Extracranial Complications of Cerebrospinal Fluid Shunt Function in Childhood Hydrocephalus

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There were 112 separate hospital evaluations in 84 patients for suspected shunt malfunction; 96 evaluations were of ventriculoperitoneal shunts, 13 were of ventriculooatrial shunts, and three were of both types of shunts. In 45 (47%) of 96 ventriculoperitoneal shunts, complications eventually led to surgical revision; 20 (44%) of these were problems of the peritoneal end and therefore peculiar to this type of shunt. Peritoneal end problems included tubing disconnection, bowel obstruction, perforation, and abdominal cerebrospinal fluid pseudocyst. Of the 13 ventriculooatrial shunts, 10 (79%) required revision; eight (61%) of these were due to problems of the atrial end. These problems included relative shortening of the tubing due to patient growth, superior vena cava thrombosis, and disconnection. Ventriculoperitoneal shunts were used most frequently and had a lower complication rate (47%). Ventriculooatrial shunts were used less often and had a higher complication rate (79%) and more serious problems.

Cerebrospinal fluid shunting systems are far from trouble-free devices, often requiring frequent surgical revision to assure continued function. In an earlier study [1], we detailed evaluation of the proximal part (ventricular end) of shunt systems by computed tomography (CT), which has become very useful in the management of pediatric hydrocephalus [2–6]. In this paper, we deal with radiologic evaluation of extracranial parts (distal ends) of shunt systems and the important complications of each of the principal shunt types.

Materials and Methods

There were 112 separate hospital evaluations of shunt function status in 84 pediatric patients under age 13 years at University of Miami School of Medicine/Jackson Memorial Hospital. All 112 evaluations involved a CT scan of the brain and routine radiography of the skull, chest, and, in ventriculoperitoneal shunts, the abdomen. Other means of radiographic evaluation of shunt function included radionuclide cisternography/shuntography (four cases) and ventriculography (one case).

In all cases, admission for shunt evaluation was predicated on some change in the patient's status that signaled possible increased intracranial pressure and prompted the question of shunt patency [1]. Shunt malfunction was recognized by papilledema, increasing head circumference, seizures, decreasing levels of consciousness, enlarged or enlarging ventricles on CT scan, difficulty pumping the small subcutaneous part of the shunt located under the scalp, or, usually, a combination of several of these criteria.

Results

Of the 112 shunt evaluations, 13 (12%) were of ventriculooatrial shunts and 96 (86%) were of ventriculoperitoneal shunts. In three instances (2%), the patient had both types of shunts.
The 112 shunt evaluations were grouped according to the shunt complications that led to surgical revision and that were peculiar to each of the two types of shunt (Table 1). Of the 13 ventriculooatrial shunts evaluated, there were 10 complications (79%). Of the 96 ventriculoperitoneal shunts evaluated, 45 cases (47%) had complications. The three cases with both ventriculooatrial and ventriculoperitoneal shunts did not need surgical revision. In each case, at least one of the shunts was working well enough to drain the ventricular system.

**Ventriculooatrial Shunts**

Ventriculooatrial shunt complications mostly involved the distal (atrial) end (eight of 10 complications). One case of ventricular end malfunction and one case of infection were excluded, since they could as easily have occurred in ventriculoperitoneal shunts and were not peculiar to the ventriculooatrial form. Routine chest radiography was used to determine the length and position of the atrial end, resulting in the discovery of shortened tubing in half of the cases (four of eight complications). As the pediatric patient grows, the constant length of the vascular tubing results in progressive elevation of the distal tip out of the region of the atrium, which is the optimal location for successful drainage of cerebrospinal fluid [7, 8]. Figure 1A shows a chest film of case 1, a 3-month-old child with functioning shunt; 8 months later, the tip of the atrial end was above the level of the atrium and the shunt functioned marginally (fig. 1B). The shunt pumped fairly well, but some ventricular enlargement was present on CT. The shunt tubing was lengthened surgically in this case, after which the patient did well. Lengthening of the atrial end is considered necessary on a prophylactic basis if the tubing becomes too short with growth of the child, and replacement of the distal end is necessary if it flips out of the atrium or kinks on itself [9].

One of the most serious complications of ventriculooatrial shunt, thrombosis of the superior vena cava [10], was found in four of our evaluations. Case 2, an older child with a ventriculooatrial shunt (fig. 2), had enlarged vessels on CT and the shunt pump depressed with some difficulty, suggesting distal shunt blockage. Attempted injection of contrast medium (Conray) toward the distal end met with obstruction; at surgical revision, the distal end was fibrosed in the superior vena cava and had to be left in place. The ventricular end was connected to a new peritoneal end shunt and the patient did well. Other potentially serious complications—disconnection of the tubing, perforation of neighboring organs [11], and infection—were not encountered in our series. Our complication rates compare favorably with other series [12].
Ventriculoperitoneal Shunts

Since they are more popular, ventriculoperitoneal shunts showed most of the complications in our series but had a complication rate of only 47%. This agrees with earlier findings that ventriculoperitoneal shunts had fewer and less serious complications than ventriculocisternal shunts [13–15]. About half (20/45 in our series) of all complications leading to surgical revision involved the distal (peritoneal) end. The other complications, 19 involving the ventricular end and six infections, were excluded as not being peculiar to the ventriculoperitoneal shunt system. Plain radiographs are most useful when the shunt tubing becomes shortened or disconnected, as observed in eight and four of our cases, respectively. Although all disconnections occurred in ventriculoperitoneal shunt patients in our series, they have also been seen in ventriculocisternal shunts, sometimes resulting in embolism of the separated end through the vascular system [11].

Case 3 is a 9-year-old child whose ventriculoperitoneal shunt became disconnected at the juncture of tubing with pumping system (fig. 3). The distal end migrated into the abdomen, and CT showed enlarged ventricles. The shunt was reconnected and worked well; his ventricles returning to normal size. Case 4 is an unusual case in which the peritoneal end of a ventriculoperitoneal shunt in an 18-month-old boy with intraventricular obstructive hydrocephalus migrated through the bowel wall and dangled from the rectum (fig. 4). CT and physical examination confirmed the shunt to be functioning well, but the distal limb had to be replaced in the peritoneal cavity. Surprisingly, the patient had no apparent infection. In addition to this case, three other instances of perforation of a viscus by a shunt tip were observed in our series, including the next case.

Case 5, a 16-month-old girl shunted for extra ventricular obstructive hydrocephalus, illustrates many of the problems with peritoneal shunts. She had a ventriculocisternal shunt initially which was revised once because of growth, then changed to a ventriculoperitoneal shunt after a shunt infection. The ventriculoperitoneal shunt was revised when she developed an acute abdomen. At laparotomy, the peritoneal end was implicated in a perforation of the gallbladder and two lacerations of the liver. Twice after this a small bowel obstruction was repaired with takedown of adhesions; on one of these occasions the shunt tubing was found to be knotted tightly around the bowel (fig. 5).

In case 6, a peritoneal end was noted to migrate into an umbilical hernia (fig. 6), causing cerebrospinal fluid subcutaneous edema. Distal ends have been known to cause peritoneal cerebrospinal fluid cysts in areas of loculated fibrosis or in cases where the greater omentum blocks the shunt tip; in our group the greater omentum was noted to have migrated to the peritoneal shunt tip and was obstructing it at laparotomy in three cases. Cerebrospinal fluid then backs up and hydrocephalus is exacerbated, requiring revision. Despite their frequency, ventriculoperitoneal shunt
complications are believed to cause less total mortality and morbidity than the intravascular complications of ventriculoatrial shunts [15–22].

Discussion

Hydrocephalus is caused by obstruction anywhere along the pathway from cerebrospinal fluid production to absorption [23, 24]. Intraventricular obstructive hydrocephalus will occur with obstruction anywhere from the lateral ventricle to the exit foramina of the fourth ventricle [24, 25]. Extraventricular obstructive hydrocephalus replaces the old term of communicating hydrocephalus and denotes obstruction to the reabsorption of cerebrospinal fluid flow by processes scarring or blocking the arachnoid granulations and the subarachnoid spaces.

The concept of shunting involves mechanical redirection of cerebrospinal fluid from an obstructed cavity to an area capable of fluid reabsorption. Just about any body cavity capable of reabsorbing fluid has been used by surgeons at one time as a repository for cerebrospinal fluid [24, 25]. Surgically, one end of a Silastic tube is placed into a lateral ventricle, usually the right, via a burr hole. A Holter, Pudenz, Portonoy, or Hakim valve will provide unidirectional flow of cerebrospinal fluid and is capable of transcuscutaneous flushing. The distal tubing is placed in a receptacle organ or cavity. If an obstruction is located intraventricularly, a shunt may be directed a short distance beneath the scalp into the cisterna magna to provide access to the normal subarachnoid reabsorptive sites. This simple bypass system, the Torkildsen procedure, is not used extensively today [26].

A popular shunting procedure in the 1950s was the ventriculoatrial shunt, in which the distal shunt was anastomosed with the free proximal end of a ureter after nephrectomy [27]. This operation has been abandoned, although the shunt had a good reputation for reliable function. Many patients with functioning ventriculoatrial shunts are still alive. The ventriculopleural shunt has been abandoned because of the often considerable resultant pleural effusion. Other abandoned sites of drainage have been to the cystic duct, Stensen's duct, thoracic duct, or fallopian tubes [26, 28]. If hydrocephalus is of the extraventricular obstructive variety, a shunting system from the lumbar subarachnoid space, subcutaneously directed around the flank and then inserted into the peritoneal cavity, can be useful. Scoliosis sometimes resulted in the growing child, whether due to previous hemilaminectomy [29] or to underlying disease such as neurofibromatosis, and its use is limited at this time. Other severe complications associated with these shunts include arachnoiditis and electrolyte imbalances.

REFERENCES