The differential diagnosis of chronic daily headaches: an algorithm-based approach

Abstract Chronic daily headaches (CDHs) refers to primary headaches that happen on at least 15 days per month, for 4 or more hours per day, for at least three consecutive months. The differential diagnosis of CDHs is challenging and should proceed in an orderly fashion. The approach begins with a search for "red flags" that suggest the possibility of a secondary headache. If secondary headaches that mimic CDHs are excluded, either on clinical grounds or through investigation, the next step is to classify the headaches based on the duration of attacks. If the attacks last less than 4 hours per day, a trigeminal autonomic cephalalgia (TAC) is likely. TACs include episodic and chronic cluster headache, episodic and chronic paroxysmal hemicrania, SUNCT, and hypnic headache. If the duration is ≥ 4 h, a CDH is likely and the differential diagnosis encompasses chronic migraine, chronic tension-type headache, new daily persistent headache and hemicrania continua. The clinical approach to diagnosing CDH is the scope of this review.

Keywords Chronic daily headache • Differential diagnosis • Strategy

Introduction

Chronic daily headaches (CDH) may be broadly or narrowly defined. In the broad sense, CDH encompasses all headaches that occur on 15 days per month or more. They are subdivided in primary CDHs (not caused by other disorders) and secondary CDHs (there is an underlying cause such as infection or tumour). The primary CDHs are further subdivided into those of long duration (≥ 4 h a day) and shorter duration (< 4 h duration) [1–3].

In its narrow and more frequently used sense, CDH is...
used to refer exclusively to “primary CDH of long duration”, a clinical syndrome defined by headaches that occur for ≥4 h a day on ≥15 days a month over >3 months in individuals without an underlying cause [1]. Unless otherwise indicated, in this paper, CDH refers to primary CDHs of long duration.

CDH is the third most common primary headache in the population, with an overall prevalence ranging from 4% to 5% [4, 5]. Because of their frequency, severity and duration, CDHs are extraordinarily burdensome to individuals and society. Furthermore, CDH is the most common reason for consultation in headache clinics.

The Silberstein and Lipton (S-L) criteria for CDH and its subtypes have been widely used and are well accepted [1]. According to these criteria, CDH is divided into four groups: chronic or transformed migraine (CM/TM), chronic tension-type headache (CTTH), new daily persistent headache (NDPH) and hemicrania continua (HC), and subclassified into subtypes with medication overuse or without medication overuse, incorporating the concept that medication overuse is often associated with the onset of CDH.

The differential diagnosis of CDHs is challenging and should proceed in an orderly fashion. Crucial elements include a thorough history, supplemented by general medical and neurological examinations, as well as laboratory testing and neuroimaging in selected patients. Herein we discuss the strategy to diagnose CDHs and their differential diagnosis. We emphasise that this strategy reflects the authors’ personal view of CDHs, as well as our personal diagnostic experience. Furthermore, the diagnostic algorithms presented herein have not been validated.

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**First step: distinguishing primary from secondary CDH**

An important first step in headache diagnosis is to distinguish primary from secondary CDH. The detailed strategies to identify or exclude secondary headaches are beyond the scope of this paper. In brief, the approach begins with a search for “red flags” that suggest the possibility of a secondary headache (Fig. 1). Once these features are identified, the physician must conduct the appropriate work-up to identify or exclude the conditions suggested by the red flag (Table 1). Once secondary headache is excluded by history, physical and neurological examination and/or appropriate testing, the next step is to diagnose the primary headache disorder [6].

Many patients with red flags do not have secondary headache disorders. Nonetheless, when red flags are present, they often merit investigation. Red flags include: (1) sudden onset of a severe headache; (2) a worsening pattern of a preexisting headache in the absence of obvious predisposing factors (e.g. medication overuse, depression); (3) headache with cancer, HIV or other systemic illness (fever, neck stiffness, cutaneous rash); (4) headache associated

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**Fig. 1** Algorithm for headache diagnosis. Modified from [25]
with focal neurologic signs other than typical visual or sensory aura; (5) headache in a patient with papilloedema; (6) moderate or severe headache triggered by cough, exertion, orgasm or Valsalva; and (7) new onset of a headache during pregnancy or post-partum.

Other less common secondary causes of intractable headache that may lead to CDH are summarised in Table 2. Detection of many of these disorders requires a systematic approach to the headache history and physical examination as well as a high index of suspicion. We will highlight some of these critical issues. For a review on this topic, the reader is referred to [7].

Red flags triggered by the temporal profile of headache

The temporal profile of the headache, and its nature and the circumstances of its onset are crucial for diagnosis. The temporal profile of a headache can provide important clues to its aetiology. A few examples are listed below:

- A rapid-onset headache may suggest a subarachnoid haemorrhage, pituitary apoplexy [8] or other intracranial catastrophes [9]. These disorders rarely cause CDH, but can result in intractability of an ongoing headache attack.

### Table 1 Red flags in the diagnosis of headache. Modified from [6]

| Red flag                             | Consider                                                                 | Possible investigation(s)                                      |
|--------------------------------------|--------------------------------------------------------------------------|----------------------------------------------------------------|
| Sudden-onset headache                | Subarachnoid haemorrhage, bleed into a mass or AVM, mass lesion (especially posterior fossa) | Neuroimaging, lumbar puncture (after neuroimaging evaluation)   |
| Worsening-pattern headache           | Mass lesion, subdural haematoma, medication overuse                      | Neuroimaging                                                   |
| Headache with cancer, HIV or other systemic illness (fever, neck stiffness, cutaneous rash) | Meningitis, encephalitis, Lyme disease, systemic infection, collagen vascular disease, arteritis | Neuroimaging, lumbar puncture, biopsy, blood tests              |
| Focal neurologic signs, or symptoms other than typical visual or sensory aura Papilloedema | Mass lesion, AVM, collagen vascular disease or Mass lesion, pseudotumour, encephalitis, meningitis | Neuroimaging, collagen vascular evaluation or Neuroimaging, lumbar puncture (after neuroimaging evaluation) |
| Triggered by cough, exertion or Valsalva | Subarachnoid haemorrhage, mass lesion | Neuroimaging, consider lumbar puncture |
| Headache during pregnancy or post-partum | Cortical vein/cranial sinus thrombosis, carotid dissection, pituitary apoplexy | Neuroimaging |

### Table 2 Secondary headaches that mimic chronic benign headache syndromes

- Headache associated with vascular disorders
- Cerebrovascular disease including carotid artery dissection and arteriovenous malformation
- Arteritis including giant cell arteritis
- Headache associated with non-vascular intracranial disorders
- Low CSF pressure syndrome (spontaneous or post-traumatic CSF “leak”)
- High CSF pressure without papilloedema
- Intracranial: Lyme disease, human immunodeficiency virus, encephalitis, fungal meningitis, etc.
- Headache associated with substances or their withdrawal
- Overuse of acute headache medications (rebound or toxic drug overuse syndromes)
- Headache associated with cranium, neck, eyes, ears, nose, sinuses, teeth, mouth or other facial or cranial structures
- Otolaryngologic disease, including chronic sphenoid sinusitis (or other sinus disease)
- Nasopharyngeal disorders, including carcinoma
- Disorders of the trigeminal nerve, including dental and oral disease, jaw pathology
- Subacute angle closure glaucoma, optic neuritis and other ocular disorders
- Occipitocervical disease, including Arnold-Chiari Malformation Type I; upper cervical joint, root or nerve (neuralgic) syndromes
- Headache associated with non-cerebral infection, metabolic or systemic disturbances
- Hepatitis, renal disease, B12 deficiency, anaemia, exposure to carbon monoxide and other toxins
- Hormonal disturbances/endocrinologic disease (oestrogen, thyroid disease, hyperprolactinaemia, etc.)
- Vasculitis/rheumatic/connective tissue disorders
- Miscellaneous
- Mediastinal and thoracic processes including angina, mass lesions, superior vena cava syndrome
• Sphenoid sinusitis may cause NDPH and may be missed radiologically unless appropriate studies are performed [10].
• Headaches that start after age 55 suggest an organic disorder for CDH, such as a mass lesion or giant cell arteritis [11]. Giant cell arteritis is often under-diagnosed, and is an important cause of preventable blindness in the elderly.

