A Ruptured Anterior Communicating Artery Aneurysm with Infra-optic Course of the Anterior Cerebral Artery: A Case Report and a Short Review

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Abstract

Infra-optic course of the anterior cerebral artery (IOA) is rare and approximately 55 cases of this anomaly have been described. We present a case of a ruptured anterior communicating artery (ACoA) aneurysm arising at the junction between the left IOA and the bilateral A2 segments, at which the right A1 segment was absent. One of the recurrent arteries of Heubner branched off directly from the aneurysmal dome, and was obstructed at aneurysmal neck clipping via an anterior interhemispheric (AIH) approach. In this report, accompanied anatomical variations and surgical approaches for ACoA aneurysms with IOA are reviewed. An IOA is frequently associated with other vascular anomalies, and the origin of functionally important recurrent arteries of Heubner is also variable. Preoperative accurate evaluation of vessel structures and the maximal exposure at surgery are very important. Pterional approach from the ipsilesional side is reportedly to be safe, but interhemispheric approach is also suggested to be effective as to full exposure to recognize the perianeurysmal anatomical structures including potential vessel anomalies.

Keywords: infra-optic course of anterior cerebral artery, anterior communicating artery aneurysm, clipping

Introduction

An infra-optic course of the anterior cerebral artery (IOA) is a rare anomaly, and there have been approximately 55 cases reported1–5 of which about half were associated with cerebral aneurysms including anterior communicating artery (ACoA) aneurysms.6,7 IOA is a developmental disorder of the anterior cerebral artery (ACA) during the embryonic period1–7. The genesis of IOA remains unclear, but this anomaly originates from the internal carotid artery (ICA) at or close to the level of origin of the ophthalmic artery, running beneath the ipsilateral optic nerve and anterior to the optic chiasm, and then turning upward between bilateral optic nerves to terminate at the ACoA. It is also characterized by associated aneurysms and various vascular anomalies. We present a case of a ruptured ACoA aneurysm associated with the IOA, and review previous literatures to discuss surgical strategy based on the anatomical structures.

Case Report

A female in her seventies was transferred to our hospital with severe headache and vomiting lasting for 4 days. The initial head computed tomography (CT) revealed subarachnoid hemorrhage (SAH) of Fisher CT group 3, and three-dimensional CT angiography and digital subtraction angiography (DSA) showed an ACoA aneurysm, of which the maximum diameter was 5.3 mm, associated with the low bifurcation of the left ICA at the level of the origin of the ophthalmic artery, the left IOA filling bilateral ACA territories, and the right A1 aplasia (Figs. 1A and 1B). The anomaly of ACA was also visible on heavily T2-weighted magnetic resonance images (Figs. 1C–1E). The patient was diagnosed with
aneurysmal SAH of Hunt and Kosnik grade 2 and World Federation of Neurological Surgeons grade 2. As endovascular treatment was considered to be inappropriate due to the irregular shape and a broad neck of the aneurysm, elective surgical clipping was performed at 12 days after the onset of SAH when a risk of delayed cerebral ischemia passed. As DSA revealed multiple recurrent arteries of Heubner (RAHs) arising around ACoA (Fig. 1B), anterior interhemispheric approach (AIH) was selected to visualize these perforators clearly. The IOA and bilateral fenestrated A2 were found during microsurgical dissection. After full exposure of the aneurysm and main perforators, it was revealed that one of two RAHs branched off directly from the aneurysmal dome and that a small cortical branch adhered to nearby the rupture point (Fig. 2A). These vessels were interrupted and the ruptured aneurysm was completely obliterated with tandem clipping using 4 Sugita Titanium II clips (17-001-01, 17-001-03, 17-001-81, and 17-001-86; Mizuho Co., Ltd, Tokyo, Japan) to avoid kinking of the bilateral A2 (Fig. 2B). Postoperative magnetic resonance images showed acute infarction in the left caudate head and internal frontal basal cortex owing to the vessels sacrifice (Fig. 2C). The patient had a mild cognitive impairment (modified Rankin scale 3) and was transferred to a rehabilitation facility 28 days after surgery.

