

1 **Clinical Practice Guideline: Dry Needling**

2
3 **Date of Implementation: April 17, 2014**

4
5 **Product: Specialty**

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7
8 **GUIDELINES**

9 American Specialty Health – Specialty (ASH) considers dry needling unproven given
10 insufficient evidence to support any conclusions related to health outcomes and benefits
11 for all indications, including but not limited to:

- 12 • Myofascial Pain Syndrome (MPS)
13 • Musculoskeletal pain; including carpal tunnel syndrome, lateral epicondylitis,
14 shoulder impingement, and others
15 • Osteoarthritis and rheumatoid arthritis
16 • Temporomandibular joint disorders

17
18 Additional clinical trials are required to determine the effectiveness of dry needling for
19 the treatment of MPS and any other condition in order to determine its benefit-risk
20 profile.

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CPT® Codes	CPT® Code Description
20560	Needle insertion(s) without injection(s); 1 or 2 muscle(s)
20561	Needle insertion(s) without injection(s); 3 or more muscles

22
23 For more information, see ASH *Techniques and Procedures Not Widely Supported as*
24 *Evidence Based (CPG 133 – S)* clinical practice guideline.

25
26 Patients must be informed verbally and in writing of the nature of any procedure or
27 treatment technique that is considered experimental/investigational or unproven, poses a
28 significant health and safety risk, and/or is scientifically implausible. If the patient
29 decides to receive such services, they must sign a *Member Billing Acknowledgment Form*
30 (for Medicare use *Advance Beneficiary Notice of Non-Coverage form*) indicating they
31 understand they are assuming financial responsibility for any service-related fees.
32 Further, the patient must sign an attestation indicating that they understand what is known
33 and unknown about, and the possible risks associated with such techniques prior to
34 receiving these services. All procedures, including those considered here, must be
35 documented in the medical record. Finally, prior to using experimental/investigational or

1 unproven procedures, those that pose a significant health and safety risk, and/or those
 2 considered scientifically implausible, it is incumbent on the practitioner to confirm that
 3 their professional liability insurance covers the use of these techniques or procedures in
 4 the event of an adverse outcome.

5 6 **DESCRIPTION/BACKGROUND**

7 Dry needling is a relatively new method of pain management in the United States. It has
 8 been performed in other countries with different variations for quite some time. There are
 9 three (3) main theoretical approaches to dry needling that are based on different
 10 hypotheses and anatomical models:

- 11 1. Myofascial trigger point
- 12 2. Radiculopathy
- 13 3. Spinal segmental sensitization

14 15 **Myofascial Trigger Point Model**

16 Myofascial trigger points (MTrPs) are defined as “hyperirritable spots in skeletal muscle
 17 associated with hypersensitive palpable nodules in a taut band” (Simons et al., 1998).
 18 These are characteristic of myofascial pain syndrome (MPS). Findings suggest that MPS
 19 is a complex form of neuromuscular dysfunction consisting of both motor and sensory
 20 abnormalities involving both the peripheral and central nervous systems (Shah and
 21 Gilliams, 2008). MTrPs are painful upon compression and can give a characteristic pain
 22 referral pattern. They can also give rise to referred tenderness, autonomic responses,
 23 motion restriction, and motor dysfunction. More specifically, trigger points are classified
 24 into active and latent trigger points. An “active” trigger point refers pain at rest, upon
 25 direct palpation, and with activity. On the other hand, “latent” trigger points are also
 26 painful upon compression but do not give off the characteristic referral pattern for the
 27 specific muscle while at rest. Identification of MTrPs by palpation (flat or pincer
 28 technique) includes the following features:

- 29 • Identification of a taut muscle band containing a discrete palpable nodule
- 30 • Focal tenderness
- 31 • Spontaneous exclamation of pain by the patient (e.g., “jump sign”, whole body
 32 movement) in response to digital pressure or dry needling
- 33 • Consistent and reproducible pattern of referred pain
- 34 • A local twitch response [LTR (muscle fasciculation)] by snapping or palpation
- 35 • Electromyogram (EMG) demonstration of end plate noise (Simons et al., 1998;
 36 Shah and Gilliams, 2008; Dommerholt and Huijbregts, 2011; Sari et al., 2012)

37
 38 Referred pain, LTR and EMG demonstration are not essential for clinical diagnosis but
 39 can be considered confirmatory observations (Dommerholt and Huijbregts, 2011). MTrPs
 40 are thought to form due to acute trauma or repetitive microtrauma, lack of exercise,
 41 nutritional deficiencies, postural faults, joint problems with dysfunctional movement
 42 patterns, proximal nerve compression and muscle spasm, muscle overload, and emotional

1 stress (Shah et al., 2008; Simons et al., 1998; Dommerholt and Huijbregts, 2011). The
2 mechanism underlying the development of MTrPs is not completely understood, but
3 recent technological advances are assisting in further understanding. MTrPs are
4 hypothesized to be a result of altered activity of the motor end plate or neuromuscular
5 junction. Changes in acetylcholine receptor activity, numbers of receptors and in
6 acetylcholinesterase (AChE) activity affect end plate activity. According to EMG studies,
7 there is an increase in the frequency of miniature end plate potential activity at the point
8 of maximum tenderness and in the neuromuscular junction end plate zone of the taut
9 band. This has been labeled as spontaneous electrical activity (SEA) and it is generated at
10 the MTrP loci and not seen elsewhere in surrounding tissue (Hubbard and Berkoff, 1993).
11 This has been confirmed by other studies (Hong and Torigoe, 1994; Gerwin and
12 Duranleau, 1997; Chen et al., 2001; Couppe et al., 2001; Simons et al., 2002; Simons and
13 Dommerholt, 2007; Dommerholt et al., 2010; Ge et al., 2011).

14
15 Shah et al. (2008) determined that several biochemical changes commonly occur at active
16 MTrPs using microdialysis sampling techniques. The findings include: excessive release
17 and elevation of acetylcholine, elevated calcitonin gene-related peptide (CGRP) levels,
18 increased bradykinin, substance P, and cytokines [tumor necrosis factor alpha (TNF- α)
19 and interleukin 1 (IL-1)], and decreased pH. The excessive acetylcholine is due to the fact
20 that acetylcholinesterase cannot function as well in an acidic environment, which was
21 also noted. These nociceptive chemicals which have been detected in abnormal high
22 concentrations in MTrPs such as bradykinin, CGRP and substance P are active in the
23 following ways: 1) bradykinin is a nociceptive agent that stimulates the release of tumor
24 necrosing factor and interleukins, some of which can stimulate further release of
25 bradykinin; 2) calcium gene-related peptide (CGRP) modulates synaptic transmission at
26 the neuromuscular junction by inhibiting the expression of AChE; and 3) substance P
27 alters the local microcirculation and vessel permeability (Shah et al., 2008). In general,
28 these chemicals create an environment of hyper-nociception and inflammation.

29
30 Researchers, Dr. Janet Travell and Dr. David Simons, are key educators of the
31 importance of myofascial pain and trigger points in musculoskeletal conditions. Simons
32 introduced the Integrated Trigger Point Hypothesis, that postulates a local energy crisis
33 resulting from the dysfunctional endplates at active loci, which brings together many of
34 these concepts. MTrPs produce spontaneous electrical activity, which is end plate noise
35 due to excessive acetylcholine. This results in muscle shortening, local ischemia,
36 sensitizing substance increase, nociceptive pain and autonomic stimulation (Simons and
37 Dommerholt, 2007). Muscle shortening or contracture compromises the local circulation,
38 causing ischemia, which has been confirmed via measurement of oxygen saturation
39 levels. This severe hypoxia in MTrPs leads to the release of sensitizing substances and
40 activates muscle nociceptors. In support of the shortened muscle concept, Wang and Yu
41 (2000) hypothesized that MTrPs are severely contracted sarcomeres whereby myosin
42 filaments get stuck in the titin gel at the Z-band. Titin is the largest protein that connects

1 the Z-band with myosin filaments within the sarcomere. Histologic studies have
2 confirmed the presence of extremely contracted sarcomeres that result in hypoxia. From
3 here, the cascade of events progresses as described above. In summary, it can be
4 concluded that MTrPs act as peripheral nociceptors that can heighten and preserve
5 sensory signals from the central nervous system. This can result in new areas of pain
6 referral via peripheral nociceptive input because these MTrPs can influence dorsal horn
7 receptors that normally only process information from remote body regions (Simons and
8 Dommerholt, 2007).

9 10 **Radiculopathy Model**

11 Dr. Chan Gunn developed the “radiculopathy model.” He also established a system for
12 the diagnosis and treatment of myofascial pain syndromes known as Intramuscular
13 Stimulation (IMS). IMS applies Cannon’s Law, which causes the muscular system to
14 display a contracted and hypersensitive state of pain and orthopedic dysfunction. Gunn
15 believed that myofascial pain is always secondary to nerve compression or irritation in
16 the form of peripheral neuropathy or radiculopathy. Therefore, myofascial pain is a result
17 of neuropathic pain in the musculoskeletal system. Features of neuropathic pain include
18 dysesthesia or deep aching, pain felt in region of sensory deficit, paroxysmal brief
19 shooting or stabbing pain, allodynia, loss of joint range or pain caused by the mechanical
20 effects of muscle shortening, autonomic symptoms, and muscle shortening in peripheral
21 and paraspinal muscles.

22
23 Theoretically, shortened muscle from the neuropathy would compress and lead to
24 “supersensitive nociceptors,” which generate pain. This theory is based on Cannon and
25 Rosenblueth’s “Law of Denervation.” This law states that the function and integrity of
26 innervated structures is dependent upon the free flow of nerve impulses to provide a
27 regulatory or trophic effect. When the flow is restricted, the innervated structures become
28 atrophic, highly irritable and sensitive. Because striated muscle is the most sensitive of
29 innervated structures, Gunn states that it is the key to myofascial pain of neuropathic
30 origin. This results in overreaction of muscle fibers to a wide variety of chemical and
31 physical inputs (Dommerholt, 2005). According to Gunn, the mechanical effects of
32 muscle shortening may result in many commonly seen musculoskeletal conditions,
33 including tendonitis and arthralgia. When considering the paraspinal musculature, muscle
34 shortening would preserve radiculopathy by disc compression, narrowing of the disc
35 space and/or application of pressure directly on the nerve root. In Gunn’s model, MTrPs
36 do not play a major role but rather the posterior and anterior rami dominate. Given the
37 segmental influence of the rami on the paraspinal and deep lumbar musculature,
38 treatment must always treat the affected area of paraspinals as well as the peripheral
39 muscles involved in the particular nerve root. Gunn assesses specific motor, sensory, and
40 trophic changes to determine which levels are affected from a neuropathic standpoint.
41 Unfortunately, Gunn’s model was not developed beyond what he theorized in 1973. Case
42 reports and review articles restating what was described above have been published but

1 much of what his theory is based on has been refuted by recent research. His major input
2 presently is the notion of segmental dysfunction and the need to consider this when
3 developing treatment interventions.

4 **Spinal Segmental Sensitization Model**

6 This model was developed by Dr. Andrew Fischer and is a combination of the previous
7 two theories; with an acknowledgment that central sensitization is often due to ongoing
8 peripheral nociceptive input. Sensitization of both peripheral and central afferents is
9 responsible for the transition from normal to abnormal pain perception in the central
10 nervous system that outlasts the actual noxious peripheral stimuli. Continual input from
11 peripheral muscle nociceptors may lead to changes in function and connections of
12 sensory dorsal horn neurons via central sensitization (Dommerholt et al., 2010;
13 Dommerholt, 2011). As an example, noxious stimuli from an active MTrP may sensitize
14 dorsal horn neurons, leading to hypersensitivity and allodynia, as well as an increased
15 area of referred pain. This results in hyperexcitation of nociceptor neurons and induces
16 apoptosis of inhibitory interneurons (Simons and Dommerholt, 2007). This noxious
17 barrage of input from the periphery results in chronic alterations in the central nervous
18 system. In this state, substance P is released at the dorsal horn and astrocytes and
19 microglia are activated and can produce cytokines (TNF- α , IL-1, IL-6) that sensitize
20 neurons and generate this hyperalgesia (Simons and Dommerholt, 2007; Watkins et al.,
21 2007). Srbely et al. (2010) tested the hypothesis that dry needle stimulation of an MTrP
22 evokes segmental anti-nociceptive effects in a double-blind RCT of 40 subjects. Results
23 demonstrated that one (1) intervention of dry needling to a single MTrP evokes short
24 term segmental anti-nociceptive effects. Authors concluded that the pain relieving effects
25 occurred due to modulation of segmental mechanisms and may be an important
26 consideration in the management of MPS (Srbely et al., 2010).

