Absence of the Common Carotid Artery: A Rare Vascular Anomaly

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Summary: Although absence of the common carotid artery (CCA) is a rare anomaly, the specific configuration in our case makes it extremely rare. Clinical, angiographic, ultrasonographic, and embryologic correlations of this anomaly are discussed after the case report.

Absence of the CCA is rare. Fewer than 25 cases have been reported in the literature, with only five angiographically proved cases. Only one case had ultrasonographic correlation (1). To the best of our knowledge, in all reported cases, the ECA originated proximal to the ICA, which makes our case an extremely rare occurrence (2–15). Absence of the CCA has no side or sex preferences (5), can occur bilaterally (5, 9), and is asymptomatic unless associated with other conditions. Among reported associations are cervical aortic arch, double aortic arch, right-sided aortic arch with aberrant left subclavian artery, and persistent trigeminal artery (5). Although in our case, the right ICA had no significant stenosis, symptomatic stenoses have been reported in association with this anomaly (11).

Case Report

Bilateral cerebral angiography was performed on a 72-year-old female patient with a medical history of hypertension, hypercholesterolemia, recent silent myocardial infarction, left hemispheric transient ischemic attacks, and a family history of stroke and coronary artery disease. The CCA was absent on the right side, but unlike the usual configuration of separate internal carotid artery (ICA) and external carotid artery (ECA), in our case, the right ICA originated proximal to the right ECA (Figs 1 and 2). No significant stenoses were found on the right side. Other findings were tortuosity of the right ICA at the level of C3–C4 (Fig 2) and hypoplasia of the A1 segment of the right anterior cerebral artery. High grade, short segment ostial stenosis of the ICA was found on the left side. The results of the examination were otherwise unremarkable. The patient then underwent stent placement of the stenotic left ICA. As part of post-stent placement workup, a bilateral carotid Doppler examination was performed, which showed satisfactory peak systolic velocities of the stented left ICA and normal pattern of spectral waveforms for the right ICA and ECA.

Discussion

Angiography

When absence of the CCA occurs on the right side, the ECA usually arises proximally from brachiocephalic artery and the ICA arises distally from the subclavian artery proximal to the origin of the vertebral artery (5). In the presence of subclavian artery stenosis between the ICA and vertebral artery, steal syndrome can occur. When the anomaly occurs on the left side, both the ECA and the ICA arise from the aortic arch, with the ECA proximal to the ICA (5). Practical consequences of this anomaly are technical difficulties in performing brachiocephalic angiogra-
phy. This anomaly can be missed during selective ICA angiography when it is presumed that the catheter tip is introduced into the ICA above the presumed level of CCA bifurcation. Unlike all reported cases, in our case, the ICA arose proximal to the ECA.

Ultrasonography

Ultrasonography is usually the first test with which to detect the possibility of an absent CCA when the sonographer cannot find normal carotid bifurcation in the neck. Intrathoracic bifurcation of the CCA also can cause the same problem. No significant difference has been reported in resistive indices of ICA and ECA in this anomaly and in normal cases, indicating that the resistive indices and spectral waveforms of the ICA and ECA are mainly determined by their supplied organs and not the CCA (1, 16).

Embryology

In a 12- to 14-mm embryo, the ductus caroticus, which connects the third and fourth branchial arches, dorsally involutes and the proximal portion of ECA forms the CCA, the third branchial arch forms the carotid sinus and proximal portion of the ICA, and the fourth branchial arch and the ductus caroticus persists (right) or, alternatively, if the fourth branchial arch involutes and the persistent third branchial arch forms a cervical aortic arch (left) (Fig 3A). In our case, however, it is likely that an extra twisting of the separate ICA and ECA occurred, causing the ECA to be distal to the ICA.

Conclusion

Absence of the CCA is rare. It is usually incidentally found at autopsy or during workup for other clinical problems; however, it may cause significant practical diagnostic and therapeutic challenges. Unlike other reported cases of an absent CCA with the ECA proximal to the ICA, in this case, the ICA arose proximal to the ECA. Reviewing the reported cases of absence of the CCA reveals that the supplied organs of the ICA and ECA and not the presence of the CCA are the main determinants for the shape of resistive indices and frequency of plaque formation and stenosis in these arteries.

References