

# Clinical Trial Protocol

## Iranian Registry of Clinical Trials

27 May 2026

### Comparative Effects of Strain Counterstain and Post-Isometric Relaxation Techniques on Pain, Range of motion and Functional Disability in Patients with Upper Cross Syndrome

#### Protocol summary

##### Study aim

To determine the comparative effects of strain counterstain and post-isometric relaxation techniques on pain, range of motion and functional disability in patients with upper cross syndrome.

##### Design

This study will be Randomized Clinical Trial, parallel-group, triple blinded

##### Settings and conduct

The Trial would be conducted in District Headquarters DHQ Hospital OKARA City

##### Participants/Inclusion and exclusion criteria

inclusion criteria □ Individuals with chronic neck pain from > 6 weeks □ Both male and female gender □ Age from 20-40 years □ Neck Pain on > 3 on numerical pain rating scale □ Individuals diagnosed with craniovertebral angle less than 50cm □ Occiput to wall distance greater than 2cm exclusion criteria □ Subjects who will have signs of recent surgery □ Whiplash injury or open wounds □ Cervical spine pathologies like radiculopathies disc herniation, spondylolisthesis, sensory changes in neck region □ Any neurological defect

##### Intervention groups

Group A will receive strain counterstain technique; the position of ease will be produced through positioning the muscle in relaxed/ shortened position, ease will be defined as where a reduction in pain at least 70% then, pressure is applied to Trp and it will be held for 90-120 seconds with 3-5 repetitions per treatment session ( three session per week for 3 weeks). Group B will receive post-isometric relaxation technique with 3-5 muscle contraction at sub maximal pain-free effort (20% of available strength) with 5-7 seconds for 5 repetitions per treatment session (three sessions per week for 3 weeks).

##### Main outcome variables

Pain; Range of motion; Functional disability

#### General information

##### Reason for update

##### Acronym

randomized clinical trial

##### IRCT registration information

IRCT registration number: **IRCT20190717044238N6**

Registration date: **2023-03-06, 1401/12/15**

Registration timing: **registered\_while\_recruiting**

Last update: **2023-03-06, 1401/12/15**

Update count: **0**

##### Registration date

2023-03-06, 1401/12/15

##### Registrant information

##### Name

Fareeha Amjad

##### Name of organization / entity

The University of Lahore

##### Country

Pakistan

##### Phone

+92 42 99200600

##### Email address

fari\_fairy22@yahoo.com

##### Recruitment status

**Recruitment complete**

##### Funding source

##### Expected recruitment start date

2023-03-01, 1401/12/10

##### Expected recruitment end date

2023-06-30, 1402/04/09

##### Actual recruitment start date

empty

##### Actual recruitment end date

empty

**Trial completion date**

empty

**Scientific title**

Comparative Effects of Strain Counterstain and Post-Isometric Relaxation Techniques on Pain, Range of motion and Functional Disability in Patients with Upper Cross Syndrome

**Public title**

Comparative Effects of Strain Counterstain and Post-Isometric Relaxation Techniques on Pain, Range of motion and Functional Disability in Patients with Upper Cross Syndrome

**Purpose**

Treatment

**Inclusion/Exclusion criteria****Inclusion criteria:**

Individuals with chronic neck pain from > 6 weeks Both male and female gender Age from 20-40 years Neck Pain on > 3 on numerical pain rating scale Individuals diagnosed with craniovertebral angle less than 50cm Occiput to wall distance greater then 2 cm

**Exclusion criteria:**

Subjects who will have signs of recent surgery Whiplash injury or open wounds Cervical spine pathologies like radiculopathies disc herniation, spondylolisthesis, sensory changes in neck region Any neurological defect

**Age**

From **20 years** old to **40 years** old

**Gender**

Both

**Phase**

3

**Groups that have been masked**

- Participant
- Outcome assessor
- Data analyser

**Sample size**

Target sample size: **70**

**Randomization (investigator's opinion)**

Randomized

**Randomization description**

As per the inclusion and exclusion criteria of the study , patients will be divided into two groups randomly by Random Number Generator table

**Blinding (investigator's opinion)**

Triple blinded

**Blinding description**

The patients taking part in the study would be blinded, they would not be able to know the group they have been allocated to, either Strain Counterstain and Post - Isometric Relaxation Techniques, The assessor of the outcomes would be blinded and lastly, out data analyzer would be blinded too, making it a triple blinded clinical trial

**Placebo**

Not used

**Assignment**

Parallel

**Other design features**

This study will be Randomized Clinical Trial, parallel-group, triple blinded ( patients, assessor and data analyzer will make it triple blinded)

**Secondary Ids**

empty

**Ethics committees****1****Ethics committee****Name of ethics committee**

Ripah College of Rehabilitation and Allied Health Sciences Lahore

**Street address**

F83G+V25, Madar-e-Millat Road, Quaid-e-Azam Industrial Estate Quaid e Azam Industrial Estate, Lahore, Punjab

