MOYAMOYA PHENOMENON IN SERBIAN POPULATION

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To the editor:

We want to provide a detailed description of the case of Moyamoya phenomenon that was recently noted in the paper about the morphology of cerebral arterial circles (CACs) in Serbian population [1]. We found that the first paper about Moyamoya phenomenon referring to Serbia was published almost twenty years ago [2].

The word “moyamoya”, taken from a Japanese dictionary, in translation means a puff of smoke; in scientific literature it refers to the network of fine collaterals around and distal to the CAC due to progressive unilateral (Moyamoya syndrome) or bilateral (Moyamoya disease) stenosis of the intracranial part of the internal carotid artery (ICA) and its proximal branches [3].

Moyamoya syndrome was found in a 55-years-old male, who died due to pulmonary thromboembolism.

The research of cadaveric brain vessels was performed at the Institute of Forensic Medicine in Niš during co-author’s (MT) academic and postdoctoral studies. An approval for doctoral investigation was granted by the

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Fig. 1. Arteries on the brain base of a male cadaver. A, Convoluted blood vessels (Moyamoya pattern) in the sphenoid part of the left middle cerebral artery (circle), and three aneurysms (arrows) in the left anterior cerebral artery. B, Brain arteries and persistent primitive anastomoses are marked on modified image. C, Outer diameters of brain arteries and anastomoses are noted in the table. C4, choroid and communicating subparts of the cerebral part of the internal carotid artery (ICA); M1, sphenoid part of the middle cerebral artery; *stenosed M1 subpart; A1, precommunicating part of the anterior cerebral artery (ACA); A2, postcommunicating part of the ACA; ACoA, anterior communicating artery; PCoA, posterior communicating artery; P1, precommunicating part of the posterior cerebral artery (PCA); P2, postcommunicating part of the PCA; P2–ICA an., P2–ICA anastomosis; **bridge artery between the left P1 and P2; BA, basilar artery; V4, infracranial part of the vertebral artery.
Research Ethics Committee (No. 01-206-1) of our Faculty of Medicine. The vessels were photographed on the brain base; their outer diameters were studied from the digital images, using the ImageJ program (http://rsb.info.nih.gov/ij/index.html). Summarizing pathoanatomical findings, we will indicate six features: 1) a partial stenosis of the left middle cerebral artery (MCA) at its sphenoid part and a presence of a network of the MCA collaterals medially to the limen insulae; 2) the presence of three (one saccular and two berry) aneurysms in the left anterior cerebral artery (ACA); 3) the presence of atheromatous plaques in the basilar artery, and internal carotid and vertebral arteries; 4) the presence of a “bridge” artery between the left precommunicating (P1) and post-communicating (P2) parts of the posterior cerebral artery; 5) the presence of the left P2–ICA anastomosis; and 6) left-right caliber asymmetry of arteries on the brain base (Fig. 1).

As cited by Scott and Smith [3], the incidence peaks were in five-year children and adults in their mid-40s and the incidence among all patients with Moyamoya in Europe appears to be about 1/10 of that observed in Japan. Case reports in the Balkan journals were related to the case of Moyamoya disease in a 13 year-old girl in Serbia [2], and the case of Moyamoya syndrome of a 71-year-old male in Slovenia [4]. In a retrospective study by Borota et al. [5] over a period of 22 years in the former Yugoslavia, Moyamoya pattern of cerebral vessels was discovered only in 31 patients of both genders. We accepted the definitions of Moyamoya syndrome and Moyamoya disease given by Scott and Smith [3], although there were different diagnoses in cited cases [2,4], or Moyamoya modification [6]. However, there was an association of many vascular abnormalities in this, as well as in the cases of Moyamoya phenomena from the Balkan peninsula [4,5] to Japan [6], and beyond.


References