Headache Management in Individuals with Brain Tumor

Ami Cuneo, MD1 Natalia Murinova, MD1

1 Department of Neurology, University of Washington, Seattle, Washington

Address for correspondence Ami Cuneo, MD, Department of Neurology, University of Washington, 1959 NE Pacific Street, Seattle, WA 98195-6182 (e-mail: amiza@uw.edu).

Abstract

Headache occurs commonly in individuals diagnosed with cerebral neoplasm. Though the features of a brain tumor-associated headache may vary, a progressive nature of headache and a change in headache phenotype from a prior primary headache disorder often are identified. Pathophysiologic mechanisms proposed for headache associated with brain tumor include headache related to traction on pain-sensitive structures, activation of central and peripheral pain processes, and complications from surgical, chemotherapeutic and/or radiotherapy treatment(s). Optimization of headache management is important for an individual’s quality of life. Treatments are based upon patient-specific goals of care and may include tumor-targeted medical and surgical interventions, as well as a multimodal headache treatment approach incorporating acute and preventive medications, nutraceuticals, neuromodulation devices, behavioral interventions, anesthetic nerve blocks, and lifestyles changes.

Keywords

► headache
► brain tumor
► headache treatment

Headaches occur commonly in the general population. Affecting more than 90% of individuals during the lifetime,1 headaches can range from an occasional nuisance to a severe, disabling condition. Within this broad clinical spectrum, most headaches are related to functional, neurochemical changes in an individual with a genetic and/or environmental predisposition. Examples of such primary headache disorders include migraine, tension-type headache, and cluster headache (►Table 1).2 In comparison, secondary headache disorders are attributed to an underlying brain pathology, such as an intracranial neoplasm, vascular condition, cerebrospinal fluid (CSF) pressure-related disorder, or other etiology (►Fig. 1).2,3

Headache attributed to intracranial neoplasm is defined by the International Classification of Headache Disorders, 3rd edition (ICHD-3) criteria.2 In an individual with a known space-occupying neoplasm, the headache must develop in temporal relation to the neoplasm, worsen or improve in parallel with progression or successful treatment of the neoplasm, respectively, and have characteristics, such as progressive headache worsened in the morning/when laying down, headache aggravated by Valsalva-like manœuvres, and/or headache accompanied by nausea and/or vomiting (►Table 2a).2 The ICHD-3 also includes definitions for headache attributed to carcinomatous meningitis and pituitary lesions (►Table 2b-c).2

Concern for an underlying brain tumor is raised frequently by individuals presenting for headache evaluation.4 However, intracranial neoplasms comprise only 1% of newly diagnosed malignancies. Furthermore, brain tumors are identified on magnetic resonance imaging (MRI) in just 1.5% of those undergoing headache workup.5,6 When present, a cerebral neoplasm is 10 times more likely to be a metastatic lesion (e.g., from lung or breast cancer, or melanoma) than a primary brain tumor.5,7,8 Though headache is rarely the only symptom of a brain tumor, in clinical practice the diagnostic discernment of a primary versus secondary headache can be challenging, due in part to the high prevalence of primary headache disorders in the general population and because features of secondary headaches can mimic primary headaches.9,10

A wide range of treatment options can be considered for headache related to cerebral neoplasm. Tumor-targeted treatments, such as surgical resection, chemotherapy, radiation...
Table 1  Characteristics of common primary headaches as compared with headache related to intracranial neoplasm, as defined by the ICHD-3 criteria

<table>
<thead>
<tr>
<th>Headache characteristics</th>
<th>Migraine</th>
<th>Tension type</th>
<th>Cluster</th>
<th>Headache attributed to intracranial neoplasm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minimum number of attacks</td>
<td>5</td>
<td>10</td>
<td>5</td>
<td>N/A</td>
</tr>
<tr>
<td>Duration (untreated or unsuccessfully treated)</td>
<td>4–72 h</td>
<td>30 min to 7 d</td>
<td>15–180 min</td>
<td>N/A</td>
</tr>
<tr>
<td>ICHD-3 diagnostic features</td>
<td>Must have ≥ 2 of 4 features: –Unilateral location –Pulsating quality –Moderate or severe intensity –Causing avoidance of routine physical activity</td>
<td>Must have ≥ 2 of 4 features: –Bilateral location –Pressing quality (non-throbbing) –Mild or moderate intensity –Not causing avoidance of routine physical activity</td>
<td>Severe or very severe unilateral, orbital, supraorbital, and/or temporal pain occurring with a frequency between 1 every other day and 8 per day</td>
<td>Requirements: –A space-occupying intracranial neoplasm has been demonstrated –Must have ≥ 2 of: –Headache developed in temporal relation to neoplasm, or led to its discovery –Headache either worsened with disease progression and/or improved in temporal relation with treatment of the neoplasm · Headache has ≥ 1 of 4: –Progressive –Worse in morning and/or when laying down –Aggravated by Valsalva –Associated with nausea and/or vomiting</td>
</tr>
<tr>
<td>Associated symptoms</td>
<td>Must have ≥ 1 of: –Light sensitivity and sound sensitivity –Nausea and/or vomiting</td>
<td>Must have both of: –No nausea or vomiting –No more than 1 of light or sound sensitivity</td>
<td>Must have ≥ 1 ipsilateral autonomic symptom (e.g., lacrimation, rhinorrhea, nasal congestion) or associated restlessness/agitation</td>
<td>Typically involves associated neurologic symptoms or signs on physical exam related to expansion of the mass. May be associated with intracranial hypertension.</td>
</tr>
<tr>
<td>Forms</td>
<td>Episodic migraine: &lt; 15 headache days per month Chronic migraine: ≥ 15 headache days per month, including ≥ 8 migraine days per month for &gt; 3 mo Migraine with aura: ≥ 2 episodes of fully reversible visual, sensory, or other symptoms lasting minutes usually associated with migraine</td>
<td>Episodic tension-type headache: &lt; 15 headache days per month Chronic tension-type headache: ≥ 15 headache days per month for &gt; 3 mo</td>
<td>Episodic cluster headache: ≥ 2 cluster periods lasting 7 d to 1 y (untreated) and separated by pain-free remission periods of ≥ 3 mo Chronic cluster headache: No remission or remission periods &lt; 3 mo for ≥ 1 y</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Note: For each headache diagnosis, the ICHD-3 criteria requires that it not be better accounted for by another ICHD-3 diagnosis.
therapy, and/or medications to reduce tumor-associated cerebral edema may be initiated on a case-by-case basis and can lead to improvement in headache. Importantly, individuals with cerebral neoplasm also may respond to standard therapies for primary headaches, based upon the patient’s tumor-associated headache phenotype and prior history of a primary headache disorder, though studies in this patient population are limited. In the last decade, new treatments for primary headaches, based upon the patient’s tumor-associated headache phenotype and prior history of a primary headache disorder, though studies in this patient population are limited.11 In the last decade, new treatments for primary headaches, based upon the patient’s tumor-associated headache phenotype and prior history of a primary headache disorder, though studies in this patient population are limited.11

