CASE REPORT

Antenatal diagnosis of aneurysmal malformation of the vein of Galen

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SUMMARY

Vein of Galen malformation (VGAM) results from an aneurysmal aberration with an arteriovenous shunting of blood. Vein of Galen aneurysmal malformations are the most frequent arteriovenous malformations in infants and fetuses. The congenital malformation develops during weeks 6–11 of fetal development. Infants often die from high-output congestive heart failure.

BACKGROUND

This case is important because vein of Galen malformation (VGAM) is a rare congenital anomaly that often increases in size with advancing age and is associated with hydrocephalus, seizures, intracranial haemorrhage, heart failure and even resultant death. So early diagnosis and treatment is important in order to avoid such consequences.

CASE PRESENTATION

A 25-year-old woman with an 8-month history of amenorrhoea presented to hospital for routine antenatal check-up and was advised sonography for fetal wellbeing. She was a second gravida with one live child and no known previous congenital malformations.

INVESTIGATIONS

Ultrasonography examination revealed that fetal maturity was 31 weeks 5 days. All fetal biometric parameters were correlating with the mother’s gestational age according to her last menstrual period. We identified a 24×15 mm solitary, oval cystic lesion in the midline in the cranium extending into the right paramedian region of the posterior fossa. The lesion appeared to be separate from the occipital horns of the lateral ventricles. Both cerebella were normal. Ventricles with choroid plexuses were normal. No other abnormalities were found in the cranium. On colour Doppler, the lesion showed characteristic bidirectional blood flow (figure 1).

No other gross congenital anomalies were detected. The fetus was found to have mild cardiomegaly with situs solitus. Sonographic evaluation of the fetal heart did not reveal any other abnormality to justify the cause of the cardiomegaly.

Diagnosis of aneurysmal malformation of vein of Galen was made based on sonographic features and on the mild cardiomegaly.

OUTCOME AND FOLLOW-UP

The patient was delivered at full term through vaginal route. Physical examination of the baby was unremarkable. The baby was called for post-natal evaluation at the radiology department on the second day. On cranial sonography, a 32×33 mm cystic lesion was found in the midline and right paramedian region in the posterior fossa showing bidirectional blood flow correlating with antenatal findings (figure 2). No other abnormalities were found in the cranium. Echocardiography showed mild cardiomegaly with otherwise normal morphology, signifying high output as the cause of the cardiomegaly.

Antenatal diagnosis of VGAM was confirmed. On discharge from hospital on the fifth post-partum day, we explained the need for CT angiography and other interventions to the relatives of

![Figure 1](http://casereports.bmj.com/figures/10.1136/bcr-2015-213785)

**Figure 1** Antenatal ultrasonography image showing cystic lesion in posterior fossa that, on Doppler, showed bidirectional colour flow—vein of Galen malformation.
the patient. For these, the patient was referred to a specialist centre. Unfortunately, the patient was lost to follow-up.

**DISCUSSION**

The vein of Galen is located under the cerebral hemispheres and drains the anterior and central regions of the brain. VGAM results from an aneurysmal aberration with arteriovenous shunting of blood due to either thalamic arteriovenous malformation or a choroidal arteriovenous fistula.

Vein of Galen aneurysmal malformations are the most frequent arteriovenous malformations in infants and fetuses. The malformation increases greatly in size with age, although the mechanism of the increase is unknown. The right side cardiac chambers and pulmonary arteries also develop mild to severe dilation. The congenital malformation develops during weeks 6–11 of fetal development.

**Frequency:** unknown  
**Race:** VGAM occurs in all races  
**Sex:** Boys and girls are affected equally  
**Age:** VGAM is a congenital malformation; therefore, it may present at birth or in early childhood. It commonly presents in the neonatal period, although it may present in early childhood. Malformations often lead to cardiac failure, cranial bruits and hydrocephaly, and subarachnoid haemorrhage and seizures in neonates. Heart failure is due to the size of the arteriovenous shunt, which can steal 80% or more of cardiac output, with large volumes of blood under high pressure returning to the right heart and pulmonary circulation, and sinus venosus atrial septal defects. It is also the most common cause of death in such patients.

**Imaging studies**

**Cranial ultrasound**

- This will help to localise or identify the lesion.
- Doppler studies can help further to understand the haemodynamics of the lesion.

**Cranial MRI and/or CT scan**

- These studies will help confirm diagnosis and define the degree of involvement. Imaging studies will also help determine any accompanying hydrocephalus. Cerebral MRI shows large flow void in the central region with enlarged straight sinus and drainage to transverse sinuses.
- MR angiography can help to delineate vascular supply. MRI venogram shows VGAM with draining veins.

**Cranial angiography:** Cranial angiography is required to define the extent of aneurysmal dilation and details for arterial feeders. Venogram shows draining vasculature for VGAM.

**Cardiac ultrasound:** This study may be indicated to assess left ventricular function.

**Medication summary**

- No specific medical therapy is available
- Seizures usually are managed with antiepileptics

**Patient care**

- Infants who undergo surgical ligation or selective embolisation should be reimaged to assess results of the therapeutic intervention
- Head circumference measurements should be obtained in patients who undergo ventriculoperitoneal shunt to treat hydrocephalus. The patient should be monitored for the development of hydrocephalus.

**Complications**

Complications usually associated with VGAMs are usually intracranial haemorrhages. Over half the patients with VGAM have a malformation that cannot be corrected. Patients frequently die in the neonatal period or in early infancy. Studies have shown that 77% of untreated cases result in mortality. Even after surgical treatment, the mortality rate remains as high as 39.4%.

**Learning points**

- Vein of Galen malformation is an uncommon congenital malformation that should be kept in consideration for diagnosis if imaging features are suggestive.
- Vein of Galen malformation may be mistaken for a cystic lesion of the brain if no colour Doppler is performed. Colour Doppler helps to differentiate cystic lesions from vascular aneurysms.
- Vein of Galen malformations are associated with hydrocephalus, seizures, intracranial haemorrhage, heart failure and death. So early diagnosis and treatment is important in order to avoid such consequences.

**Competing interests** None declared.
REFERENCES