A Pictorial Review on Reversible Splenial Lesions

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Abstract
Splenium of corpus callosum can be involved in a variety of pathologies causing reversible or irreversible damage. Magnetic resonance imaging (MRI) is a useful investigation to evaluate the same. In spite of the differing etiologies implicated, MRI findings can be quite common. We review the reversible causes of diffusion restriction involving the splenium of corpus callosum and highlight the etiopathologic mechanisms implicated in these pathologies. We further discuss these pathologies in entirety with relevant clinical and laboratory findings helping make definitive diagnosis and guiding appropriate management.

Keywords
► diffusion restriction
► reversible
► splenium

Introduction
The corpus callosum is a very thick bundle of nerve fibers with more than 300 million compact white matter fibers containing both myelinated and unmyelinated axons.¹,²

It comprises four parts:
1. the reflected anterior portion, or the rostrum;
2. the genu, or the anterior bulbar end;
3. the splenium, or the posterior rounded end; and
4. the body, which lies between the genu and the splenium.³

Splenium is the posterior part of the corpus callosum. In Greek, this word means a bandage strip tied around an injury.

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Many pathologies affect the splenium of corpus callosum that could be categorized as infective (tuberculosis, influenza), metabolic (hypoglycemia, hypernatremia), toxin induced (Marchiafava–Bignami), vascular (cerebral venous thrombosis (CVT), migraine, posterior reversible encephalopathy syndrome, PRES), autoimmune disorder (systemic lupus erythematosus, SLE), trauma (diffuse axonal injury, DAI), medications (phenytoin), and miscellaneous (malnutrition, vitamin B12 deficiency, hemolytic uremic syndrome, HUS, high-altitude cerebral edema and status epilepticus).

Classification of Etiology

I. Infective:
   – Tuberculosis
   – Influenza

II. Metabolic:
   – Hypoglycemia
   – Hypernatremia

III. Toxic:
   – Marchiafava–Bignami syndrome

IV. Vascular:
   – PRES
   – CVT
   – Migraine
   – PRES
        – SLE

V. Trauma:
   – DAI

VI. Medications:
   – Phenytoin

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Our aim is to discuss various pathologies causing diffusion restriction in the splenium that shows reversibility during the course of treatment. The vulnerability of splenium to excitotoxic injury is due to the large number of glutamate receptors and high enzymatic activity making it prone to cytotoxic edema and hence diffusion restriction. This excitotoxic edema is thought to be intramyelinic and is not neurotoxic and thus reversible in most of the cases.

The excess glutamate in case of excitotoxic injury comes from one of the three possible ways:

1. Either decreased uptake at the synapse
2. Increased release because of excessive depolarization of neuronal cell membrane
3. Leak from the axonal membrane disruption.4

**Tuberculous Meningitis**

Tuberculosis is one of the most common infectious diseases in developing countries. Disseminated tuberculosis may affect the central nervous system and lead to tuberculous meningitis (TBM). Patient generally presents with fever, altered sensorium, coma, and stroke like episodes.

Ischemic stroke is commonly seen with TBM but rarely it results in hemorrhagic infarcts. It is attributed to both arterial and venous thrombosis. Stroke in TBM occurs in 15 to 57% of patients. It is regarded as a poor prognostic predictor of TBM.

Magnetic resonance imaging (MRI) can show tuberculomas, leptomeningeal, and basal cisternal enhancement, hydrocephalus, and infarcts (Fig. 1).

Although the corpus callosum is involved in various pathologies, it is relatively immune to ischemic injury. One accepted explanation is due to its rich blood supply from both the anterior and posterior circulations. In a study by Chrysikopoulos et al, the authors described the splenium as being most commonly involved in thromboembolic causes of callosal infarction, very likely due to the predilection for posterior cerebral artery versus anterior cerebral artery territory ischemia.5

The pathophysiological mechanisms implicated in cerebral vasculitis are multiple and can include a direct action of the infectious agent on the vessel wall, blood–brain barrier damage by the cytokines, immunological involvement by inducing antigen expression on endothelial cells and forming immune complexes. In late stage, basal exudates may strangle the vessels leading to vascular narrowing and focal weakness. Vascular involvement starts in adventitia and progressively encroaches to involve the entire vessel wall constituting panarteritis tuberculosa. The uncommon complications include aneurysmal dilatation, mycotic aneurysm, thrombus formation, and venous sinus thrombosis.

**Tuberculomas**

Tuberculomas, are an encapsulated form of tuberculomas, leptomeningeal, and basal cisternal enhancement, hydrocephalus, and infarcts (Fig. 1). They are seen in the spectrum of tuberculous meningitis. Magnetic resonance imaging shows T2 hyperintensity with diffusion restriction in the splenium of corpus callosum. This suggests a vasculitic infarct due to tubercular meningitis. (D) Hyperintense signal in splenium of corpus callosum and leptomeningeal enhancement on fluid attenuated inversion recovery postcontrast. (E) A ring enhancing tuberculoma in the right cerebellar hemisphere.

