Fourth Ventricular Outflow Obstruction in an Infant with Ileal Atresia and Laryngomalacia: Endoscopic Management

Forhad H. Chowdhury¹, Mohammod Raziul Haque², Jalal Uddin Mohammad Rumi¹, Mohammad Samsul Arifin³

¹Department of Neurosurgery, National Institute of Neurosciences and Hospital, Dhaka, Bangladesh
²Department of Neurosurgery, Dhaka Medical College Hospital, Dhaka, Bangladesh
³Department of Neurosurgery, Ibn Sina Specialized Hospital, Dhanmondi, Dhaka, Bangladesh

Address for correspondence Forhad H. Chowdhury, MBBS, FCPS, MS, Department of Neurosurgery, National Institute of Neurosciences and Hospital, Shere-e-Bangla Nagar, Dhaka 1207, Bangladesh (e-mail: forhadchowdhury74@yahoo.com).

Abstract
Fourth ventricle outflow obstruction (FVOO) is a rare cause of obstructive hydrocephalus. In this study, we described a case of idiopathic FVOO with ileal atresia and laryngomalacia which was managed with endoscopic third ventriculostomy (ETV) and re-endoscopy. We also described the techniques of fenestration of Liliequist membrane and partial removal of arachnoid membrane over dorsum sella (DS) to prevent closure of fenestration and recurrence of hydrocephalus. The patient was a 4-month-old infant presented with progressively increasing head size, feeding difficulty, respiratory distress, and tense fontanel. The infant had a history of laparotomy for ileal atresia. CT scan showed panventriculomegaly due to FVOO. ETV with fenestration of Liliequist membrane was done on emergency basis. After operation, the patient improved clinically and radiologically. Four weeks later, the patient returned with recurrent hydrocephalus. Endoscopic reoperation showed closure of fenestration in arachnoid membrane (Liliequist membrane). Endoscopic refenestration with partial excision of arachnoid on DS was done. The patient again recovered radiologically and clinically till last follow-up. In idiopathic FVOO, ETV with wide fenestration of Liliequist membrane, preferably with partial removal of arachnoid on DS, may be very useful in treating hydrocephalus (HCP) and preventing recurrent HCP even in infants.

Introduction
Fourth ventricle outflow obstruction (FVOO) is a rare clinico-radiological condition, which causes obstructive hydrocephalus. In FVOO, cerebrospinal fluid (CSF) flow is blocked at the foramen Magendie and Luschka due to atresia or by a membrane in the absence of any other obstruction of CSF flow. Various terms for FVOO have been used such as fourth ventricle/ventricular outlet obstruction,¹,² fourth ventricular outflow obstruction,³ membranous obstruction of the fourth ventricle outlet,⁴ obstruction of Magendie’s and Luschka’s foramina,⁵ obstruction of fourth ventricular exit,⁶ and primary obstruction of the fourth ventricle outlets.⁷ Far distal obstructive HCP is a term that includes Dandy Walker or Arnold Chiari malformation, membranous obstruction of fourth ventricle, and intercisternal external obstruction of the CSF.⁸ The etiopathogenesis of FVOO is not well understood and may be congenital, although some cases present with a history of meningitis or intraventricular hemorrhage.
Here, we described a technical case of neonatal idiopathic FVOO with ileal atresia and laryngomalacia where initial endoscopic third ventriculostomy (ETV) was successful but re-endoscopic procedure was needed for recurrence of HCP.

**Case Presentation**

Our patient was 4 months of age, first issue of his parents (nonconsanguineous marriage) presented with progressively increasing head size, feeding difficulty, respiratory distress, and tense fontanel. The infant had a history of (H/O) laparotomy in the 1st week of his life for ileal atresia where ileal resection and ileo-ileal anastomosis with proximal ileostomy were done. Ileostomy was closed when he reached his 8th week. He was also diagnosed with a case of laryngomalacia. He had no history of birth asphyxia, head trauma, or central nervous system (CNS) infection.

The patient came to us with a previously done (10 days earlier) CT scan of head, which showed dilatation of all ventricles with ballooning of fourth ventricle, and foramen of Magendie and Luschka. The cerebellum was hypoplastic and brainstem was pushed anteriorly. There was no visible cisterna magna, premedullary cistern, pre-pontine cistern, or basal cistern. Brain parenchyma was pushed toward the periphery and there was periventricular hypodensity (►Fig. 1). So, a diagnosis of FVOO was made. A repeat CT scan was done immediately which showed all the earlier-mentioned findings in augmented form (►Fig. 2). The previously done echocardiogram and ultrasonography (USG) of whole abdomen were normal. After counseling, emergency ETV was done under G/A.

