MULTIPLE INTRACRANIAL VASCULAR ANOMALIES IN A PATIENT WITH STROKE – A CASE REPORT AND SHORT REVIEW OF LITERATURE

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INTRODUCTION

Non-saccular intracranial aneurysms are independent vascular entities, set apart from their saccular counterparts by different pathogenesis, localization, natural history and potential treatments. Flemming et al. first classified non-saccular aneurysms into 3 separate groups: fusiform aneurysms (FA), dolichoectatic aneurysms (DEA) and transitory aneurysms (TA) (1).

Fusiform aneurysms are defined by a diameter of 1.5 times that of the parent vessel, without an imagistically demonstrated neck, involving a portion of an arterial segment (1). Etiological correlations have been demonstrated with dissection, atherosclerosis, collagen and elastin metabolism disorders and infection (2). In a study by Day et al. (3) regarding middle cerebral artery (MCA) FA, a sequence of aneurysm formation and evolution has been imagined, with a central role residing in intramural dissection (3). FA comprise 3-13% of all intracranial aneurysms (2). The vertebrobasilar (VB) system has a predisposition for developing FA (1).

Dolichoectasia (DE) or dolichoectatic aneurysms are defined by uniformly dilated arteries associated with marked elongation and tortuosity (1,4). In a review of literature on DE, Gutierrez et al. identified the posterior circulation and particu-
larly the basilar artery (BA) as the most common site for the dolichoectatic process (55.5% of all cited cases affecting only the VB system) (4). Reported prevalence for DE is between 0.13 and 18% (4). Currently there is no imagistic gold standard for diagnosing DE; however, Smoker et al. (5) has suggested a series of criteria for basilar artery DE (BADE), widely utilized, pertaining to diameter, height of bifurcation and laterality compared to the clivus (5).

TA, the third variety of non-saccular aneurysms (1), refer to a superimposed focal dilatation on a previously ectatic vascular segment.

In a classification by Mizutani et al. regarding fusiform and dissecting aneurysms, FA are characterized by localized fragmented internal elastic lamina (IEL) and intimal thickening (6). Similarly, DE vessels present a disrupted IEL and atrophy of the muscle layer, widely believed to be a maladaptive vascular remodeling response to chronic arterial hypertension (4,7,8). FA and DE are associated with concomitant abdominal aortic aneurysm or ectasia and intracranial saccular aneurysms (1,8,9), which suggests a possible common pathogenesis.

Cerebral infarction and transient ischemic attack (TIA) are the most common initial symptomatology (27-40.4%), followed by compressive symptoms and subarachnoid hemorrhage (1,10).

Treatment for non-saccular aneurysms varies: if asymptomatic or in the case of initial ischemic presentation, conservative treatment is elected (2,3). Otherwise, an open microsurgical conduit might be employed (clipping wrapping, bypass (2,4)), along with endovascular techniques (coiling, parent vessel occlusion, flow-diversion (11,12)). In a recent meta-analysis on vertebrobasilar FA and DEA, natural history comprised a poor prognosis (10). Globally, non-saccular aneurysms progressed over time in nearly half the cases studied (1,13). The most frequent cause of mortality is ischemic stroke (1,10), with a high recurrence rate (1,10,13).

Encompassing most of the described vascular anomalies, we report a case of an aged male patient presenting with a cerebral ischemic event. This case report was prepared following the CARE guidelines (14).

**CASE REPORT**

We present the case of a 63-year-old male patient, with a history of arterial hypertension and dyslipidemia, who was admitted to our clinic for acute ischemic stroke. Neurologic examination revealed right-sided hemiparesis, right central facial palsy, mixt aphasia and an up-going plantar reflex.
on the right. Hospital presentation was under 2 hours from the debut of symptoms.

The emergency CT scan (not shown) revealed a spontaneously hyperdense, fusiform aneurysm at the level of the left posterior cerebral artery (PCA) (1.22 cm by 1.94 cm transverse and longitudinal diameters) (Fig. 1a); contrast enhanced-CT revealed another 2 aneurysms at the level of the right PCA (Fig 1b) and one at the level of the MCA. Intravenous alteplase administration was therefore not employed. Ultrasonography of cervico-cerebral vessels described normal flow speeds in all arteries and several atherosclerotic and calcified plaques at the level of common carotid artery bifurcation and internal carotid artery origin, on the right side.

