Myofascial Trigger Points Then and Now: A Historical and Scientific Perspective

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Abstract

The intent of this article is to discuss the evolving role of the myofascial trigger point (MTrP) in myofascial pain syndrome (MPS) from both a historical and scientific perspective. MTrPs are hard, discrete, palpable nodules in a taut band of skeletal muscle that may be spontaneously painful (i.e., active) or painful only on compression (i.e., latent). MPS is a term used to describe a pain condition that can be acute or, more commonly, chronic and involves the muscle and its surrounding connective tissue (e.g., fascia). According to Travell and Simons, MTrPs are central to the syndrome—but are they necessary? Although the clinical study of muscle pain and MTrPs has proliferated over the past two centuries, the scientific literature often seems disjointed and confusing. Unfortunately, much of the terminology, theories, concepts, and diagnostic criteria are inconsistent, incomplete, or controversial. To address these deficiencies, investigators have recently applied clinical, imaging (of skeletal muscle and brain), and biochemical analyses to systematically and objectively study the MTrP and its role in MPS. Data suggest that the soft tissue milieu around the MTrP, neurogenic inflammation, sensitization, and limbic system dysfunction may all play a role in the initiation, amplification, and perpetuation of MPS. The authors chronicle the advances that have led to the current understanding of MTrP pathophysiology and its relationship to MPS, and review the contributions of clinicians and researchers who have influenced and expanded our contemporary level of clinical knowledge and practice.

Introduction

Myofascial pain is a clinical problem that has generated interest and confusion for decades. The criteria for diagnosis and their relative importance have evolved over time. Myofascial pain is prevalent and a frequent cause of visits to primary care physicians and pain clinics [1,2]. Few people live without ever having experienced muscle pain as a result of trauma, injury, overuse, or strain. This type of pain frequently resolves in a few weeks with or without medical treatment. In some cases, however, muscle pain persists long after resolution of the injury; it may even refer to other parts of the body, usually contiguous or adjacent rather than remote. This heralds the sensitized state, one of the features of a chronic pain disorder, in which the pain itself is the pathology and requires medical intervention for its resolution.

The term “myofascial” has evolved from the view that both muscle and fascia are likely to be contributors to the symptoms [3,4]. Nomenclature from the past included “fibrositis,” which implied inflammation of the connective tissue lining muscle, along with chronic muscle pain. These terms have been replaced by the term “myofascial pain.”

For many clinicians and investigators, the finding of one or more myofascial trigger points (MTrPs) is required to ensure the diagnosis of MPS. An MTrP is a discrete, hyperirritable nodule in a taut band of skeletal muscle that is palpable and tender during physical examination (Figure 1). The pain of MPS is associated with, but may not be caused by, an active MTrP. An active MTrP is clinically associated with spontaneous pain in the immediate surrounding tissue and/or to distant sites in specific referred pain patterns. Strong digital pressure on the active MTrP exacerbates the patient’s spontaneous pain complaint and mimics the patient’s familiar pain experience. MTrPs can also be classified as latent, in which case the MTrP is physically present but not associated with a spontaneous pain complaint.
However, pressure on the latent MTrP elicits local pain at the site of the nodule. Both latent and active MTrPs can be associated with muscle dysfunction, muscle weakness, and a limited range of motion. Over the years, the necessity of the physical presence of myofascial trigger points (MTrPs) for the definition of MPS has been hotly debated.

**Historical Perspective**

Guillaume de Baillous (1538-1616) of France was one of the first to write in detail about muscle pain disorders. In 1816, the British physician Balfour associated “thickenings” and “nodular tumors” in muscle with local and regional muscle pain [5]. Various other publications contained differing descriptions and terminology, which reflects the slow evolution in the understanding of MTrPs. For example, Froriep in 1843 coined the term “Muskelschwiele” (muscle callouses) to describe what he believed was a “callus” of deposited connective tissue in patients with rheumatic disorders [6]. Subsequently, in 1904, Gowers suggested that inflammation of fibrous tissue (i.e., “fibrositis”), created the hard nodules [7]. However, the term fibrositis became discredited, as biopsy data did not substantiate an inflammatory pathology. Schade (1919) later proposed that the nodules, which he called “myogeloses,” were high-viscosity muscle colloids [8,9]. In the mid-1900s, important work was conducted independently by Michael Gutstein in Germany, Michael Kelly in Australia, and J.H. Kellgren in Britain. By injecting hypertonic saline into various anatomical structures such as fascia, tendon, and muscle in healthy volunteers, Kellgren was able to chart zones of referred pain in neighboring and distant tissue. Among others, his work influenced the U.S. physician Janet Travell, whose work on myofascial pain, dysfunction, and trigger points is arguably the most comprehensive to date. Travell and Rinzler coined the term “myofascial trigger point” in the 1950s, reflecting their finding that the nodules can be present and can refer pain to both muscle and overlying fascia [10]. The two-volume book, *Myofascial Pain and Dysfunction: The Trigger Point Manual*, which she co-authored with her colleague David Simons, represents decades of keen observation and study of myofascial pain and MTrPs.

The manual, together with more than 40 papers that Travell published on the subject, was and remains instrumental in defining and popularizing the diagnosis and treatment of MPS and MTrPs among the health care community, including physical therapists, allopathic and osteopathic physicians, chiropractors, dentists, pain specialists, massage therapists, and myofascial trigger point therapists. Among the various allopathic medical specialties, physiatrists currently have the most comprehensive working understanding of MTrPs. This is, in part, because physiatrists see MPS and the MTrP as related to muscle and musculoskeletal dysfunction. Acknowledging the lack of attention that muscle received, Simons stated, “Muscle is the orphan organ. No medical specialty claims it.” There are signs, however, that Simons’ comments, along with Travell’s myofascial pain concepts, are gaining ground in mainstream medicine.

The current use of the term “MPS” implies a specific condition that is distinguished from other soft tissue pain disorders such as fibromyalgia, tendonitis, or bursitis [11]. MPS can be both regional and widespread, sometimes with referred pain, often accompanied by increased tension and decreased flexibility. It has been reported to coincide with other diseases and syndromes associated with pain, for example, rheumatic diseases and fibromyalgia [12]. MPS has also been associated with other pain conditions including radiculopathies, joint dysfunction, disk pathology, tendonitis, cranio-mandibular dysfunction, migraines, tension type headaches, carpal tunnel syndrome, computer-related disorders, whiplash-associated disorders, spinal dysfunction, pelvic pain and other urologic syndromes, post-herpetic neuralgia, and complex regional pain syndrome [12].

