Review Article

Evidence of Physiotherapy Interventions for Patients with Chronic Neck Pain: A Systematic Review of Randomised Controlled Trials

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Chronic neck pain (CNP) is common and costly, and the effect of physiotherapeutic interventions on the condition is unclear. We reviewed the literature for evidence of effect of physiotherapy interventions on patients with CNP. Five bibliographic databases (MEDLINE, EMBASE, CINAHL, Cochrane Library, and PEDro) were systematically searched. Randomised, placebo and active-treatment-controlled trials including physiotherapy interventions for adults with CNP were selected. Data were extracted primary outcome was pain. Risk of bias was appraised. Effect of an intervention was assessed, weighted to risk of bias. 42 trials reporting on randomised comparisons of various physiotherapy interventions and control conditions were eligible for inclusion involving 3919 patients with CNP. Out of these, 23 were unclear or at high risk of bias, and their results were considered moderate- or low-quality evidence. Nineteen were at low risk of bias, and here eight trials found effect on pain of a physiotherapy intervention. Only exercise therapy, focusing on strength and endurance training, and multimodal physiotherapy, cognitive-behavioural interventions, massage, manipulations, laser therapy, and to some extent also TNS appear to have an effect on CNP. However, sufficient evidence for application of a specific physiotherapy modality or aiming at a specific patient subgroup is not available.

1. Introduction

Musculoskeletal disorders are threatening quality of life by having the potential to restrict daily activities, cause absence from work, and result in a change or discontinuation in employment. These disorders are expensive for society and for patients and are responsible for the highest number of healthy years lost [1–4]. The prevalence of chronic neck pain varies. The 12-month prevalence of pain typically ranges between 30% and 50%; the 12-month prevalence of activity-limiting pain is 1.7% to 11.5% [5]. The annual incidence of neck pain associated with whiplash varies greatly. Although 50% of whiplash victims recover in three to six months, 30% to 40% have persisting mild to moderate pain and 10% to 20% retain more severe pain [6]. It is a multifaceted phenomenon with physical impairment, psychological distress, and social dysfunction, which calls for an evidence-based, cost-effective rehabilitation treatment [7–11].

According to a Dutch study, 44% of patients with chronic neck pain visited their general practitioner (GP) with the condition during a twelve-month period; 51% of these were referred to physiotherapy treatment [12]. Knowledge of the actual effect of physiotherapy is therefore important and is anticipated to be reflected in the awareness of evidence-based practice among physiotherapists.

The Cochrane Collaboration has provided systematic reviews on the effect of massage for mechanical neck disorders [13], patient education for neck pain [14], electrotherapy
for neck pain [15], mechanical traction for neck pain with or without radiculopathy [16], and conservative treatment for whiplash [17]. The overall conclusion has been that the evidence for these treatments is low and that no definite statements on the efficacy and clinical usefulness of these treatments can be made. A further Cochrane Review on the effect of manipulation and mobilisation of neck pain found low quality evidence that cervical and thoracic manipulations may provide pain reduction [18]. An additional Cochrane Review on the effect of exercises for mechanical neck disorders concluded that the summarised evidence indicates that there is a role for exercises in the treatment of acute and chronic mechanical neck pain plus headache but that the relative benefit of each type of exercise needs extensive research [19].

However, none of these reviews have covered the majority of commonly used physiotherapy modalities in one in order to get an overview of the subject. Besides, the effect of specific physiotherapy treatments in specific subgroups of chronic pain patients is an important topic which has not yet been examined. Clinicians and policy makers need evidence from research to inform and guide clinical practice and policy. Patients and researchers also need such information to support shared decisions and to set priorities for future research.

The aim of this study was to review the literature systematically and discuss the quality of evidence of commonly used physiotherapy interventions (exercise, manual therapy, and electrotherapy) aimed at improving outcomes (on pain, function, and quality of life) important for patients with chronic neck pain [20]. Neck pain was defined as pain located in the anatomical region of the neck [21]. Pain was considered chronic if it had persisted for more than three months, as defined by the International Association of the Study of Pain.

2. Methods

We performed a systematic review of all available randomised controlled trials on the subject of physiotherapy for neck pain to determine the effects of physiotherapy interventions on pain, function, and quality of life in neck-pain patients and to explore whether beneficial effects could be explained by biases affecting individual trials [22]. Study selection, assessment of eligibility criteria, and data extraction were carried out based on a predefined, peer-reviewed protocol according to the Cochrane Collaboration's guidelines [23]. This paper was prepared in accordance with the PRISMA statement [24].

2.1. Literature Search. We searched five bibliographic databases (MEDLINE, EMBASE, CINAHL, Cochrane Library, and PEDro) from January 1990 to January 2012 with a structured, pre-defined, search strategy [25]. The search strategy was “Neck Pain AND Physiotherapy Intervention.” For neck pain, the following terms were combined with OR: “whiplash/WAD,” “neck injury,” “neck sprain/strain,” “neck ache,” “cervical sprain/strain,” “cervical disorder/syndrome,” “cervical spondylosis/itis,” “cervical osteoarthritis,” “cervicodynia,” “cervicobrachial pain/disorder/syndrome,” “myofascial pain/disorder/syndrome,” “trapezius myalgia,” “postural syndrome,” and “nonspecific neck pain.” For physiotherapy interventions, the following terms were combined with OR: “physiotherapy,” “physical therapy,” “rehabilitation,” “intervention studies,” “exercise,” “exercise therapy,” “exercise movement techniques,” “manual therapy,” “manipulative medicine,” “mobilisation/mobilization,” “musculoskeletal techniques,” and “electric/electro stimulation therapy.” All terms were searched as free text as well as keywords, where this was applicable. Limitations were human studies in the English, German, Dutch, Danish, Norwegian, and Swedish languages, in the time span of January 1990 to January 2012. To assure that the included studies followed scientifically sound methods and the data therefore were well documented, we set a limit for inclusion to publications from 1990 and onwards.

Reference lists of review articles and included studies were searched to identify other potentially eligible studies. An additional search was conducted via the scientific search machine http://www.scirus.com/, using the following search terms combined with AND: “chronic neck pain,” “physiotherapy.”

2.2. Selection Criteria. Studies were included if participants were older than 18 years of age and had chronic neck pain for more than three months (therefore considered chronic). Chronic neck pain was defined as (i) chronic whiplash-associated disorders (WAD); (ii) chronic non-specific neck pain, including work-related neck pain, myofascial neck pain, upper trapezius myalgia, chronic neck pain associated with degenerative findings with or without radicular findings, or other surrogate terms.

Eligible interventions were physiotherapy interventions commonly used in the treatment of musculoskeletal pain: (i) exercise therapy, including specific types of exercises, for example, neuromuscular training, strength training, and endurance training; (ii) manual therapy, for example, massage, manipulations, and mobilisations; (iii) electrotherapy, for example, TENS, low-level laser, or other surrogate terms. Acupuncture was not considered a physiotherapy technique since this technique is not part of physiotherapy in all countries. Comparison of the therapy had to be made with no treatment (e.g., waiting list controls), or other conservative active therapies called “care as usual,” or sham therapy. Anticipating that only a limited number of trials available used placebo/sham control, we decided also to include trials in which an active control was used as a cointervention.

To be eligible for inclusion, a study must apply at least one pain measurement prior to and following the intervention, which was an outcome considered to be of major importance to the patients. Self-reported function and disability [26], self-reported quality of life [27], objective physical function, and clinical tests were considered minor outcomes and therefore not considered necessary inclusion criteria [28–30]. Only randomised controlled trials were accepted. Exclusion criteria were studies with participants with acute or subacute...
Records identified by search strategy
\( (n = 1236) \)

Records for further assessment
\( (n = 760) \)

Full-text articles assessed for eligibility
\( (n = 151) \)

Studies included in qualitative synthesis
\( (n = 42) \)

Records excluded
\( (n = 476; 303 \) not fulfilling inclusion criteria + 173 reviews or guidelines)\)

Records excluded
\( (n = 609; 593 \) not fulfilling inclusion criteria + 16 parallel publications)\)

Full-text articles not fulfilling inclusion criteria
\( (n = 109; 11 \) not randomised/groups not comparative + 41 no chronic condition + 16 neck pain that could not be separated from data on other patient groups + 10 not commonly used physiotherapy interventions + 2 intervention not reported + 2 primary outcome not available + 4 pilot studies + 2 conference papers + 2 dissertations + 1 test sensitivity study + 18 parallel publications)\)

Figure 1: Flow diagram of the selection process of included studies.

neck pain, neck pain with definite or possible long tract signs, neck pain due to specific pathological conditions (e.g., fractures, tumours, infections, inflammatory processes, ankylosing spondylitis, and rheumatoid arthritis), and headache.

