Mutism as a consequence of callosotomy

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Transient mutism has been reported following commissurotomy and callosotomy. The cause for this mutism is unknown. A case of mutism following callosotomy is presented, and the preoperative and postoperative data on neurological, physiological, and psychological functioning are discussed. The data suggest that the mutism is not caused by general intellectual deterioration, cortical lesion, or peripheral damage affecting speech production. The syndrome may result from severing interhemispheric connections in cases where both hemispheres are required for speech production.

KEY WORDS • callosotomy • mutism • speech • hemispheric disconnection • epilepsy • corpus callosum

Commissurotomy and callosotomy have been used in the treatment of medically intractable epilepsy that is not amenable to focal surgical excision. Sunderland found in monkeys that the interhemispheric fibers are arranged anteriorly to posteriorly; the frontal lobes are connected via the rostrum and genu; the midtemporal, posterior temporal, and parietal lobe fibers pass through the body; and the occipital fibers pass through the splenium. The mesial and anterior portions of the temporal lobes are connected via another pathway, the anterior commissure.

There are fewer than 100 reported cases of callosotomy for seizure control. This procedure is palliative and produces few obvious intellectual deficits. In addition to the “disconnection syndrome,” memory quotients have been reported to be lower in postcommissurotomy patients than might be expected on the basis of intelligence quotients (IQ’s).

A major consequence of this procedure is transient mutism. Bogen reports that “in almost every case there was a time in which the patient was mute.” Wilson, et al., reported mutism in three of their first 12 cases. We present the pre- and postoperative data in a patient who suffered mutism following callosotomy. These data may be pertinent to an explanation of this phenomenon.

Case Report

This 20-year-old right-handed man had an 11-year history of seizures. He was the product of a normal pregnancy and a forceps delivery. He had delayed developmental milestones and was placed in special education classes from 6 years of age onward. His full-scale IQ at 7 years old was 73, with a verbal subscore of 91 and a performance subscore of 58.

Observed seizures began at 9 years old, with three to four episodes per week. One year later, he was noted to have one to two generalized tonic-clonic seizures per month, as well as episodes described as jerking of his head backward with momentary loss of awareness (10 to 15 per month). At the age of 12 years, he also experienced 15 epileptic drop attacks per month, requiring him to wear a helmet constantly. At 15 years of age he entered an institution for disabled children and had a normal physical and neurological examination, apart from a mild to moderate intellectual impairment. A seizure log was maintained for 3 months preoperatively by his school caretakers and revealed 94 atonic and tonic-clonic seizures. Numerous electroencephalographic (EEG) examinations showed diffuse abnormalities without a clear focus. He was admitted to our institution in August, 1980, for corpus callosotomy.

Examination. Neurological examination disclosed a cheerful cooperative boy with decreased attention
Mutism following callosotomy

FIG. 1. Preoperative electroencephalographic tracings showing 2 to 2.5/sec slow activity (predominantly anteriorly), and generalized spike and wave activity with a phase reversal frontally. Low voltage activity (20 to 25/sec) can be seen anteriorly, and 6/sec background activity bilaterally, centrally, and parietally.

span. He was able to follow two-step commands. The first through 12th cranial nerves were intact. Sensory examination was normal except that double simultaneous stimulation resulted in left-sided extinction. Motor examination and reflexes were normal. There were mild cerebellar signs.

Intellectual functioning was within the mild to moderate mental retardation range (Wechsler Adult Intelligence Scale full-scale IQ score 59, verbal score 66, performance score 43). Neuropsychological testing indicated receptive vocabulary compatible with his verbal IQ (Peabody Picture Vocabulary Test (PPVT) score 67); and memory functioning at the level of his full-scale IQ score (memory quotient 51). Analysis of memory scale subtests revealed significantly poorer functioning in visual reproduction than logical-verbal memory. This, combined with the lower performance IQ score, was interpreted as suggesting poorer functioning of the right than the left cerebral hemisphere. However, on tasks requiring sensory perceptual integration the left hemisphere appeared more impaired. Since seizures interrupted testing, the neuropsychological test scores may underestimate his abilities. Indeed, the patient was effusive and at times witty during informal discussion. For example, when asked how he felt about the extensive preoperative testing he said, “I feel as if I am going through World War II without a gun.”

