



68 (BP) of 113/77 mmHg, pulse rate 84 beats per min-  
69 ute, and normal blood sugar levels. The patient report-  
70 edly improved initially but experienced intermittent ear  
71 ringing.

72 Prior to his arrival at our facility, the patient had  
73 breakfast that morning and went out to attend to his daily  
74 activities. Later, his mother received a call from a good  
75 samaritan reporting that he suddenly experienced diz-  
76 ziness, one episode of vomiting, headache, and slurred  
77 speech. He was then taken to the local dispensary, where  
78 his blood pressure was elevated at 149/94 mmHg, and he  
79 was diclofenac for pain relief. Subsequently, he devel-  
80 oped excessive drooling and became unable to drink.  
81 Approximately ten minutes after the drooling began, he  
82 lost consciousness. There was no history of trauma. Due to  
83 the development of right-sided weakness and numbness,  
84 along with elevated blood pressure, he was referred to our  
85 higher-level facility, although no hypertensive emergency  
86 was documented at the time.

87 On arrival, the patient was unconscious with a Glasgow  
88 Coma Scale (GCS) score of 7/15. He was snoring with  
89 excessive drooling and frothing at the mouth, no pallor,  
90 jaundice, clubbing, edema, nor lymphadenopathy. Vital  
91 signs were stable. Neurologic examination showed pupils  
92 equal and reactive to light without facial asymmetry. No  
93 signs of meningeal irritation. Motor examination revealed

94 right-sided hemiparesis with decreased muscle strength. 94  
95 Muscle tone on the right side was reduced but not com-  
96 pletely flaccid. Deep tendon reflexes on the right side were  
97 diminished, and Babinski's sign was positive, indicating  
98 an upper motor neuron lesion. The left side had normal  
99 tone and reflexes. No clonus was elicited. Cranial nerve  
100 testing and Sensory testing were limited due to reduced  
101 consciousness. Gait, balance, and coordination could not  
102 be assessed due to mental status. The rest of the systemic  
103 examination were unremarkable.

104 Malaria testing via thick and thin films confirmed  
105 *Plasmodium falciparum* infection with a parasite density  
106 of 11%. This level of hyperparasitemia and the presence  
107 of neurological deficit satisfy the WHO diagnostic crite-  
108 ria for severe malaria. Hepatitis serology panel showed  
109 non-reactive results, syphilis and HIV serology tests were  
110 non-reactive, and dengue rapid tests were all negative.  
111 Additionally, an autoimmune panel was performed and  
112 showed no evidence of systemic lupus erythematosus  
113 (SLE) or other causes of vasculitis and acquired thrombo-  
114 philia (Table 1). These results helped exclude infectious  
115 and autoimmune etiologies in the differential diagnosis.  
116 Lipid profile and fasting blood glucose were also nor-  
117 mal, ruling out common metabolic risk factors for stroke.  
118 Blood cultures showed no growth.

**Table 1.** Initial laboratory tests.

Laboratory parameters	Patient's values	Normal range
Leukocyte count, x 10 <sup>9</sup> per l	13.15	4.00-10.00
Neutrophil count, x 10 <sup>9</sup> per l	10.14	2.00-7.00
Lymphocyte count, x 10 <sup>9</sup> per l	2.29	0.80-4.00
Hemoglobin, g/dl	15.0	13.0-18.8
Platelet count, x 10 <sup>9</sup> per l	266	150-350
Random blood glucose, mmol/l	6.9	4-10
Erythrocyte sedimentation rate (ESR), mm/hour	0.00	0-7.8
C-reactive protein (CRP), mg/l	27.9	0-10
Creatinine, mmol/l	48.0	62-121
Blood urea nitrogen, BUN, mmol/l	3.58	1.8-8.3
Aspartate Aminotransferase (AST), IU/l	48.8	5.0-41.0
Alanine Aminotransferase (ALT), IU/l	56.0	5.0-41.0
Serum Sodium, mmol/l	141	135-145
Serum Potassium, mmol/l	3.8	3.5-5.5
Serum Chloride, mmol/l	103	98-108
D-dimer	463	0.00-500
Prothrombin time (PT), Seconds	20.7	11-15
Internal Normalized Ration (INR)	1.83	0-1.6
Activated Partial Thromboplastin Time (aPTT)	30.1	25.8-38.8
C- ANCA, AU/ml	5.37	16-20
P-ANCA, AU/ml	1.18	< 20.0
Anti-nuclear antibodies (ANA)	Negative	-
Anti-phospholipid antibodies	Negative	-
Rheumatoid Factor	Negative	-

