MRI evaluation of pathologies affecting the corpus callosum: A pictorial essay

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Abstract
The corpus callosum is a midline cerebral structure and has a unique embryological development pattern. In this article, we describe the pathophysiology and present imaging findings of various typical/atypical conditions affecting the corpus callosum. Since many of these pathologies have characteristic appearances on magnetic resonance imaging (MRI) and their therapeutic approaches are poles apart, ranging from medical to surgical, the neuroradiologist should be well aware of them.

Key words: Corpus callosum; magnetic resonance imaging; pictorial essay

Introduction
The corpus callosum is the largest commissure consisting of tightly packed white matter tracts connecting the two cerebral hemispheres. It comprises of four parts (from anterior to posterior): (i) rostrum, (ii) genu, (iii) body, and (iv) splenium.

Various pathologies that can affect the corpus callosum and their characteristic appearances on magnetic resonance imaging (MRI) are illustrated in this pictorial essay. Pathologies of the corpus callosum result in typical symptoms of interhemispheric discoordination.

The etiological spectrum ranges from congenital, demyelinating, inflammatory, trauma to neoplasm and ischemia. Callosal abnormalities developing after ventricular decompression fall under miscellaneous conditions, which have characteristic imaging findings.

Congenital
Callosal malformations
After the formation of the massa commisuralis around the 10th week of gestation, the first callosal fibers begin to form. The formation is primarily anterior to posterior, beginning with the genu at 12th week. The rostrum is an exception, as it develops last at 18-20 weeks.[1]

Malformations range from agenesis or hypogenensis to hypoplasia. Agenesis can be complete or partial [Figures 1 and 2]. In partial agenesis, only the splenium and/or rostrum are absent. Imaging features include high-riding third ventricle, parallel and non-converging lateral ventricles, colpocephaly, and inverted Probst bundles indenting the superomedial surface of lateral ventricles.

Callosal dysgenesis is associated with other cerebral malformations such as Chiari II, migration disorders (heterotopias, lissencephaly, schizencephaly) [Figure 1], Dandy Walker malformation, holoprosencephalies [Figure 2], etc.[2]

In semi-lobar holoprosencephaly, the normal sequence of formation of the corpus callosum does not take place.[1] Apart from callosal agenesis, on sagittal images, a dorsal interhemispheric commissure or “pseudo-splenium,” which mimics the splenium, is seen in its place[3] [Figure 2].

Lipoma
As a result of abnormal persistence and differentiation of the meninx primitiva (normal resorption between 8th and 10th weeks of gestation) into lipomatous tissue, a pericallosal lipoma is formed. It generally presents with seizures.[4]

Based on the appearance of lipomas, there are two types: tubulonodular or curvilinear. Tubulonodular lipomas are located anteriorly. They measure more than 2 cm in diameter and have a high association with encephaloceles, frontal lobe anomalies, and callosal malformations. Curvilinear posterior lipomas are thinner, ribbon-like, and measure less than 1 cm in width. They usually arch around the splenium.[5]

On MRI, lesions are uniformly hyperintense on T1- and T2-weighted images [Figure 3], with signal suppression on fat-saturated images.

Disorders of myelination

Hereditary

Krabbe’s disease

Deficiency of lysosomal enzyme galactocerebroside beta-galactosidase[6] results in white matter dysmyelination associated with large macrophages. This autosomal recessive disorder is known as Krabbe’s disease or globoid cell leukodystrophy (GLD). There is an abnormal accumulation of galactocerebroside (component of mature myelin) in the macrophages, leading to the formation of large multinucleated “globoid cells.”[7]

In the early-onset type, patients develop symptoms around the age of 2 years, starting with spasticity, irritability, fever, and developmental delay. Eventually decerebrate rigidity and flaccid paralysis sets in, resulting in death.[8] This early-onset form of GLD usually involves the cerebellar white matter, thalami, dentate nuclei, and lentiform nuclei, as well as the corticospinal tracts. Callosal and white matter atrophy is usually a late manifestation.

