

Alterations of immunologic mediators during severe sepsis

Submission date 20/04/2006	Recruitment status Stopped	<input type="checkbox"/> Prospectively registered
		<input type="checkbox"/> Protocol
Registration date 13/06/2006	Overall study status Stopped	<input type="checkbox"/> Statistical analysis plan
		<input type="checkbox"/> Results
Last Edited 01/02/2019	Condition category Infections and Infestations	<input type="checkbox"/> Individual participant data
		<input type="checkbox"/> Record updated in last year

Plain English summary of protocol
Not provided at time of registration

Contact information

Type(s)
Scientific

Contact name
Dr Armin Sablotzki

Contact details
Delitzscher Str. 141
Leipzig
Germany
04129
+49 (0)341 909 2570
armin.sablotzki@sanktgeorg.de

Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number
NCT00484146

Secondary identifying numbers
LAVISS_01

Study information

Scientific Title

Alterations of immunologic mediators during severe sepsis

Study objectives

Severe sepsis induces significant changes in expression of insulin and toll-like receptors, cytokines, markers of apoptosis, and activation of t- and b-lymphocytes.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Ethics approval received from the Ethics Committee of the Saxonian Chamber of Physicians on the 21st July 2006 (ref: EK-BR-15/06-1).

Study design

Prospective, open, clinical observational study

Primary study design

Observational

Secondary study design

Cohort study

Study setting(s)

Hospital

Study type(s)

Screening

Participant information sheet

Health condition(s) or problem(s) studied

Severe sepsis

Interventions

Daily blood samples for 7 days

Intervention Type

Other

Phase

Not Specified

Primary outcome measure

Alterations of immunologic parameters

Secondary outcome measures

Not provided at time of registration

Overall study start date

01/06/2006

Completion date

31/12/2007

Reason abandoned (if study stopped)

Participant recruitment issue

Eligibility

Key inclusion criteria

1. Age 18 years or older
2. Agreement with study procedures, informed consent
3. Fulfilling 3 out of 4 criteria of a systemic inflammatory response syndrome (SIRS)
4. Suspected or proven infection
5. Two or more sepsis-induced organ dysfunctions
6. Start of first sepsis-induced organ dysfunction within the last 36 hours

Participant type(s)

Patient

Age group

Adult

Lower age limit

18 Years

Sex

Both

Target number of participants

40

Key exclusion criteria

1. Non-agreement with study procedures
2. Sign of severe sepsis for more than 36 hours
3. Chronic immuno-compromizing diseases
4. Chronic therapy with anti-inflammatory drugs
5. Non-curable cancer diseases
6. Chronic renal failure with hemodialysis
7. Pregnant or breast feeding

Date of first enrolment

01/06/2006

Date of final enrolment

31/12/2007

Locations

Countries of recruitment

Germany

Study participating centre

Delitzscher Str. 141

Leipzig

Germany

04129

Sponsor information

Organisation

Urban Clinical Center of St Georg in Leipzig (Städt. Klinikum St. Georg, Leipzig) (Germany)

Sponsor details

Delitzscher Str. 141

Leipzig

Germany

04129

+49 (0)341 909 2570

kais@sanktgeorg.de

Sponsor type

Hospital/treatment centre

Website

<http://www.sanktgeorg.de>

ROR

<https://ror.org/02y8hn179>

Funder(s)

Funder type

Hospital/treatment centre

Funder Name

Urban Clinical Center of St Georg in Leipzig (Städt. Klinikum St. Georg Leipzig) (Germany)

Results and Publications

Publication and dissemination plan

Not provided at time of registration

Intention to publish date

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration