

# **PE-Prove Study PROMUS\* Element\* Everolimus-Eluting Coronary Stent System European Post-Approval Surveillance Study**

## **A prospective, open label, multi-center observational study**

Published: 03-02-2011

Last updated: 03-05-2024

Please see pages 3-5 of the Protocol

<b>Ethical review</b>	Approved WMO
<b>Status</b>	Recruitment stopped
<b>Health condition type</b>	Coronary artery disorders
<b>Study type</b>	Observational invasive

## **Summary**

### **ID**

NL-OMON34092

### **Source**

ToetsingOnline

### **Brief title**

PE-Prove study

### **Condition**

- Coronary artery disorders

### **Synonym**

Coronary artery disease ; Narrowing of arteries

### **Research involving**

Human

## Sponsors and support

**Primary sponsor:** Boston Scientific International S.A.

**Source(s) of monetary or material Support:** Industry

## Intervention

**Keyword:** Coronary artery disease, Drug-eluting stent, Percutaneous Transluminal Coronary Angioplasty

## Outcome measures

### Primary outcome

Please see pages 3-5 of the Protocol

### Secondary outcome

Please see pages 3-5 of the Protocol

## Study description

### Background summary

Please see pages 3-5 and 9 of the Protocol

### Study objective

Please see pages 3-5 of the Protocol

### Study design

Please see pages 3-5 of the Protocol

### Study burden and risks

As all procedures are part of the routine care the only risk in participating in this study is the additional blood test.

The risks associated with providing a blood sample can include fainting, pain (a painful jab as the needle is inserted into the arm) and/or bruising. In rare cases, it is also possible for a blood clot to form or for an infection to

occur at the site where the needle was inserted.

## Contacts

### Public

Boston Scientific International S.A.

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FR

### Scientific

Boston Scientific International S.A.

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FR

## Trial sites

### Listed location countries

Netherlands

## Eligibility criteria

### Age

Adults (18-64 years)

Elderly (65 years and older)

### Inclusion criteria

All patients who are candidates for Coronary artery stenting, signed the Informed Consent Form and are eligible to receive a PROMUS\* Element\* stent will be evaluated for enrollment in this study.

### Exclusion criteria

No study-specific exclusion criteria apart from any contra-indications

## Study design

### Design

Study phase:	4
Study type:	Observational invasive
Masking:	Open (masking not used)
Control:	Uncontrolled
Primary purpose:	Other

### Recruitment

NL	
Recruitment status:	Recruitment stopped
Start date (anticipated):	21-03-2011
Enrollment:	90
Type:	Actual

## Ethics review

Approved WMO	
Date:	03-02-2011
Application type:	First submission
Review commission:	IRB Nijmegen: Independent Review Board Nijmegen (Wijchen)
Approved WMO	
Date:	28-02-2011
Application type:	Amendment
Review commission:	IRB Nijmegen: Independent Review Board Nijmegen (Wijchen)
Approved WMO	
Date:	05-05-2011
Application type:	Amendment
Review commission:	IRB Nijmegen: Independent Review Board Nijmegen

(Wijchen)

## 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
ClinicalTrials.gov	NCT01148329
CCMO	NL33913.072.10