Rehabilitation in Subjects with Chronic non-specific Low Back Pain with Sacroiliac Joint Origin: Protocol for a Systematic Review

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INTRODUCTION

Today, Low Back Pain (LBP) has become one of the most common musculoskeletal disorder in the societies with average global prevalence of 38.5% (1, 2). It is hypothesized that the 15-30% of Chronic Non-specific Low Back Pain (CNLBP) have sacroiliac joint origin (3-5). They are two main approaches dealing with sacroiliac joint impairment (6, 7). Non-invasive treatments such as physiotherapy (8-11) are the forefront of treatment in these cases (12-14). If not helpful, invasive approaches including fixation or fusion may be prescribed (15).

Since physiotherapy interventions as a non-invasive technique seems to be effective in treating chronic LBP (16), the aim of this study is to design a systematic review study seeking and comparing the effects of various interventions in this category on the pain and function of subjects that suffer from CNLBP with sacroiliac joint origin; the results will be of clinical value to determine the effectiveness of each method for this particular subgroup of LBP. If enough paper retrieved, the comparison can be made and a comprehensive therapeutic physical therapy plan may be suggested.

SUMMARY

Introduction. This systematic review protocol aims to evaluate the effect of rehabilitation for chronic non-specific low back pain with sacroiliac joint origin.

Methods. Search will be done in Pubmed, ISI Web of Science, Scopus, Clinical Key, Science Direct, Medline, Embase, PEDro, ProQuest, the Cochrane Library, PROSPERO, the MOH Thesis, MOH Articles, Magiran, and SID. Google Scholar search engine will also be used. All types of Clinical Trials, Cohort, Case-controls, Cross-sectional, Observational Descriptive, Case Report, Case Series, Ecological Studies, Systematic Reviews, thesis and dissertation in English and Persian published prior to September 2019 will be included. The articles recruiting 18 to 60 years old will be included. Considering PICO, the finally retrieved articles will be assessed qualitatively by CONSORT, STROBE, PEDro, NIH and CASP checklists. Changes in pain and function will be favorable.

Dissemination. The protocol presented in present paper will be used to summarize and qualify present literatures on conservative therapy for chronic non-specific low back pain with sacroiliac joint origin.

KEY WORDS
Rehabilitation; physical therapy; pain; function; non-specific chronic low back pain; sacroiliac joint dysfunction.
OBJECTIVES
To determine the effect of rehabilitation interventions on the pain and function in individuals suffering from CNLBP with sacroiliac joint origin.

MATERIALS AND METHODS

Trial eligibility criteria
Strict inclusion/exclusion criteria have been introduced in order to precisely protect search strategies and PICOs. The criteria are summarized below.

Study types
All study types, except Qualitative Studies, and Narrative Reviews, i.e. Clinical Trials, Cohort, Case-controls, Cross-sectional, Observational Descriptive, Case Report, Case Series, Ecological Studies, Systematic Reviews and thesis and dissertation will be included.

Participants
People between 18-60 with no regard to gender and ethnicity who suffer from non-specific CLBP of sacroiliac joint origin and received conservative/rehabilitative interventions. The study will be approved for more detailed analysis if the participants suffered from back pain not less than three months with the signs of SIJ involvement. Studies targeting nonhuman samples, professional athletes, subjects with acute LBP, symptoms persisted for less than three months will be excluded. LBP of specified origin like inflammatory diseases, spondylo-arthritis, disk hernia, spinal canal/foraminal stenosis, visceral pains, fractures and trauma, those with referral or radicular symptoms, studies on pregnant women, children (under 18 years) and elderly (over 60 years) will be excluded.

Interventions
At least one of the study groups has to undertake a rehabilitative or conservative intervention including:
- electrotherapy modalities: electrical stimulation currents (Transcutaneous Electrical Nerve Stimulation (TENS), Interferential (IF), Diadynamic, High-voltage, Russian currents, Faradic...), LASERs, ultrasound, shockwave, tecar, magnet, shortwave and microwave diathermy, infra-red radiation, hot packs, cold packs...;
- manual techniques: mobilization, manipulation, Muscle Energy Techniques (MET), soft tissue release, massage techniques, Instrumented Assisted Soft Tissue Manipulation (IASTM), visceral manipulation...;
- exercise therapy: any type of exercise including the proprioceptive neuromuscular facilitation (PNF) approaches; tapping: Kinesio Taping®, McConnell & Mulligan, elastic bandages, Prophylactic athletic taping, non-medicated taping;
- needling: dryneedling, acupuncture, electro-acupuncture;
- orthosis.