The common imaging features in the late-onset type of GLD, in patients more than 2 years of age, are involvement of the pyramidal tracts, corpus callosum, and parieto-occipital white matter, which appear hyperintense on long TR/TE images[9] [Figure 4]. The subcortical U fibers are generally spared.[10]

X-linked adrenoleukodystrophy

The leukodystrophies are a group of disorders characterized by enzyme deficiencies resulting in abnormal formation, destruction, or turnover of myelin.[11] X-linked adrenoleukodystrophy is an autosomal recessive disorder caused due to acyl-CoA synthetase deficiency, which leads to accumulation of very long chain fatty acids in tissues and plasma.[12]

Patients present during their childhood with seizures, visual–behavioral disturbances, and mental retardation. Symptoms and signs of adrenal insufficiency along with undescended testis may also exist.[13]

Of five imaging patterns described by Loes et al.[4] pattern 1 is the most common one (66% of cases) [Figure 5].
It manifests as deep white matter involvement of the parieto-occipital lobes, peritrigonal regions, and splenium of the corpus callosum. Involvement of visual and auditory pathways can also occur. These areas appear hyperintense on fluid-attenuated inversion recovery (FLAIR)/T2W images, hypointense on T1W images, and show variable post-contrast enhancement.

Inflammatory

Multiple sclerosis (MS): MS is an immune mediated, inflammatory demyelinating disorder. The relapsing-remitting form is the commonest clinical presentation, having varied imaging findings on MRI.

On MRI, the MS plaques are hypointense on T1W images and hyperintense on T2W/FLAIR images. They are thin, ovoid, flame shaped (called “Dawson’s fingers”), and oriented perpendicularly to the lateral ventricles [Figure 6B]. This morphology results from inflammatory changes in the white matter surrounding the medullary veins, thus creating dilated perivenular spaces.[13] The specificity for diagnosis increases rapidly with the involvement of the callosal-septal interface,[14] which is seen in up to 93% of patients.[15] [Figure 6A]. The corpus callosum is especially prone to demyelination in MS due to its intimate contact with lateral ventricular roofs and with small penetrating vessels.[16] As the disease progresses, the corpus callosum undergoes atrophy and becomes thinned out.[17] Amongst the newer techniques, diffusion tensor tractography can demonstrate degree of axonal loss in the regions of white matter lesions, including the corpus callosum, extremely well.[18]

“Lesion within a lesion” or the so-called “beveled” appearance is also typical. “Solid” or “open ring” form[19] of gadolinium-enhancement pattern is seen in the acute inflammatory phase, due to disruption of the blood-brain barrier [Figure 6C].
Tumefactive demyelination: The lesions appear “tumor-like” and are greater than 2 cm in size, and are often confused with gliomas and lead to unnecessary biopsies. Features highly suggestive of tumefactive demyelination are:

1. Large lesion with little mass effect and edema [Figure 7]
2. Ring-like or open-ring enhancement [Figure 7] signifying active plaque margins
3. Central dilated veins within the lesion [Figure 7]
4. Decreased perfusion within the lesions
5. Rapid resolution after steroid therapy.

Corpus callosal involvement, either primary or secondary, though not uncommon, is not pathognomonic for tumefactive demyelination. In such cases, it may mimic gliomas or lymphomas and needs to be kept in mind as a differential, to avoid unnecessary biopsies.

Marchiafava–Bignami disease
This condition affects chronic alcoholics and severely malnourished people with vitamin B12 deficiency. It causes demyelination and necrosis of the corpus callosum, with a predilection for central fibers.

In the acute phase, patients present with sudden onset of altered consciousness, seizures, ataxia, and pyramidal signs. On MRI, in the acute phase, the corpus callosum shows diffuse swelling and hyperintensity on T2W images [Figure 8]. In the chronic phase, the patients present with progressive dementia and interhemispheric disconnection. On MRI, diffuse callosal atrophy with multiple areas of focal necrosis are seen within the corpus callosum, with a predilection for the central portion [Figure 9] and sparing of the upper and lower edges. In some cases, focal hypointense areas are seen on T2W images within the corpus callosum, probably due to hemosiderin or lipid-laden macrophages. Signal changes may also be found in the subcortical and periventricular white matter.

