Trigeminal autonomic cephalalgias: fancy term or constructive change to the IHS classification?

P J Goadsby

A classification based on pathophysiology is a useful aid to differential diagnosis and effective treatment planning

For the neurologist faced with the day to day grind of clinical work a change to terminology may seem like the academics “at it again”. I will try to set out this change and illustrate a physiology that may be attractive to understand, and hopefully one that enhances, clinical practice. Appreciating the physiology of the trigeminal-autonomic reflex may make patients presenting with varying degrees of cranial autonomic activation, such as lacrimation, conjunctival injection, nasal congestion or rhinorrhoea and the like, comprehensible at the bedside.1

The trigeminal autonomic cephalalgias (TACs) is a grouping of headache syndromes recognised in the second edition of the International Headache Society (IHS) classification.2 The term was coined to reflect a part of the pathophysiology of these conditions that is a common thread—that is, excessive cranial parasympathetic activation. The TACs are defined in section III of the second edition of the classification2 and include cluster headache,1 paroxysmal hemicrania, and short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing (SUNCT).4 In an early draft, hemicrania continua was included5 but this was finally classified in section IV.1 I will briefly review the underlying physiology of the trigeminal-autonomic reflex that underpins these conditions and set out their classification and differential diagnosis. I will point out some limitations and some directions for future research. Their therapy is beyond the scope of the present paper, but it has been recently reviewed.6

PATHOPHYSIOLOGY OF TACs

Any pathophysiological construct for TACs must account for the two major shared clinical features characteristic of the various conditions that comprise this group: trigeminal distribution pain and ipsilateral cranial autonomic features.7 The pain producing innervation of the cranium projects through branches of the trigeminal and upper cervical nerves7 to the trigeminocephalic complex8 to whence nociceptive pathways project to higher centres.9 A reflex activation of the cranial parasympathetic outflow provides the efferent loop.

Experimental studies

Stimulation of the trigeminal ganglion in the cat produces cranial vasodilation and neuuropeptide release, notably calcitonin gene related peptide (CGRP) and substance P.10 The dilatation is mediated by antidromic activation of the trigeminal nerve (20% of the effect) and orthodromic activation through the cranial parasympathetic outflow via the facial (VIIth) cranial nerve, for the other 80%.11 The afferent arm of the trigeminal-parasympathetic reflex traverses the trigeminal root,11 synapses in the trigeminal nucleus and then projects to neurones of the superior salivatory nucleus in the pons.12 There is a glutamatergic excitatory receptor in the pontine synapse12 and projection via the facial nerve12 without synapse in the geniculate ganglion. The greater superficial petrosal nerve supplies classic autonomic preganglionic fibres to the sphenopalatine (pterygopalatine in humans) and otic ganglia.13 The sphenopalatine ganglion synapse involves a nicotinic ganglion that is hexamethonium sensitive.14 VIIth cranial nerve activation is associated with release of vasoactive intestinal polypeptide (VIP)15 and blocked by VIP antibodies.16 Changes in the flow of blood in the brain depend on the frequency of stimulation17,18 and are independent of cerebral metabolism.19 There is VIP in the sphenopalatine ganglion,20 as well as nitric oxide synthase, which is also involved in the vasodilator mechanism.21

Human studies

The basic science work outlined above implies an integral role for the ipsilateral trigeminal nociceptive pathways in TACs and predicts in some patients cranial parasympathetic autonomic activation. The ipsilateral autonomic features seen clinically are consistent with cranial parasympathetic activation (lacrimation, rhinorrhoea, nasal congestion, and eyelid oedema) and sympathetic hypofunction (ptosis and miosis). The latter is likely to be a neurapraxic effect of carotid wall swelling19,20 with cranial parasympathetic activation. Some degree of cranial autonomic sympathetic outflow is, therefore, a normal physiological response to cranial nociceptive input.21,22 Indeed other primary headaches, notably migraine,23,24 or patients with facial pain, such as trigeminal neuralgia,25 would be expected to have cranial autonomic activation, and they do. The distinction between the TACs and other headache syndromes is the degree of cranial autonomic activation, not its presence alone.26 This is why some patients with migraine have minor cranial autonomic activation that leads to the term cluster-migraine, when most such patients have migraine with cranial autonomic activation.

