

REVIEW ARTICLE

Headache: what to ask, how to examine, and what scales to use. Recommendations of the Spanish society of neurology's headache study group[☆]



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Abstract

Introduction: Headache is the most common neurological complaint at the different levels of the healthcare system, and clinical history and physical examination are essential in the diagnosis and treatment of these patients. With the objective of unifying the care given to patients with headache, the Spanish Society of Neurology's Headache Study Group (GECSN) has decided to establish a series of consensus recommendations to improve and guarantee adequate care in primary care, emergency services, and neurology departments.

Methods: With the aim of creating a practical document, the recommendations follow the dynamics of a medical consultation: clinical history, physical examination, and scales quantifying headache impact and disability. In addition, we provide recommendations for follow-up and managing patients' expectations of the treatment.

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PALABRAS CLAVE

Cefalea;
Anamnesis;
Exploración;
Escalas;
Educación;
Calendario;
Diario electrónico

Conclusions: With this tool, we aim to improve the care given to patients with headache in order to guarantee adequate, homogeneous care across Spain.

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¿Qué preguntar, cómo explorar y qué escalas usar en el paciente con cefalea? Recomendaciones del Grupo de Estudio de Cefalea de la Sociedad Española de Neurología

Resumen

Introducción: La cefalea es el motivo de consulta neurológico más prevalente en los distintos niveles asistenciales, donde la anamnesis y exploración son primordiales para realizar un diagnóstico y tratamiento adecuados. Con la intención de unificar la atención de esta patología, el Grupo de Estudio de Cefalea de la Sociedad Española de Neurología (GECSEN) ha decidido elaborar unas recomendaciones consensuadas para mejorar y garantizar una adecuada asistencia en Atención Primaria, Urgencias y Neurología.

Metodología: El documento es práctico, sigue el orden de la dinámica de actuación durante una consulta: anamnesis, escalas que cuantifican el impacto y la discapacidad y exploración. Además, finaliza con pautas para realizar un seguimiento adecuado y un manejo de las expectativas del paciente con el tratamiento pautado.

Conclusiones: Esperamos ofrecer una herramienta que mejore la atención al paciente con cefalea para garantizar una asistencia adecuada y homogénea a nivel nacional.

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Introduction

Headache is one of the most frequent reasons for neurological consultation. Therefore, it is essential for physicians attending these patients to be proficient in its diagnosis and treatment.

Proper history-taking and examination are fundamental,^{1,2} given that correct diagnosis and appropriate treatment may be established from the first consultation, without needing to perform any complementary examination.

With a view to ensuring the proper medical care of patients with headache, unifying clinical practice, and guaranteeing health-care equality for our patients, the Spanish Society of Neurology's Headache Study Group (GECSEN) has established the following consensus recommendations, which are intended to facilitate systematic history-taking, examination, assessment, and follow-up of patients with headache.

A table is included to facilitate their use in the very different contexts in which patients with headache are attended: primary care, emergency departments, and neurology consultations. As with other publications, the study group's Board invited young neurologists with experience in headache management to draft this consensus document, basing the recommendations on clinical experience and a review of the recent literature.

Medical history

Medical history should address the reason for consultation and the current disease (focusing on pain characteristics).³

Reason for consultation

The reason for consultation includes age, clinical course of the headache (focusing on onset and progression), and

history of headache. Emergency departments should also record the reason for seeking emergency consultation.³

Current disease**History-taking targeting pain characteristics**

Initially, patients must be allowed to spontaneously describe the characteristics of their headache. This usually only takes 1-2 minutes.^{4,5} The next stage of history-taking should comprise a structured, targeted interview (Table 1) addressing the following aspects.⁶

Temporal characteristics

It is crucial to know whether onset was sudden or insidious and, through analysing the progression of headache, whether the process is acute (< 72 hours), subacute (72 hours to 3 months), or chronic (> 3 months). We must evaluate the duration of the attack and whether pain is continuous and/or paroxysmal; in the latter case, we must establish the approximate duration. Disease progression is one of the most helpful characteristics in differential diagnosis. Pain frequency also guides assessment of the patient's level of disability and the type of treatment to be indicated. Finally, some headaches, such as cluster headache, show a clear circadian pattern.

Pain quality

Pulsating (throbbing)/pressing (tightening)/stabbing (puncture-like), sharp (perforating)/electric-shock-like.

Table 1 Key points to be addressed in medical history interviews targeting pain characteristics.**Temporal characteristics**

Age at onset
 Pain onset: sudden, gradual; time to peak
 Frequency: daily, weekly, monthly; alternating attacks and remission
 Duration: seconds, minutes, hours, days, weeks, months, years
 Timeframe: morning, evening, night; fixed time; circadian rhythm; seasonal
 Remission: sudden, progressive, persisting from onset

Quality

Pulsating, sharp, burning, lancinating, pressing

Intensity

Mild, moderate, intense
 Incompatible with daily life
 Nocturnal awakenings

Location

Focal, hemicranial, holocranial
 Frontal, occipital, orbital, vertex pain; sensation of a band around the head
 Changing location during progression

Triggering or exacerbating factors

Stress
 Sleep pattern
 Physical exercise
 Hormonal factors (menstruation, contraception)
 Diet, alcohol
 Atmospheric changes
 Valsalva manoeuvres (cough, intercourse)
 Postural changes
 Head movements
 Pressure on a trigger point

Attenuating factors

Medication, rest/sleep, decubitus positions

Accompanying symptoms and time of onset (before/simultaneous with/after pain)

Nausea/vomiting
 Phono-/photo-/osmophobia
 Photopsia, scotoma, hemianopsia, diplopia
 Hemiparesis/hemidysaesthesia
 Instability, vertigo
 Aphasia
 Dysautonomic symptoms
 Confusion/seizures/fever
 Constitutional symptoms/jaw claudication

Intensity

Numeric pain rating scale⁷: 0 (no pain) to 10 (worst possible pain). Scores of 1-3 are considered mild, 4-6 moderate, and 7-10 severe.

As pain and pain intensity are subjective, it is important to assess progression with intra-subject measures. It should also be noted that, with the exception of thunderclap headache, pain intensity usually does not correlate with the severity of the underlying cause.⁸

Location

- Circumscribed/unilateral/holocranial/nerve territory
- Radiating

Patients should be asked to precisely indicate the site of pain onset and its potential radiation to nearby areas, as well as variation or migration to other areas of the face, head, or neck. For example, pain that appears and remains on one side of the head, never crossing the midline, may be considered differently to pain initiating in the same way but that changes sides or extends to the whole head; pain circumscribed to a circular or elliptical area or coinciding with the territory of a specific nerve should also be considered differently.⁶

Triggering or exacerbating factors

Cluster headache may be triggered by alcohol or vasodilator drugs. Valsalva manoeuvres trigger or exacerbate secondary headaches, as well as certain primary headaches, such as those associated with cough, physical exercise, or intercourse. Hypnic headache exclusively appears during sleep. Low cerebrospinal fluid pressure headache typically occurs when standing, and improves with decubitus positions, unlike headache due to intracranial hypertension. Head movement exacerbates pain in such headaches as migraine and cervicogenic headache. Palpation of the emergence of the terminal branches may trigger attacks of neuralgia in the corresponding areas.

Attenuating factors

Attenuating factors are those that relieve pain, such as the type of treatment, treatment response, or head position; these may also guide us in identifying the type of headache. For example, response to indometacin is indicative of hemicrania continua, and decubitus positions typically alleviate headache secondary to intracranial hypotension.