MPS has generally, but not universally, been characterized by a physical finding, the MTrP, and a symptom cluster that lacked demonstrable pathology and attracted little research attention until recently. Unlike MPS, fibromyalgia is a widespread and symmetrically distributed pain condition associated with sleep and...
mood disturbances. By comparison, the pain of MPS is usually local or regional, distributed in a limited number of select quadrants of the body, and has traditionally been thought to present independently of mood or sleep abnormalities. Interestingly, recent studies indicate that MPS is associated with both mood and sleep disturbances [13]. However, the definition and pathogenesis of MPS is still not fully understood, and disagreement persists about whether MPS is a disease or process, rather than a syndrome.

A Contemporary Conundrum

Although the MTrP is a common physical finding, it is often an overlooked component of nonarticular musculoskeletal pain because its pathophysiology is not fully understood. Besides the use of palpation, there are currently no accepted criteria (e.g., biomarkers, electrodiagnostic testing, imaging, etc) for identifying or quantitatively describing MTrPs. In addition, diagnostic criteria are imprecise, and the full impact of MPS on life activity and function is not yet understood. To complicate this issue further, MTrPs are associated clinically with a variety of medical conditions including those of metabolic, visceral, endocrine, infectious, and psychological origin [14], and are prevalent across a wide range of musculoskeletal disorders. If the MTrP is frequently associated with other musculoskeletal pain syndromes, it would make this finding nonspecific—if it is not, it would make the MTrP specific for MPS.

Numerous clinicians over the past few centuries, from various countries and specialties, have encountered and described hard tender nodules in muscle. They have attempted to explain their etiology, tissue properties, and relationship to MPS. However, most investigations were hampered by a lack of objective diagnostic techniques that could record more than simply their presence or absence. As a result, theories of the MTrP pathogenesis, pathophysiology, and contribution to the diagnosis of MPS have been speculative.

As previously mentioned, MTrPs are found as discrete nodules within a taut band of skeletal muscle that may be spontaneously painful or painful only upon palpation. Although muscle pain displays unique clinical characteristics compared to cutaneous and neuropathic pain, the nature of the symptoms are highly dependent upon the individual’s perception of its characteristic qualities (e.g., boring, aching, sharp, etc.), intensity, distribution, and duration. The way in which individuals report their symptoms presents a challenge for standardization and validation if these are to be used as diagnostic criteria, outcome measures of improvement, and/or in clinical trials. Characteristics such as the quality of the pain, its distribution, and whether it radiates have never been required for the diagnosis of MPS.

Some investigators are reluctant to diagnose MPS without a palpable nodule and instead rely exclusively on self-reports. MTrPs are central to the process, but are they necessary? MTrPs are commonly found in asymptomatic individuals. These latent MTrPs are nodules with the same physical characteristics as active MTrPs; however, palpation is required to elicit pain. In addition, some nodules are not tender to palpation (nontender nodules) and may be found proximal to or remote from sites of pain. Although the term “MPS” is commonly used and generally accepted, it does not resolve the clinical dilemma of soft tissue pain in which the palpable nodule is nontender or no nodule is palpable (and is not explained by radiculopathy, muscle strain, etc).

In Travell and Simons’ MTrP-centered model of myofascial pain, the diagnostic criteria and relevant clinical findings can be studied only descriptively, using patient-reported outcomes. Moreover, measurements are obtained via dichotomous data (e.g., presence or absence of pain; presence or absence of MTrP) or nominal data (i.e., no nodule versus nonpainful nodule versus latent MTrP versus active MTrP). Because there is a bias toward objective findings, the sine qua non for this syndrome is the spontaneously painful nodule (i.e., active MTrP). Unfortunately, the role of the nodule in this process has not been determined. It remains unknown whether the nodule is an associated finding, whether it is a causal or pathogenic element in MPS, and whether or not its disappearance is essential for effective treatment.

In a survey conducted in 2000, the vast majority of American Pain Society members believed MPS to be a distinct clinical entity, characterized by the finding of MTrPs [15]. A growing number of pain clinics are using Travell’s pioneering techniques for the evaluation and treatment of muscle pain disorders. Nevertheless, the lack of consistent nomenclature, universally accepted diagnostic criteria, objective assessments, and conclusive biopsy findings has led to much controversy and generally poor acceptance by mainstream medicine.

The Role of Muscle

Travell and Simons methodically developed a working model based primarily on muscle anatomy and function, which evolved over years of observation and empirical testing. They consistently applied a descriptive approach and used the physical findings of painful nodules in taut muscle bands, along with the nature and distribution of pain, to establish the diagnosis. Their keen observational skills were also used to record natural history and treatment response as a way of further understanding mechanisms. Their model has proved to be extraordinarily useful to both clinicians and those suffering from pain by helping to identify the active MTrP.

By applying the clinical criteria developed by Travell and Simons, the diagnosis of myofascial pain has historically relied heavily on the clinical history and a
careful physical examination of the soft tissue by a trained clinician. Recent published findings suggest that the local milieu, as well as the nature of the tissue and whether its classification as pliable, stiff, homogeneous, or nodular may be important for evaluation and treatment response [16,17]. The contribution of the physical findings of the adjacent muscle is often considered not important; rather, it is only the presence and status of the MTrP. Accordingly, diagnosis of MPS has been tissue-specific and anatomically based on palpation of the skeletal muscle for MTrPs. Upon examination, reproduction and/or exacerbation of the patients’ spontaneous pain complaint by firm palpation of a hard, tender nodule is classically defined as an active MTrP [18]. Latent MTrPs show similar physical characteristics as active MTrPs but are painful only when palpated. However, both active and latent MTrPs are responsible for muscle stiffness, dysfunction, and restricted range of motion, as well as autonomic dysfunction, though to a lesser degree for latent MTrPs [9,19].

There have been several hypotheses, including Simons’ own work evolving from his Trigger Point Manual [18], published in 1983, that imply that MTrP development requires muscle overload and overuse. Although Simons developed these hypotheses by primarily working with a rabbit model, various researchers have supported his work since then with human studies [9].

The Cinderella Hypothesis [20] provides a possible explanation for the role of muscle in MTrP development. This hypothesis describes how musculoskeletal disorder symptoms may arise from muscle recruitment patterns during submaximal level exertions with moderate or low physical load. These types of exertions are typically utilized by occupational groups such as office workers, musicians, and dentists, in which myalgia and MTrPs have been commonly reported [21]. According to Henneman’s size principle, smaller type I muscle fibers are recruited first and de-recruited last during static muscle exertions. As a result, these “Cinderella” fibers are continuously activated and metabolically overloaded, in contrast to larger motor muscle fibers that spend more time inactivated. This property makes the “Cinderella” fibers more susceptible to muscle damage and calcium dysregulation, key factors in the formation of MTrPs [17]. A study by Treaster et al supports the Cinderella Hypothesis by demonstrating that low-level, continuous muscle contractions in office workers during 30 minutes of typing induced formation of MTrPs [21].

