Deferred Consent for Research Participation

**Title of Project:** STandard versus Accelerated initiation of Renal Replacement Therapy in Acute Kidney Injury (STARRT-AKI): A Multi-Centre, Randomized, Controlled Trial

**Principal Investigator:** [Name]

**Phone Number:** [Number]

**Co-Investigators:**

The patient named below is being enrolled in this research study by deferred consent.

When a previously incapacitated participant regains capacity, or when a substitute decision maker is found, consent shall be sought promptly for continuation in the study, and for subsequent examinations or tests related to the research study. Seeking consent prior to study participation is always preferable.

If no substitute decision maker/family member is available on site or they refuse to give consent, any data or samples collected for study purposes will be destroyed.

**Patient’s Name:** ________________________________

**Date/time assessed for enrolment:** _____/_____/______ (dd/mm/yyyy) at ____ : ______ (time)

**Reason(s) deferred consent process is used (check all that apply):**

_____ The patient is unconscious or lacks capacity to understand the risks, methods and purposes of the research study.

_____ No next of kin/substitute decision maker is available to provide consent, or attempts to contact them have been unsuccessful despite diligent and documented efforts.

_____ A substitute decision maker _________________________________ (name and relationship) has been contacted by telephone, and the purpose, methods and risks of participation in this study have been explained to the third party. While the substitute decision maker has given verbal consent for participation, written consent must be still be obtained.

_____ No relevant prior directive by the patient is known to exist.

_____ Other: ________________________________

__________________(Signature of authorized independent physician) ____________________ (Date and Time)

STARRT-AKI Trial Pro00060023 – Deferred Consent February 6, 2019