

Principal Investigator: *[Insert Last, First, Middle Name – if applicable: Study ID]*

## Dissemination Plan

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### INSTRUCTIONS FOR COMPLETING THIS TEMPLATE:

1. Insert study-specific text where included below in *green italics*.
2. Once this template has been completed:
  - a. Delete the orange instructional text
  - b. Change all text to black
3. Save the template as a PDF and upload it in Item 4.6. *Dissemination Plan of your PHS Human Subjects and Clinical Trials Information* [form](#)

In compliance with the *NIH Policy on Dissemination of NIH-Funded Clinical Trial Information* (<https://grants.nih.gov/policy-and-compliance/policy-topics/clinical-trials/reporting/nih-policy>), the project PI for the clinical trial “*[Insert study title]*” will:

- (1) Register the trial in ClinicalTrials.gov no later than 21 calendar days after enrollment of the first participant.
- (2) Submit the trial results in ClinicalTrials.gov no later than one year after the trial's primary completion date.

The project PI's home institution, *[Insert PI's institution]*, has an internal policy in place to ensure compliance with the *NIH Policy on Dissemination of NIH-Funded Clinical Trial Information* and its requirements for registration and submission of results information to ClinicalTrials.gov.

*[If the proposed project includes informed consent, include this sentence. If the project does not include informed consent, delete this sentence]:* The informed consent form(s) for this clinical trial will describe that the project and its results will be posted in ClinicalTrials.gov.