An Important Pediatric Stroke Mimic: Hemiplegic Migraine

Michael T. Jurkiewicz, PhD, MD\textsuperscript{1}, Arastoo Vossough, PhD, MD\textsuperscript{2}, Avrum N. Pollock, MD\textsuperscript{2}

\textsuperscript{1} Department of Medical Imaging, Children’s Hospital at London Health Sciences Centre, London, ON

\textsuperscript{2} Department of Radiology, Children’s Hospital of Philadelphia, Philadelphia, PA.

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Correspondence to:
Michael Jurkiewicz PhD MD
Department of Medical Imaging
Children’s Hospital at London Health Sciences Centre
London, ON, N6A 5W9
Phone: 519 685 8500 x56184
Email: michael.jurkiewicz@lhsc.on.ca
Stroke is relatively rare in children but has become increasingly recognized clinically. Hemiplegic migraine (HM) is a rare subtype of migraine, with attacks typically beginning in childhood or adolescence. Attacks are characterized by migraine headaches and motor weakness, which develop over several minutes. Hemiplegic migraine may therefore mimic acute stroke—however symptoms last less than an hour and resolve spontaneously, often without sequela\textsuperscript{1-4}. Distinction between these entities is important due to their different urgency and management. Neuroimaging is indispensable in working up patients presenting to the Emergency Department with stroke-like symptoms and can be used to distinguish between infarction and HM.

A 12-year-old boy presented with acute onset severe, left-sided headache with photophobia. After approximately two hours of constant pain, the patient’s mother noticed a right facial droop, right facial paralysis, and right arm weakness. Two hours later, physical examination in the Emergency Department identified mild right-sided facial droop, but no other weakness; a severe headache persisted. Two hours afterwards, the patient underwent an MRI to exclude infarction.

Brain MRI was unremarkable on diffusion-weighted (Figure 1A), T1-weighted, T2-weighted, and post-contrast T1-weighted images. FLAIR images revealed subtle hyperintensity involving the cortex of the left hemisphere (Figure 1B). Susceptibility weighted images demonstrated asymmetrically prominent veins throughout the left hemisphere (Figure 1C). Arterial spin
labeling perfusion images revealed hypoperfusion throughout the left hemisphere (Figure 1D). These findings were compatible with HM.

Hemiplegic migraine has two forms, familial and sporadic, which occur with equal frequency and a prevalence of approximately 1/10,000\(^5\). With no associated family history, this is a case of sporadic HM. Three known genes are associated with familial HM: CACNA1A, ATP1A2, and SCNA1, with the proline-rich transmembrane protein PRRT2 gene also implicated\(^2,6\). The pathogenesis of HM remains unclear, with conflicting theories between vascular or neuronal dysfunction suggested. The neurological deficit concomitant with both decreased and increased cerebral blood flow argues against a purely vascular mechanism.

Hemiplegic migraine may mimic an acute stroke, making clinical distinction difficult. Rapid diagnosis is vital to the urgent management of stroke patients, and neuroimaging, preferably with MRI, is essential in the work up. Brain MRI shortly after symptom onset in patients with HM demonstrates cerebral hypoperfusion\(^7,8\), with increased venous deoxyhemoglobin identified on susceptibility weighted imaging by prominent cerebral veins within the cerebral sulci on the affected side\(^4\). Although not fully understood, alterations in the blood brain barrier within the affected region is believed to result in cortical swelling and vasogenic edema, both of which contribute to FLAIR hyperintensity\(^7\). As there is no cytotoxic edema, the diffusion-weighted images remain negative, thus excluding infarction. A shift to regional hyperemia and hyperperfusion typically occurs later\(^8,9\), with subsequent near normalization of cerebral blood flow and decreased prominence of cerebral veins, which correlates with symptom resolution\(^4\).
In one recent article, it was shown that patients scanned less than 14 hours after aura onset consistently demonstrated decreased cerebral blood flow in symptom-related brain regions, while a constant increase in cerebral blood flow was seen in these regions 17 or more hours post onset. 

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References

Figure 1A. Axial diffusion weighted images show no areas of diffusion restriction, excluding stroke as a cause for the patient's symptoms.

Figure 1B. Axial FLAIR images show subtle but diffuse hyperintensity involving the cortex of the left frontal, temporal, and parietal lobes (yellow arrows) compared to the right side.

Figure 1C. Axial SWI minimum intensity projection images show increased curvilinear susceptibility effect in the sulci throughout the left hemisphere, in keeping with venous prominence.

Figure 1D. Axial arterial spin labeling relative cerebral blood flow images show decreased blood flow to the left hemisphere.