Trauma
Diffuse axonal injury
This is a significant cause of morbidity in patients with traumatic head injury due to sudden acceleration-deceleration or rotational forces, which cause axonal shear-strain deformations in the brain.

As per the grading system for DAI (according to the location of small hemorrhagic foci) by Adams in 1989, grade I involves gray-white matter interface, grade II involves the corpus callosum (posterior body and splenium, because of their dural fixity), and grade III involves the dorsolateral part of upper brainstem.

On MRI, small hemorrhages are seen as hypointense foci on T2*-weighted gradient-recalled echo (GRE) or susceptibility-weighted sequences [Figure 10]. Non-hemorrhagic lesions show restricted diffusion on diffusion-weighted imaging (DWI) and hyperintensity on long TR/TE images.

Ischemia
Infarct
The corpus callosum, despite having a rich vascular supply, shows various patterns of infarcts. It is supplied by three main arteries: The pericallosal artery, the anterior

![Figure 7 (A-D): Tumefactive demyelination. A 5-year-old male came with history of recurrent episodes of muscle weakness and seizures. Multiple round to oval lesions are seen in the left parietal and frontal lobes (arrows in A), genu (arrow in B), and body of corpus callosum. They are peripherally hypointense and centrally hyperintense on T2-weighted image (A) and show suppression on FLAIR (B) image. Their margins show partial ring enhancement on post-contrast T1-weighted image (arrow in D) and partial peripheral restricted diffusion on diffusion-weighted image (arrow in C), suggestive of active margins of demyelinating plaques.](image)

![Figure 8 (A, B): Acute Marchiafava–Bignami disease. A 36-year-old chronic alcoholic male came with acute onset of altered consciousness and seizures. On MRI, diffuse swelling of the entire corpus callosum (arrows in A, B) is seen along with increased signal on T2-weighted (A) and FLAIR (B) images.](image)
communicating artery, and the posterior pericallosal artery. The body is mainly supplied by the pericallosal branch of the anterior cerebral artery. The anterior portion of the corpus callosum is supplied by the subcallosal and medial callosal branches of the anterior communicating artery. The splenium is supplied by the posterior pericallosal branch of the posterior cerebral artery.[34]

Infarcts of the corpus callosum are generally a part of large vessel ischemia.[35] Hence, diabetes mellitus and hypertension are the major risk factors as they predispose to atherosclerosis. Focal infarct patterns [Figure 11] are uncommon as the blood supply is by small perforating vessels running perpendicular to the parent vessel, making it difficult for the emboli to enter and lodge[35] within them. The splenium is the most vulnerable, followed by the body and genu.[35]

Ischemia due to hypoxia, especially in the neonatal period, produces a typical diffuse callosal infarct pattern [Figure 12]. Such children have an unfavorable outcome.[36] On MRI, restricted diffusion on DWI is the earliest sign [Figure 11], followed by edema appearing hyperintense on T2W images and hypointense on T1W images. In chronic stages, the changes of gliosis or atrophy are seen [Figure 13].

**Tumors**

*Mixed oligodendroglioma*

Oligodendroglioma is the third most common glioma, with its incidence being 2-5% of all primary brain tumors and 5-18% of all glial tumors.[37] It commonly presents with seizure and headache in adults in the 35-45 years age group.[38]

On MRI, signal intensity of the tumors is characteristically heterogeneous, with a subcortical location. They produce overlying cortical thickening, with calvarial erosion in some cases. They have the highest frequency of calcification (50-90%) in a linear or nodular pattern.[39] Mild, ill-defined contrast enhancement is usually seen.[40] Occasionally, they arise in the frontal lobe and can cross to the contralateral hemisphere via the corpus callosum.

