Giant Aneurysm of the Azygos Pericallosal Artery: Case Report and Review of the Literature

Cahide Topsakal, M.D.,* M. Faik Ozveren, M.D.,* Fatih S. Erol, M.D.,* Mutlu Cihangiroglu, M.D.,† and Hasan Cetin, M.D.* 
*Departments of Neurosurgery and †Radiology, Firat University, School of Medicine, Elazig, Turkey


BACKGROUND
Pericallosal aneurysms are encountered less than 6.7%, and giant aneurysms among them even less. Giant azygos pericallosal artery aneurysm at the callosomarginal bifurcation is extremely rare, and our case presented herein is the second one. The case is discussed with thorough review of the literature.

METHODS
A 65-year-old woman presented with an extremely rare giant aneurysm on the azygos pericallosal artery manifesting as subarachnoid hemorrhage in World Federation of Neurosurgical Societies Grade 3. Computed tomography (CT), magnetic resonance angiography (MRA), and four vessel angiography revealed a giant azygos pericallosal artery aneurysm associated with a second aneurysm at the left M1.

RESULTS
After recovery to Grade 2, she underwent surgery via the right frontal interhemispheric approach for the azygos artery aneurysm on the 17th day after bleeding. The true dimensions of the aneurysm were greater than indicated by angiography because of partial thrombosis. Trilobulate aneurysm was carefully dissected from the surrounding structures. Postoperative cerebral angiography showed no filling of the clipped aneurysm and preservation of circulation.

CONCLUSION
The treatment of distal anterior cerebral artery aneurysms is often difficult, because of their broad-based irregular configurations and adherence to surrounding tissue, tendency to bleed irrespective of size and the coexistence of other cerebral aneurysms. However, excellent outcomes can be obtained based on thorough preoperative radiologic evaluation, including magnetic resonance imaging (MRI), and correct selection of surgical approach. © 2003 Elsevier Inc. All rights reserved.

KEY WORDS
Azygos pericallosal artery, cerebral aneurysm, cerebral angiography, distal anterior cerebral artery, pericallosal aneurysm.
matoma extending into the lateral ventricles (Figure 1). MRA revealed an aneurysmatic dilatation at the callosomarginal bifurcation of the distal ACA (Figure 2). Four-vessel angiography revealed an azygos distal ACA with a giant aneurysm located at the callosomarginal bifurcation of the azygos pericallosal artery. The aneurysm was broad based, trilobulate with irregular configuration, and $25 \times 15 \times 15$ mm in size. Another aneurysmatic dilatation was observed at the left M2 (Figure 3). Surgery was performed on the 17th day after bleeding, as soon as she recovered to WFNS Grade 2.

Right frontal paramedian craniotomy was performed across the anterior third of superior sagittal sinus and a U-shaped dural flap reflected to the left side based on the sinus to give access to the interhemispheric fissure. The self-retaining retractors were gently interplaced in the fissure between the bridging anterior and middle frontal veins. We proceeded along the shallow falx down to the lower edge by suctioning the clots. Clearing of the arachnoid adhesions between cingulate gyri exposed the single distal ACA (pericallosal) and two ramifications on both sides in the callosal cistern. We proceeded proximally, and the trilobulate giant aneurysm emerged at the callosomarginal bifurcation of pericallosal artery. The actual dimensions of the aneurysm were greater than the angiographic dimensions. Each lobule extended in a different direction, one adherent to the left cingulate gyrus at the

1 Axial computed tomography scan showing the pericallosal hematoma predominantly on the left side with widespread subarachnoid and intraventricular hemorrhage.

2 MRA revealing the azygos anterior cerebral artery associated with an aneurysm at the distal end (white arrow). A second aneurysm is seen on the left middle cerebral artery (double white arrowheads).
bleeding point, and another fully filled with thrombus extending to the base. The parent arteries were exposed proximal to aneurysm. The neck was broad based. After clearing the adhesions to the surrounding tissue and preparing the neck, we placed two Yasargil aneurysm clips without compromising the parent arteries.

No neurologic deficit developed after the surgery, and postoperative carotid angiography showed good circulation in the parent arteries without filling of the aneurysm (Figure 4). The patient recovered within hours, and did not develop any neurologic deficit all through the first month, but subsequently died of pulmonary embolus because of thrombophlebitis.