MTrPs can also develop as a result of muscle overuse in cervical and postural muscles during the performance of low-intensity activities of daily living and sedentary work [21,22]. An intriguing possible mechanism involves sustained low-level muscle contractions routinely used in tasks requiring precision and postural stability of the cervical spine and shoulder. As a result of sustained low-level contractions, a decrease in intramuscular perfusion has been postulated. Thus, it is conceivable that ischemia, hypoxia, and insufficient ATP synthesis in type I motor unit fibers may occur and are responsible for increasing acidity, Ca$^{2+}$ accumulation, and subsequent sarcomere contracture. This increased, sustained sarcomere contracture may lead to decreased intramuscular perfusion, increased ischemia, and hypoxia, a vicious cycle that may possibly lead to the development of MTrPs. As a result, several sensitizing substances may be released, leading to local and referred pain in addition to muscle tenderness, which are clinical hallmarks of MPS.

The Role of MTrPs

There exists a spectrum of physical findings and symptoms involving the nodule and surrounding soft tissue. From this perspective, the aim should be to characterize and measure the symptoms and physical findings associated with MPS and MTrPs more quantitatively and objectively using cardinal or nondichotomous data. This will help investigators to elucidate the pathogenesis and pathophysiology of MPS, and to develop better outcome measures for use in clinical treatment trials. Although the specific pathophysiological basis of MTrP development and symptomatology is unknown, several promising lines of scientific study (i.e., biochemical, tissue imaging, and somatosensory testing) as well as a recent systematic and comprehensive evaluative approach (including measures of range of motion; strength; and self-reports of pain, fatigue, mood, and health status) have revealed objective abnormalities [13,16,17,23-28].

The current gold standard for the diagnosis of MPS is the physical examination as described in The Trigger Point Manual: (1) palpation of a taut band; (2) identification of an exquisitely tender nodule (MTrP) in the taut band; and (3) reproduction of the patient’s symptomatic pain with sustained pressure. However, accurate diagnosis depends on the examiner’s clinical acumen, experience, index of suspicion, training, and palpation skills. Although there is no consensus regarding the physical findings associated with MPS, the clinical finding of a hard, palpable nodule identified as an active MTrP is generally accepted. Although it is considered the “gold standard,” digital palpation has several limitations. For instance, it lacks adequate sensitivity and specificity. It is sometimes difficult to classify the pain as spontaneous. For example, an individual may have no pain at rest, but as soon as there is movement, pain begins. These limitations make it difficult to assess treatment efficacy, enable objective study of the natural history of MTrPs, and identify the presence of MTrPs in deeper muscle fibers.

Furthermore, Tough et al reported that although many research papers on myofascial pain referenced The Trigger Point Manual for diagnostic criteria (57 of 93 in their cohort), they were used correctly by
only 12 of them [29]. In fact, in clinical practice, there are a variety of ways in which patients presenting with a regional pain complaint (and suspected of having MPS) may be evaluated and treated, depending upon the location of pain, physical findings, and results of palpation. Accordingly, beyond measuring the subjective intensity and duration of pain, the assessment of clinical outcomes is, by necessity, evolving.

One of the most important characteristics found in clinical examination that confirms the presence of an active MTrP is the local twitch response (LTR). Strumming or snapping the taut band in a direction perpendicular to muscle fibers produces a quick contraction in the muscle fibers of the taut band. The origin of the LTR is not yet fully understood, although this response may be due to altered sensory spinal processing resulting from sensitized peripheral mechanical nociceptors [19].

Electromyographic studies have revealed spontaneous electrical activity (SEA) generated at MTrP loci that was not seen in surrounding tissue [30]. Originally attributed to dysfunctional muscle spindles, the excess electrical activity was later identified as an increase in miniature endplate potentials and excessive acetylcholine (ACh) release [30]. However, there is disagreement in electromyography and physiology literature on the significance of abnormal motor endplate potentials and "endplate noise." According to Simons, investigators who lack training in examining muscles for MTrPs may misinterpret a MTrP's abnormal endplate noise as a normal finding [31,32]. Although electromyography has been used in research studies to confirm the presence of MTrPs, in clinical practice there is no advantage to using electromyography [33].

The diagnosis of MPS associated with MTrPs remains controversial, especially in the medical profession, despite the methodological improvements that have been made since the first interrater reliability study was performed in 1992 [34]. Unfortunately, recent studies are still limited by lack of rater blinding, inadequate statistical analysis of the results, and inadequate description of the study findings [35].

Other researchers emphasize the "neighborhood" of the MTrP (i.e., surrounding fascia) to explain the symptom complex and physical findings associated with MPS. Specifically, Stecco et al focus on 3 anatomical layers: the deep fascia, the layer of loose connective tissue (which houses the highest concentration of hyaluronic acid), and the layer of epimysium below it [36]. An important molecule in this system is hyaluronic acid (HA), an anionic, nonsulfated glycosaminoglycan distributed widely throughout various tissues, and one of the chief components of the extracellular matrix. Normally, HA functions as a lubricant that helps muscle fibers to glide between each other without friction. However, Stecco et al theorize that, as a result of muscle overuse or traumatic injury, the sliding layers start to produce immense amounts of HA, which then aggregate into supermolecular structures, changing its configuration, viscoelasticity, and viscosity. Because of its increased viscosity, HA can no longer function as an effective lubricant, which increases resistance in the sliding layers and leads to densification of fascia, or abnormal sliding in muscle fibers. Interference with sliding can have an impact on range of motion and can cause difficulty with movement, including quality of movement and stiffness. In addition, under abnormal conditions, the friction results in increased neural hyperstimulation (irritation), which then hypersensitizes mechanoreceptors and nociceptors embedded within densified fascia. This hypersensitization correlates with a patient's experience of pain, allodynia, paresthesia, abnormal proprioception, and altered movement. The fact that very few objective, repeatable studies have been conducted to elucidate these concepts demonstrates the constraints of our knowledge regarding the pathophysiology of MPS and our ability to measure what is relevant to the pathogenesis and pathophysiology of the MTrP. Further research is needed to determine not only the role of the MTrP but also its surrounding area [36,37].

