Vein of Galen Malformation with Cerebral Calcification: A Reversible Cause of Neurodegenerative Disease

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ABSTRACT: This paper reports two children who had mental retardation as the predominant manifestation of vein of Galen malformations. C.T. scans demonstrated extensive intracerebral calcification. Surgery reversed the progressive downhill course in one patient and may have arrested further deterioration in the other. The pathophysiological mechanisms underlying the clinical and radiographic features are discussed.

RESUME: Malformation de la veine de Galen: une cause inversible de calcification cérébrale et de retard mental progressifs. La présente communication rapporte les cas de 2 enfants présentant un retard mental comme manifestation principale de malformations de la veine de Galen. La scanographie cérébrale montrait une calcification cérébrale extensive. La chirurgie a inversé le processus de détérioration chez un patient et a peut-être arrêté sa progression chez l'autre. Les mécanismes pathophysiologiques sous-jacents aux manifestations cliniques et radiologiques sont discutées.

Vein of Galen malformations (VGM) have been postulated to result from fistular communication between diencephalic choroidal vessels and the great cerebral vein during embryogenesis. In the neonatal period the presentation is determined by the magnitude of the arteriocardiac failure and cranial bruit. Smaller shunts present in infancy, childhood, and adolescence with craniomegaly and cranial bruit, seizures, headaches, exercise syncope, and subarachnoid heamorrhage (SAH). The oldest patient reported in the literature is a 62 year old man who presented with SAH. Before 1973 the diagnosis was confirmed by cerebral angiography and while this investigation remains a prerequisite for surgical therapy, several reports have shown that VGM are demonstrable by computerized tomographic (CT) scanning.

Mental retardation (MR) is a recognized complication of VGM which has not been emphasized in the literature. This paper describes 2 patients in whom MR was the predominant clinical manifestation. Following surgery one improved dramatically; the other was left with a focal deficit but has not suffered further intellectual deterioration. In both, the VGM was demonstrated by CT but on each scan the most striking abnormality was widespread cerebral calcification.

REPORT OF CASES

Case 1: This patient was the product of an uncomplicated pregnancy, labour and delivery. His early development was normal. At 6 months of age he had a brief generalized seizure. His height, weight, and head circumference were approximately 2 standard deviations above the mean. The examination was otherwise unremarkable. There was no intracranial calcification on skull x-ray. An EEG demonstrated no abnormalities. A CT scan (unenhanced) was interpreted as normal, but in retrospect showed the VGM (Figure 1). His seizures were difficult to control but at 8 months of age his development was normal. At 12 months there was mild gross-motor delay but normal language development.

At 17 months of age he had a prolonged generalized seizure followed by a left Todd’s paralysis. On examination he had a high pitched cry, cortical blindness, marked hypotonia, hyperreflexia and a left hemiparesis. His head circumference was normal and there was no evidence of heart failure or cranial bruit. Developmentally he had regressed to a 2-3 month level with the reappearance of primitive reflexes. An unenhanced CT scan showed cortical atrophy and widespread cortical, subcortical and periventricular calcification (Figures 2 & 3). Cerebral angiography demonstrated a VGM fed predominantly by the posterior circulation. The serum calcium and phosphate were normal. Serum titres for TORCH infections were negative.

At 20 months of age his condition was essentially unchanged with his development still at a 2-3 month level. Through a right parietal-occipital craniotomy the aneurysmally dilated vein of Galen was seen to fill the
Figure 1: Case I. Unenhanced CT scan at age 6 months showing lesion in region of splenium of the corpus callosum. No cerebral calcification present.

Figure 2: Case I. Unenhanced CT scan at age 17 months showing atrophy and calcification of the cerebral cortex.

Figure 3: Case I. Unenhanced CT scan showing a dilated vein of Galen and calcification in the cerebral cortex and subcortical white matter.

Figure 4: Case I. Postoperative CT scan (unenhanced) showing bilateral subdural hygromas.
posterior cisterna ambiens. Several feeding vessels arising from the posterior cerebral and superior cerebellar arteries on both sides were clipped. This resulted in "collapse" of the brain producing a large brain-skull discrepancy (Figure 4) which was visible at the time of surgery. Bilateral subdural-peritoneal shunts were inserted.

Ten months after surgery (age 30 months) considerable improvement was evident. The Bayley Scales of Infant Development indicated that his mental development was at a 12 month level and that his psychomotor development had reached the 7 month level. Vision had returned. He was able to sit unsupported but made no stepping movements. He tended to hold his left upper extremity in a flexed, adducted posture but was able to utilize a raking grasp with the left hand. On the right he demonstrated an immature pincer grasp and could transfer objects from hand to hand. There was increased tone and hyperreflexia in both lower extremities. He used about a dozen words with meaning but was not speaking in sentences. Seizure control had improved. A repeat CT scan still showed extensive cerebral calcification (Figure 5) and almost complete resolution of the subdural hygromas.

Case 2: This boy was born at 36 weeks gestation following an uncomplicated twin pregnancy. He was noticed to have a large head at 4 weeks of age and was admitted to hospital at 6 months. His head circumference was greater than 2 standard deviations above the mean and his motor skills were delayed. The remainder of his examination was unremarkable. There was minimal cardiac enlargement on chest x-ray and a skull radiograph showed separation of the sutures but no intracranial calcification. Air ventriculography demonstrated moderate enlargement of the lateral and third ventricles but the aqueduct and fourth ventricle were normal in size, position, and shape. A ventriculoperitoneal shunt was inserted. Seizures began in the postoperative period.

At 4 years of age a CT scan showed extensive calcification throughout both cerebral hemispheres. Psychological testing at age 6 years revealed a developmental quotient of 75. His seizures were difficult to control with anticonvulsant drugs and repeated electroencephalograms showed multifocal epileptiform abnormalities. Cerebral angiography at age 7 revealed a large VGM (Figure 6). The preferential flow from all four major cerebral vessels was to the malformation. A CT scan showed extensive cortical and deep hemispheric calcification (Figure 7). The VGM appeared as a tear-drop shaped density in the region of the splenium of the corpus callosum (Figure 8). Aphasia and a right hemiparesis complicated surgical clipping of the arterial feeders.

At 11 years, he was moderately mentally retarded and had a left exotropia, left homonymous hemianopia, bilateral optic atrophy and generalized hypertonia, more marked on the right. His intellectual deficit has not progressed.

DISCUSSION

The loss of intellectual abilities which dominated the clinical picture in these two patients has not been previously emphasized as a major manifestation of VGM. Patient 1 was developmentally normal when he was examined at 6 months of age because of seizures. By 17 months he had regressed to a 3 month level with the reappearance of primitive reflexes. This marked deterioration was clearly reversed by surgery. Although still retarded at the age of 2½ years (10 months post-surgery) his development seemed to be merely delayed rather than deviated from normal. Patient 2 showed mild developmental delay at 6 months of age when evaluated for progressive macrocephaly. Air ventriculography failed to demonstrate a cause for his hydrocephalus. So it is possible that this was due to increased intracranial venous pressure. Surgery was complicated by the appearance of new focal signs and he is moderately mentally retarded but his deficits have not progressed. Thus, VGM should be considered in the differential diagnosis of infants with suspected neurodegenerative disease, especially if intracranial calcification is present.

Periventricular calcification has been reported in neonates dying from heart failure secondary to VGM. 16,17 White matter calcification has also been demonstrated in a mentally normal 20 month old child with a VGM. 13 Cortical calcification with underlying atrophy has been documented in a 2 year old with a posterior fossa dural arteriovenous malformation but no clini-
cal details were given. The two patients described here had extensive cerebral calcification. The improvement seen in case I despite persisting radiographic abnormalities suggest that calcium deposition per se is not the sole explanation for the clinical features.

In case I, the intra-operative observation of brain collapse was presumably due to a marked fall in cerebral blood volume. It is postulated that prior to surgery the arteriovenous fistula produced venous hypertension and an expanded blood volume. It is also possible that preferential flow of arterial blood to the VGM rendered much of the brain chronically hypoperfused via a steal phenomenon. The exact pathogenesis of the cerebral dysfunction and calcification in these cases is unclear, however, early intervention is important to prevent further neurological deterioration.

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REFERENCES