**Discussion**

Generally, pericallosal (distal ACA) aneurysms are rare, representing only 2 to 6.7% of all intracranial aneurysms [26,39,54,63,77,80]. Giant intracranial aneurysms (>2.5 cm) account for 3 to 13% of all intracranial aneurysms [3,11,20,41,57,69,76,79,80].
Giant pericallosal aneurysms are very rare and somehow are not mentioned in the large series of either giant aneurysms [66,69,76,77,80] or pericallosal aneurysms [23,26,54,63,76,81]. In the literature there have been only 13 cases of giant pericallosal artery aneurysms [11,14,20,21,23,45,52,55,58,59,64,73], and our case is the 14th. Association of saccular aneurysm with azygos pericallosal artery is 13% (Table 1) [1,5–7,13,14,16–20,24,27,29,31,32,35–43,46,47,51,53,57,61,62,70,72,77–79]. Giant azygos pericallosal artery aneurysm is extremely rare [19,20,46,62,78], particularly at the callosomarginal bifurcation, and out of 6 giant azygos pericallosal aneurysms, ours is the second case reported at this location.

Various degrees of fusion of the ACAs have been described, from limited or punctiform contact to a common trunk [2]. The incidence of the true azygos distal ACA (Baptista Type 3), from which all major branches are given off to both hemispheres, ranges
from 0 to 5% [2,24,32,42,43,79]. True azygos distal ACA must be distinguished angiographically from the more common (incidence 12%) so-called azygos distal ACA (Baptista Type 1), or bihemispheric pericallosal artery, in which both ACAs are present, one being hypoplastic and most of the branches to both hemispheres arising from the other. A carotid angiography with contralateral compression, rather than a bilateral carotid angiography, is more appropriate for identification of an azygos artery [5,47,51,78], allowing discrimination from the bihemispheric variant and display of the median anterior arterial vascularization and all components in greater detail [5]. The clinical importance of a correct diagnosis of azygos distal ACA arises from the presence of a single trunk, which supplies blood to both hemispheres [32,42]. Occlusion of this single trunk as a result of thromboembolic disease, vasospasm, or surgical error has disastrous effects [6,32,42]. There are other types of vascular anomalies, such as Baptista Type 2, a triple ACA with the accessory ACA or corpus callosal median artery