Literature Review

The PubMed database was used to search for previously published articles about ACoA aneurysms.
with IOA, which were treated by surgical clipping or coil embolization. The following words were searched: “infra-optic course of anterior cerebral artery,” “carotid–anterior cerebral anastomosis,” or “anterior cerebral artery anomaly” and “cerebral aneurysm.” As a result, 30 cases of ACoA aneurysms with IOA met the purpose of our review and 31 cases including the present case were analyzed regarding their anatomical features, treatment strategies, and outcomes (Table 1).\(^3\)\(^{5,9-34}\) The mean age of patients was 42.5 ± 14.6 (standard deviation) years old. The right-sided IOA was dominant (right, 22 cases; left, 2 cases; and bilateral, 7 cases). According to Wong’s classification, seven cases belonged to type I, 10 cases to type II, 12 cases to type III, one case to type IV, and one case was undetermined due to missing information.\(^2\) As to the treatment, 28 cases of ACoA aneurysms were treated by clipping via a pterional approach (22 cases), an orbitozygomatic approach (one case), or an AIH (four cases including the present case), and the information in one case was unavailable. Three cases were treated by coil embolization with a simple technique in two cases and with a

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Fig. 2 Intraoperative view of the interhemispheric approach before (illustration: A) and after aneurysmal clipping (photograph: B), and postoperative magnetic resonance imaging (MRI: C). The left IOA (single arrow), left A2 segment with fenestration (double arrows), right A2 segment with fenestration (triple arrows), dura mater of the anterior cranial fossa (single black asterisk), optic chiasm (double black asterisks), the left optic nerve (triple black asterisks), a thrombus at the rupture site of the IOA aneurysm (white asterisk), a cortical artery adhering close to the rupture site (single white arrowhead), one of two RAHs branching from the aneurysm (double white arrowheads), and another RAH branching from the supraoptic portion of the IOA (triple white arrowheads) are illustrated (A). One of two RAHs branching from the aneurysm and the cortical artery adhering close to the rupture site are disconnected and the aneurysm is completely obliterated with multiple clipping (B): the marks in the figure are the same as in A. Postoperative diffusion-weighted MRI reveals acute infarction in the left caudate head and a part of the internal capsule and the putamen (C). IOA: infra-optic course of the anterior cerebral artery, L: left, R: right, RAHs: recurrent arteries of Heubner.
Table 1 Cases of ACoA aneurysms with IOA treated with clipping or coiling