27 **Dry Needling**

29 There are several interventions for MPS and soft tissue dysfunction. Dry needling has
30 been proposed as an effective non-pharmacologic treatment that is thought to induce
31 changes in the MTrPs (Hong, 1994; Langevin, 2008; Dommerholt, 2005). Other terms
32 may be used to describe dry needling, such as intramuscular manual therapy, trigger point
33 dry needling, or intramuscular needling. According to the Virginia Board of Physical
34 Therapy Task Force on Dry Needling, “Intramuscular Manual Therapy (Dry Needling) is
35 a technique used to treat myofascial pain that uses a dry needle, without medication, that
36 is inserted into a trigger point with the goal of releasing/inactivating the trigger points and
37 relieving pain.” According to the “Intramuscular Manual Therapy (Dry Needling)
38 Resource Paper” published by the Federation of State Boards of Physical Therapy
39 (FSBPT) on March 8, 2010, “there are numerous scientific studies to support the use of
40 dry needling for a variety of conditions.” Dry needling is a technique that inserts a needle
41 without medication into a myofascial trigger point with the goal to relieve pain, increase
42 blood flow and improve function. Janet Travell, the former White House physician who

1 treated former president John F. Kennedy’s low back pain with dry needling, identified
2 trigger points as hyperirritable and sensitive palpable nodules in a taut band located
3 within skeletal muscle. Travell first described the use of MTrP injections in the treatment
4 of myofascial pain in 1942 (Travell et al., 1942). Her work led to the development of the
5 dry needling technique; differing from her injection treatment, given no substances are
6 used. In 1979, Lewit coined the term “needle effect” as the immediate analgesia that
7 occurs by the delivery of the needle into the tender spot. His study demonstrated that the
8 effectiveness of treatment was related to the intensity of pain produced at the trigger area
9 and to the accuracy with which the site of maximal tenderness was located by the needle.
10 In this paper, he also suggested upon review of techniques that the most important
11 component of the injection was the puncture of the needle and not the anesthetic used.
12 (Lewit, 1979). Since that time, other researchers have made the same finding (Simons et
13 al., 1998; Hong, 1994; Kamanli et al., 2005; Cummings and White, 2001; Ay et al.,
14 2010).

15
16 Simply stated, dry needling techniques utilize a fine gauge solid sterile needle for
17 insertion into the MTrP followed by manipulation of the needle until several LTRs are
18 induced if possible. The FDA classifies these needles as Class II medical devices ranging
19 in length from 1.5 to 130 mm. Needles are not left in situ but are removed once the MTrP
20 is inactivated. Dry needling is based on the traditional Western medical model for
21 examination and evaluation to determine a diagnosis. Western anatomy, physiology,
22 neurology, biomechanics and manual palpation and therapy skills are utilized. Red flag
23 and yellow flag recognition is also included. The site of needle insertion into MTrPs is
24 based on physical findings, although many practitioners may rely on trigger point
25 mapping to assist them. The most common sites for this treatment include neck, shoulder,
26 hip, and paraspinal musculature. The depth of needle penetration varies from superficial
27 to deep and is dependent upon the location of the targeted tissue.

28
29 More specifically, dry needling appears to have three effects: mechanical,
30 neurophysiologic, and chemical. Corrective exercises should be performed upon
31 inactivation of MTrPs (Furlan et al., 2005).

32 33 **Mechanical Effects**

34 Direct mechanical stimulation appears to induce connective tissue remodeling and
35 plasticity that interrupts the pathologic mechanism of MTrPs. Dry needling has been
36 proposed to disrupt the integrity of the motor end plate of the MTrP. Placement of the
37 needle into the shortened sarcomere may place a localized stretch on these contracted
38 structures, which may disentangle the myosin filament from the titin gel at the Z-band.
39 Through this mechanism, the resting length of the sarcomere can be achieved through
40 reduction of actin and myosin overlap. Manipulation of the needle during insertion may
41 further assist in this relaxation by winding the connective tissue up- leading to “needle
42 grasp.” Research has demonstrated that the orientation of collagen following needle

1 insertions with and without manipulation was more parallel and organized after needle
2 manipulation (Langevin et al., 2001 and 2004). As a result of the mechanical stimulation,
3 group II fibers change length, which may induce the gate control system by blocking
4 nociceptive input from the MTrP and achieving pain reduction (Baldry, 2002). The
5 mechanical pressure of the needle has also been associated with the change in electrical
6 activity observed post needling by elicitation of the LTR (Liboff, 1997). Rha et al. (2011)
7 used guided ultrasound to determine presence of LTRs and noted that in the deep back
8 musculature; often a LTR is noted on ultrasound but is not visibly seen. Researchers
9 suggest that ultrasound guidance may improve the therapeutic efficacy of trigger point
10 injection for treating MTrPs in the deep muscles (Rha et al., 2011).

11 **Neurophysiologic Effects**

12 Baldry, Gunn, and Fischer all support the neurophysiologic explanation of the effects of
13 dry needling. Baldry (2002) concludes that dry needling creates long term activation of
14 A-nerve fibers which may activate opioid mediated pain suppression. Another
15 explanation may be the activation of serotonergic and noradrenergic descending
16 inhibitory systems, which block noxious stimulus into the dorsal horn.

17 **Chemical Effects**

18
19 Shah and colleagues demonstrated that increased levels of certain chemicals, such as
20 bradykinin, substance P, CGRP, and others are reduced immediately after dry needling
21 and LTR (Shah et al., 2005, 2008; Vulfsons et al., 2012). Through real time ultrasound
22 studies, the taut band and reduced blood flow have been identified. Upon needling, the
23 hypoxic setting is alleviated with an immediate influx of blood, whereby these pain-
24 inducing chemicals can be dissipated from the area and taken up by the body (Vulfsons et
25 al., 2012; Cagnie et al., 2012; Maher et al., 2013; Turo et al., 2013; Sikdar et al., 2008,
26 2009, 2010).

27 **Dry Needling Techniques**

28
29 Travell pioneered the use of MTrP injections that eventually led to the development of
30 dry needling. There are three (3) techniques of dry needling: Superficial dry needling,
31 deep dry needling, and intramuscular electric stimulation. Typically, when the term dry
32 needling is used, it is in reference to deep dry needling. Superficial needling will be
33 specifically identified or called out because it doesn't provide the mechanical effects to
34 the muscle, nor does it have the profound biochemical effects as when an LTR is elicited
35 during deep dry needling. It targets the peripheral sensory afferents primarily and not the
36 dysfunctional motor units like deep dry needling does (Baldry, 1995). It is also performed
37 less commonly, though Baldry (2002) is a proponent of superficial dry needling except
38 when nerve root compression exists. Kalichman and Vulfsons (2010) suggest using
39 superficial dry needling when the risk of injury is increased, such as when needling over
40 the lung fields or in the presence of large blood vessels. Intramuscular electrical
41 stimulation is simply an additional technique added to deep dry needling to provide
42

1 further muscle contractions through the needle within the targeted muscle. Deep dry
 2 needling is used when mechanical stimulation or deformation of a sensitized MTrP can
 3 produce a patient’s complaint of pain. It is also necessary when the pain originates from
 4 deeper structures such as the multifidi, piriformis or supraspinatus. Also, given that dry
 5 needling is most effective when an LTR is elicited, it is important to go deep enough to
 6 promote this while confirming that the needle is placed correctly in the taut band.
 7 Interestingly, Fernández-de-Las-Peñas et al. (2022) compared the clinical effects of
 8 needling interventions eliciting local twitch responses (LTRs) versus needling without
 9 eliciting LTRs when applied to muscle trigger points (TrPs) associated with spinal pain
 10 of musculoskeletal origin. Six trials were included. The application of a needling
 11 intervention eliciting LTRs was associated with a significant reduction in pain intensity
 12 immediately after treatment when compared to the same needling intervention without
 13 elicitation of LTRs. No effect at short-term follow-up was observed. No significant
 14 differences based on elicitation or non-elicitation of LTRs were found in related disability
 15 or pressure pain thresholds. Authors concluded that low-level evidence suggests an
 16 immediate effect of obtaining LTRs during needling interventions on pain intensity, with
 17 no significant effects on related disability or pressure pain sensitivity in spinal pain
 18 disorders associated with muscle TrPs. Superficial dry needling has been found to be
 19 effective, however to a lesser extent than deep dry needling (Kalichman and Vulfsons,
 20 2010). Superficial dry needling was initially used due to concerns of causing a
 21 pneumothorax when needling a patient deeply, therefore the technique was altered so that
 22 the needle is just inserted into the tissue just overlying the MTrP and left in for a short
 23 time. Some research demonstrates that using this technique abolishes the excessive
 24 tenderness at the MTrP and alleviates the pain (Baldry, 2002; Dommerholt, 2006;
 25 Edwards and Knowles, 2003). The needling procedures can be easily combined with
 26 electrical stimulation. The best results are achieved when the needles are placed within
 27 the dermatomes corresponding to the local pathology and deep needling techniques are
 28 utilized (Couto et al., 2013; Kim et al., 2012).

30 EVIDENCE REVIEW

31 Clinical Studies

32 **Upper Quadrant Myofascial Pain Syndrome**

33 Published literature in this area has increased substantially over the recent past in
 34 attempts to identify the effectiveness and efficacy of dry needling on patients with MPS.
 35 Huang et al. (2011) evaluated outcomes in patients who have received dry needling
 36 treatments and also identified prognostic factors that may influence these outcomes.
 37 Using a prospective cohort design with 92 patients following an eight (8) week dry
 38 needling-stretching protocol for chronic musculoskeletal pain, results demonstrated
 39 reduced pain and improved quality of life. Each patient received eight (8) weekly
 40 treatments whereby accurate needling was confirmed by reproduction of pain and/or an
 41 LTR. Outcomes were measured at two (2), four (4), and eight (8) weeks. Pain reduction
 42 occurred at each point in time, with the greatest effect size at two (2) weeks. Prognostic

1 factors associated with poorer outcomes included longer duration of symptoms, repetitive
2 work, and sleep deprivation. Limitations included a lack of control group (Huang et al.,
3 2011).

4
5 In another study, Ay et al. (2010) aimed to compare the efficacy of local anesthetic
6 injection and dry needling methods on pain, cervical range of motion (ROM), and
7 depression in MPS patients. This study was designed as a prospective randomized
8 controlled study. Subjects included 80 patients diagnosed with MPS who were randomly
9 assigned into two (2) groups. One (1) group received local anesthetic injection of
10 lidocaine and the other group received dry needling to MTrPs. Both patient groups were
11 given home stretching exercises for the trapezius muscle. Significant improvements were
12 noted in pain. Outcomes were measured using the Visual Analog Scale (VAS), cervical
13 ROM, Beck Depression Scores after four (4) and 12 weeks for both groups. No
14 significant differences were noted between groups. The authors concluded that dry
15 needling was shown to be clinically and statistically beneficial in treating patients with
16 MPS of the trapezius (Ay et al., 2010). Hsieh et al. (2007) investigated changes in PPT of
17 remote MTrPs after dry needling the key active MTrP. 14 patients with bilateral shoulder
18 pain and active MTrPs in infraspinatus muscles participated in this single blinded within-
19 subject design study. An MTrP in the infraspinatus muscle on a randomly selected side
20 was dry needled, and the MTrP on the contralateral side was not and served as a control.
21 Shoulder pain intensity, shoulder internal rotation ROM, and PPT of the MTrPs in the
22 infraspinatus, anterior deltoid, and extensor carpi radialis longus muscles were measured
23 on both sides before and immediately after dry needling. Results demonstrated that both
24 active and passive ROM of shoulder internal rotation and PPT of infraspinatus MTrPs
25 were significantly increased. Pain intensity of the treated shoulder was significantly
26 reduced as well. No significant changes were noted for the control side. The authors
27 concluded this study provides evidence that inactivation of primary MTrPs inhibit the
28 activity in remote MTrPs noted in the area where pain was referred, suggesting a spinal
29 cord mechanism for this finding.