Fig. 1  Differential diagnosis of primary versus secondary etiologies of headache. *A list of other primary headaches is included in the ICHD-3 guidelines.*

Table 2a-c. ICHD-3 definitions of headache attributed to intracranial neoplasia.2

2a. Headache attributed to intracranial neoplasm
Description: Headache caused by one or more space-occupying intracranial tumours.

Diagnostic criteria:
A. Any headache fulfilling criterion C
B. A space-occupying intracranial neoplasm has been demonstrated
C. Evidence of causation demonstrated by at least two of the following:
   1. headache has developed in temporal relation to development of the neoplasm, or led to its discovery
   2. either or both of the following:
      a) headache has significantly worsened in parallel with worsening of the neoplasm
      b) headache has significantly improved in temporal relation to successful treatment of the neoplasm
   3. headache has at least one of the following four characteristics:
      a) progressive
      b) worse in the morning and/or when lying down
      c) aggravated by Valsalva-like manoeuvres
      d) accompanied by nausea and/or vomiting
D. Not better accounted for by another ICHD-3 diagnosis.

2b. Headache attributed to carcinomatous meningitis
Description: Headache caused by carcinomatous meningitis, usually accompanied by signs of encephalopathy and/or cranial nerve palsies.

Diagnostic criteria:
A. Any headache fulfilling criterion C
B. Carcinomatous meningitis (in the presence of systemic neoplasia known to be associated with carcinomatous meningitis) has been demonstrated
C. Evidence of causation demonstrated by at least two of the following:
   1. headache has developed in temporal relation to development of the carcinomatous meningitis
   2. either or both of the following:
      a) headache has significantly worsened in parallel with worsening of the carcinomatous meningitis
      b) headache has significantly improved in parallel with improvement in the carcinomatous meningitis
   3. headache is associated with cranial nerve palsies and/or encephalopathy
D. Not better accounted for by another ICHD-3 diagnosis.
headache disorders offer promise for individuals suffering from headache. Therefore, incorporation of a multimodal headache treatment approach, including evidence-based acute and preventive medications, nutraceuticals, neuromodulation devices, lifestyle modifications, and behavioral therapies, can be considered. Additional attention to diagnosis of medication overuse headache (MOH), a common, superimposed secondary headache disorder, should be paid in individuals with headache and brain tumor, as treatment of MOH may reduce central sensitization to pain and improve headache-related outcomes.12

This article reviews headache characteristics and pathophysiology in cerebral neoplasm, as well as the diagnostic evaluation and management of headache in this patient population.

Headache Features in Individuals with Brain Tumor

Traditionally, clinicians have considered certain headache features suggestive of an underlying cerebral neoplasm, including positional headache (e.g., worsened when supine/upon awakening), headache exacerbated with Valsalva-like manoeuvres, and/or headache associated with nausea and/or vomiting (—Table 2).13,14 However, these features are not specific for headache related to intracranial neoplasm.

Additional brain tumor-associated headache characteristics have been described in the last decades. Forsyth and Posner found that most of these headaches had non-specific features, were mild, and mimicked migraine.14 Valentinis et al reported intracranial neoplasm-related headaches typically involved intermittent, moderate-intensity, pressure-like pain located in the bifrontal region, were rarely associated with nausea or vomiting, and often responded to analgesics; in this study, just 5.1% of patients undergoing surgery for brain tumors had the “classic” intracranial tumor-associated headache features.15 Schankin et al suggested that nearly 40% of brain tumor patients had tension-like headache, and that dull headache occurred more frequently with glioblastoma multiforme, whereas pulsating headache was associated with meningioma.10 Pfund et al observed that tumor location coincided with the lateralization of headache in only one-third of patients.16 The progressive nature of headache, however, has been identified in 79.1% of individuals with cerebral neoplasm.16,17

Diagnostic Evaluation and Workup of Headache Associated with Brain Tumor

Initial evaluation for headache related to brain tumor includes a comprehensive headache history. The most significant risk factor for headache development in cerebral neoplasm is a prior primary headache disorder,10,18,19; at the same time, secondary headaches may masquerade as tension-type headache, migraine, trigeminal autonomic cephalalgias, or other primary headache disorders.15,20–22 Taken together, these factors can lead to a diagnostic challenge for clinicians.

Additional neurologic history may provide diagnostic clues. For instance, an alteration in headache features over time may suggest a new underlying brain pathology.10 Presence of other neurologic symptoms can suggest secondary etiology of headache. Though headache is reported in up to 71% of individuals with brain tumor,14,16,23,24 only 1 to 2% of patients with cerebral neoplasm experience headache as the sole clinical symptom.10,25,26 Cerebral neoplasm-related headache commonly is accompanied by seizure and/or other focal neurologic symptoms.27

A comprehensive neurologic exam, including evaluation for mental status changes, visual field defects, papilledema, cranial nerve dysfunction, motor and sensory abnormalities, and cerebellar dysfunction, is important. Though focal exam findings may be present, the neurologic exam may be normal due to the indolent nature of brain tumors, which allows time for compensation of neural brain networks.28 In these cases, brain tumors may be asymptomatic and clinically silent, identified incidentally on imaging.7

Table 2a-c. (Continued)

2c. Headache attributed to hypothalamic or pituitary hyper- or hyposecretion

Description: Headache caused by a pituitary adenoma and hypothalamic or pituitary hyper- or hyposecretion, usually accompanied by disorder of temperature regulation, abnormal emotional state and/or altered thirst or appetite. It remits after successful treatment of the underlying disorder.

