A rachnoid cysts are developmental anomalies involving duplication or splitting of the arachnoid membrane mostly diagnosed incidentally and was first described in 1831 by the English physician Richard Bright in 1789-1858. The large majority are often found in the middle cranial fossa with sellar/suprasellar arachnoid cyst (SSAC) comprising of 9-15%. SSAC can present with headache, endocrine dysfunction, psychomotor abnormalities, chronic hydrocephalus, vertigo, optic nerve and visual deficits and asymptotically. We report a case managed with cystocisternostomy and long term follow-up with good improvement of his visual function.

**Key Words:** arachnoid cyst, craniopharyngioma, cystoventriculostomy, cystoperitoneal shunting sella/supra sella

**Case Report**

A 35-year-old male was brought by his relatives with history of headache and progressive loss of vision for 6 months. For the last few weeks he had loss of complete vision on left side and was completely dependent for mobilization and activities of daily life-independent (ADL). Preoperative blood and endocrinology (Thyroid function [TSH 3.0mIU/ml], Prolactin [9.27 ng/ml] and Growth hormone [0.270ng/ml]) work-up was normal. Ophthalmological examination showed visual acuity in right eye as perception of light (PL) and projection of rays (PR) accurate. The visual acuity on the left eye was no perception of light (NPL). The fundus examination showed early papilloedema in right eye and severe papilloedema in left eye. Humphrey Visual field (HVF) could not be done.

Giant Sellar/Supra Sellar Arachnoid Cyst with Visual Loss: 5 Years Follow-Up Post Cystocisternostomy

Sellar/suprasellar arachnoid cyst can present with headache, endocrine dysfunction, psychomotor abnormalities, chronic hydrocephalus, vertigo, optic nerve and visual deficits and asymptotically. We report a case managed with cystocisternostomy and long term follow-up with good improvement of his visual function.
SSAC, the suprasellar types are more common with the mechanism of formation as for arachnoid cyst in other regions. The differential diagnosis of SSAC includes cystic craniopharyngiomas, pituitary cystic adenomas, Rathke’s cleft cysts, epidermoid tumours, empty sella, inflammatory cysts, and ependymal cysts. SSAC can coexist with pituitary adenoma. In addition to the symptoms mentioned above precocious puberty can be presenting sign in 10-40% secondary to damage to the ventromedial nucleus of the . Diagnosis is with help of CT or MRI showing non-enhancing hypo-dense lesions and hypo-intense on T1 and Hyper-intense on T2 with characteristics as of CSF respectively. If symptomatic, surgery is the treatment of choice and options include cyst aspiration, wall fenestration, wall excision, cystoventriculostomy, cystosubdural shunt, cystoperitoneal shunting and endoscopic transcranial or trans nasal excision. Recurrent cases may need placement of a subcutaneous reservoir or change of procedure. This case has demonstrated that symptomatic arachnoid cyst if treated early by surgical intervention can alleviate the symptoms and even improve visual outcome in those compromised. Visual and endocrinological symptom improvement have been documented by other authors after surgery.

**Conclusion**

SSAC are rare lesions that are mostly asymptomatic but can present with a variety of clinical signs and symptoms. Early surgical treatment for symptomatic lesions can result in good clinical outcome.

**References**


**Figure 1. MRI T1 and T2 image in axial and sagittal views taken in 2012 showing the SSAC (A,B), postoperative scan showing 20% residual cyst (C) and latest scan in March 2017 showing complete absence of the SSAC (D)**