We created a reliable process through consequently two reviewers who independently conducted the study selection and assessment of eligibility criteria. Similarly, two reviewers independently conducted data abstraction and assessed the risk of bias. Disagreements were resolved through consensus with a third reviewer being consulted if there was disagreement.

2.3. Data Extraction and Evidence Synthesis. Data regarding publication status, trial design, patient characteristics, treatment regimens, outcome methods, results, and funding were extracted on a standardised form using a custom-made Microsoft Excel spreadsheet.

We assessed the risk of bias by using the Cochrane Collaboration’s tool for assessing risk of bias as presented in [23]. Each of the following domains would be considered adequate—that is, presumably with a low risk of bias (i) “adequate sequence generation”; (ii) “allocation concealment”; (iii) “blinding”; (iv) “incomplete outcome data addressed”; (v) “free of selective outcome reporting”; (vi) “free of other bias (i.e., whether a study sponsor would benefit economically from a positive outcome).” Each of these key components of methodological quality was assessed on an Adequate/Unclear/Inadequate basis. We used The Cochrane Collaboration’s approach for summary assessments of the risk of bias for each important outcome across domains within a trial [23].

Due to the limited number of studies investigating each of the specific interventions, it was decided that both meta-analytical and level of evidence approaches would be inappropriate. Therefore, a narrative approach where we evaluated the study and results between groups within a trial was used to summarise the findings. To formulate conclusions, only results from trials at low risk were considered as evidence for an intervention.

3. Results

The literature search identified 4921 relevant studies (1110 from EMBASE, 1568 from MEDLINE, 1239 from CINAHL, and 491 from PEDro), of which 3685 were duplicates, leaving 1236 potentially eligible studies to be screened (see Figure 1). Following screening of titles and abstracts, 151 potentially relevant studies were identified and retrieved in full text. Finally, 42 randomised controlled trials, involving 3919 patients, fulfilled the selection criteria and were considered suitable for inclusion. The selection process and reasons for exclusions are presented in Figure 1.

3.1. Study Characteristics. Study characteristics and study results are presented under the categories exercise therapy (25 trials, 18 regarding chronic non-specific neck pain, and seven regarding chronic neck pain related to whiplash); manual therapy (six trials, all related to chronic non-specific neck pain); and electrotherapy (11 trials, all related to chronic non-specific neck pain) in Appendix A, Tables 1–4.
The trials covered the following intervention topics: (i) exercise therapy: various types of dynamic and isometric exercises, general aerobic exercises, exercises with a focus on strength, endurance, proprioception and coordination, specific neck stabilising exercises, cranio-cervical-flexion exercises, posture, behavioural graded activity, relaxation, body awareness, myo-feedback training, and multimodal physiotherapy; (ii) manual therapy: massage, manipulation, and traction; (iii) electrotherapy: laser, transcutaneous nerve stimulation (TENS), ultrasound, and repetitive magnetic stimulation (rMS).

Sham therapy or waiting list controls were used as control groups in 12 trials; 10 trials used a control group consisting of a self-management book, health-counselling, or other interventions, clearly distinguished from the active intervention group; six trials used active-treatment control reported as “treatment as usual”; active-treatment control was used in 14 trials.

Primary outcome measures were self-reported pain and/or self-reported pain and disability in 41 trials; when primary outcome measures were not reported, all outcome measures were considered. One trial had an objective test as primary outcome, yet pain was included in the secondary outcome measures.

3.2 Risk of Bias. Risk of bias is presented in Appendix B, Table 5.

Overall, the quality of reporting on methodological issues varied. Table 5 shows the judgements ("Adequate," "Unclear," and "Inadequate") for each of the domains. As can be seen, 28 of 42 trials succeeded in reporting on adequate sequence generation; 18 trials described adequate allocation concealment; four trials adequately reported on attempts to blind participants, personnel, and outcome assessors; 22 trials adequately reported on missing outcome data, using intention-to-treat analysis; three trials adequately reported on selective outcome reporting by referring to a published and available protocol for comparisons; and 25 trials adequately reported on funding and the role of funding.

The summary assessment of risk of bias revealed 19 trials at low risk of bias [34–39, 45, 47–49, 51, 55–60, 63, 64] and 23 trials as unclear or at high risk of bias [31–33, 40–44, 46, 50, 52–54, 61, 62, 65–72], and for this reason their results were not considered as evidence. Of these 19 trials at low risk of bias, 11 trials found no difference between intervention groups [34, 36–39, 45, 48, 49, 56, 58, 60], and eight trials found an effect on pain of the intervention [35, 47, 51, 55, 57, 59, 63, 64].

All studies are described in detail in Appendix A, Tables 1–4. All trials at low risk of bias, showing an effect on pain, are, furthermore, presented in the following section. According to the described criteria, the evidence for each intervention will following be summarised at the end of each section.

3.3 Effect of Physiotherapy Interventions

3.3.1 Exercise

Effect of Exercise on Pain in Patients with Chronic Nonspecific Neck Pain. As shown in Appendix A, Table 1, 18 trials examined the effect of various types of exercise in patients with chronic neck pain; nine of these were at unclear or high risk of bias [31–33, 40–44, 46], and nine were at low risk of bias [34–39, 45, 47, 48]. Seven of the trials at low risk examined the effect of different types of exercise, including proprioception exercises (eye-head coordination), cranio-cervical flexion exercises (C-CF), neck stabilisation exercises, stretching, strengthening, and behavioural graded activity programme, but did not find statistically significant difference on pain between groups following intervention [34, 36–39, 45, 48]. Two of the trials at low risk of bias succeeded in finding an effect on pain from the intervention, and for this reason, their results were considered evidence of use of exercise.

(1) Gustavsson et al. [35] examined a multicomponent pain and stress self-management group intervention (PASS) versus a control group receiving individually administered physiotherapy (IAPT). There was a statistically significant effect on ability to control pain ($P < 0.001$) and on neck-related disability (NDI) ($P < 0.001$) in favour of PASS at the 20-week followup.

(2) Ylinen et al. [47] examined three interventions: intensive isometric strength training versus lighter endurance training versus a control group. The two training groups had an additional 12-day institutional rehabilitation programme. At the 12-month followup, both neck pain and disability had decreased in both training groups compared with the control group ($P < 0.01$).

No trials with low risk of bias supported single use of proprioception exercises (eye-head co-ordination), cranio-cervical flexion exercises (C-CF), or neck stabilisation exercises for pain. No trials with low risk of bias support the use of stretching.

Effect of Exercise on Pain in Patients with Chronic Whiplash-Associated Disorder. As shown in Appendix A, Table 2, seven trials examined the effect of various types of exercise in patients with chronic WAD; three of these were at low risk of bias [49, 51, 55], and four were at unclear or high risk of bias [50, 52–54]. One of the trials at low risk of bias examined the effect of adding biofeedback training to a rehabilitation programme, but found no difference in effect between groups [49]. Two trials at low risk of bias succeeded in finding an effect on pain from the intervention, and for this reason, their results were considered evidence of use of exercise.

(1) Jull et al. [51] examined a multimodal physiotherapy programme (including exercises, education, and ergonomics) versus a self-management programme. The multimodal physiotherapy programme group attained a statistically significant greater reduction in reported neck pain and disability (NDI) posttreatment ($P = 0.04$).

(2) Stewart et al. [55] examined exercise (e.g., endurance, strength, aerobic, coordination, and cognitive behavioural therapy) versus advice alone. Exercise and advice were more effective than advice alone at
| Author            | Participants                                                                 | Interventions                                                                 | Main outcome measures | Study results on effect* of intervention on pain                                                                 |
|-------------------|-------------------------------------------------------------------------------|-------------------------------------------------------------------------------|-----------------------|---------------------------------------------------------------------------------------------------------------|
| Cunha et al. [31] | Women, aged 35–60, with diagnosed primary mechanical myogenous or arthrogenous, neck pain lasting >12 wks (N = 33) | (1) GPR group (n = 15), manual therapy for stretching fasciae for 30 min, muscle stretching in the form of global posture reeducation (GPR) for 30 min  
(2) Conventional stretching group (n = 16), manual therapy for stretching fasciae for 30 min, muscle stretching through conventional stretching exercises for 30 min  
All: two weekly physiotherapy sessions during a 6 wk period | VAS, ROM, SF-36 | There were no statistically significant differences in effect between groups after treatment and at 6 wk followup |
| Dellve et al. [32] | Women, aged 35–60, with work disability (at least 50%) and pain in the neck (diagnosed cervicobrachial pain syndrome) for at least 1 year (N = 60) | (1) Myofeedback training (n = 20), min 8 hours/wk, registered the muscle activity (EMG) of upper trapezius muscles and gave alarm if the preset level of muscular rest was not reached.  
Personal visit once/wk from a physiotherapist browsing EMG profiles with reference to diary entries  
(2) Intensive muscular strength training (n = 20), a structured 5–10 min program to be performed twice a day for 6 days/wk. A physiotherapist coached by two personal visits and additional phone calls twice/wk  
(3) Control group (n = 20)  
All: kept a diary 6 days/wk recording activities, discomfort, pain, and sleeping disturbances. All interventions lasted 1 mth | Work ability index (WAI)  
Single item on work ability, working degree, changed work ability  
Pain, NRS  
Copenhagen Psychosocial Questionnaire  
Cutlery wiping performance test, dexterity, max. grip strength | There were no statistically significant differences in effect between groups after 1 mth and at followup after 3 mths |
| Falla et al. [33] | Patients with chronic nonsevere neck pain (>3 mths), score < 16 (out of possible 50) in NDI (N = 58) | (1) Endurance-strength training of the cervical spine flexor muscles (n = 29)  
(2) Referent exercise intervention, low-load craniocervical exercise (n = 29)  
All: instruction and supervision once a wk for 6 wk, supplied with an exercise diary | EMG measures of maximum voluntary contraction force of sternocleidomastoid and anterior scalene muscle, NRS**, NDI** | There were no statistically significant differences between groups for change in pain (NRS) or disability (NDI) measured in the week immediately after intervention (week 7) |
| Griffiths et al. [34] | Chronic neck pain (diagnosed spondylosis, whiplash, nonspecific neck pain, and discogenic pain), age 18 and over (N = 74) | (1) Specific neck stabilisation exercises (n = 37) in addition to the same programme as group 2  
(2) General neck exercise programme (n = 37), posture correction technique, and active range of movement exercise  
All: max. four 30 min treatment sessions within the first 6 wks, advice to perform exercises 5–10 times daily, written sheets, after 6 wks the therapist could discharge the patient or continue | NPDS, NPQ, VAS** | There were no significant between-group differences in the NPDS at either 6 wks or 6 mths |
Table 1: Continued.

| Author                  | Participants                                                                 | Interventions                                                                                                                                  | Main outcome measures                                                                                                                   | Study results on effect* of intervention on pain                                                                                     |
|-------------------------|------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------|
| Gustavsson et al. [35]  | Patients with musculoskeletal tension-type neck pain of persistent duration (>3 mths), age 18–65 (N = 156) | (1) Multicomponent pain and stress self-management group intervention (PASS) (n = 77), relaxation training, body awareness exercises, lectures and group discussions, seven 1.5 h sessions over a 7 wk period, and a booster session after 20 wks  
(2) Control group receiving individually administered physiotherapy (IAPT) (n = 79) | Questionnaire comprising the self-efficacy scale, NDI, coping strategies questionnaire, hospital and depression scale, fear-avoidance beliefs questionnaire, and questions regarding neck pain, analgesics, and utilisation of health care | There was a statistically significant effect on ability to control pain (P < 0.001), and on neck related disability (NDI) (P < 0.001) in favour of PASS at the 20 wks followup |
| Håkkinen et al. [36]    | Nonspecific neck pain of more than 6 mths, age 25–53, pain > 29 mm on VAS (N = 101) | (1) Strength training and stretching (n = 49). Sessions once a wk for 6 wks and thereafter one session every second mth for 12 mths  
(2) Stretching group (n = 52) in a single group session instructions  
All: encouraged to perform home training regimen three times a wk and to keep weekly exercise diary | VAS, neck and shoulder disability index, NDI, ROM, isometric strength | There were no statistically significant differences in effect between groups after two and 12 months measured with VAS and NDI |
| Jordan et al. [37]      | Patients with chronic neck pain (>3 mths), nonradicular extremity pain was permitted, age 20–60 (N = 119) | (1) Intensive training of the neck and shoulder musculature (n = 40)  
(2) Individual physiotherapy treatment (n = 39)  
(3) High-velocity, low-amplitude spinal manipulation performed by a chiropractor (n = 40)  
All: above training/treatment sessions twice a wk for 6 wks, besides a single neck school group session | Self-reported disability and pain on 11-point box scales, medication use, patients perceived effect, physicians global assessment | There were no statistically significant differences in effect between groups at 4 and 12 mths followup |
| Jull et al. [38]        | Females with chronic neck pain of idiopathic or traumatic origin and abnormal measures of joint position sense (N = 64) | (1) Proprioceptive exercise intervention (n = 28)  
(2) Cranio cervical spine flexion exercise intervention (n = 30)  
All: personal instruction and supervision once a wk for 6 wks | Joint position error, NDI, NRS | There were no statistically significant differences in effect between groups measured in the week immediately after intervention (week 7) |
| Jull et al. [39]        | Females with chronic, nonsevere neck pain (>3 mths), score < 15/50 on NDI (N = 46) | (1) Cranio cervical spine flexion training (n = 23), low load  
(2) Strength training (n = 23)  
All: personal instruction and supervision once a wk for 6 wks | (NDI, NRS)*,  
EMG amplitude of deep cervical spine flexor muscles, sternocleidomastoid and anterior scalene muscle and ROM | There were no statistically significant differences in effect between groups measured in the week immediately after intervention (week 7) |
| Author                        | Participants                                                                 | Interventions                                                                                   | Main outcome measures                                                                 | Study results on effect* of intervention on pain                                      |
|-------------------------------|-------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|
| O’Leary et al. [40]           | Females with chronic neck pain (>3 mths), having in the higher end of mild to moderate pain and disability, score > 4/50 on NDI (N = 48) | (1) Cranio-cervical spine flexion coordination exercise (CCF) (n = 24)  
(2) Cervical spine flexion endurance exercise (CF) (n = 24)  
All: one experimental session | VAS                                                                                          | There were no statistically significant differences between groups on VAS                |
| Randlov et al. [41]           | Females with chronic neck/shoulder pain (>6 mths), age 18–65 (N = 77)        | (1) Light training (n = 41)  
(2) Intensive training (n = 36)  
All: three times per wk, in total 36 sessions | Pain measures with two 11-point box scales, activities of daily living, strength, endurance | There were no statistically significant differences in effect between groups after six and twelve mths followup |
| Revel et al. [42]             | Patients with chronic neck pain (>3 mths), age > 15 (N = 60)                 | (1) Rehabilitation group (n = 30), receiving common symptomatic treatment, besides eye-head exercises improving neck proprioception in individual exercise sessions twice a wk for 8 wks  
(2) Control group (n = 30), receiving only symptomatic treatment without rehabilitation | Head repositioning accuracy, VAS, medication intake, ROM                                | Significant difference between groups for the rehabilitation group on VAS pain (~21.8 ± 25.2) (P = 0.04) at 10 wk followup |
| Taimela et al. [43]           | Patients with chronic, nonspecific neck pain (>3 mths), half had local pain and half referred pain below the elbow, age 30–60 (N = 76) | (1) Active treatment (n = 25), proprioceptive exercises, relaxation and behavioural support, 24 sessions  
(2) Home regimen (n = 25), neck lecture and two sessions of practical training for home exercises and instructions for maintaining a diary  
(3) Control group (n = 26), a lecture regarding care of the neck with a recommendation to exercise | VAS, ROM, PPT                                                                               | The VAS scores after the intervention at 3 mths were significantly lower in the active treatment (22 mm) and home regimen (23 mm) groups than in the control group (39 mm) (P = 0.018) after 3 mths. No statistically significant differences between the groups were noted at 12 mths |
| Viljanen et al. [44]          | Female office workers with chronic non-specific neck pain (>12 wks), age 30–60 (N = 393) | (1) Dynamic muscle training (n = 135)  
(2) Relaxation training (n = 128)  
(3) Control group, ordinary activity (n = 130)  
Groups 1 and 2 were instructed and trained 3 times a wk for 12 wks followed by one wk of reinforcement 6 mths after randomisation | Pain rated on a scale 0 (no pain)–10 (unbearable pain), pain questionnaire | There were no statistically significant differences in effect between groups at 3, 6, and 12 mths followup |
| Vonk et al. [45]              | Patients with chronic non-specific neck pain (>3 mths), age 18–70 (N = 139)  | (1) Behaviour graded activity programme (n = 68), biopsychosocial model guided by the patient's functional abilities  
(2) Conventional exercise (n = 71), reflected usual care, exercises, massage and mobilisation and traction  
All: treatment period 9 wks | Global perceived effect, NDI, NRS                                                          | There were no statistically significant differences in effect between groups at 4, 9, 26, and 52 wks |
Table 1: Continued.

| Author          | Participants                                                                 | Interventions                                                                                           | Main outcome measures                                                                 | Study results on effect* of intervention on pain                                      |
|-----------------|------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|
| Waling et al. [46] | Women with chronic work-related trapezius myalgia (>1 ye), not on sick leave more than 1 mth during last year, age < 45 ye (N = 103) | (1) Strength training group (n = 29), loaded to allow 12 rep. maximum (RM) (2) Endurance training group (n = 28), arm-cycling intensity light (11)—somewhat hard (13) on RPE alternating with exercises loaded to 30–35 RM (3) Coordination training (n = 25). Body-awareness therapy and training. (4) Control group: nontraining. Group stress and bodily reactions due to stress were studied. Two-hour sessions once a wk for 10 wks Groups 1–3: one-hour sessions, three times a wk for 10 wks | VAS, three scales: pain-in-general, pain-at-worst, pain-at-present. Pain threshold     | Significant effect of strength training and endurance training VAS pain-at-worst after 10 wks (P < 0.05). But no difference on VAS pain-at-present or at VAS pain-at-general |
| Ylinen et al. [47] | Female office worker, age 25–53, with constant or frequently occurring neck pain of more than 6 mths. Motivated to continue working and rehabilitation (N = 180) | (1) Endurance group (n = 60), endurance training, dynamic neck exercises (2) Strength group (n = 60), strength training, high-intensity isometric neck strengthening and stabilisation exercises Groups 1 and 2: 12-day institutional rehabilitation programme with training lessons, behavioural support, 4 sessions of physical manual therapy, advice to continue exercise 3 times a wk at home (3) Control group (n = 60): 3-day institutional rehabilitation programme with recreational activities All: advice to perform aerobic exercise 3 times a wk for half an hour at home. | VAS, neck and shoulder pain and disability index, vernon neck disability index         | At the 12 mth followup, both neck pain and disability had decreased in both training groups compared with the control group (P < 0.01). Decrease Pain VAS in the endurance group: −35 ((−42)–(−28)); in the strength group: −40 ((−48)–(−32)) |
| Ylinen et al. [48] | Female, age 25–53, with constant or frequently occurring neck pain of more than 6 mths duration, pain > 44 mm on VAS (N = 125) | Crossover trial, after 4 wks (1) Manual therapy group (n = 62), low-velocity osteopathic-type mobilisation of cervical joints, traditional massage, passive stretching, two treatments a wk for 4 wks (2) Stretching exercises group (n = 63) consisted of instruction to perform neck stretching exercises at home for 4 wks | VAS, neck and shoulder pain and disability index, NDI,                                 | There were no statistically significant differences in effect between groups at the one- and three-year followup |

* In order to show an effect of an intervention and hereby support the intervention, it requires showing statistical significant difference between groups.
** Secondary outcome measure.

VAS: visual analogue scale; NRS: numerical rating scale; VNPS: verbal numeric pain scale; NPQ: Northwick Park neck pain questionnaire; NDI: neck disability index; NPDI: neck pain and disability index; NPDS: neck pain and disability scale; NPDVAS: neck pain and disability visual analogue scale; PSFS: patient specific functional scale; NPI: Northwick Park neck pain index; SF-36: short-form 36; PPT: pressure pain threshold; ROM: range of movement; RPE: rating of perceived exertion; EMG: electromyographic, HRQoL: health-related quality of life.
| Author                  | Participants                                                                 | Interventions                                                                 | Main outcome measures                                                      | Study results on effect* of intervention on pain                          |
|-------------------------|-------------------------------------------------------------------------------|-------------------------------------------------------------------------------|----------------------------------------------------------------------------|----------------------------------------------------------------------------|
| Ehrenborg and Archenholtz [49] | Patients, aged 17–58, with pain after whiplash injury (>3 mths), and referred to the pain unit for outpatient-based, interdisciplinary rehabilitation (N = 65) | (1) Biofeedback training (n = 36), eight sessions (twice/wk for four wk) while being active in a self-chosen handicraft. (2) Being active in a self-chosen handicraft on the same terms as group 1 but without biofeedback (n = 29). All: 4–6 wk rehabilitation programme consisting of a combination of education, ergonomic interventions, physical training, relaxation techniques, body awareness training, and interventions by psychologist and/or social worker if needed | Canadian occupational performance measure, Multidimensional Pain Inventory, Swedish version | There were no statistically significant differences in effect between groups at 6 mths followup |
| Fitz-Ritson [50]        | Patients with chronic pain in cervical spine musculature following motor vehicle accident (WAD), age 19–57, still having symptoms after receiving chiropractic treatments and rehabilitation exercises for >12 mths (N = 30) | (1) Continued chiropractic treatments and standard rehabilitation exercises (n = 15) (2) Continued chiropractic treatments and were advised to do “phasic neck exercises” (eye-head co-ordination) (n = 15) All: exercises 5 days a wk for 8 wks | NPI, VAS**                                                                 | The authors do not report any data on statistically significant differences between groups after 8 wk |
| Jull et al. [51]        | Patients with chronic whiplash-associated disorder (>3 mths, <2 yrs), classified WADII, age 18–65 (N = 71) | (1) Multimodal physiotherapy programme (MPT) (n = 36), low-load exercise for reeducating muscle control of the neck flexor and extensor muscles and scapular muscles, posture exercises, kinaesthetic exercises and mobilisation techniques, education including ergonomics, daily living advice, home exercise (2) Self-management programme, education, advice and exercise (SMP) (n = 35) All: intervention period 10 wks | NPI, VAS**                                                                 | The MPT group attained a statistically significant greater reduction in reported neck pain and disability (NPI) (P = 0.04), effect size 0.48, measured immediately following treatment |
| Pato et al. [52]        | Patients with whiplash injury grade I or II (Quebec Task Force Classification), with persistent neck pain or headache 6–12 mths after the accident (N = 91) | (1) Local anesthetic infiltration of tender points in the neck 2 × a wk, in 8 wks, (n = 30) (2) Physiotherapy, 2 × a wk, in 8 wks: massage, relaxation techniques of myogelotic muscles, instructed in a detailed homeprogram of isometric and low-intensity active isotonic training of neck muscles (n = 29) (3) Medication: 200 mg flurbiprofen in its slow release preparation once a day. Patients were seen twice a wk by the same study physician during the 8 wks (n = 28) All: furthermore, in each treatment group patients were randomly allocated to additional cognitive-behavioral therapy (CBT) or no CBT. CBT twice a wk for 8 wks Each session lasted 60 minutes | Subjective outcome rating (free of symptoms, improved, unchanged, worse), McGill pain questionnaire, VAS), working capacity | There were no statistically significant differences between the 3 different treatment groups measured at 8 wk and at 6 mths followup. There was a statistically significant effect in the short term in female patients in the groups with additional CBT (P = 0.024) after 8 wks of treatment in the subjective outcome, but not at 6 mths followup |
Table 2: Continued.

| Author                  | Participants                                                                 | Interventions                                                                 | Main outcome measures | Study results on effect* of intervention on pain                                                                 |
|------------------------|------------------------------------------------------------------------------|-------------------------------------------------------------------------------|-----------------------|---------------------------------------------------------------------------------------------------------------|
| Ryan [53]              | Patients with chronic WAD, duration of pain not reported (N = 103)            | (1) Strength training group (n = not reported)  
(2) Endurance training group (n = not reported)  
All: twice a wk for 8–12 wks | VAS, SF-36, strength           | There were no statistically significant differences between groups posttreatment |
| Söderlund and Lindberg [54] | Patients with chronic WAD, (>3 mths after injury), age 18–60 (N = 33)       | (1) Physiotherapy with cognitive behavioural components, learning and application of basic physical and psychological skills in everyday activities, besides physiotherapy as in group 2 (n = 16)  
(2) Physiotherapy, individualised exercises at home and/or in departments gym, various pain-relieving methods (i.e., TENS, heat) (n = 17)  
All: max. 12 individual sessions with the physiotherapist | PDI, NRS, physical measures of pain, disability, coping and self-efficacy | Results revealed no statistically significant differences between groups in self-ratings of disability or pain intensity post treatment or at 3 mths followup |
| Stewart et al. [55]    | Patients with chronic WAD (>3 mths, <12 mths), classified WAD I–III, having significant pain or disability (N = 134) | (1) Advice alone group (n = 68), received education, reassurance and encouragement to participate in light activity alone, advice given in one consultation and two follow-up phone contacts  
(2) Advice and exercise group (n = 66), individualised, progressive, submaximal programme designed to improve functional activities, endurance, strength, aerobic, speed, coordination, principles of cognitive behavioral therapy (i.e., setting goals), 12 sessions during 6 wks | Pain intensity and pain bothersomeness rated on a 0–10 box scale, PSFS | Exercise and advice were more effective than advice alone at 6 wks for all primary outcomes but not at 12 months. The effect of exercise on the 0–10 pain intensity scale was −1.1 (95% CI −1.8 to −0.3, \( P = 0.005 \)) at 6 wks and −0.2 (0.6 to −1.0, \( P = 0.59 \)) at 12 mths; on the botheromeness scale the effect was −1.0 (−1.9 to −0.2, \( P = 0.003 \)) at 6 wks and 0.3 (−0.6 to 1.3, \( P = 0.48 \)) at 12 mths |

