

The effectiveness of E-health rehabilitation for patients with knee osteoarthritis

Submission date 31/10/2025	Recruitment status No longer recruiting	<input checked="" type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
Registration date 04/11/2025	Overall study status Ongoing	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
Last Edited 05/11/2025	Condition category Musculoskeletal Diseases	<input type="checkbox"/> Individual participant data <input checked="" type="checkbox"/> Record updated in last year

Plain English summary of protocol

Background and study aims

This study aims to test whether a telerehabilitation program using a smartphone app can improve muscle strength, pain, physical performance, and adherence to exercise among people with knee osteoarthritis (KOA). Traditional home exercise programs are effective but often suffer from poor long-term adherence. Using technology like a smartphone app may help patients follow their programs more consistently through video demonstrations, reminders, and therapist communication.

Who can participate?

Adults aged 40–70 years who have been diagnosed with grade II or III knee osteoarthritis for at least 6 months, have mild to moderate pain (rated 4–7 on a 0–10 scale), can walk without assistance, can read Arabic, and can use a smartphone.

What does the study involve?

Participants will be randomly assigned to one of two groups:

App-based group: Use a smartphone application providing resistance exercises with videos, reminders, and therapist feedback.

Paper-based group: Receive the same exercises in a printed booklet.

Both groups will perform the exercises three times per week for 6 weeks. Assessments will take place before and after the 6-week program, including tests for pain, muscle strength, physical function, and exercise adherence.

What are the possible benefits and risks of participating?

Participants may experience less pain, stronger muscles, better physical performance, and improved confidence in managing their condition. Risks are minimal but may include temporary discomfort or muscle soreness after exercise.

Where is the study run from?

The study will be conducted at Al-Aqiq General Hospital in Al-Baha, Saudi Arabia.

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

December 2024 to June 2026

Who is funding the study?
Investigator initiated and funded

Who is the main contact?
Dr Rawan Aldhabi, raldhabi@kau.edu.sa

Contact information

Type(s)

Principal investigator

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Additional identifiers**Clinical Trials Information System (CTIS)**

Nil known

Protocol serial number

Nil known

Study information**Scientific Title**

Effectiveness of telerehabilitation-based resistance exercise using a smartphone application in improving muscle strength, pain, physical performance, and adherence among patients with knee osteoarthritis: a randomized controlled trial

Study objectives

Principal Objective:

To evaluate the effectiveness of telerehabilitation-based resistance exercises delivered through a smartphone app compared to paper-based exercises in improving muscle strength, pain, physical performance, and adherence in patients with knee osteoarthritis.

Hypotheses:

1. The smartphone app-based telerehabilitation program will lead to greater improvements in muscle strength compared to the paper-based program.
2. The app-based program will reduce pain and improve physical performance more effectively.
3. The app-based program will enhance adherence to exercise by increasing self-efficacy and motivation.

Ethics approval required

Ethics approval required

Ethics approval(s)

approved 09/12/2024, Scientific Research Committee at Albaha Health Cluster (6720 South Road, Building 3716, Albaha, 65784, Saudi Arabia; +966 (0)114579300; info@shc.gov.sa), ref: KFH /IRB0901202024/7

Study design

interventional single-blinded randomized controlled trial

Primary study design

Interventional

Study type(s)

Efficacy

Health condition(s) or problem(s) studied

Knee osteoarthritis (KOA)

Interventions

Participants eligible for the study will be randomly assigned in a 1:1 ratio to either the intervention group (app-based home exercise program) or the control group (paper-based home exercise program) using a computer-generated block randomization method with variable block sizes to ensure balanced allocation. Allocation will be concealed using sequentially numbered, opaque sealed envelopes prepared by an independent researcher.

App-based group: Use a smartphone application providing resistance exercises with videos, reminders, and therapist feedback.

Paper-based group: Receive the same exercises in a printed booklet.

Both groups will perform the exercises three times per week for 6 weeks. Assessments will take place before and after the 6-week program, including tests for pain, muscle strength, physical function, and exercise adherence.

Intervention Type

Behavioural

Primary outcome(s)

1. Pain is measured using the Arabic Numeric Pain Rating Scale (ANPRS) at baseline and 6 weeks post-intervention
2. Osteoarthritis symptoms, including pain, stiffness, and physical function, are measured using the Arabic version of the WOMAC at baseline and 6 weeks post-intervention
3. Quadriceps muscle strength is measured using a hand-held dynamometer (HHD, microFET®2) at baseline and 6 weeks post-intervention

Key secondary outcome(s)

1. Exercise adherence is measured using a self-reported exercise logbook, with adherence ratio calculated weekly over 6 weeks
2. Exercise self-efficacy is measured using the Arabic Exercise Self-Efficacy Scale (ESES-A) at baseline and 6 weeks post-intervention
3. Lower-limb functional strength is measured using the 30-Second Chair Stand Test (30 CST) and Five-Times Sit-to-Stand Test (FTSST) at baseline and 6 weeks post-intervention
4. Dynamic balance and mobility are measured using the Timed-Up and Go (TUG) test at baseline and 6 weeks post-intervention

Completion date

01/06/2026

Eligibility

Key inclusion criteria

1. Adults aged 40–70 years
2. Diagnosed with mild-to-moderate KOA
3. Able to use a smartphone
4. Able to perform resistance exercises independently

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

40 years

Upper age limit

70 years

Sex

All

Key exclusion criteria

1. Severe KOA or knee surgery in the past 6 months
2. Neurological or musculoskeletal disorders affecting lower limbs
3. Cognitive impairment or inability to follow instructions

Date of first enrolment

15/11/2025

Date of final enrolment

01/03/2026

Locations**Countries of recruitment**

Saudi Arabia

Study participating centre

Al-Aqiq General Hospital
3400, Al Aqiq Saudi Arabia
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65779

Sponsor information

Organisation

Scientific Research Committee of Al-Baha Health Cluster

Funder(s)

Funder type

Other

Funder Name

Investigator initiated and funded

Results and Publications

Individual participant data (IPD) sharing plan

Individual participant data (IPD) that underlie the results reported in this study will be made available upon reasonable request from the Principal Investigator (Dr Rawan Aldhabi; Raldhabi@kau.edu.sa)

IPD sharing plan summary

Available on request