A 16-month-old male presented with progressively enlarging head for 12 months, decreased appetite and drowsiness for 1 month and downgazing eye for 15 days. There was no history of fever, vomiting, seizure or any antenatal issues. On examination the child was not obeying command and had a weak cry with no obvious motor deficit. A computed tomogram (CT) was done which showed a large bifrontal multiseptate lesion with minimal hydrocephalus displacing the frontal lobes and extending across midline (Figure 1 A). A low pressure cyto-peritoneal was inserted with clinical improvement next day. The fluid analysis was negative for infection and malignancy. The child was discharged home on the seventh day and is asymptomatic for the past one year (Figure 1 B). Repeat scan at 1 year showed complete collapse of the cyst (Figure 1 C).

Intracranial cyst in children is uncommon findings in neurosurgery. These can be found in different brain compartments and may have diverse origins. Of these choroid plexus and arachnoid cysts are the most common and usually have a good prognosis. Among the intraparenchymal cysts, the periventricular pseudocysts, ependymal cyst, neuroepithelial cyst, cystic periventricular leukomalacia, and the porencephalic cystic lesions are the most representative. Other uncommon causes are hydatid cyst, intracerebral abscess, cyst with glioma and trapped lateral ventricle. The prognosis depends on the presence of associated findings and on the extent and place of the insult. The diagnosis is with CT or a Magnetic Resonance imaging. Management consist of serial follow-up if small and asymptomatic to surgery which includes fenestration- endoscopic or open, shunting of contents, complete excision or marsupilisation.

### References