The pathophysiological mechanisms involved in cerebral vasculitis are multiple and are as follows: a direct effect of the organism on the vessel wall, blood–brain barrier damage by the cytokines, antigen expression on endothelial cells and forming immune complexes by various immune-mediated reactions.

At later stage, basal exudates organized and obliterate the vessel leading to luminal narrowing. The adventitia is involved at first and progressively involves the entire vessel wall resulting in panarteritis tuberculosa.

Few rare complications include aneurysmal dilatation of vessel, mycotic aneurysm formation, and cerebral venous sinus thrombosis.6,7

**Influenza**

Influenza is one of the most common respiratory tract infections that may lead to epidemics especially in winter season and cause flu-like symptoms. Disease spectrum includes Reye syndrome, hemorrhagic/necrotizing encephalopathy, and influenza-associated encephalitis.

Multiple areas of T2 hyperintensity and restricted diffusion are seen in the spectrum of influenza-associated encephalitis/encephalopathy involving subcortical white matter, cortex, and the splenium of the corpus callosum.8 Similarly, symmetrical restricted areas may be seen in the basal ganglia, thalamus, and brainstem, cerebellum in acute necrotizing encephalopathy9 (Fig. 2).

Splenial lesion is commonly seen in influenza and probably is a result of intramyelinic edema and inflammatory infiltrates. Intramyelinic edema results in separation of myelin layers leading to reversible diffusion restriction.10
Hypoglycemia

Hypoglycemia is defined as an abnormally low plasma glucose concentration that leads to harmful effects. The plasma glucose cutoff level is less than 70 mg/dL (normal range: 70–110 mg/dL). Most common reason for hypoglycemia in this diabetic era is iatrogenic hypoglycemia, which is associated with insulin and oral hypoglycemic agents.

Glucose serves as the sole metabolic fuel for central nervous system, especially the gray matter where it is the only source of energy. Any reduction in the metabolic demand will result in symptoms ranging from palpitations, sweating, and dizziness to confusion, altered sensorium, focal deficits, seizures, and encephalopathy in severe situations.

The commonly affected areas are hippocampus, occipital and parietal cortex,pons, basal ganglia, corpus callosum, and internal capsule. On MRI, these areas show T1 hypointensity, T2/fluid attenuated inversion recovery (FLAIR) hyperintensity, and restricted diffusion on diffusion-weighted imaging (DWI) if cytotoxic edema has become evident (►Fig. 3).

Reduction in brain plasma glucose level will affect the cellular fluid mechanisms and cause intramyelinic edema. Subsequent energy depletion will lead to intracellular alkalosis, arrest of protein synthesis and at last lead to membrane ion pump failure, thus converting vasogenic edema to cytotoxic edema.

Marchiafava–Bignami Disease

Marchiafava–Bignami disease (MBD) is a rare form of toxic demyelination of the corpus callosum associated with chronic alcoholic consumption, although nutritional deficiencies have also been thought as a cause of this disease entity. It was first described by Marchiafava and Bignami in 1903. Acute MBD patients may present with generalized tonic–clonic seizures, altered sensorium, and coma. It involves not only the corpus callosum but also the hemispheric white matter.

In acute MBD, there is classical involvement of the central portion of the splenium of the corpus callosum with sparing of the dorsal and ventral layers in the form of T2 hyperintensity and diffusion restriction. On follow-up, there can be normal signal intensity of the involved region or development of cystic lesion that is suggestive of chronic changes. This diffusion restriction is caused by edema and early myelin damage that if remyelinas completely is suggestive of reversibility of the lesion (►Fig. 4).

Posterior Reversible Encephalopathy Syndrome

PRES is a neurotoxic condition that occurs in eclampsia, transplant patients on cyclosporine, autoimmune disease such as SLE or Wegener’s granulomatosis, HUS, thrombotic thrombocytopenic, and postchemotherapy. Patients initially present with headache along with altered vision, altered consciousness, and convulsion.
MRI shows T2 hyperintensity in the parietal and occipital cortex and white matter. It can also involve the frontal, parietal, and occipital lobes, with lesser involvement of the temporal lobes. Less commonly seen pattern is vasogenic edema in the watershed or anastomotic border zone (Fig. 5). Cerebral autoregulatory mechanisms work using both myogenic and neurogenic components for constant brain perfusion. In PRES, there is poor myogenic response due to passive over distention of the vessel due to high blood pressure or toxic effects on the endothelium. The posterior circulation is most vulnerable as they are poorly innervated by the sympathetic system and finally there is leakage of fluid into the interstitium and resultant vasogenic edema.

