Dear Sir,

Moyamoya disease is a clinical entity in which occlusion of intracranial large vessels, in particular carotid arteries, occurs in association with the development of an abnormal vascular network within the brain, especially in the basal ganglia. Clinical manifestations include hemiplegia of sudden onset, symptoms of intracranial bleeding and seizures [1]. Seven to 10% of cases occur with familial clustering, with variable patterns of inheritance (more often autosomal dominant with incomplete penetrance) [2, 3]. The disease, although present throughout the world, has a particularly high incidence in Eastern Asia, particularly in Japan [4], as well as within close communities all around the world, such as the Amish and Mennonites. We describe the occurrence of moyamoya disease in a member of the community of Roma gypsies.

Case Report

A 36-year-old woman presented to the hospital ward for sudden appearance of a left sensorimotor defect. The patient belonged to the ethnic group of Roma gypsies, a population originating from Eastern Europe and present in Italy as well as in many European countries, and had been living in Italy since birth. The patient’s parents had died at a young age for unknown causes. A patient’s sister had died before the age of 30 of cerebral haemorrhage (not further investigated), and a patient’s son – a heterozygote twin – had died soon after birth for unknown causes. A brain CT scan performed at admission showed several hypodense areas within the cerebral parenchyma in both hemispheres. Brain MRI confirmed the presence of multiple T1-hypointense and T2-hyperintense lesions located bilaterally in the cortex and deep white matter of the parietal and occipital lobes. The patient was then submitted to cerebral angiography and magnetic resonance angiography, which evidenced a picture consistent with moyamoya disease (fig. 1) [5]. The HLA genotype was A11, A24, B14, B15, DRB1-1, DRB1-11, DRB3. Laboratory analyses evidenced low protein...
S activity and an abnormal resistance to activated protein C. Keeping in mind the haemorrhagic risk of moyamoya disease, it was decided not to treat the patient with oral anticoagulants. Low-dose salicylate (100 mg/day) was therefore started, and the patient showed a near-complete recovery from the neurological syndrome described. Few months later she presented again for transient left superior homonymous quadrantopsia, again with spontaneous remission. No evidence of intracranial bleeding occurred on either occasion.

Discussion

Roma gypsies make part of a large population of probable Central Asian (Indian) ancestry, who migrated to Eastern Europe and further towards Western countries [6]. Within the Roma gypsies – as frequently occurs in culturally isolated communities – consanguinity between spouses is almost the rule and, as a consequence, genetically based diseases are often found. The more common diseases affecting, either directly or indirectly, the nervous system include heredodegenerative neuropathy, muscular dystrophy and coagulation disorders [7–9]. To date, this is the first report documenting the occurrence of moyamoya disease in the Roma gypsy ethnicity. The clinical and anamnestic data available, in particular the finding of a patient’s close relative with fatal cerebral haemorrhage occurring at a young age, apparently suggest the presence of familial recurrence also in our case. The genetic background is likely to be different from that characterizing the East Asian cases, in whom a high occurrence of HLA-B51 is found [10]. Unfortunately, the peculiar cultural background of Italy-resident Roma gypsies (who live in complete isolation and even in strong conflict with the autochthonous community) hindered the collection of more detailed clinical information. It cannot be excluded, however, that misdiagnosed moyamoya disease may be responsible for a number of unexplained cerebral deaths within this community.

References