30
31 Tsai et al. (2010) investigated the remote effect of dry needling on the irritability of a
32 myofascial trigger point in the upper trapezius muscle. 35 patients with unilateral active
33 MTrPs in the upper trapezius muscle were randomly divided into two (2) groups. One (1)
34 group received sham needling and the other received dry needling into MTrPs in the
35 extensor carpi radialis longus muscle. Pain, PPT, and neck ROM were measured pre- and
36 post- treatment. Results demonstrated an improvement in all parameters in the study
37 group compared to the control group. The implications of this study are that dry needling
38 a distal MTrP can reduce the irritability of a proximal MTrP. Ga et al. (2007) explored
39 whether dry needling of MTrPs with and without paraspinal needling for elderly patients
40 with MPS differ in outcomes. 40 subjects were randomized into two (2) groups. One (1)
41 received dry needling and the other groups received IMS, indicated needling of
42 corresponding segmental cervical multifidi. Outcome measures included pain rating, PPT

1 rating, and cervical ROM. Depression was also evaluated by the Geriatric Depression
2 Scale-Short Form. At 12 weeks, dry needling at both distal and proximal sites was more
3 effective in reducing pain, improving depression ratings and cervical ROM than just dry
4 needling without including proximal paraspinals (Ga et al., 2007). Shah et al. (2005) used
5 microdialysis sampling of the trapezius to measure the local biochemical milieu at
6 specific points in the upper trapezius muscle. Based on evaluation, Group 1 was
7 established as normal, Group 2 as latent, and Group 3 as active. Samples were obtained
8 before needle movement, during needle advancement and LTR, and after the LTR, for a
9 total of 15 minutes. Results demonstrated that specific chemicals (e.g., SP, CGRP,
10 bradykinin, TNF- α , IL-1) were higher than the latent and normal samples. There was no
11 overall difference between latent and normal points. At post LTR, concentrations of
12 certain chemicals, such as SP and CGRP, were lower than prior to LTR. In a second
13 study, similar sampling was done but in addition to the upper trapezius, sampling was
14 done pre- and post- needling at a remote site with no MTrPs (gastrocnemius). Findings
15 were confirmed for the upper trapezius as in the previous study, including additional
16 analysis of IL-6 and IL-8. Findings demonstrated that the active group had the largest and
17 most elevated levels, the latent group with an intermediate response and the control group
18 the lowest. Despite gastrocnemius findings showing lower concentrations, abnormalities
19 were noted. Explanations suggested were that widespread elevation of substances
20 associated with pain and inflammation follows initial, more local, MTrPs.

21
22 Similar to other studies, Tekin et al. (2013) hypothesized that dry needling is more
23 effective than sham dry needling for patients with MPS. In this prospective, double-
24 blinded, randomized controlled study, 39 subjects were randomized into two (2) groups
25 (study and sham). The treatment group received six (6) sessions of dry needling over four
26 (4) weeks. When VAS scores were compared between the groups, second and third
27 comparisons were significantly lower in the dry needling group. SF-36 scores for both the
28 physical and mental component scores were found to be significantly increased in the dry
29 needling group. This study demonstrated that dry needling treatments are effective in
30 relieving the pain and improving quality of life of patients with MPS. Pecos-Martín et al.
31 (2015) evaluated the effect of dry needling into a myofascial trigger point (MTrP) in the
32 lower trapezius muscle of patients with mechanical idiopathic neck pain. Patients ($N=72$)
33 with unilateral neck pain, neck pain for ≥ 3 months, and active trigger points in the lower
34 trapezius muscle were randomly assigned to 1 of 2 treatment groups. Dry needling in an
35 MTrP in the lower trapezius muscle, or dry needling in the lower trapezius muscle but not
36 at a MTrP. Results indicated that treatment with dry needling of the lower trapezius
37 muscle close to the MTrP showed decreases in pain and PPT as well as an improvement
38 in the degree of disability ($P<.001$) compared with the baseline and control group
39 measurements ($P<.001$). The dry-needling technique performed in the MTrP showed
40 more significant therapeutic effects ($P<.001$). Authors concluded that the application of
41 dry needling into an active MTrP of the lower trapezius muscle induces significant
42 changes in the VAS, NPQ, and PPT levels compared with the application of dry needling

1 in other locations of the same muscle in patients with mechanical neck pain. Cerezo-
2 Téllez et al. (2016) studied the effectiveness of dry needling for chronic nonspecific neck
3 pain in a randomized single-blinded, clinical trial. A total of 130 participants with
4 nonspecific neck pain presenting with active myofascial trigger points in their cervical
5 muscles were included and randomly assigned to receive: DDN plus stretching ($n = 65$)
6 or stretching only (control group [$n = 65$]). Four sessions of treatment were applied over
7 2 weeks with a 6-month follow-up after treatment. Pain intensity, mechanical
8 hyperalgesia, neck active range of motion, neck muscle strength, and perceived neck
9 disability were measured at baseline, after 2 sessions of intervention, after the
10 intervention period, and 15, 30, 90, and 180 days after the intervention. Significant and
11 clinically relevant differences were found in favor of dry needling in all the outcomes (all
12 $P < 0.001$) at both short and long-term follow-ups. Deep dry needling and passive
13 stretching is more effective than passive stretching alone in people with nonspecific neck
14 pain. According to authors, results support the use of DDN in the management of
15 myofascial pain syndrome in people with chronic nonspecific neck pain.

16
17 Gerber et al. (2016) sought to determine whether the benefits of dry needling (DN) of a-
18 MTrPs are sustained 6 weeks posttreatment. A total of 45 patients (13 male and 32
19 female) with cervical pain >3 months and a-MTrPs in the upper trapezius who completed
20 3 DN treatments and who were evaluated 6 weeks post treatment. Responders were
21 patients whose MTrP status changed from active to latent or nonpalpable nodule
22 (resolved). Secondary outcomes were pain pressure threshold (PPT), Profile of Mood
23 States, Oswestry Disability Index (ODI), MOS 36-Item Short-Form Health Survey (SF-
24 36), and cervical range of motion. In this study, there was sustained reduction of pain
25 scores after completion of DN, which is more likely with a greater drop in VAS score.
26 Patients with higher baseline VAS scores are less likely to respond to DN. Early
27 intervention toward significant pain reduction is likely to be associated with sustained
28 clinical response.

29
30 Stieven et al. (2020) sought to determine the added benefit of combining dry needling
31 with a guideline-based physical therapy treatment program consisting of exercise and
32 manual therapy on pain and disability in people with chronic neck pain. Participants were
33 randomized to receive either guideline-based physical therapy or guideline-based
34 physical therapy plus dry needling. The primary outcomes, measured at 1 month post
35 randomization, were average pain intensity in the previous 24 hours and previous week,
36 measured with a numeric pain-rating scale (0-10), and disability, measured with the Neck
37 Disability Index (0-100). The secondary outcomes were pain and disability measured at
38 3- and 6-months post randomization and global perceived effect, quality of sleep, pain
39 catastrophizing, and self-efficacy measured at 1-, 3-, and 6-months post randomization.
40 One hundred sixteen participants were recruited. Authors concluded that when combined
41 with guideline-based physical therapy for neck pain, dry needling resulted in small

1 improvements in pain only at 1 month post randomization. There was no effect on
2 disability.

3
4 Gattie et al. (2021) examined the short- and long-term effectiveness of dry needling on
5 disability, pain, and patient-perceived improvements in patients with mechanical neck
6 pain when added to a multimodal treatment program that includes manual therapy and
7 exercise. Seventy-seven adults (mean \pm SD age, 46.68 ± 14.18 years; 79% female) who
8 were referred to physical therapy with acute, subacute, or chronic mechanical neck pain
9 were randomly allocated to receive 7 multimodal treatment sessions over 4 weeks of (1)
10 dry needling, manual therapy, and exercise (needling group); or (2) sham dry needling,
11 manual therapy, and exercise (sham needling group). The primary outcome of disability
12 (Neck Disability Index score) and secondary outcomes of pain (current and 24-hour
13 average) and patient-perceived improvement were assessed at baseline and follow-ups of
14 4 weeks, 6 months, and 1 year by blinded assessors. Results showed that there were no
15 group-by-time interactions for disability, current pain, or average pain over 24 hours.
16 There were no between-group differences for global rating of change at any time point.
17 Both groups improved over time for all variables; current pain; and average pain over 24
18 hours. Authors concluded that there were no differences in outcomes between trigger
19 point dry needling and sham dry needling when added to a multimodal treatment program
20 for neck pain. Dry needling should not be part of a first-line approach to managing neck
21 pain.

22
23 Murillo et al. (2021) investigated if a single DN session of the Obliquus Capitis Inferior
24 (OCI) muscle improves head and eye movement control-related outcomes, postural
25 stability, and cervical mobility in people with neck pain. Forty people with neck pain
26 were randomly assigned to receive a single session of DN or sham needling of the OCI.
27 Cervical joint position error (JPE), cervical movement sense, standing balance and
28 oculomotor control were examined at baseline, immediately post-intervention, and at
29 one-week follow-up. Active cervical rotation range of motion and the flexion rotation test
30 were used to examine the global and upper cervical rotation mobility, respectively.
31 Analysis revealed that the DN group showed a decrease of JPE immediately post-
32 intervention compared to the sham group which was maintained at one-week follow-up.
33 No effects on standing balance or cervical movement sense were observed in both
34 groups. Upper cervical mobility showed an increase immediately after DN compared to
35 the sham group which remained stable at one-week follow-up. Both groups showed an
36 immediate increase in global cervical mobility. The results from the current study suggest
37 that a single session of DN of the OCI reduces JPE deficits and increases upper cervical
38 mobility in patients with neck pain.

40 **Shoulder Pain**

41 DiLorenzo et al. (2004) evaluated the efficacy of dry needling of MTrPs to relieve
42 hemiparetic shoulder pain resulting from CVA. 101 CVA patients entered the study and

1 randomly assigned to one (1) of two (2) groups. One (1) group received standard
2 rehabilitation and the other group received standard rehabilitation plus dry needling to the
3 shoulder and scapular musculature. Those receiving the needling reported significantly
4 less pain during sleep and physical therapy. Their sleep was also more restful, and
5 frequency and intensity of pain was reduced as well. Osborne and Gatt (2010) described
6 four (4) case reports for elite female volleyball athletes during an intense phase of
7 competition. Dry needling of scapulohumeral muscles was performed. Range of motion,
8 strength and pain were assessed before and after treatment, with a functional assessment
9 of pain immediately after playing and overhead activity, using the short form McGill Pain
10 Questionnaire. All scores were improved post-treatment and athletes were able to
11 continue overhead activities. Trigger point dry needling has been successful in treating
12 athletes with myofascial pain and impingement symptoms but with only subjective
13 improvement and not during a competitive phase. These cases support the use of dry
14 needling in elite athletes during a competitive phase with short-term pain relief and
15 improved function in shoulder injuries. Authors postulate that dry needling may help
16 maintain rotator cuff balance and strength, reducing further pain and injury. Pérez-
17 Palomares et al. (2017) investigated the effectiveness of dry needling in addition to
18 evidence-based personalized physical therapy treatment in the treatment of shoulder pain.
19 One hundred twenty patients with nonspecific shoulder pain were randomized into 2
20 parallel groups: (1) personalized, evidence-based physical therapy treatment; and (2)
21 trigger point dry needling in addition to personalized, evidence-based physical therapy
22 treatment. Patients were assessed at baseline, posttreatment, and 3-month follow-up.
23 There were no significant differences in outcome between the 2 treatment groups. Both
24 groups showed improvement over time. Authors suggested that dry needling did not offer
25 benefits in addition to personalized, evidence-based physical therapy treatment for
26 patients with nonspecific shoulder pain.

27
28 Arias-Buría et al. et al. (2018) evaluated the cost-effectiveness of the inclusion of trigger
29 point-dry needling (TrP-DN) into an exercise program for the management of
30 subacromial pain syndrome. Fifty patients with unilateral subacromial pain syndrome
31 were randomized with concealed allocation to exercise alone or exercise plus TrP-DN.
32 Both groups were asked to perform an exercise program targeting the rotator cuff
33 musculature twice daily for five weeks. Patients allocated to the exercise plus TrP-DN
34 group also received dry needling during the second and fourth sessions. Authors
35 concluded that the inclusion of TrP-DN into an exercise program was more cost-effective
36 for individuals with subacromial pain syndrome than exercise alone. From a cost-benefit
37 perspective, the inclusion of TrP-DN into multimodal management of patients with
38 subacromial pain syndrome should be considered. Pai et al. (2021) evaluated in a
39 randomized, sham-controlled study the pattern of analgesic efficacy and local sensory
40 changes of a single session of DN for MPS in patients with chronic shoulder pain.
41 Patients with chronic shoulder pain were randomized into active ($n = 20$) or sham
42 ($n = 21$) groups. A single DN was performed by a researcher blinded to group assignment

1 and pain outcomes. Pain intensity was assessed by the numeric rating score, and sensory
 2 thresholds were evaluated with a quantitative sensory testing protocol, including the area
 3 of tactile sensory abnormalities 7 days before needling, right before, and 7 days after the
 4 intervention. Results demonstrated that DN led to significant larger pain intensity
 5 reduction. Pain reduction scores were significantly different on the second day after
 6 needling and persisted so until the seventh day and were accompanied by improvement in
 7 other dimensions of pain and a decrease in the area of mechanical hyperalgesia in the
 8 active DN group alone. Authors concluded that active TP DN provides analgesic effects
 9 compared with sham and decreased the area of local mechanical hyperalgesia.