Quintner and Cohen [38] also disfavor the MTrP-driven hypotheses regarding MPS, arguing that circular reasoning has led to the construct of the MTrP. They suggest that viewing muscle and MTrPs as the primary source of MPS detracts from the possibility of nonmuscular explanations for MPS, in which MTrPs arise secondary to an underlying condition. Because the clinical characteristics of MPS are indistinguishable from peripheral neural pain, Quintner and Cohen hypothesize that a more likely explanation for the cause of MPS involves sensitization of the nervi nervorum. They suggest that the phenomenon of the MTrP, which is a critical component of MPS, is better understood as a region of secondary hyperalgesia of peripheral nerve origin, based on anatomy and physiology. Butler supported this notion, suggesting that ectopic impulse generation or abnormal impulse-generating sites and their accompanying sensitization processes in peripheral and cutaneous sensory nerves warrant reconsideration of MTrP hypotheses [39].

Although MPS is commonly thought to involve only a local muscular phenomenon of a measurably stiffer, tender nodule, the intriguing results from these studies suggest that MPS is actually a complex form of neuromuscular dysfunction associated with functional deficits and broader symptomatology. It consists of soft tissue and sensory abnormalities involving both the peripheral and central nervous systems. In addition, data suggest that neurogenic inflammation, wide dynamic range neurons, and limbic system structures likely play pivotal roles in muscle sensitization, pain chronification, somato-visceral interactions, and the objective physical findings of allodynia, hyperalgesia, and referred pain patterns [17,40,41].
Using ultrasound imaging and elastography, Sikdar et al demonstrated for the first time that there are abnormalities in the milieu of the muscle containing palpable MTrPs. They found nodular regions of hypo-echogenicity on sonography [16] (Figure 2). To investigate the mechanical properties of the muscle, an external vibration source was used to vibrate the muscle while imaging the distribution of vibration amplitude. This study demonstrated that MTrPs have diminished vibration amplitude on external vibration that is

![Figure 2. Simultaneous 2-dimensional gray-scale and color variance imaging. (A and B) Normal upper trapezius muscle. The normal muscle appears isoechoic and has uniform color variance (TIS = 0). (C and D) Muscle with a palpable myofascial trigger point (MTrP). A hypoechoic region and a well-defined focal decrease of color variance indicating a localized stiffer region is visible (TIS = 1). (E and F) Muscle with a palpable MTrP. Multiple hypoechoic regions and multiple focal nodules are visible (TIS = 2). Abbreviation: TIS, tissue imaging score. (In: Sikdar S, Shah JP, Gebreab T, et al. Novel applications of ultrasound technology to visualize and characterize myofascial trigger points and surrounding soft tissue. Arch Phys Med Rehabil 2009;90:1829-1838, used with permission).]
consistent with local regions of increased mechanical stiffness [16]. These images can be used to document the MTrP and to track it over time to assess treatment response. Sikdar et al have developed a number of methods for quantifying the images to better characterize the muscle milieu surrounding MTrPs. Their findings show that MTrPs are not necessarily associated with isolated nodular lesions but that active MTrPs are associated with substantial heterogeneity in the milieu of the muscle. In many cases, a number of lesions are visualized in close proximity. Quantification of the area of these lesions can distinguish between active and latent MTrPs and normal uninvolved muscle with high sensitivity and specificity [42]. Mechanical properties of the muscle containing MTrPs were quantitatively assessed by tracking the speed of an externally induced vibration as it propagates through tissue. This method is known as shear wave elastography. Using shear wave elastography, affected muscle in subjects with active MTrPs has been shown to be stiffer compared to palpably normal muscle [43]. Furthermore, the echogenicity and echotexture associated with active MTrPs can be quantified using image analysis methods such as entropy filtering, and provide a further source of image contrast and a method to differentiate between active MTrPs, latent MTrPs, and normal tissue [44]. Active MTrPs more often are localized near the distal fascial border of the muscle and have an irregular shape. These shapes can be characterized in 3 dimensions to reveal the complex heterogeneous nature of the muscle milieu. This lends further evidence toward an understanding of this phenomenon not just as an isolated abnormality but, rather, as a more pervasive process that has an impact on the neighborhood of the muscle and fascia. Color Doppler and spectral Doppler imaging have revealed that the neighborhood of MTrPs shows evidence of vascular remodeling that is especially pronounced in patients with acute neck pain [45].

These findings open options for future clinical research studies that could focus on identifying the mechanisms responsible for the etiology, amplification, and perpetuation of MPS. The development of successful treatment approaches depends upon identifying and targeting the underlying mechanisms of pain and dysfunction and addressing the perpetuating factors that maintain this common pain syndrome. Accordingly, researchers have explored the dynamic interaction between the nervous system and measurable variables such as sensitizing substances found in the local milieu of active MTrPs [17], the unique cortical activation observed in MPS [28], and the poor functional levels associated with MPS [13].

The Role of Pain: Peripheral and Central Sensitization

Until recently, researchers have largely relied upon Simons’ Integrated Trigger Point Hypothesis, introduced in 1999, to explain the role of peripheral sensitization. According to this hypothesis, the presence of an abnormal endplate activity augments the series of events leading to MTrP development. As hypothesized by Simons, during abnormal endplate activity, high levels of ACh are released, which travel down the sarcoplasmic reticulum and open calcium channels. When calcium binds to troponin on the muscle fibers, the muscle fibers contract. To release the contraction, ATP is required to change the conformation of the muscle fibers and actively pump calcium back into the sarcoplasmic reticulum. Thus, a lack of ATP perpetuates the sustained contracture near an abnormal endplate.

This leads to increased metabolic demands, compressed capillary circulation (which reduces blood flow, forming local hypoxic conditions), and a polarized membrane potential [9]. The increased demand for and reduced supply of ATP forms the energy crisis, which may evoke the release of neuroreactive substances and metabolic byproducts (i.e., bradykinin [BK], substance P [SP], serotonin [S-HT]) that could sensitize peripheral nociceptors [46]. Although Simons’ hypothesis explains how sensitizing neuroreactive substances are responsible for the pain associated with active MTrPs and is the most credible theory to date, it remains conjectural. Remarkably, key tenets of Simons’ Integrated Trigger Point Hypothesis overlap with the self-sustaining cycle suggested by the Cinderella Hypothesis.

