AGENESIS OF THE CORPUS CALLOSUM

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Complete or partial agenesis of the corpus callosum is uncommon especially in adults; the incidence amongst patients in institutions where numerous air encephalograms are performed is 3% (Carpenter and Druke, 1953). Reil (1812) published the first report of a case, describing the necropsy findings in a 30 year old woman of low mentality; further cases discovered at necropsy were subsequently reported (Urquardt, 1880; Richter, 1886; Bruce, 1889; Dunn, 1889; Cameron, 1917; de Large, 1925). Baker and Graves (1933) reviewed 83 cases, in a wide variety of age groups, first diagnosed at necropsy. The diagnosis during life was first made by Davidoff and Dyke (1934) using pneumo-encephalography; the pneumo-encephalographic features had been observed by Guttman (1929) but their significance was not appreciated until necropsy revealed complete agenesis. Fukai, Oguma and Sujiyama (1960) reviewed 187 cases of complete and partial agenesis; 130 cases were proven at necropsy, 57 diagnosed by pneumo-encephalography only. Since their review a further 22 cases have been reported, 7 diagnosed or confirmed at necropsy, 15 diagnosed by pneumo-encephalography only.

The function of the corpus callosum, originally regarded as the seat of the soul (Mingazzini, 1922), has been much discussed both on the evidence from cases of complete and partial agenesis and from that following surgical division in man and experimental animals. No characteristic clinical syndrome accompanies complete or partial agenesis of the corpus callosum nor are there any specific neurological or psychiatric accompaniments. In some cases of congenital agenesis there are no readily detected deficiencies; the 25th such case, which is also the 17th asymptomatic adult with complete agenesis, is here reported with a short review of the literature.

Case Report

A 65-year-old builder's labourer was admitted to hospital with a three months history of flatulence and epigastric discomfort. He claimed to have been fit and well all his life, a statement corroborated by his relation and medical records. Shortly before the onset of his symptoms, he had attended the Ophthalmic Department with some loss of vision; this was found to be due to cataracts and he was awaiting their surgical removal at the time of admission. There was no relevant family history and no noteworthy abnormality of his central nervous system or mental state.

A radiological diagnosis of carcinoma of the stomach was made and substantiated at laparotomy. Sub-total gastrectomy was performed and the diagnosis confirmed histologically, but bronchopneumonia supervened and he died on the fourth post-operative day.

Necropsy

There was no unusual feature at the site of the Polya gastrectomy; there was no residual tumour or metastasis. The lungs showed the changes of bronchopneumonia, confirmed histologically.

Brain (1173 g.): On removing the dura mater a large volume of cerebro-spinal fluid escaped from a common ventricle. There was no falx cerebri or tentorium cerebelli. The hemispheres were symmetrical and united along their supero-medial borders by a tough membrane (Fig. 1). The corpus callosum and septum lucidum were absent, there being a widely patent common ventricle (Fig. 2). On the inferior aspect, because of the absence of the anterior columns of the fornix and the anterior commissure, there was a free communication with the common ventricle (Fig. 3). There was no abnormality of the Circle of Willis or cerebral arteries. The lateral surface of the hemispheres showed a normal sulcal and gyral pattern, whilst an anterior view similarly presented a normal contour.

After fixation in 15% formal saline and radiographic examination (Fig. 4), the mid-brain was divided transversely and the cerebrum mid-sagittally. The pieces were weighed and compared with another similarly divided brain from a man of similar age and stature (Fig. 5). The lateral ventricles were large with symmetrical dilatation of the anterior and posterior horns; the inferior horns were absent. Whilst fragments of the choroid plexus were attached to the medial surfaces of the hemispheres, no fragments were attached to the walls of the lateral ventricles or their horns. An unusual reflection of the meninges joined the two cerebral hemispheres together, whilst another shorter reflection connected the occipital lobes with the cerebellum (Fig. 6).

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Cross section of this case.

**Fig. 1.**—Diagrammatic comparison of this case with the normal. The corpus callosum and septum lucidum are absent, the two hemispheres being joined along their superomedial borders by a membrane.

Cross section of normal brain.

**Fig. 2.**—View from above through the widely patent common ventricle.

The cerebral hemispheres were cut into coronal slices 1 cm. thick; this revealed a considerable deficiency in the grey and white matter. The anterior commissure, forceps minor, septum lucidum, pineal body, stria godenularis and the genu, body and splenium of the corpus callosum were not found. With the exception of slightly raised areas at the site of their attachments both the anterior and posterior columns and the body of the fornix were also absent. The basal ganglia and the attachments of the cranial nerves, including the optic commissure and tracts, were normal.

There was no abnormality on sectioning the midbrain, pons, medulla oblongata and cerebellum.

