Headache in people with epilepsy

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Abstract

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19 Epidemiological estimations indicate that individuals with epilepsy are more likely to experience headaches,

including migraine than individuals without epilepsy. Headaches can be temporally unrelated to seizures, or can

occur before, during or after an episode; seizures and migraine attacks are mostly not temporally linked. The

pathophysiological links between headaches (including migraine) and epilepsy are complex and have not yet been

fully elucidated. Correct diagnoses and appropriate treatment of headaches in individuals with epilepsy is

essential, as headaches can contribute substantially to disease burden. Here, we review the insights that have been

made into the associations between headache and epilepsy over the last 5 years, including information on the

pathophysiological mechanisms and genetic variants that link the two disorders. We also discuss the current best practice for the management of headaches co-occurring with epilepsy and highlight future challenges for this area of research.

[H1] Introduction

The hallmark of epilepsy is an enduring predisposition to seizures accompanied by neurobiological, cognitive and psychological comorbidities¹. Epileptic seizures are defined as the disruption of normal neuronal functioning owing to excessive or synchronous neuronal activity, leading to an epileptic event that is discernible by the person and/or by an observer¹. An analysis for the Global Burden of Disease Study 2016 estimated that >50 million people worldwide had active epilepsy, that is, they had continuing seizures or were receiving epilepsy treatment². The origin and cause of seizures can vary. The International League Against Epilepsy (ILAE) scheme³ classifies seizures as either "focal", meaning that seizures originate at a specific location in one hemisphere; "generalised", denoting seizures that engage bilaterally distributed networks; or "unknown", for seizures with an undefined origin. The ILAE classifies epilepsy as either "focal", "generalised", "focal and generalised", or "unknown", depending on the type of seizures that occur³. The same scheme also classifies epilepsy according to aetiology, including "structural" (for example, associated with a brain tumour or gliosis), "genetic", "metabolic" (for example, associated with mitochondrial disease), "infectious", "immune" or "unknown"³. The category "unknown" includes genetic, metabolic and structural causes that have not yet been identified.

Headaches are among the commonest disorders globally — the Global burden of Disease Study 2017 estimated that there were > 3 billion individuals with headache across 195 countries and territories⁴. The International Classification of Headache Disorders 3 (ICHD-3)⁵ distinguishes between primary headaches — including migraine, tension-type headache (TTH) and trigeminal autonomic cephalalgias — and secondary headaches, which are attributable to other disorders or substances. TTH, which affects >2 billion people globally⁴, is a poorly defined featureless headache that lacks the characteristic features of other primary headaches and is usually bilateral and pressing (non-pulsating)⁵. TTH can last for 30 minutes to seven days, is not usually aggravated by routine physical activity and is not accompanied by nausea, vomiting or photo-phobia or phonophobia⁵.

Global migraine prevalence is ~1.3 billion and the disorder is 3–4 times more common in women than men⁴. Migraine is a heterogeneous brain disorder, typically characterised by recurrent attacks of mostly severe unilateral pulsating headache lasting 4–72 hours, accompanied by nausea, vomiting and/or hypersensitivity to sensory stimuli, and a range of other sensory and cognitive symptoms⁵. In about 30% of individuals with migraine, the pain is preceded — and in rare cases accompanied or followed by — a migraine aura, consisting of transient focal neurological symptoms. Symptoms of migraine aura are usually visual but may involve tactile, motor and/or speech disturbances⁶. Some individuals have auras without headache⁷.

Here, we review the link between epilepsy and headaches, starting with the epidemiology of the two disorders. We then discuss the diagnosis and classification of headaches in epilepsy and provide an overview of the current understanding of the underlying pathophysiological mechanisms. Last, we discuss the clinical management of co-existing headaches and epilepsy. We focus on evidence published between 2015 and 2020 to provide a view of recent progress in the field, and we also provide a timeline of key publications from before 2015 (Fig. 1.).

[H1] Epidemiological evidence

Headaches, especially migraine, and epilepsy frequently co-exist in the same individuals. A meta-analysis of population-based studies of migraine in people with epilepsy published between 1996 and 2012 indicated that lifetime migraine prevalence was 52% greater in people with epilepsy than in people without epilepsy. The lifetime epilepsy prevalence was also 79% greater in people with migraine than in people without migraine. A more recent meta-analysis (including studies published between 2004 and 2019) estimated a 49% prevalence of unspecified headache among people with epilepsy. Additional evidence has confirmed the findings of these meta-analyses regarding the co-existence of epilepsy and headache (Table 1)^{10–19}. In these studies, \leq 79% of individuals with epilepsy reported experiencing headaches. The most common headache types in individuals with epilepsy were migraine (reported by \leq 25% of participants) and TTH (reported by \leq 40% of participants)^{10,13,14,16,18}. Women with epilepsy tended to report migraine more often than men with epilepsy^{11,12,16,18,20}. No clear relationship between headache type and epileptic focus location, seizure type, seizure frequency, or use of antiseizure medication was identified in these recent studies^{13,16}. One older study reported that peri-ictal headaches were ipsilateral to the epileptic focus in temporal epilepsy, but not in extra-temporal epilepsy²². Some researchers

have suggested that the association between headache and epilepsy is stronger in individuals with genetic forms of epilepsy than those with non-genetic forms, and stronger in children than in adults²³, One study reported a negative correlation between headache frequency and age of epilepsy onset¹¹ comparative meta-analytic evidence to support this finding is lacking.

[H2] Limitations of epidemiological studies

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Epidemiological studies have offered important insights into the relationship between epilepsy and headache but can be subject to biases, which might influence findings. First, the case-ascertainment method used often influences study findings, for example, studies that use self-report questionnaires tend to show a stronger association between headache and epilepsy than those that rely upon a physician's assessment⁸. This disparity might be caused by the fact that few validated instruments exist for self-diagnosis of epilepsy or headaches²⁴ studies often use their own, unvalidated instruments⁸, the accuracy of which is unknown. How questions are formulated can influence the responses; for example, the results of one study suggested that people with epilepsy were three times more likely to report headaches preceding seizures when asked closed-ended questions than when asked open-ended questions²⁵. Second is the effect of recall bias on findings²⁶. Evidence indicates that, compared with healthy individuals, individuals with a pre-existing condition are more likely to report additional symptoms²⁶. This observation might explain why individuals with epilepsy report migraine more often than individuals without epilepsy⁸. Conversely, seizures can be associated with amnesia, which would make it difficult for the individual to recall what happened just prior, during or after the seizure, thus preventing the reporting of comorbidities such as headache²⁷. Additionally, seizures are often conspicuous events and could overshadow less apparent complaints like headache, especially in children. Consequently, individuals with epilepsy might perceive headaches as "mundane" and thus not report them unless directly asked. Third, physicians might not be aware that headaches are common in individuals with epilepsy^{27–29}, which could introduce misclassification bias²⁶. This type of bias could occur when the health provider is more or less attentive to comorbidities contingent on whether the individual has a debilitating condition. A serious ailment might prompt physicians to look for other associated conditions. However, an individual might be so ill that "milder"

symptoms or diseases are overlooked or seen as part of the significant condition. We hypothesize that this bias

could explain why studies based on physician assessment show a lower association between epilepsy and headaches than studies based on self-assessment⁸.

Last, although studies that use insurance data or International Classification of Diseases codes have the advantage of physician-diagnosed data from large cohorts of individuals, the use of codes and insurance labels can be influenced by local policies. The choice of codes used might be influenced by financial or insurance-related factors, also resulting in biases. Despite these various sources of bias, epidemiological studies are essential in ascertaining the overlap between different conditions. Designing studies that are totally free of bias is impossible but bias can be reduced during the data collection phase and taken into account when interpreting results.

