Proinflammatory pathways in cervicogenic headache

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ABSTRACT
Cervicogenic headache (CEH) is a relatively common form of headache arising from the neck structures. The pathophysiology probably results from various local pain-producing factors such as intervertebral dysfunction, with a no less important role played by the frequent coexistence of a history of head traumas. This report represents a series of pathophysiological studies in CEH patients and the results achieved by pharmacological treatment of the disease. Interleukin-1 (IL-1) and Tumour Necrosis Factor α (TNF-α) exert their multifunctional biological effects by promoting and increasing the molecular events of cellular inflammation. We found that the cytokine pattern of CEH patients is similar to cluster headache - biased towards an inflammatory status. Higher levels of both IL-1β and TNF-α were detected in the sera of CEH patients than the levels in patients with migraine without aura and in healthy subjects. There were also differences between the spontaneous and mechanically worsened pain phases of CEH. Nitric oxide (NO) synthase is also activated in cervicogenic headache. No change in NO metabolites levels has been observed after NO donor administration. This behaviour is clearly different from that observed in migraine and tension headache patients.

We conclude that the high degree of cytokine and NO production in CH may depend on the differing pathophysiological mechanisms at work in CEH than in other forms of headache.

Introduction
There is now growing evidence to support the hypothesis of cervicogenic headache (CEH), an uncommon and still controversial headache disorder, as a multifactorial, strictly unilateral head pain syndrome arising from the neck structures (1). CEH has been recognized as a pain syndrome by the International Association for the Study of Pain (IASP) (1) and is currently accepted as a diagnosis in Europe and in parts of Canada, but it is almost universally rejected in the United States. CEH has not as yet been accepted by the International Headache Society as a diagnostic category (2), but on the basis of the first 1990 edition (3) new diagnostic criteria have recently been published by an ad hoc international study group (CHISG) (4).

Anatomical bases of pain
The biomechanics of the cervical spine are based upon the complex interaction between vertebral articulations, ligaments, and the intrinsic and long muscular structure of the spine itself. Additionally, muscles of the superior portion of the thorax and shoulders have a stabilizing function. The cranio cervical junction (C0-C2) is different in terms of biomechanics from the rest of the cervical spine. The most frequent movements of the head (e.g., lateral rotation up to 25 degrees and nuchal extension) are carried out in large part by segments C0/C1 and C1/C2, and the posture of the cervical column is controlled by proprioceptors situated locally in the articulations, tendons, and muscles. Also, there are close ties between the superior part of the cervical column, the vertebral artery, the nerve roots, and some of the peripheral nerves (e.g., the sinuvertebral nerve).

The C1 and C2 vertebrae are also unique in that they do not have posterior articulations and their respective cervical nerves do not emerge from the vertebral foramen or dural sheaths (as in the case of cranial nerves). The localization of some forms of headache in the neck region has always led to nosologic uncertainty and confusion. For example, certain migraine-type headaches have for years been attributed to a degenerative syndrome of the superior segment of the vertebral spine simply because the pain originated in or radiated to that area. The
Cervicogenic headache appears to be a true multifactorial condition. These factors combine to make the pain an important distinguishing feature of the cervical column at the onset of pain. Additionally, concomitance of pain elsewhere in the body is frequently reported as a characteristic element in the history of patients suffering from cervicogenic headache (14, 15). No pathologic findings have been observed on orbital phlebography in cervicogenic headache or in other forms of primary headaches which lack a vascular inflammatory etiology (16).

Clinical and diagnostic assessment of cervicogenic headache

Cervicogenic headache represents a frequent cause of pain starting from the C2/C3 dermatome with radiation to the occipital area, to vertex of the head (C2 level), the peri-oculo-frontal-auricular-temporal area, and sometimes even to the mandibular and neck regions (C3 level) (9). The pattern of pain from the neck can also follow other directions; for example, to the shoulder and the ipsilateral arm, thus mimicking a radicular type of headache. There may be some clinical variability in the type of pain, which is often continuous, non-pulsating, and non-burning, with some radicular characteristics. Additionally, there may be problems with the sense of balance and autonomic functions. Often there are periodic recurrences of paroxysmal attacks which are asymmetrically, unilaterally localized, and there is frequent concomitance of pain elsewhere in the osteoarticular apparatus. Additionally, variation of position and/or movement of the cervical column at the onset of pain are an important distinguishing feature. These factors combine to make the cervicogenic headache a true multifactorial syndrome (10).