* In order to show an effect of an intervention and hereby support the intervention, it requires showing statistical significant difference between groups.
**Secondary outcome measure.
VAS: visual analogue scale; NRS: numerical rating scale; VNPS: verbal numeric pain scale; NPQ: Northwick Park neck pain questionnaire; NDI: neck disability index; NPDI: neck pain and disability index; NPDS: neck pain and disability scale; NPDVAS: neck pain and disability visual analogue scale; PSFS: patient-specific functional scale; NPI: Northwick Park neck pain index; SF-36: short-form 36; PPT: pressure pain threshold; ROM: range of movement; RPE: rating of perceived exertion; EMG: electromyographic; HRQoL: health-related quality of life.
| Author          | Participants                                                                 | Interventions                                                                 | Main outcome measures                                                                 | Study results on effect of intervention between groups                                                                 |
|-----------------|-------------------------------------------------------------------------------|-------------------------------------------------------------------------------|--------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------|
| Bronfort et al. | Patients with mechanical neck pain lasting > 12 wks, age 20–65 (N = 191)    | (1) Spinal manipulation and low-technology rehabilitative neck exercise (n = 63)  
(2) High-technology MedX rehabilitative neck exercise (n = 60)  
(3) Spinal manipulation (n = 64)  
All: attended 20 one-hour visits during the 11 wk study | Pain rating scale (0–10), NDI, SF-36, global improvement of satisfaction with care, medication use | No statistically significant differences between groups in patient-rated outcomes at 11 wk and at 12 mth followup |
| Lau et al. [57] | Patients with a diagnosis of chronic mechanical neck pain (>3 mths), age 18–55 (N = 120) | (1) Thoracic manipulations TM, anterior-posterior approach in supine lying (n = 60)  
(2) Control (n = 60)  
All: 8 sessions infrared radiation (2/wk) for 15 min over painful site. Educational pamphlet involving active neck mobilisation, isometric neck muscle stabilisation, stretching, postural correction exercise | NPRS, NPQ, SF-36, cervical ROM, craniovertebral angle | Statistically significant differences in favour of TM post-treatment on pain intensity (P = 0.043) and NPQ (P = 0.018). Improvements were maintained at 3 and 6 mths followup |
| Martel et al. [58] | Patients with pain of mechanical origin located in the anatomical region of the neck, with or without radiation to the head, trunk, or limbs > 12 wks; between 18 and 60 yrs (N = 98) | All: spinal manipulation 10–15 treatments in 5-6 wk (symptomatic phase) after that 3 different interventions (preventive phase).  
(1) Spinal manipulation cervical and thoracic until Th4, once per month, 4 times (n = 36)  
(2) Spinal manipulation cervical and thoracic until Th4, once per mth, 4 times AND 20–30 min. home exercises 3× per wk: including range of motion exercises, 4 stretching/mobilisation, and 4 strengthening exercises. Three series of each exercise. Ten mths (n = 33)  
(3) Attention control group: no treatment (n = 29) | VAS, cervical ROM, NPDI, Bournemouth questionnaire, SF-12 questionnaire, fear-avoidance behaviour questionnaire | No statistically significant differences were found between groups |
| Sherman et al. [59] | Patients with chronic neck pain (>3 mths), age 20–64 (N = 64) | (1) Therapeutic neck massage (n = 32), commonly used Swedish and clinical massage techniques, allowed typical self-care recommendations, up to 10 treatments over a 10 wk period  
(2) Self-care book (n = 32), they were mailed a copy of a self-care book with information and recommendation | NDI, NRS | Statistically significant effect on massage after four wks measured by NDI, -2.1 (-4.00–0.03) (P = 0.047), but not in long-term followup at 10 and 26 wks |
| Sillevis et al. [60] | Patients between 18 and 65 from outpatients physiotherapy clinic with non-specific pain in the cervical and cervicothoracic region down to T4, provoked with neck movements, present for at least 3 mths (N = 100) | (1) One time thrust manipulation at T3-T4 (n = 50)  
(2) Placebo manipulation at T3-T4 (n = 50) | VAS, pupil diameter | No statistically significant differences between groups immediately after the treatment |
| Author          | Participants                                                                 | Interventions                                                                 | Main outcome measures              | Study results on effect* of intervention between groups                                                                 |
|-----------------|-------------------------------------------------------------------------------|------------------------------------------------------------------------------|-----------------------------------|----------------------------------------------------------------------------------------------------------------------|
| Yağci et al. [61] | Patients with chronic cervical myofascial pain syndrome (>6 mths), age 21–44 (𝑁 = 40) | (1) Spray-stretch technique (𝑛 = 20), ethyl chloride sprayed on muscle with trigger point in muscle stretched position, 6 sessions  
(2) Connective tissue massage (𝑛 = 20), 15 sessions  
All: followed active exercises to be carried out three times a day | VAS, pain threshold, ROM, strength, endurance                                 | No statistically significant differences between groups were found on pain posttreatment |
### Table 4: Electrotherapy—patients with chronic nonspecific neck pain.