Permitting trigeminal-parasympathetic activation

What is the basis for the cranial autonomic symptoms being so prominent in the TACs? Is it due to a central disinhibition of the trigeminal-autonomic reflex?27 Functional imaging studies—positron emission tomography studies in cluster headache13–15 and a functional magnetic resonance imaging (MRI) study in SUNCT syndrome28—has demonstrated ipsilateral posterior hypothalamic activation. Posterior hypothalamic activation seems specific to these syndromes and is not seen in episodes29–32 or chronic33 migraine, or in experimental ophthalmic trigeminal distribution headache.34 There are direct hypothalamic-trigeminal connections30 and the hypothalamus is known to have a modulatory role on the nociceptive and autonomic pathways, specifically trigeminovascular nociceptive pathways.31 Hence, cluster headache and SUNCT syndrome are probably due to an abnormality in the region of the hypothalamus (fig 1) with subsequent trigeminovascular and cranial autonomic activation. Imaging data with paroxysmal hemicrania are keenly awaited. Cranial autonomic features are not invariably linked with trigeminal pain and may persist after lesions of the trigeminal nerve.

DIFFERENTIAL DIAGNOSIS OF TACs

The TACs need to be differentiated from secondary TAC producing lesions, from
Whom multiple attacks were captured. Conjunctival injection and tearing (SUNCT) in unilateral neuralgiform headache attacks with functional MRI in a patient with short lasting blood oxygen level dependent (BOLD)-...www.jnnp.com

Table 1 Clinical features of the trigeminal autonomic cephalalgias (TACs)

<table>
<thead>
<tr>
<th></th>
<th>Cluster headache</th>
<th>Paroxysmal hemiancia</th>
<th>SUNCT syndrome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex (F:M)</td>
<td>1:4</td>
<td>2:1</td>
<td>1:2</td>
</tr>
<tr>
<td>Pain Type</td>
<td>Stabbing, boring</td>
<td>Throbbing, boring,</td>
<td>Burning, stabbing, sharp</td>
</tr>
<tr>
<td></td>
<td></td>
<td>stabbing</td>
<td></td>
</tr>
<tr>
<td>Severity</td>
<td>Severe to excruciating</td>
<td>Excruciating</td>
<td>Moderate to severe</td>
</tr>
<tr>
<td>Site</td>
<td>Orbit, temple, face</td>
<td>Orbit, temple</td>
<td>Periodoral</td>
</tr>
<tr>
<td>Attack frequency</td>
<td>1-8 daily</td>
<td>1-40/day</td>
<td>1/day-30/hour</td>
</tr>
<tr>
<td>Duration of attack</td>
<td>15-180 minutes</td>
<td>2-30 minutes</td>
<td>5-240 seconds</td>
</tr>
<tr>
<td>Autonomic features</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(prominent conjunctival injection and lacrimation)</td>
<td></td>
</tr>
<tr>
<td>Migrainous features*</td>
<td>Yes</td>
<td>Yes</td>
<td>No†</td>
</tr>
<tr>
<td>Alcohol trigger</td>
<td>Yes</td>
<td>Occasional</td>
<td>No</td>
</tr>
<tr>
<td>Indomethacin effect</td>
<td>–</td>
<td>+</td>
<td>–</td>
</tr>
</tbody>
</table>

*Nausea, photophobia (often ipsilateral to the pain) or phonophobia.
†May have photophobia ipsilateral to the pain.
SUNCT: Short lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing.

Table 2 Cluster headache

3.1 Diagnostic criteria:
A All fulfilling criteria A–D
B Severe or very severe unilateral orbital, supraorbital and/or temporal pain lasting 15-180 minutes if untreated
C Headache is accompanied by at least one of the following:
1 Ipsilateral conjunctival injection and/or lacrimation
2 Ipsilateral nasal congestion and/or rhinitis
3 Forehead and facial sweating
4 Ipsilateral eyelid oedema
5 Ipsilateral forehead and facial sweating
6 Ipsilateral miosis and/or ptosis
7 A sense of restlessness or agitation
D Attacks have a frequency from one every other day to eight per day
E Not attributed to another disorder

3.1.1 Episodic cluster headache
Description: Occurs in periods lasting seven days to one year separated by pain free periods lasting one month or more
Diagnostic criteria:
A All fulfilling criteria A–E of 3.1
B At least two cluster periods lasting from 7 to 365 days and separated by pain free remissions of one month or more

3.1.2 Chronic cluster headache
Description: Attacks occur for more than one year without remission or with remissions lasting less than one month
Diagnostic criteria:
A All alphabetical headings of 3.1
B Attacks recur for more than one year without remission periods or with remission periods lasting less than one month

For other primary headaches, migraine is the single biggest problem in the differential diagnosis of cluster headache. Migraine can cluster and despite the best intentions of the IHS classification committee short attacks do occur. Cranial autonomic symptoms are well reported, and the neuropeptide changes are the same as in cluster headache. The occurrence of attacks together does not seem to have the seasonal preponderance that is so typical of cluster headache, and this can be a useful differential diagnostic feature. I regard the term cluster-migraine as unhelpful and I am yet to see a convincing case of a distinct biological entity usefully described by this name. The criterion for the effect of movement was added to cluster headache to sharpen the difference with migraine.