<table>
<thead>
<tr>
<th>AUTHOR</th>
<th>YEAR</th>
<th>NO. OF CASES</th>
<th>LOCATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laitinen and Snellman</td>
<td>(1960)[39]</td>
<td>3</td>
<td>distal end</td>
</tr>
<tr>
<td>Pool and Potts</td>
<td>(1965)[57]</td>
<td>3</td>
<td>distal end</td>
</tr>
<tr>
<td>Le May and Gooding</td>
<td>(1966)[42]</td>
<td>1</td>
<td>proximal end</td>
</tr>
<tr>
<td>Takahashi</td>
<td>(1967)[29]</td>
<td>1</td>
<td>unknown</td>
</tr>
<tr>
<td>Okawara</td>
<td>(1968)[53]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Kitamura</td>
<td>(1970)[36]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Ishii et al</td>
<td>(1974)[27]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Kinoshita and Matsukado</td>
<td>(1975)[35]</td>
<td>2</td>
<td>1 distal, 1 middle</td>
</tr>
<tr>
<td>Yamada</td>
<td>(1975)[29]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Yamagata</td>
<td>(1975)[29]</td>
<td>1</td>
<td>proximal end</td>
</tr>
<tr>
<td>Katz et al.</td>
<td>(1978)[32]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Fankhauser and Zander</td>
<td>(1978)[13]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Nukui</td>
<td>(1978)[29]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Dennis</td>
<td>(1979)[29]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Egami</td>
<td>(1979)[29]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Kondo et al</td>
<td>(1979)[38]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Huber et al</td>
<td>(1980)[24]</td>
<td>2</td>
<td>distal end</td>
</tr>
<tr>
<td>Fujimoto et al</td>
<td>(1981)[16]</td>
<td>1</td>
<td>rt distal callosomarginal bifurcation</td>
</tr>
<tr>
<td>Nizuma et al</td>
<td>(1981)[51]</td>
<td>2</td>
<td>1 distal, 1 middle</td>
</tr>
<tr>
<td>Fukawa et al</td>
<td>(1982)[17]</td>
<td>1</td>
<td>unknown</td>
</tr>
<tr>
<td>Mochizuki et al</td>
<td>(1982)[47]</td>
<td>1</td>
<td>proximal end</td>
</tr>
<tr>
<td>Benedetti and Curri</td>
<td>(1983)[5]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Yasargil</td>
<td>(1984)[79]</td>
<td>2</td>
<td>distal end</td>
</tr>
<tr>
<td>Lau et al</td>
<td>(1984)[40]</td>
<td>1</td>
<td>distal end-callosomarginal bifurcation</td>
</tr>
<tr>
<td>Hayashi et al</td>
<td>(1985)[20]</td>
<td>2</td>
<td>distal end (giant)</td>
</tr>
<tr>
<td>Abe et al</td>
<td>(1985)[1]</td>
<td>2</td>
<td>1 distal, 1 proximal 4 distal, 1 middle, 1 callosomarginal</td>
</tr>
<tr>
<td>Kaneko et al</td>
<td>(1985)[29]</td>
<td>7</td>
<td>bifurcation, 1 proximal</td>
</tr>
<tr>
<td>Yamagami et al</td>
<td>(1986)[78]</td>
<td>1</td>
<td>distal end-callosomarginal bifurcation (giant)</td>
</tr>
<tr>
<td>Kobayashi et al</td>
<td>(1986)[37]</td>
<td>1</td>
<td>proximal end</td>
</tr>
<tr>
<td>Kato</td>
<td>(1988)[31]</td>
<td>1</td>
<td>unknown</td>
</tr>
<tr>
<td>Harada et al</td>
<td>(1987)[18]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Wisoff and Flamm</td>
<td>(1987)[77]</td>
<td>2</td>
<td>unknown</td>
</tr>
<tr>
<td>Schick and Rumbaugh</td>
<td>(1989)[61]</td>
<td>1</td>
<td>distal end</td>
</tr>
<tr>
<td>Lightfoot et al</td>
<td>(1989)[43]</td>
<td>1</td>
<td>proximal end</td>
</tr>
<tr>
<td>Mishima et al</td>
<td>(1990)[46]</td>
<td>1</td>
<td>distal end (giant)</td>
</tr>
<tr>
<td>Cinnammon et al</td>
<td>(1992)[7]</td>
<td>1</td>
<td>distal end-trifurcation</td>
</tr>
<tr>
<td>Hashizume et al</td>
<td>(1992)[19]</td>
<td>1</td>
<td>distal end (giant)</td>
</tr>
<tr>
<td>Traynelis and Dunker</td>
<td>(1992)[72]</td>
<td>1</td>
<td>distal end (3 aneurysms)</td>
</tr>
<tr>
<td>Zderkiewicz et al</td>
<td>(1992)[82]</td>
<td>1</td>
<td>not available</td>
</tr>
<tr>
<td>Suzuki et al</td>
<td>(1998)[70]</td>
<td>1</td>
<td>distal end-supracallosal</td>
</tr>
<tr>
<td>Present case</td>
<td></td>
<td>1</td>
<td>callosomarginal bifurcation (giant)</td>
</tr>
</tbody>
</table>
arising from the anterior communicating artery (AcoA) (9.6–13%) [2,79]. Baptista Types 2 and 3 are frequently observed at one of the stages of embryonic development, which involves either duplication of the common trunk or involution of the median branch [32,42,71]. Type 3 may result from the persistence of the artery of the corpus callosum with reabsorption of the two ACAs or arise from the direct fusion of the two arteries (Figure 5). Azygos pericallosal artery associated with midline malformations of the central nervous system such as agenesis of corpus callosum, holoprosencephaly [32,42,59], and association with arteriovenous malformation [18,20,24,57,59,77] suggest that the arterial anomaly might be related to embryological development. However, our patient had no associated midline malformations. Association of azygos pericallosal artery with berry aneurysms may be as high as 41.1 to 71% of the azygos pericallosal arteries [24,42,51,57,79].