10
 11 Shanmugam et al. (2021) compared the effectiveness of intramuscular electrical
 12 stimulation (IMES) combined with therapeutic exercises versus dry needling (DN)
 13 combined with therapeutic exercises in improving the clinical outcomes in patients with
 14 shoulder adhesive capsulitis (SAC). In this randomized controlled trial, IMES ($n = 45$)
 15 and DN (43) groups had received respectively IMES, and DN twice weekly for three
 16 consecutive weeks. Both groups received therapeutic exercises 1520 minutes, five days in
 17 a week during the second and third week. Pain, disability, kinesiophobia, number of
 18 active and latent MTrPs, shoulder abduction and external rotation range of motion were
 19 assessed at baseline, week-1, week-2, week-3 and follow-up at 3 months. The results
 20 demonstrate that the post intervention assessment scores of VAS, DASH, shoulder
 21 abduction and external rotation ROM, number of active and latent MTrPs and
 22 kinesiophobia were significantly improved in both groups. However, IMES group had
 23 achieved a greater improvement over DN group on the shoulder pain severity and
 24 disability, shoulder range of motion, number of active and latent MTrPs and
 25 kinesiophobia. Despite the significant statistical differences between the groups, IMES
 26 group did not achieve the minimal clinically important differences of 1.5cm and 11-
 27 points respectively for the VAS and DASH scores. No serious adverse effects occurred
 28 during the three weeks of treatment. Authors concluded that IMES combined with
 29 therapeutic exercises is an effective treatment to reduce the shoulder pain severity and
 30 upper limb disability by deactivating the active and latent MTrPs and improving the
 31 shoulder abduction and external rotation range of motion in patients with SAC.

32
 33 Dunning et al. (2020) compared the effects of spinal thrust manipulation and electrical
 34 dry needling (TMEDN group) to those of non-thrust peripheral joint/soft tissue
 35 mobilization, exercise, and interferential current (NTMEX group) on pain and disability
 36 in patients with subacromial pain syndrome (SAPS). Patients with SAPS were
 37 randomized into the TMEDN group ($n = 73$) or the NTMEX group ($n = 72$). Primary
 38 outcomes included the Shoulder Pain and Disability Index and the numeric pain-rating
 39 scale. Secondary outcomes included the global rating of change scale (GROC) and
 40 medication intake. The treatment period was 6 weeks, with follow-ups at 2 weeks, 4
 41 weeks, and 3 months. At 3 months, the TMEDN group experienced greater reductions in
 42 shoulder pain and disability compared to the NTMEX group. Effect sizes were large in

1 favor of the TMEDN group. At 3 months, a greater proportion of patients within the
2 TMEDN group achieved a successful outcome (GROC score of 5 or greater) and stopped
3 taking medication. Authors concluded that cervicothoracic and upper rib thrust
4 manipulation combined with electrical dry needling resulted in greater reductions in pain,
5 disability, and medication intake than non-thrust peripheral joint/soft tissue mobilization,
6 exercise, and interferential current in patients with SAPS. The effects were maintained at
7 3 months.

9 **Temporomandibular Dysfunction**

10 Gonzalez-Perez et al., (2012) evaluated the usefulness of dry needling in the treatment of
11 temporomandibular myofascial pain. 36 subjects with MPS in the external pterygoid
12 muscle were selected to participate. Outcome measures included pain with the visual
13 analog scale and ROM of the mandible before and after needling. Results demonstrated
14 improvement of pain and jaw movement, which continued up to six (6) months after
15 treatment. Pain reduction was more notable for those with higher intensity pain at
16 baseline. Authors concluded that dry needling to the external pterygoid MTrP is effective
17 for temporomandibular MPS. Dıraçoğlu et al. (2012) tested whether dry needling is more
18 effective than sham needling in relieving temporomandibular myofascial pain. 52
19 subjects were randomized into two (2) groups: true dry needling and sham. PPT, pain
20 ratings, and jaw opening were measured pre- and post- treatment. Results indicated that
21 dry needling appears to be an effective treatment method in relieving pain and tenderness
22 of MTrPs.

24 **Hip Pain**

25 A 2004 randomized, double-blind, placebo-controlled trial by Huguenin et al. attempted
26 to establish the effect on straight leg raise (SLR), hip internal rotation (IR), and muscle
27 pain of dry needling to the posterior hip area. 59 male athletes participated in the study
28 and randomly received either dry needling or placebo needling one (1) time to their
29 gluteal MTrPs. ROM (passive SLR and hip IR) and pain were evaluated immediately
30 after, 24 hours and 72 hours after treatment. Pain and ROM improved for both groups,
31 but the change was not different for either group. Given SLR and hip IR did not
32 demonstrate improvements, authors suggested that these tests are not valuable in
33 determining success of dry needling interventions. They suggested that patient reports of
34 response are a better indicator of success (Huguenin et al., 2004). Brennan et al. (2017)
35 investigated whether administration of dry needling (DN) is noninferior to cortisone
36 injection in reducing lateral hip pain and improving function in patients with GTPS.
37 Forty-three participants (50 hips observed), all with GTPS, were randomly assigned to a
38 group receiving cortisone injection or DN. Treatments were administered over 6 weeks,
39 and clinical outcomes were collected at baseline and at 1, 3, and 6 weeks. The primary
40 outcome measure was the numeric pain-rating scale (0-10). The secondary outcome
41 measure was the Patient-Specific Functional Scale (0-10). Authors concluded that
42 cortisone injections for GTPS did not provide greater pain relief or reduction in

1 functional limitations than DN. Data suggest that DN is a noninferior treatment
 2 alternative to cortisone injections in this patient population. Ceballos-Laita et al. (2019)
 3 sought to determine the short-term effects of DN on pain, hip ROM and physical function
 4 in patients with hip OA. Thirty patients with unilateral hip OA were randomized into two
 5 groups: DN group and sham group. Participants received three treatment sessions. The
 6 treatment was applied in active MTrPs of the iliopsoas, rectus femoris, tensor fasciae
 7 latae and gluteus minimus muscles. Pain intensity (visual analogic scale), passive hip
 8 ROM (universal goniometer and digital inclinometer) and physical function (30s chair-
 9 stand test and 20m walk test) were assessed at baseline and after the three treatment
 10 sessions. There was decreased pain intensity, increased hip ROM, and improved physical
 11 function following the DN treatment. These improvements were statistically significant
 12 ($p < 0.05$) compared to the sham group. Authors concluded that pain, hip ROM, and
 13 physical function improved after the application of DN in active MTrPs of the hip
 14 muscles in patients with hip OA.

15
 16 Ceballos-Laita et al. (2021) investigated the short-term effects of dry needling (DN) on
 17 physical function, pain, and hip muscle strength in patients with hip osteoarthritis (OA).
 18 Patients with unilateral hip OA ($N=45$) were randomly allocated to a DN group, sham
 19 DN group, or control group. Patients in the DN and sham groups received 3 treatment
 20 sessions. Three active myofascial trigger points (MTrPs) were treated in each session
 21 with DN or a sham needle procedure. The treatment was applied in active MTrPs of the
 22 iliopsoas, rectus femoris, tensor fasciae latae, and gluteus minimus muscles. Results
 23 demonstrated a significant group by time interactions for physical function, pain, and hip
 24 muscle force variables. Post hoc tests revealed a significant reduction in hip pain and
 25 significant improvements in physical function and hip muscle strength in the DN group
 26 compared with the sham and control groups. The DN group showed within- and between-
 27 groups large effect sizes. Authors concluded that DN therapy in active MTrPs of the hip
 28 muscles reduced pain and improved hip muscle strength and physical function in patients
 29 with hip OA. DN in active MTrPs of the hip muscles should be considered for the
 30 management of hip OA.

31 **Knee Conditions**

32
 33 Mayoral et al. (2013) attempted to determine whether dry needling of MTrPs is superior
 34 to placebo in the prevention of pain after total knee replacement. 40 subjects were
 35 randomized to true dry needling or sham needling. Immediately following anesthesia
 36 and before surgery started, subjects in the treatment group were dry needled in all
 37 previously diagnosed MTrPs, while the sham group received no treatment in their MTrPs.
 38 Subjects were blinded to group allocation as well as the examiner in pre-surgical and
 39 follow-up examinations performed one (1), three (3), and six (6) months after
 40 arthroplasty. Results demonstrated that subjects in the treatment group had less pain after
 41 intervention one (1) month after intervention, indicating the need for immediate post-
 42 surgery analgesics. Differences were not sustained at three (3) and six (6) month follow-

1 up examinations. In conclusion, a single dry needling treatment of MTrP under anesthesia
 2 reduced pain in the first month after knee arthroplasty, when pain was the most severe
 3 (Mayoral et al., 2013). Espí-López et al. (2017) compared the effects of adding TrP DN
 4 to a manual therapy and exercise program on pain, function, and disability in individuals
 5 with PFP. Individuals with PFP ($n = 60$) recruited from a public hospital in Valencia,
 6 Spain were randomized to manual therapy and exercises ($n = 30$) or manual therapy and
 7 exercise plus TrP DN ($n = 30$). Both groups received the same manual therapy and
 8 strengthening exercise program for 3 sessions (once a week for 3 weeks), and 1 group
 9 also received TrP DN to active TrPs within the vastus medialis and vastus lateralis
 10 muscles. The pain subscale of the Knee injury and Osteoarthritis Outcome Score (KOOS;
 11 0-100 scale) was used as the primary outcome. Secondary outcomes included other
 12 subscales of the KOOS, the Knee Society Score, the International Knee Documentation
 13 Committee Subjective Knee Evaluation Form (IKDC), and the numeric pain-rating scale.
 14 Patients were assessed at baseline and at 15-day (posttreatment) and 3-month follow-ups.
 15 At 3 months, 58 subjects (97%) completed the follow-up. No significant between-group
 16 differences (all, $P > .391$) were observed for any outcome. Both groups experienced
 17 similar moderate-to-large within-group improvements in all outcomes (standardized
 18 mean differences of 0.6 to 1.1); however, only the KOOS function in sport and recreation
 19 subscale surpassed the pre-specified minimum important change. Authors concluded that
 20 the current clinical trial suggests that the inclusion of 3 sessions of TrP DN in a manual
 21 therapy and exercise program did not result in improved outcomes for pain and disability
 22 in individuals with PFP at 3-month follow-up.

23
 24 Sánchez Romero et al. (2020) assessed the effectiveness of adding dry needling (DN) to
 25 an exercise program on pain intensity and disability in patients with knee osteoarthritis.
 26 Sixty-two patients with knee osteoarthritis were randomly allocated into one of two
 27 groups: exercise plus DN (exercise + DN; $N = 31$) or exercise plus sham DN (exercise +
 28 sham DN; $N = 31$). Participants received six sessions of either DN or sham DN over the
 29 leg muscles related to knee pain from osteoarthritis plus a supervised exercise program.
 30 Authors concluded that the inclusion of DN to an exercise program does not reduce pain
 31 or disability in patients with knee osteoarthritis.

