# Hemispheric Infarct Following a Cerebellar Hematoma: A Rare Coincidence

#### **Abstract**

Concomitant cerebral infarction developing soon after a parenchymal intracerebral hemorrhage is a rare occurrence. Usually, these remote site changes follow tumor decompression and are associated with hemorrhagic changes rather than infarcts. We report a case of a fatal malignant internal carotid territory infarct in a hospitalized patient being conservatively managed for a vermian hematoma and discuss the probable pathophysiology. Stroke physicians need to be aware that spontaneous intracerebral hematoma patients have a potential threat of developing large vessel occlusion with malignant cerebral infarcts, especially after surgical decompression. Although the exact pathogenesis is unknown, size of the clot, intraventricular hemorrhage, hydrocephalus, and aggressive reduction of blood pressure appear to be predictive factors.

**Keywords:** Hematoma, intracerebral, parenchymal, vermian

## Introduction

Remote site bleeds are often encountered after tumor decompression and haematoma evacuation, in clinical practice. However the occurence of a supratentorial infarct following a infratentorial cerebellar bleed is sparsely reported and often rarely encountered. Here we describe a case of a 73-year old patient who presented to us with a primary cerebellar haematoma which progressed to a supratentorial infarct following conservative management. We discuss the clinical dilemma, theories and management outcomes in such patients with a thorough review of literature.

## Case Report

A 73-year-old diabetic, hypertensive, male presented with a history of sudden-onset altered sensorium. On admission, nearly 4 h after the ictus, he was opening eyes to call, obeying commands, and had a slurred speech. His blood pressure (BP) was high on admission (200/120 mmHg) and his blood sugars were elevated (236 gm%). Computed tomographic (CT) scan of the brain showed a posterior fossa hematoma with early obstructive hydrocephalus [Figure 1]. He was managed conservatively as his family members were not willing for surgery. His sensorium remained intact and

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he gradually improved with conservative measures. On the 5<sup>th</sup> day after admission, his sensorium deteriorated and he developed a left-sided weakness. Repeat CT scan showed a large right hemispheric infarct [Figure 2]. He was electively ventilated, and a decompressive surgery was offered to the family who decided against surgery. In spite of elective ventilation and full decongestants, he succumbed on day 5 of the second ictus.

## Discussion

Cerebral infarction following spontaneous intracerebral hematomas (SICHs) uncommon. Wang et al. observed an incidence of 8% infarcts in their cohort of 212 patients with SICH.[1] Prabhakaran et al. detected a 22.9% prevalence of associated infarcts as evidenced by diffusion-weighted image (DWI) abnormalities in a cohort of 118 SICH patients.<sup>[2]</sup> The majority of these infarcts, however, were small subcortical and subclinical. Large vessel occlusion causing malignant cerebral infarction has been rarely reported.[3,4] We add a case of vermian hematoma with infarct to our earlier reported two cases of malignant infarction following evacuation of a supratentorial SICH.

Several mechanisms have been postulated to explain the possible occurrence of an infarct following a spontaneous hematoma.

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# R. Girish Menon, Vinod Kumar, Laskhman IK, Rajesh P. Nair

Department of Neurosurgery, Kasturba Medical College, Manipal University, Manipal, Karnataka, India

Address for correspondence:
Dr. Vinod Kumar,
Department of Neurosurgery,
Kasturba Medical College,
Manipal University,
Manipal, Karnataka, India.
E-mail: vinodneuro@gmail.com



