

This form must be completed and returned to the HSRC REC Administrator, Mr V Skosana as soon as possible but within 7 days for serious adverse events, and within 15 days for other adverse and unexpected events. One form is to be completed per participant, even if several participants are involved in a similar adverse event.

STUDY INFORMATION

STUDY NAME:

DESCRIPTION OF THE INTERVENTION:

HSRC Research Ethics Committee Number:

1. CLINIC AND PARTICIPANT INFORMATION:

CLINIC NAME:

PARTICIPANT ID:

PARTICIPANT AGE:

PARTICIPANT GENDER:

2. ADVERSE EVENT:2.1 AE REPORT TYPE: Initial ☐ Follow-Up: ☐

2.2 DATE OF ADVERSE EVENT: / / (DD/MM/YY)

2.3 ADVERSE EVENT REPORTED TO RESEARCHERS BY:

☐ Study participant returning to the site☐ By other means, specify:**2.1 COMPONENT OF STUDY, PARTICIPANT INVOLVED IN:**1 ☐ Baseline2 ☐ Six weeks3. ☐ Six months4. ☐ Other, specify:**2.2 ADVERSE EVENT SEVERITY:**1 ☐ Mild2 ☐ Moderate3 ☐ Severe4 ☐ Fatal

2. 3 ADVERSE EVENT DESCRIPTION:

PROVIDE A BRIEF DESCRIPTION OF INJURY/ADVERSE EVENT INCLUDING ANY ACTION TAKEN BY THE STUDY TEAM TO DATE ON BEHALF OF THE PARTICIPANT.

2.4 IS THE ADVERSE EVENT SERIOUS?*
 1. ☐ Yes 2. ☐ No

***SERIOUS ADVERSE EVENTS ARE CONSIDERED FATAL OR LIFE THREATENING THAT REQUIRE HOSPITALIZATION OR PROLONG EXISTING HOSPITALIZATION, OR RESULT IN PERSISTENT OR SIGNIFICANT DISABILITY**

2.5 CLASSIFICATION OF ADVERSE EVENT

- ☐ Results in death
- ☐ Is life-threatening
- ☐ Requires inpatient hospitalization or prolongation of existing hospitalization
- ☐ Results in persistent or significant disability/incapacity
- ☐ Any other experience that suggests a significant hazard, contraindication, side-effect, or precaution that may require medical or surgical intervention to prevent one of the outcomes listed above
- ☐ Events changes the risk/benefit ratio of the study

2.6 AT THE TIME OF THIS REPORT, THE ADVERSE EVENT IS:

1. ☐ Resolved (No additional follow-up necessary)
2. ☐ Unresolved (Additional follow-up necessary)

3. RESEARCH STAFF ASSESSMENT OF ADVERSE EVENT**3.1 IN YOUR JUDGEMENT, IS THE ADVERSE EVENT RELATED, POSSIBLE RELATED, UNKNOWN, OR NOT RELATED TO THE PROTOCOL?**

- | | |
|----------------------------|------------------|
| 1 <input type="checkbox"/> | Related |
| 2 <input type="checkbox"/> | Possibly Related |
| 3 <input type="checkbox"/> | Unknown |
| 4 <input type="checkbox"/> | Not related |

4. VERIFICATIONSTAFF MEMBER:

COMPLETED BY (PLEASE PRINT OR TYPE):

FIRST NAME:

LAST NAME:

DESIGNATION/ROLE ON RESEARCH PROJECT:

STAFF MEMBER SIGNATURE:

DATE: / / (DD/MM/YY)

PRINCIPAL INVESTIGATOR (PLEASE PRINT OR TYPE):

I have reviewed this AE Form for this participant and attest that the information recorded is accurate and complete.

INVESTIGATOR'S FIRST NAME:

INVESTIGATOR'S LAST NAME:

INVESTIGATOR SIGNATURE:

DATE: / / (DD/MM/YY)