| Author          | Participants                                                                 | Interventions                                                                                                                                                                                                 | Main outcome measures                      | Study results on effect* of intervention on pain                                                                 |
|-----------------|-----------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------|---------------------------------------------------------------------------------------------------------------|
| Altan et al. [62] | Patients with chronic cervical myofascial pain syndrome (≥3 mths), having tender points (N = 53) | (1) Laser treatment (n = 23), applied over four trigger points bilat., frequency 1000 Hz for 2 min over each point. Laser parameters: infrared 27 GaAs diode, 904 nm, frequency range 5–7000 Hz, max power of 27 W, 50 W, or 27 × 4 W was used  
(2) Placebo, sham laser treatment (n = 25)  
All: treatment once a day for 10 days during a period of 14 days, instructed to perform isometric exercises and stretching at home | VAS, algometric measurements, ROM                                                                 | There were no significant differences between groups immediately after (wk 2) and at 12 wks followup |
| Chiu et al. [63]  | Patients with chronic intermittent neck pain (≥3 mths), age 20–70 (N = 218)    | (1) TENS group (n = 73): infrared radiation, advice on neck care, TENS to the neck region for 30 min. TENS parameters: dual-channel TENS unit (130 Z), continuous 150 μs square pulses at 80 Hz, four surface electrodes, intensity of TENS was adjusted to produce a tingling sensation  
(2) Exercise group (n = 67): infrared radiation, advice on neck care, intensive neck exercise programme  
(3) Control group (n = 78): infrared radiation, advice on neck care  
All: two sessions a wk for six wks | VNPS, NPQ, NPI, strength                                                                 | There were no statistically significant differences between the three groups on VNPS pain after 6 wk and at 6 mths followup, but the TENS group and the exercise group had a significantly better improvement in NPQ than that of the control group (P = 0.034 and P = 0.02, resp.) after 6 wks and at 6 mths followup |
| Chow et al. [64]   | Patients with chronic neck pain (≥3 mths), age > 18 (N = 90)                | (1) Laser treatment (n = 45), applied to tender points for 30 s per point with up to 50 points being treated. Laser parameters: class 3B, diolase devices, wavelength 830 nm, power of 300 mW in continuous wave mode at a power density of 0.67 W/cm²  
(2) Sham laser treatment (n = 45)  
All: 14 treatments over 7 wks | VAS                                                                 | The improvement in raw VAS was statistically significantly greater in the laser-treatment group than in the sham laser treatment group (~2.7 compared with +0.3, P < 0.001), at 12 wk followup |
| Dundar et al. [65]  | Patients with chronic cervical myofascial pain, having spot tenderness along taut band, age 20–60 (N = 64) | (1) Laser treatment (n = 32), applied over three trigger points bilat., frequency 1,000 Hz for 2 min over each point, power output 58 mW/cm² by 1,000 Hz. Dose per point 7 J, total per treatment 42 J. Laser parameters: infrared Ga-As-Al diode, wavelength 830 nm, max power output of 450 mW  
(2) Placebo, sham laser (n = 32)  
All: once a day for 15 days during 3 wks, instructed in daily isometric exercise and stretching exercise | VAS, ROM, NDI                                                                                   | There were no statistically significant differences between groups after 4 wks                                                                 |
| Author           | Participants                                                                 | Interventions                                                                 | Main outcome measures | Study results on effect of intervention on pain                                                                 |
|------------------|------------------------------------------------------------------------------|------------------------------------------------------------------------------|-----------------------|---------------------------------------------------------------------------------------------------------------|
| Esenye et al. [66] | Patients with chronic myofascial trigger points (duration 6 months to 7 yrs) in one side of the upper trapezius muscles (N = 102) | (1) Ultrasound therapy (n = 36), dose 15 W/cm², 6 min., 10 sessions  
(2) Trigger point injections (1% lidocaine) (n = 36)  
(3) Control (n = 30)  
All: neck-stretching exercises | VAS, PPT, ROM | Statistically significant and equal reduction in VAS pain from ultrasound and injection groups compared with controls (P < 0.001) after treatment and at 3 mth followup. There were no statistically significant differences in outcome measures between groups 1 and 2 |
| Gam et al. [67]   | Patients with chronic trigger points in the neck and with an intensity disturbing normal daily activity, age 18–60 (N = 67) | (1) Ultrasound, massage, exercise (n = 18), dose 100 Hz, pulse = 2: 8, 3 W/cm², 3 min  
(2) Sham ultrasound, massage, exercise (n = 22)  
(3) Control group (n = 18)  
Groups 1 and 2 were treated 2 sessions per wk in 4 wks | VAS, measure of trigger points | There were no significant differences between groups post treatment and at 6 mth followup |
| Gur et al. [68]   | Patients with chronic myofascial pain syndrome in the neck (>1yr), affecting quality of life, with 1–10 tender points in shoulder girdle (N = 60) | (1) Laser treatment (n = 30), 2 J/cm² at each trigger point (max. 20 J/cm²). Laser parameters: Ga-As laser, 20 W max output per pulse, 904 nm, 200 nanoseconds max duration pulse, 2.8 kHz pulse frequency, 11.2 mW average power, 1 cm² surface  
(2) Placebo, sham laser treatment (n = 30)  
All: treatment 3 min at each triggerpoint, 5 times a wk for 2 wks, instructed in correct posture, ergonomics and to avoid activity exacerbated pain | NPDS, VAS | Statistically significant difference on pain in favour of laser treatment at 2nd wk and 3rd wk on pain VAS (2nd wk: VAS pain at rest 3.11 ± 2.29, P = 0.01; VAS pain at movement 2.67 ± 2.58, P = 0.01) and NPDS, and at 12 wk followup maintained at NPDS (41.14 ± 28.34) (P = 0.01) |
| Özdemir et al. [69] | Patients with chronic neck pain related to osteoarthritis (N = 60) | (1) Low-level therapy (n = 30), applied to 12 points, 0.90 J for each 1 cm², each point for 15 s. Laser parameters: endolaser 476, Ga-As, Al, power output of 50 mW, wavelength 830 nm, diameter beam 1 mm., 0.90 J for each 1 cm²  
(2) Placebo, sham laser (n = 30)  
All: treatment in 10 consecutive days | VAS, physician assessment of pressure pain, angle of lordosis, ROM, NPDS | The authors did not report any data on statistically significant differences on pain between groups after treatment |
| Seidel and Uhlemann [70] | Patients with chronic cervical pain syndrome (>6 mths) (N = 51) | (1) Placebo, sham laser treatment (n = 13)  
(2) Laser treatment (n = 12), output 7 mW, stimulation to meridian points, 1 min per point, totally 15 points. Laser parameters: cw-IR-GaAlAs-Laser, 830 nm, Lasotronic, energy density 0 J/cm²; 21 J/cm²; 90 J/cm², irradiation area 0.02 cm², laser skin difference 8 mm  
(3) Laser treatment (n = 13), output 30 mW, stimulation to meridian points, 1 min per point, totally 15 points. Laser parameters: cw-IR-GaAlAs-Laser, 830 nm, Lasotronic, energy density 0 J/cm²; 21 J/cm²; 90 J/cm², irradiation area 0.02 cm², laser skin difference 8 mm  
(4) Needle acupuncture (n = 13)  
All: 8 treatments in 4 wk | VAS, PPT, ROM | The authors did not report any data on statistically significant differences on pain between groups after 4 wk |
| Author          | Participants                                                                 | Interventions                                                                 | Main outcome measures | Study results on effect\* of intervention on pain                                                                 |
|-----------------|-------------------------------------------------------------------------------|-------------------------------------------------------------------------------|-----------------------|------------------------------------------------------------------------------------------------------------------|
| Smania et al. [71] | Patients with chronic myofascial pain syndrome of the superior trapezius muscle (and in no other muscle), age 18–80 (N = 53) | (1) Repetitive magnetic stimulation (rMS) (n = 17), stimulation to trigger points with figure-eight-shaped coil until coil temperature reached 40 degrees and then replaced by circular coil, pulsed magnetic stimuli (4000) each 20 min session in 5-second trains at 20 Hz separated by 25-second pause  
(2) Transcutaneous electrical stimulation (TENS) (n = 18), 100 Hz, pulse width 250 μs, asymmetrical rectangular biphasic waveform, intensity set to patients comfort until significant local sensation  
(3) Placebo (n = 18), sham-ultrasound therapy  
All: treatment sessions, 2 times a wk for 2 wks | NPDVAS, VAS, PPT, ROM | The rMS group and the TENS group showed a statistically significant improvement in the NPDVAS compared to the placebo group; differences to placebo group in NPDVAS, rMS group: pre-post P < 0.01; pre-1 mth P < 0.001; pre-3 mths P = 0.038. Differences to placebo group in NPDVAS, TENS group: pre-post P < 0.01; no difference in pre-1 mths and pre-3 mths test. Difference in effect on NPDVAS between rMS and TNS in favour of rMS only in pre-1 mths test, P < 0.001, and in pre-3 mths test, P < 0.001 |
| Thorsen et al. [72] | Female laboratory workers with chronic pain (>1 yr) from neck and shoulder girdle, pain affecting the quality of work or daily living, 1–10 tender points, age 18–65 yrs (N = 49) | Crossover study, 6 sessions over 2 wks followed by one wk pause before 6 new treatment sessions over 2 wks in other group.  
(1) Laser treatment (n = 25) 0.9 J per treated point max 9 J per treatment. Laser parameters: endolaser 465 class no. B, 830 nm ± 0.5 nm, 30 mW, GaAlAs diode, beam divergence 4 degrees, probehead 2.5 mm²  
(2) Placebo, sham laser treatment (n = 22)  
All: 6 sessions over a 2 wk period | VAS | There were no statistically significant differences between groups post treatment |

\* In order to show an effect of an intervention and hereby support the intervention, it requires showing statistical significant difference between groups.  
VAS: visual analogue scale; NRS: numerical rating scale; VNPS: verbal numeric pain scale; NPQ: Northwick Park neck pain questionnaire; NDI: neck disability index; NPDI: neck pain and disability index; NPDS: neck pain and disability scale; NPDVAS: neck pain and disability visual analogue scale; PSFS: patient-specific functional scale; NPI: Northwick Park neck pain index; SF-36: short-form 36; PPT: pressure pain threshold; ROM: range of movement; RPE: rating of perceived exertion; EMG: electromyographic; HRQoL: health-related quality of life.
Table 5: (a) Exercise therapy, (b) manual therapy, and (c) electrotherapy.