Author	Age/sex	ried cases of remote site bleed for Primary diagnosis	Site of bleed	Management	Outcome
Konig et al.	56/male	Meningioma Meningioma	Cerebellar bleed	Conservative therapy	Dead
van Calenbergh	42/female	· ·	Cerebellar bleed	EVD	Dead
	59/female		Cerebellar bleed	EVD	Good
	58/male	Metastasis of a keratinizing	Cerebellar bleed	Conservative therapy	Good
van Calchoeign	30/maic	epithelioma	Cerebellar bleed	Conservative therapy	Good
Kuroda et al.	63/male	Pituitary tumor	Cerebellar bleed	EVD, VPS	Good
	72/male	Tuberculum sellae meningioma	Cerebellar bleed	VPS, decompressive surgery	Good
	58/female	Sphenoid ridge meningioma	Cerebellar bleed	Conservative therapy	Good
Brisman et al.	73/male	Tuberculum sellae meningioma	Cerebellar bleed	NA	Good
Papanastassiou et al.	54/female	Suprasellar meningioma	Cerebellar bleed	EVD, decompressive surgery	Disabled
Cloft et al.	47/male	Sphenoid ridge meningioma	Cerebellar bleed	NA	Good
Tomii et al.	37/male	Craniopharyngioma	Cerebellar bleed	Conservative therapy	Good
Friedman et al.	64/male	Metastasis	Cerebellar bleed	Conservative therapy	Good
	36/male	Glioma	Cerebellar bleed	EVD	Good
	53/male	Glioma	Cerebellar bleed	Conservative therapy	Disabled
	47/female	Schwannoma	Cerebellar bleed	Conservative therapy	Good
		Tuberculum sellae meningioma	Cerebellar bleed	Conservative therapy	Disabled
	34/male	Craniopharyngioma	Cerebellar bleed	Conservative therapy	Good
	55/male	Metastasis	Cerebellar bleed	Conservative therapy	Good
	36/male	Glioma	Cerebellar bleed	EVD	Good
	53/male	Glioma	Cerebellar bleed	Conservative therapy	Disabled
	47/female	Schwannoma	Cerebellar bleed	Conservative therapy	Good
	47/female	Tuberculum sellae meningioma	Cerebellar bleed	Conservative therapy	Disabled
Honegger et al.	54/male	Intraventricular meningioma	Cerebellar bleed	Decompressive surgery	Disabled
	28/male	Ganglioglioma	Cerebellar bleed	NA	Good
	33/male	Astrocytoma	Cerebellar bleed	NA	Good
Marquardt <i>et al</i> .	31/male	Histiocytoma	Cerebellar bleed	EVD, decompressive surgery	Disabled
	42/male	Glioma	Cerebellar bleed	Conservative therapy	Good
	73/male	Glioma	Cerebellar bleed	Conservative therapy	Disabled
	44/male	Glioma	Cerebellar bleed	EVD	Disabled
Siu et al.	64/male	Temporal tumor	Cerebellar bleed	EVD	Dead
Brockmann et al.	58/female	Temporal meningioma	Cerebellar bleed	Conservative therapy	Good
Yang et al.	15/male	Pleomorphic xanthoastrocytoma	Cerebellar bleed	Conservative therapy	Good
Amini et al.	36/female	Oligodendroglioma	Cerebellar bleed	Conservative therapy	Good
	53/male	Glioblastoma	Cerebellar bleed	Conservative therapy	Good
Sasani et al.	14/male	Dysembryoplastic neuroepithelial tumor	Cerebellar bleed	Conservative therapy	Good
Mandonnet et al.	49/male	Meningioma	Cerebellar bleed	Decompressive surgery	Good
Rezazadeh et al.	60/male	Meningioma	Cerebellar bleed	Conservative therapy	Good
Paul et al.	23/male	Xanthoastrocytoma	Cerebellar bleed	NA	NA
Huang et al.	45/female	Sphenoid ridge meningioma	Cerebellar bleed	Conservative therapy	Good
	66/female		Cerebellar bleed	Conservative therapy	Good
	18/male	Suprasellar tumor	Cerebellar bleed	Conservative therapy	Good
	59/female	Oculomotor nerve tumor	Cerebellar bleed	Conservative therapy	Good
	65/male	Meningioma	Cerebellar bleed	Conservative therapy	Good
Dincer et al.	43/male	Astrocytoma	Cerebellar bleed	Conservative therapy	Good
	49/female	Sphenoid ridge meningioma	Cerebellar bleed	Conservative therapy	Good
	44/male	Oligodendroglioma	Cerebellar bleed	Conservative therapy	Good
		Astrocytoma	Cerebellar bleed	Conservative therapy	Good
Hara et al.	44/male	Anaplastic oligoastrocytoma	Cerebellar bleed	Conservative therapy	Good

Contd...