[H2] A bidirectional relationship

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Whether epilepsy and headaches have a "bidirectional" association — meaning that the occurrence of one influences the onset of the other and vice versa — remains unknown. To date, most studies of the association between epilepsy and headaches have been cross-sectional, so do not allow for such assessments. To assert that a relationship between two conditions is bidirectional, a precise determination of condition B's onset in relation to condition A is required, and thus costly and labour-intensive longitudinal studies are needed. One such study evaluated the risk of developing subsequent epilepsy when first diagnosed with migraine and found that individuals with migraine and those who had migraine and sleep disorders, cognitive disorders, anxiety or depression were more likely to develop epilepsy than healthy individuals³⁰. This cohort was followed-up for a mean period of 12 years, and the relative risk of developing epilepsy was found to be 2.3 times higher in men than in women³¹. Risk was increased by older age, low-income status and comorbidities, especially head trauma. For example, the risk of developing epilepsy was 4.6 times higher in men with migraine and a history of head trauma than in men with migraine and no history of head trauma³¹. These studies are longitudinal, but only assessed the risk of developing epilepsy in people with migraine and do not provide information on whether or not the relationship is truly bidirectional. Multi-centre prospective, long-term studies with clear diagnostic criteria will be vital to shed light on the complex relationship between epilepsy and headache and help identify individuals at risk of developing severe or chronic forms of either condition.

[H1] Diagnosis and classification

Headaches that co-occur with epilepsy can be classified according to their temporal relationship to seizures (Fig.

- 2). Interictal headaches occur > 24 hours before and > 72 hours after epileptic seizures. Peri-ictal headaches, including migraine, occur shortly before, during or just after an epileptic seizure and can present a diagnostic challenge. The distinction between epilepsy and peri-ictal headaches is often apparent, the conditions can sometimes overlap either temporally or in terms of symptoms. These temporally classified types of headache (pre-ictal, post-ictal, ictal and interictal headache) can occur in the same individual (table 1).
- Accurate classification of epilepsy and headache is important for initiating adequate, timely and appropriate treatment and requires a good description of the symptoms and their temporal relationships. The ILAE seizure classification scheme does not include a class of seizures with symptoms that overlap with headaches. However, the ICHD-3 includes several categories of seizure-related headaches⁵ (Box 1): migraine aura-triggered seizure (section 1.4.4), ictal epileptic headache (section 7.6.1) and post-ictal headache (section 7.6.2).

[H2] Pre-ictal headaches

Headaches that occur < 24 hours before a seizure and last until seizure onset have been defined as pre-ictal¹¹. According to the ICHD-3⁵, the existence of pre-ictal headaches is controversial⁵, even though they have been reported in several studies³²⁻³⁵. The issue is that an EEG recording of the headache event is mandatory for the diagnosis of pre-ictal headache — for a headache to be pre-ictal, it must not be accompanied by ictal epileptic discharges on the EEG — and the studies cited above did not include an EEG recording of the event³²⁻³⁵. Headache concomitant with ictal epileptic discharges should be classified as ictal epileptic headache (see below). A classification of pre-ictal headache is not given in the ICHD-3⁵, but the comments section calls for more studies to establish the existence, prevalence and features of this type of headache. The results of cohort studies suggest that possible pre-ictal headaches (without EEG confirmation) occur in 1–10% of people with epilepsy^{10,12–15,19,21} — (Table 1) the headache is migraine-like in 30–60% of these individuals and tension-type in ~20% ^{10–15,17,19,21}. In a video-EEG study, 25 of 831 (6.3%) individuals with epilepsy reported pre-ictal headache without epileptic discharges on the EEG¹⁷. Five had "headache as a seizure aura", which should be classified as "ictal epileptic headache", see below¹⁷.

[H2] Migraine-aura triggered seizures

The term aura is used to describe subjective precursory symptoms of seizures and migraine headaches; however, it refers to different phenomena in the context of migraine or epilepsy. The ICHD-3⁵ defines aura as "recurrent attacks, lasting (5–60) minutes, of unilateral fully reversible visual, sensory, motor or other central nervous system symptoms that usually develop gradually and are usually followed by headache and associated migraine symptoms." (Box 2). In contrast, a report by the ILAE Task Force on Classification and Terminology describes aura as "A subjective ictal phenomenon that, in a given individual, may precede an observable seizure; if alone, constitutes a sensory seizure." An epileptic aura is confirmed by epileptic discharges on EEG and is part of the seizure³⁶. Some epileptic auras do not have a visible EEG correlate as they can be very focal, occupying such a small cortical area that the spatial resolution of surface EEG is insufficient to detect them³⁷.

In migraine, no consistent EEG abnormalities are observed during the aura and headache phase^{38,39}. Studies have found either slow waves, attenuation of background activity amplitude or the presence of normal EEG patterns during migraine aura^{38,40}. During attacks of hemiplegic migraine and migraine with disturbed consciousness, abnormal EEG patterns with unilateral or bilateral delta activity have been recorded⁴⁰. The EEG has no diagnostic

with epilepsy and comorbid headache⁴¹.

In rare cases, a migraine-like aura can occur immediately before a seizure⁵. The ICHD-3 refers to seizures that occur during or < 1 hour after the end of a migraine with aura attack as "A seizure triggered by an attack of migraine with aura". These seizures are sometimes referred to as migralepsy⁵. Visual symptoms and hallucinations are hallmarks of migraine aura and occipital epilepsy, making it difficult to distinguish between the two conditions. In a meta-analysis published in 2019, the most common visual symptoms of migraine aura reported were foggy and/or blurred vision, zigzag or jagged lines, scotoma, phosphenes and flickering light⁴². (Table 2) The symptoms of occipital epilepsy are elementary and visual hallucinations or illusions; blindness; palinopsia and sensory hallucinations of ocular movements; ocular pain and oculomotor symptoms, including deviation of the eyes; and nystagmus and repetitive eyelid closure or fluttering⁴³. The duration of symptoms is the most helpful feature for differentiating between migraine-related aura and occipital epilepsy⁴⁴: the median duration of migraine aura is ~25 minutes, whereas epileptic visual hallucinations last < 1 minute⁴⁵. The hallmark of migraine aura is a slowly progressive centrifugal or centripetal scotoma that expands over 10–60 minutes^{5,42};

value in migraine (or headaches)³⁸, but is mandatory for diagnosis of epilepsy, which also applies to individuals

a feature not described by people with occipital epilepsy^{43,45}. In migraine, visual symptoms are almost always lateralised⁵. Similarly, event-associated nausea, vomiting, photophobia and phonophobia occur more often in migraine with aura than in occipital epilepsy⁴⁵. Clinically, the simultaneous occurrence of positive and negative phenomena is more suggestive of a migraine aura than of epilepsy^{5,43,45}.