One-week follow-up CT scan revealed an ischemic, subacute hypodensity affecting the left occipito-parahypocampal and thalamic regions, consistent with an ischemic stroke in the deep left PCA territory (Fig. 1).

Furthermore, based on Smoker et al. criteria (5) (Table 1), we established the diagnosis of VBDE (Fig. 2), in addition to the aneurysms previously identified.

### TABLE 1. Smoker et al. criteria (5)

<table>
<thead>
<tr>
<th>BA diameter &gt;4.5 mm at the level of midpons</th>
<th>Laterality of BA</th>
<th>Height of BA bifurcation</th>
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<tr>
<td></td>
<td>1. Midline throughout</td>
<td>0. At or below the dorsum sellae</td>
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<tr>
<td></td>
<td>2. Medial to the lateral margin of the clivus</td>
<td>1. Within suprasellar cistern</td>
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<tr>
<td></td>
<td>3. Lateral to the lateral margin of the clivus</td>
<td>2. At the 3rd ventricle floor</td>
</tr>
<tr>
<td></td>
<td>4. At the level of the cerebello-pontine angle</td>
<td>3. Indenting and elevating the floor of the 3rd ventricle</td>
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Cerebral angiography (Fig. 3) revealed two aneurysmal dilatations at the level of the P2 segment of the right PCA, along with VBDE and absence of contrast at the level of the left PCA (thrombosed aneurysm). Furthermore, the left common carotid artery stemmed from the innominate artery (not shown).

The thrombosed aneurysm was not amenable to interventional treatment, nor did the other vascular...
anomalies present an indication of interventional therapy. Therefore, the patient received conservative treatment with anti-platelet medication and high-dose statin. The patient’s condition gradually improved and he partially regained the mobility of his right upper and lower limbs, named objects, formulated simple sentences and stood upright by his own at bedside. Upon discharge he was included in a neuro-rehabilitation program and clinical and imagistic follow-up plan.

**DISCUSSION**

Our case revealed multiple vascular anomalies in a male patient presenting with ischemic stroke. Concerning DE, the diagnosis was based on the aforementioned Smoker et al. criteria (5). In the published literature the application of these criteria resulted in a prevalence of 7.7-17% in series of stroke patients (7). More recently, in a multiethnic cohort of 718 stroke-free patients, the authors proposed a novel method for diagnosing intracranial artery DE (IADE): total cranial-volume adjusted artery diameter (9). Resulting prevalence was 4.7% for BADE (vs. 15.5% by using the previous criteria, with a cutoff used for defining BADE of 5.1 mm).

Concerning the posterior circulation, PCA aneurysms are rare, with an estimated prevalence of 1% of all intracranial aneurysms (3,15,16); this, corroborated with a reduced prevalence of the fusiform subtype resulted in several cases of PCA FA being documented (3,15).

Multiple literature reviews have demonstrated a constant correlation of non-saccular aneurysms with several risk factors: old age, male gender and arterial hypertension. In addition, a statistical link has been identified between IADE and cerebral small vessel disease and other ectatic vascular anomalies. In a recent meta-analysis including 877 patients with VB non-saccular aneurysms, mean age was between 55-75 years, 72% were male participants and 67% had arterial hypertension (10). In a review by Gutierrez et al. of 14 studies on VBDE (4), mean age was 63 years, 64% were male and 66% were hypertensive.

In a review by Brutto et al., patients with stroke and IADE had lacunar infarctions, severe leukoaraiosis and microbleeds in 36-42% of cases compared to patients with stroke, but without IADE (17-19%) (7). In a series by Flemming et al., 18% of patients with non-saccular aneurysms had an abdominal aortic aneurysms and 10% had concomitant intracranial saccular aneurysms (1). In 22 cases of PCA aneurysms, Beute et al. identified 12 cases with concomitant intracranial vascular anomalies (15). These associations suggest a common pathogenetic pathway, with consequences on different vascular territories. Our patient presented all the previously mentioned risk factors, as well as small

**FIGURE 3. Angiography of the VB system: oblique (a.) and anteroposterior (b.) views, showing VBDE, two (fusiform and saccular) right PCA aneurysms (yellow arrows), lack of contrast at the level of the left PCA P2 segment (white arrow); in addition, a probable transitional aneurysm can be observed at the level of the BA (green arrow).**
vessel disease and two saccular aneurysms. However, he was not further investigated concerning a potential aortic abnormality.

