# Pancreatitis and early omega-3-fatty acid infusion for reduction of organ failure and mortality: a multicenter randomized controlled trial (PLANCTON)

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This study has been transitioned to CTIS with ID 2023-505220-57-03 check the CTIS register for the current data. The PLANCTON trial will investigate the effect of early intravenous OM-3 FAs on new onset organ failure and mortality in patients with...

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

**Health condition type** Exocrine pancreas conditions

Study type Interventional

## **Summary**

#### ID

**NL-OMON51853** 

#### **Source**

ToetsingOnline

#### **Brief title**

PLANCTON trial

#### Condition

• Exocrine pancreas conditions

#### Synonym

Inflammation of the pancreas

#### Research involving

Human

## **Sponsors and support**

**Primary sponsor:** Radboud Universitair Medisch Centrum

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**Source(s) of monetary or material Support:** Fresenius Kabi (bedrijf),Fresenius Medical Care

#### Intervention

Keyword: Acute pancreatitis, Mortality, Omega-3 fatty acids, Organ failure

#### **Outcome measures**

#### **Primary outcome**

Endpoints will be evaluated after the 180 days study period. The primary endpoint is a composite endpoint of new onset of organ failure (organ failure not present at randomization) and mortality.

#### **Secondary outcome**

Secondary endpoints are individual components of the composite endpoint, severe complications (([infected] pancreas necrosis, sepsis, pneumonia or cholangitis), quality of life, costs effectiveness, number of (surgical, endoscopic or radiologic) interventions, length of hospital and ICU stay.

# **Study description**

### **Background summary**

Acute pancreatitis (AP) is the most common gastrointestinal disorder requiring acute hospitalization. About 20% of all patients will develop severe acute pancreatitis (SAP) marked by a pro-inflammatory response and characterized by massive release of cytokines, which can cause the systemic inflammatory response syndrome (SIRS). SIRS increases the risk on (multi) organ failure and contributes to a mortality up to 30%.

Intravenous omega-3 fatty acids (OM-3 FAs) induce the production of anti-inflammatory cytokines and hereby ameliorate the inflammatory response. We hypothesize that the anti-inflammatory function of OM-3 FAs could attenuate SIRS and decrease the severity of SAP resulting in less (severe) organ failure and ultimately a lower mortality. The clinical implications of this mechanism is shown in a recent meta-analysis describing reduced mortality by the use of OM-3 FAs in 4 randomized trials in patients with acute pancreatitis. However,

the evidence was of low quality and a large multicenter trial on OM-3 FAs in predicted SAP is currently lacking. This could provide definitive proof for the beneficial effect of OM-3 FAs in acute pancreatitis.

#### Study objective

This study has been transitioned to CTIS with ID 2023-505220-57-03 check the CTIS register for the current data.

The PLANCTON trial will investigate the effect of early intravenous OM-3 FAs on new onset organ failure and mortality in patients with predicted SAP.

## Study design

A multicenter randomized controlled trial

#### Intervention

Intravenous administration of a lipid emulsion (0.2g/kg/day) with OM-3 FAs, started within 24hrs of diagnosis of predicted SAP and within 72hrs after onset of symptoms of AP, for a total of 7 days.

#### Study burden and risks

The burden for participants in this study is limited. The risk of OM-3 FAs administration is estimated to be negligible because (serious) adverse events were not described in 14 randomized controlled trials in 551 patients (9 trials in 322 patients with sepsis and 5 trials in 229 patients with acute pancreatitis). Additionally, the known side effects of OM-3 FAs are rare (e.g. lipid overload syndrome and prolonged bleeding time) or of relative limited clinical importance (i.e. the taste of fish). The parenteral administration of OM-3 FAs and questionnaires can be marked as a (small) burden in addition to standard medical care. The benefit for (future) patients treated with OM-3 FAs could be substantial with a reduction in new onset organ failure and mortality.

## **Contacts**

#### **Public**

Radboud Universitair Medisch Centrum

Geert Grooteplein Zuid 10 Nijmegen 6525 GA NL

#### **Scientific**

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Radboud Universitair Medisch Centrum

Geert Grooteplein Zuid 10 Nijmegen 6525 GA NL

## **Trial sites**

#### **Listed location countries**

**Netherlands** 

# **Eligibility criteria**

#### Age

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

#### Inclusion criteria

- Predicted severe acute pancreatitis
- >=18 years old
- First episode of acute pancreatitis
- <24 hours after diagnosis of acute pancreatitis</li>
- <72 hours after onset of symptoms of acute pancreatitis</li>
- Able to read and/or understand the study procedures
- Able to give informed consent (or their legal representatives)

#### **Exclusion criteria**

- Intake of any OM-3 FAs-, krill and/or algae supplements in the week prior to complaints
- Participation in another intervention study for AP
- Organ failure on admission (Modified Marshall score >2)
- Recurrent pancreatitis
- Chronic pancreatitis
- o Defined by the MANNHEIM criteria53
- Known allergy to fish oil, seafood, soja or egg products
- History or existing hyperlipidemia (laboratory proven triglycerides > 10.0 mmol/l)
- History of (severe) liver failure
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- Impaired lipid metabolism may lead to accumulation of fatty acids in the blood, increasing risk of adverse events.
- o Based on coagulation Factor V level or INR > 3 (without anti-coagulation by vitamine K)
- Ketoacidosis
- Acute thrombo-embolic disease
- Pregnancy or lactation
- Recent (< 6 months) myocardial infarction or strokeKnown coagulations disorders

(e.g. Factor V Leiden, thrombocytopenia, etc.)

- Pancreatitis due to a (suspected) periampullary/ampullary or bile duct malignancy
- Other known or suspected malignancy that may interfere with the outcome(s) and/or execution of the PLANCTON trial
- Post ERCP-pancreatitis due to a (suspected) malignancy
- Patient is classified as moribund or expected to die within 24hours o The intervention will not be able to affect this patient and is therefore useless to expose these patients.

# Study design

## **Design**

Study phase: 3

Study type: Interventional

Intervention model: Parallel

Allocation: Randomized controlled trial

Masking: Open (masking not used)

**Primary purpose:** Treatment

#### Recruitment

NL

Recruitment status: Recruiting
Start date (anticipated): 15-07-2022

Enrollment: 212

Type: Actual

## Medical products/devices used

Product type: Medicine

Brand name: Omegaven

Generic name: Omegaven

Registration: Yes - NL outside intended use

## **Ethics review**

Approved WMO

Date: 29-03-2022

Application type: First submission

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 09-05-2022

Application type: First submission

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 28-07-2022

Application type: Amendment

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 26-09-2022

Application type: Amendment

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 03-10-2022

Application type: Amendment

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 18-10-2022

Application type: Amendment

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 15-11-2022

Application type: Amendment

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 05-12-2022

Application type: Amendment

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 11-01-2023

Application type: Amendment

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 08-02-2023

Application type: Amendment

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 15-02-2023

Application type: Amendment

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 16-05-2023

Application type: Amendment

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

# **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

EU-CTR CTIS2023-505220-57-03 EudraCT EUCTR2022-000474-26-NL Register ID

CCMO NL80570.091.22

Other Nummer volgt, ingediend bij ISRTCN