Accompanying symptoms

Phono-/photophobia and nausea are common symptoms during migraine attacks, although they may also appear in secondary headaches. Osmophobia and gustatory alterations may be even more specific to this type of headache.⁹ Diagnostic certainty is greater in patients presenting transient focal symptoms (scotoma; fortification spectrum; progressive, self-limited sensory or language syndromes) compatible with typical migraine aura or such premonitory symptoms as asthenia, neck stiffness, hyperphagia, or yawning.^{10,11} We should also enquire about oculofacial

Pain is usually perceived as pulsating in migraine, pressing in tension-type headache, sharp in cluster headache, burning and continuous in epicranial neuralgias, and electric-shock—like in neuralgias involving major trunks and in short-lasting unilateral neuralgiform headache with conjunctival injection and tearing (SUNCT)/short-lasting unilateral neuralgiform headache with autonomic symptoms (SUNA). However, pain quality is not decisive in diagnosis, and careful attention must be paid to the remaining pain characteristics. In many cases, patients do not apply any of these descriptors to their pain.⁶

Table 2 Warning signs and symptoms of secondary headache.

	Type of warning sign/symptom
Pain characteristics	Sudden or explosive onset, with or without associated effort or Valsalva manoeuvre Recent onset with progressive increase in intensity or frequency Recent, considerable change in characteristics of previous headache Progressive increase in intensity/frequency and loss of response to previously effective analgesics Clear, consistent worsening with postural changes Nocturnal awakenings due to headache Strictly unilateral location
Clinical characteristics	Age of onset > 50 years, especially if associated with rheumatic symptoms, jaw claudication, or constitutional syndrome Presence of neoplasia and/or immunosuppression Increased risk of haemorrhage (eg, in patients receiving anticoagulants) Association with fever without focus, especially in patients with meningeal syndrome Vomiting not attributable to primary headache (migraine), or projectile vomiting Focal neurological signs Headache associated with seizures Visual alterations suggestive of papilloedema

Table 3 Signs and symptoms that may be associated with both primary and secondary headaches.

Sign/symptom	Primary headache	Secondary headache
Pulsating pain, worsening with Valsalva manoeuvres, nausea and/or vomiting	Migraine	Meningitis Hypertensive crisis/PRES Cerebral venous sinus thrombosis
Atypical visual aura	Migraine with aura	Stroke HaNDL syndrome
Ptosis, miosis	Trigeminal autonomic cephalalgia	Carotid artery dissection Cavernous sinus syndromes
Orbital pain, red eye, pupillary alterations	Trigeminal autonomic cephalalgia	Angle-closure glaucoma Uveitis Cavernous sinus syndromes
Holocranial, pressing pain of mild to moderate intensity	Tension-type headache	Highly non-specific. Multiple processes
Pain in temporal region	Myofascial syndrome	Temporal arteritis TMJ disorders
Exclusively nocturnal awakenings due to headache	Hypnic headache	Nocturnal hypertension

HaNDL: transient headache and neurological deficits with cerebrospinal fluid lymphocytosis; PRES: posterior reversible encephalopathy syndrome; TMJ: temporomandibular joint.

Adapted from Irimia.³

autonomic signs and symptoms ipsilateral to pain: in the correct clinical context, this guides the diagnosis of trigeminal autonomic cephalalgias, although these symptoms have also been described in association with migraine-type headache. Secondary headaches may also be accompanied by focal neurological signs, altered level of consciousness, fever, or seizures. Intracranial hypertension may manifest with tinnitus or temporary darkening of vision. Giant cell arteritis may be associated with constitutional and rheumatological symptoms or jaw claudication.

Secondary headache should initially be ruled out based on the signs and/or symptoms of pain itself, as well as the clinical characteristics associated with headache (Table 2).

However, certain signs and symptoms can be associated with both primary and secondary headache (Table 3).

History-taking targeting patient characteristics

This part of the medical history interview should address the following areas (Table 4):

Personal history

It is important to gather information about the patient's clinical history, as some health conditions, such as sleep

Table 4 Key points to be addressed in targeted medical history interviews with patients with headache.

