

PROTECT EU

A multinational registry on preventive left ventricular support with Impella in anatomically high-risk PCI treated patients





We are very pleased to announce you the FIRST PATIENT ENROLLED! (23.02.2023)



5

6

4 ACTIVE CENTERS

OSPEDALE SAN RAFFAELE0 PTS ENROLLEDPOLICLINICO SAN DONATO2 PTS ENROLLEDAOU PADOVA1 PTS ENROLLEDAO SAN CAMILLO FORLANINI0 PTS ENROLLED

SUBMISSION STATUS

CONTRACT NEGOTIATION

> AWAITING FOR EC OPINION



Prof. Giuseppe Tarantini

Principal Investigators



Dott.ssa Alaide Chieffo



RITERI

Prospective, non-interventional, multi-center international study

PRIMARY ENDPOINT:

Composite endpoint of all-cause death, cerebrovascular accident, myocardial infarction, repeat revascularization at 90 days.

OBJECTIVE:

To evaluate 90 days outcomes in high risk-PCI patients treated with a preventive strategy with Impella CP

System.

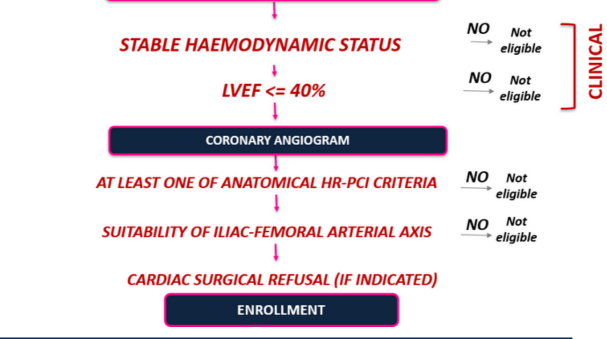
TIMELINE

First subject enrolled on 23.02.2023

FLOW CHART

Enrollment target: 859 patients Enrollment period: 1 year





Principal Investigators

Prof. Giuseppe Tarantini & Dott.ssa Alaide Chieffo



Committee members

Writing and steering committe



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CORELAB



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STUDY SPONSOR

Fondazione per la ricerca e l'innovazione in cardiologia

CRO

SCTB

Service for Clinical Trials and Biometrics UNIT OF BIOSTATISTICS, EPIDEMIOLOGY AND PUBLIC HEALTH Department of Cardiac, Thoracic, Vascular Sciences and Public Health **University of Padova**



Prof. Giuseppe Tarantini



Principal Investigators



GUIDELINES

We recommend you to follow the general guidelines described below:

- Upload the signed informed consent by the patient in the trial master files in redcap before starting data compilation in the eCRF
- Angiographic images must be uploaded in the "procedure file upload" section of the eCRF, before uploading, the images must be anonymized as indicated in the chart
- The follow-ups scheduled at 90 days and 1 year can also be performed by telephone, we recommend that you respect the execution windows reported for each patient in the corresponding follow-up chart