Diagnostic criteria:

A. Any headache fulfilling criterion C
B. Hypothalamic or pituitary hyper- or hyposecretion associated with pituitary adenoma has been demonstrated
C. Evidence of causation demonstrated by at least two of the following:
   1. headache has developed in temporal relation to onset of hypothalamic or pituitary hyper- or hyposecretion
   2. either or both of the following:
      a) headache has significantly worsened in parallel with worsening of the hypothalamic or pituitary hyper- or hyposecretion
      b) headache has significantly improved in parallel with improvement in the hypothalamic or pituitary hyper- or hyposecretion
   3. headache is associated with at least one of the following:
      a) disorder of temperature regulation
      b) abnormal emotional state
      c) altered thirst and/or appetite
D. Not better accounted for by another ICHD-3 diagnosis.
In gathering the history and performing the neurologic exam, attention should be paid to the identification of “red flags.” The SNNOOP10 criteria addresses risk factors for secondary etiologies of headache, such as systemic symptoms (e.g., fever); history of neoplasm or neurologic findings on exam; sudden-onset headache; older age (> 50 years); pattern change from prior headaches; positional headaches; headaches precipitated by cough, sneeze, or Valsalva-like manoeuvres; papilledema; pregnancy/post-partum; progressive symptoms; painful eye with autonomic features; immunosuppression history; and use of acute analgesics associated with MOH (Table 3). If any of these “red flags” is present, further workup to evaluate for secondary headache should be considered (Fig. 2).

Table 3 The SNNOOP10 criteria includes select red flags associated with secondary etiologies of headache.

<table>
<thead>
<tr>
<th>Red Flag</th>
<th>Examples of Clinical Signs, Symptoms, and/or Comorbidities</th>
<th>Select Related Secondary Headache Etiologies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systemic</td>
<td>Fever, chills, sweats, weight loss, systemic disease, immunocompromised status, pregnancy/puerperium</td>
<td>Secondary headaches associated with malignancy, infection/opportunist infection, inflammatory condition, and pregnancy (e.g., cerebral venous sinus thrombosis, reversible cerebral vasoconstriction syndrome (RCVS), posterior reversible encephalopathy syndrome)</td>
</tr>
<tr>
<td>Neurologic deficit</td>
<td>Focal neurologic signs on exam (e.g., papilledema, cranial nerve deficits, numbness, tingling, weakness, altered cognition) and/or seizure</td>
<td>Secondary headaches related to brain tumor (primary versus metastatic), infection (e.g., abscess, leptomeningeal disease, meningoecephalitis), vascular disorders, inflammatory/autoimmune conditions</td>
</tr>
<tr>
<td>Onset sudden</td>
<td>“Thunderclap headache,” defined as a rapid onset headache achieving maximal intensity within 1 minute</td>
<td>Subarachnoid hemorrhage, other vascular disorders, RCVS, pituitary apoplexy, arterial dissection, intracranial hypotension, Chiari malformation, hypertensive crisis, complicated sinusitis, giant cell arteritis, brain tumor</td>
</tr>
<tr>
<td>Older age</td>
<td>Age &gt; 50 years</td>
<td>Giant cell arteritis, brain tumor, vascular disorders, non-vascular intracranial disorders (e.g., inflammatory condition)</td>
</tr>
<tr>
<td>Pattern change and/or other ‘p’ features</td>
<td>New headache, change in headache frequency, severity, duration, location, and/or associated features; progressive headache; any new headache phenotype; headaches associated with position, precipitated by cough, sneeze, or exercise, painful eye +/- vision change, post-traumatic onset of headache, painkiller medication usage</td>
<td>Brain tumor, vascular and non-vascular (e.g., inflammatory) intracranial disorders, CSF-related disorders (e.g., intracranial hypotension, intracranial hypertension), posterior fossa malformation, Chiari malformation, acute angle closure glaucoma, subdural or epidural hematoma, medication overuse headache, medication side effect</td>
</tr>
</tbody>
</table>


Fig. 2 A decision-making algorithm for further diagnostic workup, based on the presence of SNNOOP10 criteria risk factors for secondary headache.
previously diagnosed with brain tumor, repeat imaging should be considered if headache features change, if new neurologic symptoms or focal signs on exam develop, or if seizure, cognitive change, and/or signs of intracranial hypertension are identified.13

Though the chance of finding a brain tumor on imaging in a headache patient with no history of malignancy is low,22,29 MRI or computed tomography (CT) of the brain can help identify space-occupying cerebral lesions. MRI is superior to CT both in terms of imaging resolution and visualization of the brain’s posterior fossa, whereas CT is more widely available and can be helpful in evaluating for a hemorrhagic component of tumor and/or associated cerebral edema. MRI spectroscopy can analyze the chemical composition of the lesion to help distinguish primary brain tumors from non-neoplastic etiologies (e.g., demyelinating disease, abscess, or stroke), but cannot diagnose tumor type. Histological evaluation from tumor biopsy or resection sample is required for confirmatory diagnosis.30 If neurologic history, exam, and/or imaging raise concern for carcinomatous meningitis, CSF studies including cytology and flow cytometry are obtained. Labs (e.g., inflammatory markers), opening pressure during lumbar puncture, and additional imaging studies (e.g., head and neck vessel imaging) can help identify non-neoplastic secondary etiologies of headache.