<i>Personal history</i>
Medication allergies
Vascular risk factors
Drug habits
Craniofacial disorders (TMJ disorders, etc)
Heart disease/asthma/nephrolithiasis/glaucoma
Mood alterations
Sleep alterations (drowsiness, insomnia, bruxism, SAHS)
Head trauma
Phase in the menstrual cycle, oral contraceptives
<i>Family history</i>
<i>Social factors</i>
Type of work/shifts
Number of children and socioeconomic situation
<i>Factors affecting headache transformation</i>
Medication overuse
Body mass index
Mood alterations
Sleep cycle alterations
Caffeine consumption
Chronic stress
Sedentary lifestyle
<i>Medication</i>
Current or previous treatments
Maximum dose reached/duration
Effectiveness/adverse reactions

SAHS: sleep apnoea-hypopnoea syndrome; TMJ: temporomandibular joint.

apnoea-hypopnoea syndrome, may play a role in headache. The patient's history may also influence the treatments we can use, as some drugs may exacerbate underlying conditions (for instance, flunarizine can exacerbate depression and may cause weight gain). In other cases, we may be able to select a treatment with a dual effect; for example, in hypertensive patients, we may use beta-blockers, lisinopril, or candesartan to treat headache.

Family history of headache

We should ask about family history of headaches resembling the patient's, particularly among close relatives. Migraine is an example of a primary headache that can affect several members of the same family, with heritability estimated at 40% to 60% (higher levels in migraine with aura); therefore, family members of patients with migraine present higher risk than the general population.¹² Tension-type headache, and especially chronic forms, also tends to aggregate in families.¹³ Many patients with cluster headache report family history of headache, including migraine, with increased risk among first-degree relatives.¹⁴

Social factors

We must determine whether any current or previous conflict or stressful situation may be triggering headache, particularly chronic headache. Personal, family, social, or

work-related problems can occasionally play a role in the form of presentation of headache.⁶ Pain may also cause considerable disability; we should therefore enquire about this possibility and its potential impact on patients' mood and cognitive performance.

Factors affecting headache transformation

Physicians should enquire about factors potentially leading to changes in pain frequency, and even chronic headache.^{15,16} The most frequent risk factors are listed in Table 4.

Medication

It is helpful to determine what drugs the patient has used to control headache, and which were the most and the least effective (it is also important to record the maximum dose reached and the actual duration of treatment). We should also consider whether headache may be caused by some drug the patient is using. Drugs liable to worsen the clinical course of migraine include those used for hormone replacement therapy and oral contraception, vitamin A, and retinoic acid derivatives. Both migraine and cluster headache may be exacerbated by the use of such vasodilators as nitrites, minoxidil, and nifedipine. Chronic use of medications including opiates, barbiturates, caffeine, ergotamines, or a combination of these may cause rebound or withdrawal headache.¹⁷ It is also important to enquire about the use of oral anticoagulants, given the risk of secondary headaches or with a view to performing procedures during the consultation.

Recommended scales

The instruments we recommend are self-administered questionnaires with validated Spanish-language versions that may be used for adult patients and for both clinical and research purposes. They aim to quantify disability, headache impact, comorbidities, and other variables of interest, as well as symptom improvement and worsening.

Disability assessment

The Migraine Disability Assessment (MIDAS) questionnaire¹⁸ is broadly validated and has been translated into multiple languages. It comprises 5 items evaluating the impact of headache on work/school performance, domestic work, and social activities over a 3-month period. The total score indicates the patient's grade of disability, from I to IV. Grade IV was recently subdivided into grades IVa and IVb in order to better characterise the disability associated with chronic migraine.

Another tool for measuring the functional impact of headache is the 6-item Headache Impact Test (HIT-6).¹⁹ The scale assesses the frequency of severe pain, limitations in daily activities (including work, school, and social activities), desire to lie down, fatigue, irritability, and difficulty concentrating. It is also useful for assessing treatment

Table 5 Characteristics of the main self-administered scales for assessing patients with headache.