Severe vasogenic edema may progress to cytotoxic edema. DWI hyperintensity is seen in PRES with either low, normal, or elevated apparent diffusion coefficient (ADC) values; such variable ADC values are attributed to a phenomenon called pseudonormalization that results from intravoxel averaging of both cytotoxic and vasogenic edema in areas affected by PRES. Restricted diffusion in cytotoxic edema causes decrease in ADC values and vasogenic edema causes elevation of the ADC values, the effects of cytotoxic edema and vasogenic edema cancel each other or one of the effects overrides the other when the two are combined at the subvoxel level. Restricted diffusion in PRES is irreversible when the effects of cytotoxic edema begin to dominate and ischemia or infarct may result from it.

According to Ay et al, diffusion restriction is seen in areas of massive unchecked vasogenic edema due to increased tissue pressure causing abnormality in the microcirculation and there by cytotoxic edema.

Cerebral Venous Thrombosis

CVT is a condition of focal or diffuse occlusion of venous channels of the brain that includes cortical vein thrombosis and deep cerebral vein thrombosis. CVT, generally a disease of the young, is comparatively rare to arterial stroke. Its clinical evolution is different from other stroke subtypes with symptoms varying from headaches, nausea, vomiting to decreased and altered conscious with signs of papilloedema, cranial nerve palsies, or focal neurological deficits. Pregnancy, malignancy, dehydration, and prothrombotic hematological conditions may predispose to CVT. Most commonly used method for diagnosis of CVT is time of flight MR venography. In approximately half of all patients increased T2 signal intensity associated with CVT is seen with diffusion weighted images showing restricted diffusion. MRI with GRE sequences is sensitive for parenchymal hemorrhages that are flame-shaped irregular zones of lobar hemorrhage typically with cortical and subcortical extension. Restricted diffusion in splenium is seen in cases of internal cerebral vein and straight sinus thrombosis (Fig. 6).

Increase in back pressure results in reversal of flow and increase in velocity in the venous sinuses leading to blood-brain barrier disruption contributing to cytotoxic edema. Neovascularization due to opening of new channels and recanalization of the occluded veins cause accelerated myelination, leading to reversible splenial lesion.

Systemic Lupus Erythematosus

SLE is a complex autoimmune connective tissue disease with multisystem involvement ranging from kidney, hematopoietic systems, joints, skin to central as well as peripheral nervous with overall increased female predilection. The disease commonly presents with neurological symptoms in the form of headache, limb weakness, and seizure and sometimes with hemiparesis, cranial nerve, and sensory involvement. Autoantibodies against host antigens produce inflammatory process and noninflammatory thickness of small vessels with intima proliferation, small vessel obstruction and infarcts, embolism or hemorrhage due to vasculitis.
MRI shows various patterns of disease in the form of white matter hyperintensity that may be small focal area ovoid in shape suggestive of microinfarct, large area in the form of infarct and area of hyperintensity in the cortical gray matter with atrophy of the brain that is disproportionate to the duration of the disease as patients are on steroid\textsuperscript{23} (►Fig. 7).

SLE causes small vessel arteriopathy with reduced flow of blood to the brain and altering the adenosine triphosphate (ATP) pump mechanism. This causes fluid accumulation in the intracellular compartment, leading to diffusion restriction.\textsuperscript{24}

Diffuse Axonal Injury

DAI encompasses a vast majority of intra-axial lesions in traumatic closed head injury from rotational acceleration and deceleration forces producing diffuse shear-strain with significant morbidity and mortality.

T2 hyperintense lesions with diffusion restriction are seen usually at the gray-white junction (grade 1 injury), in the corpus callosum (grade 2 injury) and at the dorsolateral aspect of the upper brain stem (grade 3 injury). Direct contact between the skull and brain parenchyma leads to cortical contusion in the temporal and frontal lobes with small areas of hemorrhage identified by signal blooming on gradient-echo images. Involvement of splenium is not uncommon in DAI and renders poor prognosis (►Fig. 8).\textsuperscript{25}

Traumatic brain injury causes parenchymal damage that may be due to the loss of calcium and potassium homeostasis and release of excitotoxic amino acids, free radicals, and tissue acidosis causing neurotoxic edema that may be the cause of extended period of decreased ADC values in traumatic brain injury.\textsuperscript{26}

Phenytoin

Antiepileptic medications are commonly used for seizure disorders. Therapeutic blood level (normal is 10–20 ng/dL) monitoring of drug is essential to prevent toxic effects of antiepileptic medications. These patients present with nausea, confusion, nystagmus, gait disturbances, coma, and seizures.

Focal brain lesion in the splenium of the corpus callosum is observed with normal or increased blood levels or sudden withdrawal of antiepileptic medications, such as phenytoin, vigabatrin, and carbamazepine (►Fig. 9).