| Author               | Agenda                                                                 | Sequence generation | Allocation concealment | Blinding of participants, personnel, and outcome assessors | Incomplete outcome data | Selective outcome reporting | Other sources of bias | Result of summary assessment of risk of bias |
|----------------------|------------------------------------------------------------------------|---------------------|------------------------|-----------------------------------------------------------|-------------------------|-----------------------------|------------------------|------------------------------------------|
| Cunha et al. [31]    | Global posture reeducation + stretching                                | Adequate            | Unclear                | Inadequate                                                | Inadequate              | Unclear                     | Unclear                | High                                     |
| Dellve et al. [32]   | Myofeedback training + intensive strength training                     | Unclear             | Unclear                | Inadequate                                                | Inadequate              | Unclear                     | Unclear                | Low                                      |
| Ehrenborg and Archenholtz [49] | Biofeedback training + interdisciplinary rehabilitation         | Adequate            | Adequate                | Inadequate                                                | Adequate                | Unclear                     | Unclear                | Low                                      |
| Falla et al. [33]    | Endurance-strength exercise                                           | Adequate            | Unclear                | Inadequate                                                | Adequate                | Unclear                     | Adequate              | Unear                                    |
| Fitz-Ritson [50]     | Proprrioception, eye-head-neck coordination                           | Inadequate          | Inadequate             | Inadequate                                                | Inadequate              | Unclear                     | Unclear                | High                                     |
| Griffiths et al. [34] | Specific neck stabilisation exercises + general exercises               | Adequate            | Adequate                | Inadequate                                                | Adequate                | Unclear                     | Adequate              | Low                                      |
| Gustavsson et al. [35] | Multicomponent pain and stress self-management group intervention + individual physiotherapy | Adequate            | Adequate                | Inadequate                                                | Adequate                | Unclear                     | Adequate              | Low                                      |
| Häkkinen et al. [36] | Strength training + stretching                                         | Adequate            | Adequate                | Inadequate                                                | Adequate                | Unclear                     | Adequate              | Low                                      |
| Jordan et al. [37]   | Intensive training + physiotherapy + chiropractic manipulation         | Adequate            | Adequate                | Inadequate                                                | Unclear                 | Unclear                     | Adequate              | Low                                      |
| Jull et al. [38]     | Proprioception, eye-head coordination + cranio-cervical flexion        | Adequate            | Adequate                | Inadequate                                                | Inadequate              | Unclear                     | Adequate              | Low                                      |
| Jull et al. [51]     | Multimodal physiotherapy programme                                     | Adequate            | Adequate                | Inadequate                                                | Adequate                | Unclear                     | Adequate              | Low                                      |
| Jull et al. [39]     | Cranio-cervical flexion exercise + strength exercises                  | Adequate            | Unclear                | Inadequate                                                | Adequate                | Unclear                     | Adequate              | Low                                      |
| O’Leary et al. [40]  | Cranio-cervical flexion + cervical flexion endurance training          | Unclear             | Unclear                 | Inadequate                                                | Unclear                 | Unclear                     | Adequate              | Unclear                                  |
| Pato et al. [52]     | Cognitive behavioural therapy + multimodal physiotherapy + infiltration + medication | Unclear             | Unclear                 | Inadequate                                                | Unclear                 | Unclear                     | Unclear              | Unear                                    |
| Randlov et al. [41]  | Intensive training + light training                                     | Adequate            | Unclear                | Inadequate                                                | Unclear                 | Unclear                     | Adequate              | Unclear                                  |
| Revel et al. [42]    | Proprioception, eye-head-neck coordination                             | Unclear             | Unclear                 | Inadequate                                                | Unclear                 | Unclear                     | Adequate              | Unclear                                  |
| Ryan [53]            | Strength training + endurance training                                 | Unclear             | Unclear                 | Inadequate                                                | Unclear                 | Unclear                     | Adequate              | Low                                      |
| Söderlund and Lindberg [54] | Cognitive behavioural programme                                        | Unclear             | Unclear                 | Inadequate                                                | Adequate                | Unclear                     | Adequate              | Unclear                                  |
| Stewart et al. [55]  | Exercise                                                                | Adequate            | Adequate                | Inadequate                                                | Adequate                | Adequate                     | Adequate            | Low                                      |
| Taimela et al. [43]  | Multimodal proprioceptive training + home exercises                    | Unclear             | Unclear                 | Inadequate                                                | Adequate                | Adequate                     | Adequate            | Unear                                    |
| Viljanen et al. [44] | Dynamic muscle training + relaxation training                          | Adequate            | Unclear                | Inadequate                                                | Adequate                | Adequate                     | Adequate            | Unclear                                  |
| Vonk et al. [45]     | Behavioural graded activity + exercise                                 | Adequate            | Adequate                | Inadequate                                                | Adequate                | Adequate                     | Adequate            | Low                                      |
| Waling et al. [46]   | Strength + endurance + coordination training                           | Unclear             | Unclear                 | Inadequate                                                | Unclear                 | Adequate                     | Adequate            | High                                     |
| Ylinen et al. [47]   | Intensive strength training + lighter endurance training               | Adequate            | Adequate                | Inadequate                                                | Adequate                | Unclear                     | Adequate            | Low                                      |
| Ylinen et al. [48]   | Stretching exercises + manual therapy                                  | Adequate            | Adequate                | Inadequate                                                | Adequate                | Unclear                     | Adequate            | Low                                      |
| Author                  | Agenda                                      | Sequence generation | Allocation concealment | Blinding of participants, personnel, and outcome assessors | Incomplete outcome data | Selective outcome reporting | Other sources of bias | Result of summary assessment of risk of bias |
|------------------------|---------------------------------------------|---------------------|------------------------|----------------------------------------------------------|------------------------|-----------------------------|-------------------------|-------------------------------------------|
| Bronfort et al. [56]   | Manipulation + exercise                     | Adequate            | Adequate               | Inadequate                                              | Adequate              | Unclear                     | Adequate               | Low                                       |
| Lau et al. [57]        | Thoracic manipulation                       | Adequate            | Adequate               | Inadequate                                              | Adequate              | Unclear                     | Unclear                | Low                                       |
| Martel et al. [58]     | Spinal manipulation + home exercise         | Adequate            | Adequate               | Inadequate                                              | Adequate              | Adequate                    | Adequate               | Low                                       |
| Sherman et al. [59]    | Massage                                     | Adequate            | Adequate               | Inadequate                                              | Adequate              | Unclear                     | Adequate               | Low                                       |
| Sillevis et al. [60]   | Thoracic manipulation                       | Adequate            | Adequate               | Inadequate                                              | Adequate              | Unclear                     | Unclear                | Low                                       |
| Yağcı et al. [61]      | Connective tissue massage + spray-stretch technique | Unclear            | Unclear               | Inadequate                                              | Unclear              | Unclear                     | Unclear                | Unclear                                   |
| Altan et al. [62]      | Laser                                       | Unclear             | Unclear               | Adequate                                                | Inadequate            | Unclear                     | Unclear                | Unclear                                   |
| Chiu et al. [63]       | TENS                                        | Adequate            | Adequate               | Inadequate                                              | Adequate              | Unclear                     | Adequate               | Low                                       |
| Chow et al. [64]       | Laser                                       | Adequate            | Adequate               | Inadequate                                              | Adequate              | Unclear                     | Unclear                | Low                                       |
| Dundar et al. [65]     | Laser                                       | Adequate            | Unclear               | Inadequate                                              | Adequate              | Unclear                     | Unclear                | Low                                       |
| Esenye et al. [66]     | Ultrasound                                  | Adequate            | Unclear               | Inadequate                                              | Adequate              | Unclear                     | Unclear                | Unclear                                   |
| Gam et al. [67]        | Ultrasound                                  | Adequate            | Unclear               | Inadequate                                              | Adequate              | Unclear                     | Adequate               | Unclear                                   |
| Gur et al. [68]        | Laser                                       | Adequate            | Unclear               | Inadequate                                              | Unclear              | Unclear                     | Unclear                | Unclear                                   |
| Ozdemire et al. [69]   | Laser                                       | Adequate            | Unclear               | Inadequate                                              | Unclear              | Unclear                     | Unclear                | Unclear                                   |
| Seidel and Uhlemann [70]| Laser                                       | Adequate            | Unclear               | Inadequate                                              | Unclear              | Unclear                     | Unclear                | Unclear                                   |
| Smania et al. [71]     | rMS + TENS                                  | Adequate            | Unclear               | Inadequate                                              | Unclear              | Adequate                    | Unclear                | Unclear                                   |
| Thorsen et al. [72]    | Laser                                       | Unclear             | Unclear               | Adequate                                                | Inadequate            | Unclear                     | Adequate               | Unclear                                   |
6 weeks on pain intensity scale ($P = 0.005$) and on a bothersomeness scale at 6 weeks ($P = 0.003$) and at 12 months ($P = 0.003$).

No trials at low risk of bias support the use of EMG biofeedback.

3.3.2. Manual Therapy

Effect of Manual Therapy on Pain in Patients with Chronic Nonspecific Neck Pain. As shown in Appendix A, Table 3, six trials examined the effect of various types of manual therapy in patients with chronic non-specific neck pain [56–61]. One of the trials was at unclear risk of bias, and for that reason not considered evidence [61]. Three trials at low risk of bias examining the effect of spinal manipulations found no difference between groups [56, 58, 60]. Two trials succeeded in finding an effect on pain from the intervention. Both trials had a low risk of bias, and for this reason, their results were considered evidence of use of manual therapy.

(1) Lau et al. [57] examined thoracic manipulation versus a control group. They found statistically significant differences in favour of thoracic manipulation post-treatment on pain intensity ($P = 0.043$) and pain and disability (NPQ) ($P = 0.018$). Improvements were maintained at 3, and 6-month followup.

(2) Sherman et al. [59] examined massage versus a self-care book. They found statistically significant effect on massage following four weeks of treatment on neck pain and disability (NDI) ($P = 0.047$), but not at long-term followup at 10 and 26 weeks.

No trials at low risk of bias support the use of traction.

3.3.3. Electrotherapy

Effect of Electrotherapy on Pain in Patients with Chronic Nonspecific Neck Pain. As shown in Appendix A, Table 4, 11 trials examined the effect of various types of electrotherapy in patients with chronic non-specific neck pain; two of these were at low risk of bias [63, 64], and nine were at unclear or high risk of bias [62, 65–72]. The two trials at low risk of bias both succeeded in demonstrating an effect on pain from this type of intervention and for this reason; their results were considered evidence of use of electrotherapy.

(1) Chiu et al. [63] examined three interventions: TENS versus exercise versus a control group. There were no statistically significant differences between the three groups on pain (VNPS) after 6-week and at 6-month followup, but the TENS group and the exercise group had a significantly better improvement in neck pain and disability (NPQ) than the control group ($P = 0.034$ and $P = 0.02$, resp.) after 6-week, and at 6-month followup.

(2) Chow et al. [64] examined laser versus sham laser treatment. The improvement in VAS was statistically significantly greater in the laser treatment group than in the sham laser treatment group ($-2.7$ compared with $+0.3$, $P < 0.001$) at 12-week followup.

No trials at low risk of bias support the use of ultrasound therapy. No trials at low risk of bias support the use of rMS.

4. Discussion

4.1. General Interpretation. In this review, we assessed the effect of various interventions for the treatment of chronic neck pain and evaluated the methodological quality of the trials. Our findings emphasise the importance of taking the risk of bias into consideration when evaluating the evidence of an intervention.

Trials varied substantially regarding their internal validity, although the methodological quality of the RCTs in general appeared to be somewhat low with an unclear or high risk of bias. We identified various methodological flaws that may have implications for the internal validity of the trials and consequently may result in biased outcomes. Key domains in this context were randomisation, blinding, and incomplete outcome data.

Our evaluation also exposes a widespread use of within-group analyses, claiming statistically nonsignificant results to be beneficial. Results were frequently analysed and reported as if they were uncontrolled within-group studies, which consequently led to misinterpretation of results. To some extent this may be due to the absence of a control group in many trials, and the use of an active treatment as a comparative group makes the “proof” of a truly statistically significant effect more difficult to find. We believe that attention should be paid to inadequate interpretation of a trial result when authors inadequately interpret lack of difference in terms of efficacy [73–76].

4.2. Effect of Physiotherapy on Chronic Neck Pain. Overall, the evidence of effect of physiotherapy for chronic neck pain is strengthened. Yet, for some of the treatments offered, no definite effect and clinical usefulness can be shown. This does not necessarily implicate that these treatments have no effect, only that the present evidence is not sufficient.

Physiotherapy interventions for chronic neck pain showing the strongest support for an effect on pain are strength and endurance training (supported by two trials by Stewart et al. [55] and by Ylinen et al. [47], treating patients with chronic WAD and patients with chronic non-specific neck pain, resp.). In patients with chronic WAD, multimodal physiotherapy was also shown to have a beneficial effect by one trial by Jull et al. [51]. In patients with chronic non-specific neck pain, the use of cognitive/behavioural components in exercise was supported by one trial by Gustavsson et al. [35]. In regard to manual therapy, massage seems to have an effect on pain in patients with chronic non-specific neck pain, supported by one trial by Sherman et al. [59], and thoracic manipulation seems to have an effect on pain, supported by one trial by Lau et al. [57]. Within the area of electrotherapy, both laser therapy and TNS seem to have an effect on pain in patients with chronic non-specific neck pain. This was supported by one trial by Chow et al. [64] and one trial...
We too found evidence that thoracic manipulations may have an effect on pain [57]. Regarding low-level laser therapy, our findings are consistent with the findings of a review by Chow et al. [81], who found that low-level laser therapy reduces pain in patients with chronic neck pain. A Cochrane Review from 2007 [17] on the effect of conservative treatment for whiplash concluded that clearly effective treatments are not found for treatment of acute, subacute, or chronic symptoms. The findings of our newer review do support multimodal physiotherapy and mixed exercise programmes for chronic WAD. The explanation for the difference may be that the Cochrane Review by Verhagen in 2007 was not updated after January 2007, and our findings are based on more recent research, namely, two trials published later in 2007 [51, 55]. A more recent review by Teasell et al. [82] found evidence to suggest that exercise programmes are the most effective noninvasive treatments for patients with chronic WAD. Our findings give support to the use of cognitive-behavioural element, and to pain and stress self-management. This is in discrepancy to another Cochrane Review by Gross et al. [14] on patient education for neck pain, concluding that there is no strong evidence for the effectiveness of educational interventions in various neck disorders. This difference may be due to the use of only single-modal trials in their review rather than multimodal trials as used in the current review.

4.3. Strengths and Weaknesses of Review Procedures. To our knowledge, this is the first systematic review on interventions for chronic neck pain addressing the majority of commonly used physiotherapeutic modalities in one study, in order to get an overview of the subject area.

The search strategy and selection criteria we used were quite strict and easy to apply and according to normal procedures for conducting systematic reviews [23]. Yet the following limitations of the literature search may have introduced a bias: some relevant trials may have been missed if they used other keywords, although this is not very likely. We had limitations in language, and this may have led to missing studies from countries in Eastern Europe with a tradition of physiotherapy research, like Poland. We decided to limit our search from 1990 to January 2012. This was due to physiotherapists prior to this time not being trained in scientific methods and also that RCTs were rare. Studies earlier than 1990 would in general not be following a strict protocol like the ones used for RCTs, but at best be longitudinal cohort studies.

The quality assessment was presented in a reproducible manner. However, the results may be affected by our emphases during filtering methods for synthesis evidence. We might have chosen to exclude all trials with insufficient reporting on allocation sequence and allocation concealment. However, we chose not to, since this would have left us with very few trials to assess. We assessed risk of bias, requiring a convincing mechanism to be described in order for a trial to be classified as “adequate.” Our approach to this problem was to assume that the quality was inadequate unless information on the contrary was provided, and in doing so, we might have misclassified well-conducted but badly reported trials.
The present review succeeded in a subgroup assessment of physiotherapy treatment for chronic non-specific neck pain and for chronic WAD. Yet the first group was very wide due to the mixed conditions in the group of participants. The various interventions were considered to be complex, multifaceted, and with various co-interventions, and by classifying them into intervention groups according to—what we believed to be—the trial's agenda, we may have misclassified some. On the other hand, the often used combined therapies also highlight a fundamental problem when assessing effect of specific and single physiotherapy modalities. Another issue is the quality of the intervention since the interventions were administered in different ways and in different settings. It is reasonable to expect that the way in which they were administered including the dose-response relationship could have influenced the outcome. It would have been interesting and very relevant to examine this. Herbert and Bo [83] emphasise that researchers carrying out systematic reviews should routinely examine the quality of interventions.

4.4. Future Directions. We need to know which patients will benefit from which intervention, built on well-conducted and well-reported trials, considering subgroups of patients with chronic neck pain, in order to support recommended evidence-based decisions and to set priorities for future research. We also request future trial investigators to consider to what extent co-interventions are valuable, in addition to possible confounders. Another issue to consider is the extent to which the control groups ought to be given care and attention to the same extent as the intervention groups.

Appendices

A. Study Characteristics and Study Results

See Tables 1–4.

B. Risk of Bias

See Table 5.

Conflict of Interests

The authors report no conflict of interests.

Authors’ Contribution

P. Damgaard contributed to the conception, design, and writing of the study protocol and the design of search strategies; she located and obtained trial reports, helped to select and assess trials, conducted the data analysis, and drafted and approved the final paper. E. M. Bartels contributed to the conception of the study protocol and the design of search strategies; she helped to locate and obtain trial reports, and revised and approved the final paper. I. Ris helped to select and assess trials, contributed to the data analysis, and revised and approved the final paper. R. Christensen contributed to the conception, design, and writing of the study protocol, conducted data analysis, and revised and approved the final paper. All authors acted as guarantors for the paper. B. Juul-Kristensen contributed to the conception, design, and writing of the study protocol and the design of search strategies; she helped to select and assess trials and revised and approved the final paper.

Disclosure

The funding organisations had no role in any aspect of the study, the paper, or the decision to publish.

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