



Vice President for Research
Institutional Review Board for Health Sciences Research

Chair, Rita Basu, MD

December 13, 2022

To NCATS N3C Data Access Committee,

This letter is being provided as an update to an earlier letter given the availability of new types of data in the National COVID Cohort Collaborative (N3C) enclave. We understand that data in the N3C enclave includes a limited dataset of health record data from contributing sites (currently >75 sites) with site ID anonymized. It now also includes Patient Privacy Preserving Linked (PPRL) data such as CMS data and death data, and public datasets such as census data. As before, the Institutional Review Board for Health Sciences Research (IRB-HSR) at the University of Virginia (UVA) has determined that research studies using ONLY data available in the N3C enclave (including health record data, PPRL data, and public datasets) will be considered non-human subject research if the following criteria are met:

1. Study does not require submission to the FDA.
2. All data collected for the project, were entirely collected for purposes other than this project.
 - a. The investigator will not obtain information through intervention or interaction with the individual.
 - b. The investigator will not obtain data for this project via a process like a chart review where identifiable health information will be viewed, used or analyzed.
3. The research does not require access to any data that are readily identifiable and/or contain HIPAA identifiers exceeding the criteria of a Limited Dataset per HIPAA regulations. In addition, the research will not involve importing additional information into the dataset that would make the data within the N3C identifiable per HIPAA regulations. The research does not require access to any individuals as subjects of the research.
4. A contract is in place covering institutional access to N3C data that includes HIPAA Data Use Agreement (DUA) language. The researchers will each submit a Data Use Request (DUR) or DUR Collaborator form for new access to data within N3C. As part of the DUR the researcher will attest to having reviewed the institutional DUA as well as agreeing to the N3C Data User Code of Conduct.

In cases where the criteria above **are not met**, IRB review or oversight will be required since the project may then meet the federal definitions of human subject research.

Sincerely,

DocuSigned by:

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Eileen Sembrowich

Interim Director, UVA IRB for Health Sciences Research