32 33 **Low Back Pain**

34 Koppenhaver et al. (2015) explored the literature for associations between demographic,
 35 patient history, and physical examination variables and short-term improvement in self-
 36 reported disability following dry needling therapy performed on individuals with low
 37 back pain (LBP). Seventy-two volunteers with mechanical LBP participated in the study.
 38 Potential prognostic factors were collected from baseline questionnaires, patient history,
 39 and physical examination tests. Treatment consisted of dry needling to the lumbar
 40 multifidus muscles bilaterally, administered during a single treatment session.
 41 Improvement was based on percent change on the Oswestry Disability Index at 1 week.
 42 Authors concluded that increased LBP with the multifidus lift test was the strongest

1 predictor of improved disability after dry needling, suggesting that the finding of pain
 2 during muscle contraction should be studied in future dry needling studies. Wang et al.
 3 (2022) investigated the effects of electrical dry needling (DN) plus corticosteroid
 4 injection (CSI) on pain, physical function, and global change in patients with
 5 osteoarthritis of the knee (KOA). Sixty patients with KOA were randomly assigned to the
 6 electrical dry needling plus corticosteroid injection (electrical-DN+CSI) group or CSI
 7 group. The CSI group received glucocorticoid injection only once during the trial, and the
 8 electrical-DN+CSI group received glucocorticoid injection combined with 4 sessions of
 9 electrical-DN. The primary outcome was the numerical rating scale at 3 months. The
 10 secondary outcomes were the Western Ontario and McMaster Universities Osteoarthritis
 11 Index, the time to complete the Timed Up and Go test, and the score of the global rating
 12 of change scale at 3 months. Baseline characteristics and measurements were similar in
 13 the 2 groups. The group by time interaction effect was significant for all variables
 14 ($P<.05$). The electrical-DN+CSI group obtained a more significant reduction in pain
 15 intensity and more significant improvement in dysfunction than the CSI group at 3
 16 months ($P<.05$). The median global rating of change score for the CSI group was +3
 17 (somewhat better), and that for the electrical-DN+CSI group was +4 (moderately better).
 18 Authors concluded that electrical-DN therapy at myofascial trigger points combined with
 19 CSI is more effective at alleviating pain, improving dysfunction, and creating global
 20 change than CSI alone for patients with KOA. Electrical-DN may be an essential part of
 21 treatment for KOA rehabilitation.

22 23 **Heel Pain**

24 Cotchett et al. (2010) reviewed the current evidence for the effectiveness of dry needling
 25 and/or injections of MTrPs associated with plantar heel pain. They included trials where
 26 participants diagnosed with plantar heel pain were treated with dry needling and/or
 27 injections (local anesthetics, steroids, Botulinum toxin A, and saline) alone or in
 28 combination with acupuncture. They determined limited evidence for the effectiveness of
 29 dry needling and/or injections of MTrPs associated with plantar heel pain. However,
 30 given the heterogeneity and poor quality of included studies, definitive conclusions
 31 cannot be made. Cotchett et al. (2014) evaluated the effectiveness of dry needling for
 32 plantar heel pain. Study participants were 84 patients with plantar heel pain of at least 1
 33 month's duration. Participants were randomly assigned to receive real or sham trigger
 34 point dry needling. The intervention consisted of 1 treatment per week for 6 weeks.
 35 Participants were followed for 12 weeks. At the primary end point of 6 weeks, significant
 36 effects favored real dry needling over sham dry needling for pain (adjusted mean
 37 difference: VAS first-step pain= -14.4 mm, 95% CI= -23.5 to -5.2; FHSQ foot pain=10.0
 38 points, 95% CI=1.0 to 19.1), although the between-group difference was lower than the
 39 minimal important difference. The number needed to treat at 6 weeks was 4 (95% CI=2
 40 to 12). The frequency of minor transitory adverse events was significantly greater in the
 41 real dry needling group (70 real dry needling appointments [32%] compared with only 1
 42 sham dry needling appointment [$<1\%$]). Authors concluded that dry needling provided

1 statistically significant reductions in plantar heel pain, but the magnitude of this effect
 2 should be considered against the frequency of minor transitory adverse events. Dunning
 3 et al. (2018) compared the effects of adding electrical dry needling into a program of
 4 manual therapy, exercise and ultrasound on pain, function and related-disability in
 5 individuals with plantar fasciitis (PF). One hundred and eleven participants ($n = 111$)
 6 with plantar fasciitis were randomized to receive electrical dry needling, manual therapy,
 7 exercise and ultrasound ($n = 58$) or manual therapy, exercise and ultrasound ($n = 53$). The
 8 primary outcome was first-step pain in the morning as measured by the Numeric Pain
 9 Rating Scale (NPRS). Secondary outcomes included resting foot pain (NPRS), pain
 10 during activity (NPRS), the Lower Extremity Functional Scale (LEFS), the Foot
 11 Functional Index (FFI), medication intake, and the Global Rating of Change (GROC).
 12 The treatment period was 4 weeks with follow-up assessments at 1 week, 4 weeks, and 3
 13 months after the first treatment session. Both groups received 6 sessions of impairment-
 14 based manual therapy directed to the lower limb, self-stretching of the plantar fascia and
 15 the Achilles tendon, strengthening exercises for the intrinsic muscles of the foot, and
 16 therapeutic ultrasound. In addition, the dry needling group also received 6 sessions of
 17 electrical dry needling using a standardized 8-point protocol for 20 minutes. Authors
 18 concluded that the inclusion of electrical dry needling into a program of manual therapy,
 19 exercise and ultrasound was more effective for improving pain, function and related-
 20 disability than the application of manual therapy, exercise and ultrasound alone in
 21 individuals with PF at mid-term (3 months).

22 **Fibromyalgia**

23 Casanueva et al. (2013) evaluated the short-term efficacy of dry needling for patients
 24 diagnosed with fibromyalgia. 120 patients were randomly selected into two (2) groups
 25 (control and dry needling). Dry needling treatments included weekly one (1) hour
 26 sessions for six (6) weeks. At the end of the treatment, the dry needling group showed
 27 significant differences in most tests, including pain, fatigue SF-36 pain rating, myalgic
 28 scores, PPTs and global subjective improvement. In conclusion, patients severely
 29 affected by fibromyalgia can obtain short-term improvements following weekly dry
 30 needling for six (6) weeks. Castro Sánchez et al. (2019) compared the effectiveness of
 31 dry needling versus myofascial release on myofascial trigger points pain in cervical
 32 muscles, quality of life, impact of symptoms pain, quality of sleep, anxiety, depression,
 33 and fatigue in patients with fibromyalgia syndrome. Sixty-four subjects with
 34 fibromyalgia were randomly assigned to a dry needling group or a myofascial release
 35 group. Pain pressure thresholds of myofascial trigger points were evaluated in the
 36 cervical muscles. In addition, quality of life, impact of fibromyalgia symptoms, quality of
 37 sleep, intensity of pain, anxiety and depression symptoms, impact of fatigue at baseline
 38 and post treatment after four weeks of intervention were evaluated. Authors reported that
 39 dry needling therapy showed higher improvements in comparison with myofascial release
 40 therapy for pain pressure thresholds, the components of quality of life of physical role,
 41 body pain, vitality and social function, as well as the total impact of FMS symptoms,
 42

1 quality of sleep, state and trait anxiety, hospital anxiety-depression, general pain intensity
2 and fatigue. Implications for rehabilitation They concluded that dry needling therapy
3 reduces myofascial trigger point pain in the short term in patients with fibromyalgia
4 syndrome. This therapeutic approach improves anxiety, depression, fatigue symptoms,
5 quality of life, and sleep after treatment. Dry needling and myofascial release therapies
6 decrease intensity of pain, and the impact of fibromyalgia symptoms in this population.
7 These intervention approaches should be considered in an independent manner as
8 complementary therapies within a multidisciplinary setting.

9 **Headache**

10 Gildir et al. (2019) aimed to explore the effectiveness of trigger point dry needling in
11 patients with chronic tension-type headache in reducing headache frequency, intensity
12 and duration, and improvement of health-related quality of life. One hundred sixty
13 participants were randomly assigned to one of two treatment groups for dry needling or
14 sham dry needling, delivered in 3 sessions a week for 2 weeks. The dry needling was
15 applied in active trigger points located in the musculature of the head and the neck. The
16 sham dry needling procedure was applied into the adipose tissue located at any area
17 where an active trigger point was absent. The primary outcome measurement was the
18 headache intensity. In the dry needling group, intensity, frequency and duration of
19 headache, and the scores of Short Form-36 subscales were significantly improved after
20 treatment ($P < .05$). In the dry needling group, all the effect sizes for headache variables
21 were large. Authors concluded that results of this clinical trial suggest that trigger point
22 dry needling in patients with chronic tension-type headache is effective and safe in
23 reducing headache intensity, frequency and duration, and increasing health-related
24 quality of life.

25
26
27 Mousavi-Khatir et al. (2022) compared the long-term effect of adding real or sham dry
28 needling with conventional physiotherapy in cervicogenic headache. Sixty-nine patients
29 with cervicogenic headache were included in this study. Patients were randomly assigned
30 into a control group ($n = 23$) receiving conventional physical therapy; a dry needling
31 group ($n = 23$) receiving conventional physical therapy and dry needling on the cervical
32 muscles; placebo needling group ($n = 23$) receiving conventional physical therapy and
33 superficial dry needling at a point away from the trigger point. The primary outcome was
34 the headache intensity and frequency. Neck disability, deep cervical flexor performance,
35 and range of motion were secondary outcomes. Outcomes were assessed immediately
36 after treatment and 1, 3, and 6 months later. Sixty-five patients were finally included in
37 the analysis. Headache intensity and neck disability decreased significantly more in the
38 dry needling compared to sham and control groups after treatment and during all follow-
39 ups. The frequency of headaches also reduced more in the dry needling than in control
40 and sham groups, but it did not reach statistical significance. Higher cervical range of
41 motion and enhancement of deep cervical flexors performance was also observed in the
42 dry needling compared to sham and control groups. Authors concluded that dry needling

1 has a positive effect on pain and disability reduction, cervical range of motion, and deep
2 cervical flexor muscles performance in patients with cervicogenic headache and active
3 trigger points, although the clinical relevance of the results was small.

4 **Review Articles**

5 **Upper Quadrant MPS**

6 Cummings and White (2001) authored a review article on needling therapies in the
7 management of MTrP pain. Randomized controlled trials (RCTs) in which some form of
8 needling therapy was used to treat MPS were selected for inclusion. A total of 23 papers
9 were chosen based on specific method, quality and outcome parameters. Trials that
10 compared different injectable substances or dry needling to other injectable substances
11 found that the effect was independent of the substance injected, with a dependence upon
12 the actual needling procedure. The review, however, did not find rigorous evidence to
13 confirm that needling therapies have an effect beyond placebo for MTrP pain. Authors do
14 express a caveat being that only one (1) trial identified whether an LTR was noted and as
15 stated earlier, achieving an LTR improves results. Because all groups in which MTrPs
16 were directly needling demonstrated marked improvement, further research is needed to
17 investigate whether needling has an effect beyond placebo. Tough et al., (2009) reviewed
18 the current evidence on needling without injection. They included studies where at least
19 one (1) group were treated by needling directly into the MTrP and where the control was
20 either no treatment, or usual care, indirect local dry needling or some form of placebo
21 intervention. Seven (7) studies were included. One (1) study concluded that direct dry
22 needling was superior to no intervention. Combining these studies ($n=134$), needling was
23 not found to be significantly superior to placebo; however, marked statistical
24 heterogeneity was present. In conclusion, there is limited evidence deriving from one (1)
25 study that deep needling directly into myofascial trigger points has an overall treatment
26 effect when compared with standard care. Limited sample size and poor quality supports
27 the need for improved trials. In 2011, the American Physical Therapy Association
28 (APTA) performed a synthesis and evaluation of the related literature. Based on specified
29 search criteria, 154 articles were identified. Articles were reviewed to determine those
30 appropriate for individual expert review. The remaining 46 individual studies were
31 reviewed by a member expert in research analysis using a standardized review form.
32 These 46 studies were reviewed using a rating scale from 0-5, with 5 indicating the
33 highest level of quality and highest level of support for dry needling. The median quality
34 of the research was 3; the median support of dry needling was 2. Of the 23 RCTs, the
35 median quality of the research was 4; the median support of dry needling was 3.

