

# Optimizing management of musculoskeletal pain disorders in primary physiotherapy care

<b>Submission date</b> 05/01/2021	<b>Recruitment status</b> No longer recruiting	<input checked="" type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
<b>Registration date</b> 18/01/2021	<b>Overall study status</b> Completed	<input checked="" type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
<b>Last Edited</b> 04/12/2024	<b>Condition category</b> Musculoskeletal Diseases	<input type="checkbox"/> Individual participant data <input type="checkbox"/> Record updated in last year

## Plain English summary of protocol

### Background and study aims

Musculoskeletal disorders (MSDs) are the number one cause of years lived with disability and reduced health worldwide. In Norway, every fourth patient in primary care suffers from MSDs. Treatment effects are however modest and knowledge of best practice limited. The SupportPrim project will address these challenges in two main steps:

Firstly, to optimize person-centered care, we will employ methods from artificial intelligence in terms of Case-Based Reasoning to build a clinical decision support system (cDSS) based on patient data already collected in primary care physiotherapy. Case-Based Reasoning aims to solve new problems based on the solutions to similar problems in the past. In other words, previous MSD cases will be used to help similar cases in the future, just as humans learn from their own experience. We will then assess the efficacy of the cDSS in physiotherapy practice in a cluster-randomized controlled trial.

Secondly, this effort will be expanded to general practice by implementing The STarT MSK screening Tool as the basis for stratified care for MSD patients. The efficacy of stratified care will be assessed in a cluster-randomized controlled trial in general practice.

### Who can participate?

Adults over 18 years, presenting to a primary care physiotherapist with musculoskeletal pain disorder in any of these areas; shoulder, neck, upper/low back, hip, knee or with multisite pain as primary contact reason. Also 40 physiotherapists will participate.

### What does the study involve?

The physiotherapists will be randomly allocated to receive access to the cDSS to aid their physiotherapy practice or to continue giving treatment as usual. Patients will be treated by the physiotherapists and data will be regularly collected for 12 months to assess if the cDSS is contributing to improved outcomes for patients.

### What are the possible benefits and risks of participating?

Benefits for those in the intervention group is access to a comprehensive overview of the patient's characteristics and reported symptoms at initiation of treatment, as well as treatment advice (on-screen) based on previous, similar patients with successful outcome, which is to be used in a co-decision process between therapist and patient for optimal management of the

current patient.

Participation in the project does not entail any risks or disadvantages in relation to ordinary physiotherapy treatment, as management is at the discretion of the therapist.

Where is the study run from?

Norwegian University of Science and Technology in collaboration with 40 physiotherapists at different private primary care clinics throughout Norway

When is the study starting and how long is it expected to run for?

February 2019 to September 2022

Who is funding the study?

Norges Forskningsråd (Norwegian Research Council)

Who is the main contact?

Dr Ingebrigt Meisingset, Ingebrigt.meisingset@ntnu.no

## Contact information

### Type(s)

Scientific

### Contact name

Dr Ingebrigt Meisingset

### Contact details

Department of Public Health and Nursing  
Faculty of Medicine and Health Science  
Norwegian University of Technology and Science  
Trondheim  
Norway  
N-7491

-  
Ingebrigt.meisingset@ntnu.no

## Additional identifiers

### Clinical Trials Information System (CTIS)

Nil known

### Protocol serial number

Project number NRC: 303331

## Study information

### Scientific Title

An AI-based clinical decision support system for personalized care of common musculoskeletal pain disorders in primary physiotherapy care

### Acronym

SupportPrim

### **Study objectives**

Relative to usual care, we hypothesize that personalized care provided by a clinical decision support system founded in artificial intelligence results in better globally perceived effects and improved function among patients with common musculoskeletal disorders in primary physiotherapy care.

### **Ethics approval required**

Old ethics approval format

### **Ethics approval(s)**

Approved 08/09/2020, Regional Committee for Medical Research Ethics - Mid Norway (NTNU /REK midt, Det medisinske fakultet, Postboks 8905, 7491 Trondheim, Norway; +47 73 59 75 11; rek-midt@mh.ntnu.no), ref. 493080

### **Study design**

Multicentre cluster-randomized controlled trial where individual physiotherapists serve as clusters

### **Primary study design**

Interventional

### **Study type(s)**

Treatment

### **Health condition(s) or problem(s) studied**

Improved treatment outcome for patients with common musculoskeletal disorders such as neck, shoulder, low back, hip, knee and multisite pain

### **Interventions**

To implement personalized care, we will employ innovative methods from artificial intelligence (AI) in terms of Case-Based Reasoning (CBR) to build a clinical decision support system (cDSS) based on patient data already collected in primary care physiotherapy. The CBR methodology will be used to match new patients to previous, similar patients with successful outcome, i.e. it enables a patient-centered intervention based on what has been successful in previous, similar cases. Results from the CBR will be displayed in a clinical dashboard, i.e. the cDSS, for shared decision-making and optimal management of new patients with common musculoskeletal disorders.

Physiotherapists in the intervention group will be provided access to the cDSS, while physiotherapist randomized to the usual care group will carry on patient management as usual without access to the cDSS.