	Use	Disease	No. items	Administration time
<i>Disability</i>				
MIDAS	Workplace, home, social	Migraine	5 (+2)	< 5 min
HIT-6	Everyday activities	Migraine	6	< 5 min
WPAI	Everyday and workplace activities	Non-specific	6	< 5 min
<i>Quality of life</i>				
MSQ	Quality of life limitations	Migraine	14	5 min
SF-36	General health status	Non-specific	36	10 min
SF-12	General health status	Non-specific	12	< 5 min
EQ-5D	Health-related	Non-specific	6	< 5 min
<i>Psychiatric comorbidities</i>				
HADS	Anxiety/depression screening	Non-specific	14	< 5 min
BDI-II	Degree of depression	Non-specific	21	5-10 min
PHQ-9	Depression screening and degree	Non-specific	9	< 5 min
STAI	Degree of anxiety	Non-specific	40	10 min
BAI	Anxiety screening	Non-specific	21	5-10 min
<i>Other</i>				
Short IPAQ	Physical activity	Non-specific	9	5-10 min
PSQI	Sleep quality	Non-specific	19	5-10 min
ISI	Insomnia	Non-specific	5	< 5 min
PSS	Self-perceived stress	Non-specific	14	5-10 min
ASC-12	Allodynia	Migraine	12	5 min
CAPS	Autonomic symptoms	Migraine	5	< 5 min

ASC-12: Allodynia Symptom Checklist; BAI: Beck Anxiety Inventory; BDI-II: Beck Depression Inventory; CAPS: cranial autonomic parasympathetic symptoms; EQ-5D: European Quality of Life–5 Dimensions; HADS: Hospital Anxiety and Depression Scale; HIT-6: Headache Impact Test; IPAQ: International Physical Activity Questionnaire; ISI: Insomnia Severity Index; MIDAS: Migraine Disability Assessment; MSQ: Migraine-Specific Quality of Life Questionnaire; PHQ-9: 9-item Patient Health Questionnaire; PSQI: Pittsburgh Sleep Quality Index; PSS: Perceived Stress Scale; SF: Short Form; STAI: State-Trait Anxiety Inventory; WPAI: Work Productivity and Activity Impairment Questionnaire.

response: a reduction of 2 or 3 points after 4 weeks of treatment is correlated with clinically significant improvement. It can also be used to evaluate patients with chronic migraine or chronic daily headache.

Finally, the Work Productivity and Activity Impairment Questionnaire (WPAI),²⁰ including 6 items, assesses limitations in everyday activities and performance at work. While this scale was not designed specifically for patients with headache or pain, an adapted version was created for this purpose.

Quality of life assessment

In addition to the reduction in disability, improvements in quality of life are also used to evaluate patients' response to certain treatments and interventions. The Migraine-Specific Quality of Life Questionnaire (MSQ)²¹ is ideal for assessing limitations on quality of life and treatment effectiveness. It evaluates the impact of migraine on 3 domains: socialising, work-related activities, and emotions related to migraine. In version 2.1, the content of the questionnaire was improved for greater clarity, and it was shortened to improve its ease of administration.

While it is not specific to migraine, the most widely used instrument for assessing quality of life is the 36-item Short-Form Health Survey (SF-36),²² which addresses physical functioning, physical role, bodily pain, general health, vitality, social functioning, emotional role, and mental health. It includes 36 questions addressing patients' experience over

the previous 4 weeks. An abbreviated version, the SF-12,²³ covers all 8 domains but does not provide specific scores for each.

Another widely used scale, which is also not specific to migraine, is the European Quality of Life–5 Dimensions (EQ-5D) questionnaire,²⁴ which evaluates health-related quality of life. The first part of the questionnaire includes 5 questions covering 5 dimensions (mobility, self-care, usual activities, pain/discomfort, and anxiety/depression), and the second is a visual analogue scale on which the patient is asked to rate their health from 1 to 100.

Assessment of psychiatric comorbidities

It is useful to have tools for screening or assessing the need to treat psychiatric disorders associated with headache. The Hospital Anxiety and Depression Scale (HADS)²⁵ is a widely used screening tool comprising 14 items: 7 assessing anxiety (HADS-A) and 7 assessing depression (HADS-D). Scores are classified as normal, borderline, or abnormal.

