

## Data Verification Plan HOT-ICU

## "Handling Oxygenation Targets in the Intensive Care Unit" EudraCT-no. 2017-00632-34

Level I: Systematic data verification of all data collected. Applies to the first 3 trial participants and hereafter on every 10<sup>th</sup> randomly chosen trial participant. Level II: All trial participants.

Data	eCRF Code/page	I	II	Comments
Informed consent		х	х	
Consent stated in medical record		Х	х	
Inclusion criteria	S1-S7	Х	х	
Exclusion criteria	S8-S19	х		
Randomisation	S20-S25	Х	Х	
Baseline				
General Patient Information	B1-B6	Х		B4: Check only for presence of data.
Respiratory support	B7-B7d	Х		Appears in the eCRF if S4 = "Yes".
Arterial Blood Gas	B8-B11	Х		Last value before randomisation.
Acute Illness	B12-B18	Х		B18: Check only for presence of data.
Sofa score	B19-B26	Х		B19, B20 and B25: Check only for presence of data.
Chronic co-morbidities	B27-B30a	х		B28 and B30a: Check only for presence of data.



Data	eCRF Code/page	ı	II	Comments		
Daily registration						
Respiratory support	D1	X <sup>1</sup>	х	D1: Invasive mechanical ventilation, NIV or CPAP (intermittent CPAP = less than one hour is not regarded as respiratory support).		
Use of prone position, inhaled vasodilators or ECMO	D1a-D1c	x <sup>1</sup>				
06.00h – 05.59h	D2-D5b	x <sup>1</sup>				
Respiratory status 08:00	D6-D7a4	x <sup>1</sup>				
Remaining organ systems	D8	x <sup>1</sup>				
Remaining organ systems	D9, D10 and D14	X <sup>1</sup>				
SAE/secondary outcome (Defined as acute myocardial ischaemia, ischemic stroke, intestinal ischaemia or a new incidence of shock)	D11-D13	X	Х	Hospital records are reviewed for the first 7 days of admission. In case of an SAE, check eCRF for correct registration.		
$X^1$ : Check the first 7 daily forms and hereafter every $5^{th}$ daily form						
Discharge from ICU	DR1-DR3	Х				
Readmission	DR4-DR5	Х		Within 90 days from randomisation		
Withdrawal	W1-W3b	х		NOTE! It should be stated in the patient's medical record whether consent is given to keep up data registration when written consent for trial participation is retracted or not given		
Follow up/primary outcome after 90 days		х	х			