Prevalence of unmet target LDL-C recommendations in very high risk patients despite high intensity lipid modifying therapy

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To assess the prevalence of LDL-C >1.8 mMol/L in a subgroup of very high risk patients with ASCVD, who remain at a very high-residual risk for ACS, despite treatment with high-intensity statins in combination with ezetimibe. This subgroup of very...

Ethical review Approved WMO

Status Recruitment stopped

Health condition type Arteriosclerosis, stenosis, vascular insufficiency and necrosis

Study type Interventional

Summary

ID

NL-OMON55816

Source

ToetsingOnline

Brief title

LDL-C in very high risk patients

Condition

• Arteriosclerosis, stenosis, vascular insufficiency and necrosis

Synonym

Hyperlipidaemia/ high cholesterol post myocardial infarction

Research involving

Human

Sponsors and support

Primary sponsor: WCN (Werkgroep Cardiologische centra Nederland)

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Source(s) of monetary or material Support: Bedrijf; Sanofi, Sanofi-aventis

Intervention

Keyword: PENELOPE

Outcome measures

Primary outcome

Study endpoint:

- Proportion of patients with LDL-C >1.8 mMol/L during stepwise incremental lipid modifying therapy with respectively a statin, statin + ezetimibe.

Secondary outcome

Study parameters:

- Medical history: ASCVD; T2DM; developing allergies or intolerances to alirocumab, ezetimibe or statins
- LDL-C plasma levels at each consecutive step
- Optional: non-HDL plasma levels at each consecutive step
- Prescription preference for atorvastatin or rosuvastatin to achieve "high intensity statin therapy" (HIST)
- % of patients that tolerates sustained HIST on atorvastatin or on rosuvastatin
- % of patients that does NOT tolerate HIST with atorvastatin, but does tolerate HIST when switching to rosuvastatin
- % of patients that does NOT tolerate HIST with rosuvastatin, but does tolerate HIST when switching to atorvastatin
- % of patients that reaches target LDL <=1.8 mMol/l while NOT on HIST
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- % of patients that compared to baseline achieves 50% LDL reduction ([1] patients not on HIST; [2] patients on HIST; [3] patients on HIST + ezetimibe; [4] patients on HIST + ezetimibe + alirocumab)
- % of patients that compared to baseline achieves 50% LDL reduction AND LDL
 >1.8 OR LDL <=1.8 (patient-groups 1-4)
- non-HDL-C levels in patients with LDL <=1.8 mMol/l and in patients with LDL
 >1.8 mMol/l, with or without alirocumab
- Non-HDL-C levels in patients with LDL <=1.8 mMol/l ánd triglycerides >2 mMol/l, with or without alirocumab
- Prescription preference when atorvastatin 1dd40mg does not meet the LDL-target
- Creatinine kinase in patients with statin intolerance
- % of patients with an LDL <=1.8 mMol/l after one year on HIST
- % of patients with an LDL <= 1.8 mMol/l after one year on HIST + ezetimibe
- % of patients with an LDL <=1.8 mMol/l after one year on HIST + ezetimibe + alirocumab

Components of the TIMI Risk Score for Secondary Prevention (TRS 2 P):

- CHF, hypertension, age (>=75 yr), diabetes, prior stroke, prior CABG, peripheral artery disease, eGFR <60, smoking [12]

Study description

Background summary

Hyperlipidaemia is one of the important risk factors in developing cardiovascular disease. LDL-C of <1.8 mMol/L or a reduction of at least 50% is

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recommended for very high risk patients. Different classes of lipid- modifying drugs are available. In patients with hypercholesterolaemia or combined hyperlipidaemia, statin mono-therapy is first choice therapy. In a large proportion of patients on statin mono-therapy, however, therapeutical target LDL-C levels are not reached, and ezetimibe should be added. If target LDL-C levels are unmet despite statin-ezetimibe combination therapy, a PCSK9 inhibitor (PCSK9-i) may be added.

In real world care, the proportion of patients with atherosclerotic cardiovascular disease (ASCVD) not reaching target LDL-C levels despite consecutive therapy with statin mono-therapy, statin + ezetimibe or statin + ezetimibe + PCSK9-i is unknown.

In the Netherlands and many other European countries, reimbursement of the PCSK9-i is restricted to a subgroup of *very high risk patients* not reaching target LDL-C levels despite statin-ezetimibe combi-therapy.

In the present study, the prevalence of very high risk patients with ASCVD who remain at very high-residual risk for an acute coronary syndrome (ACS), defined as an LDL-C >1.8 mmMol/L (or non-HDL >2.6 mMol/l) will be analysed by a prospective, stepwise implementation of high intensity statin mono-therapy, followed by high intensity statin + ezetimibe combination-therapy if LDL-C still >1.8 mMol/L after 4 weeks of statin mono-therapy. Lipid levels will be measured 4 weeks after initiation of each step. A subset of patients whose LDL-C remains >1.8 mMol/L despite this intervention, will be treated with the PCSK9 inhibitor Alirocumab, 75 or 150 mg, and the efficacy of this therapy will be measured by lipid levels after 4 weeks of therapy.

Study objective

To assess the prevalence of LDL-C >1.8 mMol/L in a subgroup of very high risk patients with ASCVD, who remain at a very high-residual risk for ACS, despite treatment with high-intensity statins in combination with ezetimibe. This subgroup of very high risk patients is defined as patients with a history of ASCVD and/or diabetes mellitus type II (T2DM), and a new type I ST elevation or non-ST elevation myocardial infarction ([N]STEMI).

Study design

A prospective, open label, stepwise cohort study of consecutive patients admitted for type I STEMI or NSTEMI, characterized by a rise and/or fall of troponin with at least one value above the 99th percentile upper reference limit, and a history of ASCVD and/ or T2DM.

Patients with an LDL-C<=1.8 mMol/L at baseline will be registered, but not included in the study. All patients with an LDL-C>1.8 mMol/L, with or without

therapy with statins and/or ezetimibe at baseline, will be treated with high-intensity statin therapy (i.e., atorvastatin >=40 mg or rosuvastatin >=20 mg or the maximum tolerated dose of a statin) for a period of 4 weeks. Lower statin doses are acceptable for patients with a valid reason for not using high intensity doses (advanced age and high frailty score, low body weight, drug-drug interaction). Patients with known statin-attributed muscle symptoms or who develop statin-attributed muscle symptoms during the study will be treated following the therapeutic flow-chart for management of patients with statin-associated muscle symptoms of the EAS Consensus Panel.

After 4 weeks, lipids are measured. If LDL-C >1.8 mMol/L, ezetimibe 10 mg is added on top of statin therapy. Patients with documented statin intolerance to at least three different statins, as defined by the EAS Consensus Panel, will be treated with 10 mg ezetimibe mono-therapy. Four weeks later lipids are measured, and if LDL-C >1.8 mMol/L, alirocumab will be added on top of current treatment with a statin and ezetimibe, in accordance with the following dosing-schedule:

- if 1.8alirocumab 75 mg, or alirocumab 150 mg will be added
- if 2.6<=LDL-C<3.6 mMol/L, at the investigator*s discretion alirocumab 75 mg or alirocumab 150 mg will be added
- if LDL-C>=3.6 mMol/L, alirocumab 150 mg will be added Two weeks after the second dose of alirocumab, lipids are measured.

Intervention

Patients are treated according to cholesterol treatment guidelines. The choice of PCSK-9 inhibitor is directed by the protocol. When a PCSK-9 inhibitor needs to be prescribted, alicobumab is the drug of choice.

There are no other interventions in this study,

Study burden and risks

There are no additional risks or burden for study participants.

The burden for the patient is limited to the drawing of blood samples (at 4, 8 and 12 weeks and 12 months), all of which are standard of care. For this study no extra visits or physical examinations are required.

Contacts

Public

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Scientific

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Trial sites

Listed location countries

Netherlands

Eligibility criteria

Age

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

Inclusion criteria

- Admission because of type I (N)STEMI, and
- History of ASCVD (i.e., cerebrovascular disease (Transient ischemic attack, cerebral infarction, amaurosis fugax, retinal infarction), Coronary artery disease (unstable Angina pectoris, MI, ACS, coronary revascularization (coronary angioplasty or surgical procedure for coronary bypass)), Peripheral artery disease (Symptomatic and documented obstruction of an distal extremity artery or surgical operation (percutaneous transluminal angioplasty, bypass or amputation), and/or a history of T2DM.

Exclusion criteria

- Age <18 years
- Age >70 years and a Clinical Frailty Score >3.
- o To measure the frailty score, the validated Dutch translation of the Canadian Study of Health and Aging (CSHA) Clinical Frailty Scale will be used (table 2)
- Pregnancy and lactating women
- Known intolerance for alirocumab
- Active PCSK9-i therapy
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- Participation in lipid modifying drug trials
- Life expectancy <1 yr.

Study design

Design

Study phase: 4

Study type: Interventional

Masking: Open (masking not used)

Control: Uncontrolled

Primary purpose: Other

Recruitment

NL

Recruitment status: Recruitment stopped

Start date (anticipated): 15-01-2019

Enrollment: 1000

Type: Actual

Ethics review

Approved WMO

Date: 11-01-2019

Application type: First submission

Review commission: MEC-U: Medical Research Ethics Committees United

(Nieuwegein)

Approved WMO

Date: 14-02-2019
Application type: Amendment

Review commission: MEC-U: Medical Research Ethics Committees United

(Nieuwegein)

Approved WMO

Date: 19-06-2019

Application type: Amendment

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Review commission: MEC-U: Medical Research Ethics Committees United

(Nieuwegein)

Approved WMO

Date: 11-09-2019

Application type: Amendment

Review commission: MEC-U: Medical Research Ethics Committees United

(Nieuwegein)

Approved WMO

Date: 03-12-2019

Application type: Amendment

Review commission: MEC-U: Medical Research Ethics Committees United

(Nieuwegein)

Approved WMO

Date: 23-09-2020

Application type: Amendment

Review commission: MEC-U: Medical Research Ethics Committees United

(Nieuwegein)

Approved WMO

Date: 23-09-2021

Application type: Amendment

Review commission: MEC-U: Medical Research Ethics Committees United

(Nieuwegein)

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 CCMO

ID

NL66879.100.18