**Fig. 3.**—View from below. This shows the free communication with the common ventricle due to the absence of the anterior columns of the fornix and the anterior commissure.
**Fig. 4.**—Radiograph demonstrating the wide common ventricle and its large lateral extensions.

**Fig. 5.**—Diagrams to show the relative sizes of the left hemisphere, right hemisphere and cerebellum (including brain stem).

**Fig. 6.**—Diagram of the medial surface to show the reflection of the meninges joining the two cerebral hemispheres and that between the occipital lobes and the cerebellar hemispheres.
Embryology

The corpus callosum develops from the lamina terminalis where a thickening develops during the third or fourth month of foetal life. This thickening is on the dorsal and cephalic aspects of the hippocampal commissure which has just formed; the inferior part of the thickening becomes the anterior commissure while the remainder forms the corpus callosum (Davidoff and Epstein, 1950).

Growth of the corpus callosum first occurs in a cephalic direction, then dorsally, to form the rostrum and genu. In the final stage growth is in a caudal direction, the corpus carrying the hippocampal commissure on its undersurface. The septum lucidum represents pieces of each cerebral hemisphere which are separated from the remainder of the hemispheres by the growth of the corpus callosum and the formation of the fornix. Development is complete by the end of the fifth month (van Epps, 1953; Sheldon and Pegman, 1953).

The anatomical defects seen in the various types of agenesis are determined by the stage at which normal development is interrupted (Bruce, 1889). Arrest within the first three weeks of foetal life results in complete fusion of the hemispheres and ventricles with no distinct commissures between the hemispheres (Mosberg and Voris, 1954; Marburg, 1949; Slager, Kelly and Wagner, 1957). If arrest occurs at any stage between the end of the first and the beginning of the fourth month, there is complete agenesis of the corpus callosum, septum lucidum, anterior commissure and hippocampal commissure (Jackson, Sinclair and Belber, 1960). Should development terminate late in the fourth month the anterior commissure, rostrum and genu will be present but not the body or splenium of the corpus callosum. Termination of development after the beginning of the fifth month restricts any defect to the body and splenium of the corpus callosum.

On this basis the defects in our case are attributable to arrest in the second or third months of foetal life.

Aetiology

In this case of congenital agenesis, as in most others reported, no causative agent is known. Carpenter and Drukmiller suggested that the basic defect was impaired metabolism due to anoxia; Sander (1868) considered mal-development of the anterior callosal artery with inadequate vascularisation of the neuro-pore to be a cause. However, to attribute age-

ness to a defect in the embryonic blood supply or in the development of the lamina terminalis is merely to suggest the mechanism without determining the fundamental aetiology. Various agents have been suggested.

1. Genetic defect: Naiman and Clark (1955) reported two cases of complete agenesis in siblings.

2. Intra-uterine infection or toxæmia of pregnancy (Wilson, 1954).

3. Transient barriers in embryonic life such as a cyst within the ventricular system (de Morsier and Mozer, 1935; Oftedal, 1959).

Discussion

Complete or partial agenesis of the corpus callosum is diagnosed most commonly in the youngest age groups with no significant difference in incidence between the sexes (Bunts and Chaffee, 1944; Carpenter, 1954). Bunts and Chaffee, in a series of 15 cases, reported 12 under 10 years of age and commented that the majority of cases developed symptoms before this age. Carpenter and Drukmiller reviewed 45 cases with a similar finding; in 25 the onset of symptoms occurred under two years of age, in eight under 10 years of age, and in only two cases reported by them did the onset of symptoms occur after 20 years of age. Hankinson and Amadore (1957) reviewed a further 14 cases and divided them into two groups. There were 11 infants some of whom presented with failure to thrive or with periodic vomiting and irritability so that a subdural hæmatoma or other intra-cranial lesion was considered; the others presented with rapidly or gradually rising intra-cranial pressure. The remaining three cases all presented as epileptics of late onset, the oldest being 46 years. In a series of 111 cases, in whom the age of onset of symptoms was known, Slager, Kelly and Wagner reported 45 cases where the onset of symptoms occurred under the age of 15 years; in the other 66 cases only 19 developed their initial symptoms over the age of 40 years. They commented that severe associated cerebral anomalies seemed to preclude long survival in the majority of cases of agenesis recorded in children. However, complete agenesis has been reported in an 84-year-old female (Kirschbaum, 1947) and partial agenesis in a 73-year-old male (Slager, Kelly and Wagner).

Anomalies frequently reported with complete or partial agenesis of the corpus callosum include dilatation of the posterior horns of the lateral ventricles with thinning of their walls as in our case, non-union of the calcanea and
parieto-occipital sulci by interposition of a superficial gyrus, and radial arrangement of the sulci on the medial surface of the brain as seen in the foetus. Total absence of the septum lucidum and hippocampal commissure is found with complete agenesis though, unlike our case, the body and columns of the fornix are usually present (Baker and Graves, 1933). Other anomalies often reported include cranial nerve defects, incomplete separation of the frontal lobes, hydrocephalus, porencephaly, arrhinencephaly, microcephaly and dilatation of the cisterna magna (Bouchier, 1957). In some cases the anterior commissure is larger than normal (Norman, 1958).

