Migraine madness: recurrent psychosis after migraine

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Abstract
A 69 year old man with longstanding migraine with aura had four episodes of psychosis lasting 7–28 days during a 17 year period. During attacks he had formed visual hallucination and delusions, including reduplicative paramnesia. His mother was similarly affected. His EEG showed symmetrical frontal delta waves. The time course and EEG changes are similar to acute confusional migraine. The reduplicative paramnesia suggests a focal non-dominant hemisphere dysfunction.

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A wide variety of auras have been described in classical migraine (migraine with aura).1 Visual phenomena both positive and negative are frequent, usually lasting less than one hour and occurring either before, coincident with or at different times from the headache.2,3 Much less common are aura that involve higher mental function4 such as confusional states, particularly seen in children,5 temporal lobe phenomena,6 and dysphasia.7

Dysphrenic migraine is a term used to refer to mental changes or "psychic alterations" associated with migraine.8 The term has been used to encompass a wide range of phenomena including twilight states, hallucinations, somnambulism, suicidal attempts, ideas of reference, religious and persecutory delusions, confusional states and stupor.9 Psychosis is rarely reported6 7 and the form of the psychosis is rarely described in detail. We describe a man with recurrent episodes of psychosis with prominent reduplicative paramnesia (a delusion that familiar people, places or objects have been replaced by a double) associated with migraine.

Case report
A 69 year old man with longstanding migraine with aura presented with persecutory beliefs and formed visual hallucinations. His wife reported that 3 days before he had been normal. He had then gone to bed complaining of severe bifrontal headache associated with vomiting. The next day the headache continued. On waking on the third day the headache had resolved and he complained that his wife and brother-in-law were changing before his eyes, that their arms lengthened and they each had only one eye. He said familiar things looked strange, for example he thought the original house had been replaced by a cleverly produced copy. The next day he started seeing threatening words written on the walls and had refused to enter certain rooms because of evil in them. He had at times seemed confused and had fallen 5–6 times. He had no abnormal movements.

He had suffered from migraine with aura since the age of 16. The aura usually began with tingling in both lips and dysarthria followed by tingling of his left arm, followed by flashing lights in both eyes and after about 30 minutes a right sided throbbing headache. These attacks had occurred at 3–6 monthly intervals. At the age of 53 he had a migrainous headache followed within 24 hours by an episode of paranoid psychosis that lasted 3 weeks with full recovery. Two years later, when well, he had an EEG which showed a low voltage post-central activity at 6–8 Hz with a moderate amount of slower waves at 4–6 Hz bilaterally. Some episodic anterior delta waves were seen on overbreathing.

At 64 he was admitted to a psychiatric hospital with a similar episode. This began with a migrainous headache and was followed, within 24 hours, by formed persecutory visual and auditory hallucinations. He called the police twice in one evening when he saw 100 burglars in the front garden. He was initially disorientated in time. He returned to normal after 7 days. An EEG performed 1 month after the attack was similar to that performed when he was 55 except that there were more frequent episodes of bisynchronous 2–3 Hz delta activity at 20–100uV. The delta waves were not seen on an EEG 3 months later. At the age of 65 he had a further episode of paranoid psychosis within 24 hours of a migrainous headache that lasted 14 days.

He had been treated for hypothyroidism since the age of 65. His mother had migraine with aura and recurrent episodes of migraine followed by paranoid hallucinatory episodes. He had no history of epilepsy and drank alcohol infrequently.

On examination he was agitated and with a perplexed facial expression. His mood was suspicious but otherwise euthymic. His speech was disjointed with paraphasia and neologisms (for example, I have a telephonic expression on the brain). He experienced well formed colourful mobile visual hallucinations, worse in dim illumination, with perceptual distortions and illusions. These included seeing his hospital bed being swallowed up by the floor, micropsia and macropsia, for
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temporal relationship. The episodes of psychosis have only occurred following migraine and were transient. Between attacks there is full recovery with insight. In addition there have been no features suggesting an alternative aetiology for the headaches or the psychoses, notably he had no history of epilepsy or alcohol abuse.

The description of a similar pattern of migraine with aura in the patient's mother, with the development of migraine followed by psychosis in the later part of her life suggests a link between the psychosis and migraine. It also suggests that it may be familial.

The types of the hallucinations described by our patient included some visual phenomena not infrequently seen in migraine aura, such as micropsia and macropsia as well as formed visual hallucinations which are rare.

The time course of the visual hallucinations and delusions in our patient are unusual in following the headache and lasting for several days.

Previously reported migraine psychoses have had a similar time course. Klee reported four patients with psychoses, with visual and auditory hallucinations and in one a distortion of body perceptions (she believed her legs to have been cut off) that lasted a few days. The episodes of psychoses followed the migraine attack and lasted several days. Eight out of nine members of a family with hemiplegic migraine described by Feeley et al had either a psychotic episode or confusional states, both of which lasted several days and followed migraine attacks.

A similar time relationship with the migraine attack is seen in acute confusional migraine when the confusion follows the migraine and may last for up to 5 days. The EEG abnormalities of intermittent frontal delta waves seen in this patient during and shortly after attacks resolved between attacks. Similar EEG findings have been reported in young patients with juvenile acute confusional migraine.

One other phenomenon of particular interest is that of reduplicative paramnesia and delusional misidentification. Reduplicative paramnesia is a term coined by Pick to describe the delusion that a familiar person, object, place or even the patient's self was replaced by a double. The most well known variants of reduplicative paramnesia are delusional misidentification syndromes that involve the recognition of people. In Capgras' syndrome patients believe a familiar person is replaced by an identical double. In the syndrome of intertematymorphosis the patient believes someones appearance has radically changed to correspond with the appearance of someone else. In another syndrome of delusional misidentification, the Frégoli syndrome, the patient misidentifies strangers as being a familiar person in disguise. Reduplicative paramnesia of inanimate objects is less often recognised. Our patient believed that his house had been replaced, thought his surroundings had been transported to an exactly similar site at a different place.
location and failed to believe that his wife was his wife. The first two are delusions of replacement of objects and surroundings and it is not clear whether the last represents prosopagnosia or delusional misidentification.

Capgras' syndrome has been described previously in two patients with migraine. Both patients had common migraine and developed a single episode of delusional misidentification following a migraine attack or an episode of vomiting associated with a right hemiparesis attributed to vertebrobasilar migraine, though apparently without headache. We are not aware of reports of other delusional misidentification syndromes, such as the Frégoli syndrome or interm termedorphosis, in migraine.

It has been proposed that reduplicative paramnesia arises from a disconnection between the non-dominant visual association cortex and the limbic structures associated with memory. This provides a potential discrete site for the disturbance of function that occurred in our patient and is in keeping with the mild neuropsychological deficit found. The pathophysiology of the disturbance is unknown but has many features in common with other delayed disturbances of higher function such as acute confusional migraine and hemiplegic migraine.

6 Klee A. A clinical study of migraine with particular reference to the most severe cases. Copenhagen: Munksgaard, 1968.
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