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Study protocol for a randomised controlled multicentre study: the Foraminotomy ACDF Cost-Effectiveness Trial (FACET) in patients with cervical radiculopathy

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ABSTRACT

Introduction: Cervical radiculopathy due to discogenic or spondylotic stenosis of the neuroforamen can be surgically treated by an anterior discectomy with fusion (ACDF) or a posterior foraminotomy (FOR). Most surgeons prefer ACDF, although there are indications that FOR is as effective as ACDF, has a lower complication rate and is less expensive. A head-to-head comparison of the 2 surgical techniques in a randomised controlled trial has not yet been performed. The study objectives of the Foraminotomy ACDF Cost-Effectiveness Trial (FACET) study are to compare clinical outcomes, complication rates and cost-effectiveness of FOR to ACDF.

Methods and analysis: The FACET study is a prospective randomised controlled trial conducted in 7 medical centres in the Netherlands. The follow-up period is 2 years. The main inclusion criterion is a radiculopathy of the C4, C5, C6 or C7 nerve root, due to a single-level isolated cervical foraminal stenosis caused by a soft disc and/or osteophytic component, requiring operative decompression. A sample size of 308 patients is required to test the hypothesis of clinical non-inferiority of FOR versus ACDF. Primary outcomes are: 'operative success', the measured decrease in radiculopathy assessed by the visual analogue scale and 'patient success', assessed by the modified Odom’s criteria. Secondary outcomes are: Work Ability Index (single-item WAI), quality of life (EuroQol 5 Dimensions 5 level Survey, EQ-5D-5L), Neck Disability Index (NDI) and complications. An economic evaluation will assess cost-effectiveness. In addition, a budget impact analysis will be performed.

Ethics and dissemination: Ethical approval was obtained from the Institutional Ethics Committee of the University Medical Center Groningen. Results of this study will be disseminated through national and international papers. The participants and relevant patient support groups will be informed about the results of the study.

Trial registration number: NTR5536, pre-results.

Strengths and limitations of this study

- The Foraminotomy ACDF Cost-Effectiveness Trial (FACET) study is the first randomised controlled trial investigating anterior discectomy with fusion versus foraminotomy for cervical foraminal pathology.
- Methodological strengths include an appropriate sample size based on a non-inferiority design, seven participating medical centres and an independent institute facilitating the randomisation system and monitoring activities.
- The study is performed with care as usual; therefore, the final results of this study could be generalisable to usual clinical practice.
- With respect to the evaluation of spinal fusion and instrumentation failure, the relatively short follow-up period of 2 years could be a limitation of this study.
- In this study, it is not feasible to blind the participant and the surgeon to the allocated treatment.

INTRODUCTION

Background and rationale

Cervical radiculopathy has a serious impact on a patient’s quality of life (QoL). Many patients with a cervical radiculopathy are in their productive phase of life and are part of their community’s labour force. Therefore, temporary disability due to the radiculopathy may result in a major loss of productivity.

A cervical radicular syndrome (CRS) constitutes pain, paresis and/or paraesthesia in the distribution area of the corresponding nerve root. It can be caused by nerve root compression due to soft disc protrusion or by a spondylotic stenosis of the neuroforamen. The annual incidence of a CRS is
estimated at 0.8 per 1000 inhabitants. The initial treatment is primarily conservative. The revised Dutch CRS guideline (2010) states that a CRS existing for longer than 2 months, and not responding to conservative treatment in that time course, can be considered an indication for surgical treatment.2

Various surgical approaches for the treatment of cervical radiculopathy have been described. Spurling and Scoville6 reported the successful removal of the cervical disc with a hemilaminectomy in 1944. In the following years, the techniques further evolved into a keyhole foraminotomy (FOR) being popularised by Frykholm.4

4. Indirect costs of FOR seem lower, which may be related to a faster rehabilitation and an earlier resumption of work. However, this assumption is based on two retrospective studies comparing the cost-effectiveness of FOR to ACDF with plating. A comparison to ACDF without plating has not yet been made.20 21

In summary, a high-quality study comparing the effectiveness of FOR to ACDF has not yet been performed. Patients, the medical community and stakeholders would benefit from a study that puts both procedures to the test. The results of this study could be of use for the update/revision of the (inter)national guidelines for the treatment of a CRS.