The overlapping features of migraine aura and occipital seizures means that diagnosis requires a detailed description of the subjective symptoms, and pre-ictal and ictal EEG recordings. The absence of epileptiform abnormalities when the symptoms are present is the gold standard for ruling out an epileptic origin. The lack of epileptic EEG abnormalities during the migraine aura phase is essential for diagnosing migraine aura-triggered seizure. Experts doubt the existence of migraine aura-triggered seizures 46-48 as pre-ictal and ictal EEG recordings often confirm an epileptic rather than a migraineous origin of the symptoms. For example, in one EEG study, 16 out of a cohort of 4,600 children diagnosed with epilepsy had an epileptic seizure < 1 hour after a presumed migraine attack. These children had focal or generalized ictal EEG abnormalities during the migraine phase, indicating an epileptic origin of the migraine-like symptoms⁴⁶. In a more recent study involving a large cohort of individuals with epilepsy, three participants (<1%) reported epileptic seizures within an hour of an attack of migraine with aura. Two of these individuals were diagnosed with occipital epilepsy — the migraine-like aura was interpreted as an occipital seizure — and the third was diagnosed with epilepsy secondary to systematic lupus erythematosus⁴⁹. In a case report, two individuals presented with visual auras lasting 13–17 minutes, followed by a forceful turning of the head and, in one individual, a generalised tonic-clonic seizure⁴⁸. EEG recordings showed a left occipital seizure in the first individual and a right parietal-occipital seizure in the other individual. We observed a similar presentation in one of our patients, who presented with headache accompanied by epileptic discharges on the EEG (Supplementary video 1). These individuals, in whom epileptic discharges accompany the visual symptoms and headaches on the EEG, should receive a diagnosis of ictal epileptic headache (see below), not migraine aura-triggered seizures, highlighting the challenges involved in diagnosing these conditions.

[H2] Ictal epileptic headache

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A headache accompanied by epileptic abnormalities on the EEG is classified as an "ictal epileptic headache" by the ICHD-3⁵. The headache should develop simultaneously with the seizure, and either be ipsilateral to the ictal discharge and/or show a substantial reduction in severity immediately after the seizure has terminated. Ictal

epileptic headache can be accompanied or followed by other epileptic manifestations, such as motor, sensory or autonomic signs⁵⁰. If 'pure' or 'isolated' ictal epileptic headache is the only manifestation of a seizure, it requires a differential diagnosis from other types of headache. In the ICHD-3 'hemicrania epileptica' signifies a rare variant of ictal epileptic headache, characterised by headache that is ipsilateral to ictal EEG paroxysms⁵. The precise definitions of the terms 'hemicrania epileptica' and 'ictal epileptic headache' have, however, been extensively debated^{27,29,51–53}. Indeed, the ICHD-3 begins the definition of hemicrania epileptic with "if confirmed to exist", indicating the difficulties involved in confirming this diagnosis — EEG recordings are rarely performed in individuals with isolated headache. However, a video-EEG study did identify two instances of hemicrania epileptica¹⁷ People with ictal epileptic headache can have interictal abnormalities on the EEG⁵³. The diagnosis is confirmed by the presence of epileptiform patterns on the ictal EEG; however, as these abnormalities can occur with different types of lesional and non-lesional epilepsy, there is no unique EEG pattern linked to ictal epileptic headache^{27,53}. Persistent ictal epileptic headache can occur in non-convulsive status epilepticus and in some individuals the headache only resolves after intravenous administration of anti-seizure medication²⁷. Some researchers have suggested that an ability of anti-seizure medication to resolve the headache and the epileptic discharges on the EEG should be added as a diagnostic criterion for ictal epileptic headache^{51,54}. Our view is that, owing to potential pharmacokinetic and pharmacodynamic differences between individuals, a response to treatment should not be part of a clinical definition. EEG recordings have little diagnostic value in the majority of individuals with isolated headaches, including migraines, so are rarely performed in this group of people³⁸. Therefore, ictal epileptic headache, although rare, is probably underdiagnosed. For example, one study reported that out of 831 people with epilepsy and peri-ictal headaches who underwent video-EEG monitoring, six had "headache as an aura of a seizure", along with epileptic discharges on the EEG¹⁷. Therefore, these headaches should be classified as ictal epileptic headache⁵. The headaches lasted <35s in all cases, which is also suggestive of ictal events¹⁷. A systematic review published in 2017 analysed 32 cases of reported ictal epileptic headache and found that the headache can be migraine-like or tension-type, and the location of the pain can vary⁵³. The headaches occurred in children and adults and affected

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the sexes equally. Evidence from this and other studies indicates that the epileptic focus and EEG features of ictal epileptic headaches are heterogeneous 52,53,55.

As in other focal epilepsies, in some individuals with ictal epileptic headache, epileptic abnormalities can only be detected with intracranial electrodes, suggesting a deep epileptic focus⁵⁶. Ictal epileptic headache was identified in just five people in a retrospective review of 8,800 video-EEG recordings of 4,800 individuals with epilepsy⁵⁷. Three of these five individuals had lesions in the left posterior regions, whereas the other two had generalised genetic or idiopathic epilepsy. A descriptive study of 47 people with epilepsy or unusual headache identified 22 individuals reporting headaches during seizures¹⁹. This high prevalence was attributed to the use of self-reports, and the absence of an objective tool to evaluate headache characteristics and accurately define the timing of headache onset relative to the seizure¹⁹. EEG recordings confirmed ictal headache in two individuals¹⁹. These studies and the definitions given in the ICHD-3 highlight the overlap between headaches and epilepsy. Atypical headaches — especially those with an abrupt onset and ending, or those that do not respond to analgesic treatment — should suggest to the clinician the possibility of an epileptic origin warranting an ictal EEG recording, especially if other suggestive features, such as a family history of epilepsy, are present. Paroxysmal episodes with visual signs can point to migraine with aura or epilepsy, and require detailed history taking. EEG recordings, ideally with concomitant video and encompassing the pre-ictal and ictal phase, are mandatory to support these challenging differential diagnoses and should be performed when the clinician has even the slightest suspicion that the headaches have an epileptic origin⁵⁸.

[H2] Post-ictal headaches

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Post-ictal headache is defined as a headache caused by an epileptic seizure, occurring < 3 hours after the end of the seizure event and remitting spontaneously < 72 hours after seizure termination⁵. Evidence indicates that post-ictal headache occurs in < 45% of individuals with epilepsy (Table 1), making it the most common type of peri-ictal headache^{10–17,19,21}. In \sim 50% of individuals with post-ictal headache, the headache is migraine-like (Table 1)^{10–12,14–17}. The results of a meta-analysis published in 2019 indicated that of individuals with epilepsy, one third experience post-ictal headache and 16% experience post-ictal migraine⁵⁹. Interestingly, in people with focal epilepsy, post-ictal headache is more common in those with occipital epilepsy than those with epilepsy originating

in the frontal or temporal lobes⁴⁹. Post-ictal headache is also more common after convulsive seizures than after non-convulsive seizures³⁵.

[H1] Pathophysiology of headache disorders in epilepsy

Comparing the pathophysiology of seizures and headache could help uncover the mechanisms underlying the observed associations between these two disorders. A neuronal excitation/inhibition imbalance is thought to contribute to attack susceptibility in epilepsy and migraine^{60–62}. The link between hyperexcitability, seizures and cortical spreading depolarisation — the neurobiological correlate of the migraine aura and a putative trigger of migraine attacks — provides a mechanistic framework for some, but not all, of the clinical observations of headache in epilepsy (Box 2; Fig. 3).

[H2] Mechanisms underlying seizures and headaches

Epilepsy is characterised by a temporary disruption of neurological function caused by seizures, which spread across neuronal networks within seconds and are typically associated with hypersynchronous activity on EEG recordings⁶³. This neuronal network synchronisation is thought to be caused by neuronal hyperexcitability⁶⁴, which is likely to result from multiple factors. These factors include perinatal insults, impaired mitochondrial function and mutations in genes encoding ion channels or transporters that are involved in glutamatergic or GABAergic neuronal transmission or glial buffering capacity^{65–70}.

