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Transient Motor Aphasia following Metrizamide Myelography

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Metrizamide, a water-soluble iodinated contrast agent for myelography, has been widely used in the United States for more than 4 years. Although acute generalized encephalopathy with confusion has been recognized as a complication of metrizamide myelography, aphasia, with the connotation of focality that it entails, has seldom been reported [1, 2]. We describe two patients with a transient motor aphasia clearly related to metrizamide myelography.

Case Reports

Case 1

A 56-year-old, right-handed house painter complained of intermittent low back and leg pain. He had mild coronary artery disease and hypertension, treated for 8 years with alpha-methylidopa and hydrochlorothiazide.

After premedication with 50 mg of meperidine, lumbar myelography with metrizamide (15 ml of 190 mg/ml) was performed without difficulty. After the myelogram he was kept flat in bed. About 6–8 hr later he complained of decreased hearing and light-headedness. He then became nauseated, vomited, and perspired heavily for 1 hr. At this time his language became abnormal. He followed verbal commands well, but his speech was nonfluent. He had considerable anoma to confrontation, verbal perseveration, and impaired repetition. Motor and sensory examinations were normal. Blood pressure readings were unremarkable. The entire episode resolved over 24–30 hr. Afterward he vividly remembered having had difficulty thinking of the words he wanted to say.

The episode was misinterpreted as a focal transient ischemic attack of the left hemisphere. Pencerebral angiography revealed a 30% stenosis of the origin of the left internal carotid artery. Mild stenotic changes were also noted in the right carotid siphon, proximal portion of the angular branch of the left middle cerebral artery, and perimesencephalic portion of the left posterior cerebral artery. Contrast clearance in the left frontoparietal operculum was slightly delayed. Subsequently, an atheromatous plaque was removed through a left carotid endarterectomy, and the patient was discharged.

His original back pain and leg pain persisted 9 months after the first myelogram, and he was readmitted to the orthopedic service. It was elected to repeat a metrizamide lumbar myelogram, this time without premedication. Again the myelogram was obtained without complication; afterward he lay flat in bed. About 8 hr postmyelography a sequence of events identical to the one described above occurred, of about the same duration.

Computed tomography (CT) and electroencephalography (EEG) performed after the patient made a complete recovery were normal. Regional cerebral blood flow, studied by the xenon-133 inhalation method (Novo Cerebrograph), was abnormal. Gray-matter blood flow averaged 55.1 ml/min/100 g in both hemispheres. The distribution pattern showed two areas of relative hypoperfusion, one in the right frontal region and one along the left Sylvian region. Within these two regions, perfusion was 40–53 ml/min/100 g (fig. 1). Blood flow in the left Sylvian region was 10%–22% lower than measured in the corresponding region of the right hemisphere.

Case 2

A 57-year-old woman complained of progressive gait disturbance for about 5 months. Neurological examination revealed mild spastic paraparesis. Cervical myelography was performed using 10 ml of metrizamide (290 mg/ml) injected in the lumbar region. During the procedure, metrizamide entered the cranial cavity; radiography showed slight opacification of the basal cisterns. The cervical myelogram was normal. Headache developed 7 hr after myelography, and she vomited several times. When examined 15 hr after myelography, she had a severe nonfuntion aphasia with marked anoma, but she followed instructions readily. In addition, she had a left supranuclear facial palsy, with impairment of voluntary facial movements; automatic facial movements were spared. EEG showed bilateral slowing of the background rhythm, with episodic bursts of delta activity, most pronounced over the left perisylvian region. She gradually improved, and by 24 hr after myelography her language had reverted to normal and the left facial weakness had disappeared. Repeated EEG was normal.

Discussion

The most frequently reported complications of metrizamide myelography are headache and other meningeal symptoms [3–5]. Transient memory loss and depression [6], as well as long-lasting confusion related to hydrocephalus [7],
metrizamide with a brain already damaged by a vascular abnormality. Neither of their patients, however, was evaluated by angiography or regional cerebral blood flow studies. Of interest, the cerebral blood flow study in our case 1 showed diminished perfusion of the left perisylvian region.

Several studies have documented the reproducibility of cerebral blood flow measurements obtained with equipment similar to that used for the study of case 1 [18–20]. Interhemispheric differences are particularly reproducible. Considering the normative data (71 volunteers aged 15–69; Wilson D, unpublished observations) from the laboratory that determined cerebral blood flow for our first patient, the low perfusion recorded over his left sylvian region is unlikely to be related to chance. In our patient, interhemispheric differences for the sylvian detectors were 12.7 ml/min/100 g for detector 6 (normals, 3 ml; SD, 5) and 10.7 ml/min/100 g for detector 10 (normals, 1.7 ml; SD, 5.6). Clustering of abnormal values to a group of detectors enhances the reliability of this finding. Although the cerebral blood flow measurements were performed after the patient had already had myelography, it is most likely that the same situation of decreased perfusion of the left perisylvian region was present before myelography. Thus, this case seems to support the view of Böker et al.

There is a controversy about the action of iodinated contrast materials upon the vasculature of the brain. Foltz et al. [21] reported transient severe vasospasm of the pial arterioles in monkeys after the intracarotid injection of contrast material [21]. Their technique involved direct visualization of the cortex, but as Hilal [22] noted, systemic blood pressure during the procedure was not recorded. Other authors were unable to duplicate their findings. [23, 24]. In a recent study arteriolar constriction was observed in the rabbit omentum 5 min after the intravenous injection of methylglucamine diatrizoate 60% [25]. Although no similar studies are available after the intrathoracic injection of metrizamide, it could be postulated that the contrast material, in contact with the adventitia of the pial vessels, may cause transient vasospasm. In our case 1, the left perisylvian region, with a diminished basal cerebral blood flow due to focal atheromatous disease of intracranial arteries, might have been affected more than the rest of the brain by such a process. An alternate explanation need not postulate vasospasm. Position emission tomography has shown that viable areas of the brain may have impaired blood flow but normal oxygen metabolism [26]. Such ischemic, but essentially metabolically normal areas may be rendered functionally incompetent by an increased metabolic load. Water-soluble contrast materials are known to enhance the evoked corticospinal responses [12]. Hypersynchrony and spikes observed in the EEG of patients after metrizamide myelography would also suggest transiently increased neuronal activity [9, 15]. Moreover, metrizamide in high concentration competes with glucose for hexokinase [27]. An area taxed by relative ischemia may not be able to meet these increased metabolic demands, and focal neurologic signs ensue.

Systemic hypotension might be postulated to explain a transient neurologic deficit related to myelography. Nevertheless, systemic hypotension has seldom been reported after metrizamide myelography [5, 28, 29]. In our cases,
blood pressure remained normal during the postmyelography episodes.

Although metrizamide is the best tolerated water-soluble contrast material available for myelography, it may cause an acute encephalopathy. Occasionally metrizamide encephalopathy may present with signs suggestive of a focal lesion, as exemplified by our cases.

REFERENCES