Regarding depressive symptoms, the Beck Depression Inventory (BDI)²⁶ is a 21-item questionnaire covering all the diagnostic criteria for depression listed in the fourth edition of the Diagnostic and Statistical Manual of Mental Disorders, and is the most indicated instrument for assessing the severity of depression. Each item is scored from 0 to 3, and the total score is divided into 4 categories: absent or minimal, mild, moderate, or severe. The 9-item Patient Health Questionnaire (PHQ-9)²⁷ may be useful for diagnosing depressive

Table 6 Key procedures in the examination of patients with headache.**General examination****Head and facial examination**

Ophthalmoscopy

Inspection

Trophic/colouration changes

Trigeminal autonomic signs

Palpation

Allodynia; areas of hyper-/hypoalgesia

Pericranial nerves: supra-/infraorbital, GON, LON, mental, supra-/infratrochlear; lacrimal, auriculotemporal

Tenderness test

Trochlear region

Paranasal sinuses

Craniocervical muscles: assess trigger points

Temporomandibular joint

Temporal pulse

Cervical flexion-rotation test

GON: greater occipital nerve; LON: lesser occipital nerve.

symptoms and for monitoring their intensity and treatment response. It includes 9 items scored from 0 to 3; patients are classified as having no depression, minor depressive disorder, or major depressive disorder, according to their total score.

The State-Trait Anxiety Inventory (STAI)²⁸ measures the presence and severity of symptoms of anxiety. It includes 2 subscales comprising 20 questions each: the state subscale addresses the patient's current state (at this moment"), whereas the trait subscale focuses more on trends (generally"). Finally, the Beck Anxiety Inventory (BAI)²⁹ is useful for screening for anxiety: this 21-item questionnaire assesses somatic symptoms of the condition.

Other instruments

When evaluating a patient with headache, we may wish to quantify other symptoms or comorbidities potentially influencing the patient's functional status. While reviewing all such scales is beyond the scope of the present study, Table 5 includes some of the most useful.^{30–35}

Examination of patients with headache

Patient examination should be systematic and include both a general neurological examination and a specific examination of the head (Table 6).

General examination

Special attention should be paid to signs that may present with secondary headache, such as meningeal signs or focal neurological signs.

Head and facial examination**Ophthalmoscopy**

Ophthalmoscopy should always be performed. It is particularly important to identify the optic disc, which measures approximately 1.5 cm and is located to the nasal side and appears pale pink in colour, with a rounded shape and well-defined edges. The emergence of vessels and venous pulsation should be evaluated, and any exudate or haemorrhage identified.

Optic disc oedema or pallor should also be assessed. While the term papilloedema refers specifically to inflammation of the optic disc secondary to increased intracranial pressure, oedema of the optic disc may also be caused by other factors. Cases have also been described of intracranial hypertension syndrome without papilloedema.³⁶ Presence of venous pulsation is an additional sign indicating intracranial pressure below 20 cm H₂O.

Inspection

During the inspection, we must assess changes in colouration or trophic changes (eg, in patients with nummular headache) and establish the presence of trigeminal autonomic signs, such as ocular hyperaemia, tearing, ptosis, miosis, or rhinorrhoea. Examination may reveal findings that are not completely specific but are highly suggestive, such as asymmetrical findings in patients with hemicrania continua.

Palpation

- Cranial allodynia should be assessed with gentle palpation of the parietal region. Subsequently, different symmetrical points on both sides of the head should be compared to identify areas of increased sensitivity or hyperalgesia.
- Pericranial nerves: diffuse cranial allodynia must always be ruled out first. The areas most frequently evaluated are:

Supraorbital nerve: emerges from the supraorbital foramen, in the medial third of the superciliary arch.³⁷

Infraorbital nerve: follows a straight line on the vertical plane along the zygomatic arch from the supraorbital area.³⁸

Greater occipital nerve: on an imaginary line from the occiput to the mastoid process, the nerve emerges where the middle third meets the medial third. It is usually easy to determine the course of the nerve by applying firm pressure along this imaginary line and asking the patient to locate the most painful area.³⁹

Lesser occipital nerve: located laterally to the greater occipital nerve.