119 Radiographic imaging included a non-contrast brain CT  
 120 scan, which was normal. However, further evaluation with  
 121 non-contrast brain MRI, showed right cerebellar, medulla,  
 122 and pontine acute multiple ischemic infarcts without hem-  
 123 orrhage (Figure 1). Echocardiography demonstrated nor-  
 124 mal cardiac structure and function without any evidence of  
 125 atrial thrombus or patent foramen ovale. An electrocardio-  
 126 gram showed a normal sinus rhythm without arrhythmias.

127 These findings excluded common causes of stroke,  
 128 supporting a diagnosis of malaria-associated ischemic  
 129 stroke as a neurological complication of *P. falciparum*  
 130 infection in this young adult patient.

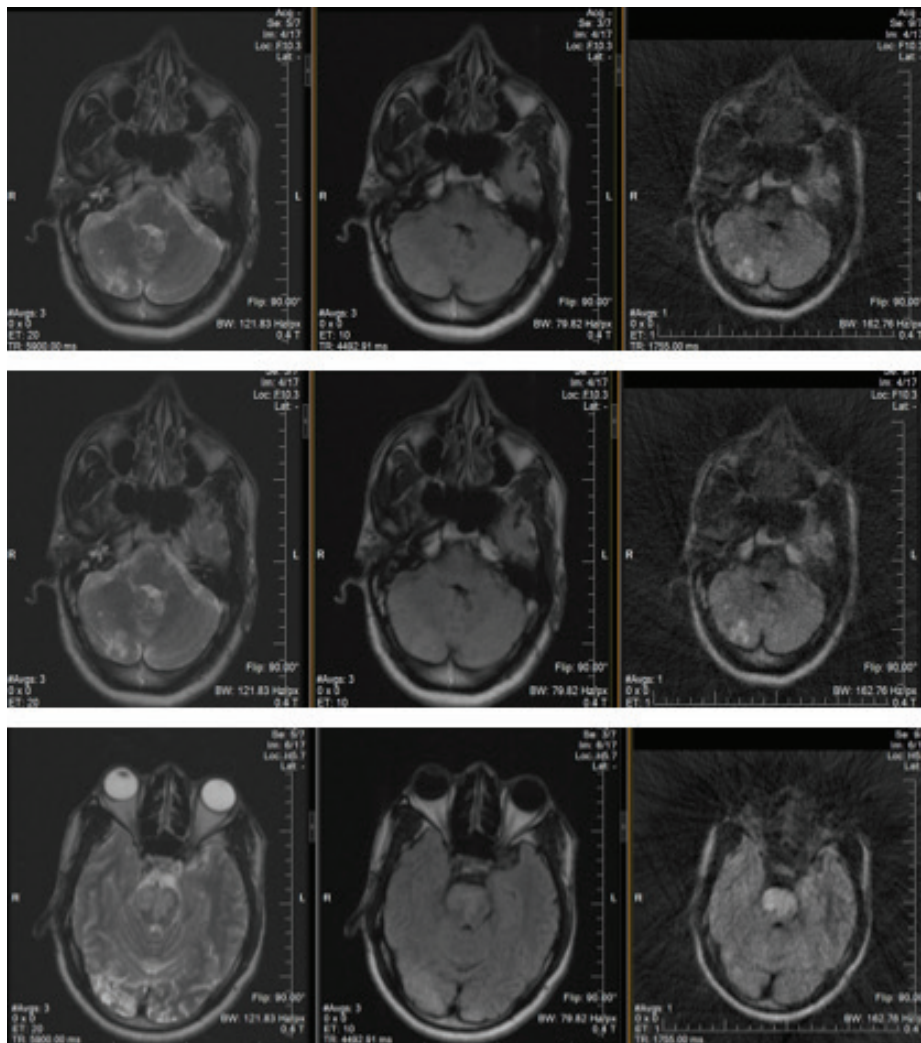
131 The switch from oral artemether-lumefantrine to IV  
 132 artesunate was necessitated by the patient’s transition from  
 133 uncomplicated to severe malaria; therefore, the patient  
 134 was started on IV artesunate in accordance with WHO  
 135 protocols. Enoxaparin was administered at a prophylac-  
 136 tic dose initially, as CT scan results were inconclusive.  
 137 Prophylactic enoxaparin was started due to the patient’s

