known predisposing factors, but the majority of cases are spontaneous or related to minor/trivial trauma.2

ICAD pathophysiology lies in a compromise of structural integrity of the arterial wall through the formation of an intimal tear, resulting in stenosis, occlusion, intraluminal thrombus or extravascular haemorrhage. Most cases comprise extracranial arteries but, rarely, intracranial arteries are affected.2

ICAD usually presents as a triad of headache/neck pain, Horner syndrome or cerebral ischaemic symptoms. However, most patients do not show the complete triad and may exhibit less common manifestations such as cranial nerve palsies.2

Cranial nerve palsies are described in around 12% of spontaneous extracranial ICAD cases, mainly as a syndrome of lower cranial nerve palsies (IX, X, XI and XII). Isolated III (oculomotor) nerve palsy is rare, described in only 2 of 155 patients with ICAD in a case series.3

Nevertheless, the mechanism underlying these cranial neuropathies is still not fully understood.4–6 Direct compression by an enlarged dissected vessel is proposed to be responsible for lower cranial nerve palsies. However, concerning the III cranial pair, an anatomic proximity with ICA is only found within the cavernous sinus.

We present a case report on ICAD presented as III nerve palsy to emphasise the importance of considering a cervicocerebral dissection in less typical clinical presentations.

CASE PRESENTATION

A 58-year-old man came to our emergency department (ED) reporting sudden-onset persistent binocular double vision which had appeared 3 days before. He denied headache, cervical pain and trauma history.

He had medical history of well-controlled diabetes, hypertension and dyslipidaemia. Family medical history was irrelevant for stroke or connective tissue disorders.

On initial ED evaluation the patient was found to be afebrile with a blood pressure of 149/78 mm Hg and a regular pulse of 75 bpm. General physical examination was normal. He showed no signs of ocular erythema, proptosis or palpable orbital mass. On neurological examination he was alert, oriented, with no evidence of higher function abnormalities. The patient presented isocoric and normally reactive pupils, partial left ptosis and, although with no obvious limitation in eye movements, the patient consistently mentioned binocular horizontal diplopia in dextroversion. There was no evidence of abnormalities involving other cranial pairs, muscle tonus and strength, osteotendinous reflexes, cutaneoplantar reflexes, sensation, coordination and gait.

INVESTIGATIONS

Laboratory screening: normal haemogram, ionogram, liver, renal, thyroid function, sedimentation rate; glycated haemoglobin 5.3%; total cholesterol 215 mg/dL, low-density lipoprotein cholesterol 153 mg/dL, high-density lipoprotein cholesterol 46 mg/dL, triglycerides 80 mg/dL; hepatitis B and C, HIV and borrelia serology negative; treponema pallidum hemagglutinations assay/Venerale Disease Research Laboratory negative.

The ECG had sinus rhythm.

Transhoracic echocardiogram was normal.

Brain non-contrast CT scan obtained in ED showed no signs of infarction.

Brain MRI angiography revealed a crescent-shaped mural hyperintensity in left ICA at skull base that extends to the intrapetrous segment with reduced calibre and flow, acquiring his normal pattern in the intracavernous portion suggesting a left ICAD. Brain MRI did not show evidence of parenchymal signal abnormalities (figures 1 and 2).
DIFFERENTIAL DIAGNOSIS

Addressing a sudden-onset persistent horizontal diplopia with clinical signs of incomplete pupil-sparing oculomotor nerve palsy (partial left ptosis and diplopia maximal at dextroversion), the main differential diagnosis includes stroke and ischaemic oculomotor mononeuropathy.

The sudden onset and presence of vascular risk factors (arterial hypertension, diabetes and dyslipidaemia) favours an ischaemic aetiology. An infarct involving oculomotor nucleus is rare and clinically different from III cranial nerve palsy. Paired oculomotor nuclear complexes are located in dorsal midbrain at superior colliculus level. An infarct involving the caudal subnucleus located in midline may result in bilateral ptosis as this nucleus provides innervation for bilateral *levator palpebrae superioris*. As already described, our patient had a unilateral ptosis that did not support this hypothesis.