Others: mental nerve, supra- and infratrochlear nerves, lacrimal nerve, and auriculotemporal nerve.⁴⁰

During palpation, one or 2 fingers (generally the second and third) should be used for support, while increasing pressure is applied along the course of the nerve. This support is important to prevent the finger from slipping into the eye. One clinical detail that often indicates hypersensitivity of a nerve is the jump sign, whereby the patient will wince and move the head/body to pull away from the pressure. It is also important to be aware that pressure applied to a nerve will cause an unpleasant and even painful sensation, even in

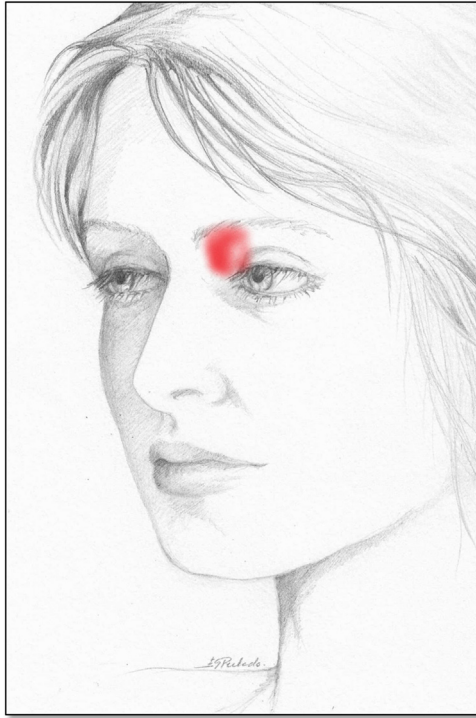


Figure 1 Painful area described by patients with involvement of the trochlear region.

healthy individuals. This part of the examination also provides the opportunity to evaluate such treatment options as anaesthetic nerve block.⁴¹

- **Tenderness test:** this test is mainly used for tension-type headache, and enables us to establish the presence of pericranial hypersensitivity.
- **Trochlear region:** located above the lacrimal caruncle, between the eye and the frontal bone (Fig. 1). Palpation should be used to ascertain the existence of clear hypersensitivity, which is typically exacerbated by vertical movement of the eye (which involves the superior oblique muscle). It should be noted that trochleitis is defined as the inflammation of this region, and specifically the trochlea of the superior oblique nerve; trochleodynia refers to the typical pain in this region without inflammation.
- **Paranasal sinuses:** during palpation of the paranasal sinuses, we must bear in mind that the supratrochlear and supraorbital nerves run close to this area; the pain evoked in these nerves is much more focal.
- **Palpation of the main craniocervical muscles:** it is essential to search for trigger points, or hyperirritable spots in the muscle, which may cause local pain or a pattern of referred pain when the muscle is active. Tender points (areas in which palpation provokes intense pain) are identifiable by a clear tendency to recoil.⁴² Patients with temporomandibular joint disorders often present trigger points in the masseter and muscles of the temporal area. In these patients, clicking sounds and pain upon palpation of the temporomandibular joint and disproportionate wear of the cusps of the teeth should alert us to the possibility of temporomandibular joint dysfunction syndrome.

- **Bilateral palpation of the temporal pulse:** this is important in elderly patients with suspected temporal arteritis.

Cervical flexion-rotation test

This test should be performed in patients with predominantly posterior headache. From behind the seated patient, the examining physician must flex the patient's neck vertically by around 10°, then rotate the head on the horizontal plane to determine whether the patient is able to rotate the head by at least 70° in both directions. Absence of asymmetry or reproduction of the pain that led the patient to consult can reasonably be considered to rule out cervicogenic headache.

Recommendations for patient education, expectation management, and follow-up

After the pharmacological treatment has been explained, clear instructions should be given to track pain at home with the use of headache calendars and diaries; patient education should also be provided to resolve any questions and manage patients' expectations.

Headache calendars and diaries

The difference between these tools is that while headache diaries record the semiology of each episode in greater detail, and are mainly useful in early stages of assessment, calendars track the chronological distribution of episodes, and help in identifying aggravating factors and evaluating the effectiveness of preventive treatments without risk of recall bias.^{43–45}

The key points to be recorded are:

- Pain days, indicating intensity (mild/moderate/severe)
- Disability days
- Symptomatic treatment days (number of pills)
- Menstruation.

