Venous Hemorrhagic Infarct Leading to Delayed Brain Abscess Formation: A Case Report

Anshu Warade1  Alay V. Khandhar1 Charulata Sankhla2 Ketan I. Desai1

1 Department of Neurosurgery and Gamma Knife Radiosurgery, P. D. Hinduja National Hospital and Research Center, Mumbai, India 2 Department of Neurology, P. D. Hinduja National Hospital and Research Center, Mumbai, India

Indian J Neurosurg

Address for correspondence Anshu Warade, DNB (Neurosurgery), Department of Neurosurgery & Gamma Knife Radiosurgery, P.D. Hinduja National Hospital and Research Center, Mumbai 400016, India (e-mail: dranshuwarade@gmail.com).

Abstract

Keywords Venous sinus thrombosis is a common neurovascular problem with multifactorial etiology, infection being one of the common causes. Cerebral abscess causing thrombophlebitis and venous sinus occlusion is a known entity. In contrast, venous hemorrhagic infarct leading to abscess formation is extremely uncommon. We report a rare case of such delayed abscess formation in a venous hemorrhagic infarct secondary to superior sagittal sinus occlusion.

A 52-year-old man arrived in the emergency room with repeated episodes of generalized tonic-clonic seizures, right hemiparesis of power grade 1 on the Medical Research Council (MRC) scale and pure motor aphasia (Broca's aphasia). An urgent computed tomography (CT) scan of the brain revealed a large patchy area of hemorrhage in the left frontoparietal region of the brain. Magnetic resonance imaging (MRI) scan along with magnetic resonance venography (MRV) of the brain (► Fig. 1A) confirmed hemorrhagic venous infarct in the left frontoparietal region of the brain secondary to occlusion of the anterior one-third superior sagittal sinus (SSS) and the anterior portion of the middle one-third SSS (► Fig. 1B). He was treated with low-molecular-weight heparin, antiedema measures, and antiepileptics medications, and was discharged on oral warfarin. At discharge, his motor aphasia had improved and could verbalize monosyllabic words with the right hemiparesis of power grade 3 on the MRC scale.

A month later, he came to the emergency department with altered sensorium and right hemiplegia. CT scan of the brain (► Fig. 1C) showed complete resolution of the hemorrhage with severe white matter edema in the left frontoparietal region of the brain. There was mass effect with a midline shift to the right of 16 mm. He was treated with antiedema measures. Over the period of 1-month, he became alert and had right hemiparesis of MRC power grade 3.

He presented again 4 months later with multiple episodes of right-sided focal seizures with hemiplegia. There was a history of high-grade fever for the last 1 week. MRI of the brain demonstrated two contiguous ring-enhancing lesions in the same area of previous bleed revealing diffusion restriction suggestive of abscess (► Fig. 1D, E). A thorough checkup was done including 2D echocardiography, CT scan of the chest, ultrasonography of the abdomen, and blood cultures. All these investigations failed to demonstrate any septic focus elsewhere in the body. Dental and ENT (ear, nose, and throat) examination revealed no abnormality. He was operated on and a craniotomy drainage of the left frontoparietal abscess was performed. After draining the pus, the thick fibrous capsule was also radically excised (► Fig. 1F). The pus culture failed to demonstrate any growth of organisms. Histopathological examination (► Fig. 1G, H) revealed foamy macrophages, multinucleated giant cells, and fibrous scarring suggestive of chronic abscess. At discharge, the patient was ambulatory without support with right hemiparesis of MRC power grade 3. A 6-week course of antibiotics—linezolid and ceftriaxone—were given. At 6 weeks of follow-up, his hemiparesis had improved (grade 4 MRC) and motor aphasia had completely recovered. A follow-up contrast MRI of the brain showed near-complete resolution of the abscess with a small area of enhancement in the left deep frontal region (► Fig. 1I, J).
Cerebral venous thrombosis is a rare cerebrovascular disorder affecting 5 per million people and accounting for nearly 0.5% of all strokes.\(^1\) It presents commonly with headache, seizures, altered sensorium, focal neurological deficit, and raised intracranial pressure. Thrombosis of the transverse sinus and SSS is more common than that of the straight sinus, cortical veins, and deep venous sinuses.\(^2\) Hemorrhagic infarction due to venous thrombosis is a fairly known entity and approximately 30 to 40% of cerebral venous thrombosis present with intracranial hemorrhage.\(^3\)

Risk factors for development of cerebral venous thrombosis are prothrombotic conditions, namely, antithrombin III deficiency, protein C and S deficiency; hematological disorders like polycythemia and thrombocytopenia; pregnancy; oral contraceptives; and parameningeal infections of the ear, paranasal sinuses or mastoid sinus, dental, oral, or of neck origin.\(^3\)

Cerebral venous thrombosis secondary to brain abscess is a known entity with the primary source of infection being otitis media, mastoiditis, or cardiac origin, and occurs due to thrombophlebitis as a result of infection in the venous sinus, leading to venous sinus occlusion.\(^4\) On the contrary, in our case, there was abscess formation in the hemorrhagic infarct due to venous sinus thrombosis.

Brain abscess complicating an arterial infarct not associated with either surgical or endovascular procedure is rare and only 12 such cases have been reported.\(^5\) However, only one case of venous infarct complicated by intracranial abscess with source of infection being cutaneous hand infection has been reported in the literature.\(^5\) In our patient, all investigations done failed to reveal any primary source of infection. He presented to us with a large intracranial abscess 4 months after the primary insult (\(\text{Table 1}\)).

We propose that the formation of such abscess can be due to blood–brain barrier disruption of the involved region due to hemorrhage, thereby making the area vulnerable to seeding of infection from a distant unknown source, which is similar in case of arterial infarct. Another hypothesis is that it can be due to persistent inflammatory response to blood products or an autoimmune response to the long-standing inflammation causing liquefactive necrosis of the involved neural tissue, further leading to the formation of abscess.
The diagnosis of such abscess may remain concealed as it occurs in the same area of infarct, leading to similar neurological deficits. One should have a high index of suspicion to diagnose such cases when the patient presents with worsening neurology associated with fever. Timely diagnosis and management of these cases is warranted to prevent further worsening of the neurological deficits and to achieve a good outcome.

**Conclusion**

Cerebral abscess causing cerebral venous thrombosis is a known entity, but venous hemorrhagic infarct complicated by brain abscess is extremely rare. A diagnosis of such atypical brain abscess should be considered if the patient presents lately with deteriorating neurological deficits along with fever. Regular follow-up with serial imaging is highly recommended, which leads to early diagnosis and favorable outcome.

**Conflict of Interest**

None declared.

**References**