The committee hoped this would draw attention to the fact that most cluster headache patients feel restless or agitated, whereas most migraine patients are quiescent, as IHS-1 recognised. In clinical practice, this symptom, and the periodicity, are extremely helpful in differential diagnosis. The other feature of cluster headache, and this is a feature of TACs when compared with migraine, is that patients with TACs often complain of unilateral, homolateral photophobia, whereas patients with migraine more often complain of bilateral photophobia. Bilateral photophobia in patients with TAC could be speculated to occur in about 25% purely by the chance of them having some migrainous biology. The TACs themselves (table 1) can often be differentiated by their attack...
length. This is certainly true when comparing cluster headache with SUNCT/short lasting unilateral neuralgiform headache attacks with cranial autonomic symptoms (SUNA). The IHS criteria for TACs does betray an uncomfortable biological naivety with regard to the timing. The A, C, D, E/F criteria are rather similar for each TAC (tables 2–4). It seems neat in some way to have SUNCT be up to four minutes long, paroxysmal hemicrania from two to 30 minutes and cluster headache from 15 minutes onwards. The overlap seems minimal. It almost goes without saying that this must be wrong in absolute terms, biology rarely provides such neatness, but it does provide a useful way to identify cases of sufficiently similarity to make biologically meaningful studies.

**CHALLENGES FOR THE TACs**

The classification and biology of the TACs have come a long way in a short time. The syndromes are well established, and although rare compared with migraine they are sufficiently common, with cluster headache affecting about 0.2% of the population, to demand a neurological and headache specialist’s attention. There are some particular issues of classification that are not currently clear.

**Cluster headache**

A patient with a first attack of cluster headache is now simply classified as cluster headache (3.1). This takes the top-down view—that is, diagnose what you can and fill in the detail as available. Such cases are unsuitable for almost any study except natural history studies where they are ideally the starting point. A similar problem is how to refer to patients who have one type of TAC, typically an episodic form, and then evolve to the chronic form. The old classification differentiated primary from secondary chronic cluster headache depending on whether there was a period of episodic headache first. This argument would apply equally to chronic paroxysmal hemicrania. There seems little evidence that the clinical characteristics or therapeutic behaviour of primary or secondary chronic cluster headache are different, and the terminology secondary in headache parlance generally implies an underlying pathology. Moreover, the main clinical imperative when the timing alters would be review, perhaps with investigation, but this is a generic principle in headache management. For the moment the distinction has been dropped.

**Paroxysmal hemicrania (PH)**

The diagnosis of PH by the IHS criteria requires a response to indometacin. This is very difficult. It is not clear what the basis for the indometacin effect is, although it is perfectly clear that the effect is clinically very meaningful (table 5). Patients with PH who are treated with indometacin have an almost unbelievably spectacular resolution. This response seems so distinct that reserving the diagnosis of PH for these patients seems reasonable. Given varying sensitivity to indometacin, we have seen a requirement for a single dose given first thing in the morning of 300 mg indometacin to produce a complete response—perhaps there are unrecognised dosing requirements. There is certainly a timing requirement and again we have seen patients turn off, but only after 10 days at the dose of 275 mg daily.

**SUNCT**

For SUNCT the most immediate challenge must be to define the phenotype properly. We have seen patients who fulfil criteria for SUNA (table 6) but not SUNCT (see table 4). Typically the eye is not red, but we have also seen, for example external auditory canal swelling and periaural flushing as the sole cranial autonomic symptom, as has been reported for PH. It seems possible, given the relative proportion of patients with cluster headache who have lacrimation and conjunctival injection as compared with other cranial autonomic symptoms, that these symptoms are for some reason biologically more likely. This is supported by the same relative changes being seen in experimentally induced head pain.
Thus research criteria for a more encompassing syndrome are proposed (see Table 6).

### CONCLUSION

The TACs represent a great success story in headache. From a classification point of view, the syndromes share much biology so their agglomeration in section III draws attention to them and to the trigemino-parasympathetic reflex. It is highly desirable that headache classification moves to a more biological and pathophysiological basis and the TACs are a step in that direction. The TACs also represent excellent clinical opportunities to take a careful history and offer effective therapy to otherwise highly disabled, suffering patients. Lastly, further investigations of the TACs are bound to illuminate physiological processes whose understanding will be useful to the range of primary headache syndromes.

**J Neurol Neurosurg Psychiatry** 2004; 76: 301–305
doi: 10.1136/jnp.2004.036012

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Competing interests: P J G is a Wellcome Trust Senior Research Fellow.

### REFERENCES


Cholinesterase inhibitors

Action of cholinesterase inhibitors in patients’ brains

K Herholz

Cholinesterase inhibitors in patients’ brains

Cholinesterase (ChE) inhibitors are the only class of drug that has consistently shown improvement in cognitive function in patients with mild to moderate Alzheimer’s disease. Unfortunately, improvement is generally rather small. Recent clinical trials in patients with mild to moderate Alzheimer’s disease concluded that benefits were “below minimally relevant thresholds.”

