Form C Request for Full Institutional Review Board Review

Cover Sheet

This form is used to request the Institutional Review Board to take action regarding human subjects research which does NOT meet the criteria for EXEMPT FROM REVIEW or EXPEDITED REVIEW.

Primary Researcher	
Signed:	Date:
from the IRB if the research is still in progress 12 mon	
	l principles required for the protection of human subjects. I agree to immediately anges in this research to the IRB Chair. I agree to request a Continuing Review
ethical responsibilities expected by my profession. I ha	we included below all information and documents as required by the University
Agreement to follow IRR procedures My signature be	elow indicates that I understand and agree to adhere to relevant principles and
Identify all sources of funding:	
Anticipated last date of contact with participants:	
Anticipated date of first contact with participants:	
Start date of research: If human subjects are involved:	Completion date of research:
Canad Juda of managed in	Completion data of managed
Title of Research:	
Project Director:	
I have read this request and verify that it is an	n accurate description of the proposed research:
Is this Project Supported by any funds extern	al to Delta State University?
Campus Phone Number:	
Mailing Address:	
Project Director:	

Send to: Institutional Review Board, Chair; 233 Kent Wyatt Hall, Delta State University, Cleveland, MS 38733

Fax: 662-846-4313

Phone: 662-846-4010

Revised September 23, 2025

Email: irb@deltastate.edu

Narrative

(For items that do not apply please mark N/A)

- 1. Brief statement of project goals/research questions
- 2. Research Protocol including:
 - Research procedures
 - Description of the subject population
 - Recruitment procedures
 - Length of research procedure and time commitment required of subjects
 - Location of the study
 - Methodology
 - Description of who will gather the data and how they are/will be trained
 - Sources of funding
 - Special circumstances
- 3. Benefits to the subject or to others
- 4. Risks -
 - Describe the possible risks, discomforts, and inconvenience to the subjects and the precautions that will be taken to minimize them (include physical, psychological and social risks).
 - Describe appropriate controls, screenings methods, follow-up procedures.
 - Describe what constitutes termination from the study before its completion.
 - Describe how confidentiality will be maintained including confidentiality of data collection and who will have access to the data.
- 5. Informed Consent. Describe the procedures that will be used in obtaining informed consent, keeping in mind that informed consent is a process, not just a form.

REMINDER:

WITH YOUR REQUEST TO IRB, YOU MUST SUBMIT ALL RELEVANT DOCUMENTS. FOR YOUR ASSISTANCE, A LIST OF MINIMUM REQUIREMENTS IS PROVIDED AS A CHECKLIST

Informed consent documents/forms. Ensure that informed consent addresses at the	minimum the following:
Purpose and description of the research	_
Amount of time required of the subject	
Voluntary participation	
Confidentiality of data	
Contact information of the researcher	
Information concerning the IRB	
Any additional and necessary documents such as:	
Survey/research instruments	
Copy of Consent and Assent forms	
Communication that will be provided to parents and/or external organization	ons such as schools, clinics, etc
List of external organizations that will be contacted	
Current certification (NIH)	
attached (OR) on file	