



# Institutional Review Board/IRB-00002563 Continuing Review

IRB-00002563

Date: \_\_\_\_\_ IRB # \_\_\_\_\_

Principal Investigator: \_\_\_\_\_

Department: \_\_\_\_\_ Phone # \_\_\_\_\_

Research Coordinator: \_\_\_\_\_ Phone # \_\_\_\_\_

Project Title:

### INVESTIGATOR COMPLETE:

- 1. Is it an Expedited Review?..... Yes No
- 2. Date of Original Approval: \_\_\_\_\_
- 3. Date of Last Approval (if applicable): \_\_\_\_\_
- 4. Is project completed? (all research done, enrollment close & patients follow-up complete) ... Yes No  
If yes, complete questions 7-13. If no, complete questions 5-13.
- 5. Is the research permanently closed to the enrollment of new subjects?. ..... Yes No
- 6. Have all participants completed all research-related interventions?..... Yes No
- 7. Give a brief statement of the progress of the project (i.e., number of subjects entered, length of time since study initiated, length of time projected to go on, etc.)
  
- 8. Have any subjects withdrawn or been removed from the research due to AE or safety concern? ..... Yes No  
If yes, state how many have withdrawn and describe the circumstances:
  
- 9. Have there been any complaints/concerns about the research since the last IRB review? ..... Yes No  
If yes, please report the complaints/concerns and your response/action.
  
- 10. Any unanticipated problems involving risks not detailed in your original application? ..... Yes No
- 11. Any serious adverse events (SAE) at your site since last review? ..... Yes No
- 12. Any SAE, safety update, action letter, IND or other at another site since last review?..... Yes No
- 13. Any unexpected mortality or morbidity since last review? ..... Yes No

If yes to any of the above (7-13), please provide a detailed summary below. Need to write a more detailed summary about the adverse events, problems, etc. (i.e., the types of adverse events, numbers, trends).

**Example:** There were 6 adverse events during the past year, four of which occurred in patients at other sites and 2 occurred at this site. Three of the events were constipation thought to be related to the study drug; two were shortness of breath thought not to be related to the study drug; and one was a petechial rash thought to be possibly related.)

**Also, make certain adverse events/mortality reports have been submitted and are on file in the IRB office.**

14. Any protocol changes, consent form changes or amendments since last review?..... Yes No  
If yes, list below (make certain protocol changes/amendments have been submitted and are on file in the IRB office).

15. Number of subjects enrolled since last review: \_\_\_\_\_

16. Total number of subjects enrolled since project initiated: \_\_\_\_\_

17. Have all PIs involved with the research completed the IRB Education Requirements? ..... Yes No  
(Education requirements must be completed before the IRB can grant continued approval for the research project).

**A copy of the current informed consent document (with, if possible, the IRB Approval dated stamp) must be submitted with this report.**

\_\_\_\_\_  
Signature of Principal Investigator Date

**IRB USE ONLY:**

Continue approval based on Expedited Review  
Continue approval based on Full Board Review  
Suspend approval, pending investigation

\_\_\_\_\_  
Signature of IRB Chairperson/Designee Date

Next Review required before \_\_\_\_\_

Completed by: \_\_\_\_\_