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| **Progress Report for Expedited Studies****This study is exempt from continuing review because of one of the following conditions:**  **1. The study is NOT regulated by the FDA under 21 CFR 312 (drugs) or 21 CFR 812 (devices),**  **2. The study is eligible for expedited review, i.e. research activity is limited to the activities described in categories 1 to 7 of the OHRP Expedited Review Categories, or**  **3. Research has advanced to involve only one or both:** **a. Data analysis, including analysis of identifiable private information or identifiable biospecimen, and/or** **b. Accessing follow-up clinical data from procedures that subject would undergo as part of clinical care; not specifically for research.****Therefore, only a progress report must be filed.****\*Note: Eliminating the continuing review requirement does NOT apply to FDA-regulated research** |

**Return this form to:**

Baptist IRB Baptist.IRB@bmhcc.org (Baptist IRB Memphis)

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| Date of Submission:       | Baptist-IRB Number:       | Principal Investigator:       |
| Title of Study:       |

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| General Information |
| 1. Date initially approved:
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| 1. Study site:
 |
| 1. Person completing this form:
 |
| 1. **Report of Activity: If necessary, use separate sheets to explain any answers**
 |
|  | When the protocol was initially approved, how many subjects were authorized for enrollment?       NA       |
| 1. **Study Personnel & Study Site**
 |
|  | Have any changes occurred in the professional personnel participating in the study? If Yes, please explain.       | [ ]  Yes | [ ]  No |
|  | Have the licenses, certifications or professional privileges of any personnel participating in the study been restricted or modified? If Yes**,** please explain.       | [ ]  Yes | [ ]  No |

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| 1. **Study Status and Supporting Documentation**
*
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|  | Are subjects still being enrolled or records being reviewed?  | [ ]  Yes | [ ]  No |
|  | Is the study permanently closed to enrollment AND subjects ARE NOT receiving active study treatment BUT undergoing follow-up assessment only? [ ]  Yes [ ]  No   |
| (c) |

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| Data Analysis only. | [ ]  Yes | [ ]  No |

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| **Statement of Principal Investigator: I have reviewed this Progress Report. To the best of my knowledge, the information provided is accurate.** |
| **Principal Investigator (please print)** |  |
| **Signature of Principal Investigator** | **Date** |