Chronicity factors of temporomandibular disorders: a critical review of the literature

Abstract: Facial pain often persists long after any identifiable organic pathology has healed. Moreover, in a subgroup of patients with temporomandibular disorder (TMD), no treatment is effective. Knowledge of factors associated with persistent pain in TMD could help identify personalized treatment approaches. Therefore, we conducted a critical review of the literature for the period from January 2000 to December 2013 to identify factors related to TMD development and persistence. The literature findings showed that chronic TMD is marked by psychological distress (somatization and depression, affective distress, fear of pain, fear of movement, and catastrophizing) and characteristics of pain amplification (hyperalgesia and allodynia). Furthermore, these factors seem to interact in TMD development. In addition, our review demonstrates that upregulation of the serotonergic pathway, sleep problems, and gene polymorphisms influence the chronicity of TMD. We conclude that psychological distress and pain amplification contribute to chronic TMD development, and that interactions among these factors complicate pain management. These findings emphasize the importance of multidisciplinary assistance in TMD treatment.

Keywords: Craniomandibular Disorders; Chronic Pain; Facial Pain.

Introduction

Temporomandibular disorder (TMD)-associated pain is the third most prevalent chronic pain condition worldwide, after tension headaches and back pain.¹ Chronic facial pain, including pain associated with TMD, is most often caused by myoarthropathy of the masticatory system.² Despite the prevalence of TMD, factors involved in the transition from the acute to the chronic phase of the disorder remain unclear.

Chronic TMD occurs in a subgroup of TMD patients who, unlike other TMD patients, do not respond to treatment. These “nonresponders” have higher depression, pessimism, and catastrophizing scores and lower self-efficacy and coping scores than their peers.³ Researchers have proposed a heuristic model of causal influences that contribute to the onset and persistence of TMD and related conditions.⁴ This model includes two principal intermediate phenotypes: psychological distress and pain amplification.⁵ Psychological distress is believed to result from the discomfort and frustration associated with the disorder, and it has a bidirectional role in TMD pain.⁶ Pain amplification refers to alterations in peripheral and central nervous system processes that have
the net effect of amplifying the perceptual response to nociceptive stimuli (e.g., hyperalgesia, allodynia). Pain of this nature and preoccupation with pain often lead to major distress, suffering, and functional disability, which are associated with inappropriate use of medical services and high-cost insurance claims. Furthermore, in patients with chronic TMD, it is difficult to determine what type of care will or will not work for each patient. Knowledge of factors associated with TMD persistence and treatment approaches that emphasize flexibility to meet a patient’s individual needs could represent a new direction in the treatment of TMD-related pain. Therefore, the objective of our literature review was to identify factors related to TMD development and persistence.

**Methodology**

A literature search was performed of electronic bibliographic databases (Medline, PubMed, and LILACS) for the period between January 2000 and December 2013. The following keywords were used: temporomandibular disorders and chronic facial pain combined with: catastrophizing, coping behavior, emotional stress, somatization disorder, affective disorders, depression, hyperalgesia, pain threshold, central sensitization, or sleep disorders.

Two independent reviewers read the obtained abstracts and, by consensus, selected articles that met the following inclusion criteria: patients with TMD diagnosis and chronic facial pain. The reviewers read the selected articles and evaluated them for inclusion in the literature review. Additional inclusion criteria for the research articles were as follows: investigations of the relationship between psychological distress and TMD, and investigations about pain amplification in TMD.

All research articles included in the review had a sample population aged 18 years or older with subjects of both sexes. Of the initial 977 abstracts found (524 related to psychological distress and 453 related to pain amplification), 789 were excluded. Excluded abstracts were those of repeated studies and studies with unrelated scopes. The remaining 188 articles were read in their entirety. Then, 150 studies were excluded because they were not clearly related to the review topics. Most of the 38 selected studies had cross-sectional study designs (only one study had a prospective design).

**Results**

**Psychological Distress**

In TMD, psychosocial factors have been associated with the severity of clinical symptoms and pain chronification. Multiple psychological factors have been implicated as potential risk factors for the development of painful TMD. Main literature findings about the relationship between psychological distress and TMD are presented in Table 1.