Aneurysms not associated with azygos artery usually occur at the level of the genu of the corpus callosum, where the pericallosal artery sharply arches posteriorly, i.e., the callosomarginal bifurcation 69 to 78% [21,39,77,80] and then the frontopolar artery, whereas very proximal and distal locations are rare, and frontobasal is the rarest [11,21]. Aneurysms associated with azygos distal ACA are usually located at the distal end of the azygos pericallosal artery [7,16,24,32,35,36,38,39,42], generally at the callosomarginal bifurcation [6,16,35,51], as in our case, or trifurcation. The proximal location is rare [42]. Associated vascular anomalies and the variations in the circle of Willis [33,67] with marked effects on circulatory dynamics may be involved in the formation of the aneurysm [37,42,51,54,61,62]. Geometric changes from the symmetrical to asymmetrical arterial junction develop higher shear stress than critical values and the stagnation point at the junction [75]. The branching geometry determines significant increase of shear stress at the branching sites of cerebral arteries in patients with aneurysm of the DACA compared to normal controls [60]. The burden of blood flow from the
bilateral A1 on the end of anomalous azygos artery is responsible for aneurysm development [51].

In an arterial network, the pressures and shear stresses that develop along the outer wall of a curved artery, and the apex of an arterial bifurcation create a hemodynamic state that promotes saccular aneurysm formation [15,60]. Additionally, geometrical relation between aneurysm and parent vessels is the primary factor governing the intraaneurysmal flow pattern [68]. For uneven branch flow, flow activity inside the aneurysm and the stresses acting on the aneurysmal wall increase with increasing bifurcation angle, and may cause aneurysm rupture [44]. It is less susceptible to thrombus formation, because both sides of the aneurysm neck are subjected to increased states of hemodynamic stress, one arising from the pressure and the other from the shear on the side of the larger and the smaller daughter branches, respectively [15]. On the other hand, for evenly distributed branch flow, as is valid in our case, the intraaneurysmal flow is sluggish and therefore, prone to thrombosis for all studied bifurcation angles [44]. In our case, the broad-based and giant aneurysm was trilobulate, with every lobule oriented in a different direction. The largest lobule was directed to the left and embedded into the left cingulate gyri, and full of thrombus, so the dimensions were underestimated on MRA and DSA [58,74].

Giant pericallosal aneurysms are encountered very seldom, possibly because of the tendency to early rupture and treatment before enlargement [63]. Giant aneurysms tend to thrombose and are invisible on angiography [58]. Therefore, MRI is superior to CT and DSA for evaluating the characteristics of aneurysms and provides additional information about the condition of the surrounding brain [45,48,59]. MR imaging shows giant aneurysm as a large, round, nonhomogeneous intensity area, with a patent lumen appearing as signal void and thrombosis as hyperintense areas. More recently thrombosed aneurysm appears as more homogeneous with a density similar to that of coagulated blood and therefore, may be indistinguishable from hematoma on CT, allowing the membrane of the hematoma to mimic a tumor [40,45]. In our case, because we could not provide intraoperative MRI, we were confronted with problems all through the surgery because of the underestimation of its exact dimensions. Hematoma is seen in 60.9 to 67% of cases of distal ACA aneurysm, of which half are callosal hematoma, and more than half of these cases are associated with intraventricular hematoma [28,80]. In our case, the callosal hematoma extended into the lateral ventricles.

Patients with distal ACA aneurysm have multiple aneurysms at rates of 5–55% [21,54,63,77,80]. Most cases (68–83%) are located on the middle cerebral artery and at the contralateral ACA (mirror aneurysm) [39,48,50,65,77,80], and 10.4% on the AcoA [54]. In our case, a second aneurysm was found on the left M2.