This information may be noted on a paper record (which is not always available, potentially leading to less complete data or forgotten days) or electronically. Advances in information technology have led to the creation of digital tools specifically designed to be used simultaneously by patients and physicians.⁴⁶ Specific diaries have also been designed to improve the semiological description of migraine aura and postdromic symptoms.^{47,48} While they are of limited clinical value, and patients must be instructed in their use, they can be helpful in the differential diagnosis of doubtful cases. These tools also allow for better communication between physician and patient.^{49–51}

Patient education

Several studies have shown that behavioural interventions are well accepted by patients⁵²; interventions aiming to improve lifestyle and adherence to preventive treatment are associated with better clinical outcomes.⁵³ Table 7 shows

Table 7 Education measures for patients with migraine and other primary headaches.

<i>General measures</i>	<i>Stress</i>
Avoidance of trigger factors	Relaxation techniques
Avoidance of irritating stimuli	Biofeedback
Minimal use of screens	Behavioural interventions
Prevention of brain trauma	Cognitive-behavioural therapy
<i>Sleep</i>	<i>Physical exercise</i>
Regular sleep pattern	Increased baseline activity level
8 h' sleep per day	Aerobic exercise
Dinner 4 h before bed	Moderate intensity
Avoidance of fluids for 2 h before bed	Regular frequency
Elimination of naps	Adapted to physical fitness
Avoidance of screens in bed	Progressive onset
Visualisation techniques	Proper hydration
<i>Diet</i>	<i>Overweight and obesity</i>
Balanced, varied, moderate diet	Dietary interventions
5 meals per day	Physical exercise
Avoidance of prolonged fasting	Behavioural therapy
2.5 L water/day	Evaluation of pharmacological treatments
No more than 2 coffees per day on a regular basis	Evaluation of bariatric surgery
Minimal alcohol consumption	Avoidance of drugs that can cause weight gain
<i>Acute treatment</i>	<i>Preventive treatment</i>
Individualised	Non-pharmacological measures
Early	Compliance with daily regime
Appropriate administration route	Effectiveness to be assessed at 3 months
Additional dose in event of recurrence	Treatment to be maintained for 6-12 months
Treatment of accompanying symptoms	Notification of potential adverse effects
Emergency department visit if pain is refractory to treatment	Contact with medical team
Avoidance of opioids and combinations of drugs	
Avoidance of symptomatic medication overuse	

Adapted from Torres-Ferrus and Pozo-Rosich.⁵⁵

some standard recommendations to help in the education of patients with headache.

As well as empowering patients, new technology has enabled them to improve their self-care. The perceived benefits to patients of applications and websites for the follow-up of chronic headache include better self-awareness, the ability to share data with physicians and to review historical data, and better control of their disorder.⁵⁴

Many online resources with patient information are available in English, although some Spanish-language resources also exist. Such resources include:

Mobile applications including calendars, such as MyMigraines and MyMigraines Pro (Guerrero-Peral and colleagues), which also offer the possibility for the patient's neurologist to view the data in real time; and MigraineBuddy, which also assists in identifying potential trigger factors including atmospheric changes in the patient's location.

The website midolordecabeza.org (Pozo-Rosich and colleagues) is aimed both at patients and at professionals, and allows for the creation of calendars, the application of scales for follow-up (data may be sent to the physician to optimise history-taking in consultations), a forum for sharing experiences, and a blog with relevant news, run by specialists. It also provides information on access to research projects.

The websites neurodidacta.es and neurowikia.es also provide information for patients and physicians on headache and other neurological disorders.

Conclusions

Rigorous, systematic history-taking and examination are essential in the diagnosis and treatment of headache, particularly given that complementary examinations are not diagnostic, and are only able to diagnose some secondary headaches. The training of primary care and emergency department physicians, as well as neurologists, is not always appropriate. Although these recommendations are no substitute for a rotation at a headache unit, we hope they will provide basic tools for performing holistic, targeted patient interviews, systematising and unifying the care provided to patients with headache at the national level.

Conflicts of interest

The authors have no conflicts of interest to declare.

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