**Global Measures of Psychological Function**

Somatization and depression are examples of global psychological symptoms. A recent study, which examined TMD patients of different cultures from widely separated clinical sites, reported a prevalence of severe somatization of 28.5%. Prevalence of severe depression increased with the rate of pain-related impairment, ranging from 16.7% in TMD patients with no disability to 53.8% in patients with high disability and severely limiting impairments. However, it is unclear whether depression and somatization are derived from chronic pain or whether they are risk factors for the development of chronic pain.

**Affective Distress and Psychosocial Stress**

Anxiety is a relatively permanent state of worry and nervousness characterized by physical symptoms, which are usually accompanied by compulsive behaviors or panic attacks. Anxiety levels are correlated with facial pain, and TMD patients who are more anxious seem to be at greater risk of developing chronic pain. Anxiety sensitivity is defined as the fear of anxiety symptoms (e.g., palpitations, dizziness, gastrointestinal upset). Elevated levels of anxiety about pain and fear of pain contribute to disability and interfere with life activities.

If pain is interpreted as threatening (pain catastrophizing), then pain-related fear evolves. Pain-related fear leads to avoidance/escape, followed by disability, disuse, and depression. Depression maintains pain experiences, thereby fueling a vicious circle of increasing fear and avoidance. In patients who do not catastrophize, pain-related fear is unlikely to occur. These patients are expected to confront daily activities rapidly, leading to fast recovery. Among the psychological determinants of TMD persistence,
there is increasing evidence that fear and fear of movement are predictive of future perceived disability. An analysis using the Tampa Scale for Kinesiophobia for TMD (TSK-TMD) showed that TMD functional problems were strongly associated with activity avoidance, but not with somatic focus.  

Coping and Catastrophizing

Coping strategies are defined as constantly changing cognitive and behavioral efforts to manage specific external and/or internal demands. Catastrophizing is defined as expecting or worrying about major negative consequences of a situation, even one of minor importance. Recent findings in chronic TMD patients indicated that treatment nonresponders, who accounted for 16% of the sample, reported more psychiatric symptoms, poorer coping, and higher levels of catastrophizing than patients who responded to treatment.  

Pain Amplification

Main findings from the literature about pain amplification are presented in Table 2. Multiple bodily pain conditions in TMD have been associated with generalized alterations in pain processing. However, it is not fully understood which parts of the peripheral or central nervous systems play a role when hyperalgesia becomes maladaptive rather than protective. Moreover, for reasons still unknown, TMD can manifest as localized pain or in conjunction with widespread pain.  

One study suggested that primary insomnia might share a common substrate underlying central sensitivity or play a causal role in the development of hyperalgesia in TMD patients. Researchers have distinguished between TMD as a regional or widespread pain syndrome on the basis of the identification of a “sensitive” TMD subgroup that had symptoms resembling fibromyalgia and differed from an “insensitive” TMD subgroup. Mechanisms contributing to pain amplification include decreased function in pain inhibitory systems and enhancement of pain facilitatory pathways. Pain amplification may be both an inherited trait and a phenotype that can develop over time in response to emergent biological processes or environmental exposures.  

People with localized TMD differed from healthy controls in their allelic frequency of single nucleotide polymorphisms that mapped to a serotonergic receptor pathway, suggesting that upregulation of the serotonergic pathway may play a role in this condition. Consistent with this possibility,
individuals with localized TMD reported less depressive symptoms compared to patients with TMD and widespread pain. Another study proposed that negative affect and genetics (serotonin transporter polymorphisms) lead to disrupted sleep via an increase in stress reactivity. The interaction of these variables led to an increase in learned negative associations, which, in turn, increased the likelihood of developing poor sleep and insomnia, which are common TMD phenotypes.

Discussion
This review of the literature shows that chronic TMD is marked by psychological distress and pain amplification, and that these factors appear to interact with each other. Moreover, upregulation of the serotonergic pathway, sleep problems, and gene polymorphisms influence the development of chronic TMD.