**Figure 9 (A, B):** Chronic Marchiafava–Bignami disease. A 51-year-old male, chronic alcoholic, presented with ataxia and dementia. MRI shows atrophy of the corpus callosum along with areas of focal necrosis in the genu and body which appear hypointense on T1 FLAIR image (arrows in A) and hyperintense on T2-weighted image (arrows in B).

**Figure 10 (A, B):** Diffuse axonal injury (DAI). A 30-year-old male presented with head injury and loss of consciousness. MRI shows multiple foci of altered signal intensity in the body and splenium of the corpus callosum, showing restricted diffusion (arrows in A) on diffusion-weighted image (A) and blooming (arrow in B) on GRE image (B), signifying hemorrhagic DAI.

**Figure 11 (A, B):** Acute splenial infarct. A 25-year-old female came in an unconscious state, with a history of hanging. A focal area of restricted diffusion is noted in the splenium of the corpus callosum (arrows in A, B) on diffusion-weighted (A) and apparent diffusion coefficient (B) images. It was interpreted as an acute infarct.

**Figure 12 (A-D):** Diffuse hypoxia. A 13-day-old baby with asphyxia and respiratory distress at birth presented with decreased tone in all four limbs. On imaging, restricted diffusion is seen in the entire corpus callosum (arrows in A-D) on diffusion-weighted (A, C) and apparent diffusion coefficient images (B, D). This atypical infarct pattern was attributed to diffuse hypoxia.
Histologically, “mixed” tumors also exist with astrocytes as their co-component. These oligoastrocytomas show a slightly higher incidence of contrast enhancement[41]; however, they are indistinguishable from pure oligodendrogliomas on MRI alone[42] [Figure 14]. Pure oligodendrogliomas show longer time to progression and survival period as compared to oligoastrocytomas, on administration of procarbazine, lomustine and vincristine (PCV) chemotherapy.[43] However in practice, it is more important to differentiate between these oligodendrogial tumors and astrocytomas, as the latter show poor response to PCV chemotherapy.[44,45]

**Glioblastoma multiforme (GBM)**

GBM (WHO grade IV astrocytoma) is the most common primary intracranial malignancy, accounting for 12-15% of all intracranial neoplasms.[46] It generally originates in the cerebral white matter and spreads along the corpus callosum in a “butterfly” pattern [Figure 15A] to the opposite side, thus demonstrating its aggressive nature. Subependymal spread and CSF seeding are the other routes of tumor spread.[47] It may also originate in any part of the corpus callosum with an exophytic extension into the lateral ventricles, wherein it may be difficult to be differentiated from a primary intraventricular tumor. However, in such cases, one must look for the location of the tumor bulk, white matter edema, and enhancement,[47] as these findings favor GBM.

It shows marked perilesional edema and heterogeneity within due to areas of necrosis, hemorrhage, and hypercellularity. It also shows intense inhomogeneous post-contrast enhancement in its solid portion as a result of neoangiogenesis.[48]

**Lymphoma**

Primary cerebral lymphoma (PCL) represents 1-2% of all intracranial neoplasms and is particularly seen in patients with AIDS[49] and other immunodeficiency states. Almost all cases of PCL are of the non-Hodgkin’s type.[50]

PCLs have a predilection for the meninges and ependyma and are typically found in the deep basal ganglia, periventricular regions, and corpus callosum[50,51] [Figure 15B].

Histologically, non-cleaved small-cell tumors are more commonly found in the corpus callosum,[52] Their imaging appearances, especially those of AIDS-related lymphomas, are similar to GBM; however, PCLs are more diffuse, generally multiple, less necrotic, have a lower incidence of intratumoral hemorrhage and a lesser degree of perilesional edema. Contrast enhancement pattern is more homogeneous in immunocompetent patients, whereas in patients with AIDS, it shows ring enhancement pattern due to higher degree of central necrosis.[53] Pathologically, the infiltrative edges of the lesions extend along perivascular spaces and infiltrate blood vessel walls; hence, post-contrast enhancement along perivascular spaces is highly suggestive of lymphoma.[53] Rapid regression is seen after steroid administration; however, the prognosis is poor due to high recurrence rates, which are as high as 60% after radiotherapy.[53]