Unlike seizures, headaches are not associated with hypersynchronous EEG activity, except in the case of headaches with an epileptic origin^{5,71,72}. Headache is thought to result from activation of the trigeminovascular system, which involves meningeal nociceptive afferents from trigeminal ganglion sensory neurons, the brainstem trigeminal cervical complex (TCC), and thalamocortical areas contributing to the sensation of pain^{73,74}. Several factors can activate the trigeminovascular system at the meningeal level. These factors include the build-up of diffusible substances such as extracellular K⁺ and H⁺ (leading to low pH), release of vasoactive mediators such as calcitonin gene-related peptide (CGRP) or substance P, as well as inflammatory mechanisms^{74–76}. The results of preclinical studies in rodents indicate that the trigeminovascular system can become activated by cortical spreading depolarisation^{77,78} and that this activation involves inflammatory cascades^{79,80}. These observations

suggest that cortical spreading depolarisation during migraine aura might initiate headache⁸¹(Fig.3; but see also Box 2).

Meningeal vasodilation has been cited as trigger for trigeminovascular system activation, in line with the ancient 'vascular theory' of migraine, but more recent evidence suggests that changes to cerebral blood flow during a migraine attack are an accompanying phenomenon induced by trigeminal nerve activation⁸². In addition to the release of vasoactive substances from trigeminal nociceptive afferents, cerebral vasodilation could also result from activation of cardiovascular nuclei in the brainstem⁷⁴. Neuroimaging studies have identified functional changes in the thalamic nuclei and brainstem, hypothalamus, frontal cortex, anterior cingulate cortex, basal ganglia, and insula during headache generation^{83,84}. Connectivity changes in some of these regions have also been observed outside of and during attacks, as have changes affecting other regions such as the pons and somatosensory cortex^{85–89}. Within this larger 'head pain matrix', hyperexcitability at any level could contribute to headache initiation^{74,76,90}.

[H2] Interictal headaches

General brain hyperexcitability in people with epilepsy⁶⁴ might, even in the absence of seizures, lower the activation thresholds of brain regions that are part of the trigeminovascular system, resulting in interictal headaches. This hyperexcitability can be a result of genetic mutations that affect neurotransmission (see section on overlapping genetics below)⁹¹. Studies in transgenic mouse models of migraine have identified an association between migraine-causing mutations and inflammatory changes^{92,93}, which might also contribute to trigeminovascular system activation. In migraine, effects of exogenous triggers such as light or stress, food or sleep deprivation, and systemic fluctuations in sex hormones are hypothesized to contribute to attack initiation via the dysregulation of cortical and (hypo)thalamic pathways^{74,76,94-101}. For example, in rats, bright-light stress causes cortical activation⁹⁶, and sleep deprivation is associated with reduced brain glycogen levels and enhanced susceptibility to cortical spreading depolarization^{97,98}. As hyperexcitability seems to contribute to the lowered threshold to headache triggers in migraine^{74,76}, this could be hypothesized to also lead to an increased propensity for interictal headaches to occur in people with epilepsy.

[H2] Pre-ictal headaches

Brain parenchymal inflammation has been shown to promote seizure initiation in rodent models^{102,103}. One mechanism underlying this inflammatory response involves the neuronal release of brain high mobility group box 1 (HMBG1) as a result of brain hyperexcitability¹⁰⁴. In migraine headaches, activation of the trigeminovascular system by cortical spreading depolarization was shown to activate inflammatory cascades, including neuronal release of HMBG1, resulting in meningeal nociceptive activation⁷⁹. It could be hypothesized that cortical network hyperexcitability, if maintained below the thresholds for eliciting epileptiform discharges and sensorimotor manifestations, could lead to trigeminovascular system activation via neuronal HMBG1 release. At the subcortical level, pre-ictal hyperexcitability can affect central autonomic circuits, including hypothalamic and brainstem areas¹⁰⁵, and projections to the limbic system¹⁰⁶. Given the involvement of these areas in the development of head pain^{73,74}, pre-ictal hyperexcitability in these regions could be hypothesized to elicit head pain before the development of widespread seizure activity.

[H2] Migraine-aura triggered seizures

The occurrence of a migraine aura before a seizure suggests an underlying cortical spreading depolarisation, followed by epileptiform activity. This sequence of events has been observed in preclinical studies, in which spreading depolarisation increased epileptic activity in rat brain slices¹⁰⁷, as well as in resected human epileptic brain tissue^{107–110}. Evidence indicates that suppression of inhibitory GABA function can contribute to this increase in epileptic activity^{107,110}. Given the scarcity of clinical evidence for migraine aura-triggered seizures, this sequence of events is likely to be rare in humans. Indeed, the results of a preclinical study found that spreading depolarisation protected rat cortical networks from expressing seizure activity¹¹¹.

[H2] Ictal epileptic headache

Multiple mechanisms could be responsible for ictal epileptic headache, including seizure-related changes in the trigeminovascular system and in pain-causing brain regions. The cortical projections responsible for head pain are likely to be widespread, involving primary sensory areas and the central autonomic network, that is, the thalamus, hypothalamus, insula, anterior cingulate cortex, medial prefrontal cortex, precuneus, amygdala, hippocampus and other parts of the limbic system^{54,72,112,113}. A study in people with epilepsy evaluated participants' responses to direct electrical stimulation of the cortex during pre-surgical evaluation and showed that pain responses were scarce (observed for 1.4% of the stimulated sites). Pain was only triggered by stimulation

of the medial parietal operculum and posterior insula¹¹⁴. This deep localisation of several pain areas might explain why, in some individuals, the electrophysiological correlate of ictal epileptic headache is only recorded using depth electrodes. However, seizures with a confirmed origin in the parietal operculum and posterior insula lead to pain sensations in the limbs contralateral to the epileptic focus and do not always lead to head or facial pain¹¹⁵. It is hypothesized that seizure activity in autonomous areas could cause direct neuronal activation of the brainstem trigeminocervical complex⁵⁴ resulting in headache^{54,112,113}, but direct evidence for this mechanism occurring in ictal epileptic headache is lacking.

A case series identified a multitude of EEG patterns in ictal epileptic headache^{52,53} suggesting that this form of headache is associated with different seizure types and localisations. As was suggested for pre-ictal headache, the mechanisms underlying ictal epileptic headache might also involve inflammatory changes caused by enhanced network excitability during seizures. However, in ictal epileptic headache, the timing of events triggering the trigeminovascular system occurs in parallel to the expression of symptomatic seizures and epileptiform EEG bursts. Increased cerebral blood flow during the pre-ictal and ictal period has also been suggested as a possible trigger of the trigeminovascular system, resulting in headache during seizures³³. However, we do not consider this to be plausible as the results of magnetic resonance angiography studies in people with migraine indicate that arterial dilatation is an effect of headache, as opposed to a cause 116,117. One such study found no evidence of arterial dilatation during migraine at all¹¹⁸. Indeed, the historical view of vasodilation as a cause of migraine headaches has now effectively been excluded 74,82. In addition to the release of vasodilating substances from trigeminal nerve endings, vasodilation might also result from increased activity of the trigeminovascular system brainstem nuclei inducing vascular changes such as enhanced cerebral blood flow⁷⁴. These observations suggest that an ictal epileptic headache is likely to result from direct activation of trigeminovascular system brainstem regions involved in headache generation, or seizure-related parenchymal changes triggering the activation of the trigeminovascular system.

[H2] Post-ictal headaches

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Evidence from preclinical studies in rats indicates that seizures can be followed by spreading depolarisation^{119–122}; however, post-ictal spreading depolarisation has not been observed in humans (except for studies in individuals with brain damage^{123,124}) suggesting that this mechanism is not responsible for post-ictal headache.

Experimental evidence also indicates that neurons do not remain depolarised after the termination of tonic—clonic seizures, but instead become hyperpolarized¹²⁵(Box 2). This post-ictal neuronal silencing is sudden and widespread, instead of spreading¹²⁶. Preclinical studies indicate that the mechanisms underlying post-ictal silencing are multifactorial^{126,127}, including astrocytic adenosine release¹²⁸, acidosis and hypoxia-related vesicular transmitter depletion^{128,129}, none of which have been implicated in the initiation of spreading depolarization. There is no clinical evidence that post-ictal spreading depolarization contributes to post-ictal neuronal silencing (Box 2). In people with epilepsy, levels of adenosine were found to be enhanced post-ictally up to 18 minutes after seizures¹³⁰, and post-ictal acidosis is evidenced from postictal hypercapnia¹³¹ and enhanced plasma levels of lactate¹³². Clinical evidence for post-ictal neurotransmitter depletion is lacking¹³³. Analysis of neocortical tissue from individuals with chronic epilepsy and a rat model of epilepsy suggested that the low likelihood of spreading depolarisation in epileptic tissue results from intrinsic changes in GABAergic transmission¹³⁴.

Evidence from studies in rodent brain slices indicates that, even in the absence of post-ictal spreading depolarisation, excessive neuronal network activation during seizures can lead to trigeminovascular system activation via mechanisms such as the build-up of K⁺, acidosis and neuronal release of CGRP during or shortly after a seizure^{135–137}. On the basis of evidence from preclinical studies, activation of meningeal nociceptive fibres by such compounds would be expected to lead to perception of headache by thalamocortical activation within 10–30 minutes⁷⁴, in line with a post-ictal phenomenon. Inflammatory changes also occur during seizures¹⁰², for example, neuronal release of HMBG1 was shown to occur within an hour of seizure initiation in animal models¹⁰⁴. It is possible that following seizures, these enhanced HMBG1 levels activate the trigeminovascular system (similar to the activation after spreading depolarization observed in experimental studies) causing post-ictal headache, although this hypothesis has not yet been tested in animals or humans. Last, seizures can yield post-ictal hypoperfusion as shown in rodent¹³⁸ and some clinical epilepsy studies^{139,140}. The resulting hypoxia¹³⁸ might be sufficient to trigger headache mechanisms as occurs in hypoxia-induced migraine attacks¹⁴¹.

[H1] Overlapping genetics

Variants in > 200 genes have been identified as causing or enhancing the risk of specific types of epilepsy¹⁴².

Some monogenic forms of epilepsy exist, but for other epilepsies the genetic risk is complex and polygenic 143.

Juvenile myoclonic epilepsy has both a monogenetic and a complex genetic origin. In one study, 70% of people with this form of epilepsy reported a family history of migraine, almost twice as many as in an age-matched and sex-matched control group, suggesting an overlap in genetic risk between juvenile myoclonic epilepsy and migraine ¹⁴⁴. Some specific genes have also been associated with both epilepsy and migraine^{66,145}. This commonality is most evident in familial hemiplegic migraine (FHM), which is an autosomal dominant subtype of migraine with aura, characterised by a transient hemiparesis during the aura and headache characteristics that are identical to those observed in common forms of migraine 146,147. Three genes have been associated with FHM: CACNAIA, which is located on chromosome 19p13 and encodes a subunit of neuronal voltage-gated Ca²⁺ channel 2.1 (Ca_V2.1)¹⁴⁸: ATP1A2 ¹⁴⁹, which is located on chromosome 1g23 and encodes the α2 subunit of the glial Na⁺/K⁺-ATPase; and SCNIA¹⁵⁰, which is located on chromosome 2q24 and encodes a subunit of neuronal voltage-gated sodium channel 1.1 (Na_V1.1). These three genes form the basis for the definition of three subtypes of FHM: mutations in CACNAIA cause FHM1, mutations in ATP1A2 cause FHM2 and mutation in SCNIA cause FHM3. For all three forms of FHM, specific mutations have been linked to specific presentations of migraine and epilepsy^{147,150–153}. In FHM1 the 'mild' R192Q mutation in CACNAIA causes hemiplegic migraine without epileptic features¹⁴⁸, whereas the more severe S218L mutation can also cause seizures¹⁵². In FHM2, novel missense mutations in ATP1A2 can result in the co-occurrence of migraine and childhood epilepsy 151. In FHM3, different mutations in SCNIA have been be associated with either childhood epilepsy¹⁵⁰ or generalised tonic-clonic seizures¹⁵⁴. One study found that, in people with epilepsy and FHM3, generalized seizures occurred independently from hemiplegic migraine attacks¹⁵⁴, suggesting that FHM and epilepsy share common molecular pathways. Functional studies of FHM-associated mutations in vitro and in transgenic animal models have provided preclinical evidence that epilepsy and migraine result from partially overlapping genetic mechanisms ^{155,156}. These involve alterations to neuronal and glial ion transport, resulting in network mechanisms hyperexcitability^{61,146,155,157,158}. Transgenic knock-in mice carrying the human FHM1-causing S218L mutation mimic the phenotype observed in humans with the mutation and display spontaneous or cortical spreading depolarisation-induced generalized seizures 159,160. Results from in vitro studies suggest that the susceptibility for generalised seizures in FHM1 S218L mice is related to strongly enhanced excitatory transmission, resulting in

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excessive recruitment of excitatory and inhibitory neuronal networks 161,162. In FHM3, the spectrum of Na_V1.1 defects seems complex, and both gain-of-function and loss-of-function effects of mutations in SCNIA have been reported 163,164. The identification of gain-of-function effects of FHM3-associated mutations contrasts with the loss-of-function mutations in SCN1A that are associated with Dravet Syndrome and cause impaired firing of inhibitory interneurons¹⁶⁵. Computational work indicates that dynamic changes in the activity of genetically affected excitatory and inhibitory neuronal networks, and associated changes in ion activity determine whether neuronal hyperexcitability may result in a seizure, a cortical spreading depolarisation, or both 166 (Box 2). This observation underscores the complexity of predicting the functional outcome of shared genetic defects between epilepsy and migraine. Truncating deletions in the PRRT2 gene, which encodes a proline-rich transmembrane protein, were identified in a small number of people with (hemiplegic) migraine 167,168, as a result of which PRRT2 was put forward as the fourth FHM-associated gene. However, the same and similar PRRT2 deletions have been identified in people with paroxysmal kinesigenic dyskinesia, benign familial infantile convulsions and infantile convulsion choreoathetosis without signs of migraine 147. Therefore, the relationship between PRRT2 and migraine does not seem to be precise. A missense mutation in the SLC1A3 gene, which encodes the glutamate transporter EAAT1 that is important in removing glutamate from the synaptic cleft¹⁶⁹, has been associated with severe episodes of ataxia, epileptic seizures and hemiplegic migraine that can be explained by impaired glutamate transport¹⁶⁹. Other genetic findings associated with features of epilepsy and migraine include mutations in POLG and C10orF2, which encode mitochondrial DNA polymerase¹⁷⁰ and Twinkle helicase¹⁷¹, respectively, and are involved in the maintenance of neuronal and glial energy supply. Some evidence suggests that mutations in mitochondrial genes associated with MELAS syndrome can predispose individuals to dysfunctional oxidative brain metabolism, explaining the cooccurrence of migraine-like episodes and epilepsy features in individuals with this syndrome^{172,173}. The genetic associations between polygenic forms of epilepsy and migraine remain unclear. However, a greater

prevalence of migraine has been observed among family members of people with non-acquired focal epilepsy or

generalised epilepsy than in the general population¹⁷⁴, indicating a shared genetic susceptibility to both conditions.

The results of a large-scale genome-wide association study identified a correlation between variants associated

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with migraine, especially migraine with aura, and variants associated with epilepsy; however, this correlation did not reach statistical significance¹⁷⁵.