Stimulated by Simons’ work, investigators have sought to provide objective evidence underlying the role of peripheral and central sensitization by using various lines of study, including histological, neurophysiological, biochemical, and somatosensory. For example, Shah et al hypothesize that local tissue injury (with concomitant elevation of various inflammatory mediators, catecholamines, neurogenic peptides, and cytokines) leads to sensitization of the nociceptor terminal (i.e., peripheral sensitization) [26]. In addition, small-fiber, unmyelinated afferents have been found to exhibit retrograde, neurosecretory properties similar to sympathetic fibers, involving a process known as neurogenic inflammation (Figure 3). Therefore, in the presence of persistent nociceptive bombardment (e.g., from an active MTrP), the dorsal root ganglion will release SP and calcitonin gene-related peptide (CGRP) antidromically into the peripheral tissue. The peripheral secretion of these substances can lead to a cascade of events, including the degradation of local mast cells, local vasodilation, plasma extravasation, and the development of a sensitizing biochemical mixture, which may underlie the clinical findings of active MTrPs [47]. This process of neurogenic inflammation leads to the enhanced release of endogenous substances, such as BK, 5-HT, norepinephrine, nerve growth factor, and adenosine. The release of these substances leads to local allodynia and hyperalgesia, and exacerbates local tissue tenderness, causing an active MTrP to become...
Efficiency through activation of previously silent expanded referral pain patterns is increased synaptic peripheral stimulus. A possible explanation for in the central nervous system that outlasts the noxious transition from normal to aberrant pain perception are maximal at several hours after an initial noxious stimulation, and bolster functional changes after oxygenase 2 (COX-2) induction in dorsal horn neurons, Metabolic and gene induction changes, such as cyclo-

Long-lasting alterations in the central nervous system. Do latent MTrPs play a role in central sensitization? How is it possible for these lesions to refer pain to distant locations? According to Mense, two assumptions must be made to explain the mechanism of latent MTrPs. First, these latent MTrPs send nociceptive, subthreshold signals to the dorsal horn of the spinal cord. This would effectively sensitize the central nervous system without the perception of pain, as is characteristic of latent MTrPs. Second, ineffective synapses exist within the dorsal horn. In this way, nociceptors from muscle containing the MTrP have connections to dorsal horn neurons, innervating remote muscle regions [55].

Subthreshold potentials and ineffective synapses could serve to explain the sensory phenomenon of latent MTrPs. Taken together, it could be hypothesized that the characteristics of latent MTrPs occur through a series of events. Although latent MTrPs are not spontaneously

Even more painful and tender. The continual bombardment of primary afferent activity over time can lead to abnormal function and structural changes in the dorsal root ganglia and dorsal horn neurons. This is known as central sensitization, and clinical manifestations include allodynia, hyperalgesia, temporal summation of pain [48], and expansion of the receptive field of pain [17].

There is a biochemical basis to explain the development of peripheral and central sensitization in muscle pain. Continuous activation of muscle nociceptors leads to the co-release of L-glutamate and SP at the presynaptic terminals of the dorsal horn. This leads to maximal opening of calcium-permeable ion channels, which hyperexcites nociceptive neurons and causes apoptosis of inhibitory interneurons [25]. Consequently, a persistent noxious barrage from the periphery can create long-lasting alterations in the central nervous system. Metabolic and gene induction changes, such as cyclooxygenase 2 (COX-2) induction in dorsal horn neurons, are maximal at several hours after an initial noxious stimulation, and bolster functional changes after peripheral tissue injury [49].

Sensitization of primary afferents is responsible for the transition from normal to aberrant pain perception in the central nervous system that outlasts the noxious peripheral stimulus. A possible explanation for expanded referral pain patterns is increased synaptic efficiency through activation of previously silent (ineffective) synapses at the dorsal horn. This concept of opening previously ineffective connections was demonstrated in a rat myositis model. Experimentally induced inflammation unmasked receptive fields remote from the original receptive field, indicating that dorsal horn connectivity expanded beyond the original neurons involved in nociceptive transmission [50]. In this study, nociceptive input resulted in central hyperexcitability, and this finding helps to explain referred pain patterns common to MPS.

Central sensitization may also facilitate additional responses from other receptive fields as a result of convergent somatic and visceral input at the dorsal horn [51] via wide dynamic range (WDR) neurons. Furthermore, afferent fibers have the ability to sprout new spinal terminals that broaden synaptic contacts at the dorsal horn and may also contribute to expanded pain-receptive fields [52]. This change in functional connectivity may occur within a few hours, even before metabolic and genetic alterations occur in dorsal horn neurons [53]. After activating WDR neurons, afferent input from active MTrPs then ascends the spinothalamic tract to reach higher brain centers. In addition to activating the thalamus, muscle afferent input preferentially activates the limbic system (i.e., the anterior cingulate gyrus, insula, and amygdala), which plays a critical role in modulating muscle pain and the emotional or affective component of persistent pain [54]. Increased activity in the limbic system leads to greater fear, anxiety, and stress. Furthermore, Niddam et al demonstrated increased limbic system (i.e., anterior insula) activity in patients with upper trapezius MPS [28].

**Sensitization Without Pain**

Latent MTrPs are not associated with spontaneous pain; however, they cause local and possibly referred pain upon deep palpation. This begs questions of their underlying pathophysiology and connectivity to the central nervous system. Do latent MTrPs play a role in central sensitization? How is it possible for these lesions to refer pain to distant locations? According to Mense, two assumptions must be made to explain the mechanism of latent MTrPs. First, these latent MTrPs send nociceptive, subthreshold signals to the dorsal horn of the spinal cord. This would effectively sensitize the central nervous system without the perception of pain, as is characteristic of latent MTrPs. Second, ineffective synapses exist within the dorsal horn. In this way, nociceptors from muscle containing the MTrP have connections to dorsal horn neurons, innervating remote muscle regions [55].

Subthreshold potentials and ineffective synapses could serve to explain the sensory phenomenon of latent MTrPs. Taken together, it could be hypothesized that the characteristics of latent MTrPs occur through a series of events. Although latent MTrPs are not spontaneously
painful, they send excitatory, subthreshold potentials that sensitize the dorsal horn. Sensitization of the dorsal horn opens previously ineffective synapses to distant muscle sites. As a result of sensitization, palpation of the latent MTrP induces pain locally and to distant sites (i.e., referred pain) upon the opening of previously ineffective synapses. Thereby, following muscle palpation, pain is experienced at the site of the palpated latent MTrP and in distant, seemingly unrelated muscle [55].

Human studies have shown that central sensitization can occur without the experience of acute pain. Intramuscular injection of nerve growth factor (NGF) results in allodynia and hyperalgesia, both manifestations of muscular injection of nerve growth factor (NGF) results can occur without the experience of acute pain. Intraspinal potentials [57,58]. The thalamus and limbic system are critically involved in the subjective experience of pain. Since subthreshold potentials provide nociceptive input without subsequent action potentials into higher brain centers, there is no perception of pain.