**Infective**

**Reversible lesions in the splenium of corpus callosum**

Reversible lesions in the splenium of the corpus callosum (SCC) [Figure 16], caused by various infective...
agents such as influenza, rotavirus, mumps, *Escherichia coli*, and adenovirus have been documented.\[^{54}\] The changes become apparent on MRI from as early as 2\(^{nd}\) day of onset of symptoms and CSF findings are usually normal or nonspecific. Patients with motor deficits show signal abnormalities in the deep cerebral white matter.

There are other pathologies that can cause transient abnormal signals in the SCC. These include certain antiepileptic drugs (AED), epilepsy itself, posterior reversible encephalopathy syndrome,\[^{55}\] mild encephalitis/encephalopathy with reversible isolated SCC lesion (MERS),\[^{56}\] hypoglycemia, multiple sclerosis, hemolytic–uremic syndrome with encephalopathy, AIDS dementia complex, etc.\[^{57}\]

Over the years, many hypotheses have been suggested to explain these transient lesions of the SCC. In 1999, Kim *et al.*\[^{59}\] proposed reversible demyelination as the reason, probably due to antiepileptic drug toxicity. AED like primidone and carbamazepine were found to reduce plasma arginine vasopressin values by Krause *et al.*\[^{58}\] in 1983. In 2005, Gurtler *et al.*\[^{59}\] proposed that abrupt stoppage of chronic antiepileptic therapy could lead to ischemia and resultant cytotoxic edema, as a result of fluctuations in the arginine–vasopressin system.

In 2004, Tada *et al.*\[^{56}\] studied patients with encephalitis and concomitant reversible lesions in SCC. They postulated that viral antigens or antibodies induced by them could have an increased affinity for the receptors on splenial axons, resulting in inflammatory infiltrates. Hacket *et al.*\[^{60}\] studied cerebral edema occurring at high altitudes and observed similar changes in SCC. They advocated that vasogenic edema involving white matter was the reason for signal changes. This was also concluded by Conti *et al.*\[^{61}\] in 2007 in their study on the transient focal lesion in SCC on MRI, with an attempt to explain underlying pathophysiology.

However, in 2003, Oster *et al.*\[^{62}\] showed reduced apparent diffusion coefficient (ADC) values in these reversible splenial lesions on MRI. They suggested that repeated and excessive electrical discharges along the commissural fibers during seizures caused transient changes in energy metabolism and ionic transport, resulting in rapidly resolving intramyelinic edema.

Therefore, given its transient and reversible nature, this entity needs to be diagnosed accurately in order to avoid unnecessary invasive diagnostic and therapeutic measures.

**Tuberculosis**

India, with its tropical climate and high rate of HIV prevalence, has a large patient population afflicted with tuberculosis (TB). TB is shown to have many typical and atypical presentations.
Besides meningitis, the other form of presentation is granuloma formation. This is the most common form of parenchymal central nervous system (CNS) TB. CNS spread usually occurs via the hematogenous route from a primary focus in the lungs or gastrointestinal tract. The tuberculomas can also evolve from spread of infection through CSF via cortical veins and small penetrating arteries into the adjacent neuroparenchyma.[63] The corpus callosum, a rare site of tuberculoma formation [Figure 17], can be affected through both these routes. In such a case, it may mimic a glioma.[64] Advanced MR techniques are useful in such cases to accurately differentiate between them. On MR spectroscopy, tuberculomas show characteristic prominent lipid and lactate peaks with absent amino acid resonances. Lipid resonances are seen at 0.9, 1.3, 2.0, 2.8, and 3.7 ppm, due to the high lipid content of the central caseous material.[65] Gliomas typically show elevated choline-to-creatine ratio and reduced N-acetylaspartate-to-creatine ratio, indicating cellular proliferation and neuronal destruction.[66] High-grade gliomas show elevated lipid levels at 0.3 and 1.9 ppm, while low-grade gliomas show elevated myoinositol levels at 3.5 ppm.[66]