| Author (year)    | Age (years) / sex | Rupture state | Side of IOA | Wong’s classification | Associated anomaly                                                                 | Associated aneurysm | Branch from aneurysm | Surgical approach | Endovascular treatment | Postoperative course |
|------------------|-------------------|---------------|-------------|-----------------------|------------------------------------------------------------------------------------|----------------------|----------------------|---------------------|------------------------|---------------------|
| Teal (1973)      | 41 / M            | SAH           | Rt          | III                   | Persistent primitive trigeminal artery, Rt PICA arising from distal cervical ICA | –                    | –                    | –                   | –                      | –                   |
| Nutik (1976)     | 22 / F            | SAH           | Lt          | I                     | Lt AICA arising from cavernous ICA                                                 | –                    | –                    | Lt pterional         | –                      | Uneventful           |
| Kessler (1979)   | 23 / M            | SAH           | Rt          | I                     | –                                                                                  | –                    | –                    | Anterior interhemispheric | –                      | Regrowth, but uneventful |
| Lehman (1980)    | 23 / M            | SAH           | Bilateral   | II                    | Median artery of corpus callosum, PCoA arising from MCA, peripheral moyamoya vessels, aortic coarctation | –                    | –                    | Rt pterional         | –                      | Uneventful           |
| Bernini (1982)   | 50 / F            | SAH           | Rt          | I                     | PCA arising from carotid siphon                                                    | –                    | –                    | Rt pterional         | –                      | Uneventful           |
| Senter (1982)    | 48 / M            | SAH           | Rt          | III                   | –                                                                                  | –                    | –                    | Rt pterional         | –                      | Uneventful           |
| Fujimoto (1983)  | 60 / F            | SAH           | Rt          | II                    | Plexiform network of ACoA                                                         | –                    | –                    | Rt pterional         | –                      | Uneventful           |
| Sasaki (1984)    | 32 / F            | SAH           | Rt          | –                     | –                                                                                  | –                    | –                    | Rt pterional         | –                      | –                   |
| Rosenorn (1985)  | 55 / F            | SAH           | Rt          | III                   | –                                                                                  | –                    | –                    | Rt pterional         | –                      | Uneventful           |
| Klein (1987)     | 43 / F            | SAH           | Bilateral   | II                    | –                                                                                  | Rt IOA aneurysm      | –                    | Rt pterional         | –                      | Uneventful           |
| Bollar (1988)    | 61 / F            | SAH           | Rt          | III                   | –                                                                                  | –                    | –                    | Rt pterional         | –                      | –                   |
| Odake (1988)     | 56 / M            | SAH           | Rt          | III                   | –                                                                                  | –                    | –                    | Rt pterional         | –                      | Rerupture, dead       |
| Takeshita (1991) | 40 / M            | SAH           | Rt          | II                    | Unpaired pericallosal artery, fenestration of VA, duplication of bilateral SCA, Rt OA arising from ICA | –                    | –                    | Anterior interhemispheric | –                      | –                   |
| Onishi (1992)    | 37 / M            | SAH           | Rt          | I                     | Skull dysplasia, short stature                                                    | –                    | –                    | Rt pterional         | –                      | Uneventful           |
| Ladzinski (1997) | 39 / F            | SAH           | Rt          | IV                    | –                                                                                  | –                    | –                    | Rt pterional         | –                      | Transient oculomotor palsy |
| Ogura (1998)     | 58 / F            | SAH, ACoA aneurysm unruptured | Bilateral  | II (bilateral)        | Bilateral MCA aneurysm (Rt, ruptured)                                            | –                    | –                    | Anterior interhemispheric | –                      | –                   |
| Author (year) | Age (years) / sex | Rupture state | Side of IOA | Wong’s classification | Associated anomaly | Associated aneurysm | Branch from aneurysm | Surgical approach | Endovascular treatment | Postoperative course |
|--------------|------------------|---------------|-------------|----------------------|-------------------|-------------------|-------------------|----------------|---------------------|-------------------|
| Spinatto (1999) | 30 / M | SAH, ACoA aneurysm unruptured | Rt | III | – | Rt ruptured MCA aneurysm | – | Rt pterional | – | Uneventful |
| Hilliard (2002) | 30 / F | SAH | Rt | I | – | – | – | – | Coiling (simple technique) | Uneventful |
| Al-Qahtani (2003) | 11 / M | SAH | Rt | I | – | – | – | – | Coiling (simple technique) | IOA occluded, but uneventful |
| Kilic (2005) | 38 / F | SAH | Bilateral | I | Fenestration of ACoA | – | – | Lt pterional | – | Uneventful |
| McLaughlin (2007) | 34 / F | Unruptured | Bilateral | II (bilateral) | Abnormal gyral segmentation | – | – | Lt pterional | – | Uneventful |
| Yurt (2008) | 35 / M | SAH | Rt | III | – | – | – | Rt pterional | – | Uneventful |
| Cheol (2010) | 28 / F | Unruptured | Bilateral | II (bilateral) | – | Lt MCA aneurysm | – | Lt pterional | – | – |
| Wong (2010) | 45 / M | SAH | Rt | III | Median anterior cerebral artery, fenestration of MCA, neither of the ophthalmic arteries from ICA | – | – | Rt pterional | – | Uneventful |
| Turkoglu (2011) | 53 / F | SAH | Rt | III | Rt PICA arising from ICA, Bilateral SCA arising from PCA | Rt A1 aneurysm | – | Rt orbitozygomatic | – | Uneventful |
| Kang (2012) | 59 / F | SAH | Rt | III | Agenesis of Lt ICA | – | – | – | Coiling (stent-assisted technique) | Uneventful |
| Chua (2014) | – | Unruptured | Bilateral | II (bilateral) | – | – | – | Rt pterional | – | Uneventful |
| Kheyreddin (2019) | 44 / M | SAH | Rt | III | – | – | – | Rt pterional | – | – |
| Nandish (2019) | 42 / M | SAH | Rt | II | – | – | – | Rt pterional | – | – |
| present case (2020) | 77 / F | SAH | Lt | III | Bilateral fenestrated A2 | – | Recurrent artery of Heubner | – | – | Mild cognitive dysfunction |