**Operation ETV**

Under general anesthesia with endotracheal intubation, patient was positioned in supine position. Through the precoronal right-lateral part of the anterior fontanel, the lateral ventricle was entered. Through the right foramen of Monro, the third ventricle was reached. After identification of mammillary bodies, infundibular recess, basilar bifurcation, and dorsum sella ([DS] through the ventricular floor), a fenestration was made very carefully with endoscopic ventricular forceps in the midline floor of the third ventricle between DS and basilar artery (BA). Then, the space between DS and BA was increased due to CSF entry from ventricle and that made the rest of the operation more feasible. The fenestration was enlarged with 3F Fogarty catheter balloon. The thick layer of arachnoid band was found attached to the DS to BA, which was also widely fenestrated and partially excised (►Fig. 3). Clean pulsating CSF flow was seen through the fenestration. Endoscope was removed after inspection of

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**Fig. 1** CT scan of head that was done 10 days earlier to presentation to us: (A) sagittal section, (B-F) axial sections showing panventriculomegaly due to FVOO with periventricular edema. Arrow marks in (A) showing fourth ventricular enlargement with absent of cisterna magna, indicating foramen Magendie atresia. Arrow marks in (B) showing enlargement of lateral recesses of fourth ventricle as well as absent cerebellopontine angle cisterns, indicating foramen Luschka atresia. FVOO, fourth ventricular outflow obstruction.

**Fig. 2** CT scan of head at presentation: (A, B) sagittal sections and (C-F) axial sections, showing augmented form of panventriculomegaly due to FVOO with increased periventricular edema. Arrow marks in (A, B) showing fourth ventricular enlargement with absence of cisterna magna, indicating foramen Magendie atresia. FVOO, fourth ventricular outflow obstruction.

**Fig. 3** Peroperative pictures of 1st operation ETV: thick floating arachnoid layer anterior to BA and posterior to clivus and DS, as indicated by arrow marks (which reunited [seen during 2nd endoscopy] and hydrocephalus recurred). BA, basilar artery; DS, dorsum sella; ETV, endoscopic third ventriculostomy.
cerebral aqueduct (which was open and enlarged) and wound closed accordingly.

Postoperatively, the patient became stable with respiration and breast feeding. His fontanel became lax. CT scan of head on 1st postoperative day showed decrease size of ventricles, including fourth ventricle with CSF in subarachnoid spaces which was more marked on the right side (Fig. 4A, B).

After 4 weeks of ETV, the patient again developed feeding and respiratory difficulties with tense anterior fontanel. Repeat CT scan showed ventricular re-enlargement including fourth ventricle with periventricular hypodensity; brain parenchyma was pushed toward periphery with drained subarachnoid spaces (Fig. 5). We decided to reoperate on him after counseling with parents regarding the necessity of shunt operation if re-endoscopy fails.

**Re-endoscopic Operation**

Under general anesthesia with endotracheal intubation patient in supine position, ventricle was reached through the previous route. Endoscopic stoma in the floor of the third ventricle was patent but there was no flow through the stoma. Arachnoid membranes were uniformly reunited to a membrane between DS and brainstem (including BA; Fig. 6). With the boring action of the tip of the Fogarty catheter, we perforated the membrane between clivus and BA. With the help of endoscopic grasping forceps, we stripped off and partially removed the arachnoid membrane from DS, clivus and...
arachnoid membrane anterior to the BA, taking great care not to injure the vessels and neurostructures, including the third and sixth cranial nerves (►Fig. 7–9). After making a big hole in the reunited membrane, endoscope was removed and the wound was closed accordingly.

Postoperatively, the patient’s respiration became stable with respiration and breast feeding was restarted. His fontanel became lax. CT scan of head on the fourth postoperative day showed decrease size of ventricles, including fourth ventricle with CSF in subarachnoid spaces, and there was marked reduced periventricular hypodensity (►Fig. 10). Then, the patient was symptom-free with normal milestone of development till last follow-up (i.e., 14 months after operation).

Discussion

FVOO is a very rare cause of obstructive HCP. FVOO usually occurs in pediatric age group and may be congenital, but can occur in adults also. There is no sex differentiation. Mohanty et al. reported a case series of 22 patients with FVOO; of these, 10 patients had a medical history, three had suffered intraventricular hemorrhage, and seven patients had infections, including tubercular meningitis, bacterial infection, or prolonged and unexplained fever. The rest of the 12 cases were idiopathic FVOO. Head injury can be a cause of membranous obstruction of CSF and can cause FVOO.

Diagnostic Modalities for FVOO

The pattern of ventricular enlargement in FVOO is named panventriculomegaly or tetraventricular HCP. Dilatation or large CSF collection of the foramina of Magendie and Luschka is a characteristic radiological finding in cases of FVOO. However, it is very difficult to confirm the presence of a membranous obstruction at ventricular outflow via conventional magnetic resonance imaging (MRI). High-resolution, three-dimensional constructive interference with steady state sequence on 3T MRI may be able to detect obstructive membranes, but this may not be possible in all cases.