DWI provides us with ADC values, which are inversely proportional to cellular density[67] and cytotoxic edema. In a study by Gupta et al. conducted in 2005,[68] tuberculomas with T2-hyperintense cores were found to have ADC values ranging from 0.8 ± 0.08 × 10⁻³ mm²/sec to 0.74 ± 0.13 × 10⁻³ mm²/sec. In a study by Yang et al. conducted in 2002[69] on 17 cases of proven cerebral gliomas, they found that the mean ADC values ranged from 0.92 ± 0.27 × 10⁻³ mm²/sec to 1.28 ± 0.15 × 10⁻³ mm²/sec in the solid, non-necrotic, non-cystic components of the tumors. Hence, ADC values of necrotic tuberculomas are comparatively lesser than those of gliomas. Also, ADC values in the peritumoral region of an infiltrative glioma are lower than those of a tuberculoma, as this region contains nests of tumor cells.[70]

Magnetization transfer (MT) has recently improved lesion detection and characterization in TB. It is directly proportional to the protein or macromolecular content of the lesion.[71] Lipids form a large part of caseating tuberculomas and they show no MT.[72] Hence, the MT ratios of tuberculomas are significantly lower than those of gliomas.[73]

Perfusion imaging helps in assessing tumor-related angiogenesis. The leaky nature of these vessels is reflected in the high relative tumor blood volume (rTBV).[66] In comparison, tuberculomas have very low rTBV values due to the absence of angiogenesis.

Tuberculomas can occur anywhere in the brain, most commonly at the gray–white matter junction and basal ganglia, and depending on their signal characteristics and enhancement patterns, they can be classified as non-caseating and caseating tuberculomas. Caseating tuberculomas can be further classified as liquefied and non-liquefied.[74]

Intra-axial abscess formation is another mode of presentation of intracranial TB. On MRI, it appears as centrally necrotic space-occupying lesion with intense peripheral rim enhancement on post-contrast images. Surrounding perilesional edema is also seen.[75]

**Aspergillosis**

Immunocompromised patients are at an increased risk for fungal infection of the CNS. Amongst the fungal pathogens, *Candida* and *Aspergillus* are the main causative organisms of cerebral abscess in these patients.[76] The respiratory system is the primary site for infection via inhalation of spores. In the *Aspergillus* genus, *Aspergillus fumigatus* is the most common infecting agent. It can access the CNS via the hematogenous route from the lungs, or directly from the paranasal sinuses and nasal cavities. Rarely, direct inoculation may take place due to trauma or surgical procedures.[77]

Nonspecific symptoms of headache, lethargy, hemiparesis, and seizures in an immunocompromised patient should raise the suspicion of cerebral aspergillosis.[78]

A distinct feature of *Aspergillus* is its angioinvasive nature, which frequently causes early cerebral infarction, with or without hemorrhage.[77] From the parent artery, the fungi have a tendency to occlude the origins of the smaller perforating arteries.[79] Hence, the common sites of infection include the corpus callosum, basal ganglia, thalam, and brainstem, along with the gray–white matter junction.[79]

Three common imaging patterns are recognized:

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**Figure 17:** Tuberculosis. A 40-year-old male, K/C/O pulmonary tuberculosis + anti-Koch’s therapy (AKT) defaulter, came with fever, headache, and disorientation. Post-contrast T1-weighted image shows a ring-enhancing lesion in the body of the corpus callosum (arrow) suggestive of a tuberculoma.
1. Ill-defined cortical–subcortical lesions appearing hyperintense on T2W images and hypointense on T1W images
2. Multiple ring-enhancing abscesses [Figure 18]
3. Abnormal dural enhancement adjacent to the primary source of infection (e.g., paranasal sinuses).