The Evolving Concept of Pain Perception

Simons’ Energy Crisis Hypothesis and later his Integrated Hypothesis considered the local milieu of the MTrP, taut band, endogenous muscle contracture, and presence of sensitizing substances to explain local muscle tenderness and pain associated with active MTrPs. However, these hypotheses did not consider the prevailing theories of pain processing such as the Gate Control Theory of Pain by Melzack and Wall [59], even though Simons and Travell were familiar with them. The Gate Control Theory was a breakthrough concept in pain research because it suggested that the pain experience was a dynamic one, in which afferent nociceptive signaling could be modified and influenced by neurons in the CNS acting as inhibitory or excitatory gates. According to the theory, large myelinated Aβ sensory afferents synapse on inhibitory interneurons located primarily in the substantia gelatinosa of lamina II in the dorsal horn. Specifically, activation of these sensory afferents can inhibit the activation of second-order neurons that receive input from the smaller nociceptor fibers (Figure 4). In addition, supraspinal inputs can also modulate pain perception.

These advances in understanding the pathophysiology of chronic pain had implications for MPS. Travell and Simons were well aware of the mechanistic aspects of the Gate Control Theory of Pain, as well as some of the implications with respect to central sensitization and gene regulation. However, the integration of this theory and their clinical observations did not occur until newer theories of pain perception and modulation were described. These involved gene regulation, receptor expression, and depolarization thresholds. In particular, more recent findings and data suggest that other processes such as neurogenic inflammation, sensitization of wide dynamic range neurons, and limbic system dysfunction may play a role in the initiation, amplification, and perpetuation of MPS.

Another important concept that Simons and Travell did not consider in their theories was the dynamic balance between supraspinal descending facilitation and inhibition, and its influence on pain perception. The relative amount of descending facilitation versus inhibition modulates the perception of pain from a normal to an aberrant state. The rostral ventral medulla (RVM) is a critical relay area between the periaqueductal gray and the spinal cord which functions in the descending pain control system. The RVM contains a population of ON cells and OFF cells, which can either increase or decrease the level of pain, respectively. It does so through projections that modulate activity in the dorsal horn. Following initial tissue injury, the ON cells serve a useful and protective purpose designed to prevent further damage. Under ordinary circumstances, tissue healing would lead to a decrease in ON cell activity and an increase in OFF cell activity. However, in chronic musculoskeletal pain conditions, there appears to be an overall shift to a decrease in inhibition, presumably due to an imbalance of ON cell and OFF cell activity [60]. Disrupted descending inhibition in chronic musculoskeletal pain may lead to an increased pain sensitivity of muscle tissue [61].

Evaluation and Treatment

The history of soft tissue pain treatments follows the clinical trends of those physicians and soft tissue
specialists (e.g., massage therapists, MTrP therapists) most likely to evaluate and treat MPS, and not necessarily researchers. Hence, early literature about therapeutics for MPS was mainly descriptive. Publications included recommendations for the use of postural exercise, heat, cold, stretch, fluoromethane spray, electrical stimulation, needling, and acupuncture, to mention a few [62].

In her autobiography Office Hours: Day and Night, Travell describes her approach to the diagnosis of MPS and selection among treatment options. She demonstrates her keen clinical acumen to judge treatment effectiveness, using a trial-and-error empirical approach [63]. In The Trigger Point Manual, Travell and Simons worked together to provide a more systematic approach. They stressed the importance of the physical examination, including thorough palpation of muscle and surrounding tissue to identify MTrPs. The MTrP requires training and experience to identify and was the target for their treatment [9]. They often relied on pressure pain threshold to supplement their palpation findings to distinguish a latent from an active MTrP. Today, several other methods including intramuscular needling, surface electromyography-guided assessment, infrared thermography, ultrasound, and laser Doppler flowmetry are used in an attempt to objectify MTrP findings [64]. Nevertheless, palpation by an experienced clinician remains the gold standard in MTrP identification.

Travell and Simons were not the first to identify and develop a treatment for MTrPs. They were, however, among the first to recognize the relationship of the trigger point to MPS and clinical soft tissue pain syndromes. They proposed that deactivating the trigger point was an essential component in successfully treating the pain syndrome [9].

One of various techniques that Travell and Simons relied on to treat MTrPs was injection with local anesthetics. Travell was greatly influenced by Kellgren’s work using injections with procaine to treat “myalgia” [65]. The injections were beneficial because they could reach muscles that could not be stretched during manual therapies (e.g., sternalis) and had the longest analgesic effect of any treatment at the time. However, in the 1930s-1940s, the use of local anesthetics raised several concerns, including muscle necrosis, fatal anaphylactic shock, and dose-related toxic effects that resulted from multiple treatments using cumulative doses. Travell and Simons stressed the importance of using a low dose and taking precautionary measures by having a tourniquet, intravenous diazepam, equipment for artificial respiration, and cardiac defibrillator available during the injection [9,62]. Although Travell and Simons used this technique, they were concerned about the risks of anaphylactic shock in a susceptible person. Accordingly, they often preferred to deactivate MTrPs by spray and stretch [9], which involves spraying the overlying skin with fluoromethane, introducing a sudden sensory stimulus that distracts the patient from the discomfort associated with stretching the affected muscles [9] (Figure 5). One of the benefits of spray and stretch is that it allows many muscles to be treated in a short period of time. Simons and Travell hypothesized that this method was effective because it targeted an “energy crisis” in the region of the MTrP. Stretching the muscles would lengthen the sarcomere and reduce overlap between actin and myosin molecules, decreasing the need for ATP and breaking the vicious cycle of the energy crisis [19]. Simons and Travell were enthusiastic about this technique, calling it the “workhorse” of myofascial therapy. However, in the event that the MTrP was unresponsive to spray and stretch, injections with anesthetics were still used.

In 1955, Sola and Kuitert introduced saline injections to deactivate MTrPs, which lacked the risks of local anesthetics [66]. Interestingly, Frost et al discovered that, in a double-blinded comparison, 80% of patients...
reported pain relief with saline injections, compared to 52% of patients with injections of mepivacaine [67].

Naturally, the effectiveness of needling without anesthetics lead to the question of whether saline was even necessary for deactivating MTrPs. In 1979, Lewit was one of the first physicians to try needling without the use of anesthetic or even saline solution, a technique that became known as dry needling [68]. Given the size of the hypodermic needle and the invasive technique used, this form of dry needling was quite painful, which deterred others from using it instead of injections with procaine. However, Lewit found that the effectiveness of dry needling was related to both the severity of the pain and the precision by which the needle is inserted in relation to the MTrP [68]. In a study of 241 chronic myofascial pain patients with various pain sites, Lewit found that dry needling caused immediate analgesia in nearly 87% of cases. For example, more than 31% of patients had analgesia that was permanent, whereas 20% had several months of pain relief, 22% several weeks, and 14% had no relief at all [68]. Currently, clinicians use acupuncture needles to minimize pain and tissue injury, and have found that inserting a needle into the general area of the MTrP as opposed to directly into the MTrP can have the same therapeutic effect [62] (Figure 6).