Red flags triggered by concurrent events or provoking activity

Concurrent events and headache triggers may give clues to diagnosis, and some of them are listed below:
• Chronic headaches (NDPH) that occur in the peripartum period may be due to dural sinus thrombosis [12].
• Orthostatic CDH suggests a low CSF pressure headache (from a spontaneous CSF leak, a previous lumbar puncture, or an epidural block) [13].
• CDH exacerbated by straining, coughing or sneezing suggests a hindbrain malformation, occipitocervical junction disorder or increased intracranial pressure [14].
• CDH that is worse in the morning in the absence of medication overuse, that may suggest raised intracranial pressure. CDH with medication overuse is often worse in the morning.
• In contrast, subjects with CDH that is better in the morning and worse at night that may suggest low CSF volume headache [13].
• Similarly, CDHs that are worse on awakening sometimes occur with obstructive sleep apnoea [13].
• Increasing pain while upright, and neck pain and headache provoked by flexion or extension of the neck might suggest an occipitocervical junction disorder or a systemic medical illness [15–17].
• Search for opportunistic infections, including toxoplasmosis and cryptococcal meningitis, in patients who may have HIV infection or HIV risk factors [18].
• A previous mild to moderate head or neck injury is frequently overlooked in the standard history, despite the fact that it can render an individual refractory to standard headache treatments [19]. A diagnosis of post-traumatic headache may lead to additional treatment including trigger point injections and facet joint and cervical nerve blocks.
• The history often neglects recent dental procedures. Root canals, tooth extractions or bite disturbances may provoke or be associated with the development of CDH due to brain abscess [20, 21].
• Consider ocular disturbances, such as subacute angle closure glaucoma, or infection, if the CDH is worse in the periorbital region [22].
• Nasal blockage, drainage, pus or pressure may suggest sinus disease [23]. Nasopharyngeal carcinoma can produce chronic head and face pain and can be identified by detailed, expert examination of the nasopharynx or neuroimaging [24].

In the absence of secondary CDH, the clinician proceeds to diagnose a primary CDH disorder. If the headache is atypical or difficult to classify, the possibility of secondary headache should be reconsidered, although the modifying effect of any treatment being taken should be kept in mind. Additionally, lack of response to appropriate treatment given for enough time may also be considered a red flag.

Second step: classifying primary CDH based on duration and frequency (Fig. 2)

Basically, in a patient where we excluded, either by clini-
The chronic daily headaches of shorter duration (Fig. 3)

As discussed previously, headaches that happen on more than 15 days per month, for less than 4 hours per day, are usually not referred to as CDHs. They belong to a group called the trigeminal autonomic cephalalgias (TACs) [26]. The TACs include the episodic and chronic cluster headache, episodic and chronic paroxysmal hemicrania, short unilateral neuralgiform headache with conjunctival injection and tearing (SUNCT) syndrome, and hypnic headache. The TACs are characterised by unilateral pain in the trigeminal distribution, with ipsilateral autonomic features. The most common disorder in this family is cluster headache (CH). The pain of CH is described variously as sharp, boring, drilling, knife-like, piercing or stabbing, in contrast to the pulsating pain of migraine. It usually peaks in 10–15 min but remains excruciatingly intense for an average of 1 h within a duration range of 15–180 min. During this pain, patients find it difficult to lie still, exhibiting often marked agitation and restlessness, and autonomic features are usually obvious. After an attack, the patient remains exhausted for some time [26, 27].

Like CH, the paroxysmal hemicranias are characterised by unilateral attacks of trigeminal pain and autonomic features. In contrast with CH, the paroxysmal hemicranias have three main features: (1) more frequent (more than five per day); (2) short duration (2–30 min); and (3) absolute response to therapeutic doses of indomethacin. They are rare [28, 29].

The third TAC, SUNCT, is a very rare primary headache. The diagnostic criteria require at least 20 high-frequency attacks (3–200 per day) of unilateral orbital, supraorbital or temporal stabbing or pulsating pain, lasting 5–240 s and accompanied by ipsilateral conjunctival injection and lacrimation. The attacks are characteristically dramatic, with moderately severe pain peaking in intensity within 3 s and prominent tearing [30].