36
37
38 Kietrys et al. (2013) performed a systematic review and meta-analysis to identify the
39 effectiveness of dry needling in reducing pain for patients with MPS of the upper quarter.
40 Four (4) separate meta-analyses were performed: (1) dry needling compared to sham or
41 control, immediate effects; (2) dry needling compared to sham or control, four (4) weeks;
42 (3) dry needling compared to other treatments, immediate effects; (4) dry needling

1 compared to other treatments, four (4) weeks. Based on the best current available
2 evidence, the authors recommend dry needling, compared to sham or placebo, for
3 decreasing pain (immediately after treatment and at four (4) weeks) in patients with upper
4 quarter MPS. However, due to the small number of high-quality RCTs published to date,
5 additional well-designed studies are needed. Cagnie et al. (2015) described the effects of
6 ischemic compression and dry needling on trigger points in the upper trapezius muscle in
7 patients with neck pain and compare these two interventions with other therapeutic
8 interventions aiming to inactivate trigger points. Fifteen randomized controlled trials
9 were included in this systematic review. There is moderate evidence for ischemic
10 compression and strong evidence for dry needling to have a positive effect on pain
11 intensity. This pain decrease is greater compared with active range of motion exercises
12 (ischemic compression) and no or placebo intervention (ischemic compression and dry
13 needling) but similar to other therapeutic approaches. There is moderate evidence that
14 both ischemic compression and dry needling increase side-bending range of motion, with
15 similar effects compared with lidocaine injection. There is weak evidence regarding its
16 effects on functionality and quality-of-life. Authors reported that based on this systematic
17 review, ischemic compression and dry needling can both be recommended in the
18 treatment of neck pain patients with trigger points in the upper trapezius muscle.
19 Additional research with high-quality study designs is needed to develop more conclusive
20 evidence. Liu et al., (2015) evaluated current evidence of the effectiveness of dry
21 needling of MTrPs associated with neck and shoulder pain. The results suggested that
22 compared with control/sham, dry needling of MTrPs was effective in the short term
23 (immediately to 3 days) and medium term; however, wet needling, when a substance is
24 injected (including lidocaine) was superior to dry needling in relieving MTrP pain in the
25 medium term. Other therapies (including physiotherapy) were more effective than dry
26 needling in treating MTrP pain in the medium term.

27
28 Navarro-Santana et al. (2020) evaluated the effect of dry needling alone as compared to
29 sham needling, no intervention, or other physical interventions applied over trigger points
30 (TrPs) related with neck pain symptoms. Randomized controlled trials including one
31 group receiving dry needling for TrPs associated with neck pain were identified in
32 electronic databases. Outcomes included pain intensity, pain-related disability, pressure
33 pain thresholds, and cervical range of motion. Results demonstrated dry needling reduced
34 pain immediately after and at short-term when compared with sham/placebo/waiting
35 list/other form of dry needling and, also, at short-term compared with manual therapy. No
36 differences in comparison with other physical therapy interventions were observed. An
37 effect on pain-related disability at the short-term was found when comparing dry needling
38 with sham/placebo/waiting list/other form of dry needling but not with manual therapy or
39 other interventions. Dry needling was effective for improving pressure pain thresholds
40 immediately after the intervention. No effect on cervical range of motion of dry needling
41 against either comparative group was found. No between-treatment effect was observed
42 in any outcome at mid-term. Low to moderate evidence suggests that dry needling can be

1 effective for improving pain intensity and pain-related disability in individuals with neck
2 pain symptoms associated with TrPs at the short-term. No significant effects on pressure
3 pain sensitivity or cervical range of motion were observed.
4

5 **Lower Quarter MPS**

6 Morihisa et al. (2016) assessed and provided a summary on the current literature for the
7 use of dry needling as an intervention for lower quarter trigger points in patients with
8 various orthopedic conditions. This review of current literature suggests that dry needling
9 is effective in reducing pain associated with lower quarter trigger points in the short-term.
10 However, the findings suggest that dry needling does not have a positive effect on
11 function, quality of life, depression, range of motion, or strength. Further high-quality
12 research with long-term follow-up investigating the effect of dry needling in comparison
13 to and in conjunction with other interventions is needed to determine the optimal use of
14 dry needling in treating patients with lower quarter trigger points. Khan et al. (2021)
15 explored the current evidence on effects of trigger point dry needling as a treatment
16 strategy on pain and range of motion among subjects with lower extremity myofascial
17 trigger areas. Of the 564 articles initially found 10 (33.3%) were selected for final
18 assessment. All the 10 (100%) studies documented improvement in the pain over time
19 with dry needling strategy. None of the studies targeted any other outcome, like anxiety
20 and sleep disturbances, related with myofascial trigger points. Authors concluded that on
21 the basis of the best evidence available, dry needling seemed to be effective in pain
22 reduction related to lower extremity myofascial trigger points. Evidence also suggested
23 that there was not much positive effect of myofascial trigger point dry needling on
24 depression, anxiety, muscular strength and quality of life.
25

26 **Low Back Pain**

27 In 2005, Furlan et al. updated a systematic review on acupuncture and dry needling for
28 low back pain using the framework of the Cochrane Collaboration. Studies included in
29 this review were RCTs of acupuncture where needling was involved and RCTs of dry
30 needling of adults with non-specific acute, subacute, or chronic low back pain. 35 studies
31 were included for a total of 2861 patients. The majority of these patients experienced
32 chronic low back pain. Two (2) of these studies had fatal flaws and were not included. Of
33 the remaining 33 trials, 14 were of higher quality and 19 of lower methodologic quality.
34 No blinding was done in any of the trials. In 28 trials, similar timing of outcome
35 measurements occurred, but the quality of reporting was variable. This resulted in an
36 inability to judge many aspects of the trials. Limiting discussion to dry needling, efficacy
37 and effectiveness at trigger and motor points shows variable results. Evidence is limited
38 that superficial needling inserted at MTrPs is better than placebo TENS. There is limited
39 evidence that adding dry needling to standard physical therapy, occupational therapy or
40 industrial assessments is better than standard care alone at the short (between one (1)
41 week and three (3) months after end of sessions) and intermediate term follow up
42 (between three (3) months and one (1) year after end of sessions). There is moderate

1 evidence that there is no difference between a session of dry needling and injection of
2 lidocaine and/or steroid. In identifying this data, evidence shows that deep needling is
3 more effective at short term follow up than superficial needling for chronic low back
4 pain. Also, distal point needling is no different from local lumbar area needling for
5 measures of pain, function and ROM. It also appears that needle retention for about 10
6 minutes is better than immediate removal. Some dry needling practitioners have adopted
7 this technique. Authors conclude that although dry needling appears to be a useful
8 adjunct to other therapies for chronic low back pain, no clear recommendations can be
9 made due to poor quality of studies. There is insufficient evidence supporting its use for
10 acute low back pain. They also note that although methodologic quality has improved
11 over the past several years, it is still poor.

12
13 Liu et al. (2018) evaluated the current evidence of the effectiveness of dry needling of
14 myofascial trigger points (MTrPs) associated with low back pain (LBP). A total of 11
15 RCTs involving 802 patients were included in the meta-analysis. Results suggested that
16 compared with other treatments, dry needling of MTrPs was more effective in alleviating
17 the intensity of LBP and functional disability; however, the significant effects of dry
18 needling plus other treatments on pain intensity could be superior to dry needling alone
19 for LBP at post-intervention. Authors concluded that moderate evidence showed that dry
20 needling of MTrPs, especially if associated with other therapies, could be recommended
21 to relieve the intensity of LBP at post-intervention; however, the clinical superiority of
22 dry needling in improving functional disability and its follow-up effects still remain
23 unclear. Hu et al. (2018) evaluated the efficacy and safety of dry needling for treating
24 LBP. Sixteen RCTs were included and the risk of bias assessment of them was “high” or
25 “unclear” for most domains. Meta-analysis results suggested that DN was more effective
26 than acupuncture in alleviating pain intensity and functional disability at postintervention,
27 while its efficacy on pain and disability at follow-up was only equal to acupuncture.
28 However, compared with other treatments (laser, physical therapy, other combined
29 treatments, etc.), it remained uncertain whether the efficacy of DN was superior or equal
30 because the results of included studies were mixed. Authors concluded that compared
31 with acupuncture and sham needling, DN is more effective for alleviating pain and
32 disability at postintervention in LBP, while its effectiveness on pain and disability at
33 follow-up was equal to acupuncture. Besides, it remains uncertain whether the efficacy of
34 DN is superior to other treatments. Nevertheless, considering the overall “high” or
35 “unclear” risk of bias of studies, all current evidence is not robust to draw a firm
36 conclusion regarding the efficacy and safety of DN for LBP. Future RCTs with rigorous
37 methodologies are required to confirm findings.

38
39 Radi et al. (2023) completed an evidence summary on the effectiveness of dry needling
40 for low back pain. They concluded that a comprehensive treatment program that includes
41 dry needling may provide some benefit in decreasing pain scores and perceived disability
42 vs. standard physical therapy (PT) and home PT in the short term. However, this

1 improvement is small, and the clinical significance is questionable. (Strength of
 2 Recommendation: B, randomized controlled trials [RCTs].) Additional research is needed
 3 to determine the best regimens to augment dry needling.

4 5 **Knee Pain**

6 Rahou-El-Bachiri et al. (2020) evaluated the effect of trigger point dry needling alone or
 7 as an adjunct with other interventions on pain and related disability in people with knee
 8 pain. Ten studies (six patellofemoral pain, two knee osteoarthritis, two post-surgery knee
 9 pain) were included. The risk of bias was generally low, but the heterogeneity and the
 10 imprecision of the results downgraded the level of evidence. Authors concluded that low
 11 to moderate evidence suggests a positive effect of trigger point dry needling on pain and
 12 related disability in patellofemoral pain, but not knee osteoarthritis or post-surgery knee
 13 pain, at short-term. More high-quality trials investigating long-term effects are clearly
 14 needed.

15 16 **Shoulder**

17 Hall et al. (2018) completed a systematic review and meta-analysis on patients with upper
 18 extremity pain and dysfunction. Eleven randomized trials involving 496 participants were
 19 appraised. Authors concluded that there is very low evidence to support the use of TDN
 20 in the shoulder region for treating patients with upper extremity pain or dysfunction. Two
 21 studies reported adverse effects to TDN interventions. Most common adverse effects
 22 included bruising, bleeding, and pain during or after treatment. Navarro-Santana et al.
 23 (2021) evaluated the effects of trigger point (TrP) dry needling alone or as an adjunct to
 24 other interventions on pain intensity and related disability in nontraumatic shoulder pain.
 25 The search identified 551 publications with 6 trials eligible for inclusion. Results
 26 demonstrated there was moderate-quality evidence that TrP dry needling reduces
 27 shoulder pain intensity with a small effect and low-quality evidence that TrP dry needling
 28 improves related disability with a large effect compared with a comparison group. The
 29 effects on pain were only found at short term. The Cochrane Risk of Bias was generally
 30 low, but the heterogeneity of the results downgraded the evidence level. Authors
 31 concluded that moderate- to low-quality evidence suggests positive effects of TrP dry
 32 needling for pain intensity (small effect) and pain-related disability (large effect) in
 33 nontraumatic shoulder pain of musculoskeletal origin, mostly at short term.

34
35 Para-García et al. (2022) examined the effects of dry needling alone or in combination
 36 with exercise therapy for reducing pain and disability in people with subacromial pain
 37 syndrome in a systematic review and Meta-Analysis. Five RCTs (n = 315) were included
 38 in the meta-analysis and qualitative analysis. Results determined that dry needling alone
 39 or combined with exercise therapy showed improvements in pain in the short-term and
 40 mid-term compared to a range of interventions. However, no differences were shown for
 41 disability at short-term and mid-term. Dry needling alone or in combination with exercise
 42 therapy may result in a slight reduction in pain in the short-term and mid-term. However,

1 the evidence about the effect of this therapy on disability in the short- or mid-term is very
 2 uncertain compared to the range of interventions analyzed in this systematic review.
 3 Griswold et al. (2023) evaluated the evidence for the effectiveness of various applications
 4 of dry needling (DN) combined with other conservative treatments for subacromial pain
 5 syndrome (SAPS) in a systematic review with meta-analysis. Eight studies were selected.
 6 All eight studies involving 10 comparisons were included in the analyses (N = 538). Dry
 7 needling performed in combination with other conservative interventions produced
 8 favorable outcomes at all time points for pain and disability. Standard mean differences
 9 ranged from -0.57 (moderate) to -1.29 (large) for pain and -0.69 (moderate) to -1.07
 10 (large) for disability, favoring groups receiving DN in addition to conservative treatment.
 11 Four of the eight studies were rated as having unclear or high risk of bias. Authors
 12 concluded that this meta-analysis suggests that various applications of DN performed
 13 with other conservative interventions are more effective than conservative treatment
 14 alone for reducing pain and disability in patients with SAPS. Direct-comparison studies
 15 are needed to determine whether one application of DN is superior to another.