[H1] Clinical management

[H2] Impact and diagnosis

The results of a cross-sectional study indicated that ~50% of individuals with headache and epilepsy report the headaches as severe²¹. Headaches linked to epilepsy negatively affect quality of life²¹. A study at an epilepsy clinic found that depression and anxiety were linked to the presence of headache¹⁵. Postictal headaches, in particular, were associated with depression and suicidality. The first step for successfully managing any condition is a correct diagnosis. The results of a Dutch questionnaire-based study found that neurologists underestimate the occurrence of headache among individuals with epilepsy²⁸. This observation suggests that increased awareness among neurologists of the association between epilepsy and headache is required. Atypical or persistent headaches not responding to standard treatment should suggest a possible epileptic origin, warranting an EEG-recording during the symptomatic (headache or possible migraine aura) phase. We are not aware of published guidelines on managing headaches in people with epilepsy, so we summarize the current practice below, providing suggestions for managing headaches in people with epilepsy based on the currently available evidence and our expertise.

[H2] Management of headaches in epilepsy

Physicians managing the care of individuals with epilepsy should actively enquire about ictal, pre-ictal, and postictal headaches. An EEG recording of a headache event is mandatory to ascertain whether or not headaches have an epileptic origin, especially in the case of atypical, short-lasting and/or peri-ictal headaches ^{19,45}. Interictal and peri-ictal headaches that the individual reports as moderate or intense, once correctly diagnosed, should be treated with analgesics. If migraine is diagnosed concomitantly with epilepsy or vice-versa, an anti-seizure medication that also has proven efficacy for migraine should be prescribed whenever possible to avoid polypharmacy and possible drug—drug interactions ^{176,177}. The anti-seizure medications topiramate and valproate are approved for treatment of migraine by the FDA and European Medicines Agency ^{178–180}. However, topiramate and valproate can be teratogenic, so neither is suitable for treating women of child-bearing age^{181–183} unless there is no other

especially for migraine¹⁸⁴.

Paradoxically, headaches are a common (>10%) adverse-effect of anti-seizure medication, and are most often associated with carbamazepine, phenytoin, lamotrigine and levetiracetam¹⁸⁵. When evaluating headache in epilepsy, the possibility of an adverse effect of medication should be considered. Lower doses of topiramate, valproate or lamotrigine are used for the treatment of migraine than for the treatment of epilepsy, but people with migraine still seem to be more prone to the adverse effects of these medications than people with epilepsy¹⁸⁶. People with migraine or migraine and epilepsy are also more likely to discontinue medication than those with epilepsy alone¹⁸⁶. Medications used for migraine have not been associated with seizures. Individuals with pharmacoresistant focal epilepsy can benefit from a resection of the epileptic focus; 34%–74% become seizure-free following the procedure¹⁸⁷. However, in one study 12% of participants who underwent the procedure subsequently developed chronic headaches, which persisted for > 1 year after surgery¹⁸⁸.

effective treatment available¹⁷⁹. Other anti-seizure medications, such as lamotrigine, can be used off-label,

[H2] Novel pharmacological therapies

Novel pharmacological therapies for migraine include those that target calcitonin gene-related peptide (CGRP), a trigeminal sensory neuropeptide that is expressed in neuronal tissue and distributed in discrete areas of the central and peripheral nervous system¹⁸⁹. Although the precise mechanisms are unknown, activation of the trigeminovascular system seems to be associated with the increased release of CGRP from C-fibres in the trigeminal ganglion. Upon its release, CGRP binds to its receptor on $A\delta$ -fibres, leading to pain perception¹⁹⁰. The results of clinical trials of CGRP-inhibiting drugs in migraine have shown an efficacy that is superior to placebo, and generally good tolerability¹⁹¹, making these drugs an attractive new avenue for acute and prophylactic treatment of migraine. CGRP-inhibiting drugs hold particular promise for individuals with difficult-to-treat migraine, who have high unmet needs and few treatment options^{191–193}. CGRP has vasodilatory effects and is important for blood pressure regulation^{189,194} and the long-term effects of CGRP-inhibition, especially in individuals with cardiovascular comorbidities, are still unknown¹⁹⁵. Interestingly, the results of a study published in 2018 indicate that the new anti-seizure medication perampanel, which acts on glutamatergic

AMPA receptors, inhibits CGRP release in rat brainstem¹⁹⁶. This observation suggests that perampanel could, in theory, be effective in treating peri-ictal headaches, although this has not been investigated yet.

Cannabidiol has received considerable media attention^{197–199} after a case report indicated that it can reduce seizure frequency in individuals with epilepsy²⁰⁰. The results of clinical trials in Dravet syndrome ^{201–203} and Lennox–Gastaut syndrome^{204,205} suggest that cannabidiol is more effective than placebo in reducing the frequency of convulsive and drop seizures²⁰⁶. Additional open-label studies of cannabidiol in other types of epilepsy are ongoing^{207–209}. An oral cannabidiol solution has been approved by the FDA²¹⁰ and the European Medicines Agency²¹¹ for treatment of seizures in children aged 2 years and older with Dravet syndrome and Lennox–Gastaut syndrome, two rare forms of severe epilepsy. One trial to assess the effect of cannabis on migraine is ongoing²¹² and another is planned²¹³.

[H2] Non-pharmacological approaches

A meta-analysis of studies on transcranial magnetic stimulation (TMS) found that low-frequency TMS was associated with a reduction in seizure frequency in 30% of participants with treatment-resistant epilepsy²¹⁴. The studies included in this analysis were, however, relatively small and heterogeneous, so more evidence to support this approach is needed. A systematic review of TMS for the treatment of headache disorders found that stimulation was associated with reduced headache frequency, duration, intensity and medication use; however, few studies reported TMS-associated changes greater than those observed with sham treatment²¹⁵. Several studies have found an association between treatment with single-pulse TMS and a reduction in headache days and medication use in individuals with migraine with aura²¹⁶⁻²¹⁸. This evidence led the FDA to approve a single-pulse TMS device for the acute treatment of this type of migraine²¹⁹. Evidence from a study using a rat model of migraine suggests that the effect of TMS on headache involves the suppression of cortical excitability, including the cortical spreading depolarisation that underlies the aura phase²²⁰. Clinical trials have found non-invasive stimulation of the trigeminal nerve to be moderately effective for acute migraine treatment²²¹ and prevention²²². Non-invasive stimulation of the vagus nerve was highly effective for acute migraine treatment²²³ but ineffective for migraine prevention²²⁴. In three small randomized controlled trials (n<150 in each study) this form of vagus nerve stimulation was also shown to be effective in drug-resistant epilepsy ²²⁵⁻²²⁷.

Evidence is emerging that therapeutic education, including the provision of information on lifestyle factors such as sleep and alcohol consumption as well as behavioural, self-management and mind-body approaches can have beneficial effects for individuals with chronic conditions, including headache^{228,229}, migraine^{230–232} and epilepsy^{233,234}. Although therapeutic education approaches do not cure these conditions, they can help individuals cope with the associated psychological burden. The ILAE recently recommended the widespread implementation of such techniques for people with epilepsy²³³.

[H1] Conclusions and future challenges

Clear evidence exists for an association between headaches and epilepsy. The results of studies published in the last five years have confirmed that headaches, especially migraines, often co-occur with epilepsy. This observation is in keeping with the growing body of evidence that comorbidity and multi-morbidity are common in neurological conditions^{235,236}. Highlighting this overlap during neurological and medical training should help neurologists and general physicians be more attentive to the association between headaches and epilepsy. The gap between headache and epilepsy classifications highlights the need for closer collaboration between specialists, within departments and between professional bodies such as the ILAE and IHS. Such partnership could lead to the development of standardised questionnaires to aid the diagnosis of headache in epilepsy and guidelines on the management of comorbid headache and epilepsy. These diagnostic tools and guidelines will help improve the treatment, care, and management of these complex conditions.