In addition to dry needling, clinicians in the mid-20th century started experimenting with the use of acupuncture, after learning that the Chinese were using acupuncture analgesia to suppress surgically evoked pain [62]. In the 1970s, interest in acupuncture surged, and even President Nixon and his personal physician enthusiastically supported its use. Although Melzack and Wall found that acupuncture points for pain and MTrP sites share a close spatial relationship (71% overall correspondence), MTrPs and acupuncture points are not the same phenomena [69]. For example, MTrPs, unlike acupuncture points, are not immutable, and thus MTrPs sites, as outlined in The Trigger Point Manual, serve mainly as a guide for where a clinician should start looking for MTrPs. Furthermore, MTrPs are palpable tender nodules, whereas acupuncture points are not palpable, necessarily tender, or nodular [18]. Because of the lack of sound scientific studies, interest in acupuncture quickly declined, but continues to be further studied as a topic with important MTrP implications today.

In the 1980s, use of fluoromethane spray, a topical anesthetic, ceased because of its toxic effect on the ozone layer and its highly flammable properties that led to accidental death [70]. This occurrence spurred the development of various alternative treatments. For example, the counterstrain method (see online video supplement), a common positional release technique, is shown to be effective at reducing pain and improving function. Ischemic compression, which aims to equalize the length of the sarcomeres, has also been shown to decrease pain sensitivity. Another method is transverse friction massage, which, when combined with exercise, has been shown to increase flexibility along with function [64]. All three of these techniques stretch the muscles, which is the aim of any manual therapy. In addition, Travell and Simons stressed the importance of moist heat to relax the underlying muscles and to diminish the tension caused by the MTrPs. They also educated patients on proper postural positioning while sitting, standing, and reading to avoid sustained contraction or prolonged shortening of muscles [9].

During this time, Travell and Simons also explored the role of medications with respect to pain relief (codeine, aspirin, anti-inflammatory drugs), muscular relaxation (diazepam), sleep (antihistamines), and post-therapy soreness (anti-inflammatory action), although their findings were largely empirical. Travell and Simons noted that many of the side effects for these drugs were often greater than symptoms of MPS. In cases where the drug seemed effective, there was always the possibility of a placebo effect [9].

In the 1980s and 1990s, there was significant interest in technologies such as transcutaneous electrical nerve stimulation (TENS), ultrasound, and laser for the treatment of soft tissue pain. Researchers debate the effectiveness of laser and ultrasound for the deactivation of MTrPs, but generally agree that these technologies are effective for pain management. Biofeedback and other relaxation techniques like hypnotherapy, are also available to help patients manage their pain by training them to regain control of their pain condition.

It is important to note that clinicians have been treating MPS based on what they believe is its underlying pathophysiology and what are considered safe and effective treatments. For example, Travell and Simons primarily used the spray and stretch technique because they attributed MPS to muscle overload, whereas Sola and Kuitert [66] and Lewit [68] used saline injections and dry needling, respectively, because they believed that their methods mechanically disrupted the dysfunctional endplates located near the MTrP. Both of these theories are still considered plausible and are discussed today.

Although the etiology of MPS and the pathophysiology of MTrPs are not yet fully understood, some investigators are suggesting that treatments should focus not only on the MTrP but also on the surrounding environment (e.g., fascia, connective tissue) [36,55]. The biochemical contributors to pain are very important. The role of muscle, fascia, and their cellular components are also important factors to both MPS and the formation of the MTrP. This thinking has led clinicians to try to reduce the size of the MTrP, correct underlying contributors to the pain, and restore the normal working relationship between the muscles of the affected functional units [71].

According to Dommerholt, all treatments fall into one of these two categories or both: a pain-control phase
and a deep conditioning phase. During the pain-control phase, trigger points are deactivated, improving circulation, decreasing pathological nociceptive activity, and eliminating the abnormal biomechanical force patterns. During the deep conditioning phase, the intra- and inter-tissue mobility of the functional unit is improved, which may include specific muscle stretches, neurodynamic mobilizations, joint mobilizations, orthotics, and strengthening muscle [72].

Current approaches for management of MPS include pharmacological and nonpharmacological interventions. Among the pharmacological approaches are anti-inflammatory, analgesic, and narcotic medications, topical creams, and trigger point injections, which are now safer and more effective. Nonpharmacological interventions include manual therapies, which continue to include post-isometric relaxation, counterstrain method [73], trigger point compression, muscle energy techniques, and myotherapy [72], along with other treatments such as laser therapy [74], dry needling, and massage [35,75].

According to Simons, stretching and strengthening of the affected muscles is important for any treatment [18]. Although many of the manual treatment methods stay the same or are only slightly modified (all include some form of mechanical pressure), it is the underlying
theory as to why they are effective that continues to evolve with further study. Modalities and manual treatments are often clinically effective for deactivating active MTrPs and desensitizing sensitized spinal segments, and are commonly used as a first line of treatment before more invasive therapies are attempted. Although a number of recent reviews and meta-analyses have focused on needling, the effectiveness of manual therapy should not be overlooked, and may possibly be just as effective as needling [76].

Among the invasive therapies, the scientific articles report mixed results. Generally, dry needling, anesthetic injection, steroids, and botulinum toxin-A [BTA] of the trigger point have all been shown to provide pain relief [71,77-81]. Regardless of the method used, there is considerable agreement that elicitation of an LTR produces more immediate and long-lasting pain relief than no elicitation of an LTR [81-86], although some still believe that eliciting an LTR is not necessary for improvement. Nevertheless, within minutes of a single induced LTR, Shah et al found that the initially elevated levels of SP and CGRP within the active MTrP in the upper trapezius muscle decreased to levels approaching that of normal, uninvolved muscle tissue. Although the mechanism of an LTR is unknown, the reduction of these biochemicals in the local muscle area may be due to a small, localized increase in blood flow and/or nociceptor and mechanistic changes associated with an augmented inflammatory response [87]. Although treatment options for soft tissue pain have not changed dramatically, researchers today have certainly discovered better ways of categorizing and analyzing the clinical data that they collect and determining whether a treatment is effective. For example, since Travell and Simons’ time, researchers have begun to use classifications such as latent, “nonpainful

| Table 1 | Comparison of Travell and Simons’ Contributions to the Contemporary Understanding of Myofascial Trigger Points [9,16,17,19,25,26,32,44,68,87-94] |

