# Pressure Guidewire Comparative In Patients Study

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Compare the stability of the FFR measurement and the handling performance of the St. Jude

Pressure Wire X compared to the Opsens OptoWire Deux.

**Ethical review** Approved WMO **Status** Recruiting

**Health condition type** Coronary artery disorders **Study type** Observational invasive

# **Summary**

#### ID

NL-OMON45355

Source

ToetsingOnline

**Brief title** 

FFR COMP-IP-02 study

## **Condition**

Coronary artery disorders

## **Synonym**

Coronary artery disease, Intravascular blood pressure

## Research involving

Human

# **Sponsors and support**

**Primary sponsor:** Jeroen Bosch Ziekenhuis

Source(s) of monetary or material Support: Opsens Inc., TOP Medical BV Elsloo

#### Intervention

**Keyword:** Diagnostic angiography, Drift, Fractional flow reserve (FFR), intravascular blood

## **Outcome measures**

## **Primary outcome**

The FFR measurement reliability is assessed by two measures. First reliability is assessed by tracking and recording the drift caused by the pressure guidewire residing within the coronary artery. This is determined by the value of Pd/Pa before and after the procedure when pulled back just outside the guiding catheter.

Secondly, FFR reliability is assessed by the stability of Pd/Pa during constant hyperemia.

## **Secondary outcome**

The wire handling performance is assessed by scoring both wires by the same investigator in the same patient. The following characteristics are scored on a 5 point scale: torquability, steerability, pushability and support. The wire performance is also assessed by measuring the time it takes to reach and cross the lesion.

# **Study description**

#### **Background summary**

Fractional Flow Reserve (FFR) is an index that is used to determine the functional significance of coronary artery disease (CAD) during cardiac catheterization. FFR requires measurement of aortic and distal coronary pressure. The latter is usually determined by a pressure sensor mounted guide wire. The challenge with current pressure guidewires is that they measure pressure using piezo-resistive pressure sensors that are sensitive to moisture

which influences the stability of the transmitted pressure signal. Potentially, unreliable measurements with consequent erroneous clinical decisions could be made. Moreover, wire handling with these pressure mounted guide wires is different from regular guidewires due to the design. This fact limits adoption of functional assessment of CAD in the cathlab. Therefore, there is a clinical need for a FFR wire providing reliable pressure measurement and whose performance would be closer to standard angioplasty-wire.

## **Study objective**

Compare the stability of the FFR measurement and the handling performance of the St. Jude Pressure Wire X compared to the Opsens OptoWire Deux.

## Study design

Observational study, two-center, two-arm study

## Study burden and risks

In this study the small additional small risk arises from the elongation of the procedure due to the repetition of the FFR measurement. FFR measurement is standard of care and has a Class 1A recommendation for lesion assessment pre-percutaneous coronary intervention in the European Society of Cardiology guidelines for revascularization. The risks of a FFR measurement are small in experienced hands and almost always temporary. There is no direct benefit for the patient when participating in this study.

# **Contacts**

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# **Trial sites**

## **Listed location countries**

**Netherlands** 

# **Eligibility criteria**

#### Age

Adults (18-64 years) Elderly (65 years and older)

## Inclusion criteria

Patients with at least one lesion indicated for FFR

## **Exclusion criteria**

More than 3 lesions indicated for FFR measurements Type C lesions Lesions with angiographic 'haziness' or suspected to contain thrombus

# Study design

# **Design**

Study type: Observational invasive

Intervention model: Other

Allocation: Non-randomized controlled trial

Masking: Open (masking not used)

Control: Active

Primary purpose: Diagnostic

## Recruitment

NL

Recruitment status: Recruiting

Start date (anticipated): 16-03-2018

Enrollment: 20

Type: Actual

## Medical products/devices used

Generic name: Optical FFR Guidewire

Registration: Yes - CE intended use

# **Ethics review**

Approved WMO

Date: 19-12-2017

Application type: First submission

Review commission: METC Brabant (Tilburg)

# **Study registrations**

# Followed up by the following (possibly more current) registration

No registrations found.

# Other (possibly less up-to-date) registrations in this register

No registrations found.

# In other registers

Register ID

CCMO NL60721.028.17