Cranial arterial dolichoectasia (CADE) is angiopathy that is characterized by dilation and elongation of cranial arteries. The prevalence of dolichoectasia has been estimated to be 0.05%–0.06%.1,2 The main etiologic factors of dolichoectasia are atherosclerosis, defects or destruction of the internal elastic, and hypertension.2-5 Dolichoectasia preferentially involves the intracranial vertebral and basilar arteries. Involvement of carotid arteries and branches of the carotid arteries is relatively rare.6 Dolichoectasia is frequently associated with ischemia and rarely with intracranial hemorrhage.9 Intracranial hemorrhages may be caused by the rupture of associated aneurysm, rupture of perforating blood vessels originating from ectatic arteries, or dissection of an ectatic artery.2,7,8 Dissections of the cranial arteries are sometimes misinterpreted as duplications or fenestrations of these arteries. We present a case of vertebral, basilar, and bilateral carotid dolichoectasia associated with an unusual type of dissection of the petrous segment of the internal carotid artery (ICA) revealed by conventional angiography.

Case Report
A 58-year-old male smoker, with arterial hypertension, was admitted to the hospital because of a sudden onset of right-sided weakness and pronounced dysphasia. The blood pressure on admission measured 150/100 mmHg. Emergency CT showed intracerebral hematoma localized in the left thalamus, the posterior limb of the internal capsule, and the left lenticular nucleus, as well as the extension of the hematoma into the left lateral ventricle (Fig 1A). The same study showed fusiform dilation of the supraclinoid segments of both ICAs. The finding was more pronounced on the right side. The supraclinoid segment of the right ICA measured 9 mm and the supraclinoid segment of the left ICA measured 5 mm, whereas the diameter of the basilar artery measured 7 mm (Fig 1B, -C). The bone carotid channels appeared to be normal (Fig 1D). The patient was referred for angiographic evaluation. The aortic arch was slightly elongated and dilated. The branches originating from the aortic arch were moderately tortuous and had normal lumen. The left ICA was tortuous and had irregular lumen with 2 fusiform dilations localized, respectively, distally to the bifurcation of the common carotid artery and proximally to the inlet of carotid canal. From this point the left internal carotid artery had 2 lumena that coursed parallel through the carotid canal (Fig 2). Both limbs had smooth contours but slightly irregular, partly fusiform dilated lumen. These limbs were reconnected at the level of the C3 segment. The supraclinoid segment of the ICA was fusiform dilated. The right ICA was dilated distally to the bifurcation of the common carotid artery and distally to the origin of the ophthalmic artery (Fig 3). This fusiform widening involved supraclinoid segment up to the bifurcation of the ICA. Anterior and middle cerebral arteries had normal lumena and branching on both sides. The basilar artery was tortuous and moderately dilated (Fig 4). The branches of vertebral and basilar arteries displayed slightly reduced lumen but normal pattern of branching.

The patient was treated conservatively and referred to rehabilitation. Symptoms related to the bleeding have partly regressed, and 3 months after discharge the patient is doing well.

Discussion
CADE is an angiopathy that is characterized by dilation, elongation, and tortuosity of the brain arteries, most frequently distal vertebral arteries, the basilar artery, or distal ICAs.2,4 Because the dilation seems to be the most important feature, the condition is now referred to as dilative arteriopathy.6 Dolichoectasia that involves both the vertebrobasilar and carotid systems is rare entity and is seen in only 0.5% of patients suffering a first cerebral infarction caused by this angiopathy.9 Despite the observation that approximately 1 of 8 imaged brains has some increase in the length and diameter of intracranial arteries, the prevalence of dolichoectasia is relative low and varies between 0.05% and 0.06% depending on the source.1,2,6 According to Greenfield’s Pathology, a basilar artery lumen >4.5 mm is considered to be ectatic.10 Some authors, however, emphasize that there are no generally accepted quantitative criteria for dilative arteriopathy.6,11,12 Extremely dilated and elongated intracranial arteries are referred to as fusiform or giant fusiform aneurysms. According to Drake and Peerless, large S-shaped aneurysms of the vertebrobasilar system in a patient with hypertensive arteriopathy have been known as the classic form of the giant fusiform aneurysm.17 In our case, the process was bilateral and asymmetrical, involving extracranial and intracranial segments of both ICAs and basilar artery. The lumena of all these arteries were >4.5 mm but did not fit the criteria for fusiform or giant fusiform aneurysm.

Dolichoectasia is commonly seen in patients with advanced cerebral atherosclerosis and arterial hypertension.2-5 Besides atherosclerosis and hypertension, there are several factors and conditions that have been implicated in the pathogenesis of dolichoectasia, including a defect or destruction of the internal elastic lamina, Ehlers-Danlos syndrome,14 Marfan syndrome,15 and tuberous sclerosis.16 The significance of thin-
ning and disruption of internal elastic lamina in the process of elongation and dilatation of intracranial arteries has been emphasized by several authors. The results of recently published studies indicate that the relationship between these pathologic changes and atherosclerosis in the pathogenesis of dolichoectasia remains unclear. The same studies clearly showed that dilatative arteriopathy was associated with increasing age, male sex, arterial hypertension, and history of myocardial infarction. In our case, the patient is a smoker and has been treated for hypertension since his 30s. The pathologic background of the dolichoectasia in this patient cannot be determined on the basis of available image material. The shape of the arterial lumen with dilation, elongation, and smooth irregularities of the contours strongly suggest atherosclerotic changes of the wall. The possible presence of thinning and discontinuity of internal elastic lamina, as well as atrophic or fibrotic tunica media that could have contributed to the dilation and elongation of arteries, however, cannot be excluded.

There are several clinical manifestations of the dolichoectasia: compression on cranial nerves or brain stem, obstructive hydrocephalus, and ischemia. Ischemic lesions may possibly be caused by emboli derived from thrombosis or fragments of plaques in the walls of the enlarged arterial segment. The occurrence of spontaneous intracranial hemorrhages is less common than that of ischemia in patients with dolichoectasia. Intracranial hemorrhages may be caused by the dissection of ectatic artery, rupture of associated aneurysm, or rupture of stretched perforating arteries exposed to chronic hypertension. The shape, size, and localization of the hematoma revealed in our case fitted the criteria for hypertensive bleeding caused by rupture of perforating arteries. Angiography of the left ICA showed ordinary lumen of both anterior and middle cerebral arteries as well as a normal pattern of branching of these arteries. We did not notice changes in course or lumen of the perforating arteries that could have indicated pathologic process related to the dolichoectasia. Of course, possible microscopic defects of the walls of these arteries sometimes associated with dolichoectasia cannot be excluded.

The most remarkable finding in our case was the double lumen of C2 segment of the left ICA. In general, a double lumen of one artery may represent duplication or fenestration of one artery or dissection of its wall with 2 patent lumena. The anatomic variant consisting of 2 arteries running parallel instead of one is referred to as duplication. Fenestration is a duplication of a shorter segment of an artery. Duplications and fenestrations are frequently seen in the vertebrobasilar system but very rarely in the carotid system. There have been only 7 reported cases of duplication or fenestration of the ICA prior to 2004: 3 cases of fenestration of the supraclinoid segment, a single case of multichannel, short-segment duplication of the petrous segment of ICA, and 3 cases of duplication of the C1 segment of ICA. The main features of all of these cases, except the multichannel fenestration of the C2 segment, are parallel but clearly separate courses, smooth contours, and almost equal diameters of lumena. Gailloud et al reported 6 cases of “pseudofenestration” of ICA. The common feature of all these cases was relatively short involvement of the cervical ICA with pronounced asymmetry of the limbs of fenestration. Additional common features are irregularity...
of the contours and segmental narrowing and enlargement of the lumen. The authors pointed out that such changes are not typical in fenestration of arteries in other locations. On the other hand, described features are commonly associated with arterial dissection. Gailloud et al pointed out that review of the 4 previously reported cases of cervical ICA fenestration revealed similar findings. In all of these cases, asymmetry of the limbs, pronounced irregularity of the contours, segmental narrowing, and enlargement of the lumen of the ICA were described as additional findings to the fenestration. Gailloud et al ended their article with the conclusion that all described cases represent segmental arterial dissection with a double lumen sign rather than an anatomic variant.

Review of the medical records did not reveal any trauma that could have caused dissection of the ICA in our patient. Moreover, the localization and extent of the double lumen is not typical for dissection caused by trauma. Typical traumatic dissection of the ICA is located immediately bellow the skull base, at the entrance of the carotid canal. Only 4 cases of isolated atraumatic dissections of the petrous segment of the ICA have been published to date. All of these cases were symptomatic with radiologically confirmed disturbances of the blood flow in the dissected segment of the ICA. The absence of symptoms in our case suggests a very slowly progressing dissection that has not produced significant flow disturbances. One can assume that the combination of hypertension, weakness of the arterial wall, and the shearing forces of the blood flow at the entrance of carotid channel could have promoted a dissection. One can also hypothesize that distribution of flow forces in the limited space of the relatively narrow carotid channel could have contributed to such an unusual form of 2 patent arterial channels.

In conclusion, we have presented a case of a relatively rare form of CADE associated with a previously unreported form of dissection of the petrous segment of the ICA. The knowledge of such rare entities may have importance in diagnosis and treatment of vascular lesions of the brain.

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