

# New camera-based instrument to measure neck movement: its validity and sex differences in healthy adults

<b>Submission date</b> 12/08/2024	<b>Recruitment status</b> No longer recruiting	<input type="checkbox"/> Prospectively registered
<b>Registration date</b> 05/09/2024	<b>Overall study status</b> Completed	<input type="checkbox"/> Protocol
<b>Last Edited</b> 28/10/2025	<b>Condition category</b> Musculoskeletal Diseases	<input type="checkbox"/> Statistical analysis plan
		<input checked="" type="checkbox"/> Results
		<input checked="" type="checkbox"/> Individual participant data

## Plain English summary of protocol

### Background and study aims

Neck pain is a common musculoskeletal problem that can limit daily activities and cause substantial healthcare costs. Accurate measurement of neck motion is essential for diagnosis and rehabilitation. This study aimed to evaluate RomIX, a new photogrammetric instrument developed by the research team to measure cervical spine motion objectively and automatically. The study examined how reliable and valid RomIX is compared with the standard manual goniometer and also explored differences in neck mobility between male and female.

### Who can participate?

Healthy adults aged 18 years or older with no history of neck or shoulder disorders, no pain (VAS = 0), and minimal neck disability (NDI  $\leq$  4).

### What does the study involve?

Participants stood in the RomIX system and performed six active neck movements: flexion, extension, right and left side bending, and right and left rotation. Each movement was held for five seconds. The same participants were also measured using a manual digital goniometer by two independent clinicians. The results from both methods were compared to test RomIX's reliability and validity. A larger group of 480 asymptomatic adults was assessed using RomIX to establish normative reference data and investigate sex-based differences in cervical motion.

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

Participants contributed to validating a new instrument that may help clinicians assess neck function more accurately and efficiently. The procedure was non-invasive and safe, with no adverse effects reported. RomIX may provide a faster, marker-free, and more objective method for cervical motion assessment in clinical and rehabilitation settings.

### Where is the study run from?

Ho Chi Minh City Hospital for Rehabilitation – Occupational Diseases, Vietnam

When is the study starting and how long is it expected to run for?  
May 2023 to September 2024

Who is funding the study?  
Korea International Cooperation Agency (KOICA) under the project "Education and Research Capacity Building Project at University of Medicine and Pharmacy at Ho Chi Minh City" (Project No. 2021-00020-3).

Who is the main contact?  
Dr Nguyen Huu Duc Minh, MSc, MD (University of Medicine and Pharmacy at Ho Chi Minh City),  
nhdminh@ump.edu.vn

## Contact information

### Type(s)

Public, Scientific, Principal investigator

### Contact name

Mr Minh Nguyen Huu Duc

### ORCID ID

<https://orcid.org/0009-0000-6185-9554>

### Contact details

145 Nhat Tao Street, Ward 8, District 10  
Ho Chi Minh City  
Viet Nam  
700910  
+84 983276267  
nhdminh@ump.edu.vn

## Additional identifiers

### Clinical Trials Information System (CTIS)

Nil known

### ClinicalTrials.gov (NCT)

Nil known

### Protocol serial number

No. 2021-00020-3

## Study information

### Scientific Title

Novel photogrammetric instrument to measure cervical spine range of motion: test-retest reliability, construct validity, and sex-based assessment through a cross-sectional study on asymptomatic adults

### Study objectives

Current Study objectives as of 27/10/2025:

To examine test-retest reliability, construct validity, and measurement error of RomIX, a photogrammetric instrument developed by the authors, to measure active cervical spine range of motion (CROM) compared to manual goniometry.

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Previous Study objectives:

This study aims to examine the reliability and validity of the RomIX, a photogrammetric scanner instrument, to measure cervical range of motion (ROM) compared with manual goniometry

### **Ethics approval required**

Ethics approval required

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

1. approved 27/07/2023, Ethics committee of University of Medicine and Pharmacy at Ho Chi Minh City (217 Hong Bang Street, Ward 11, District 5, Ho Chi Minh City, 748000, Viet Nam; +84 (0) 2838558411; hanhchinh@ump.edu.vn), ref: No. 704/HDDD-DHYD

2. approved 27/07/2023, Ethics committee of Ho Chi Minh City Hospital for Rehabilitation – Occupational Diseases (313 Au Duong Lan Street, Ward 2, District 8, Ho Chi Minh City, 751000, Viet Nam; +84 (0)2838569147; bvphcn.syt@tphcm.gov.vn), ref: No. 16/HDDD-BVPHCN-DTBNN

### **Study design**

Cross-sectional observational clinimetric study"

### **Primary study design**

Observational

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

Diagnostic

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

Evaluation of active cervical spine range of motion in asymptomatic adults using a novel photogrammetric instrument as foundation for cervical spondylosis screening and diagnosis

### **Interventions**

Current interventions as of 27/10/2025:

A total of 530 asymptomatic adults participated in the measurement sessions assessing active cervical spine range of motion in six directions: flexion, extension, right and left side bending, and right and left rotation. Measurements were performed in standing position using both manual digital goniometry and the RomIX photogrammetric instrument.