Cervicogenic headache appears to be based more on a functional disorder than on degenerative lesions of the cervical spine (11). Even though clinical information in the past has attributed the pain observed in these regions to osteochondritic/osteoarthritic type problems, various studies have shown that there are no radiological (X-ray, CT) correlations between cervicogenic headache and cervical degenerative lesions (12, 13). Furthermore, it is important to remember that degenerative lesions of the cervical spine are to be found more frequently in the lower and mid-cervical rather than in the upper cervical region, probably because the latter is associated with fewer degenerative risk factors (for example, there is no intervertebral disc between C0/C1 and C1/C2) (12).

Symptoms that may distinguish cervicogenic headaches from certain other common headache syndromes include: a female predominance, the strictly unilateral localization of pain (neck, temporal, orbital and frontal regions), and a distinct reduction in the range of movement of the cervical column. Also, the intensity of pain is often severe in contrast to the radicular syndromes. These clinical symptoms appear to be linked to a very low pain pressure threshold in the occipital part of the head in these patients, with reproduction of pain in the site of pain predominance (14, 15). No pathologic findings have been observed on orbital phlebography in cervicogenic headache or in other forms of primary headaches which lack a vascular inflammatory etiology (16).

Previous trauma in the head and neck region is frequently reported as a characteristic element in the history of patients suffering from cervicogenic headache (10), a fact which may create diagnostic confusion. However, post-traumatic headache or the “postconcussion syndrome” generally have a different clinical picture. The differential diagnosis of this form of headache creates difficulties because of signs and symptoms similar to various other pain syndromes located within the same region. Neri-Barré-Lieou syndrome should, however, be considered as completely distinct from cervicogenic headache because of the prominence of altered autonomic function, if this syndrome can be said to represent a true clinical entity (17, 18). Another pain syndrome with cervicocervical localization that should be considered distinct from the cervicogenic headache is Arnold’s neuralgia, which effectively responds to local steroid therapy (19, 20).

The differential diagnosis of cervicogenic headache includes other common entities such as tension headache and migraine (21), and a rare SUNCT syndrome (a short-lasting, unilateral, neuralgiform headache and tension-type headache) (22). In fact, while the localization of pain in the neck may or may not be present in tension headache, it represents a starting point in cervicogenic headache. In tension headache, the pain can occur on a daily basis and may not be localized in the neck region. There may be a “weight” or pressure-like sensation, and often there is the sensation of a tight skullcap. These clinical characteristics of tension headache are not found in cervicogenic headache. A strict and consistent unilaterality of pain is very rare in migraine with or without aura (23), while it is the rule in cervicogenic headache.

Another form of headache which may be clinically separated from cervicogenic headache is that associated with the Chiari type I malformation (24). The presence of dizziness is often the dominant feature in Chiari patients. However, herniation of the lowest intervertebral cervical disc(s) can also produce a unilateral headache and radiating pain in the ipsilateral arm mimicking cervicogenic headache (25).

The pattern of response of cervicogenic headache to various therapeutic interventions, as well as the findings on certain diagnostic tests, serve to mark this headache as a distinct entity. These clinical characteristics and diagnostic criteria can be used to define similarities and differences with the various other identified headache syndromes.

Proinflammatory cytokines and cervicogenic headache

No evidence of inflammation has yet been demonstrated in CEH, although the administration of epidural corticosteroid (C7) seems to have a short-term clinical effectiveness (26, 27). The diagnostic value of this blockade is supported by the existence of numerous sensitive but non-motor connections between C2 and
Role of nitric oxide in cervicogenic headache

The pivotal role of nitric oxide (NO) was recently suggested in other forms of headaches, such as migraine and cluster headache (34-36). In migraine, NO reduces the cytokine-induced expression of a number of pathophysiological effector molecules characteristic of endothelial activation. Pain crises induced in migraine patients by administering an NO donor (NOD) reduces the monocyte expression of intercellular adhesion molecule 1 (ICAM-1) (37), as well as of endothelial leukocyte adhesion molecule 1 (ELAM-1) (38). Conversely, NO generation (assayed as nitrite levels) appears to be upregulated progressively in migraine from the basal condition to pain.

**Role of nitric oxide in cervicogenic headache**

The nature of the pain in CEH has not yet been defined. However, the enhanced production of both IL-1β and TNF-α in CEH, as shown in the present experiment, could represent a specific signal from the immune system resulting in the subsequent activation of the well-known links existing between immunopeptides and neuropeptides, such as Substance P and calcitonin-gene-related peptide (10). Moreover, the increased level of IL-1β and TNF-α that we measured during the mechanically-induced CEH attacks could be stress-related, but this would not account for the high values detected during the inter-ictal phase. Although the role of these pro-inflammatory cytokines remains to be determined, both IL-1β and TNF-α may promote hyperalgesia in CEH. Furthermore, the anatomical derivation of blood sampling in CEH (extracranial circulation) reinforces the observed differences with migraine (intracranial vessels) (28) and could help to define CEH diagnostically as an inflammatory headache.