Baker and Graves observed that the mentality of patients with agenesis varied considerably but was fairly well correlated with the severity of the associated anomalies. They suggested that the various symptoms reported in many cases of complete or partial agenesis were a manifestation of the accompanying anomalies rather than of the defect in the corpus callosum. This is supported by cases where the defect follows haemorrhage, neoplasm or surgery; the symptoms probably result from damage to adjacent areas (Kirschbaum, 1947; Greenblatt and Anderson, 1953).

Considerable controversy has occurred regarding the function of the corpus callosum and it is of interest that of 209 reports of complete and partial agenesis 24 were asymptomatic during life. The only significant common feature of these latter cases, which vary in age from 11 months to 76 years, is the accidental discovery of complete or partial agenesis at necropsy.

A definite function for the corpus callosum was suggested by the loss of memory of topography, with visual and tactile agnosia, after the surgical division of the posterior half of the corpus callosum during the removal of a colloid cyst from the third ventricle (Trescher and Ford, 1937). Further evidence of a function was the apraxia attributed to involvement of the callosal fibres by an aneurysm abutting on the corpus callosum (Sweet, 1941).

Evidence against a definite function for the corpus callosum was, for many years, considerable. Cameron (1917) and Dandy (1936) found no abnormality in patients in whom the corpus callosum was surgically divided. Van Wagenen and Yorke Herren (1940) divided the corpus callosum of 10 epileptic patients to prevent generalisation of the fits and found that section of the commissural pathways contained in the corpus callosum could be performed without any untoward effect. Intensive studies were made on these patients and six others, the pre-operative and post-operative findings being carefully compared (Akelaitis, 1941; Smith and Akelaitis, 1942); no evidence of any significant abnormality was found. Other workers found that complete or partial division of the corpus callosum caused dyspraxia in the subordinate or dominant hand only when there was also damage to the subordinate or dominant hemisphere (Akelaitis, Riksteen, Yorke Herren and Van Wagenen, 1942). These authors and later Myers (1956), concluded that the bilateral functions such as motor activity are integrated through commissural pathways at levels below the corpus callosum.

During the past ten years numerous investigations of the processes of inter-hemispheric transfer of learning have demonstrated definite functions for the corpus callosum. Myers (1956; 1961) studied the role of this structure in the transfer of visual learning in cats whose optic chiasma and corpus callosum had previously been divided. Cats trained with one eye masked were unable to remember with the second eye what they had learned with the first eye. Furthermore the second eye could be used to learn the complete reverse of what the cat had been trained to do with the first; this indicated a complete functional independence of the separated hemispheres, a conclusion supported by Sperry, Stamm and Miner (1956). Sperry (1958), studying somesthetic and motor learning in similarly treated monkeys, found that in this species the independence of the separated hemispheres was less clear cut, there sometimes being transfer of learning from one hand to the other. Further investigation into the occurrence of such transfer showed that the corpus callosum when present did play an important transfer role between the hemispheres (Glickstein and Sperry, 1960). Nevertheless it is well established that either forelimb can be controlled from a single hemisphere in a split-brain cat or monkey. Myers (1962), using split-brain animals taught various tasks by presenting one task to one eye and another to the other eye, demonstrated that by careful selection of the tasks used, the absence of interaction, facilitation of learning or interference between the hemispheres could be demonstrated.

The function of the corpus callosum is believed to be that of supplementing the activity of each cerebral hemisphere with both complementary and contrary information about what is occurring in the other hemisphere rather than effecting a bilateral spread of information;
in persons with complete absence of the corpus callosum the organisation of the brain is undoubtedly very different from the normal but much work needs to be done in developing experimental psychological tests on the lines suggested by animal experiments before defects in inter-hemispheric transfer can be detected.

Summary

Agenesis of the corpus callosum is an uncommon abnormality; excluding the case reported, there are some 209 examples of complete or partial agenesis in the literature. All age groups are represented, but many cases are diagnosed in infancy or childhood because of severe symptoms or early death, both attributed to associated anomalies of the brain and to the defect in the corpus callosum. There is, however, a group of 25 asymptomatic cases of complete or partial agenesis; of these 17, including the present case, are adults with complete agenesis. The aetiology of most cases is unknown. The function of the corpus callosum is believed to be that of supplementing the activity of each hemisphere with information from the other hemisphere rather than bringing about a bilateral spread of information; tests for the detection of abnormalities in inter-hemispheric transfer have yet to be evolved.

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