Table 1: Contd									
Author	Age/sex	Primary diagnosis	Site of bleed	Management	Outcome				
Landeiro <i>et al</i> .	45/male	Ruptured Acom	Cerebellar bleed	Conservative therapy	Good				
	58/female	Left frontal convexity meningioma	Cerebellar bleed	Right FTOZ and 3 <sup>rd</sup> nerve decompression	Good				
	42/female	Left cavernous sinus meningioma	Cerebellar bleed	Conservative therapy	Good				
Nam et al.	61/male	L3-L4, L4-L5 canal stenosis	Bilateral cerebellar	SOC	Good				
Chadduck	59/male	Cervical canal stenosis	Left cerebellar	SOC + EVD	Good				
Mikawa et al.	75/male	Cervical fusion	Bilateral cerebellar	SOC + EVD	Good				
Andrews and Koci	36/male	Lumbar spinal scoliosis	Bilateral cerebellar	EVD	Good				
Gabel et al.	40/female	Lumbar disc	Right cerebellar	SOC + EVD	Good				
	57/female	Lumbar spondylolisthesis	Bilateral cerebellar	EVD	Good				
Satake et al.	62/male	Cervical intramedullary tumor	Cerebellar	SOC	Good				
Morandi <i>et al</i> .	34/male	Cervical schwannoma	Right cerebellar+left temporal	Conservative therapy	Good				
Friedman et al.	43/male	Thoracic herniated disc	Right cerebellar	Conservative therapy	Good				
	56/male	Lumbar spinal stenosis	Bilateral cerebellar	Conservative therapy	Good				
Thomas et al.	38/female	Thoracic tumor	Left cerebellar+right temporal	Conservative therapy	Good				
Karaeminogullari et al.	73/female	Lumbar spinal stenosis	Left cerebellar	SOC	Good				
Brackmann et al.	52/male	Lumbar spondylolisthesis	Bilateral cerebellar	EVD	Good				
Kanya et al.	48/female	Lumbar herniated disc	Bilateral cerebellar	Conservative therapy	Good				
Calisaneller et al.	67/female	Lumbar spondylolisthesis	Bilateral cerebellar	Conservative therapy	Good				
Agrawal et al.	47/female	Posterior fossa meningioma	Diffuse SAH	Conservative therapy	Good				
Garg et al.	28/female	Right sylvian fissure arachnoid cyst	Bilateral frontal EDH	Craniotomy and evacuation of EDH	Good				
	50/female	Left vestibular schwannoma	Diffuse SAH	VPS	Good				
	20/female	Left subdural hygroma	Supratentorial diffuse hematoma	Conservative therapy	Good				
	38/female	Right vestibular schwannoma	Left parieto-temporal EDH	Craniotomy and evacuation of hematoma	Good				
	64/male	Cerebellar hematoma	Left fronto-parieto-temporal SDH	Craniotomy and evacuation of hematoma	Good				
	13/female	Left lateral ventricular neurocytoma	Right frontal EDH	Craniotomy and evacuation of hematoma	Good				

 $SAH-Subarachnoid\ hemorrhage;\ EDH-Extradural\ hemorrhage;\ VPS-Ventriculoperitoneal\ shunt;\ EVD-External\ ventricular\ drainage;\ NA-Not\ available;\ FTOZ-Fronto-temporo-orbito-zygomatic;\ SOC-Suboccipital\ craniectomy;\ Acom-Anterior\ communicating\ artery$ 

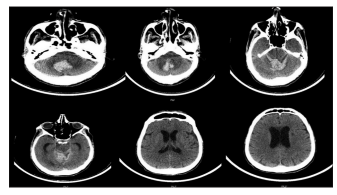


Figure 1: Computed tomography brain showing vermian bleed

The majority of such infarcts are small and subcortical, conforming to border-zone or single perforator territories. Few recent studies have observed subclinical DWI lesions in 15% of patients with acute intracerebral hematoma (ICH) attributable to cerebral amyloid angiopathy. Since both