Comparators
Studies that compare the effect of aforementioned interventions with a control group (without treatment), sham group (placebo treatment), healthy group (of matched healthy subjects) or those comparing two or more interventions will be included.

Outcome measures
Studies will be included that the experimental (case) group and the control group were established, and the related monitoring data were introduced. Pain and function will be the primary outcome measures if are reported by valid scale or devices. Pain will be assessed by the Numerical Rating Scale (NRS), Visual Analogue Scale (VAS), Pressure Pain Threshold (PPT), McGill Pain Questionnaire, pain provocation tests. Function will be measured by Roland-Morris Disability Questionnaire, Oswestry Disability Index, or clinical/functional tests like active single leg raise, reverse single leg raise. Other tools may be also considered according to the included studies.

Search methods to identify studies
Articles will be accessed from international (Pubmed, ISI Web of Science, Scopus, Clinical Key, Science Direct, Medline, Embase, PEDro, ProQuest, the Cochrane Library, PROSPERO) and national (MOH Thesis, MOH Articles, Magiran, and SID) databases. Google Scholar search engine will also be searched.

Although narrative reviews and qualitative studies will not be targeted in current project, their references will be checked through Cross Reference. The main key words will be rehabilitation, conservaive, physical therapy, pain, func-
tion, non-specific chronic low back pain, sacroiliac that will be updated with study progression. The search strategy for accessing articles will be like the following search query and covers PICO.

Nonspecific AND chronic AND (“low back pain” OR (low AND back AND pain) OR “back ache”) AND (“sacroiliac joint*” OR (sacroiliac AND joint) OR “sacroiliac”) AND (“electric stimulation*” OR (electric AND stimulation) OR “electrical stimulation*”) AND (TENS OR “Transcutaneous Electrical Nerve Stimulation” OR Interferential OR Diadynamic OR High Voltage OR Russian OR Faradic OR LASERs OR ultrasound OR shockwave OR tecar OR magnet OR shortwave OR microwave OR diathermy OR infra-red OR hot pack OR cold packs OR (“manual techniques*” OR (manual AND technique)) OR mobilization OR manipulation OR (“muscle energy*” OR (muscle AND energy)) OR (“soft tissue release*” OR (“soft tissue release*” AND release)) OR (“visceral manipulation*” OR (visceral AND manipulation)) OR (“instrumented assisted soft tissue manipulation*” OR (“instrumented assisted*” AND “soft tissue manipulation*”) OR exercise OR (“proprioceptive neuromuscular facilitation*” OR (proprioceptive AND neuromuscular AND facilitation)) OR taping OR needling OR acupuncture OR orthosis) AND (“control group*” OR ((placebo or unrealistic) AND (treatment OR therapy*)) AND Function*) in TITLE/SUMMARY/KEY WORDS.

P: nonspecific AND chronic AND “low back pain” OR (low AND back AND pain) OR “back ache”).

I: (“electric stimulation*” OR (electric AND stimulation) OR “electrical stimulation*”) AND (TENS OR “Transcutaneous Electrical Nerve Stimulation” OR Interferential OR Diadynamic OR High Voltage OR Russian OR Faradic OR LASERs OR ultrasound OR shockwave OR tecar OR magnet OR shortwave OR microwave OR diathermy OR infra-red OR hot pack OR cold packs OR (“manual techniques*” OR (manual AND technique)) OR mobilization OR manipulation OR (“muscle energy*” OR (muscle AND energy)) OR (“soft tissue release*” OR (“soft tissue release*” AND release)) OR (“visceral manipulation*” OR (visceral AND manipulation)) OR (“instrumented assisted soft tissue manipulation*” OR (“instrumented assisted*” AND “soft tissue manipulation*”) OR exercise OR (“proprioceptive neuromuscular facilitation*” OR (proprioceptive AND neuromuscular AND facilitation)) OR taping OR needling OR acupuncture OR orthosis).

C: “control group*” OR ((placebo or unrealistic) AND (treatment OR therapy*)).

O: Therapy OR Treatment OR function*.