Preoperative EEG showed irregular, anteriorly predominant bilateral spike and wave activity at a rate of 2 to 2½/sec. The background activity was 6/sec (Fig. 1). Computerized tomography (CT) scans showed moderate cortical atrophy, vermic atrophy, and mildly enlarged ventricles (Fig. 2). Positron emission tomography (PET) revealed a region of hypometabolism extending throughout the left temporal lobe.19

Pancerebral arteriography, performed in conjunction with the intracarotid amobarbital study, was normal. There was filling of both anterior cerebral arteries when each internal carotid artery was injected. Only ipsilateral middle cerebral arteries were visualized. The posterior cerebral arteries were visualized when the right vertebral artery was injected. The Mayo Clinic procedure was followed for the amobarbital study.9 Each
Convulsions were observed at monthly intervals beginning the 3rd day following left hemispheric injection and for 5 minutes 56 seconds following left hemispheric injection.

Operation. Corpus callosotomy was performed when the patient was 18 years old. He was placed in the supine, semi-sitting position. The surgical technique followed the procedure described by Wilson, et al.,38,40 in which the corpus callosum is divided without entering the third ventricle. The hippocampal commissure was sectioned, but the anterior commissure was not divided. To divide the genu, rostrum, and anterior portion of the body of the callosum the neck was extended and a coronal incision was placed 9 cm posterior to the nasion, starting 3 cm to the left of midline and extending 8 cm to the right of midline. Two burr holes were placed over the sagittal sinus, 4 cm apart, with the anterior burr hole placed approximately at the intercept of a tangent of the A2 segment of the anterior cerebral artery with the skull. A craniotomy, which extended just to the left of the sagittal sinus and 4 cm to the right, was made with the neurotome. The dura was opened and the medial aspect of the right frontal lobe was retracted laterally to expose the midportion of the body of the corpus callosum. This portion was approached first because the falx was complete posteriorly and the pericallosal arteries could be identified prior to dissecting adhesions between the cingulate gyri. Under × 16 magnification and 300 mm focal length, the corpus callosum was divided with No. 5 and No. 6 French suction devices down to, but not through the blue-appearing ependymal lining of the roof of the ventricle. Working forward between the pericallosal arteries, the genu was divided and the subarachnoid space around the A2 segment of the anterior cerebral arteries was entered. Section was completed where the genu disappeared into the rostrum. Similar techniques were used to section the splenium and posterior body. With the patient’s neck flexed, a 10 cm coronal incision was placed 4 cm behind the midpoint between the nasion and inion, permitting a 5-cm craniotomy which extended just to the left of midline. Retraction of the right parietal lobe permitted recovery of a cottonoid, which had been placed at the posterior extent of the first dissection, and continuation of the section posteriorly. Callosal fibers were then divided posteriorly until the arachnoid membrane overlying the vein of Galen and the cerebellum were seen. In this case, the cingulate gyri may have been entered and fornices may have been touched. (In more recent procedures, we have sectioned the splenium and posterior body through the anterior craniotomy.) Dexamethasone (Decadron, 10 mg) was given every 6 hours for 12 hours preoperatively and for 3 days postoperatively.

Postoperative Course. The patient was seizure-free for the first 6 postoperative months. Five nocturnal convulsions were observed at monthly intervals between the 7th and 12th postoperative months. Six diurnal seizures were seen during the 2nd postoperative year (about one every other month), characterized by shaking of the right arm and leg and stiffening of the left side.

Snout and bilateral grasp reflexes were seen only during the 1st postoperative day and Babinski signs persisted for 3 months. A disconnection syndrome was demonstrated through testing on the 8th postoperative day (the patient was not testable during the 1st postoperative week). With vision obscured, the patient was able to identify (by writing with his right hand) objects placed in his right hand, but not in his left hand. He was unable to identify by writing (with either hand) objects placed in his left hand. However, he could recognize objects through selecting them, by left-hand palpation, from an array of objects. The disconnection syndrome persists to the present time. On the 4th day, he had a buccofacial apraxia and was unable to move his mouth on command to smile, frown, whistle, stick out his tongue, or blow out his cheeks. The apraxia lasted for 16 months. An otolaryngological examination at 3 months and at 1 year showed normal hearing with normal reflex movement of palate and vocal cords. However, no voluntary movement of either was produced upon commands to gag or to pronounce "ee." Transverse and coronal CT sections through the brain were obtained 6 weeks postoperatively without contrast administration. There was a midline lucency which extended from the genu of the corpus callosum posteriorly to the splenium, and probably represented postoperative changes resulting from sectioning the callosum. No parenchymal hemispheric abnormalities were demonstrated (see Fig. 3). Positron emission tomography showed symmetrical metabolism in the temporal lobe regions that had shown left-sided hypometabolism preoperatively.19

The EEG performed 6 weeks and 1 year postoperatively showed low- to medium-voltage 8/sec activity posteriorly and 5 to 6/sec activity centrally and parietally, with the right more predominant than the left. There were no rhythmic slow-wave bursts (Fig. 4). During sleep, spikes occurred independently on the right and the left with a frontal predominance.