Hypnic headache is a primary headache disorder of the elderly, characterised by short-lived attacks (typically 30 min) of nocturnal head pain that awakens the patient at a constant time each night, in many cases on more nights.

Fig. 3 Short duration headaches
than not. It does not occur outside sleep. Hypnic headache is usually bilateral (though unilaterality does not exclude the diagnosis) and usually mild to moderate, very different from the unilateral orbital or periorbital knife-like intense pain of cluster headache. Autonomic features are absent [31, 32].

Chronic daily headaches of long duration (Fig. 4)

There are four CDHs of long duration, which are discussed here. According to the S-L criteria, all subtypes are subdivided into with or without medication overuse. According to the Second Edition of the International Classification of the Headache Disorders (ICHD-2), a CDH with medication overuse should be classified as medication overuse headache (MOH) regardless of the phenotype. Therefore, in the following section we first describe the four subtypes of the CDH and then separately describe MOH.

**Chronic or transformed migraine**

The terms chronic migraine (CM) and transformed migraine (TM) are frequently used interchangeably, although since the publication of ICHD-2 this is no longer appropriate [33]. Although the two are related, they are different clinical entities with specific definitions.

Patients with TM/CM typically have a past history of migraine. It is more frequent in women with a past history of migraine without aura. Subjects usually report a process of transformation over months or years, and as headache increases in frequency, associated symptoms become less severe and frequent. The process of transformation frequently ends in a pattern of daily or nearly daily headache that resembles chronic tension-type headache, with some attacks of full-migraine superimposed [1–3, 34, 35]. In the clinical setting, migraine transformation most often is related to acute medication overuse, but transformation may occur without overuse. In the more general population, most cases of TM are not related to medication overuse [36]. Multiple risk factors may be involved in these cases.

The S-L criteria identify three potential links to migraine in TM: (A) a prior history of ICHD defined migraine; (B) a period of escalating headache frequency; and (C) concurrent superimposed attacks of migraine that fulfil the IHS criteria. TM is subdivided into with or without medication overuse. If frequent migraine develops *de novo* in a previously headache-free subject, the diagnosis is new daily persistent headache according to S-L criteria.

The ICHD-2 criteria for CM initially required 15 or more days of migraine per month [33], but these criteria were shown to be too restrictive for clinical practice and research.

![Fig. 4 CDHs of long duration](image-url)
As a consequence, the ICHD-2 criteria for CM (ICHD-2R) were recently revised in the Appendix of the classification (Table 3) [42]. CM-R is now defined by CDH plus 8 or more days of migraine as evidenced by headaches that meet ICHD-2 criteria for migraine without aura or headache with response to acute treatment with a migraine-specific medication (triptan or ergotamine compound). As previously stated, if subjects are overusing acute medication they should be classified as having MOH [42, 44].

A recent study compared the criteria for CM-R and TM in a headache specialty clinic. Of 158 patients with TM, just 5.6% met the old ICHD-2 criteria for CM. According to the ICHD-2R, a total of 92.4% met the revised criteria for CM ($p<0.001$ vs. ICHD-2), which represents a significant improvement. Subjects with TM and medication overuse should be classified as MOH (Table 4), and not CM+, according to the ICHD-2R. Nonetheless, the same study assessed the proportion of them who had ≥8 days of migraine per month. Of the 399 individuals with TM+, just 10.2% could be classified as CM+ in the ICHD-2. However, most (349, 86.9%) had ≥8 days of migraine per month and could be classified as MOH and probable CM in the ICHD-2R ($p<0.001$ vs. ICHD-2). The study concluded that the new criteria for CM address most of the criticism towards the ICHD-2 and should be adopted in clinical practice and research. In the population where use of specific acute migraine medications is less common, the agreement between ICHD-2R CM and TM may be less robust [43].

**CM/TM in adolescents and adults**

Differences exist regarding the clinical presentation of TM in adolescents and adults. Most adults with TM have fewer than 15 days of full blown migraine per month and more days of headache resembling tension-type headache than of migraine. In contrast, TM in adolescents is replete with migraine attacks. Medication overuse is more common in adults (84.0%) than in adolescents (58.9%) [45].