Duration of treatment and number of treatments is at the discretion of each therapist and it is up to each therapist to terminate the treatment. Patients are followed up for 12 months for the purposes of data collection.

### **Randomisation**

We will randomise the therapists (clusters) in 1:1 ratio to the control and intervention groups (automated computerized procedure).

## Intervention Type

Other

### Primary outcome(s)

1. Patients' assessment of their condition measured by Global perceived effect (GPE), a seven-point Likert scale, at 3, 6 and 12 months. The GPE scale will be dichotomized as "improved" (score 1-2) or "unchanged/worse" (score 3-7).
2. Clinically important improvement in function, measured by the Patient Specific Function Scale (PSFS; 0-10), where a 30% change is defined as a Minimal clinically important difference (MCID). PSFS will be measured at baseline, 2, 4, 8 and 12 weeks

### Key secondary outcome(s)

At baseline and 3, 6 and 12 month follow-up secondary outcomes include:

1. Pain intensity (NRS; 0-10)
2. Pain drawing, number of pain sites
3. Pain mapping; how often, continuous (of and on), daily variation
4. EQ-5D-5L (quality of life), five items
5. 15D (health-related quality of life, two items; sleep and vitality)
6. Musculoskeletal Health Questionnaire (MSK-HQ)
7. Number of treatments by 3 months (patient records)
8. Patient-specific functional scale (PSFS; 0-10)
9. Work ability (single item from Work Ability Index; current work ability compared with life-time best)
10. Work status/sick leave (patient records)
11. Medication (patient records)
12. Patient-therapist relationship (3 questions); overall satisfaction, belief in therapist's competency, and communication
13. Benefits of physiotherapy and expectations to physiotherapist (2 questions)
14. Adherence to treatment plan (5 alternatives)
15. Goal achievement (not, partly, or fully achieved) (patient records)
16. Most important reason for success/non-success (text), at 3 months only
17. Still receiving physiotherapy after 3, 6 and 12 months (yes/no)
18. Physical activity, 3 questions; frequency, intensity and duration
19. Hopkins Symptom Check List, HSCL-10 (emotional distress)
20. Pain Self efficacy, 2 questions (from PSEQ 2-item)
21. Cost-utility based on patients deductibles and costs covered by the Norwegian Health Economics Administration (HELFO) and utility measured by EQ-5D

At 2, 4 and 8 weeks secondary outcomes collected by SMS (text-messages):

1. Patient-specific functional scale
2. Work ability
3. Global perceived effect score
4. Pain intensity, at 2 and 8 weeks only
5. Pain self-efficacy, at 2 and 8 weeks only

### Completion date

01/09/2022

## Eligibility

**Key inclusion criteria**

1. Patients presenting to a primary care physiotherapist with musculoskeletal pain disorder in any of these areas; shoulder, neck, upper/low back, hip, knee or with multisite pain as primary contact reason.
2. Age above 18 years
3. Physiotherapists with the patient participants in their current patient population

**Participant type(s)**

Patient

**Healthy volunteers allowed**

No

**Age group**

Adult

**Lower age limit**

18 years

**Sex**

All

**Total final enrolment**

298

**Key exclusion criteria**

1. Reduced cognitive function or skills in Norwegian (impeding reading, speaking or comprehension of Norwegian language)
2. Neurological diagnosis (e.g. multiple sclerosis, stroke, ALS, Parkinson, dementia)
3. On-going cancer
4. Pregnancy or pregnancy-related disorders
5. Scheduled for surgery or recent surgery or fracture within the last 6 months

**Date of first enrolment**

01/02/2021

**Date of final enrolment**

30/11/2021

**Locations****Countries of recruitment**

Norway

**Study participating centre**

**Norwegian University of Science and Technology in collaboration with 40 physiotherapists at different private primary care clinics throughout Norway**

Faculty of Medicine and Health Sciences  
Department of Public Health and Nursing

P.O.Box 8905  
Trondheim  
Norway  
N-7491

## Sponsor information

### Organisation

Norwegian University of Science and Technology

### ROR

<https://ror.org/05xg72x27>

## Funder(s)

### Funder type

Government

### Funder Name

Norges Forskningsråd

### Alternative Name(s)

Forskningsrådet, Norwegian Research Council, Research Council of Norway, The Research Council of Norway

### Funding Body Type

Government organisation

### Funding Body Subtype

National government

### Location

Norway

## Results and Publications

### Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request. Dr Ingebrigt Meisingset, Ingebrigt.meisingset@ntnu.no. Anonymized clustered and participant-level data can be shared after acceptance of publications of main results, providing a relevant research question and ethical approval of the study protocol is documented, including whether data sharing is covered by the

current, approved consent from participants. Collaborative research efforts and work is appreciated.

## IPD sharing plan summary

Available on request

### Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Participant information sheet</a>			04/02/2021	No	Yes
<a href="#">Participant information sheet</a>			04/02/2021	No	Yes
<a href="#">Statistical Analysis Plan</a>	version 1.0	14/06/2022	01/07/2022	No	No
<a href="#">Statistical Analysis Plan</a>	version 2.0	01/03/2023	01/03/2023	No	No
<a href="#">Study website</a>	Study website	11/11/2025	11/11/2025	No	Yes