The voiding position study (RRSCE)

Submission date

20/12/2005

Recruitment statusNo longer recruiting

Overall study status

Registration date

Completed

20/12/2005 **Condition category** Urological and Genital

Last Edited 19/12/2008

Diseases

Retrospectively registered

? Protocol not yet added

? SAP not yet added

Results not yet added and study completed for more than 2 years

? Raw data not yet added

Study completed

Plain English Summary

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

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Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Protocol/serial number

NTR146; 17

Study information

Scientific Title

The voiding position study - a replicated randomised single case experiment

Acronym

RRSCE

Study hypothesis

The position in which men with lower urinary tract symptoms (LUTS) are voiding may have an influence on the quality of the voiding. The Dutch guidelines for the treatment of LUTS in general practice recommend patients to void in a sitting position. It is unclear whether this advice is correct. The (little) available evidence suggests it may even be wrong.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Received from the local medical ethics committee

Study design

Randomised active controlled crossover two-arm trial

Primary study design

Interventional

Secondary study design

Randomised controlled trial

Study setting(s)

Other

Study type(s)

Quality of life

Participant information sheet

Condition

Lower urinary tract symptoms (LUTS)

Interventions

Two positions in which men can void are compared:

- 1. Sitting
- 2. Standing

Intervention Type

Other

Phase

Not Specified

Primary outcome measure

Peak urinary flow rate

Secondary outcome measures

- 1. Mean urinary flow rate
- 2. Mean micturition time
- 3. Experienced differences

Overall study start date

01/08/2004

Overall study end date

22/03/2005

Eligibility

Participant inclusion criteria

- 1. Men
- 2. Aged greater than or equal to 55 years
- 3. Moderate LUTS, i.e. total score on the International Prostate Symptom Score 8 19

Participant type(s)

Patient

Age group

Adult

Sex

Male

Target number of participants

21

Participant exclusion criteria

- 1. History of surgery for LUTS, e.g. prostatectomy
- 2. Current use of medication that may influence voiding, e.g. diuretics, anticholinergic medication, alpha-blockers, 5-alpha-reductase inhibitors

Recruitment start date

01/08/2004

Recruitment end date

22/03/2005

Locations

Countries of recruitment

Netherlands

Study participating centre Maastricht University Maastricht Netherlands

Sponsor information

Organisation

6200 MD

Care and Public Health Research Institute (CAPHRI) (The Netherlands)

Sponsor details

University Maastricht P.O. Box 616 Maastricht Netherlands 6200 MD +31 (0)43 388 2446 e.habets@caphri.unimaas.nl

Sponsor type

Research organisation

Website

http://www.caphri.nl/

ROR

https://ror.org/02jz4aj89

Funder(s)

Funder type

Research organisation

Funder Name

The Netherlands Organisation for Health Research and Development (ZonMw) (The Netherlands)

Results and Publications

Publication and dissemination plan

Not provided at time of registration

Intention to publish date
Individual participant data (IPD) sharing plan

IPD sharing plan summaryNot provided at time of registration