Lastly, since chronic pain in the cervical spine (at the cervical zygapophyseal joint level) seems to be a very common problem after whiplash injury (32, 33) our results could be explained as an inflammatory consequence of local traumatic events.

**Fig. 1.** Mean serum values (± SE) of interleukin-1β (IL-1β) and tumour necrosis factor-α (TNF-α) in patients with cervicogenic headache (CEH) (n = 15), patients with migraine without aura (MWA) (n = 15), and in controls (C) (n = 15) (NPAR1WAY and Duncan's t-test). For details, see text.

**Fig. 2.** Comparative values of serum nitrite (NO₂⁻) accumulation in cervicogenic headache (CEH) patients studied during both spontaneous pain (CEH IN) and nitric oxide donor (CEH NOD)-elicited pain, and 15 min after the administration of i.v. acetylsalicylic acid (ASA) 1000 mg (CEH ASA). The control group consisted of healthy subjects (C), NOD migraine patients (MWA), and chronic tension headache patients (CTH). *P < 0.02; **P < 0.001.
attack (39), but only during the pain phase in cluster headache (36).

We found that CEH patients showed a more marked activation of the NO pathway than that reported in migraine or in cluster headache (40); moreover, no difference in NO release was found between a spontaneous CEH attack and CEH pain elicited by NOD administration (Figs. 2-4). Interestingly, pulse intravenous administration of acetylsalicylic acid 1 g (ASA) dramatically reduced the NO levels in CEH patients (Figs. 2-4). Furthermore, the visual analogue scale (VAS) of pain was not affected by the administration of NOD to CEH patients and showed a marked reduction after ASA administration (Fig. 5). Lastly, two subgroups of CEH patients may be distinguished with different amounts of nitrite accumulation and different disease activity (VAS scores) (Fig. 6).

The hypothesis that upregulated NO synthase activity may be related to an activation of the NO-ergic vascular endothelial system is widely accepted for both migraine and cluster headache, partly on the basis of the demonstrated effectiveness of drugs acting on the cerebral vasculature, such as triptans or hyperbaric oxygen (41, 42). Since the cerebral blood flow velocity has been shown to be unchanged during the pain phase of CEH (43) and in addition oxygen, ergotamine and sumatriptan have been proven to be ineffective in CEH (44), the upregulation of the NO-ergic system that occurs during this pain phase cannot be accounted for by a cerebrovascular dysfunction. The role of NO in various cytotoxic and inflammatory processes is diverse. Although numerous reports contend that NO can mediate or increase oxidative injury, a strong case can be made that in fact NO is an antioxidant. The evidence so far suggests that protection against oxidative stress requires submicromolar fluxes of NO, which are achievable under normal physiological conditions. However, at higher concentrations of NO the effects of the various reactive nitrogen oxide species (RNOS) may trigger other mechanisms which can lead to tissue injury and consequently to localized pain (45).

Summarizing these concepts, within a putative cervicogenic headache pathophysiology, the observed increase in NO formation in the presence of reactive oxygen species (ROS) may act jointly with IL-1ß and TNF-α (31) as deleterious, pro-inflammatory, pain-producing agents. In fact, in addition to familiar inflammatory mediators, such as the prostaglandins and bradykinin, potentially important pronociceptive roles have been proposed for a variety of “exotic” species, including the cytokines and nitric oxide (46). Their activation plays a key role in the induction of neuronal sensitization, a process underlying prolonged painful states, such as may be the case with cervicogenic headache.

Lastly, the observed influence on RNOS after ASA administration could be due to a direct inhibition of the NO synthase (NOS) / cyclooxygenase type 1 and 2 (COX-1 COX-2) ratio (47). The inducible isofoms of these two enzymes both have a deleterious effect in various inflammatory diseases (48), among which cervicogenic headache should be included.
Conclusions
In summary, cervicogenic headache seems to represent an ideal experimental model for the study of the multifactorial mechanisms which lie at the origin of headaches stemming from the neck. A growing body of evidence indicates that cervicogenic headache is a distinct nosological entity, clearly distinguishable from both migraine and tension headache (49).

In the future, when the parameters for the clinical, diagnostic and pathophysiological definition of cervicogenic headache have been validated, it will finally be possible to give the proper weight to this under-recognized type of headache (50).

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