The abscesses appear hypointense on T1W images, with areas of hyperintensity peripherally or centrally, corresponding to hemorrhage. On T2W images, they appear centrally hyperintense with a peripheral hypointense rim. In a study by Zinreich et al. conducted in 1988, this rim of low signal intensity was attributed to fungal concretions containing iron, manganese, and magnesium that are found in abundance in the fungal hyphae at the periphery.

Hemorrhages occur in approximately 25% of the lesions and are seen as areas of blooming on gradient echo T2W images. Enhancement may range from being peripheral and mild in immune suppressed patients to ring-enhancing lesions in patients with normal lymphocyte and white blood cell counts. Rarely, CNS aspergillosis may also present with a mycotic aneurysm.

Miscellaneous

Post shunt decompression in chronic hydrocephalus

Post-shunt decompression in patients of long-standing hydrocephalus yields typical MRI findings in the corpus callosum.

Prolonged lateral ventriculomegaly causes stretching and impingement of the corpus callosum against the rigid falk cerebri, resulting in chronic ischemia. Rapid decompression of this long-standing hydrocephalus results in transcallosal demyelination.

On MRI, the corpus callosum appears redundant and slightly increased in size, with signal changes appearing hypointense on T1W images and hyperintense on T2W images. Its rostral margin appears irregular due to segmental tethering by branches of the pericallosal artery [Figure 19]. The asymptomatic nature of these changes is assumed to be due to the relative sparing of the splenium.

Cytotoxic/vasogenic edema due to sinus thrombosis

Cerebral venous sinus thrombosis commonly presents as an acute neurological deterioration with symptoms such as headache, seizures, etc. Its etiology is varied with common causes being pregnancy or puerperal period, infection, dehydration, oral contraceptives, hypercoagulable states, and trauma.

The superior sagittal sinus is the most commonly affected, followed by transverse and straight sinuses. On comparison with dural sinus thrombosis, deep venous system thrombosis has a poorer clinical course and outcome, with a high incidence of coma and death. The deep venous system consists mainly of the vein of Galen, internal cerebral veins, the basal vein of Rosenthal, and medullary and subependymal veins. This system drains the inferior frontal lobe, deep white matter of the parietal and temporal lobes, corpus callosum, thalamus, basal ganglia, and the upper brainstem.

The appearance of the venous thrombus on T1W and T2W images depends on the stage of thrombus formation at the time of imaging. With the formation of paramagnetic products such as deoxyhemoglobin and intracellular methemoglobin within the thrombus, GRE sequences show blooming in the region of the sinuses due to magnetic susceptibility, making them highly specific [Figure 20].

Figure 19 (A, B): Post shunt decompression of chronic hydrocephalus. An 8-year-old female, K/C/O tectal glioma (arrow in B, FLAIR image) causing obstructive hydrocephalus, underwent an MRI for follow-up 7 days after palliative shunt decompression. The corpus callosum (arrows in A) appears redundant with irregular rostral margins and focal signal abnormalities, which appear hypointense on T1-weighted (A) and hyperintense on FLAIR images.
By estimation of ADC values on diffusion-weighted sequences, we can differentiate between cytotoxic and vasogenic edema. Cytotoxic edema shows reduced ADC values (0.33 × 10⁻³ mm²/s to 0.58 × 10⁻³ mm²/s) due to disabling of the Na⁺/K⁺ pump. Raised ADC values (around 1.1 × 10⁻³ mm²/s to 1.955 × 10⁻³ mm²/s) are seen in vasogenic edema, possibly due to raised dural sinus pressure. Both these patterns of edema can be seen together, with or without hemorrhage.

It has also been observed that in patients diagnosed with venous sinus thrombosis along with reduced ADC values, the changes of edema have resolved.

Hence, the term “infarct” should be discouraged, and should only be used if signal changes are irreversible and gliosis is demonstrated on follow-up scans.

Conclusion

A number of pathologies affecting the corpus callosum, typical and atypical, have thus been illustrated in this pictorial essay. The various conditions must be kept in mind in order to make an accurate diagnosis, which is essential for appropriate therapy, whether medical or surgical.

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