

INSTITUTIONAL REVIEW BOARD

1204 Marie Mount Hall College Park, MD 20742-5125 TEL 301.405.4212 FAX 301.314.1475 irb@umd.edu www.umresearch.umd.edu/IRB

DATE:	January 19, 2023
TO: FROM:	Sarah Silberman, BA, MS University of Maryland College Park (UMCP) IRB
PROJECT TITLE:	[1983197-1] Sexual Misconduct in Academia
SUBMISSION TYPE:	New Project
ACTION: APPROVAL DATE: EXPIRATION DATE: REVIEW TYPE:	APPROVED January 19, 2023 January 18, 2024 Expedited Review
REVIEW CATEGORY:	Expedited review category # 7. Waiver of Consent Documentation, 45CFR46.117(c).

Thank you for your submission of New Project materials for this project. The University of Maryland College Park (UMCP) IRB has APPROVED your submission. This approval is based on an appropriate risk/benefit ratio and a project design wherein the risks have been minimized. All research must be conducted in accordance with this approved submission.

Prior to final approval of this project scientific review was completed by the IRB Member reviewer

This submission has received Expedited Review based on the applicable federal regulations. This project has been determined to be a MINIMAL RISK project. Based on the risks, this project requires continuing review by this committee on an annual basis. Please use the appropriate forms for this procedure. Your documentation for continuing review must be received with sufficient time for review and continued approval before the expiration date of January 18, 2024.

Please remember that informed consent is a process beginning with a description of the project and insurance of participant understanding followed by a signed consent form. Informed consent must continue throughout the project via a dialogue between the researcher and research participant. Unless a consent waiver or alteration has been approved, Federal regulations require that each participant receives a copy of the consent document.

Please note that any revision to previously approved materials must be approved by this committee prior to initiation. Please use the appropriate Amendment forms for this procedure.

All UNANTICIPATED PROBLEMS involving risks to subjects or others (UPIRSOs) and SERIOUS and UNEXPECTED adverse events must be reported promptly to this office. Please use the appropriate reporting forms for this procedure. All FDA and sponsor reporting requirements should also be followed. All NON-COMPLIANCE issues or COMPLAINTS regarding this project must be reported promptly to this office.

Please note that all research records must be retained for a minimum of seven years after the completion of the project.

If you have any questions, please contact the IRB Office at 301-405-4212 or irb@umd.edu. Please include your project title and reference number in all correspondence with this committee.

This letter has been electronically signed in accordance with all applicable regulations, and a copy is retained within University of Maryland College Park (UMCP) IRB's records.



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DATE:	January 22, 2024
TO: FROM:	Sarah Silberman, BA, MS University of Maryland College Park (UMCP) IRB
PROJECT TITLE:	[1983197-4] Sexual Misconduct in Academia
SUBMISSION TYPE:	Continuing Review/Progress Report
ACTION: APPROVAL DATE: EXPIRATION DATE: REVIEW TYPE:	APPROVED January 22, 2024 January 18, 2025 Expedited Review
REVIEW CATEGORY:	Expedited review category #7. Waiver of consent documentation 45CFR46.117(c).

Thank you for your submission of Continuing Review/Progress Report materials for this project. The University of Maryland College Park (UMCP) IRB has APPROVED your submission. This approval is based on an appropriate risk/benefit ratio and a project design wherein the risks have been minimized. All research must be conducted in accordance with this approved submission.

Prior to final approval of this project scientific review was completed by the IRB Member reviewer

This submission has received Expedited Review based on the applicable federal regulations. This project has been determined to be a MINIMAL RISK project. Based on the risks, this project requires continuing review by this committee on an annual basis. Please use the appropriate forms for this procedure. Your documentation for continuing review must be received with sufficient time for review and continued approval before the expiration date of January 18, 2025.

Please remember that informed consent is a process beginning with a description of the project and insurance of participant understanding followed by a signed consent form. Informed consent must continue throughout the project via a dialogue between the researcher and research participant. Unless a consent waiver or alteration has been approved, Federal regulations require that each participant receives a copy of the consent document.

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