Aims and hypotheses

The Foraminotomy ACDF Cost-Effectiveness Trial (FACET) study is designed as a high-quality study to analyse the cost-effectiveness of the FOR technique compared with ACDF in patients with a cervical monoradiculopathy caused by a discogenic and/or osteophytic foraminal stenosis.

The primary hypotheses are:

▸ The effectiveness of the FOR technique is non-inferior compared with the ACDF technique.

▸ The FOR technique is cost-effective compared with the ACDF technique.

The secondary hypotheses are:

▸ The FOR technique will have a lower complication rate compared with the ACDF technique.

▸ The FOR technique will have lower direct and indirect costs compared with the ACDF technique.

▸ The FOR technique is associated with more neck pain in the first 30 days after the surgical procedure.
Trial design
The FACET study is a nationwide, prospective, multicentre, investigator-blinded randomised controlled trial with a follow-up of 2 years. A total of 308 participants will be included in seven medical centres in the Netherlands during 2 years. Both the FOR (experimental group) and the ACDF (active control) are established surgical techniques. A non-inferiority trial design was chosen to show whether FOR has at least as much efficacy as the ACDF technique or is worse by an amount <10% with regard to the primary outcome parameters.

METHODS AND ANALYSIS

Study setting
The study will focus on patients with a monosegmental radicular syndrome due to a lateral or foraminal herniated disc or osteophyte, with compression of the C4, C5, C6 or C7 nerve root. Patients will be informed about the study at the outpatient clinics of the participating hospitals. Participating neurosurgeons are from the University Medical Center Groningen, Martini Hospital Groningen, Radboud University Medical Center Nijmegen, Canisius Wilhelmina Hospital Nijmegen, Zuyderland Medical Center Heerlen, Medical Center Haaglanden and Medisch Spectrum Twente Enschede. Two centres are university teaching hospitals, and five are large regional medical centres. The participating hospitals were selected because of the high volume of spinal cases and their neurosurgeons’ familiarity with the two procedures under investigation. The participating centres are evenly distributed over the Netherlands, which will result in a study population that reflects the population of the entire country.

Patient inclusion criteria
In order to be eligible to participate in this study, a participant must meet all of the following criteria:

- Age between 18 and 80 years.
- Single-level isolated cervical foraminal stenosis due to a soft disc or osteophytic component causing radiculopathy of C4, C5, C6 or C7 and requiring decompression of neuroforamen. (Foraminal stenosis due to a soft disc component is defined as: 2/3 of the total discal component is located intraforaminally and a maximum of 1/3 of the total discogenic component is located medially within the spinal canal. Radiculopathy is defined as pain, paresis and/or paraesthesia in corresponding nerve root distribution areas of C4, C5, C6 or C7, and must include at least arm or shoulder pain with a minimum of 30 mm on a 100 mm visual analogue scale (VAS).)
- Failure to respond to conservative treatment for 8 weeks or, during these 8 weeks, progressive signs or symptoms of nerve root compression.
- Discogenic or spondylotic foraminal stenosis (determined by MRI or CT, with or without an oblique X-ray of the cervical spine) at the anatomical level that correlates with the clinical symptoms.
- Psychologically and physically able to comply with the study-related procedures.
- Sufficient mastery of the Dutch language to fill out the questionnaires.
- Signed and dated informed consent form, prior to any study-related procedures.

Investigational treatment

*ACDF technique.*
Microsurgical discectomy is performed by a ventral approach described by Smith and Robinson. The procedure can be executed with microscope or loupe magnification. The content of the intervertebral disc is exposed and removed. The posterior ligament is incised and removed with rongeurs. Disc fragments that are extruded into the neuroforamen are removed. If an osteophytic component is present, this part of the unco- vertebral joint is reduced. An intervertebral spacer (cage or PMMA) is inserted into the intervertebral disc space. No additional plate fixation is used.