**Phenotype of CM/TM changes over time**

In a recent study of 402 subjects with TM, the proportion of migraine attacks decreased with age (with a proportional increase of tension-type headache attacks), from 71% below the age of 30 years to 22% at age 60 or above. It was higher in those with shorter interval from the onset of migraine to the onset of CDH (less than 5 years, $p=0.003$), and in those with a more recent onset of CDH (less than 6 years, $p<0.0001$). These findings suggest that CM (15 or more days of migraine per month) is the first stage of migraine chronicity in most patients. Subsequently, the frequency of clearcut migraine attacks diminishes. Thus, CM may be viewed as the earlier stage of TM, and both are different evolutive stages of a chronic migrainous process [46].

The concept that early in the process of transformation

![Table 3 Diagnostic criteria for primary chronic daily headaches according to the International Classification of Headache Disorders (2006) Revised Criteria and the Silberstein-Lipton Criteria](image)

| Silberstein-Lipton From 1996 | ICHD-2R |
|-----------------------------|---------|
| **Transformed migraine**    | Chronic migraine |
| A. Daily or almost daily (>15 days a month) head pain for >1 month | **Diagnostic criteria:** |
| B. Average headache duration of >4 h per day (if untreated) | Headache ≥15 days/month for >3 months. |
| C. At least one of the following: | Occurring in a patient with at least 5 prior migraine attacks. |
| 1. History of episodic migraine meeting any IHS criteria 1.1–1.6 | On ≥8 days per month, for at least three months, headache fills |
| 2. History of increasing headache frequency with decreasing | criteria C1 and C2 |
| severity of migraineous features over at least 3 months | |
| 3. Headache at some time meets IHS criteria for migraine | |
| 1.1–1.6 other than duration | 1. **Unilateral** |
| D. Does not meet criteria for new daily persistent headache (4.7) | **Throbbing** |
| or hemicrania continua (4.8) | **Moderate or severe** |
|                               | **Aggravated by physical activity** |
|                               | **Nausea and/or vomiting** |
|                               | **Photophobia and phonophobia** |
|                               | 2. Treated or relieved with triptans or ergotamine compounds. |
|                               | D. No medication overuse and not attributable to another causative disorder. |

*HIS, International Headache Society*
most headache days fill criteria for migraine, and as disease evolves, the headache attacks become less typical, has also been supported by an adolescent study, where those with recent-onset CDH were much more likely to have 15 or more migraine days per month (74.5% vs. 25.8%, \(p<0.001\)) [47].

Because the phenotype of CM/TM changes over time, CM is frequently misdiagnosed as CTTH.

*Chronic tension-type headache*

CTTH represents half of the CDH cases found in population studies, but just a small fraction of those found in specialty clinics [48]. It is the only CDH that the ICHD-I considered as a single diagnosis. The criteria remained little changed in the ICHD-II. Despite the fact that ICHD-I and ICHD-II criteria distinguish between episodic and CTTH, the latter headache type remains surprisingly poorly studied. This can be partially explained by the confusion between CTTH and TM with a low frequency of superimposed migraine attacks.

The prevalence of CTTH is markedly lower than that of episodic tension-type headache (ETTH), the most prevalent primary headache disorder. Its 1-year prevalence estimates range from 1.7% to 2.2% [48, 49]. The female preponderance of CTTH is greater than that of ETTH but lower than that of migraine. In the USA, the prevalence of CTTH was reported to be 2.8 in women and 1.4 in men, with an overall gender prevalence ratio of 2.0. The prevalence of CTTH increases with age [50].

The distinguishing pain features of CTTH are bilateral location, nonpulsating quality, mild to moderate intensity and lack of aggravation by routine physical activity. The pain is unaccompanied by nausea, although just one of photophobia or phonophobia does not exclude the diagnosis. In other words, CTTH is defined by *what it is not* (i.e., migraine) [51].

The S-L criteria reserve the diagnosis of CTTH if the CDH develops abruptly (*de novo*). It is inferred that this is also true in the ICHD-II criteria, although this feature of evolution is described in the notes but is not part of the formal criteria.

**New daily persistent headache**

CDH may begin without a history of evolution from episodic headache. NDPH is characterised by the relatively abrupt onset of an unrelenting primary CDH; i.e., a patient without a previous headache syndrome develops a CDH that does not remit. The patient’s remembering the circumstances, place and date that the headache began is a pathognomonic feature. It is the new onset of this primary daily headache that is the most important feature [52]. The clinical features of the pain are not considered in making the diagnosis, which simply requires absence of history of evolution from migraine or ETTH [1]. The ICHD-II addresses NDPH as a single diagnosis in those with a new-onset CDH that resembles CTTH. A new-onset CDH with migrainous features cannot be classified as NDPH according to the ICHD-II criteria whereas the S-L classification allows this diagnosis in patients with headache features of migraine or ETTH if the disorder arises abruptly.

**Table 4 Classification of the medication overuse subtypes according to the ICHD-2**

| ICHD-2 code | Diagnosis                                      | Amount of medication                                          |
|-------------|------------------------------------------------|--------------------------------------------------------------|
| 8.1         | Ergotamine overuse headache                    | Ergotamine intake on ≥10 days per month on a regular basis for >3 months |
| 8.2         | Triptan overuse headache                       | Triptan intake (any formulation) on ≥10 days per month on a regular basis for >3 months |
| 8.3         | Analgesic overuse headache                     | Intake of simple analgesics on ≥15 days per month on a regular basis for >3 months |
| 8.4         | Opioid overuse headache                        | Opioid intake on ≥10 days per month on a regular basis for >3 months |
| 8.5         | Combination analgesic overuse headache         | Intake of combination analgesic medications on ≥10 days per month on a regular basis for >3 months |
| 8.6         | MOH attributed to combination of acute medications | Intake of any combination of ergotamine, triptans, analgesics, and/or opioids on ≥10 days per month on a regular basis for >3 months without overuse of any single class |
| 8.7         | Headache attributed to other medication overuse | Regular overuse for >3 months of a medication other than those described above |
| 8.8         | Probable MOH                                   | Overused medication has not yet been withdrawn or medication overuse has ceased within the last 2 months but headache has not so far resolved or has reverted to its previous pattern |
Hemicrania continua

HC is probably the most frequently unrecognised primary headache. HC is a chronic, unilateral pain, commonly mistaken for TM [53, 54]. Both disorders are characterised by chronic unilateral pain with superimposed painful exacerbations. In HC, the painful exacerbations are often associated with ipsilateral autonomic features such as conjunctival injection, lacrimation and ptosis. In TM, exacerbations are more typically accompanied by nausea, photophobia and phonophobia. In addition, patients with HC usually do not have antecedent history of episodic migraine. In TM, attacks increase in frequency over time. If the headaches are longstanding, the patient may not remember how they began. Though pain fluctuates in HC, it does not usually have the morning and end-of-dosing-interval pattern of exacerbations typical of TM. It is advisable to offer to patients with unilateral CDH a therapeutic trial with indomethacin prior to other intervention (doses of up to 225 mg/day for three to four days).

Medication overuse headache

The ICHD-II classifies MOH under Chapter 8 (headache attributed to a substance or its withdrawal), under the topic 8.2 (MOH). MOH is subdivided into seven groups, plus an additional topic for probable MOH [42]. The diagnostic criteria and the critical amount of consumption of medication are displayed in Table 4. According to the ICHD-II, a diagnosis of MOH is basically established when two situations are fulfilled: 1. The consumption of acute medication is beyond a critical dose; 2. There is a CDH.

Although the traditional criteria required demonstration of improvement after detoxification, the revised criteria for MOH do not. It just requires a CDH and overuse of medication. Furthermore, the ICHD-II does not suggest that subsidiary investigation is required in all cases but requires that, at least clinically, the provider exclude secondary disorders other than medication overuse.

Conclusions

The differential diagnosis of CDH is challenging and requires a systematic approach. Herein we presented an algorithm-based approach to the differential diagnosis of the CDHs, which should help physicians interested in headache to move forward quickly and safely when assessing patients with daily or almost daily headaches.

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