# A Prospective, Nonrandomized, Noninterventional Study to Compare Nexfin CO-trek Cardiac Output with Thermodilution Cardiac Output

Published: 30-08-2013 Last updated: 24-04-2024

Demonstrate that noninvasive monitoring of cardiac output with the ccNexfin System is comparable to that with thermodilution cardiac output.

Ethical reviewApproved WMOStatusRecruitment stoppedHealth condition typeOther condition

**Study type** Observational non invasive

# **Summary**

#### ID

NL-OMON38805

#### **Source**

ToetsingOnline

#### **Brief title**

Nexfin/TD Clinical Study

#### Condition

Other condition

#### **Synonym**

N.A.

#### **Health condition**

specifieke aandoeningen zijn niet van belang voor het onderzoek; daar een clinische standaard methode voor cardiac output bepaling aanwezig moet zijn als referentie, zal het onderzoek worden uitgevoerd in high-risk surgery Research involving

Human

Sponsors and support

**Primary sponsor:** Edwards Lifesciences BMEYE

Source(s) of monetary or material Support: Edwards Lifesciences LLC

Intervention

**Keyword:** Cardiac Output, ccNexfin, Comparison, Noninvasive

**Outcome measures** 

**Primary outcome** 

Noninvasive monitoring of CO with the ccNexfin System is comparable to TD as

determined by a bias less than 0.6 L/min.

**Secondary outcome** 

Comparability of both methods as determined by Bland-Altman analysis (Bias and

Percentage Error). Precision of Nexfin CO-trek versus TD will be determined.

The Pearson correlation coefficient for the CO pairs of both methods will be

assessed. In patients where Trendelenburg / reverse Trendelenburg positions are

clinically required, changes in CO due to these interventions on hemodynamics

will be measured to demonstrate concordance of devices in case of repeated

measurement. Fluid administration, use of vasoactive and inotropic drugs will

be recorded when available to the Investigator.

**Study description** 

**Background summary** 

Noninvasive measurement of cardiac output using ccNexfin allows monitoring of almost any patient in the OR. Consequently, fluids for individual patients may

2 - A Prospective, Nonrandomized, Noninterventional Study to Compare Nexfin CO-trek ... 18-05-2025

be managed more appropriately, leading to less post-operative complications and morbidity.

#### Study objective

Demonstrate that noninvasive monitoring of cardiac output with the ccNexfin System is comparable to that with thermodilution cardiac output.

#### Study design

Prospective, nonrandomized, noninterventional validation study.

#### Study burden and risks

Burden and risks are negligible. The study is not group related.

## **Contacts**

#### **Public**

**Edwards Lifesciences BMEYE** 

Hoogoorddreef, Centerpoint 1, 4e verdieping 60 Amsterdam Zuidoost 1101 BE NL

#### Scientific

Edwards Lifesciences BMEYE

Hoogoorddreef, Centerpoint 1, 4e verdieping 60 Amsterdam Zuidoost 1101 BE NL

## **Trial sites**

#### **Listed location countries**

**Netherlands** 

# **Eligibility criteria**

#### Age

3 - A Prospective, Nonrandomized, Noninterventional Study to Compare Nexfin CO-trek ... 18-05-2025

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

#### Inclusion criteria

Subjects will be included if they meet the following criteria:

- 1. Subjects must be at least 18 years of age
- 2. Subjects must give signed written informed consent
- 3. Subjects\* height and weight must be accurately obtained prior to study start.

#### **Exclusion criteria**

Subjects will be excluded if any of these items exist:

- 1. Aortic or tricuspid valve regurgitation
- 2. Aortic stenosis or aneurysms
- 3. History of uncontrolled cardiac arrhythmia
- 4. Any peripheral vascular disease or conditions such as Raynaud\*s disease or Buerger\*s disease
- 5. Insufficient perfusion of the digits
- 6. Inability to place the finger cuff appropriately due to subject anatomy or condition
- 7. Known pregnancy
- 8. Patients being treated with an intra-aortic balloon pump
- 9. Patient is currently participating in an investigational drug or another device study that clinically interferes with the study endpoints

# Study design

### **Design**

**Study type:** Observational non invasive

Masking: Open (masking not used)

Control: Uncontrolled

Primary purpose: Other

#### Recruitment

NL

Recruitment status: Recruitment stopped

Start date (anticipated): 11-12-2013

Enrollment: 40

Type:	Actua

# **Ethics review**

Approved WMO

Date: 30-08-2013

Application type: First submission

Review commission: METC Amsterdam UMC

Approved WMO

Date: 26-11-2013

Application type: Amendment

Review commission: METC Amsterdam UMC

# **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 NL44270.018.13