Physiological distress has been related to masticatory function, which could explain its influence on TMD chronicity. Catastrophizing measurements have been linked to greater levels of depression, activity interference, and perceived jaw interference in TMD patients. However, no process variable (catastrophizing or coping) was associated with the objective measurement of jaw impairment. Beliefs and catastrophizing explained significant portions of the variance in nonmasticatory jaw activity (e.g., laughing and yawning) limitations, but none of these factors were associated with masticatory jaw activity (e.g., eating an apple) limitations.

There were also significant positive correlations between depression and jaw amplitude, and stress and jaw velocity for standardized chewing, but not for habitual chewing. This finding suggests that psychological factors, which manifest in depression and stress, play a role in the association between pain and motor activity.

Myalgia patients may experience more stressful life circumstances and a greater negative effect of illness. These patients showed more severe depressive and nonspecific physical symptoms than patients with internal derangement. The myogenous pain subgroup had significantly higher somatization and depression scores than the normal and arthrogenous pain subgroups.

Although pain is the major complaint of TMD patients, the presence of pain-independent functional problems is especially associated with higher levels of fear of movement. Pain-related fear is more disabling than pain itself and is related to poor behavioral performance. On the other hand, patients with a positive affect, social support, adequate treatment adherence, and spirituality felt better regarding their disease conditions and, consequently, had a better quality of life. Pain beliefs are important predictors of treatment outcome and should be considered in the management of TMD patients.

A prospective study showed that two risk factors for elevated TMD incidence were greater numbers of comorbid pain conditions and a greater

| Author | Measurements | Study Design | Main Findings |
|--------|--------------|--------------|---------------|
| Smith et al. (2009) | Laboratory measures of pain sensitivity and polysomnography | Cross-sectional. N = 53 myofascial TMD. | Primary insomnia was associated with hyperalgesia at a non-orofacial site. |
| Pfau et al. (2009) | Tender point, and quantitative sensory testing profiles. | Case-control N = 23 TMD. N = 18 fibromyalgia. | They found an insensitive subgroup resembling healthy controls and a sensitive TMD subgroup resembling fibromyalgia patients. |
| Park et al. (2010) | Thermal pain sensitivity threshold. | Case-control N = 36 normal and 39 TMD. | TMD patients were more sensitive to thermal pain. |
| Sipilä et al. (2011) | Muscle and Joint standardized palpation. | Cross-sectional. N = 6,227 TMD. | Masticatory muscle pain and TMD joint pain was associated with back, neck and shoulder pain. |
| Chen et al. (2012) | Heat and pressure pain thresholds (PPT). | Case-control N = 76 TMD with widespread body pain (WPT), N = 83 TMD, N = 181 controls. | TMD with WPT presented with reduced PPT in cranial and extracranial regions compared to TMD without WPT. Heat pain tolerance was slightly lower. |
extent of nonspecific orofacial symptoms. Other baseline risk factors were preexisting bodily pain, heightened somatic awareness, and greater extent of pain in response to examiners’ palpation of the head, neck, and body. Other study showed that measures of catastrophizing and active pain coping, which are well-established constructs associated with chronic pain, were not significant predictors of the onset of TMD, but they could play a role in the perpetuation of TMD symptoms.

Taken together, physiological distress, sleep problems, upregulation of the serotonergic pathway, and gene polymorphisms could act as chronicity factors for TMD and pain amplification because they also act as pain-perpetuating factors. These literature findings are in line with the multifactorial etiology of chronic facial pain, and shift the perspective away from a local etiology towards a more central etiology, with dysregulations in the stress and pain-modulating systems. Therefore, to understand how a person responds to persistent pain, it is important not only to examine the physical parameters, but also to consider factors such as cognition, coping strategies, life events, and personality.

**Conclusion**

The chronic TMD process is marked by psychological distress and pain amplification, and these factors appear to interact with one another. We conclude that psychological distress factors (e.g., somatization, catastrophizing, and depression), poor sleep, and genetic polymorphisms related to generalized alterations in pain processing are more commonly associated with TMD development and persistence than mechanical factors (e.g., clenching). Therefore, these factors have been the focus of current research.

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