On the background of this confusing situation, studies are particularly welcome that provide clues as to how ChE inhibitors exert their moderate effect in patients and how we could increase their efficacy. In this issue, such information is provided in a study by Bohnen et al., which measured the actual inhibition of cortical acetylcholine esterase (AChE) activity by donepezil in vivo and studied the correlation of the degree of inhibition with the cognitive effects. Several observations were made that indicate directions for improving therapy.

The inhibition of cortical AChE activity by donepezil at the recommended dose of 10 mg daily was rather low (on average 16–24% depending on cortical regions) and it varied considerably among patients. Although somewhat higher values had been measured with a slightly different tracer by other authors cited in the paper, inhibition of human cerebral AChE is much less than observed in peripheral blood, which is in contrast to findings in rats. Thus, dosage, pharmacokinetics, or specific binding of the drug to human cerebral AChE appear to be suboptimal, and this had not become evident during preclinical and clinical phases of drug development and testing.

The study also indicates that the degree of cerebral AChE inhibition makes a clinical difference because it was significantly correlated with measures of executive function and attention. This indicates that it could indeed be worthwhile to increase inhibition in selected patients, e.g. by higher dosage if side effects permit. It is expected that similar positron emission tomography (PET) studies will be performed to measure inhibition of cerebral butyrylcholinesterase (BChE) for selection and development of drugs that achieve higher effective levels of acetylcholine by additional inhibition of this degradation pathway.

Another interesting aspect is that the relatively small inhibition effects were observed in temporal and parietal association cortex—structures that are thought to be of pivotal importance for episodic and semantic memory— that did not benefit significantly from treatment in this and other studies. One would wish to see similar studies with other ChE inhibitors to determine whether this is a property of the entire class of drugs.

It is gratifying that such direct in vivo assessments of pharmacological action are happening now, which means that we do not depend solely on large trials with clinical outcome measures that are, of course, of utmost clinical importance but often tell very little about the mechanisms that explain interindividual variation. One can hope that this will ultimately provide rational means to improve treatment of individuals, which is in the primary interest of patients and doctors.

REFERENCES

Recreational cannabis use: not so harmless!
D Deplanque

Cannabis and stroke

Cannabis is currently the most widely used illicit drug in Western populations. The question of whether or not it should remain prohibited is under debate in many European countries. Possible adverse health effects play an important role in this debate. The classical anxiolytic, sedative, analgesic, and psychedelic properties of cannabis are well known, but it has recently been suggested that it may also induce cerebrovascular disease. Now for the first time, Mateo et al have shown (this issue, see page 435) that a causal relation with cerebrovascular events is highly plausible. The major argument in favour of this is that the events only occurred during periods when the patient was consuming cannabis, and this is a major criterion of adverse drug reaction monitoring.

Several possible mechanisms are discussed by the authors, the most likely being a drug induced vasculopathy. Others include postural hypotension with secondary impairment of the autoregulation of cerebral blood flow, atrial fibrillation, other arrhythmias, and vasospasm. Though concomitant alcohol consumption may be a confounder, the absence of any proven cause, particularly the absence of haemodynamic or cardiac abnormalities, and the presence of multiple infarcts associated with narrowing of the intracranial arteries strongly suggest a toxic vasculopathy. The long duration of these alterations argues in favour of a drug induced vasculitis rather than vasospasm. Such a vasculitis has already been described in peripheral vessels as a result of chronic cannabis consumption, with many similarities to Buerger’s disease. In the present case, the patient was not a regular user and both the arterial abnormalities and the time course of the disease suggest an immuno-allergic vasculitis.

The systemic effects of natural cannabis compounds (Δ9-tetrahydrocannabinol, Δ8-tetrahydrocannabinol, cannabidiol, and cannabidiol) are primarily mediated by the activation of cannabinoid receptors (CB1 and CB2), which are present in various tissues including the brain, cells of the immune system, blood vessels, and heart. Apart from the vasoactive properties of cannabinoids, it is also possible that a central nervous system vasculitis could result from a dysfunction of the immune system. It remains to be determined whether the immunomodulatory effects of cannabinoids could be involved as a mechanism of cannabis induced vasculitis through their action on CB2 receptors. The very low frequency of such complications may reflect a genetic predisposition in a few individuals. Better identification of patients with cannabis related strokes may lead to epidemiological case-control studies that include genomic investigations. In that recreational cannabis use appears not to be as harmless as was thought, there is a need to improve public information. The therapeutic potential of cannabis and its derivatives should be rigorously evaluated and the benefit to risk ratio taken into account before authorising their medical use.

doi: 10.1136/jnnp.2004.048405

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Competing interests: none declared

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