138 deranged coagulation profile (Table 1) and high risk of  
 139 venous thromboembolism.

140 Once an ischemic stroke was confirmed via brain MRI,  
 141 antiplatelet therapy with aspirin daily was initiated, and  
 142 the enoxaparin was discontinued. Supportive stroke care  
 143 included airway protection, seizure prophylaxis with  
 144 Levetiracetam, vitamin B6, and vitamin B12.

145 This combined approach targets both the malaria infec-  
 146 tion and ischemic stroke, optimizing neurologic recovery  
 147 and reducing mortality risk.

148 The patient responded well to the treatment regimen.  
 149 Repeat blood smears, performed after the completion of  
 150 the artesunate course, confirmed the clearance of para-  
 151 sitemia. Neurological function gradually improved, with  
 152 partial recovery observed in both motor and cognitive  
 153 abilities. The patient was discharged two weeks later with  
 154 a Glasgow Coma Scale (GCS) score of 11/15. Close moni-  
 155 toring for potential complications and ongoing neurore-  
 156 habilitation remain in place.



**Figure 1.** MRI of the brain showing multiple T2/FLAIR hyperintensities with restricted diffusion involving the right cerebellar hemisphere, medulla, and pons, consistent with acute multiple ischemic infarcts consistent with a small vessel/lacunar distribution. The absence of watershed (border-zone) infarcts effectively ruled out systemic hypotension or low-flow states as the primary cause of the stroke.

## 158 Discussion

159 Nervous system involvement in malaria primarily presents  
160 as cerebral malaria or other neurological symptoms [5].  
161 Stroke, whether hemorrhagic or ischemic, is an uncom-  
162 mon but recognized complication [2,3]. Malaria-induced  
163 stroke in young patients is a rare but serious complication  
164 that results from multifactorial pathophysiological mech-  
165 anisms [6]. The primary mechanism involves mechanical  
166 obstruction of cerebral vessels by parasitized red blood  
167 cells, particularly in infections caused by *Plasmodium*  
168 *falciparum*, leading to local ischemia and infarction [7].  
169 In addition to this mechanical blockage, inflammatory  
170 responses induced by the parasite contribute to endothe-  
171 lial damage and a systemic hypercoagulable state, fur-  
172 ther increasing the risk of cerebral thrombosis and stroke  
173 [4,5]. These processes impair cerebral blood flow and  
174 trigger neurological deficits, as seen in ischemic stroke  
175 presentations. The clinical recognition of malaria-associ-  
176 ated stroke is challenging due to symptom overlap with  
177 cerebral malaria and other neurological complications of  
178 malaria, underscoring the need for heightened diagnostic  
179 vigilance in endemic areas. When evaluating a patient  
180 with stroke, infectious and tropical diseases should be  
181 considered in the differential diagnosis, and neuroimag-  
182 ing plays a critical role in making an accurate diagnosis  
183 [8-10]. Management includes prompt initiation of effec-  
184 tive antimalarial therapy, supportive care, and antithrom-  
185 botic treatment, while carefully balancing risks such as  
186 thrombocytopenia and hemorrhage [9,10]. Early diagno-  
187 sis and comprehensive treatment are critical to improving  
188 outcomes and reducing the risk of long-term neurological  
189 impairment in young patients suffering from malaria-in-  
190 duced stroke.