ACoA: anterior communicating artery, AICA: anterior inferior cerebellar artery, ICA: internal carotid artery, IOA: infra-optic course of the anterior cerebral artery, Lt: left, MCA: middle cerebral artery, Oa: occipital artery, PCA: posterior cerebral artery, PCoA: posterior communicating artery, PICA: posterior inferior cerebellar artery, Rt: right, SAH: subarachnoid hemorrhage, SCA: superior cerebellar artery, VA: vertebral artery, Wong’s classification: type I, normal ICA bifurcation with unilateral or bilateral IOA, type II, unilateral or bilateral IOA without supraoptic anterior cerebral artery, type III, unilateral IOA without contralateral A1, type IV, accessory anterior cerebral artery variant with infra-optic course.
Discussion

IOA is a known but rarely reported anomaly. In 1959, Robinson first described this ACA anomaly in an autopsy case. Approximately 55 cases of this anomaly have been described as carotid-ACA anastomosis, anomalous origin of ACA, or IOA. Three major hypotheses have been proposed concerning the development of the IOA. Robinson reported the first case of this anomaly as the persistent enlargement of the presellar arterial plexus including the prechiasmatic branch of the ophthalmic artery, the superior hypophyseal branch of the ICA, and the chiasmal branch of the ACA. Isherwood and Dutton hypothesized this anomaly as the persistence of an anastomotic loop between the primitive dorsal and ventral ophthalmic arteries. Bosma explained IOA as the persistence of the embryonic anastomosis between the primitive maxillary artery and the primitive olfactory artery, which is a precursor of ACA.

Wong et al. reviewed 44 cases of IOA including their cases, and classified the various patterns of A1 and IOA configurations into four types according to the presence of the ipsilateral supraoptic A1 and the contralateral A1; type 1 (12 cases), normal bifurcations of the middle cerebral artery and the ACA with unilateral (10 cases) or bilateral (2 cases) IOA; type II (14 cases), unilateral (8 cases), or bilateral (6 cases) IOA with no ipsilateral supraoptic A1 associated with ICA bifurcation at the level of the origin of the ophthalmic artery; type III (10 cases), unilateral IOA with no contralateral A1; and type IV (1 case), an accessory ACA variant with the infraoptic course. The data of the remaining 7 cases were not adequate. In their series, the frequency of associated cerebral aneurysms was 59% (50% in type I, 79% in type II, and 70% in type III), and ACA aneurysms were the most common (100% in type I-associated aneurysms, 46% in type II-associated aneurysms, and 67% in type III-associated aneurysms). Based on Padget’s and Moffat’s articles, Wong et al. proposed the hypothesis of the development of IOA: that is, a variety of faulty embryogenesis of caudomedial and rostolateral arterial plexuses around the optic vesicle (Padget’s stage 1) could lead to type 1 to 4 IOA configuration during Padget’s stage 3.

We reviewed 31 cases of ACoA aneurysms with unilateral or bilateral IOA including our case that were treated by surgical clipping or coil embolization from published literatures (Table 1). ACoA aneurysms were accompanied by all patterns of IOA according to the Wong’s classification. As to the surgical approach, 23 of 27 cases of ACoA aneurysms were treated by clipping via a pterional approach and an orbitozygomatic approach. The side of a pterional approach was basically the same side as that of IOA, and relatively high-positioned ACoA aneurysms tended to be operated via an orbitozygomatic approach or an AIH including our case. In the present case, complete obliteration of the aneurysm could not avoid the sacrifice of the RAH and resulted in cerebral ischemia in the RAH territory, causing mild cognitive dysfunction. The diverse vascular anomalies around the ACoA associated with IOA may mean highly various branching patterns of functionally important ACA perforators such as the RAH, the hypothalamic artery, and the medial lenticulostriate arteries. Besides, RAH was reported to be multiple in 28% of 2300 cadaveric cases by Bonasia et al. Therefore, it is important to carefully evaluate the perianeurysmal anatomical structures before surgical and endovascular treatment. Surgical clipping via a pterional approach has been reported to be safe, but an AIH is also suggested to be reasonable and effective as to full exposure of the perianeurysmal anatomical structures including potential anomalies of the ACA, ACoA, and its perforators. Alternatively, endovascular treatment of ACoA aneurysms with IOA may be equally feasible depending on their accessibility and morphology, although the importance and limitation of precise detection and preservation of these vessels remain as with surgical treatment.

Conflicts of Interest Disclosure

The authors declare that they have no conflict of interest.

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