

Clinical Trial Protocol

Iranian Registry of Clinical Trials

30 Jul 2026

Comparison of the effect of Tamsulosin, Solifenacin and their combination in the treatment of double J stent-related urinary symptoms in patients with ureteral stones

Protocol summary

Study aim

to evaluate the effect of Tamsulosin, Solifenacin and their combination in the treatment of double J stent-related urinary tract symptoms in patients with ureteral stones

Design

A phase 3, single-blinded, sham-controlled randomized clinical trial (RCT) with the parallel-groups design, on 80 patients. Simple individual randomization was done using random numbers table and sealed envelopes.

Settings and conduct

The study was performed in the Ghaem hospital, Mashhad, Iran. It was a single-blind RCT in which the research participants were blinded to the randomization. Each group of patients received the allocated medication following transurethral lithotripsy and double J stent placement. The Ureteral Stent Symptoms Questionnaire (USSQ) questionnaire was used to evaluate the double J stent-related urinary tract symptoms after 2 weeks of stent placement.

Participants/Inclusion and exclusion criteria

Inclusion criteria include the following: age 20 to 60 years, candidate for unilateral stent implantation following TUL (trans ureteral lithotripsy) and informed consent. Exclusion criteria include: history of surgery on the urinary system, urethroscopy or bilateral double j implantation, history of LUTS symptoms following BPH or other urinary tract diseases such as neurogenic bladder and overactive bladder, concomitant use of anti-adrenergic and anti-urinary drugs Cholinergic, pregnancy, drug allergies and finally the occurrence of major complications following surgery

Intervention groups

patients are randomized into 4 groups receiving the following medications respectively: Tamsulosin 0.4 mg once daily, Solifenacin 5 mg once daily, Tamsulosin + Solifenacin with the same dosage, and placebo

Main outcome variables

double J stent-related lower urinary tract symptoms

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20200831048564N1**

Registration date: **2020-09-17, 1399/06/27**

Registration timing: **retrospective**

Last update: **2020-09-17, 1399/06/27**

Update count: **0**

Registration date

2020-09-17, 1399/06/27

Registrant information

Name

Mohammad Rezaei Moghaddam

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 51 3607 2499

Email address

rezaeimm911@mums.ac.ir

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2019-09-23, 1398/07/01

Expected recruitment end date

2020-07-22, 1399/05/01

Actual recruitment start date

2019-09-23, 1398/07/01

Actual recruitment end date

2020-07-22, 1399/05/01

Trial completion date
2020-07-22, 1399/05/01

Scientific title
Comparison of the effect of Tamsulosin, Solifenacin and their combination in the treatment of double J stent-related urinary symptoms in patients with ureteral stones

Public title
The effect of Tamsulosin and Solifenacin in the treatment of double J stent-related urinary symptoms

Purpose
Treatment

Inclusion/Exclusion criteria
Inclusion criteria:
patients with ureteral stones undergoing unilateral transurethral lithotripsy
Exclusion criteria:
History of surgery on the urinary system Performing urethroscopy or double j implantation bilaterally History of LUTS symptoms following BPH Other diseases of the urinary system such as neurogenic bladder and overactive bladder Concomitant use of anti-adrenergic and anticholinergic drugs pregnancy Drug Allergy Occurrence of major complications following surgery

Age
From **20 years** old to **60 years** old

Gender
Both

Phase
3

Groups that have been masked
• Participant

Sample size
Target sample size: **80**
More than 1 sample in each individual
Number of samples in each individual: **20**
4 groups: 20 patients in each group
Actual sample size reached: **20**

Randomization (investigator's opinion)
Randomized

Randomization description
simple individual randomization using sealed envelopes. In order to random allocation concealment by SNOSE method First, a random sequence assignment was created using the random number table method that created numbers from 00 to 99. Then based on the total sample size of the research, which is 80 people, 80 envelopes with aluminum wrappers (in order not to clarify the contents of the envelopes) were prepared and each of the random sequences created was recorded on a card and the cards were placed in envelopes. The order was placed. In order to maintain a random sequence on the outer surface of the envelopes, numbering was performed in the same way that A, B, C, and D drug groups were assigned in this envelopes. Accordingly, numbers between 00 to 24 were assigned to A group, number 25 to 49 B group, number 50 to 74 C group, and number 75 to 99 D group. Finally, the letter envelopes were glued and placed in a box, respectively. At the

beginning of the registration of participants, based on the order of entry of the participants who met the inclusion criteria, one of the envelopes was opened and the assigned group of the participant was revealed.

Blinding (investigator's opinion)
Single blinded

Blinding description
Research participants were blinded to the randomization. To maintain blindness, identical-looking medications and placebo without name were used.

Placebo
Used

Assignment
Parallel

Other design features
A phase 3, single-blinded, sham-controlled randomized clinical trial (RCT) with the parallel-groups design of 80 patients. Simple individual randomization was done using sealed envelopes.