To improve our understanding of the nature of the association between epilepsy and headache, and to establish

the direction of this association, thorough longitudinal studies in large, multi-centric cohorts will be vital. Additional research efforts aimed at elucidating the pathophysiological mechanisms underlying headache in epilepsy and improving the management of these conditions are also needed. Although the pathophysiological mechanisms underlying epilepsy and migraine are highly complex, animal models of comorbidity^{103,237} will help uncover the mechanistic links between activation of the trigeminovascular system and epilepsy.

In conclusion, headaches, and epilepsy are not separate disease entities but seem to be symptoms of altered neuronal network excitability. Ultimately, it will be important to elucidate the various, likely multifactorial,

causes underlying the different epilepsy–headache constellations thus enabling the development of aetiological diagnostic classifications and corresponding therapies.

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Review criteria

We searched PubMed for articles with the MeSH terms and keywords "headache", "migraine" "epilepsy" and "treatment" in the title, abstract or keywords. The search focused on primary studies published in the last 5 years (April 2015 – April 2020). Additional articles were identified from the authors' own files and from chosen bibliographies. The articles in this Review were included at the authors' discretion on the basis of originality and relevance of the publication. Selected key works from before 2015 are shown in figure 1.

The authors affirm that human research participants provided informed consent for publication of the video in Supplementary Video 1.

Key points

Informed consent

- The lifetime prevalence of migraine is 52% greater in individuals with epilepsy than in individuals with epilepsy.
 - The symptoms of epilepsy and headache can present diagnostic challenges; a detailed history and EEG recording of the epileptic and/or headache event are important for classification and management.
 - Enhanced neuronal excitability might be the mechanistic link between headaches and seizures.
- Several genetic mutations can cause epilepsy and migraine, but the genetic association between polygenic form of epilepsy and migraine remains unclear.

 Novel therapies include calcitonin gene-related peptide-blocking drugs for migraine and neuromodulative non-pharmacological approaches for migraine and epilepsy; behavioural and selfmanagement approaches are increasing in popularity.

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Study	Cohort size and type	Case ascertainm ent	Number reporting headache								
			Total	Pre- ictal	Ictal	Post-ictal	Inter- ictal	Inte r- ictal and pre- ictal	Pre - icta l and pos t- icta l	Post- ictal and inter- ictal	Pre- ictal , post - ictal and inte r- ictal
Begasse de Dhaem 2019 ¹⁸	349 (209 female); new-onset focal epilepsy	Validated headache questionnair e (ICHD)	74 (21.2 %) migrai ne	NA	NA	NA	NA	NA	NA	NA	NA
Çililer 2017 ¹⁰	349 (190 female); consecutive epilepsy cases (69 partial seizures; 209 generalised seizures; 71 secondary generalised seizures)	Interview with questionnair e (ICHD-2)	152 (94 MI; 60 TTH; 43 U)	19 (12 MI; 4 TTH; 3 U)	NA	82 (30 MI; 25 TTH, 27 U)	17 (8 MI; 7TTH, 2 U)	NA	33	26	16
Hofstra 2015 ¹³	255 (126 female); cross-sectional	Questionnai re, ICHD-2 criteria	186 (65 MI; 97 TTH; 15 U)	3	NA	28	92	NA	NA	NA	NA
Kim 2016 ¹⁷	831 (391 female); consecutive video EEG cases (775 partial seizures; 55 generalised seizures)	Epileptic aura description, follow-up by phone interview (457 no aura; 374 with aura)	NA	25 (all partial seizure s)	6 (2 hemicra nia epileptic a, 4 R- TLE, 1 L-TLE, 1 Central seizure)	257 (238 partial ^b ; 18 generalis ed)	NA	NA	NA	NA	NA
Mainieri 2015 ¹²	388 (209 female); consecutive cases with epilepsy (101 generalised epilepsy; 280 focal epilepsy; 7 U)	Self-report and structured interview	209	26 (16 MI; 5 TTH; 5 other)	3	74 (37 MI; 30 TTH)	188 (102 MI ^d ; 74 TTH; 2 cluster ; 9 U)	NA	NA	NA	NA
Mameniški enė 2016 ²¹	289 (172 female); adults with epilepsy treated in epilepsy center	Self-report and structured interview	233 (69 MI, 85TT H, 79 other)	23	1	46	218 (69 MI, 85 TTH, 52 other	NA	NA	NA	NA

Mutlu 2018 ¹⁴	420°; consecutive outpatient cases	Interview (ICHD)	111 (63 MI)	29 (9 MI)	NA	32 (5 MI)	83 (58 MI)	15 (5 MI)	17 (3 MI)	NA	NA
Salma 2019 ¹⁹	47 (28 female); cases with epilepsy or unusual headache (33 focal epilepsy; 6 generalised epilepsy; 8 U)	Interview (ICHD)	37	2	22 (5 isolated IEH ^a)	10 (focal seizures)	15	NA	NA	NA	NA
Seo 2016 ¹⁵	177 (85 female); consecutive individuals with epilepsy diagnosis	Interview	73	3 (1 MI)	NA	48 (17 MI; 24 TTH; 7 U	34	NA	NA	NA	NA
Wang 2014 ¹¹	1109 (502 female) (856 partial seizures; 195 generalised seizures; 58 unclassified seizures)	Questionnai re, then interview (ICHD)	667	59 (38 MI)	NA	469 (314 MI)	231 (139 MI)	NA	9	45 (interict al migrain e)	9
Whealy 2019 ¹⁶	120 (67 female);epile psy monitoring unit	Questionnai re (ICHD 3)	NA	NA	NA	75 (15 definite MI; 23 probable MI; 10 definite TTH; 3 probable TTH; 24 U)	97 (22 definit e MI; 26 probab le MI; 14 definit e TTH; 13 probab le TTH; 22 U)	NA	NA	NA	NA

Table includes only studies published between 2014 and 2019 that were not included in the two meta-analyses^{8,9}, except for the studies highlighted in grey. ^a associated with focal onset, most often temporal lobe, ^b discrepancy in the original study, ^c Sex of participants not reported. ^d of which, 6 with aura. ICHD=International Classification of Headache Disorders. TTH=tension type headache, U= unclassified, TLE=temporal lobe epilepsy, FLE=frontal lobe epilepsy, OLE=occipital lobe epilepsy; MI, migraine.

Table 2 | Features of migraine aura and occipital seizures

Feature	Migraine	Occipital lobe seizure	1149
Main symptoms	Foggy or blurred vision	Visual hallucinations	
	Zigzag or jagged lines	Visual illusions	
	Scotoma	Blindness	
	Phosphenes	Palinopsia	
	Flickering light	Sensory hallucinations of ocular	
		movement	
		Ocular pain	
		Nystagmus, eyelid closure and/or	
		fluttering	
Duration	1060 minutes	<1 minute	
Progression	Centrifugal or centripetal	No centrifugal or centripetal	
	progression of visual	progression of visual symptoms	
	symptoms		
Accompanying symptoms (e.g.	Common	Rare	
nausea, vomiting, photo-			
phonophobia)			

Figure 1 | A selection of key publications on headache in epilepsy from before 2015.

This timeline shows milestone publications in the field of headache in epilepsy. We selected publications that were particularly notable, for example, the first publication to report a specific finding, or a publication that had a large influence on subsequent research. The first reports of an overlap between epilepsy and headache were published at the end of the 19th century. From the 1960's onward, epilepsy was increasingly seen as a systemic disorder with many comorbidities. Technical advances in the 1980's spurred on research in this area, including studies that used animal models, in vitro approaches and depth electrodes in patients. From the early 2000's, there was an increased interest in the molecular mechanisms of anti-seizure medication and their effect on associated conditions such as migraine, and in the molecular genetics of epilepsy and migraine.

Figure 2 | A timeline showing the different types of peri-ictal headaches.

The timing of pre-ictal, ictal and post-ictal headaches is shown in relation to the seizure. Pre-ictal headaches occur < 24 hours before a seizure and last until seizure onset. Ictal headaches develop simultaneously with the seizure. Post-ictal headaches occur < 3 hours after the end of the seizure event and remit spontaneously < 72 hours after seizure termination. Specific types of seizure-related headaches are also illustrated, including migraine as seizure trigger, hemicrania epileptica and headache as seizure aura.

Figure 3 | Putative pathophysiological mechanisms linking seizures and headache. a | Hyperexcitability in epilepsy often involves impaired GABAergic transmission, facilitating hypersynchronous seizure bursts. In migraine, hyperexcitability seems to be largely the result of enhanced glutamatergic transmission, which could facilitate pain pathway activation via inflammatory changes and calcitonin gene-related peptide (CGRP) release in the absence or presence of CSD. In migraine, GABAergic transmission seems to be unaltered or could be dynamically enhanced, as indicated by the results of preclinical studies on the effects of mutations associated with familial hemiplegic migraine (FHM) type 3. Strongly enhanced glutamatergic transmission in migraine resulting from pathogenic mutations, as is known to occur in FHM type 1, will increase the likelihood of co-morbid epilepsy. b | Cortical spreading depolarization (CSD) is likely to be the neurophysiological mechanism underlying

migraine aura. CSD could also trigger migraine headache originating in the trigeminovascular system. CSD consists of a slowly propagating wave of network depolarization that is presumably caused by cortical hyperexcitability. CSD-associated increases in the concentration of potentially noxious molecules, including K⁺ and H⁺ (i.e. low pH), in the extracellular space could reach pial, arachnoid, and dural surfaces and activate perivascular sensory afferents from trigeminal ganglion (TG) neurons. Inflammatory changes, involving neuronal release of high mobility group protein 1 (HMBG1) following CSD-induced pannexin channel opening, provide a mechanistic link between CSD and pain pathway activation. Signals from activated meningeal nociceptors are relayed through TG nerve processes to the brainstem trigeminal cervical complex (TCC) and subsequently to thalamic and cortical areas (including cingulate cortex, CC) and produce sensations of pain. Adapted from Chen et al, Cephalalgia 2019 and Ferrari et al Lancet Neurology 2015.

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- **Box 1** | ICHD-3 diagnostic criteria relevant to epilepsy
- Migraine aura-triggered seizure (ICHD-3 code 1.4.4.)
- A. A seizure fulfilling diagnostic criteria for one type of epileptic attack and criterion B below
- B. Occurring in a patient with 1.2 Migraine with aura, and during or within one hour after an attack of migraine with aura
- 1195 C. Not better accounted for by another ICHD-3 diagnosis.
- While migraine-like headaches are quite frequently seen in the epileptic post-ictal period, sometimes a seizure occurs during or following a migraine attack.
- This phenomenon, sometimes referred to as migralepsy, is a rare event, originally described in patients with 1.2

 Migraine with aura. Evidence of an association with Migraine without aura is lacking.
 - Ictal epileptic headache (ICHD-3 code 7.6.1)
- A. Any headache fulfilling criterion C
- B. The patient is having a partial epileptic seizure

C. Evidence of causation demonstrated by both of the following: 1203 1. headache has developed simultaneously with onset of the partial seizure 1204 2. either or both of the following: a) headache is ipsilateral to the ictal discharge. b) headache significantly 1205 improves or remits immediately after the partial seizure has terminated 1206 D. Not better accounted for by another ICHD-3 diagnosis. 1207 Hemicrania epileptica (ICHD-3 code 7.6.1.) 1208 (if confirmed to exist) is a very rare variant of 7.6.1 Ictal epileptic headache characterized by ipsilateral location of headache 1209 and ictal EEG paroxysms. 1210 Postictal headache (ICHD-3 code 7.6.2) 1211 A. Any headache fulfilling criterion C 1212 B. The patient has recently had a partial or generalized epileptic seizure 1213 C. Evidence of causation demonstrated by both of the following: 1214 1. headache has developed within three hours after the epileptic seizure has terminated 1215 2. headache has resolved within 72 hours after the epileptic seizure has terminated 1216 D. Not better accounted for by another ICHD-3 diagnosis. 1217 1218 Box 2 Spreading depolarization and seizures – a missing link underlying headache in epilepsy? 1219

Migraine aura⁵ is likely to be caused by cortical spreading depolarisation, a slow-spreading (\sim 2–6 mm per min) wave of neuronal and glial depolarisation followed by neuronal silencing (evident from suppression of local field potential (LFP) or EEG activity) lasting a couple of minutes^{238–240}. Neuronal hyperexcitability predisposes to spreading depolarisation and seizures, and might be a key shared mechanism of epilepsy and migraine^{158,241}. Changes in ion concentration can shift neurons towards moderate depolarisation leading to synchronous epileptiform firing (part a of the figure), or — if extracellular K⁺ ([K⁺]_{out}) rises above \sim 12 mM — towards near-complete depolarisation, yielding spreading depolarisation^{122,166} (part b of the figure shows a hypothetical seizure

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followed by spreading depolarisation). Silencing of bioelectrical activity during spreading depolarisation is caused by sustained neuronal depolarisation that exceeds the inactivation threshold for ion channels, thus preventing action potentials¹²³. Conversely, post-ictal suppression in the absence of spreading depolarisation is associated with neuronal hyperpolarisation 136. Spreading depolarisation-related suppression should not be confused with post-ictal generalised EEG suppression (PGES)²⁴², which is an immediate (within 30 seconds) complete suppression of EEG activity following a seizure^{243,244}. Clinically, PGES appears non-spreading²⁴⁵, lasts up to 338 seconds (mean 46 seconds) and is associated with motionlessness²⁴⁶, whereas changes in perception associated with migraine aura last $\sim 20-30$ minutes¹²³. Preclinical work has indicated that network suppression by spreading depolarisation prevents seizures¹¹¹, suggesting that post-ictal spreading depolarisation constitutes an intrinsic seizure-termination process. The link between spreading depolarisation and headache remains unclear. In rodents, spreading depolarisation activates the trigeminovascular system at the meningeal level^{77,247} (Fig. 3) and might affect the brainstem via a corticotrigeminal projection⁷⁴. How the trigeminovascular system is activated in humans remains unclear, and cortical spreading depolarisation could be one of many triggers⁷⁴. No clear evidence exists that spreading depolarisation occurs in association with epileptic discharges in humans outside of trauma or stroke^{123,124}. When cortical spreading depolarisation was observed in individuals with ischemic stroke, headaches were not reported²⁴⁸. Research in rodents indicates that the excessive network activity during seizures and associated increases in extracellular K⁺, H⁺ and inflammatory changes might be sufficient to activate the trigeminovascular system without the need of a spreading depolarisation ¹⁰³.

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Part A adapted from REF¹³⁶. Part B is a stylized representation of the changes that are thought to occur during post-ictal spreading depolarization^{122,158}.

Supplementary Video 1 | Video-EEG recording of an individual with ictal epileptic headache