





National Institutes of Health Bethesda, Maryland 20892

**DATE:** December 6, 2022

**TO:** Christeenna Iraheta

National Library of Medicine

**FROM:** NIH Senior Official for Privacy

**SUBJECT:** Applicability of the Privacy Act: "Information Program on Clinical Trials:

Maintaining a Registry and Results Databank (NLM)"

I have reviewed the National Library of Medicine (NLM) submission to OMB under the Paperwork Reduction Act which proposes to renew the existing clearance for the ClinicalTrials.gov website, which is the largest and most comprehensive clinical trial registry and results database in the world.

The information collection will enable compliance with statutory requirements of Section 801 of the Food and Drug Administration (FDA) Amendments Act of 2007 (FDAAA) and satisfy the purposes of the original clinical trial information collection that was established to comply with the FDA Modernization Act of 1997 (FDAMA). The information collection is necessary to allow researchers and organizations who are not subject to FDAAA or FDAMA to voluntarily register trials and other clinical studies. The information collection will provide patients, family members, clinicians, and researchers with timely access to up-to-date information about clinical trials, other types of clinical studies and their results. Alone or when combined with collected results information, the information collection can contribute to better-informed decisions about medical treatment, and reduce inadvertent and unnecessary duplication of clinical research studies.

Information is collected via electronic submission to the ClinicalTrials.gov Protocol Registration System, available at <a href="https://clinicaltrials.gov/ct2/manage-recs/register">https://clinicaltrials.gov/ct2/manage-recs/register</a>. The expanded clinical trials registry provides basic information about the studies, their implementation, and how to enroll. The results portion of the databank summarizes the outcomes of the trial and helps ensure that scientists have access to the latest scientific information about potential treatments for disease, as much of this information is not published in the scientific literature. The information can help scientists to better plan new research projects and avoid duplication that can expose human volunteers to unnecessary risks. It can also ensure that treatment decisions are based on a more complete set of scientific evidence.

Page 2 Applicability of the Privacy Act: "Information Program on Clinical Trials: Maintaining a Registry and Results Databank (NLM)"

I have determined that the Privacy Act will not apply to this information collection. Although personally identifiable information will be collected, none of the data elements will be retrieved to identify the individual. If you have any questions, please contact me at (301) 402-6201.

Celeste Dade-Vinson

NIH Senior Official for Privacy

Celeste Dade-Vinson