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Mashhad University of Medical Sciences

Street address

Mashhad, Park Mellat Square, University Campus, Mashhad University of Medical Sciences

City

Mashhad

Province

Razavi Khorasan

Postal code

9188911111

Approval date

2018-12-18, 1397/09/27

Ethics committee reference number

IR.MUMS.MEDICAL.REC.1397.580

Health conditions studied

1

Description of health condition studied

lower urinary tract symptoms, ureteral stone

ICD-10 code

N20.9

ICD-10 code description

Urinary calculus, unspecified

Primary outcomes

1

Description

ureteral stent urinary symptoms

Timepoint

2 weeks after intervention

Method of measurement

Based on Ureteral Stent Symptoms Questionnaire (USSQ)

Secondary outcomes

empty

Intervention groups**1****Description**

Intervention group: 1; cap Tamsulosin 0.4 mg once daily for 2 weeks

Category

Treatment - Drugs

2**Description**

Intervention group: 2; Tab Solifenacin 5 mg once daily for 2 weeks

Category

Treatment - Drugs

3**Description**

Intervention group:3, cap including Tamsulosin 0.4 mg + Solifenacin 5 mg, once daily for 2 weeks

Category

Treatment - Drugs

4**Description**

Control group:Tab Placebo (identical-looking with the medication) once daily for 2 weeks

Category

Treatment - Drugs

Recruitment centers**1****Recruitment center****Name of recruitment center**

Department of Urology, Mashhad University of Medical Sciences

Full name of responsible person

Dr Alireza Akavan Rezayat

Street address

Mashhad, Ahmadabad Boulevard, Ghaem Hospital

City

Mashhad

Province

Razavi Khorasan

Postal code

9188911111

Phone

+98 51 3840 0000

Email

Rezaeimm911@mums.ac.ir

Sponsors / Funding sources**1****Sponsor****Name of organization / entity**

Mashhad University of Medical Sciences

Full name of responsible person

Dr Mohsen Tafaghodi

Street address

Mashhad, University Street, Qureshi building

City

Mashhad

Province

Razavi Khorasan

Postal code

91375345

Phone

+98 51 3841 1538

Email

vcresearch@mums.ac.ir

Web page address

<http://www.mums.ac.ir/research>

Grant name**Grant code / Reference number****Is the source of funding the same sponsor organization/entity?**

Yes

Title of funding source

Mashhad University of Medical Sciences

Proportion provided by this source

100

Public or private sector

Public

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**

Mashhad University of Medical Sciences

Full name of responsible person

Dr Alireza Akhavan Rezayat

Position

Assistant Professor

Latest degree

Specialist

Other areas of specialty/work

Urology

Street address

Mashhad, Ahmadabad Blvd., Ghaem Hospital,

Department of Urology

City

Mashhad
Province
Razavi Khorasan
Postal code
9176999311
Phone
+98 51 3840 0000
Email
akhavara@mums.ac.ir

Person responsible for scientific inquiries

Contact

Name of organization / entity
Mashhad University of Medical Sciences
Full name of responsible person
Dr Alireza Akhavan Rezayat
Position
Assistant Professor
Latest degree
Specialist
Other areas of specialty/work
Urology
Street address
Mashhad,Ahmadabad St,Ghaem Hospital,Department of Urology
City
Mashhad
Province
Razavi Khorasan
Postal code
9176999311
Phone
+98 51 3840 0000
Email
akhavara@mums.ac.ir

Person responsible for updating data

Contact

Name of organization / entity
Mashhad University of Medical Sciences
Full name of responsible person
Dr Alireza Akhavan Rezayat
Position
Assistant Professor
Latest degree
Specialist
Other areas of specialty/work

Urology
Street address
Mashhad, Ahmadabad St,Ghaem Hospital,Department of Urology
City
Mashhad
Province
Razavi Khorasan
Postal code
9176999311
Phone
+98 51 3840 0000
Email
akhavara@mums.ac.ir

Sharing plan

Deidentified Individual Participant Data Set (IPD)

Undecided - It is not yet known if there will be a plan to make this available

Study Protocol

Yes - There is a plan to make this available

Statistical Analysis Plan

No - There is not a plan to make this available

Informed Consent Form

No - There is not a plan to make this available

Clinical Study Report

No - There is not a plan to make this available

Analytic Code

Not applicable

Data Dictionary

Not applicable

Title and more details about the data/document

All data is potentially shareable after unidentified individuals

When the data will become available and for how long

Access period starts 6 months after the results are published

To whom data/document is available

Researchers of medical universities

Under which criteria data/document could be used

The results of this study are intended to help the medical community to help treat patients

From where data/document is obtainable

Ghaem Hospital, Department of Urology
akhavara@mums.ac.ir

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

Please request by email

Comments