The research process will be conducted independently by two researchers (SIL and TSM) and their results will be compared each week. For all included articles, the search in the reference list (Hand Search) will also be performed. If the full text of the article is not found, the researchers will email the authors or the editor of the journal three times. If the reply was not satisfying, that article will be excluded. To find grey sources, special search in their related databases including registries of clinical trials (i.e. http://www.irct.ir/), http://www.trialcentral.com/, http://www.proquest.com/, http://www.gateway.com/worldwide/ will be done.

### Study selection

Any article published before the end of September 2019 (Shahrivar 9th, 1398 Persian Calendar) will be potentially suitable. The search will be extended three years before the publication of the first article in the field of each intervention type. Search results and Reference lists will be transferred into the citation manager software. Duplicates and those marked as irrelevant will be ignored by screening titles and summaries. The full text of the remaining articles will be reviewed in detail. Under supervision of peers, i.e. ZSR, FB and AR, the whole procedure will run by two researchers (SIL and TSM) who are blind to each other’s work. Any ambiguity or controversy will be discussed in consensus.

**Figure 1** summarizes the study selection flow according to the PRISMA flow diagram.

### Data extraction

The databases mentioned previously will be searched. To determine the inter-rater agreement, screening of PubMed title/summaries will be done independently by two researchers (SIL and TSM) as supervised by ZSR, AR and FB. The reference list in included articles types (qualitative studies and narrative reviews) will be checked by Cross Referencing. The search line will be revised and key words will be updated as the project progresses. Key words from included studies will be merged in the search line. The references list of all accepted studies will be checked using Hand Search strategy. If the researchers did not access the full text of any article, the corresponding/first author or the editorial board of the publishing journal will be emailed thrice. If not effective, the article will be excluded.

The screening of the title/summaries will be performed independently by SIL and TSM and duplicated articles or unrelated ones will be excluded. They also will report the number of the articles retrieved from each database in a flowchart. The whole search results will be transferred into a citation manager. The screening will be start over every three months for validating fast exclusions.

The research team (SIL, ZSR, FB, AR and TSM) will criticize the included studies according to their full-text inde-
pendently. Final decision on whether a study should be accepted for quality and quantity analysis will be made according to inclusion/exclusion criteria in a consensus. Reasons for the exclusion will be reported. Data from the approved full-texts will be categorized in an Excel sheet (data extraction) according to publication indices (the author(s), title, publication year, journal, country), participants, study design, sample size, randomization, allocation concealment, blinding, intervention, control intervention, main outcomes, adverse effects, follow-up, withdrawals and results. If needed, more information will be requested from original authors. PI (ZSR) supervises the procedure. Any ambiguity or controversy in any phase will be illuminated in the consensus.

Quality assessment
Supervised by PI (ZSR), each approved article will be qualified in expert consensus with regard to its design by Consort, STROBE, PEDro, CASP and NHLBI checklists. Every checklist will be scored. Articles that get at least 50% of total score of one checklist will be approved quantitative analysis. The articles’ quality will be considered as high (75%), medium (50-75%), low (25-50%), poor (< 25%) based on the scores they gained by each single checklist. Any ranking dissimilarities will be discussed in expert consensus. PEDro is an 11-item scale for clinical trials with “plus” (well addressed items) or “minus” (not-localized item) marks (18). In addition, as the identical, internationally accepted standard for qualification of clinical trials (19), the CONSORT checklist will also be used in present work. For more detailed assessment, the appropriate CONSORT extension may also be used (20).

The STROBE designed separate evaluation checklists to qualify case-control studies, cohort and other designs of studies (21, 22). On the other hand, some researchers believe that the STROBE checklists are not for formal quality assessment like the procedure required in a systematic review and recommend Study Quality Assessment Tools proposed by National Heart, Lung, and Blood Institute (NHLBI) instead (23). Both STROBE and NHLBI checklists were be administered in present study. CASP also introduced specific checklist for each study designs (24) and will be used for determining the evidence ranking. For inclusive qualification of all article types, TIDieR checklist will be administered beside the main checklist.

Figure 1. PRISMA 2009 Flow diagram (17) of the articles selection process.
Measure of Treatment Effects
Mean Difference (MD) with a 95% CI will be the format of choice for reporting continuous outcomes (like pain scales). If other forms of reports are present in articles, they will be covert into MD. For dichotomous or binary data (i.e. adverse events), a risk ratio (RR) with a 95% CI will be calculated.

Missing data
If there was a potential of missing data, the original research teams will be contacted. If they do not reply properly, only available data will be analyzed.

