

Zoetermeer Study: double-blind randomised, placebo-controlled clinical study to investigate the effects of daily oral atamestane (100 mg /day) and dehydroepiandrosterone (50 mg/day) alone and in a combined regimen on physical frailty and quality of life in 100 elderly male volunteers over a treatment period of 36 weeks

Submission date 20/12/2005	Recruitment status No longer recruiting	☐ Prospectively registered ☐ Protocol
Registration date 20/12/2005	Overall study status Completed	☐ Statistical analysis plan ☐ Results
Last Edited 17/09/2008	Condition category Other	☐ Individual participant data ☐ Record updated in last year

Plain English summary of protocol
Not provided at time of registration

Contact information

Type(s)
Scientific

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Additional identifiers

Protocol serial number

Study information

Scientific Title

Study objectives

The study hypothesis is that that daily oral atamestane (100 mg/day), dehydroepiandrosterone (50 mg/day) alone and the combined regimen improve physical frailty, muscle strength and functional performance compared to placebo.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Ethics approval received from the local medical ethics committee

Study design

Randomised, double blind, placebo controlled, parallel group trial

Primary study design

Interventional

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Physical frailty

Interventions

1. Atamestane (100 mg/day)
2. Dehydroepiandrosterone (50 mg/day)
3. Combined regimen of atamestane (100 mg/day) and dehydroepiandrosterone (50 mg/day)

Intervention Type

Drug

Phase

Not Specified

Drug/device/biological/vaccine name(s)

Atamestane, dehydroepiandrosterone

Primary outcome(s)

1. Isometric grip strength
2. Leg extension power
3. Physical performance (according to Guralnik)

Key secondary outcome(s))

1. Activities of Daily Living
2. Quality of life
3. Mini Mental State Examination
4. Body composition
5. Bone density of hip
6. Bone metabolism
7. Hormonal parameters total testosterone, dehydroepiandrosterone (DHEA), dehydroepiandrosterone sulphate (DHEAS), oestradiol, oestrone, sex hormone binding globulin (SHBG), insulin-like growth factor 1 (IGF-1), IGF-binding proteins (IGFBP), IGF-binding protein 3 (IGFBP3)
8. Glucose, insulin HbA1c
9. Immunological parameters (lymphocyte sub-populations and surface markers)
10. Lipid metabolism (high density lipoprotein [HDL], low density lipoprotein [LDL], triglycerides, cholesterol)
11. Carotid intima-media thickness

Completion date

31/08/1997

Eligibility

Key inclusion criteria

1. Men
2. 70 years or older
3. Participant in previous cross-sectional study among 400 men
4. Low performance score on isometric grip strength (IGS) and leg extensor power (LEP) test compared to mean of 400 men in cross-sectional study

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Senior

Sex

Male

Key exclusion criteria

1. Severe arthropathic deformation of knee joint severely limiting mobility
2. Severe systemic disease interfering with conduct of study or interpretation of results
3. Abnormal lab functions from preceding cross-sectional study considered clinically significant and giving suspicion of specific organ dysfunction
4. Myocardial infarction within 6 months prior to first visit or clinical evidence of congestive heart failure
5. History of stroke or transient ischaemic attacks (TIAs)
6. Sitting systolic blood pressure of 200 mmHg or higher or diastolic blood pressure of 105 mmHg or higher at any of pretreatment visits
7. Active malignant disease with significant impact of physical condition

8. History of prostatic cancer
9. Diabetes mellitus treated with insulin
10. Preexisting signs of abnormal liver function with clinical significance
11. History of alcohol abuse within last 2 years
12. Participation in another clinical trial or systemic administration of an investigational drug within the last 3 months prior to start of study

Date of first enrolment

01/01/1996

Date of final enrolment

31/08/1997

Locations

Countries of recruitment

Netherlands

Study participating centre

Erasmus Medical Center

Amsterdam

Netherlands

3015 GD

Sponsor information

Organisation

Erasmus Medical Centre (Netherlands)

ROR

<https://ror.org/018906e22>

Funder(s)

Funder type

Industry

Funder Name

Schering AG (Germany) - Strategic Business Unit Fertility Control and Hormone Therapy (SBU FC /HT)

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration