FES: FESS Effectiveness Study: a multi-centre randomised controlled trial studying the effectiveness of functional endoscopic sinus surgery (FESS) in adult patients with chronic rhinosinusitis/nasal polyps unresponsive to medical therapy

Submission date 14/02/2006	Recruitment status No longer recruiting	 Prospectively registered Protocol
Registration date 14/02/2006	Overall study status Completed	 Statistical analysis plan Results
Last Edited 14/08/2009	Condition category Ear, Nose and Throat	 Individual participant data Record updated in last year

Plain English summary of protocol

Not provided at time of registration

Study website http://www.fess.nl

Contact information

Type(s) Scientific

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Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers NTR558

Study information

Scientific Title

Acronym FES

Study objectives

FESS is effective: giving significant reduction of symptoms. The indication for FESS must be based on the symptoms of the patient and its duration, computed tomography (CT) scan abnormalities and/or nasal endoscopic abnormalities, and a history of adequate conservative treatment.

Ethics approval required

Old ethics approval format

Ethics approval(s) Received from local medical ethics committee

Study design

Multicentre randomised open label active controlled parallel group trial

Primary study design Interventional

Secondary study design Randomised controlled trial

Study setting(s) Hospital

Study type(s) Treatment

Participant information sheet

Health condition(s) or problem(s) studied Chronic rhinosinusitis (CRS), nasal polyps (NP)

Interventions

The intervention to be investigated is Functional Endoscopic Sinus Surgery (FESS). One treatment arm will receive FESS plus a standardised form of medical treatment. The control group will receive standardised medical treatment. The standardised medical treatment is topical steroids for mild CRS (without NP). In moderate/severe disease a long-term antibiotic is added. The therapy for NP will be corticosteroids. For mild NP therapy is a spray, for moderate disease therapy is a spray and drops, and for severe disease therapy is oral steroids with drops. Specific details are in the protocol.

Intervention Type

Other

Phase Not Specified

Primary outcome measure

The primary outcome measure is a validated disease-specific quality of life questionnaire: SNOT-20.

Secondary outcome measures

The secondary endpoint is re-evaluation of the indication for FESS. Another secondary endpoint will be the standardised evaluation of the nasendoscopy and the CT-scan. For the efficiency assessment 2 secondary endpoints will be evaluated: days of sick-leave and a work-productivity questionnaire.

Overall study start date

01/01/2006

Completion date

01/01/2008

Eligibility

Key inclusion criteria

1. Males or females aged 18 years old can participate

2. Diagnosis CRS with/without NP (definition according to the European Position Paper on Rhinosinusitis and Nasal Polyposis [EPOS])

3. Prior treatment as defined in the treatment scheme of the protocol for at least 12 weeks

- 4. No prior sinus surgery
- 5. Indication for FESS, both criteria must be met:

5.1. RSOM-31 (add score of magnitude of questions 1, 2, 4, 22 result >9)

5.2. CT score >3 on 1 side at least, judged on a CT-scan made prior to visit 1 and made less than 4 months ago; Lund/Mackay scoring

6. Written informed consent

Participant type(s)

Patient

Age group

Adult

Lower age limit

18 Years

Sex

Both

Target number of participants

160

Key exclusion criteria

1. Cystic fibrosis

- 2. Gross immunodeficiency (congenital or acquired)
- 3. Congenital mucociliary problems e.g. primary ciliary dyskinesia (PCD)
- 4. Non-invasive fungal balls and invasive fungal disease
- 5. Systemic vasculitis and granulomatous diseases
- 6. Patients who have any serious or unstable concurrent disease
- 7. Any structural nasal abnormalities (other than polyps or chronic sinusitis) e.g. severe nasal septum deviation
- 8. Rhinosurgery during the past 6 weeks
- 9. Systemic steroids 4 weeks before the study

10. Medication affecting nasal mucosa (cyclosporin, ß-blocker, ACE inhibitors, non-steroidal antiinflammatory drugs [NSAIDs], reserpine, guanethidinge, phenolomine, methyldopa, alfaadrenoceptor antagonist and chlorpromazine)

- 11. Medication other than trial medication
- 12. Females who are pregnant or lactating

13. Inability to follow the instructions within this protocol or known inability to attend ALL clinical visits within the intervals stated

Date of first enrolment

01/01/2006

Date of final enrolment

01/01/2008

Locations

Countries of recruitment Netherlands

Study participating centre Academic Medical Center, Amsterdam Almere Netherlands 1100 DD

Sponsor information

Organisation Academic Medical Centre (AMC) (Netherlands)

Sponsor details

ENT Department P.O. Box 22660 Amsterdam Netherlands 1100 DD

Sponsor type Hospital/treatment centre

ROR https://ror.org/03t4gr691

Funder(s)

Funder type Hospital/treatment centre

Funder Name

Academic Medical Centre (AMC) (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 summary

Not provided at time of registration