INTRODUCTION
According to WHO direct estimate, India measures highest number of cases of snake bites worldwide with a mortality of 35,000 to 50,000 cases per year[1]. Predominantly, bites are more frequent among males, in the lower limb and during rainy season[2]. Envenomation occurs in 10-80% of the snake bites[1]. Viper being the most common cause of bite[1-4], local envenomation is the frequent manifestation, followed by haemostatic abnormalities and neurotoxicity[2,3,4]. Coagulopathy, if present, is diagnostic of viper bites in South Asia[1]. Viper bites have been associated rarely with cerebrovascular accidents, most commonly due to hemorrhagic and rarely due to infarct[5]. Ischemic stroke following viper bite is rare. Most common and serious central nervous system complication following snake bite is intracranial hemorrhage. Ischemic stroke commonly involves anterior circulation. Here we describe a polyvalent anti-venom treated patient with multiple infarcts in right temporal and in cerebellar hemisphere.

CASE REPORT
We report a case of posterior circulation ischemic infarction following viper bite in a previously healthy man. A 75-year-old healthy man with history of DM type 2 on OHA was admitted with history of snake bite on his right foot. The snake was identified as Russell’s viper, as per the descriptions given by relatives. A few minutes after the bite, patient noticed minimal swelling over the right foot and over ankle joint. Two hours later, patient deteriorated and became unconscious. Patient was taken to local hospital. CT scan brain was done which was normal. There was no history of convulsion or any bleeding manifestations. Patient was hemodynamically stable. He was treated with 30 vials of polyvalent anti-snake venom (ASV).

Key words: Snakebite; Viper; Infarct; Ischemic stroke
Paul R et al. Ischemic stroke after viper bite

Next day, after 24 hrs of snake bite patient developed weakness of right upper limb and lower limb. Repeat CT scan brain was done. Patchy hypodensities in bilateral cerebellar lobes, the superior vermis and the right inferior temporo occipital lobe were observed (Figure 1). Patient referred to our hospital for further management. On arrival patient was drowsy with weakness of right upper and lower limb, pupils were bilaterally reacting. Chest X-ray showed bilateral infiltrates. Complete blood count showed leukocytosis, PT, aPTT were prolonged, mildly elevated total bilirubin and creatinine (Table 1), 2D echo was normal. In view of increased breathing effort patient was intubated and kept on ventilator support. Tracheostomy was performed in view of early weaning, pooling of secretions and weak cough reflex. Patient was treated with prophylactic antibiotics, sedation, ventilation and physiotherapy. Patient responded to treatment and discharged in stable condition.

DISCUSSION

Viper bite is the most frequent snake bite in the Indian subcontinent. The presentation of envenomation by viper bite leads to local envenomation, followed by abnormal coagulation[6]. The various toxins present in the viper venom can be categorized both as procoagulant and anticoagulant. The toxins with well established procoagulant/platelet aggregating properties are cerastobin[9], factor IVa[7], cerastocytin[8], cerastotin[9], and afacytin[10]. These various protein products have thrombin like enzymatic activity. Different toxins activate different parts of the coagulation cascade[6-10]. There activity is inhibited by monoclonal antibodies against GP1b or GPIIb/IIIa or thrombin receptor.

Disseminated intravascular coagulation and hypotension are also the associated risk factors in viper bite patients. Disseminated intravascular coagulation can be a cause of neurological disorder, largely due to vessel occlusion[11]. The toxin itself can cause vasospasm which can lead to a cerebrovascular accident.

Reported case reports of infarct following snake bite are majorly due to viper bite. Most of the patients had features of local envenomation with disseminated intravascular coagulation. The infarct commonly involved the anterior circulation, with hemiparesis being the frequent presentation[12-15]. As a matter of fact, cerebral infarction after snake envenomation is a complex multifactorial mechanism. It includes direct cardiotoxic effects of venom leading to dysrrhythmias. This may cause cardiac thromboembolism and hyperviscosity due to hypovolemia and hypoperfusion secondary to hypotension. In a study by Thomas, et al[16] it has been demonstrated that administration of the anti-snake venom within six hours of the viper bite, may prevent associated complications.

CONCLUSION

After viper bite, neurological manifestations can be attributed to various reasons, such as toxin induced vasculitis, procoagulant effect, endothelial damage, disseminated intravascular coagulation and hypotension. Needless to say, in a snake bite prone country like India, it is necessary to evaluate the incidence of ischemic stroke or neurological disorder and possible viper bite. The early detection of...
the viper bite and administration of anti-snake venom can prevent this rare but devastating complication.

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