In general, distal ACA aneurysms have a worse prognosis than those of other sites of the anterior circulation [63] since they are fragile and rupture prematurely during exposure, resulting in a higher morbidity than expected from the angiographic appearance and location [8]. Patients with distal ACA aneurysm associated with multiple aneurysms suffer bleeding in 30 to 50% of cases [25]. Bleeding occurs irrespective of their size because of the lack of resistant arachnoid membranes at the level of the pericallosal cisterns [54,63]. Distal ACA aneurysm presents special difficulties for the surgeon because of the association of the azygos ACA and multiple aneurysms, including the narrow working space in the interhemispheric fissure and the callosal cistern, dense adhesions between the cingulate gyri, a broad-based and/or a sclerotic neck in the aneurysms, difficulty in identifying the parent artery, and the fixed dome on the pial layer limiting lateral retraction of the right hemisphere to a few millimeters [80]. We experienced similar difficulties during surgery [21,48,54,80]. Surgery of aneurysms at the origin of the callosomarginal artery is technically difficult, because this portion of the ACA lies beneath the corpus callosum. Cingulate resection for pericallosal aneurysms [22] or partial callosal resection, either along the commissure or genu [39], is recommended to gain control of the ACA proximal to the origin of callosomarginal artery without causing disconnection syndrome [9,45,72]. Lateral right frontal approach with amputation of the frontal pole avoids traumatic maneuvers on the medial surface of the frontal lobes [5]. Good results were obtained with anterior frontal trephine and interhemispheric dissection [34]. We favored unilateral interhemispheric exposure [8,18,20,70] to treat the pericallosal aneurysm and left the other aneurysm for further operation to reduce the risk of mortality [8]. We approached from the distal to proximal direction, not to confuse the parent arteries.

Handling giant pericallosal aneurysms are rare, particularly in association with azygos pericallosal artery. Preoperative difficulties include the tendency to bleed irrespective of size and invisibility on angiography, and surgical difficulties include the broad base and adherence to the surrounding tissues, which make clipping cumbersome. Preoperative radiologic examinations are extremely impor-
tant for good surgery. They include examinations not only of the internal shape on angiograms, but also the outer shape of aneurysm demonstrated by MRI, because inside of a large aneurysm partial thrombosis, like in this case, is frequent. Proper preoperative evaluation and surgical approach with the use of microsurgical techniques and minimal retraction of the brain and preservation of the bridging veins will reduce the risk of postoperative mortality and morbidity to an acceptable level [63].

REFERENCES

34. Keogh AJ, Sharma RR, Vanner GK. Partial callosal
74. Turtz A, Allen D, Koenigsberg R, Goldman HW. Non-


COMMENTARY

The authors report the successful surgical management of a patient with a ruptured giant pericallosal artery aneurysm associated with an azygos pericallosal artery. This is a very unusual diagnosis, and the authors’ illustrations as well as artistic renderings point out the relevant anatomy very nicely. As the authors note, MRI studies are invaluable in determining whether portions of an unusual aneurysm are thrombotic or not. This information can be extremely beneficial, as one would typically be planning for periods of temporary arterial occlusion. Giant aneurysms of the anterior cerebral branches are particularly difficult because the parent vessel is so small. If the aneurysm is thick walled, it can be extremely difficult to leave enough lumen for orthograde flow through the parent vessel. Intraoperative angiography can be quite helpful in this regard.

H. Hunt Batjer, M.D., F.A.C.S.
Department of Neurological Surgery
Northwestern University
Chicago, Illinois

This is a very good case report of the treatment of a patient with a very rare lesion, a giant aneurysm of an azygos pericallosal artery. The authors review their evaluation and treatment of this patient and should be congratulated for their surgical result despite the patient’s ultimate death from nonsurgical causes.

They provide a thorough review of the relevant anatomical, biophysical, and epidemiological factors relevant to these lesions. In particular, their discussion of the anatomical variants of the distal anterior cerebral arteries is excellent.

The high incidence of multiple aneurysms in these patients fits currently held hypotheses that shear stress plays a major role in the origin of these and other berry aneurysms. Perhaps more interesting is the fact that giant aneurysms are rare in this setting, where there is almost certainly high shear. This low incidence of giant aneurysms supports the idea that while shear is important in aneurysm initiation, other mechanisms are more important in aneurysm growth.

Phillip Dickey, M.D.
Department of Neurosurgery

Purushothaman Kailasnath, Ph.D.
Department of Diagnostic Radiology
Yale University

The campaign of prosecutions of nursing homes for quality will probably spill over to hospitals and physicians’ practices, according to Jim Sheehan, Assoc. U.S. Attorney for the Eastern District of Pennsylvania (Medicare Compliance Alert 8/4/03).

—AAPS News, Volume 59, No. 9, September 2003