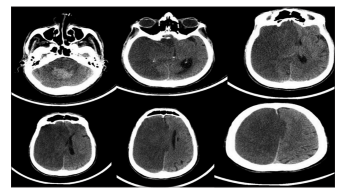


Figure 2: Computed tomography brain showing massive right internal carotid artery territory infarct

ischemic and hemorrhagic stroke share common risk factors and certain common pathogenic mechanisms, it is possible that an ischemic stroke may simply be a co-occurrence in the presence of common risk factors. Hypotension due to attempted aggressive reduction of BP following a hypertensive bleed may precipitate an ischemic event. Surgery for evacuation of the hematoma may be associated with intraoperative hemodynamic instability, which can result in an ischemic event. Though unlikely, infarcts following craniotomy may also be attributable to iatrogenic compression of vascular structures during craniotomy or through the durotomy defects. Large hematomas with surrounding edema can theoretically directly compress the adjacent cerebral vessels causing ischemia. Infection with persistent fever, dehydration, and electrolytic imbalance can all result in a hypercoagulable state following a bleed and can result in ischemia. One another postulated mechanism for ischemia is the massive release of blood and blood breakdown products into the CSF and subsequent inflammatory changes in the smooth muscle of the large cerebral arteries. Cerebral vasospasm, due to the presence of concomitant intraventricular hemorrhage or after indirect vessel manipulation at the time of craniotomy. is another suspected mechanism for ischemia. After acute brain injury, autoregulation may be abolished such that cerebral blood flow is linearly related to cerebral perfusion pressure. Aggressive BP lowering beyond the lower limits of cerebral autoregulation might induce cerebral ischemia in chronic hypertensive ICH patients. Our patient did not undergo surgery and the infarct and hematoma were located in different compartments. He did develop features of early sepsis while under treatment for vermian hematoma. Sepsis with an associated hypercoagulable state may be the precipitating factor resulting in the infarct.

Attempts have been made to predict the risk of developing concomitant infarcts in patients with SICH. Wang *et al.* observed that the presence of intraventricular hemorrhage (IVH), hydrocephalus, the volume of intracranial hematoma, and neurosurgical intervention were important predictors of infarction of which IVH had the most statistical significance. [1] Similarly, Prabhakaran *et al.* observed that the factors independently associated with DWI abnormality were a prior ischemic stroke, lowering of mean arterial pressure by over 40%, and craniotomy

for ICH evacuation.<sup>[2]</sup> Our patient did have the evidence of fourth ventricular blood but did not satisfy most of the above criteria.

The rarity of such complications makes it less likely to appear in our clinical diagnosis and we feel one should consider this scenario as well. The remote site hemorrhages which are often encountered after tumor or hematoma decompressive surgery are shown in Table 1.

The exact etiopathogenesis of these large infarcts remains uncertain and the management of such large infarcts needs to be on similar lines as for any ischemic infarct. The outcome, however, is usually grave.

## **Conclusion**

Stroke physicians need to be aware that SICH patients have a potential threat of developing large vessel occlusion with malignant cerebral infarcts, especially after surgical decompression. Although the exact pathogenesis is unknown, size of the clot, IVH, hydrocephalus, and aggressive reduction of BP appear to be predictive factors.

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### **Conflicts of interest**

There are no conflicts of interest.

## References

- Wang HC, Lin WC, Yang TM, Lin YJ, Chen WF, Tsai NW, et al. Risk factors for acute symptomatic cerebral infarctions after spontaneous supratentorial intra-cerebral hemorrhage. J Neurol 2009;256:1281-7.
- Prabhakaran S, Gupta R, Ouyang B, John S, Temes RE, Mohammad Y, et al. Acute brain infarcts after spontaneous intracerebral hemorrhage: A diffusion-weighted imaging study. Stroke 2010;41:89-94.
- Agarwal A, Satish Kumar S, Umamaheshwara RV. Massive cerebral infarction following evacuation of intracerebral hematoma. Rom J Neurol 2014;13:141-3.
- Kim CH, Kim JS. Development of cerebral infarction shortly after intracerebral hemorrhage. Eur Neurol 2007;57:145-9.