A Collet–Sicard syndrome due to internal carotid artery dissection associated with cerebral amyloid angiopathy–related inflammation

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Abstract

Background: Cerebral amyloid angiopathy–related inflammation is a rare condition with approximately 100 reported cases. Its clinical manifestations are varied. We report here a novel presentation of this disease.

Case presentation: A 61-year-old Caucasian man presented with rapidly progressive paralysis of the IX, X, XI and XII right cranial nerves associated with right central facial nerve palsy. Brain computed tomography angiography and cerebral catheter angiography found a focal fusiform enlargement of the distal cervical portion of the right internal carotid artery, related to a pseudo-aneurysm suggesting an evolution of a dissection and intra-cranial vessel dysplasia. Brain magnetic resonance imaging showed multiple asymmetrical subcortical regions of hyperintensity on T2 fluid-attenuated inversion recovery sequences. Punctiform cortical hyposignals on T2-weighted gradient echo magnetic resonance imaging sequences were mostly congruent with the white matter hyperintensities. There was a decreased cerebral perfusion at the frontal hyperintense fluid-attenuated inversion recovery region. Spectrometry identified a lactate–lipid peak. A brain biopsy showed intravascular amyloid deposits. Corticosteroid therapy was initiated, leading to a dramatic improvement of both clinical condition and magnetic resonance imaging brain lesions.

Conclusion: This case report suggests that extra-cranial vasculitis and dysplasia can exceptionally be found in patients satisfying cerebral amyloid angiopathy–related inflammation criteria.

Keywords
Cerebral amyloid angiopathy, cerebral amyloid angiopathy–related inflammation, vasculitis, dissection, Collet–Sicard syndrome

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Background

The term “cerebral amyloid angiopathy–related inflammation” (CAA-ri) was introduced to refer to inflammatory phenomena in relation to amyloid β protein deposits on cortical cerebral and meningeal vessel walls.1 Collet–Sicard syndrome, on the other hand, consists of unilateral IX–XII cranial nerve lesions.2 Both of these entities are extremely rare, with less than 100 cases described for each.1,2 We report a unique presentation of CAA-ri associated with Collet–Sicard syndrome.

Case presentation

A 61-year-old right-handed man without particular history, or risk factors presented for otorhinolaryngology consultation...
after 48 h cervicalgia, dysphonia, swallowing and cognitive disorders. Examination found a right central facial palsy, a nasal voice, a rightward deviation of the tongue on protrac-

tion, an asymmetric soft palate and hanging curtain sign affecting the right side. Nasal endoscopy found right hemi-

pharyngeal and hemilaryngeal paresis and right pyriform

sinus stasis. No headache was reported. The patient showed

slowed ideomotor response without detectable cognitive

impairment, on usual cognitive battery (MMSE = 29/30, nor-

mal Batterie Rapide d’Efficience Frontale (BREF), Dubois

and clock-drawing tests).

Brain computed tomography (CT) scan found frontal and

left temporal cortical and subcortical edema. On brain CT

angiography, there was an ectatic aspect of the distal branch of the right middle cerebral artery (Figure 1(a)). Cerebral catheter angiography found a focal fusiform enlargement of the distal cervical portion of the right internal carotid artery, related to a pseudo-aneurysm suggesting an evolution of a dissection, above which a string of beads suggesting a dysplastic artery (Figure 1(b)). Brain MRI: (c) T2-weighted FLAIR sequence, axonal slice: bilateral frontal lobe hypersignal. Moderate mass effect. (d) T2-weighted gradient echo sequence: numerous punctiform cortical hyposignals adjacent to white matter infiltration regions. (e) Spectrometry, TE (echo time) 35 ms: normal NAA/Cr and Cho/Cr ratios, slight lipid–lactate peak at 1.33 ppm. (f) Color map, perfusion-weighted sequence: reduced rCBV in FLAIR hypersignal region. (g) Control MRI after 3 months of corticotherapy: T2-weighted FLAIR sequence, axonal slice, showing clear regression of white matter hypersignal, compared to initial images. Right frontal cerebral biopsy (arrow on (c)): (h) arachnoid and cortical vessels stained by Congo red. Polarized light: green birefringence of amyloid deposits.

Figure 1. Complementary examinations: Brain CT angiography (a) and brain arteriography (b): On brain CT angiography, there was an ectatic aspect of the distal branch of the right middle cerebral artery (arrow on (a)). Cerebral catheter angiography found a focal fusiform enlargement of the distal cervical portion of the right internal carotid artery, related to a pseudo-aneurysm suggesting an evolution of a dissection, above which a string of beads suggesting a dysplastic artery (arrow on (b)). Brain MRI: (c) T2-weighted FLAIR sequence, axonal slice: bilateral frontal lobe hypersignal. Moderate mass effect. (d) T2-weighted gradient echo sequence: numerous punctiform cortical hyposignals adjacent to white matter infiltration regions. (e) Spectrometry, TE (echo time) 35 ms: normal NAA/Cr and Cho/Cr ratios, slight lipid–lactate peak at 1.33 ppm. (f) Color map, perfusion-weighted sequence: reduced rCBV in FLAIR hypersignal region. (g) Control MRI after 3 months of corticotherapy: T2-weighted FLAIR sequence, axonal slice, showing clear regression of white matter hypersignal, compared to initial images. Right frontal cerebral biopsy (arrow on (c)): (h) arachnoid and cortical vessels stained by Congo red. Polarized light: green birefringence of amyloid deposits.
normal at 10.05 ng/L, β-amyloid 1-42 was low at 436 ng/L (Alzheimer’s threshold, >700 ng/L) and β-amyloid 1-42/1-40 ratio was 0.043 (Alzheimer’s threshold, >0.075), in favor of amyloid deposits. Thoraco-abdomino-pelvic CT was normal. A positron emission tomography (PET)-scan was performed and was normal with no arguments for aortitis and vasculitis.