Statistical Methods

Data synthesis
If the number of homogenous studies was sufficient, data synthesis will be conducted using Stata software 8.0 (Stata Corporation, College Station, TX). If possible, meta-analysis will be considered using RevMan (Review Manager Software, Version 5.3; Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration, 2014, The Cochrane Collaboration, Oxford, England). The RR and MD with the 95% CI will be determined for dichotomous and continuous data respectively. Heterogeneity assessment will be carried out using the Q test and the I² index. If $I^2 \leq 50\%$, fixed-effects model will be used for calculating the RR and MD; the Mantel-Haenszel random-effects model will be applied, and aggregate participant data will be used. Nonetheless, the random-effects model will be of choice. If quantitative synthesis is not applicable, the results will be discussed descriptively. The procedure will be performed by two researchers (SIL and TSM) independently. Again, any disagreements will be resolved through consensus.

Assessment of heterogeneity
For calculating the heterogeneity, the $I^2$ and $\chi^2$ tests will be used. In $I^2$ analysis the cut off will be set as 50%. If $I^2 > 50\%$, subgroup analysis will be run to highlight the potential factors.

Subgroup and sensitivity analysis
In order to determine the heterogeneity among included studies, subgroup analysis that categorizes each intervention mode according to its frequency and/or timing, type of control, countries and different outcomes may be considered. Then, if the heterogeneity persists or if studies with incomplete results were included, the sensitivity analysis will be done with omitting low quality articles. The meta-analysis will be developed again and the results of these two meta-analyses will be matched and discussed in terms of the sample size, strength of evidence and influence on the pooled effect size.

Assessment of reporting biases
Risk of bias will be assessed by two reviewers (SIL and TSM) using the Cochrane Collaboration’s tool. Disagreements will be resolved in consensus. Reaching appropriate number of studies for qualitative analysis (at least 10 per intervention), funnel plots will be developed for analyzing the publication bias. Besides, the effect of possible selective reporting, reporting deviations from the original protocols, effect of protocol compliance and adherence will be tracked.

DISCUSSION
This systematic review will provide a comprehensive search to retrieve all existing evidences concerning the effects of physiotherapy interventions in CNLBP with sacroiliac joint origin. The inclusion and exclusion criteria provide a reasonable base to assure that these effects will be discussable in temporal spectrum from immediate to very long term effects upon the time intervals of follow ups in the included studies. The main reason for conducting this review was to summarize clinical value of physiotherapy in treatment of these subjects with regard to evidence hierarchy and indicate their strengths. For best internal validity, various check-lists will be administered for each study design and articles scoring will be scheduled in a peers’ consensus. The study will provide data for developing rehabilitative guidelines, apprise insurance coverage and standard protocols of physical therapy planning. According to the retrieved articles, cost-effectiveness and best practice of various physiotherapy intervention may be judged. However, it should be kept in mind that the main challenge in rehabilitation of CNLBP with sacroiliac origin is the accuracy of diagnosis that is not easily confirmed based on clinical examination alone. This fact needed to be appropriately dealt with in the original studies included in the review. We will try to collect all existing studies in this field covering all study designs and all physiotherapy interventions subheadings. The review results will also highlight the existing research and clinical gaps to conduct future researches. As the search has been started right now, it seems that there is not any study available concerning some interventions. In addition, meta-analysis will be applicable only if the retrieved articles were not heterogeneous.

ETHICS
The study has been funded and ethically approved by Isfahan University of Medical Sciences (Ethics Code: IR.MUI.REC.1397.335) as a part of a thesis for Master’s Degree in Physical Therapy by Taraneh Shahmahmoodi (Registration
The sponsor has no role in data collection, analysis of the data and drafting the manuscript. The study meets the ethical standards of the journal of Muscle, Ligament, and Tendon Journal (25).

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This study will be developed with the financial support and ethically approval by Isfahan University of Medical Sciences (Ethics Code: IR.MUL.REC.1397.335) as a part of a thesis for Master’s Degree in Physical Therapy by Taranesh Shahmahmoodi (Registration code: 297122). The protocol has been registered in International Prospective Register of Systematic Reviews (CRD42020121383). The sponsor will not play a role in data collection, analysis of the data and drafting the manuscript.

CONFLICT OF INTERESTS
The authors declare that they have no conflict of interests.

REFERENCES