191 The patient's GCS of 7, right-sided flaccid hemipares-  
192 is with diminished reflexes, positive Babinski sign, and  
193 sensory deficit contribute to an approximate score of 15  
194 points, reflecting severity. This classification helps guide  
195 clinical treatment decisions and prognosis. A more com-  
196 prehensive assessment of visual fields, language, dysar-  
197 thria, and neglect could refine this score further.

## 198 Conclusion

199 Malaria-induced stroke, although rare, should be sus-  
200 pected in patients from endemic regions, even in the  
201 absence of traditional stroke risk factors. The pathogen-  
202 esis is multifactorial, primarily involving mechanical  
203 obstruction of cerebral vessels by parasitized red blood  
204 cells and an accompanying inflammatory and hyperco-  
205agulable state. Early diagnosis and prompt antimalarial  
206 treatment, along with supportive care, are essential to  
207 improving outcomes and minimizing long-term neuro-  
208 logical deficits. This case provides significant insight  
209 by demonstrating that malaria-associated stroke can  
210 occur in adults with high parasitemia, manifesting as a

multiple-territory ischemic stroke that may not be ini- 211  
tially detectable on CT scan. 212

### What is new?

213  
214 This case report highlights a young, otherwise healthy  
215 32-year-old male developing an ischemic stroke shortly  
216 after severe malaria infection. It emphasizes the associa-  
217 tion between malaria and stroke in adult patients without  
218 traditional risk factors, expanding awareness of this poten-  
219 tial complication. The report underscores the importance  
220 of considering malaria as a differential diagnosis in young  
221 stroke patients in endemic areas and discusses the manage-  
222 ment implications, encouraging early diagnosis to improve  
223 outcomes.

### List of abbreviations

C-ANCA	Cytoplasmic Anti-Neutrophil Cytoplasmic	225
	Antibodies	226
CT	Computed Tomography	227
DVT	Deep Venous Thrombosis	228
ECG	Electrocardiogram	229
ESR	Erythrocyte Sedimentation Rate	230
FBC	Full Blood Count	231
ICU	Intensive Care Unit	232
IV	Intravenous	233
MRI	Magnetic Resonance Imaging	234
P-ANCA	Perinuclear Anti-Neutrophil Cytoplasmic	235
	Antibodies,	236
PCR	Polymerase Chain Reaction	237

### Conflict of Interests

238 The authors declare that there is no conflict of interest regard-  
239 ing the publication of this article. 240

### Funding

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### Consent for publication

243 Written informed consent was obtained from the next kin of  
244 the patient (since the patient is unable to provide consent for  
245 publication) 246

### Ethical Approval

247 Ethical approval is not required at our institution to publish an  
248 anonymous case report. 249

### Take-home message

250 In malaria-endemic regions, clinicians should maintain a high  
251 index of suspicion for stroke in young patients presenting with  
252 acute neurological symptoms following malaria infection, as  
253 early recognition and prompt treatment are critical to improv-  
254 ing outcomes and preventing long-term neurological disability. 255

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308 **Summary of case**

1	Patient (gender, age)	32 years, Male
2	Final diagnosis	Cerebral malaria with multifocal ischemic stroke
3	Symptoms	dizziness, vomiting, headache, slurred speech, drooling, right-sided weakness and numbness
4	Clinical investigations	Laboratory tests, immunological assays and radiographic tests
5	Medications	Artesunate, aspirin, vitamin B6, vitamin B 12, enoxaparin, levetiracetam
6	Clinical procedure	None
7	Specialty	Neurology