Seizure activity effects of medications act on the arginine–vasopressin system and glutamate receptors, leading to excitotoxic injury and reversible cytotoxic edema in astrocytes or myelin sheaths in the splenium, and propagate to the contralateral hemisphere.\textsuperscript{4}

Another mechanism suggested is by interfering the intestinal absorption of dietary folate, causing folate deficiency leading to encephalopathy, cerebellar atrophy, myelopathy, and peripheral neuropathy.\textsuperscript{27}

Fig. 7 A 48-year-old female presented with systemic lupus erythematosus. T2-weighted axial magnetic resonance imaging reveals (A) bilateral periventricular discrete hyperintense foci with areas of diffusion restriction (B, C) seen in splenium suggestive of vasculitic infarct and other regions of old sequelae to vasculitis. There is generalized cerebral atrophy.

Fig. 8 (A, B) T2 axial image shows hyperintensity with diffusion restriction (B) in the splenium and a hypointense foci in the left half. This represents hemorrhagic foci in a case of diffuse axonal injury. (C, D) T2 and diffusion-weighted imaging follow-up images of the same patient after 9 months show reversibility of the hyperintensity and the restricted diffusion in the splenium.

Fig. 9 Known case of chronic seizures on eptoin with blood eptoin level of 50 µg/mL. T2-weighted axial magnetic resonance imaging (A) and (B) fluid attenuated inversion recovery coronal image show hyperintensity in the central region of the splenium corresponding diffusion restriction (C, D). Follow-up images (E, F) of T2 and diffusion-weighted imaging of the same patient after 1 year show reversibility of the hyperintensity and the restricted diffusion in the splenium.
Epilepsy

Epilepsy is a disorder that causes multiple and recurrent seizures due to recurrent episodes of paroxysmal brain dysfunction from a sudden, disorderly, and excessive neuronal discharge resulting in transient episodes of sensory or motor neurological dysfunction or psychic dysfunction.

There is a transient increase in signal intensity with swelling on peri-ictal T2-weighted and diffusion-weighted images involving the cortical gray matter and subcortical white matter specifically bilateral cingulate gyri, parieto-occipital, parahippocampal, hippocampus region. Hippocampal atrophy after recurrent seizures has been seen in many cases.

There has been a transient T2/FLAIR hyperintensity with gyral swelling and restricted diffusion, specifically the bilateral cingulate gyri, parahippocampal, hippocampal, and parieto-occipital region. Follow-up MRI with recurrent seizures had shown hippocampal atrophy (►Fig. 10).28

Reversible splenial lesions showing diffusion restriction are not uncommon in generalized tonic–clonic seizures. The cause for diffusion restriction is repeated excessive activity of commissural projections from temporal structure, leading to transient disturbance of energy metabolism and ionic transport.29

Migraine

Migraine is a very common condition associated with severe unilateral debilitating headaches accompanied with an aura in some cases with nausea, photophobia, vomiting, and rarely presents with acute neurologic deficits, which may pose diagnostic difficulties.

Fig. 10 A 32-year-old female presented with multiple episodes of generalized tonic–clonic seizures. On T2-weighted axial magnetic resonance imaging, (A) shows hyperintensity in the central fibers of the splenium of corpus callosum. (B) Diffusion restriction is seen in the central fibers of the splenium of corpus callosum. (C–E) Image of the same patient after 6 weeks shows complete reversibility of the T2-weighted hyperintensity and restricted diffusion seen in the splenium in the prior study.

Many causes have been postulated but changes in cerebral vessel reactivity explain the symptoms with initial vasoconstriction of the cerebral vasculature and cerebral ischemia. This results in T2 hyperintensity in the centrum semiovale similar to small vessel ischemic changes. Later, cerebral vasodilatation results in the pulsatile headache (►Fig. 11).30

These patients also showed a reduction in the gray matter density especially in the bilateral frontal and temporal lobes as compared with healthy controls.31 There is a significant increase in risk of silent infarct especially in the posterior circulatory territory of the brain, thus at times affecting the splenium.32

Transient vasoconstriction in migraine is more prolonged in posterior circulation including the splenium causing depletion of ATP and poor functioning of NA+/K+ pump. This causes cellular edema and reduction in the extracellular compartment, leading to transient diffusion restriction.24

Conclusion

Isolated splenial lesion showing diffusion restriction is generally reversible. There are many differentials of reversible splenial lesion and reversibility itself suggests better prognosis. Good clinicoradiological correlation and relevant laboratory investigations help in determining the etiopathogenesis of reversible causes and guide appropriate patient management.

If only single/solitary lesion is seen in splenium, it is almost always reversible and there is no need for further follow-up. Multiple lesions may require follow-up depending on the clinical setting of the individual condition.

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Conflict of Interest
The authors have no conflict of interest.

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