

**Memorial Health University Medical Center  
Institutional Review Board**

**SUBMISSION OF SERIOUS ADVERSE EVENT REPORT FOR SPONSORED STUDY**

**Study Title:**

**MHUMC Study Number (assigned by IRB):**

**Principal Investigator Information:**

Name: \_\_\_\_\_ Phone: \_\_\_\_\_  
Address: \_\_\_\_\_ Email: \_\_\_\_\_

**Study Coordinator:**

**DETAILS OF EVENT**

Local  Non-local/IND Safety Report  Not specific to this study IND or IDE #:

Initial  Follow-up  Final (If follow-up or final, provide initial SAE #: \_\_\_\_\_ )

Date of Event: \_\_\_\_\_ Date of Initial Awareness by Research Team: \_\_\_\_\_

**Type of Event:**

Hospitalized  Life-threatening  Other Serious  Persistent Disability  Congenital Anomaly  Death

**Event Relationship to Study Drug:**

Related  Probably Related  Possibly Related  Not Related  Probably Not Related  Undetermined

**Subject Information:** Age: \_\_\_\_\_ Gender:  Male  Female ID: \_\_\_\_\_

**Status in Study:**  Continued  Discontinued  Tx on hold/Interrupted  Died  Unknown

**Brief Description of Event:**

Is this type of event expected for this particular study?  Yes  No

**As a result of this SAE, does the Informed Consent need to be revised to include new risk?**

No  Yes (attach copy)

\_\_\_\_\_  
Principal Investigator Date

**\*\*\* Section below for IRB administrative use only \*\*\***

Accepted as information.

SAE #: \_\_\_\_\_

\_\_\_\_\_  
IRB Chair

\_\_\_\_\_  
Date

IRB Agenda: \_\_\_\_\_