Right frontal cerebral biopsy (arrow on Figure 1(c)) found thick-walled arachnoid and cortical vessels, without perivascular inflammatory infiltrate, granuloma or giant cells. Specific staining revealed congophilic amyloid deposits (Figure 1(h)). Genotype analysis showed that the patient was homozygous for the ε4 variant of apolipoprotein E.

The patient received 1 g intravenous methylprednisolone, after cerebral biopsy was made, for 3 consecutive days, then 1 mg/kg corticosteroids daily, progressively tapered over 6 months. At 3-month follow-up, the facial palsy had regressed, the conversational voice was of good quality, swallowing was restored and there was no more cognitive disorders. After 6-month treatment, CSF analysis was normal, tau protein levels had decreased to 241 ng/L, and β-amyloid 1-40 and 1-42 levels were stable, respectively, 10.37 and 443 ng/L. Control brain MRI showed clear regression of FLAIR hyper-signals (Figure 1(g)).

**Conclusion**

Our patient presented a peripheral right palsy of the last four cranial nerves (IX, X, XI, XII), constituting Collet–Sicard syndrome. This syndrome, also named syndrome of the Foramen Lacerum, could be explain here by the aspect of carotid dissection of the skull base level. Our patient also had central neurological involvement, including right central facial palsy and slowed ideomotor response which could be consistent with CAA-ri.

MRI showed white matter abnormalities with multiple micro-bleeding, immediately suggestive of CAA-ri. CAA-ri was also based on behavioral disorder, ε4 homozygosity and amyloid deposits in vessels on pathologic examination.

According to CAA-ri diagnostic criteria, the diagnosis of this patient is probable as all clinical and radiological criteria are met, but not certain as no perivascular or parietal inflammation was found together with amyloid deposits. The absence of inflammatory infiltrate in this case can be explained by the right site of the biopsy, which was not performed in predominant left brain MRI inflammatory area. Since infectious and other causes were ruled out, patient was treated with immunosuppressive therapy as recommended.

CAA-ri is divided into two subtypes: inflammatory cerebral amyloid angiopathy (ICAA) and amyloid-β-related angiitis (ABRA), according to histopathology. ICAA is characterized by perivascular inflammation and ABRA by a vasculitic transmural and intramural infiltration. ICAA and ABRA are considered by some authors as subsets of primary angiitis of the central nervous system (PACNS). The diagnosis of ICAA is more likely for our case due to the dramatic response to corticosteroids alone, the absence of enhancement of lesions on MRI and the ApoE genotype ε4/ε4. In 2011, Di Francesco’s et al. hypothesized that inflammation in CAA-ri is an autoimmune response specific to amyloid protein deposits and showed an intrathecal synthesis of anti-β-amyloid antibodies reduced after corticosteroids therapy. Similarly, it was also described that β-amyloid 1-40, 1-42, tau and phospho-tau-181 proteins rates in the CSF decrease after immunosuppressive treatment, as we observed in this case.

Spectroscopy found a lipid–lactate peak compatible with cell hypoxia. There was hypoperfusion in the inflammatory region, which was compatible with vasculitis. The literature contains little description of CNS vasculitis aspects on spectroscopy or perfusion: case reports show diminished rCBV on perfusion and elevated glutamate–glutamine and/or lactate–lipid peaks on spectrometry in PACNS. To our knowledge, there are no published descriptions of CAA-ri profiles in MRI perfusion. We suggest here the interest of spectrometry and perfusion profiles for the study of CAA-ri.

To our knowledge, the present paper is the first description of dissection or dysplasia aspect associated with CAA-ri. It is unlikely that these two rare events are unrelated in the present case, given the chronology of clinical central and peripheral presentation associated with concomitant brain and vessels imaging abnormalities.

Carotid dissection is a very rare complication of vasculitis and was reported in only few case reports. To our knowledge, only one observation of carotid dissection during the course of PACNS has been previously reported. In the present case, it can be hypothesized that carotid dissection was caused by extensive inflammation of the arterial wall, secondary to deposition of amyloid protein in the wall of cortical and meningeal vessels. A study to determine factors associated with dissection in giant cell arteritis found a higher frequency of active aortitis in patients with dissection, suggesting the role of active inflammation in the occurrence of dissection in patients with vasculitis.

This case report suggests that extra-cranial vasculitis and dysplasia can exceptionally be found in patients satisfying CAA-ri criteria. In future observational studies of CAA-ri patients, dedicated angiography could be performed to determine the frequency, characteristics and clinical significance of arterial abnormalities.

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Ethics and consent to participate
Written informed consent for publication of their clinical details and clinical images was obtained from the patient. A copy of the consent form is available for review by the editor of this journal.

Ethics approval
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