**City**

Lahore

**Postal code**

54000

**Approval date**

2023-01-02, 1401/10/12

**Ethics committee reference number**

REC/RCR& AHS/23/0110

**Health conditions studied****1****Description of health condition studied**

Upper Cross Syndrome

**ICD-10 code****ICD-10 code description****Primary outcomes****1****Description**

Functional Disability

**Timepoint**

Total intervention protocol will be given for three weeks of duration, 3 sessions per week for three weeks, Outcomes will be assessed at Baseline, at the end of 3 week( last treatment session) and at the end of 6th week

**Method of measurement**

NDI is the self-report questionnaire that is designed to determine how the neck pain affects a patient's daily life and the disability of patients with neck pain. It consists of 10 questions that ask about ADLS. More the score, greater was the disability. Sample questionnaire is attached at the end. Four sections relate to subjective symptoms, and the remaining 6 sections relate to activities of daily living. Each section is scored from 0 to 5 points, giving a maximum score of 50. The total score of the NDI ranges from 0 to 50 points

## Secondary outcomes

### 1

#### Description

Pain

#### Timepoint

Total intervention protocol will be given for three weeks of duration, 3 sessions per week for three weeks, Outcomes will be assessed at Baseline, at the end of 3 week (last treatment session) and at the end of 6th week

#### Method of measurement

Patient level of pain will be assessed using Numerical Pain Rating Scale ( NPRS) This scale ranges from 0 to 10. 0 indicates “no pain” and 10 indicates “worst pain”

### 2

#### Description

Range of Motion

#### Timepoint

Total intervention protocol will be given for three weeks of duration, 3 sessions per week for three weeks, Outcomes will be assessed at Baseline, at the end of 3 week (last treatment session) and at the end of 6th week

#### Method of measurement

Changes from the Baseline ROM range of Motion of Cervical spine will be taken with the Help of universal Goniometer

## Intervention groups

### 1

#### Description

Intervention group: group A will receive strain counterstain technique; the position of ease will be produced through positioning the muscle in relaxed/ shortened position, ease will be defined as where a reduction in pain at least 70% then, pressure is applied to Trp and it will be held for 90-120 seconds with 3-5 repetitions per treatment session ( three session per week for 3 weeks)

#### Category

Treatment - Other

### 2

#### Description

Intervention group: group B will receive post-isometric relaxation technique with 3-5 muscle contraction at sub maximal pain-free effort (20% of available strength) with 5-7 seconds for 5 repetitions per treatment session (three sessions per week for 3 weeks).

#### Category

Treatment - Other

## Recruitment centers

### 1

#### Recruitment center

#### Name of recruitment center

DHQ hospital Okara

#### Full name of responsible person

Dr Zaheer Abbas khan

#### Street address

RC6R+QVF, Eid Gah Rd okara, Punjab

#### City

okara

#### Postal code

56300

#### Phone

+92 322 5536578

#### Email

zeeokg@gmail.com

## Sponsors / Funding sources

### 1

#### Sponsor

##### Name of organization / entity

Ripah international university

##### Full name of responsible person

Dr. Fareeha Amjad

##### Street address

Ripah international university Maddar e Millat Road, Quaid e Azam industrial Estate Lahore, Punjab

##### City

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##### Postal code

54000

##### Phone

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##### Email

fareeha.amjad@ripah.edu.pk

#### Grant name

#### Grant code / Reference number

#### Is the source of funding the same sponsor organization/entity?

No

#### Title of funding source

Ripah international university

#### Proportion provided by this source

100

#### Public or private sector

Private

#### Domestic or foreign origin

Domestic

#### Category of foreign source of funding

empty

#### Country of origin

#### Type of organization providing the funding

Academic

## Person responsible for general inquiries

#### Contact

##### Name of organization / entity

Ripah international university

##### Full name of responsible person

Anila Ramzan

##### Position

student  
**Latest degree**  
Master  
**Other areas of specialty/work**  
Physiotherapy  
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## Person responsible for scientific inquiries

### Contact

**Name of organization / entity**  
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Assistant professor  
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## Person responsible for updating data

### Contact

**Name of organization / entity**  
Ripah International University  
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Anila Ramzan  
**Position**

student  
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Physiotherapy  
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56130  
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## Sharing plan

### Deidentified Individual Participant Data Set (IPD)

Yes - There is a plan to make this available

### Study Protocol

Yes - There is a plan to make this available

### Statistical Analysis Plan

Yes - There is a plan to make this available

### Informed Consent Form

Yes - There is a plan to make this available

### Clinical Study Report

Yes - There is a plan to make this available

### Analytic Code

Yes - There is a plan to make this available

### Data Dictionary

Yes - There is a plan to make this available

### Title and more details about the data/document

All collected identified IPD

### When the data will become available and for how long

Data will be available after the completion of study and will remain available till 6 months

### To whom data/document is available

Data will be available for other people almost 6 months after the completion of study

### Under which criteria data/document could be used

The data/document could be used by communicating with the principle investigator Hadiqa Naeem on email address hadiqa621@gmail.com

### From where data/document is obtainable

Hadiqa Naeem , hadiqa621@gmail.com

### What processes are involved for a request to access data/document

The data/document could be used by communicating with the principle investigator Hadiqa Naeem on email address hadiqa621@gmail.com

### Comments