Our patient presented with an initial ischemic event, caused by left PCA aneurysm thrombosis. In a study by Flemming et al. (1), ischemic stroke and transient ischemic attack were the most frequent clinical presentations (27%), followed by compressive symptoms (22%); however, most non-saccular aneurysms (DE in particular) were incidental imagistic findings (1). Ischemic symptoms were the initial presentation in almost 31% of FA in the series by Day et al. (3) and remained the major initial symptomatology in a recent meta-analysis (40.4%) (10). Day et al. (3) proposed a process consisting of several steps leading to spontaneous FA formation. Pathogenesis revolves around intramural hemorrhage following arterial dissection. The authors considered subintimal dissection as being responsible for the progressive luminal narrowing altering local hemodynamics and creating a thrombogenic milieu.

VB non-saccular aneurysms have, in general, a poor natural history (1,10). In a series of 159 patients, almost one third of deaths were related to ischemic events. The 1-, 5- and 10-year risk of ischemic stroke after diagnosis was 2.7%, 11.3% and 15.9%, respectively. Predictors of cerebral ischemia due to VB non-saccular aneurysms were symptomatic aneurysms at presentation, TA and prior ischemic events. Nearly half of the cases studied demonstrated diameter growth at follow-up (1).

In a series of 156 patients with DE by Passero and Rossi et al. (13), with a mean follow-up of 11.7 years, VBDE progressed in 43% of the patients. Cumulative risk of a first recurrent stroke was 56% at 10 years and 79% at 15 years. Stroke was the most common cause of death, accounting for 40% of the deaths in this series.

In a recent meta-analysis (10), the authors considered FA and DEA as separate vascular entities, due to different natural history: FA had a superior rupture rate (3%/year vs. almost no risk) and a superior growth rate (12%/year vs. 3%/year for DEA).

Taking into account all concomitant anomalies, our patient mandates a thorough clinical and imagistic follow-up, focusing on both cerebral and extra-cerebral sites.

In our patient treatment was conservative with anti-platelet agent, statin and efficient control of blood pressure; this was due to an initial ischemic event and to the lack of symptomatology concerning the other vascular incidental findings. Intravenous infusion of alteplase was not attempted owing to the multiple concomitant intracranial vascular anomalies. In 22 cases of FA treated by Park et al. with different therapies (wrapping, clipping, resection, proximal occlusion, bypass and combinations of these), the authors concluded that asymptomatic aneurysms as well as stenotic or occluding lesions should be treated conservatively (2). However, in a review on DE, the authors conclude that there is no specific treatment and no controlled trial attesting efficacy of either aspirin or warfarin (4).

In a series of 102 MCA FA (3), the authors advocate for maximal interventional treatment (occlusion of parent vessel with trapping of the aneurysm, resection and vascular bypass). Moreover, endovascular therapies are increasingly utilized, with Ciceri et al. and Taqi et al. successfully treating a small series of mostly saccular PCA aneurysms by means of endovascular coiling (11,16); the efficiency of these methods was also confirmed in a review by Ayad et al. (12) for posterior circulation FA (proposing flow-diversion in addition to stenting and coiling).

**CONCLUSIONS**

Non-saccular aneurysms present a different clinical behavior and challenge compared to saccular aneurysms, frequently leading to ischemic events. Association of VBDE and FA with concomitant saccular aneurysms may orient towards a common vessel wall pathogenetic mechanism within different territories, mandating thorough imagistic investigation (cerebral and aortic). Owing to a poor natural history of non-saccular aneurysms, patients require periodic clinical and imagistic follow-up.

**Consent**

Signed consent for the publishing of the patient’s clinical and imagistic data was obtained from a close relative.
REFERENCES