The most sensitive diagnostic method is computed tomography (CT) ventriculography, with the injection of contrast medium through a ventricular catheter. Serial CT images after injection will show collected contrast medium in the outlets of the fourth ventricle and subsequent blockage of its diffusion to the prepontine cistern.
The use of MRI instead of CT as the diagnostic modality for FVOO is recommended to avoid exposure to radiation.\(^6\)

Phase-contrast MRI,\(^6,7\) cine MRI,\(^1,4,6,7,10,11,17\) or radioisotope cisternogram\(^1\) can be used to see the CSF dynamics.

Direct endoscopic inspection of the fourth ventricle, in case the aqueduct is sufficiently expanded, is another diagnostic option.\(^2\) Although this technique needs to be done under general anesthesia and carries a risk of damaging the midbrain around the aqueduct, it has been recently reported to be relatively safe.\(^2,13,18,19\) When MRI is suggestive of FVOO, this technique can allow simultaneous diagnosis and treatment.

Mohanty et al\(^2\) reported the entire success rate of ETV for FVOO is 65% (13 successes in 20 cases). Although they did not evaluate the success rates of primary and secondary FVOOs separately, they speculated that failure was attributable to CSF malabsorption as a result of prior meningitis or intraventricular hemorrhage.

Oertel et al\(^1\) reported the results of ETV in 20 cases of far distal obstructive HCP where most of the cases were with Dandy Walker or Arnold Chiari malformations; however, there were four cases considered as secondary FVOO. Two of the four patients (50%) were successfully treated by ETV, while the remaining two patients required early shunting. The success rate of ETV in primary FVOO is 75 to 100%.\(^1,4,6,8,17,20\)

Most of the failures of ETV for treating FVOO occur within 6 weeks of surgery and that subsequent endoscopic re-exploration revealed patency at the fenestration site.\(^2\) In some cases, the recurrent HCP can differ. The recurrence occurs later than 6 weeks and stomal stenosis or obliteration can be the cause, which can be successfully treated with endoscopic re-expansion. Indeed, this suggests that the recurrence of HCP observed in such case is not attributable to CSF malabsorption. In such cases (where recurrent HC does not appear to be caused by malabsorption), repeated ETV would be an effective treatment option.\(^21\)

Direct fenestration of membranous obstruction at the fourth ventricle outlets is another previously reported treatment option.\(^1,18,19\) However, its usefulness is still unclear because it was used in combination with ETV in most cases.

Regarding ETV in infant, some series showed that failure rate can be very high and can be even higher in neonates.\(^1,7,10,20,22\) But at the same time, more recent series showed more success of ETV in infants and neonates.\(^2,23\)

Our case was a unique case of primary FVOO with classical CT findings in neonatal age associated with other congenital conditions such as ileal atresia and laryngomalacia.

During ETV, we preferred it to ventriculo-pleural (V-P) shunt as:

1. HCP was of an obstructive variety.
2. Simple, quick, and less traumatic.
3. It is effective even in neonates.\(^22\)
4. H/O laparotomy and ileostomy in a small abdomen where shunt insertion may be difficult with failure and complication.
5. V-P or ventriculo-atrial shunts may cause more respiratory problem (as patient had laryngomalacia).

6. Shunt is standby procedure when ETV fails.

But there were some concerns such as the following:

Result of ETV in FVOO is not well known.

Failure chance of ETV is more in neonates.

No or very minimum space between brain stem and DS and clivus.

Chance of thick Liliequist membrane.

To work in a very narrow deep space, with important surrounding vessels and microvessels.

Chances of damage to third and sixth cranial nerves.

During second endoscopic procedure, we preferred it as:

1. To confirm the cause–where we found stoma was alright but arachnoid reunited to form a membrane that obstructed CSF flow.
2. If cause is found that can be treated simultaneously (here, we found the cause and treated it with endoscopy).
3. Shunt is standby procedure.

Before second endoscopy, we reviewed the 1st operation’s video, where we noticed a large thick flap/band of arachnoid remained flapping in front of BA and posterior to clivus that reunited with the DS arachnoid membrane and surrounding arachnoids to stop CSF flow (►Figs. 3 and 6). We thought if we were removing (partially /totally) or coagulating this “culprit” arachnoid flap with removal of the arachnoid on DS, we could prevent recurrence of HCP.

So, during re-endoscopic procedure when we found “the thought culprit arachnoid flap” reunited, we perforated it and removed it partially (►Figs. 7–9). At the same time, we also removed the arachnoids on DS. We think, arachnoid on DS has a very important role in recurrence of HCP after ETV with Liliequist membrane penetration, especially in infants.

**Conclusion**

From a single case, it was difficult to make a conclusion but, in idiopathic FVOO, ETV with wide fenestration of Liliequist membrane, preferably with partial removal (or judicial coagulation) along with partial removal of arachnoid on DS, may be very useful in treating HCP and preventing recurrent HCP even in infants.

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**Conflict of Interest**

None declared.

**References**

Endoscopic Management of Neonatal FVOO  Chowdhury et al.