16 **Neck**

17 The Orthopaedic Section of the American Physical Therapy Association (APTA)
 18 published a revision of the neck pain clinical practice guideline (Blanpied et al., 2017).
 19 Authors suggest that for individuals with chronic neck pain with mobility deficits,
 20 clinicians should provide a multimodal approach of the following:
 21

- 22 • Thoracic manipulation and cervical manipulation or mobilization
- 23 • Mixed exercise for cervical/scapulothoracic regions: neuromuscular exercise (e.g.,
 24 coordination, proprioception, and postural training), stretching, strengthening,
 25 endurance training, aerobic conditioning, and cognitive affective elements
- 26 • Dry needling, laser, or intermittent mechanical/manual traction

27
 28 The Royal Dutch Society for Physical Therapy (KNGF) issued a clinical practice
 29 guideline for physical therapists that addresses the assessment and treatment of patients
 30 with nonspecific neck pain, including cervical radiculopathy, in Dutch primary care (Bier
 31 et al., 2018). Recommendations were based on a review of published systematic reviews.
 32 The physical therapist is advised not to use dry needling, low-level laser, electrotherapy,
 33 ultrasound, traction, and/or a cervical collar.

34 **Headache**

35 France et al. (2014) sought to determine the evidence supporting the use of dry needling
 36 in addition to conventional physiotherapy in the management of tension-type and
 37 cervicogenic headache. Only three relevant studies were identified and all three showed
 38 statistically significant improvements following dry needling, but no significant
 39 differences between groups. Only one study reported on headache frequency or intensity,
 40 reporting a 45 mm improvement in VAS score following the addition of dry needling to
 41 conventional physiotherapy. Two studies showed significant improvements with dry
 42

1 needling over 4-5 weeks of treatment. No adverse events were reported. Authors
2 concluded that literature suggests that while there is insufficient evidence to strongly
3 advocate for the use of dry needling, it may be a useful addition to conventional
4 physiotherapy in headache management. Further research with a stronger methodological
5 design is required.

6
7 Pourahmadi et al. (2021) assessed the effectiveness of dry needling on headache pain
8 intensity and related disability in patients with tension-type headache (TTH),
9 cervicogenic headache (CGH), or migraine. Of 2715 identified studies, 11 randomized
10 clinical trials were eligible for qualitative synthesis and 9 for meta-analysis. Only 4 trials
11 were of high quality. Very low-quality evidence suggested that dry needling is not
12 statistically better than other interventions for improving headache pain intensity in the
13 short term in patients with TTH, CGH, or mixed headache (TTH and migraine). Dry
14 needling provided significantly greater improvement in related disability in the short term
15 in patients with TTH and CGH. The synthesis of results showed that dry needling could
16 significantly improve headache frequency, health-related quality of life, trigger point
17 tenderness, and cervical range of motion in TTH and CGH. Authors concluded that dry
18 needling produces similar effects to other interventions for short-term headache pain
19 relief, whereas dry needling seems to be better than other therapies for improvement in
20 related disability in the short term.

21
22 Vázquez-Justes et al. (2022) reviewed the level of evidence for DN in patients with
23 headache. Of a total of 136 studies, they selected 8 randomised clinical trials published
24 between 1994 and 2019, including a total of 577 patients. Two studies evaluated patients
25 with cervicogenic headache, 2 evaluated patients with tension-type headache, one study
26 assessed patients with migraine, and the remaining 3 evaluated patients with mixed-type
27 headache (tension-type headache/migraine). Quality ratings ranged from low (3/10) to
28 high (7/10). The effectiveness of DN was similar to that of the other interventions. DN
29 was associated with significant improvements in functional and sensory outcomes.
30 Authors concluded that dry needling should be considered for the treatment of headache,
31 and may be applied either alone or in combination with pharmacological treatments.

32
33 Kamonseki et al. (2022) systematically reviewed the evidence about the effectiveness of
34 manual therapy (MT) on pain intensity, frequency and impact of headache in individuals
35 with tension-type headache (TTH). Fifteen studies were included with a total sample of
36 1131 individuals. High velocity and low amplitude techniques were not superior to no
37 treatment on reducing pain intensity (low evidence) and frequency (moderate evidence).
38 Soft tissue interventions were superior to no treatment on reducing pain intensity (low
39 evidence) and frequency of pain (low evidence). Dry needling was superior to no
40 treatment on reducing pain intensity (moderate evidence) and frequency (moderate
41 evidence). Soft tissue interventions were not superior to no treatment and other
42 treatments on the impact of headache. Authors concluded that soft tissue interventions

1 and dry needling can be used to improve pain intensity and frequency in patients with
2 tension type headache. High velocity and low amplitude thrust manipulations were not
3 effective for improving pain intensity and frequency in patients with tension type
4 headache.

6 **All Body Regions**

7 Boyles et al. (2015), sought to determine the effectiveness of TDN based on high-quality
8 RCTs for all body regions. The majority of high-quality studies included in this review
9 showed measurable benefit from TDN for MTrPs in multiple body areas, suggesting
10 broad applicability of TDN treatment for multiple muscle groups. Rodríguez- Mansilla et
11 al. (2016) summarized the literature about the effectiveness of dry needling (DN) on
12 relieving pain and increasing range of motion (ROM) in individuals with myofascial pain
13 syndrome (MPS). Authors concluded that DN was less effective on decreasing pain
14 comparing to the placebo group. Other treatments were more effective than DN on
15 reducing pain after 3-4 weeks. However, on increasing ROM, DN was more effective
16 comparing to that of placebo group, but less than other treatments. Gattie et al. (2017)
17 examine the short- and long-term effectiveness of dry needling delivered by a physical
18 therapist for any musculoskeletal pain condition. After screening, 13 were included. Eight
19 meta-analyses were performed. In the immediate to 12-week follow-up period, studies
20 provided evidence that dry needling may decrease pain and increase pressure pain
21 threshold when compared to control/sham or other treatment. At 6 to 12 months, dry
22 needling was favored for decreasing pain, but the treatment effect was not statistically
23 significant. Dry needling, when compared to control/sham treatment, provides a
24 statistically significant effect on functional outcomes, but not when compared to other
25 treatments. Authors concluded that very low-quality to moderate-quality evidence
26 suggests that dry needling performed by physical therapists is more effective than no
27 treatment, sham dry needling, and other treatments for reducing pain and improving
28 pressure pain threshold in patients presenting with musculoskeletal pain in the immediate
29 to 12-week follow-up period. Low-quality evidence suggests superior outcomes with dry
30 needling for functional outcomes when compared to no treatment or sham needling.
31 However, no difference in functional outcomes exists when compared to other physical
32 therapy treatments. Evidence of long-term benefit of dry needling is currently lacking.
33 Espejo-Antúnez et al. (2017) examined the effectiveness of dry needling in the treatment
34 of myofascial trigger points and to explore the impact of specific aspects of the technique
35 on its effectiveness. Fifteen studies were included in this systematic review. The main
36 outcomes that were measured were pain, range of motion, disability, depression and
37 quality of life. The results suggest that dry needling is effective in the short term for pain
38 relief, increase range of motion and improve quality of life when compared to no
39 intervention/sham/placebo. There is insufficient evidence on its effect on disability,
40 analgesic medication intake and sleep quality. Authors state that despite some evidence
41 for a positive effect in the short term, further randomized clinical trials of high

1 methodological quality, using standardized procedures for the application of dry needling
2 are needed.

3
4 Sánchez-Infante et al. (2021) sought to determine the short-, medium-, and long-term
5 effectiveness of dry needling (DN) applied by physical therapists to myofascial trigger
6 points for the treatment of pain via systematic review and meta-analysis. The initial
7 search identified 1771 articles. After the selection, 102 articles were assessed for
8 eligibility; 42 of these articles measuring pain were used for the meta-analysis. Four
9 meta-analyses were performed according to the follow-up period from the last reported
10 treatment. This meta-analysis found a large effect to decrease pain within 72 hours, a
11 moderate effect in 1 to 3 weeks, a large effect in 4 to 12 weeks, and a large effect in 13 to
12 24 weeks. The risk of bias was generally low; however, the heterogeneity of the results
13 downgraded the level of evidence. Authors concluded that low-quality evidence that the
14 immediate to 72-hour (large) effect, 4- to 12-week (large) effect, 13- to 24-week (large)
15 effect, and moderate-quality 1- to 3-week (moderate) effect suggested that DN performed
16 by physical therapists was more effective than no treatment, sham DN, and other
17 therapies for reducing pain.

18
19 Sousa Filho et al. (2021) compared the effects of corticosteroid injection (CSI) and dry
20 needling (DN) for musculoskeletal conditions at short-, medium-, and long-term follow-
21 up. Six studies were included ($n = 384$ participants). Four musculoskeletal conditions
22 were investigated. There is very low-quality evidence that CSI is superior to DN for
23 reducing heel pain (plantar fasciitis) and lateral elbow pain at short- and medium-term
24 follow-up, but not for myofascial pain and greater trochanteric pain. There is very low-
25 quality evidence that DN is more effective than CSI at long-term follow-up for reducing
26 pain in people with plantar fasciitis and lateral epicondylitis. Very low-certainty evidence
27 shows that there is no difference between DN and CSI for disability at short-term follow-
28 up. One study showed that CSI is superior to DN at medium-term follow-up and another
29 observed that DN is superior to CSI for reducing disability at long-term. Authors
30 concluded that there are no differences between DN and CSI in pain or disability for
31 myofascial pain and greater trochanteric pain syndrome. Very-low certainty evidence
32 suggests that CSI is superior to DN at shorter follow-up periods, whereas DN seems to be
33 more effective than CSI at longer follow-up durations for improving pain in plantar
34 fasciitis and lateral epicondylitis. Large RCTs with higher methodological quality are
35 needed in order to draw more incisive conclusions.

36
37 Valera-Calero et al. (2022) investigated the efficacy of dry needling and acupuncture in
38 patients with FM regarding pain, function and disability in both the short and the long
39 term. A total of 25 studies addressed randomized controlled trial studies evaluating
40 efficacy data of dry needling or/and acupuncture treatments to improve pain, fatigue,
41 sleep disturbance and impaired quality of life and/or daily function. Most studies had an
42 acceptable methodological quality. Four studies assessed the effect of dry needling, and

1 twenty-one studies assessed the effect of acupuncture. In general, both interventions
 2 improved pain, anxiety, depression, fatigue, stiffness, quality of sleep and quality of life.
 3 However, both techniques were not compared in any study. Acupuncture and dry
 4 needling therapies seems to be effective in patients with FM, since both reduced pain
 5 pressure thresholds, anxiety, depression, fatigue, sleep disturbances and disability in the
 6 short term. It is still required to compare both techniques and their application in the long
 7 term.

9 **Tendinopathy**

10 Krey et al. (2015) summarized the best available evidence to determine if tendon
 11 needling is an effective treatment for tendinopathy. The studies that were included in this
 12 review suggest that tendon needling improves patient reported outcomes in patients with
 13 tendinopathy. In 2 studies evaluating tendon needling in lateral epicondylitis, one
 14 showed an improvement in a subjective visual analogue scale score of 34% (significant
 15 change > 25%) from baseline at 6 months. The other showed an improvement of 56.1%
 16 in a visual analogue scale score from baseline. In 1 study evaluating tendon needling in
 17 addition to eccentric therapy for Achilles tendinosis, the subjective Victorian Institute of
 18 Sport Assessment-Achilles (VISA-A) score improved by 19.9 (significant change > 10)
 19 (95% CI, 13.6-26.2) from baseline. In 1 study evaluating tendon needling in rotator cuff
 20 tendinosis, the subjective shoulder pain and disability index showed statistically
 21 significant improvement from baseline at 6 months ($P < 0.05$). Authors concluded that
 22 the evidence suggests that tendon needling improves patient-reported outcome measures
 23 in patients with tendinopathy. Stoychev et al. (2020) reviewed the use of dry needling as
 24 a treatment modality for tendinopathy. The effectiveness of dry needling for treatment of
 25 tendinopathy has been evaluated in 3 systematic reviews, 7 randomized controlled trials,
 26 and 6 cohort studies. The following sites were studied: wrist common extensor origin,
 27 patellar tendon, rotator cuff, and tendons around the greater trochanter. There was
 28 considerable heterogeneity of the needling techniques, and the studies were inconsistent
 29 about the therapy used after the procedure. Most systematic reviews and randomized
 30 controlled trials supported the effectiveness of tendon needling. There was a statistically
 31 significant improvement in the patient-reported symptoms in most studies. Some studies
 32 reported an objective improvement assessed by ultrasound. Two studies reported
 33 complications. Authors concluded that current research provides initial support for the
 34 efficacy of dry needling for tendinopathy treatment. In further high-quality studies,
 35 tendon dry needling should be used as an active intervention and compared with
 36 appropriate sham interventions. Studies that compare the different protocols of tendon
 37 dry needling are also needed.

