**INVESTIGATOR SITE HEADED PAPER**

**Sepsis Trials in Critical Care - SepTiC**

**Retrospective Information Summary and Consent form**

You are being invited to continue to participate in our research study. Sepsis is when the body overreacts to an infection and starts to damage itself. This study is trying to find the best treatments for sepsis. This form provides summary information on the study, the treatments we are using, what has happened and what will happen if you decide to continue as well as the risks and benefits of taking part.

Please take time to read the following information carefully and do talk about it with other people if you would like to. Ask us if there is anything that you do not understand or if you would like more information. You can take time to think about whether you wish to continue taking part in the study.

**What are the treatments?**

We are testing three different treatments in this study: -

1. Diagnostic Trial – this involves a PCR test that quickly tells us what bacteria may be causing the infection and may help us decide which antibiotics to give.
2. Fluid Trial – giving fluid is routine when treating sepsis, but there is no clear guide on how best to provide the treatment, so we will test how much fluid should be given and if medication should be given to remove any build-up of extra fluid.
3. GM-CSF Trial – this drug can help the immune system fight infection by making more white blood cells.

These treatment options listed above also include a ‘standard care’ or ‘placebo’ option. A computer randomly selects which treatment options you have received. This means you might not have received any of these treatments, even if you choose to continue to participate in the SepTiC trial. How patients recover over time is then compared between the different treatment options to work out which treatments are best.

**What do I need to do?**

Treating sepsis quickly is important and therefore you were included in the trial when you were very unwell and lacked the capacity to discuss the study. We have spoken to a relative / friend or an independent doctor about your taking part. You may already have received treatment and a few extra blood samples may have been collected.

You do not need to do anything for the study while you are in hospital. We will collect data from your hospital records and other NHS linked data. We may contact you 3 months later to see how you are and would like to contact you 6 months later with a short telephone call to ask about your quality of life, and wellbeing.

Participation into the SepTiC trial is voluntary. All patients, including those that do not wish to participate in SepTiC, will receive the best available standard of care available at this hospital. You can choose not to take part in the study any more.

More information about the study including how we use patient data and privacy, legalities and insurance of the study, risks and benefits, how to make a complaint and how to find out the results of the study can be found in our patient information sheet. (A written copy is available or is available online at [www.septictrial.co.uk](http://www.septictrial.co.uk))

If you are happy to proceed, please complete the attached consent form.

**Site Contact Information**

Investigator name:-

Site Contact details:-

**Study Contact Information**

Please contact The **SepTiC** Trial team using the following contact details:

Name: The **SepTiC** Trial Team

24hr Telephone: 0207 5949725

Email: septic@imperial.ac.uk

Website: [www.septictrial.co.uk](http://www.septictrial.co.uk/)

Thank you very much for continuing to take part in this study!

A copy of the written information and signed Informed Consent form will be given to you to keep.

**Consent Form for Participants – Regained capacity**

**Full Title of Project:** Sepsis Trials in Critical Care - **SepTiC**

|  |  |
| --- | --- |
| **Site number:**  |  |
| **Patient Number:** |  |
| **Patient Name:** |  |
| **Name of Principal Investigator:** |  |

**Please initial box**

|  |  |
| --- | --- |
| 1. I confirm that I have read and understand this document and have read/received a copy of the appropriate patient information sheet and privacy notice for **SepTiC.**
 |  |
| 1. I confirm I am happy to consent to continue to participate in the following trials:

Diagnostic and Fluid Trial GM-CSF Trial.  |  |
| 1. I confirm that I understand the **SepTiC** study and I have had the opportunity to ask questions which have been answered fully.
 |  |
| 1. I understand that my participation is voluntary, and I am free to withdraw at any time, without giving any reason and without my legal rights nor treatment / healthcare being affected.
 |  |
| 1. I understand that sections of any of my medical notes may be looked at by responsible individuals from Imperial College London, from the NHS Trust or from regulatory authorities where it is relevant to my taking part in this research.
 |  |
| 1. I give consent for information collected about me to be used to support other research or in the development of a new test, medication, or treatment by an academic institution or commercial company in the future, including those outside of the United Kingdom (which Imperial has ensured will keep this information secure).

  |  |
| 1. I give consent for samples (blood samples) collected from me to be used to support other research or in the development of a new test, medication, or treatment by an academic institution or commercial company in the future, including those outside of the United Kingdom (which Imperial has ensured will keep this information secure).

  |  |
| 1. I understand that blood samples and / or data collected from me are a gift donated to Imperial College and that I will not personally benefit financially if this research leads to an invention and/or the successful development of a new test, medication treatment, product or service.
 |  |
| 1. I give consent to being contacted about the possibility to take part in other research studies.
 |  |
| 1. I consent to allow the use of data already collected in the trial, as well as ongoing data collection and follow up information to be obtained from my medical records at 3, 6 months and 12 months after my inclusion
 |  |
| 1. I agree that my medical records and other personal data generated during the study may be examined by representatives of the sponsor (Imperial College London), by people working on behalf of the sponsor, and by representatives of Regulatory authorities, ICNARC, NHS Digital and SICSAG where it is relevant to my taking part in this research.
 |  |
| 1. I agree to my blood samples being used to undertake genetic research which may have the potential to generate data that can be tracked back to me
 |  |
| 1. I consent to continue to take part in **SepTiC**
 |  |

\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_ \_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_ \_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_

Name of patient Signature Date

\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_ \_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_ \_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_

Name of staff member Signature Date

*(Listed on delegation log)*

Original for Principal Investigator, 1 copy for participant; 1 copy for hospital notes

To ensure confidence in the process and minimise risk of loss, all consent forms must be printed, presented, and stored in double sided format