Routine imaging for all headache patients is not required due to the high prevalence of primary headache disorders in the general population in combination with a similar rate of incidental findings of brain tumor in individuals with or without headache.21,32 Regular performance of extensive workup for secondary etiology of headache in all patients would subject individuals to invasive testing, risk of false-positive diagnoses with incidental findings, and contribute to a high economic burden with limited diagnostic yield.32–34 An exception includes headaches with features of the trigeminal autonomic cephalalgias. Lesions in the posterior fossa, pituitary, trigeminal nerve, and other locations, as well as cerebral venous sinus thrombosis, have been associated with headaches with phenotypes of hemiplegic migraine,35 paroxysmal hemicrania,36 cluster headache,20,37 and/or short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing or cranial autonomic symptoms (SUNCT or SUNA, respectively).38–40 Therefore, MRI of the brain with and without contrast, as well as MR angiography of the head and neck, should be considered in individuals with these headache characteristics.31

Pathophysiology of Brain Tumor-Related Headache Pain

Several mechanisms for cerebral neoplasm-related headache pain have been hypothesized. The traction theory proposes brain tumor headache is related to compression and distension of pain-sensitive structures from tumor growth, surrounding edema, or secondary hemorrhage.13,41 Though the brain parenchyma itself lacks pain receptors, several intracranial and extracranial structures are pain-sensitive, including blood vessels, cranial nerves with afferent pain fibers (e.g., the trigeminal nerve), cervical nerves (e.g., C1 and C2), dura mater, periosteum, and extra-cranial tissue.13,42–45

Increased intracranial pressure can cause traction on pain-sensitive structures and trigger headache, as well.14,46 Intracranial pressure may increase in the setting of a rapidly expanding tumor with associated cerebral edema within the cranial vault.13,14,16 In addition, daily fluctuations in intracranial pressure can cause headache. For example, headache may be more pronounced upon awakening. Mechanistically, when in the supine position overnight, the diminished gravity-assisted venous drainage can contribute to cerebral edema. At the same time, a normal sleep-related increase in arterial pressure of carbon dioxide can lead to vasodilatation, followed by a rise in intracranial pressure and headache.44 In other cases, intraventricular, midline, and posterior fossa tumors may block CSF drainage, leading to increased intracranial pressure and traction upon periventricular pain-sensitive structures. For some individuals, a sudden-onset, paroxysmal headache triggered by change in body position or Valsalva-like manoeuvre may occur due to transient blockage of CSF flow at the foramen of Monro (i.e., from a colloid cyst or other tumor of the 3rd ventricle).13,44 Despite these factors, the risk of any particular cerebral neoplasm on headache is variable. Tumors in identical locations may have differing associations with headache,22,24 and tumors without apparent direct mass effect on pain sensitive structures have been associated with headache in some cases.13,16

Peripheral sensitization to pain also can contribute to headache pain. Prolonged irradiation of pain-sensitive structures from increased intracranial pressure can lead to release of pro-inflammatory neuropeptides at sites of inflammation, thereby exacerbating vascular edema, causing infiltration of immune cells, and leading to neurogenic inflammation with release of pro-pain substances, such as calcitonin gene-related peptide (CGRP) and substance P.44 The increased risk of rare headache phenotypes in individuals with secretory pituitary adenomas suggests that hormones (e.g., prolactin and growth hormone) may lead to biochemical changes in the hypothalamic-pituitary axis that contribute to headache, as well.3,47

Another proposed mechanism for headache in individuals with brain tumor includes central sensitization to pain from prolonged irradiation from pain-sensitive pericranial structures, as well as from dysfunction of the efferent, pain-inhibition pathways that descend from brainstem nuclei on trigeminocephalic and spinal nerves.13,17 Dysregulation of these descending pain inhibition pathways may contribute to prolonged headache pain.13,44

Headache additionally can occur as a direct acute or chronic complication of brain tumor treatment, such as radiation therapy, chemotherapy, or neurosurgical intervention.24 For example, during or immediately following radiation therapy, patients may develop acute injury (e.g., steroid-responsive damage to capillaries and leakage with edema), which can manifest clinically with headache, nausea, and
vomiting.\textsuperscript{48} Though the central nervous system (CNS) is protected in part by the blood–brain barrier, the CNS remains vulnerable to toxicity from some chemotherapies (e.g., retinoids, intrathecal methotrexate causing aseptic meningitis), resulting in headache among other neurologic symptoms, including encephalopathy, seizures, cerebrovascular complications, vision changes, cerebellar dysfunction, and neuropathy.\textsuperscript{39,50} While surgical intervention may be necessary in some cases of brain tumor, craniotomy can be associated with an independent risk of headache development predominantly at the surgical site due to mechanical disruption of the scalp’s sensory innervation from nerves, including the occipital nerve, trigeminal nerve, and sensory branches of the 2nd and 3rd cervical nerves (\textsuperscript{-Fig. 3}).\textsuperscript{18,46} Other secondary etiologies (e.g., CSF leak, hydrocephalus, cerebral hemorrhage, and/or meningoencephalitis) can lead to post-craniotomy headache in some individuals.\textsuperscript{19}

### Treatments for Headache Related to Cerebral Neoplasm and Intracranial Hypertension

A range of treatments may be used for headache management in patients with brain tumor. An individualized approach including patient-specific therapies should be pursued. Interventions targeting the brain tumor itself (e.g., neurosurgery, chemotherapy, radiation therapy, and/or management of intracranial hypertension) may be considered based upon the tumor's type, location, malignancy-potential, and extent of disease, as well as the patient's functional status and age.\textsuperscript{24,51}

Corticosteroids are recommended for treatment of headache from intracranial hypertension from tumor-related cerebral edema.\textsuperscript{13} This medication class is thought to decrease tumor-associated vasogenic edema via reduction of capillary permeability at the blood–brain barrier.\textsuperscript{52} The most potent corticosteroid, dexamethasone, has a relatively long half-life (36–72 hours) compared with prednisone (2–3 hours). Dexamethasone reaches complete efficacy at 24 to 72 hours and has limited mineralocorticoid effect.\textsuperscript{24} Though no standardized protocol for dexamethasone therapy exists, typically dexamethasone 4 mg daily can adequately control headache.\textsuperscript{25} Higher doses can be used, though they may be associated with dose-related toxicity (e.g., hyperglycemia, steroid-induced myopathy, avascular necrosis, infection, insomnia, and psychiatric disturbance) and have not been shown to be more effective.\textsuperscript{53} The role for corticosteroids in headache treatment for individuals with cerebral neoplasm without associated brain edema or mass effect remains controversial.\textsuperscript{13} Other treatments for cerebral edema, such as osmotic agents (e.g., mannitol) or diuretics (e.g., furosemide), do not have established efficacy for headache management.\textsuperscript{13}

For individuals with hydrocephalus related to CSF obstruction, ventricular drainage and/or ventricular shunting can be considered, if within the patient's goals of care.\textsuperscript{54} However, the incidence of shunt malfunction is elevated in patients with brain tumor compared with those experiencing hydrocephalus from other etiologies\textsuperscript{55} and the risk of early shunt failure after craniotomy in individuals with cerebral neoplasm is high (20%).\textsuperscript{56} Though not studied directly in hydrocephalus related to cerebral neoplasm, carbonic anhydrase inhibitors (e.g., topiramate or acetazolamide) may decrease CSF production in hydrocephalus, as established in other conditions, such as idiopathic intracranial hypertension.\textsuperscript{57,58}

For most patients with cerebral neoplasm-related headache, treatment with tumor debulking and/or decompression results in improvement of headache symptoms.\textsuperscript{16,59} In one study of 164 patients with brain tumor and headache, Pfund et al showed that only 8% of patients continued to experience headache 3 months postoperatively.\textsuperscript{16} In a 2018 study, headache prevalence was found to decrease from 52% preoperatively to 30% at 6 months postoperatively in brain tumor patients.\textsuperscript{59} Though neurosurgical intervention can help headache, in some individuals craniotomy itself may contribute to the development of a new, post-operative headache.\textsuperscript{60} Historically, craniotomy was considered less painful than other surgeries, due to fewer pain receptors in the dura and insensitivity of the brain parenchyma to pain.\textsuperscript{61,62} This, however, has been debated. More recent studies have suggested that craniotomy is associated with significant pain in up to 60 to 84% of individuals, often located at the surgical site due to disruption of the pericranial muscle and soft tissue, as well as injury to cranial nerves.\textsuperscript{18,60,63–65} and eventually, central sensitization to pain may develop.\textsuperscript{60} Suboccipital craniotomies, particularly using the retrosigmoid approach, are associated more frequently with chronic headache, with up to 66% of patients experiencing persistent headaches 3 years postoperatively.\textsuperscript{11,66}

New-onset or worsening headache also may occur as an adverse event related to certain chemotherapeutic agents (e.g., temozolomide, bromocriptine, bevacizumab, withdrawal of corticosteroids, and others),\textsuperscript{11,24,67} radiation therapy (particularly whole brain radiation),\textsuperscript{24} or as a symptom of a medical complication from these treatments (e.g., headache related to aseptic meningitis from intrathecal chemotherapy,
or from radiation encephalopathy, which can present with headache at 1–6 months post-radiation. Alternatively, palliative radiotherapy has been shown to decrease headache severity in 41% of individuals with metastatic cerebral neoplasm. In combination with tumor-specific treatments, a range of medication and non-medication approaches for acute and preventive symptomatic management of headache may be employed for patients undergoing surgical intervention, as well as for those for whom surgery is not indicated.

**Symptomatic Management of Post-Craniotomy Headache and Tumor-Related Headache in the Outpatient Setting**

**Acute Treatment**

In the post-craniotomy patient, headache may affect both neurosurgical recovery and quality of life. Fifty to 90% of individuals may require medication for pain relief post-craniotomy, however headache often is undertreated. This may be related in part to a dearth of evidence and consensus regarding treatment for post-operative pain following cranial neurosurgery.

Traditionally, medications used in the acute, post-surgical period have included acetaminophen, nonsteroidal anti-inflammatory drugs (NSAIDs), and opioids. Subcutaneous injection of sumatriptan, a medication used commonly as an abortive treatment for migraine, has been shown to be beneficial in a small study of individuals with post-craniotomy headache following microvascular decompression; in another study, subcutaneous sumatriptan was associated with decreased opioid use in both adult and pediatric populations following cranial surgery. Other treatments used in the acute, post-surgical setting include gabapentin and regional anesthetic nerve blocks.

Though acetaminophen, NSAIDs, triptans, and opioids typically are used in moderation in the short-term post-surgically, certain risks should be considered. NSAIDs, which cause impaired platelet aggregation, have been associated with post-surgical intracranial hemorrhage and also confer increased risk of post-operative renal failure. Opioids, though commonly used in the post-surgical setting, can be sedating and compromise the neurologic exam during the acute period in which mental status, as well as motor and sensory function, is assessed. Furthermore, frequent, prolonged use of opioids, acetaminophen, NSAIDs, and triptans can result in development of central sensitization to pain and MOH. Defined by the ICHD-3 criteria as transformation of episodic primary headaches into chronic headaches occurring more than 15 days per month (Table 4) in the setting of frequent use of abortive medications, a role for MOH in exacerbating secondary headaches, such as headache related to cerebral neoplasm, has been postulated. Therefore, the diagnosis and prevention of MOH should be addressed early in a patient’s clinical course. The variable influences of different medication classes upon the risk of development of MOH should be considered from the onset of headache symptoms. Opioids should be avoided when possible, due to risk of opioid-induced hyperalgesia and MOH; even low-frequency use of opioids and barbiturates (e.g., 8 or 5 days per month, respectively) can contribute to MOH. The use of NSAIDs and Tylenol should be limited to no more than 14 days per month in combination to reduce the risk of development of MOH. Triptans also can be associated with MOH and require limited use (no more than 9 days per month). If acute analgesics are required, the practitioner and patient should tally the number of days per month of use of any culprit medication(s) at each visit; a general guideline is to limit the total use of these medications to 8–9 days per month. Treatment for MOH includes withdrawal from the culprit medications and use of alternative pain medications in the acute setting. For example, gabapentin, which can be used for acute postoperative pain, has been associated with decreased analgesic consumption after surgery, though risks of delayed tracheal extubation and increased sedation may occur.

Regional nerve blocks using bupivacaine, ropivacaine, and other anesthetics injected at the scalp can decrease pain severity after craniotomy and may exert a long-lasting benefit without causing MOH.

For all patients with headache related to cerebral neoplasm, whether or not craniotomy has been performed, standard acute treatments targeting both the brain tumor-

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**Table 4 ICHD-3 definition of medication overuse headache**

<table>
<thead>
<tr>
<th>Medication overuse headache</th>
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<tbody>
<tr>
<td><strong>Description</strong></td>
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<tr>
<td><strong>Diagnostic criteria</strong></td>
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</table>

*Medications that can contribute to MOH include NSAIDs, acetaminophen, combination medications that include caffeine, triptans, opioids, ergotamine, butalbital, lasmiditan, and others.*
Table 5 Preventive medication treatments for migraine and tension-type headache phenotypes

<table>
<thead>
<tr>
<th>Preventive medication</th>
<th>Dosing range</th>
<th>Initial dose(^a)</th>
<th>Use for migraine vs tension type headache</th>
<th>Possible benefit for comorbidities</th>
<th>Side effects</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Antiepileptics</strong></td>
<td></td>
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<tr>
<td>Topiramate (PO)(^a)</td>
<td>25 mg/d to 200 mg/d</td>
<td>25–50 mg/d</td>
<td>Migraine Tension-type (limited evidence)</td>
<td>Weight loss</td>
<td>Paresthesia, memory impairment, dizziness, drowsiness, glaucoma, nephrolithiasis</td>
</tr>
<tr>
<td>Valproate (PO)(^a)</td>
<td>500 mg/d to 1,000 mg/d</td>
<td>250 mg twice daily</td>
<td>Migraine</td>
<td>Mood stabilization Epilepsy</td>
<td>Alopecia, dizziness, drowsiness, hematologic effects, hepatotoxicity, hyperammonemia, delayed hypersensitivity reactions, pancreatitis, weight gain, Teratogenic (^a)</td>
</tr>
<tr>
<td><strong>Antidepressants</strong></td>
<td></td>
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<tr>
<td>Amitriptyline (PO)(^b)</td>
<td>10 mg/d to 150 mg/d</td>
<td>10 mg/d</td>
<td>Migraine Tension-type</td>
<td>Depression Neuropathic pain Fibromyalgia</td>
<td>Anti-cholinergic effects (e.g., constipation, blurred vision, urinary retention), increased risk of bleeding if used with anti-platelet agents/anticoagulants, cardiac conduction abnormalities, arrhythmias, dizziness, drowsiness, cognitive dysfunction, orthostatic hypotension</td>
</tr>
<tr>
<td>Venlafaxine (PO)(^b)</td>
<td>70 mg/d to 225 mg/d</td>
<td>37.5 mg/d</td>
<td>Migraine</td>
<td>Anxiety Depression Neuropathic pain</td>
<td>Activation of mania/hypomania, hyponatremia, serotonin syndrome, sexual dysfunction, suicidal ideation, weight loss, nausea</td>
</tr>
<tr>
<td><strong>Anti-hypertensives</strong></td>
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<tr>
<td><strong>Beta-blockers</strong></td>
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<tr>
<td>Metoprolol (PO)(^a)</td>
<td>25 mg twice daily to 200 mg/d</td>
<td>25 mg twice daily</td>
<td>Migraine</td>
<td>Hypertension, cardiac arrhythmias, heart failure</td>
<td>Bradyarrhythmia, bronchospasm, hypoglycemia, sleep disturbance, depression, dizziness, fatigue</td>
</tr>
<tr>
<td>Propranolol (PO)(^a)</td>
<td>10 mg twice daily to 40 mg twice daily</td>
<td>10 mg twice daily</td>
<td>Migraine</td>
<td>Hypertension, cardiac arrhythmias, anxiety, essential tremor, postural orthostatic tachycardia syndrome</td>
<td></td>
</tr>
<tr>
<td>Timolol (PO)(^a)</td>
<td>5 mg/day to 15 mg twice daily</td>
<td>5 mg/d</td>
<td>Migraine</td>
<td>Hypertension, cardiac arrhythmias</td>
<td></td>
</tr>
<tr>
<td>Atenolol (PO)(^b)</td>
<td>25 mg/d to 100 mg/d</td>
<td>25 mg/d</td>
<td>Migraine</td>
<td>Hypertension, cardiac arrhythmias</td>
<td></td>
</tr>
<tr>
<td>Nadolol (PO)(^b)</td>
<td>20 mg/day to 240 mg/d</td>
<td>20 mg/d</td>
<td>Migraine</td>
<td>Angina, cardiac arrhythmias, hypertension</td>
<td>Drowsiness, insomnia, bradycardia, cardiac failure, hypotension</td>
</tr>
</tbody>
</table>

(Continued)
<table>
<thead>
<tr>
<th>Preventive medication</th>
<th>Dosing range</th>
<th>Initial dose</th>
<th>Use for migraine vs</th>
<th>Possible benefit for</th>
<th>Side effects</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>a</td>
<td>tension type headache</td>
<td>comorbidities</td>
<td></td>
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<tr>
<td><strong>Angiotensin-II receptor blocker</strong></td>
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<tr>
<td>Candesartan (PO)</td>
<td>4 mg/d to 16 mg/d</td>
<td>4 mg/d or 8 mg/d</td>
<td>Migraine</td>
<td>Hypertension, heart failure</td>
<td>Hypotension, renal function abnormality, dizziness, hyperkalemia, back pain, upper respiratory infection</td>
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<tr>
<td><strong>Angiotensin-converting enzyme inhibitor</strong></td>
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<tr>
<td>Lisinopril (PO)</td>
<td>10 mg/d to 20 mg/d</td>
<td>10 mg/d</td>
<td>Migraine</td>
<td>Acute coronary syndrome, hypertension, proteinuric chronic kidney disease</td>
<td>Acute kidney injury, angioedema, cough, hyperkalemia, hypotension, syncope</td>
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<tr>
<td><strong>Glutamate antagonist</strong></td>
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<tr>
<td>Memantine (PO)</td>
<td>5 mg/d to 10 mg twice daily</td>
<td>5 mg/d</td>
<td>Migraine Tension-type Posttraumatic</td>
<td>Dementia, neurocognitive toxicity of whole brain irradiation</td>
<td>Hypertension, hypotension, dizziness, headache, agitation, delusion, hallucination</td>
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<tr>
<td><strong>CGRP monoclonal antibodies</strong></td>
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<tr>
<td>Erenumab (SQ)</td>
<td>70 mg monthly or 140 mg monthly</td>
<td>70 mg monthly</td>
<td>Migraine</td>
<td>–</td>
<td>Hypertension, constipation, antibody formation, injection site reaction, hypersensitivity reaction</td>
</tr>
<tr>
<td>Fremanezumab (SQ)</td>
<td>225 mg monthly or 675 mg every 3 mo</td>
<td>225 mg monthly or 675 mg every 3 mo</td>
<td>Migraine</td>
<td>–</td>
<td>Antibody formation, injection site reaction</td>
</tr>
<tr>
<td>Galcanezumab (SQ)</td>
<td>240 mg first month, followed by 120 mg monthly</td>
<td>240 mg first month</td>
<td>Migraine</td>
<td>–</td>
<td></td>
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<tr>
<td>Eptinezumab (IV)</td>
<td>100 mg every 3 mo or 300 mg every 3 mo</td>
<td>100 mg every 3 mo or 300 mg every 3 mo</td>
<td>Migraine</td>
<td>–</td>
<td>Antibody formation, injection-site reaction, nausea, fatigue</td>
</tr>
<tr>
<td><strong>CGRP antagonists (gepants)</strong></td>
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<tr>
<td>Atogepant (PO)</td>
<td>10 mg/d, 30 mg/d, or 60 mg/d</td>
<td>10 mg/d or 30 mg/d</td>
<td>Migraine</td>
<td>–</td>
<td>Weight loss, constipation, drowsiness, fatigue, elevated liver enzymes</td>
</tr>
<tr>
<td>Rimegepant (ODT)</td>
<td>75 mg every other day</td>
<td>75 mg every other day</td>
<td>Migraine</td>
<td>–</td>
<td>Abdominal pain, nausea, rash, hypersensitivity reaction, dyspnea</td>
</tr>
<tr>
<td>OnabotulinumtoxinA (IM)</td>
<td>155 units distributed evenly in 31 sites, or up to 195 units</td>
<td>155 units distributed evenly in 31 sites, or up to 195 units</td>
<td>Chronic migraine only</td>
<td>–</td>
<td>Hypertension, infection, and/or pain at injection site, exacerbation of headache, facial weakness, myasthenia, neck pain, muscle pain or spasm, blepharoptosis, bronchitis</td>
</tr>
</tbody>
</table>
related headache phenotype and any other underlying primary or secondary headache disorder(s) should be considered.\(^{13,60}\) If a headache related to intracranial neoplasm and/or post-craniotomy headache has migraine features, acute treatments for migraine can be used. Some of the newest abortive medications developed for treatment of migraine include gepants (e.g., rimegepant, ubrogepant, and zavegepant). Antagonists of CGRP, gepants are not thought to cause MOH.\(^{50,81}\) Compared with triptans, gepants have additional benefits including lack of vasoconstrictive properties, fewer side effects, and are not associated with adverse outcomes in individuals with cardiovascular risk factors.\(^{82,83}\) Though gepants have not been studied specifically in patients with cerebral neoplasm or post-craniotomy headache, these medications may be considered in individuals with headache with migraine features. Lasmiditan, another newer acute migraine medication, similarly does not cause vasoconstriction; however, it has more potential side effects (e.g., dizziness, drowsiness, euphoria) and may contribute to MOH.\(^{80}\)

Non-medication approaches used for the acute treatment of migrainous headaches may also be considered as adjunctive treatment. Several safe and effective neuromodulation devices that stimulate structures involved in the pathophysiology of headache pain (e.g., the trigeminal nerve, vagus nerve, and/or occipital nerve) have been developed for migraine. Though these devices have not been studied specifically in headache related to cerebral neoplasm, they may be considered as acute treatment, particularly if the headache has features of migraine or, in some cases, of cluster headache. Furthermore, when implemented as an acute treatment strategy, neuromodulation has been found to decrease the need for acute analgesics.\(^{84,85}\) Hot packs and cold packs applied to the head also can help headache pain acutely.\(^{18}\)

**Preventive Treatment**

Though gradual resolution of post-craniotomy headache may occur over time without the need for significant medical treatment for most patients,\(^{18,63}\) Schankin et al found that 32% of individuals continue to experience moderate-severe headache pain for more than 6 months following craniotomy.\(^{86}\) Despite this, there is a striking paucity of randomized control trials and other literature surrounding the risks and benefits of preventive treatments for individuals with symptomatic post-craniotomy and/or cerebral neoplasm-related headache.\(^{60}\) Practically, headache providers may employ evidence-based therapies targeting the headache phenotype (e.g., migraine, tension-type), with the understanding that further trials are needed to understand the benefits and risks of these treatments in individuals with brain tumor.\(^{13,60,87}\)

The aims of preventive medication for headache include decreasing headache attack frequency, severity, and duration, as well as headache-related disability. Adequate preventive treatment additionally can improve responsiveness to acute medication treatment.\(^{88}\) Initiation of preventive medication for headache can be considered for individuals experiencing at least four headache days per month.\(^{88,89}\) Treatments for migraine-like and tension-like headaches in individuals with brain tumor may include conventional, evidence-based migraine therapies that also help with neuropathic pain, such as antidepressants, anticonvulsants, and others (\(\approx\) Table 5). Though the primary goal is to treat headaches effectively, a patient’s medical and psychiatric comorbidities should be considered when choosing a preventive medication.
The antidepressant medication category includes selective serotonin and norepinephrine reuptake inhibitors (SSRIs and SNRIs), as well as tricyclic antidepressants (TCAs). SNRIs, such as venlafaxine and duloxetine, are considered probably effective as treatment for migraine and tension-type headache, though they have not been studied specifically in individuals with cerebral neoplasm. SNRIs can help improve comorbidities of neuropathic pain, anxiety, and depression. Dosing for most preventive medications for headache should start low with gradual titration to higher doses to reduce the risk of side effects, which can lead to premature discontinuation. SNRIs may be associated with gastrointestinal side effects. TCAs, such as nortriptyline and amitriptyline, are effective treatments for migraine and tension-type headache. A common side effect of TCAs is reduction in sleep-onset latency, which may be used to the patient’s advantage if insomnia is a medical comorbidity.

Several anticonvulsant medications can be considered, as well. Gabapentin, an anticonvulsant with limited evidence for the treatment of migraine-type and tension-type headache, can be used for the treatment of neuropathic pain, anxiety, and focal-onset seizures. In a retrospective study, gabapentin was found to improve postoperative pain and decrease opioid use in post-craniotomy patients, and has limited evidence for the treatment of tension-type headache. Gabapentin’s wide range of dosing from 100 mg daily to 1,200 mg three times daily allows for convenient gradations in titration when weighing benefit versus risk of side effects (e.g., drowsiness or dizziness). Pregabalin, a gabapentinoid derivative, has similar anticonvulsant and anti-pain properties. In a meta-analysis, pregabalin was not shown to decrease post-surgical pain; however, perioperative pregabalin was associated with reduced opioid consumption after surgery. Topiramate is considered a highly effective treatment for the prevention of migraine and has limited evidence for the treatment of tension-type headache. Topiramate has been used for post-craniotomy pain, particularly when the pain has neuropathic features and/or there is trigeminal nerve involvement.

Memantine is a glutamate antagonist that has been used increasingly for the prevention of migraine and tension-type and posttraumatic headache. In one study, memantine was found to have a short window of titration (3 days) to full benefit, while also demonstrating a minimal side effect profile. Memantine additionally has been shown to help associated cognitive dysfunction in individuals receiving whole brain radiation.

The new CGRP monoclonal antibodies (e.g., erenumab, fremanezumab, galcanezumab, and eptinezumab) and gepants (e.g., atogepant, rimegepant) for the prevention of migraine headache have not been studied in individuals with post-craniotomy headache or headache related to brain tumor. These can be considered in individuals with migraine-phenotype headache. CGRP receptor antibodies also may have a neuroprotective role, and the possibility that CGRP antibodies could be used as an anti-tumor agent has been proposed. With this uncertainty, the decision to start an anti-CGRP medication in individuals with migraine and brain tumor should be considered a risk–benefit analysis, made on a case-by-case basis.

Other medical interventions, such as muscle relaxants (e.g., tizanidine) and nerve blocks (e.g., occipital nerve block, cervical trigger point injections), may be used in individuals with migraine and tension-type headache, and can be considered for symptomatic benefit in patients with headache related to cerebral neoplasm. OnabotulinumtoxinA injection, an effective treatment for chronic migraine, has been shown in several case series to improve post-craniotomy headache, and can be considered if the individual’s headache meets chronic migraine diagnostic criteria.

Non-pharmacologic treatments for migraine, tension-type headache, and cluster headache can be considered in a patient with the appropriate headache phenotype. Nutraceuticals, such as riboflavin, magnesium, coenzyme Q10, and feverfew, have variable evidence for the preventive treatment of migraine. Involved in cellular energy production, riboflavin is a well-tolerated and cost-effective supplement that has been shown in several randomized controlled trials to reduce migraine frequency, though other studies have failed to show benefit. Low magnesium levels associated with mitochondrial dysfunction have been identified in migraine. Two randomized control trials have suggested benefit from daily magnesium supplementation for migraine prevention, while other studies have shown no benefit. Coenzyme Q10 supplementation is thought to have protective and supportive roles for mitochondrial function, and is considered a possibly effective migraine treatment. Though butterbur has strong evidence in migraine prevention, it is not recommended due to risk of hepatotoxicity.

Several neuromodulation devices, as described above, have been approved for migraine and cluster headache prevention, and may be considered in patients with these headache phenotypes. Attention should be paid to whether the patient has history of seizure, as this may influence choice of neuromodulation device. In addition, physical therapy and massage therapy help with contributory neck tension. Behavioral interventions, such as biofeedback, cognitive behavioral therapy, and relaxation techniques, have grade A evidence as preventive treatment for migraine and can reduce headache-related outcomes by 35 to 55%, even in individuals without comorbid anxiety and depression. Lifestyle modifications, such as exercising frequently, maintaining healthy nutrition, managing obesity, and obtaining good quality sleep, have evidence in the treatment of migraine.

Comorbidities associated with headache exacerbation should be assessed. Individuals living with cerebral neoplasm commonly experience anxiety and depression, and further depressive symptoms may develop after brain surgery. In migraine, anxiety and depression have been associated with increased headache-related disability, and interventions targeting these psychiatric comorbidities may lead to improvement in disability. Other comorbidities, such as musculoskeletal conditions, temporomandibular disorder, chronic pain, and seizure, should be evaluated, as
some treatments may help both headache and these conditions.65

Conclusions

Though most of the general population will experience a headache at some time, cerebral neoplasm as the cause of headache is rare. Individuals presenting with progressive headache, focal neurological signs or symptoms, and those with other features of the SNNOOP10 criteria warrant consideration of additional workup for secondary etiology of headache.

Risk factors for the development of headache in individuals with cerebral neoplasm include a prior history of headache and a larger tumor burden with midline shift.24 Structure and function-related pain mechanisms (e.g., from compression of pain-sensitive structures and/or intracranial hypertension) may contribute to headache.

An individualized headache treatment approach should be developed using shared decision-making and consideration of the patient's goals of care. Patients should be monitored for tumor-related intracranial hypertension and treated appropriately with dexamethasone or neurosurgical intervention, if indicated. Though headache due to cerebral neoplasm often improves with neurosurgical intervention, many patients continue to experience chronic headaches after craniotomy and are at risk of undertreatment. At the same time, an underlying diagnosis of MOH should be identified early on, and treatment for this condition should be implemented so that optimal benefit of headache-related treatments can be achieved.

For individuals with headache related to brain tumor, a multimodal treatment approach incorporating pharmacologic and nonpharmacologic strategies should be used. Though a paucity of literature regarding acute and preventive treatments for headache related to cerebral neoplasm and post-craniotomy headache exists, symptomatic treatment may be based upon conventional therapies for the patient's headache phenotype, including acute and preventive medications, nutraceuticals, neuromodulation devices, behavioral interventions, anesthetic nerve blocks, and lifestyle changes. Additionally, an individual's headache-related comorbidities should be addressed, while understanding the patient's social determinants of health and prioritizing patient function and quality of life.

Conflict of Interest

None declared.

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