39 Navarro-Santana et al. (2020) evaluated the effect of dry needling alone or combined
 40 with other treatment interventions on pain, related-disability, pressure pain sensitivity,
 41 and strength in people with lateral epicondylalgia of musculoskeletal origin in a meta-
 42 analysis. Seven studies including 320 patients with lateral epicondylalgia were included.

1 Authors concluded that low to moderate evidence suggests a positive effect of dry
 2 needling for pain, pain-related disability, pressure pain sensitivity and strength at short-
 3 term in patients with lateral epicondylalgia of musculoskeletal origin. Jayaseelan et al.
 4 (2021) systematically reviewed the utilization and effects of DN for tendinopathy. After
 5 screening 462 articles, 10 studies met inclusion criteria. Study designs included case
 6 reports, case series, and randomized clinical trials. DN was used in isolation in 3/10
 7 studies and as part of a multimodal approach in 7/10 studies. DN was associated with
 8 improved pain, function, muscle performance and perceived improvement in each study
 9 evaluating the relevant outcome. Authors concluded that DN may be a useful adjunctive
 10 treatment in the conservative management of tendinopathy, although its discrete effect is
 11 unclear. Very low-quality evidence and methodological limitations suggest further
 12 investigation is warranted.

13
 14 Giorgi et al. (2022) summarized the best available evidence on the use of DN and
 15 exercise combined to treat tendinopathy. Seven studies met the inclusion and exclusion
 16 criteria. Current evidence supports the use of DN combined with therapeutic exercises,
 17 especially those including eccentric exercises, can improve pain and function for various
 18 tendinopathies. However, limited evidence exists regarding specific therapeutic
 19 interventions to be combined with DN. Authors concluded that there is moderate, level B
 20 evidence to suggest the use of DN techniques targeted at the tendon and combined with
 21 eccentric therapeutic exercise to improve pain and functional outcomes for
 22 tendinopathies. Nuhmani et al. (2023) evaluated the best available evidence on the
 23 effectiveness of DN in the management of tendinopathy. Seven randomized control trials
 24 were selected. To be included in the current systematic review, the study had to be an
 25 RCT conducted on human participants, which investigated the effect of the DN technique
 26 on the management of tendinopathies. A total of 357 participants were enrolled in the
 27 seven included studies, which were on greater trochanteric pain syndrome, lateral
 28 epicondylitis, supraspinatus tendinopathy and Achilles tendinopathy. DN was compared
 29 with various interventions, including platelet-rich plasma injection, autologous blood
 30 injection and non-steroidal anti-inflammatory medication. All the selected studies
 31 reported a significant positive effect of DN on pain intensity and other outcome
 32 measures, such as patient-specific functional score, disability index, range of motion and
 33 health-related quality of life. Authors concluded that these results indicate that DN
 34 appears to be as effective as other treatment methods at relieving pain and other
 35 symptoms of tendinopathy immediately after treatment and up to 6 months. DN can be
 36 considered among the many options available for the management of tendinopathy.

37 **Heel Pain**

38
 39 He et al. (2017) conducted this meta-analysis to evaluate the effect of MTrP needling in
 40 patients with plantar heel pain. Extensive literature search yielded 1,941 articles, of
 41 which only seven RCTs met the inclusion criteria and were included in this meta-
 42 analysis. Authors determined that MTrP needling effectively reduced the heel pain due to

1 plantar fasciitis. However, considering the potential limitations in this study, more large-
 2 scale, adequately powered, good-quality placebo-controlled trials are needed to provide
 3 more trustworthy evidence in this area. Llurda-Almuzara et al. (2021) evaluated the
 4 effects of dry needling over trigger points associated with plantar heel pain on pain
 5 intensity and related disability or function in a meta-analysis. The search identified 297
 6 publications, with six trials eligible for inclusion. The meta-analysis found low-quality
 7 evidence that trigger point dry needling reduces pain intensity in the short term and
 8 moderate-quality evidence that it improves pain intensity and related disability in the long
 9 term, as compared with a comparison group. The risk of bias of the trials was generally
 10 low, but the heterogeneity of the results downgraded the level of evidence. Authors
 11 concluded that moderate- to low-quality evidence suggests a positive effect of trigger
 12 point dry needling for improving pain intensity and pain-related disability in the short
 13 term and long term, respectively, in patients with plantar heel pain of musculoskeletal
 14 origin. The present results should be considered with caution because of the small
 15 number of trials.

16 **Orofacial Pain**

17 Vier et al. (2019) systematically reviewed the effects of dry needling on orofacial pain of
 18 myofascial origin in patients with temporomandibular joint dysfunction. Seven trials
 19 were considered eligible. There was discrepancy among dry needling treatment protocols.
 20 Meta-analysis showed that dry needling is better than other interventions for pain
 21 intensity as well as than sham therapy on pressure pain threshold, but there is very low-
 22 quality evidence and a small effect size. There were no statistically significant differences
 23 in other outcomes. Authors concluded that clinicians could use dry needling for the
 24 treatment of temporomandibular joint dysfunction, nevertheless, due the low quality of
 25 evidence and high risk of bias of some included studies, larger and low risk of bias trials
 26 are needed to assess the effects of dry needling on orofacial pain associated with
 27 temporomandibular joint dysfunction. Al-Morraissi et al. (2020) completed a network
 28 meta-analysis (NMA) of randomized clinical trials (RCTs) aiming to compare the
 29 treatment outcome of dry needling, acupuncture or wet needling using different
 30 substances in managing myofascial pain of the masticatory muscles (TMD-M). Twenty-
 31 one RCTs involving 959 patients were included. The quality of evidence of the included
 32 studies was low or very low. Authors concluded that based on this NMA, one can
 33 conclude that the effectiveness of needling therapy did not depend on needling type (dry
 34 or wet) or needling substance. This NMA did not provide enough support for any of the
 35 needling therapies for TMD-M.
 36

37 **Spasticity**

38 Bynum et al. (2021) examined existing studies on dry needling for spasticity and range of
 39 motion (ROM) and discusses its potential for use as an occupational therapy intervention.
 40 Authors noted that strong evidence was found to support the use of dry needling to
 41 decrease spasticity and increase ROM. They concluded that this systematic review
 42

1 suggests that dry needling is an effective physical agent modality to decrease spasticity
 2 and increase ROM, both of which are potentially beneficial to functional outcomes.
 3 Fernández-de-Las-Peñas et al. (2021) evaluated the effects of muscle dry needling alone
 4 or combined with other interventions on post-stroke spasticity (muscle tone), related pain,
 5 motor function, and pressure sensitivity. Seven studies (three within the lower extremity,
 6 four in the upper extremity) were included. The meta-analysis found significantly large
 7 effect sizes of dry needling for reducing spasticity, post-stroke pain, and pressure pain
 8 sensitivity as compared with a comparative group at short-term follow-up. The effect on
 9 spasticity was found mainly in the lower extremity at short-term follow-up. No effect on
 10 spasticity was seen at 4 weeks. No significant effect on motor function was observed. The
 11 risk of bias was generally low, but the imprecision of the results downgraded the level of
 12 evidence. Authors concluded that moderate evidence suggests a positive effect of dry
 13 needling on spasticity (muscle tone) in the lower extremity in post-stroke patients. The
 14 effects on related pain and motor function are inconclusive. Valencia-Chulián et al.
 15 (2020) summarized the available evidence about the effectiveness of deep dry needling
 16 (DN) on spasticity, pain-related outcomes, and range-of-movement (ROM) in adults after
 17 stroke. A total of sixteen studies, 7 of which were RCTs, were selected. All studies
 18 generally reported an improvement of spasticity level, pain intensity, and ROM after the
 19 use of DN, alone or combined with other interventions, in stroke survivors. Authors
 20 concluded that the management of adults after stroke with DN may impact positively on
 21 spasticity, pain, and ROM. However, there was significant heterogeneity across trials in
 22 terms of sample size, control groups, treated muscles, and outcome measures, and a meta-
 23 analysis was not feasible.

24 **DRY NEEDLING SAFETY**

26 Serious adverse events are rare with dry needling. Serious events include infection,
 27 internal bleeding, and pneumothorax. Other mild events include nausea, dizziness,
 28 faintness, somato-emotional responses, aggravation of symptoms, bruising, post-needle
 29 soreness, and bleeding. To reduce risk of infection, standard precautions should be
 30 followed by all practitioners. Use of gloves, sterile needles, appropriate needle
 31 placement, skin cleansing, and sharps management are important.

32
 33 Absolute contraindications include:

- 34 • Patient with needle phobia or an unwilling patient due to fear or patient beliefs
- 35 • Inability to give consent — age-related, communication, cognitive
- 36 • History of reaction to needling (or injection) in the past
- 37 • Medical emergency
- 38 • Into a muscle or area in patients on anticoagulant therapy or with
- 39 thrombocytopenia, where hemostasis by palpation cannot be carried out
- 40 appropriately (e.g., psoas, tibialis posterior)
- 41 • Into an area or limb with lymphedema due to increased risk of infection or after
- 42 surgical lymphectomy

1 Relative contraindications or precautions include:

- 2 • Abnormal bleeding tendency
- 3 • Compromised immune system
- 4 • Vascular disease
- 5 • Diabetes
- 6 • Pregnancy
- 7 • Frail patients
- 8 • Epilepsy
- 9 • Medications (e.g., anti-coagulants)
- 10 • Psychological status (e.g., schizophrenic or intoxicated patient)

11
 12 Boyce et al. (2020) reported on the type of adverse events associated with the utilization
 13 of therapeutic dry needling (TDN). Four hundred and twenty physical therapists
 14 participated in this study. Information related to minor and major adverse events that
 15 occurred during 20,464 TDN treatment sessions was collected. Each physical therapist
 16 respondent was asked to fill out two weekly self-reported electronic surveys over a six-
 17 week period. One survey was related to “minor adverse events” (i.e., pain, bleeding,
 18 bruising), while the other was related to “major adverse events” (i.e., pneumothorax,
 19 excessive bleeding, prolonged aggravation). Following the six-week period, descriptive
 20 statistics were used to describe the adverse events (AE) associated with TDN and
 21 calculate the frequencies of those events. A total of 7,531 minor AE’s were reported,
 22 indicating that 36.7% of the reported TDN treatments resulted in a minor AE. The top
 23 three minor AE’s were bleeding (16%), bruising (7.7%), and pain during dry needling
 24 (5.9 %). The average ratio of minor AE’s for all respondents across all weeks was 0.53 or
 25 approximately one event for every two patients. Twenty major AE’s were reported out of
 26 the 20,494 treatments for a rate of <0.1% (1 per 1,024 TDN treatments). No associations
 27 were noted between the frequency of adverse events and the number of patients treated,
 28 practitioner age, level of education, years in practice, level of training or months
 29 experience with dry needling. Authors concluded that expected minor AE’s such as mild
 30 bleeding, bruising, and pain during TDN were common and major AE’s were rare.
 31 Physical therapists and other medical practitioners need to be aware of the risks of TDN.
 32 Based on the findings of this study the overall risk of a major adverse event during TDN
 33 is small.

34 35 **PRACTITIONER SCOPE AND TRAINING**

36 Practitioners should practice only in the areas in which they are competent based on their
 37 education, training and experience. Levels of education, experience, and proficiency may
 38 vary among individual practitioners. It is ethically and legally incumbent on a practitioner
 39 to determine where they have the knowledge and skills necessary to perform such
 40 services and whether the services are within their scope of practice.

1 It is best practice for the practitioner to appropriately render services to a patient only if
 2 they are trained, equally skilled, and adequately competent to deliver a service compared
 3 to others trained to perform the same procedure. If the service would be most
 4 competently delivered by another health care practitioner who has more skill and
 5 training, it would be best practice to refer the patient to the more expert practitioner.

6
 7 Best practice can be defined as a clinical, scientific, or professional technique, method, or
 8 process that is typically evidence-based and consensus driven and is recognized by a
 9 majority of professionals in a particular field as more effective at delivering a particular
 10 outcome than any other practice (Joint Commission International Accreditation Standards
 11 for Hospitals, 2020).

12
 13 Depending on the practitioner’s scope of practice, training, and experience, a member’s
 14 condition and/or symptoms during examination or the course of treatment may indicate
 15 the need for referral to another practitioner or even emergency care. In such cases it is
 16 prudent for the practitioner to refer the member for appropriate co-management (e.g., to
 17 their primary care physician) or if immediate emergency care is warranted, to contact 911
 18 as appropriate. See the *Managing Medical Emergencies (CPG 159 – S)* clinical practice
 19 guideline for information.

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