*FOR technique.*
All patients are operated in the prone position with the head in a three-point head holder. After fluoroscopy to determine the correct level, a vertical midline incision is made, and the ipsilateral lamina and facet joints are exposed. After a second confirmation of the correct level, a partial hemilaminectomy and FOR of the involved level is performed with the use of an operating microscope or loupe magnification. The amount of reduction of the facet joint is determined by the extent of...
of the foraminal stenosis. In cases of soft disc compressions, the proximal root is visualised adequately and mobilised to allow the removal of the compressing disc material. However, complete removal of the soft disc component is not strictly necessary and the decision to remove the soft component will be left to the surgeon’s preference. In cases of an osteophytic foraminal stenosis, bony decompression of the proximal root is performed. The surgeon has the choice to carefully coagulate and divide the venous plexus that covers the nerve sheath.

**Main study parameters**

The primary study parameters are ‘operative success’ and ‘patient success’.

‘Operative success’ will be the postoperative decrease in radiculopathy assessed by the VAS for self-reported arm pain, between the patients operated with the FOR technique and with the ACDF technique during 24 months of follow-up. We will consider the decrease in radiculopathy clinically relevant when the postoperative decrease is more than 41 mm on the VAS for arm pain.

‘Patient success’ will be assessed by the modified Odom’s criteria, which address physical symptoms and socioeconomic status. We will consider the first and second categories (‘excellent’ and ‘good’) as a successful outcome of the operation.

The distinction between ‘patient success’ and ‘operative success’ is made because a patient could have full improvement of the radiculopathy causing arm pain, but would not be satisfied with the operation because he or she is not able to perform his or her daily activities, for example, because of a persistent sensibility disorder in the fingertips.

**Secondary study parameters are:**

1. Differences in postoperative work ability during 24 months of follow-up.
2. Differences in postoperative QoL during 24 months of follow-up.
3. Changes and differences in neck pain in the short term after the procedure (weekly assessed in the first 6 weeks postoperative) and during the 24 months of follow-up.
4. Differences in neck disability during 24 months of follow-up.
5. Type of complications and complication rates in the short-term (30 days) and long-term (24 months) periods.
7. Budget impact (extrapolated to 5 years).

**Participant timeline**

The inclusion of participants has started on 1 December 2015. The total follow-up is 2 years. At baseline (ie, at enrolment, before the surgical procedure) and at 6, 26, 52, 78 and 104 weeks after the surgical procedure, participants will fill out web-based questionnaires. These questionnaires take ~30 min to fill in. A short questionnaire, addressing only the arm and neck pain (VAS), will be used weekly from the surgical intervention until the visit to the outpatient clinic after 6 weeks.

The participant will visit the outpatient clinic 6 weeks after the surgical procedure, in line with standard care. An independent interviewer will contact the participants by telephone at 26, 52, 78 and 104 weeks to assess Odom’s criteria. For a schematic diagram, see table 1.

**Sample size calculation**

The sample size analysis is based on the manuscript of Dohrmann and Hsieh. Based on this review of cohort studies, an overall success rate of 87% for both groups was detected. Based on the current guidelines of the CRS, there is evidence that FOR is not worse in comparison with ACDF concerning decrease in arm pain.

Therefore, the assumption is that there will not be a statistically significant difference between the two surgical techniques concerning the primary outcome measure (relief of arm pain). With the aforementioned success rate in mind, an $\alpha$ of 0.05, power of 0.8 and a non-inferiority margin ($\delta$) of 10%, the authors conducted a sample size calculation. Including 10% drop outs, a total sample size of 308 participants was calculated, which means 154 participants per treatment arm.

**Recruitment**

Enrolment will take place at the outpatient clinics of the participating medical centres. When a patient with a CRS is indicated for a surgical treatment, the attending neurosurgeon judges if the patient is eligible for the trial. If the patient fulfils the inclusion and exclusion criteria, he or she is informed about the possibility to participate in the study.

To determine if the inclusion rate of 308 participants is feasible, we asked 25 patients (with a cervical radiculopathy due to a foraminal compression) in one of the outpatient clinics whether they would participate in a surgical randomised controlled trial. More than 80% of the patients were willing to participate. Furthermore, it was calculated that with a low inclusion rate of 30% each year, the authors will be able to include 308 participants in 2 years. For 7 participating centres, this implies an inclusion rate of ~2 participants per centre per month.

**Randomisation and treatment allocation**

Participants are randomised to either FOR or ACDF. Randomisation will be executed per participant per centre, by web-based block randomisation after the informed consent procedure is fulfilled. An independent institute, the Trial Coordination Center of the University Medical Center Groningen, will facilitate the randomisation.

**Blinding**

In this trial, blinding of the participant or the surgeon is not feasible. The data analysis will be performed with blinded data. Assessment of Odom’s criteria will be...
performed by an independent interviewer who is blinded to the treatment allocation.

**Data collection methods**

Preoperative history is obtained from standard care procedures and includes length, weight, number of months/years of neck and arm pain, signs and symptoms, other significant illnesses, pain medication (use of non-steroidal anti-inflammatory drugs) and smoking history. Information about the operative procedure will be obtained from the medical record of the participant and will include date and type of procedure, which level was operated, use of implants and the occurrence of complications during the operative procedure.

At baseline, discharge and 6 weeks after the procedure, a clinical evaluation will be performed. Information will be recorded on signs and symptoms, reflexes, sensibility and strength.

**Outcome measurements**

**Primary outcomes:**

‘Operative success’ will be the postoperative decrease in radiculopathy assessed by theVAS for self-reported arm pain, between the patients operated with the FOR technique and with the ACDF technique during 24 months of follow-up. The VAS is a well-known and validated outcome measurement. It is provided to the participants as a horizontal scale ranging from 0 to 100 mm.

‘Patient success’ will be assessed by the modified Odom’s criteria, which address physical symptoms and socioeconomic status. Odom’s criteria have been widely used in studies regarding different cervical procedures and assess the improvement of the physical symptoms and the ability to perform daily activities. Although Odom’s criteria have not been thoroughly validated yet, they have been correlated with the VAS arm pain (p=0.002) and VAS neck pain (p<0.0005). There is also a correlation between Odom’s criteria and two QoL assessments for spinal disorders: the Million VAS (p<0.0005) and the Oswestry Disability Index (p<0.0005).

In the original article, Odom’s criteria are defined differently for each item. Therefore, we modified Odom’s criteria in analogue terms. The investigator will categorise the participant, with regard to his or her symptoms and neurological status. We will consider the first and second categories (‘excellent’ and ‘good’) as a successful outcome of the operation.

**Modified Odom’s criteria**

1. Excellent: no symptoms referable to cervical disease. Able to perform daily activities without limitations.
2. Good: moderate symptoms referable to cervical disease. Able to perform daily activities without significant limitations.
4. Poor: no improvement or aggravation of symptoms referable to cervical disease. Not able to perform daily activities.

Secondary outcomes:

QoL will be assessed with the EuroQol 5 Dimensions 5 level Survey (EQ-5D-5L). This survey ranks the QoL over five dimensions in a five-level rating scale. The scores can be converted in quality-adjusted life years (QALYs). An improvement of 0.24 QALY will be considered clinically relevant. For neck disability, the Neck Disability Index (NDI) will be used. This questionnaire consists of 10 items, which can be ranked from 0 to 5. The sum of these scores can be doubled to get a percentage. An improvement of 17.3% will be considered clinically relevant. Work ability will be assessed with the single-item Work Ability Index (WAI). A participant rates his or her current ability to work, as opposed to the lifetime best ability, on a scale from 1 to 10.

Cost-effectiveness and the budget impact analysis (BIA) are calculated by means of questionnaires for medical consumption and productivity costs. The National Health Care Institute of the Netherlands mentions these questionnaires as the preferred measurement tools for calculating economic evaluations. The questionnaires will be provided to the participant in a secure online environment. If a participant does not have access to the internet, paper questionnaires will be sent.

Withdrawal of individual participants

Participants can discontinue the participation of the study at any time for any reason. The investigator can decide to withdraw a participant from the study for urgent medical reasons (not related to the treatments under investigation).

The aim is to include 308 participants within 2 years. When a participant withdraws from the study, he or she will not be replaced since a 10% loss of inclusion is accounted for within the power calculations of the trial.

If participants withdraw after they have received the operative treatment, their permission will be asked to use the information from their medical files. Data of participants after withdrawal will be used and analysed with the intention-to-treat principle. A per-protocol analysis will be performed as well.

Data management

Data will be recorded in a web-based data capture system (OpenClinica), which is hosted by the Trial Coordination Center of the University Medical Center Groningen. This system is customised and has an audit trail facility. Ranges and programmed validation checks are implemented in the system in order to aid reliable data entry.

Statistical methods

Primary study parameters

The primary end point (postoperative decrease of self-reported arm pain (VAS)) will be defined as ‘operative success’. As a second primary end point, Odom’s criteria will be assessed. Odom’s criteria will be defined as ‘patient success’. Both items will be analysed as appropriate depending on data distribution with a one-sided 0.05 level of significance (non-inferiority). We consider the FOR technique to be non-inferior if the technique has at least as much efficacy (decrease in VAS arm pain and the good/excellent scores on Odom’s criteria) as the ACDF technique or is worse by an amount <10%.

Detailed descriptive statistics will be provided for the data collected and 95% CIs will be calculated for all relevant estimates. Clinical follow-up data will be analysed by analysis of covariance or generalised model alternatives for categorical or semiquantitative data. Changes within the treatment groups over time, as well as differences between groups, will be assessed by intention-to-treat analyses. Also, the primary analysis will follow the per-protocol principle. Sensitivity analysis will be provided to evaluate robustness of the results with regard to unexpected circumstances, for example, the impact of ‘crossover’ and centre-related effects.

Secondary study parameters

Secondary end points will be analysed in an exploratory manner at a two-sided significance level of 5%. Safety and tolerability parameters will be analysed descriptively. Analysis of time-dependent probabilities of critical events will be performed using the Kaplan-Meier method. Furthermore, multivariate event analyses will be performed using Cox proportional hazard regression models. In addition, for the purpose of a supportive sensitivity analysis, multiple imputation procedures will be applied.

The cost-effectiveness analysis will be performed alongside the clinical trial to assess the cost-effectiveness of FOR versus ACDF. There will be two separate outcome measures for the cost-effectiveness analysis, resulting in two incremental cost-effectiveness ratios for FOR as compared with ACDF. The first is incremental costs per extra percentage of patients with arm pain relief, while the second is incremental costs per QALY gained. EQ-5D-5L scores will be converted into health state utilities using the Dutch value set. These health state utilities range between 0.446 and 1, with a higher utility indicating a better health-related QoL. The utilities will be multiplied by follow-up time spent in that particular health state (area under the curve) to eventually convert into QALYs.

The analysis will be performed taking a societal perspective. The time horizon will be equivalent to the full
follow-up of the clinical study, which is 24 months. According to pharmacoeconomic guidelines, discounting will be applied for costs (4%) and effects (1.5%) in the second year. A number of sensitivity analyses will be performed to identify the impact of variables, such as the costs of the FOR and ACDF procedures, resource use and effect size, on cost-effectiveness. A cost-effectiveness acceptability curve will be constructed, based on bootstrap simulations, showing the probability of FOR being cost-effective compared with ACDF at varying levels of the willingness to pay, for either one additional percentage of patients with arm pain relief or one additional QALY.

Based on the results of the clinical study and the cost-effectiveness analysis, a BIA will be performed to inform decision makers on the financial consequences of implementing FOR as treatment of first choice for cervical foraminal soft disc/osteophytic disease in the Dutch healthcare system. The BIA will be performed according to the principles of the International Society for Pharmacoeconomics and Outcomes Research (ISPOR) task force and from the perspective of the government as well as a third party payer’s/healthcare insurer’s perspective. The trial results will be extrapolated, by means of a simple model, from a time horizon of 24 months to 5 years, and for the entire Dutch population concerned. The extrapolation will assume a constant incidence of cervical foraminal soft disc/osteophytic disease. We expect that the proportion of cases eligible for surgical intervention will be stable over time as well. Therefore, the extrapolation will be linear. Sensitivity analyses will be performed on relevant parameters such as the eventual substitution rate of FOR versus ACDF (may not be 100%), the uptake of FOR with time, costs of the procedures and other cost items. We will assume that current usual care already consists of a mix of ACDF and FOR, that is, 90% and 10%, respectively.

Monitoring
The study will be monitored by the Trial Coordination Center of the University Medical Center Groningen. This independent institute will examine the execution of the study procedures and verify source data by random sampling in the seven participating medical centres. At least one monitoring visit per year per centre will be conducted. All protocol deviations will be documented. The exact details of the monitoring will be documented in a monitoring plan in accordance with the guidelines for a low-risk to moderate-risk study.

The investigational treatments are widely accepted and frequently performed surgical procedures by neurosurgeons and orthopaedic surgeons. Participating in the study does not cause any additional risk to the participant. Therefore, the investigators decided to not establish a data monitoring committee. In the Netherlands, liability insurance is available for all participants as insurance is mandatory in the conduction of clinical trials.

Box 1  List of adverse events for the FACET study

- Death
- Thrombosis
- Pulmonary embolism
- Urinary retention
- Postoperative bleeding/haematoma
- Postoperative wound infection
- Nerve root injury
- Dural tear
- Horner’s syndrome
- Blood loss from the wound
- Urinary infection
- Spinal cord injury
- Oesophagus injury
- Hoarseness
- Pneumonia
- Reoperation at the index level
- Reoperation at adjacent level(s)
- Other

During the complete period of the study, all adverse events will be reported. Adverse events are defined as any undesirable experience occurring to a participant during the study, whether or not considered related to intervention. The events that will be assessed are listed in box 1.

The definition of serious adverse events is in line with the guidelines of the International Council on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use (ICH). Every adverse event will be categorised as serious if it:

- Results in death;
- Is life threatening (at the time of the event);
- Requires hospitalisation or prolongation of existing inpatients’ hospitalisation;
- Results in persistent or significant disability or incapacity;
- Is a congenital anomaly or birth defect;
- Concerns any other important medical event that may not result in death, be life threatening or require hospitalisation may be considered a serious adverse experience when, based on appropriate medical judgement, the event may jeopardise the participant or may require an intervention to prevent one of the outcomes listed above’.

If a serious adverse event occurs, the local investigator is obliged to give direct notice (<24 hours) to the sponsor. According to the Medical Research Involving Human Subjects Act, the sponsor has to report the event in 7–15 days to the reviewing committee.

ETHICS
This study will be conducted in accordance with the Declaration of Helsinki and in compliance with the Medical Research Involving Human Subjects Act.
Ethical approval has been granted from the Institutional Ethics Committee of the initiating medical centre on 5 November 2015.

Consent
A participant’s information folder in adjusted linguistics is offered to all participants containing information about the study aims, procedures and the risks of both surgical techniques. In a conversation with the attending physician, questions regarding the study will be answered. If the patient is fully informed and willing to participate, the informed consent form is signed. The patient is provided with ample time to consider his or her decision and to ask for additional information. No maximum amount of days is set; patients can consider their decision until the surgical treatment is scheduled.

Confidentiality
The data of the participant will be recorded and analysed without any personal identifiers, by using coded information. The source documents and identification lists will be archived in a secured facility per centre. Permission for accessing data will be documented per investigator.

Dissemination
Results of this study will be disseminated through national and international papers. The participants and relevant patient support groups will be informed about the results of the study.

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Contributors
JMAK and RS initiated the study. JMAK, RS, RJMG, MFR, ADiVA, JMCdV and PCCAIV designed the FACET study. JMAK, GATL-L, RS and ADiVA constructed the first draft of the protocol. AEBH and JMAK further edited the protocol conform SPIRIT guidelines. RHMAB; MRG, JK, HvS, NH and MA critically reviewed the protocol and the study procedures. All authors have given final approval for the manuscript.

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Competing interests
None declared.

Ethics approval
Medical Ethics Committee University Medical Center Groningen, 5 November 2015.

Provenance and peer review
Not commissioned; externally peer reviewed.

Data sharing statement
For access to the data set, a formal request should be sent to the FACET study group. The request will only be considered when the primary results of the study have been published.

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