A 38-year-old woman presented with a right-sided throbbing headache associated with visual symptoms, nausea, vomiting, photophobia, and phonophobia. She had experienced migraines since her teenage years characterized by monthly unilateral headache associated with nausea and photophobia. The current headache had started with symptoms she recognized as her regular monthly migraine, but it had uncharacteristically persisted for 3 days. She also described a persistent aura of “pixelated 3-dimensional objects” and “colored pinwheels,” which she had not experienced with her previous headaches. Two days into the headache, she had developed a defect in her left visual field and had narrowly missed hitting a tree on that side while driving. At the time of presentation to us, she had already undergone a head CT, which was reported normal, and she was prescribed prednisone, sumatriptan, and oxycodone for management of status migraine, without benefit.

Our initial evaluation found an obese woman wearing dark glasses due to photophobia. The neurologic examination revealed visual acuity of 20/25 bilaterally, no papilledema, left homonymous hemianopsia, and a horizontal nystagmus with a fast component to the left. Laboratory workup was significant for anemia, ketonuria secondary to vomiting and dehydration, and positive antinuclear antibody (1:80) with speckled pattern. The remainder of her hematologic workup was normal.

A brain MRI showed thickened cortex in the right occipital lobe on T2 fluid-attenuated inversion recovery (FLAIR) (figure, A) as well as hyperintense signal on diffusion-weighted imaging (DWI) (figure, B) with subtle corresponding apparent diffusion coefficient hypointensity, isolated to the same cortex. Isolated cortical involvement with sparing of the white matter suggested that the findings were unlikely to be due to arterial ischemic infarction. Her visual symptoms worsened, and she variously described them as a “kaleidoscope-like effect,” with “yellow visual fields” and “right side of my brother’s body is melting away.” Her persistent complex visual symptoms with colored 3-dimensional figures and distortions (metamorphopsia) were more typical for visual hallucinations of occipital epilepsy rather than migrainous aura. EEG was obtained and she was found to be in partial status epilepticus with an ictal focus in the right occipital and parietal cortices (figure, C). She was treated with lorazepam and loaded with levetiracetam with significant improvement in the headache over the next 24 hours. Topiramate was added to the regimen as a dual agent for seizure and migraine prophylaxis. She was discharged home headache-free but with a mild residual hemianopsia. This fully resolved at 3-month follow-up, and a repeat MRI showed complete resolution of the previously noted findings. Levetiracetam was withdrawn at follow-up and she was maintained on topiramate monotherapy.

The final diagnosis was partial status epilepticus from the right occipitoparietal cortex with ictal hemianopsia and hemicrania epileptica. It is debatable whether the seizure was precipitated by migraine, constituting migralepsy, or her migraines were always an epileptic phenomenon. Topiramate was chosen for prophylaxis because it is a dual agent against both epilepsy and migraine.

PEARS

Hemicrania epileptica (HE) is recognized as an ipsilateral headache phenomenon (in the International Classification of Headache Disorders [ICHD] classification) with migrainous features but occurring as an ictal manifestation of focal epileptic seizures.1

Diagnostic criteria2:
A. Any headache, fulfilling criterion C.
B. Patient experiencing a partial epileptic seizure.
C. Evidence of causation demonstrated by both of the following:
1. Headache has developed simultaneously with onset of a partial seizure.
2. Either or both of the following:
   a. Headache has significantly improved immediately after the partial seizure has terminated.
   b. Headache is ipsilateral to ictal discharge.
D. Not better accounted for by another ICHD-3 diagnosis.

Parisi et al. identified 5 potential HE cases in the literature from 1988. In these cases, migraine...
appeared to be the sole manifestation of nonconvulsive status epilepticus; 4 patients showed partial status in occipital lobes in the EEG.1 None of these patients met the ICHD-2 diagnostic criteria for HE. Seizure-induced neuroimaging changes were not described in these cases. Our patient had a persistent headache with migrainous features ipsilateral to the side of ictal activity that improved after appropriate antiepileptics were started and was not better accounted by any other ICHD-3 diagnosis; therefore, this classifies as hemicrania epileptica. Our patient’s headache was prolonged as she was in focal status and she also had neuroimaging changes secondary to focal status epilepticus.

OY-STERs The typical migrainous visual aura starts in the central visual field described as flickering, uncolored, zigzag lines, then moving toward the periphery leaving a scotoma. The aura usually lasts more than 4 minutes and less than 30 minutes. Visual hallucinations secondary to occipital seizures are more complex, mainly colored and circular in nature, occurring in the periphery of the temporal contralateral visual field, becoming larger and moving horizontally toward the ipsilateral visual field. Complex visual hallucinations with distorted imagery can occur when the visual association areas are involved. These ictal visual symptoms are shorter in duration, lasting 30 seconds to 3 minutes.3–5

EEG should be considered in all patients with headache who exhibit both positive and negative visual symptoms and the positive visual symptoms have the complex characteristics described above for ictal visual symptoms. Evidence of cortical edema or thickening on FLAIR and restricted diffusion on DWI limited to the cortex with sparing of white matter may be attributed to ongoing focal seizures and is one of the described transient peri-ictal MRI abnormalities.6

AUTHOR CONTRIBUTIONS
Dr. Sheikh designed, drafted, and revised the manuscript and created the figure. Dr. Georgsson revised and edited the manuscript. Dr. Marks edited the manuscript for intellectual content and approved all final changes.

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REFERENCES

Figure Right occipital seizure on EEG and resultant changes on MRI

T2 fluid-attenuated inversion recovery showing thickening of cortex in right occipital lobe (A), restricted diffusion in right occipital cortex (B), and EEG showing seizure from right occipital and parietal cortices (C).


Pearls & Oy-sters: Hemicrania epileptica: Unfolding the mystery of an unremitting migraine
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