The child with jerking eyes and gait

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MRI IMAGES OF BRAIN IN JOUBERT SYNDROME

Joubert syndrome is a rare genetic disorder inherited as an autosomal recessive trait with an incidence of approximately 1 in 100 000 children. Primarily it affects balance and coordination and is characterised by the partial or complete absence of cerebellar vermis and dysplastic development of the midbrain.

MRI of the brain shows characteristic *wisdom tooth appearance* (figure 1) of the midbrain, secondary to the lack of decussation of the superior cerebellar peduncle which appears elongated along with a decreased diameter of the midbrain and a deep interpeduncular cistern. A *bat wing appearance* (figure 2) of the fourth ventricle is caused by the absence of the cerebellar vermis. The lateral ventricles and corpus callosum are normal.

This syndrome was first described by Marie Joubert,1 in 1969 in four siblings and one sporadic case that exhibited episodic hyperpnoea, abnormal eye movements, ataxia and mental retardation with cerebellar vermian agenesis and the midbrain hindbrain malformation.

The importance of this syndrome is related to the outcome and the potential complications that occur. A follow-up study of 19 children with Joubert syndrome showed neuromotor disability, developmental retardation or early death.2

These patients are sensitive to the respiratory depressant effects of anaesthetic agents which therefore must be avoided or need close perioperative respiratory monitoring.

Learning points

▸ Joubert syndrome is an inherited condition associated with developmental retardation, neuromotor disability and early death.

▸ Patients are extremely sensitive to the depressant effects of anaesthetic agents and require close perioperative respiratory monitoring.

▸ It is easy to identify the classical features on MRI brain. Antenatal diagnosis in subsequent pregnancies with ultrasound is possible.

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REFERENCES