<table>
<thead>
<tr>
<th>Understanding of MTrPs based on Travell and Simons’ work</th>
<th>Contemporary understanding of MTrPs based on scientific evidence</th>
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</thead>
<tbody>
<tr>
<td><strong>MTrP Characteristics</strong></td>
<td><strong>MTrP Characteristics</strong></td>
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<tr>
<td>A systematic description of physical and electrodiagnostic findings:</td>
<td>Objective measures of abnormal physical findings suggesting MTrP pathophysiology:</td>
</tr>
<tr>
<td>• Defined MTrP as “a hyperirritable locus within a taut band of skeletal muscle . . . [that] is painful on compression and can evoke characteristic referred pain and autonomic phenomena”</td>
<td>• Biochemical findings indicate local and remote inflammation, and local acidic milieu</td>
</tr>
<tr>
<td>• Differentiated active from latent MTrPs</td>
<td>• Biochemical and physical findings implicate local sensitization</td>
</tr>
<tr>
<td>• Codified criteria for identifying MTrPs in the evaluation of pain</td>
<td>• Oxygenation studies indicate local regions of hypoxia</td>
</tr>
<tr>
<td>• Used electrodiagnostic studies to demonstrate abnormal activity, indicating involvement of the neuromuscular junction</td>
<td>• Imaging studies indicate local regions of muscle stiffness</td>
</tr>
<tr>
<td><strong>Relation to MPS</strong></td>
<td><strong>Evidence implicates abnormalities of the myofascial neighborhood beyond the MTrP</strong></td>
</tr>
<tr>
<td>MTrP causes MPS symptomatology:</td>
<td><strong>Relationship between MTrP and MPS has not yet been determined:</strong></td>
</tr>
<tr>
<td>• MTrP associates with focal pain and hyperirritability</td>
<td>• Patients may have MPS without MTrPs, and MTrPs without MPS</td>
</tr>
<tr>
<td>• MTrP presents with pain radiation</td>
<td>• MTrP may or may not present with pain radiation</td>
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<tr>
<td>• MTrP perturbation produces local twitch response</td>
<td>• MTrP perturbation does not always produce local twitch response</td>
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<tr>
<td><strong>Clinical Evaluation</strong></td>
<td><strong>Clinical Evaluation</strong></td>
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<tr>
<td>Clinical case series propose tentative link between symptoms and physical findings:</td>
<td>Clinical studies and trials establish link between symptoms and physical findings:</td>
</tr>
<tr>
<td>• A specific collection of symptoms is associated with MTrPs, including regional pain, decreased flexibility, and clinical signs of allodynia and hyperalgesia</td>
<td>• Mechanisms of muscle nociception, sensitization, and pain have been well documented</td>
</tr>
<tr>
<td>• Stereotypical patterns of referred pain are associated with MTrPs in different muscles</td>
<td>• Biochemical studies link painful MTrPs with muscle nociception, sensitization, and pain</td>
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<tr>
<td><strong>Treatment</strong></td>
<td><strong>Treatment</strong></td>
</tr>
<tr>
<td>Treatments target MTrPs to reduce pain:</td>
<td>Treatments target MTrP to reduce pain and to improve symptoms and function:</td>
</tr>
<tr>
<td>• Spray and stretch</td>
<td>• Manual manipulation</td>
</tr>
<tr>
<td>• Deep massage</td>
<td>• Dry needling</td>
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<tr>
<td>• Anesthetic injections</td>
<td>• Transcutaneous electrical nerve stimulation</td>
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<tr>
<td>• Pharmacological agents</td>
<td>• Ultrasound</td>
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<tr>
<td><strong>Outcome Measures</strong></td>
<td><strong>Outcome Measures</strong></td>
</tr>
<tr>
<td>Treatment of MTrP leads to:</td>
<td>Treatment of MTrP leads to:</td>
</tr>
<tr>
<td>• Improvement of pain</td>
<td>• Improved pain</td>
</tr>
<tr>
<td>• Increased flexibility (anecdotally)</td>
<td>• Decreased tenderness</td>
</tr>
<tr>
<td></td>
<td>• Increased range of motion</td>
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<td></td>
<td>• Improved quality of life</td>
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pallbable,” and “painful, but no nodule” to categorize MPS. Gerber et al have also begun to assess the effect of treatment on other aspects besides pain such as quality of life, function, disability, sleep, mood, and range of motion [13]. Clinicians are shifting the focus from not only pain relief and increasing function, but to improving the patient’s quality of life as well.

Although many practitioners can attest to improvement in pain levels of MPS, it is measured using self-reports of pain levels before and after treatment. To date, the number of randomized, placebo-controlled trials is few, and most of them have small numbers of participants. In addition, because they rely exclusively on self-reports, there remains uncertainty about the validity of the findings. Thus, although a variety of pharmacological and nonpharmacological treatments have shown efficacy, studies with proper size and quantitative outcome measures need to be performed.

Conclusion

To date, the pathogenesis and pathophysiology of MTrPs and their role in MPS remain unknown. Data have been published suggesting that MPS is a pain syndrome that can be acute or chronic, and that it involves muscle and fascia. The MTrP remains central to its diagnosis, and possibly its successful treatment. New methods of describing and imaging the MTrP as well as the milieu of the MTrP have suggested that there are a variety of objective findings associated with the syndrome and active MTrPs. Table 1 compares the contributions of Travell and Simons to the contemporary understanding of the MTrP.

Earlier theories for the pathogenesis of MTrPs and MPS, including muscle overuse and mechanical difficulties, remain current and have been neither proved nor disproved.

Current data suggest that active MTrPs are associated with a high symptom burden and a negative impact on function, both physical and psychosocial. Gerber et al demonstrated that dry needling improves pain and changes the status of the active MTrP to either a non-spontaneously tender nodule or its resolution. Change in MTrP status is associated with a significant reduction in pain, improved mood, function, and level of disability [88]. However, the mechanism through which this occurs has not yet been determined.

Many questions remain to be answered. For example, what is the etiology and pathophysiology of MPS? What is the role of the MTrP in the pathogenesis of MPS? Is the resolution of the MTrP required for clinical response? What is the mechanism by which the pain state begins, evolves, and persists? Although the presence of pro-inflammatory and noxious biochemicals has been established, what are the levels of anti-inflammatory substances, analgesic substances, and muscle metabolites in the local biochemical milieu of muscle with and without MTrPs? How does a tender nodule progress to a myofascial pain syndrome? Which musculoskeletal tissues are involved, what are their properties, and how do these change in response to treatment? These are some of the questions that researchers must address in the future. Proper treatment of MPS requires identification and targeting of the mechanisms and pathophysiology of perpetuating factors to obtain sustained relief.

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