Regarding the hypothesis of an ischaemic mononeuropathy, it is well known that oculomotor is the most frequent cranial nerve palsy associated with diabetes, affecting predominantly older patients with long-term diabetes. The clinical presentation has a sudden onset, usually spares pupillary involvement and can be accompanied with retro-orbital pain.

TREATMENT

Addressing a cervicocerebral artery dissection, the current accepted therapy remains antithrombotic medication (anticoagulation or antiaggregation) and there is no consistent medical evidence to guide therapy. We opted for antiaggregation therapy with 150 mg aspirin/day.

OUTCOME AND FOLLOW-UP

After 3 months, the patient was asymptomatic and showed no abnormality in neurological examination. Brain CT confirmed ICA recanalisation and cervical doppler ultrasonography showed normal morphological and haemodynamic features.

Owing to the presence of vascular risk factors he maintains antiaggregation therapy.

DISCUSSION

The patient reported a horizontal binocular diplopia maximal when looking to the right, which, in association with left ptosis, is consistent with a left oculomotor nerve palsy. Although we were not able to observe any limitation in ocular movements, this does not exclude III nerve palsy.

It has been reported that some ICAD cases are actually asymptomatic. We do not support the hypothesis that a left oculomotor nerve palsy and a left intrapetrous ICAD are simply a manifestation of two different conditions (an ischaemic mononeuropathy and an asymptomatic ICAD). Thus, the patient has type 2 diabetes with excellent glycaemic control on oral antidiabetics and does not show evidence of microvascular/macrovascular involvement of target organs. Moreover, the patient denied ocular pain that often accompanies an ischaemic neuropathy.

Anatomically, the distal part of the III nerve is supplied by the inferior cavernous sinus and meningohipophyseal arteries, both branches from the intracavernous segment of ICA. Thus, it has been proposed that an embolic and/or haemodynamic event causing hypoperfusion of oculomotor *vasa nervorum* may be responsible for III paresis in the setting of ICAD involving extracavernous segments.4

ICAD presented with oculomotor nerve palsy is rare and there are few cases described in the literature. Hegde *et al*5 described a case of a 60-year-old woman with headache and evidence of a pupil-sparing oculomotor nerve palsy, which started 3 weeks after a frontal trivial trauma, with MRI features of dissection involving petrous segment of the left ICA.

There are also reports of cases involving extracranial ICA segment. Campos *et al*6 described a case of a 50-year-old man who began having an occipital headache and developed diplopia days later, with objective impairment in right eye adduction and upward gaze and slight pupillary dilation with MR angiography, evidence of a dissection of right extracranial ICA. Nizam *et al*7 described a case of a 24-year-old man presenting a right hemianopsia, hemiparesis and hemihypesthesia 2 days after the onset of a right complete oculomotor nerve palsy, with MRI evidence of a dissection involving extracranial left ICA and an acute infarct in left middle cerebral artery.

Figure 1  MR angiography showing irregularities in left internal carotid artery lumen.

Figure 2  Fat-suppressed T1-weighted MRI showing a crescent-shaped haematoma within the wall of the left internal carotid artery, adjacent to the eccentric lumen.
Learning points

▸ Internal carotid artery dissection (ICAD) cases may present without the more common clinical signs, such as Horner syndrome, headache/neck pain or focal neurological ischaemic signs.
▸ Although rare, cranial nerve palsies can be the only manifestation of an ICAD.
▸ Considering the small group of patients presenting cranial nerve palsies in the setting of an ICAD, oculomotor nerve involvement is extremely rare.
▸ The hypothesis of a cervicocerebral dissection should be considered when addressing a young patient with cranial nerve palsy.

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