The xenon-133 inhalation method for measuring regional cerebral blood flow was performed 2 weeks preoperatively and 6 weeks postoperatively.25,26,28 Flow was measured using 16 sodium iodide scintillation detectors placed over eight homotopic regions of each hemisphere. Preoperative flow was low in all regions. Postoperative flow was at a level comparable to age- and sex-matched normal individuals, with the exception of lower than normal flow in precentral regions.

The patient was mute for the first 16 postoperative months, except for eight brief one-sentence outbursts of speech. For example, on the 4th postoperative day, when electrode paste was placed on his head, he said “this smells like s--.” On the 7th postoperative day, when a nurse commented that he was looking “good”...
Mutism following callosotomy

he responded "that's what everybody is telling me." Sixteen months postoperatively he started to whisper names of objects and his own name, and to repeat simple words. Articulation was difficult and the sound was breathy (produced with audible exhalation) without apparent paraphasias or other aphasias. The buccofacial apraxia improved, except for difficulties with rapid alternating lateral movements of the tongue without using lip support. Currently the patient speaks spontaneously and fluently, but his speech is somewhat pressured and choppy.

The patient received a standardized neuropsychological evaluation 4 months after surgery. Because of his mutism, only nonverbal tests were administered. The patient showed slight improvement on performance IQ (score 48) and a dramatic improvement in measures of perceptual and motor functions (although these were still impaired bilaterally relative to age-matched populations). The patient was tested again 15 months postoperatively. This time he was asked to respond to the verbal tests in writing. Results of this procedure clearly revealed retention of verbal IQ (score 64), and a substantial further improvement of performance IQ (score 58). Memory quotient was essentially unchanged (score 57).

Two years postoperatively, the patient continued to show intellectual improvement (full-scale IQ score 62, verbal IQ score 68, performance IQ score 60). His receptive vocabulary was somewhat poorer (PPVT standard score 55). Sensorimotor tasks also showed continued improvement.

Discussion

Corpus callosotomy has reduced the number of seizures in this patient. It also produced transient mutism. For the first 16 postoperative months, his speech production was limited to a few sentences or statements. He comprehended written and verbal information and was able to communicate by writing and gesturing. The results of the pre- and postoperative observations and

Fig. 3. Electroencephalographic tracings obtained 6 weeks postoperatively while the patient was awake, showing a background activity of 8/sec.

Fig. 4. Computerized tomography scans without contrast material 6 weeks postoperatively at the same levels as in Fig. 2. Left: A lucent defect is visualized in the genu (arrow) and splenium (arrowhead) of the corpus callosum. Right: A lucent defect extends through the body of the corpus callosum from the genu to the splenium (arrows). The frontal horn and body of the lateral ventricles are slightly smaller than preoperatively.
testing may provide a step toward some understanding of possible mechanisms responsible for the postoperative mutism.

The patient’s neurological and neuropsychological functioning remained the same or improved following surgery. Thus, it is unlikely that the mutism was a concomitant of general intellectual deterioration due to direct cortical damage. The syndrome does not resemble apraxia, nor does it resemble mutism following lesions of the supplementary motor cortex. However, on the eight occasions when the patient spoke, his sentences were clear and grammatically correct. By the time speech production had virtually stopped (11th postoperative day), the snout and the grasp reflexes had cleared. The Babinski signs disappeared 3 months after surgery, but the mutism remained. Thus, it is unlikely that these phenomena result from a common cortical mechanism such as bilateral supplementary motor region destruction. In addition, the postoperative CT scan did not reveal edema or structural lesions except for the destruction of the corpus callosum. Finally, the EEG did not show any significant focal slowing or subclinical seizure activity. Regional cerebral blood flow improved postoperatively to normal in all regions except the frontal lobes, where it increased but did not reach normal levels. It is unclear why cutting the corpus callosum is followed by a relative decrease in blood flow in the frontal regions (both middle and anterior cerebral artery territories). However, the reduction in frontal blood flow may indicate a physiological abnormality in regions that include structures important for speech production. Apparently, the neural mechanisms subserving speech were suppressed, not destroyed.

For a full evaluation of individual differences in linguistic organization, more cases are needed, complete with sufficient preoperative data. It should be noted that in this case the intracarotid amobarbital study indicated slow return of speech following each injection. It may be that early cerebral insult or delayed language acquisition resulted in bilateral speech representation making it difficult for the left hemisphere to support speech without input from the right. Thus, in patients in whom postoperative mutism occurs, speech production may require interhemispheric interaction.

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Mutism following commissurotomy