For RomIX, each participant stood at the center of the instrument and performed maximal active neck movements in each direction, holding for 5 seconds per movement. A single rater recorded all directions three times under identical conditions to determine test-retest reliability. The full RomIX procedure lasted approximately 5 minutes, with a 5-minute rest between repetitions. For manual goniometry, two independent clinicians performed all assessments following

standardized protocols for flexion/extension, side bending, and rotation. Inter-rater reliability was evaluated by comparing results between the two clinicians, who were blinded to each other's data. Each goniometric session took about 10 minutes, with a 5-minute break between repetitions.

All conditions, including participant posture, device alignment, and measurement environment, were standardized to ensure reproducibility and to minimize measurement error.

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Previous interventions as of 09/09/2024:

530 enrolled participants underwent the measurement sessions, where their neck range of motion in six directions (flexion, extension, right/left side bending, right/left rotation) was recorded using manual goniometry and the photogrammetric instrument. Each participant was examined using both methods, with the total duration of involvement being 15 minutes. The total duration of observation is 1 week, with a follow-up period of 2 weeks.

Previous interventions:

50 enrolled participants underwent the measurement sessions, where their neck range of motion in six directions (flexion, extension, right/left side bending, right/left rotation) was recorded using manual goniometry and the photogrammetric instrument. Each participant was examined by both methods, respectively, with the total duration of involvement being 15 minutes. The total duration of observation is 1 week, with a follow-up period of 2 weeks.

## **Intervention Type**

Device

## **Phase**

Phase II/III

## **Drug/device/biological/vaccine name(s)**

RomIX (photogrammetric instrument)

## **Primary outcome(s)**

Current primary outcome measure as of 27/10/2025:

Active cervical spine range of motion in six directions (flexion, extension, right/left side bending, right/left rotation) by manual goniometry and the RomIX photogrammetric instrument at baseline

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Previous primary outcome measure:

Cervical spine range of motion measured in six directions (flexion, extension, right/left side bending, right/left rotation) by goniometry and the the photogrammetric instrument at baseline

## **Key secondary outcome(s)**

Current secondary outcome measures as of 27/10/2025:

Intraclass correlation coefficient (ICC), standard error of measurement (SEM), minimal detectable change (MDC), and Pearson's correlation coefficient (r) from active cervical spine range of motion at baseline

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Previous secondary outcome measures:

Intraclass correlation (ICC), standard error of measurement (SEM), minimal detectable change (MDC), and Pearson's correlation coefficient (r) calculated from neck range-of-motion at baseline

**Completion date**

30/09/2024

## Eligibility

**Key inclusion criteria**

Current key inclusion criteria as of 27/10/2025:

Eligibility criteria for participants were adults aged 18 years or older, regardless of sex or occupation, who were actively participating in study, work, and normal daily activities. Inclusion criteria required that participants had a Visual Analog Scale (VAS) pain score of 0, a Neck Disability Index (NDI) score of  $\leq 4$ , full cognitive and behavioral awareness, and voluntary agreement to participate.

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Previous key inclusion criteria:

1. Participants are 18 years of age or older, regardless of gender or occupation.
2. Participants are capable of performing normal studying or working activities.
3. Participants have full cognitive and behavioral capacity.
4. Participants voluntarily agree to participate in the study.

**Participant type(s)**

Healthy volunteer

**Healthy volunteers allowed**

No

**Age group**

Adult

**Lower age limit**

18 years

**Upper age limit**

60 years

**Sex**

All

## **Total final enrolment**

530

## **Key exclusion criteria**

Current key exclusion criteria as of 27/10/2025:

Exclusion criteria included current or recent neck, shoulder, and/or headache symptoms within the previous 3 months; history of cervical spine or shoulder disorders; neurological and/or rheumatic conditions; and symptoms of nerve compression or radiculopathy.

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Previous key exclusion criteria:

1. Participants report or complain of neck, shoulder and/or headache pain in the previous month.
2. Participants have history of neck and/or shoulder disorders, including trauma and fractures, history of neurological and/or rheumatic disorders.
3. Participants have history of other relevant medical conditions.

## **Date of first enrolment**

01/06/2023

## **Date of final enrolment**

30/09/2024

## **Locations**

### **Countries of recruitment**

Viet Nam

### **Study participating centre**

**Ho Chi Minh City Hospital for Rehabilitation – Occupational Diseases**

313 Au Duong Lan Street, Chanh Hung Ward

Ho Chi Minh City

Viet Nam

751000

## **Sponsor information**

### **Organisation**

University of Medicine and Pharmacy at Ho Chi Minh City

## **Funder(s)**

**Funder type**

Government

**Funder Name**

Korea International Cooperation Agency

**Alternative Name(s)**

KOICA

**Funding Body Type**

Government organisation

**Funding Body Subtype**

National government

**Location**

Korea, South

## Results and Publications

**Individual participant data (IPD) sharing plan**

The datasets generated during and/or analysed during the current study will be available upon request from the corresponding author (Dr Nguyen Huu Duc Minh, [nhdminh@ump.edu.vn](mailto:nhdminh@ump.edu.vn))

**IPD sharing plan summary**

Available on request

**Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Results article</a>			28/10/2025	Yes	No
<a href="#">Dataset</a>			28/10/2025	No	No
<a href="#">Other files</a>	Basic results (Phase 1 only)	09/10/2024	09/10/2024	No	No
<a href="#">Participant information sheet</a>			05/09/2024	No	Yes
<a href="#">Participant information sheet</a>			09/09/2024	No	Yes
<a href="#">Study website</a>	Study website	11/11/2025	11/11/2025	No	Yes