ATTACHMENT 2a: NCCDPHP DMP Template

OMB NO: 0920-1301

Exp. Date: 6/30/2023

NCCDPHP Data Management Plan (DMP)

for use by applicants and awardees of contracts and notices of funding opportunity (NOFOs)

Background: The DMP should be developed during the project planning phase prior to the initiation of collecting or generating **public health data** and regularly updated as plans evolve. The DMP will be evaluated by CDC for completeness and quality at the time of application submission, award, or submission of the evaluation plan; at least annually thereafter; and when the project approaches termination. **Public health data** means digitally recorded factual material commonly accepted in the scientific community as a basis for public health findings, conclusions, and implementation. It does <u>not</u> include grantee progress reports, administrative data, preliminary analyses, drafts of scientific papers, plans for future research, reports, communications with colleagues, or physical objects, such as laboratory notebooks or specimens. In most cases, acquisition of secondary data does not require a DMP. For projects in which CDC aggregates, analyzes, and disseminates awardees' data, CDC may choose to develop the DMP. If the applicant or awardee believes that their project does not meet the criteria for submission of a DMP, the applicant/awardee must provide a justification.

ata Management Plan Requirement: Required Not Required Reason not required:	-
hase of project (check one): \square New Application \square New Award \square Evaluation Plan \square Continuation	า 🗆 Fina
ate:	
OFO or Contract name:	
OFO or Contract (solicitation or award) Number:	
wardee Name or Number:	

Description of the Data

In the following table, identify the data to be generated or collected for the public health dataset. If the project involves more than one, repeat this table for each public health dataset. All cells should be filled with brief answers. Expand cells as needed. Where necessary, state "n/a" or "plans pending"; however, the final DMP must have substantive responses for all cells

Dataset Title	Study Type/Design	Frequency of Data Collection	Data Collection Timeframe	Where Data Will Be Maintained During Study Period	Responsible Person / Contact Information
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Describe the content (topics, variables) of the data		
Description of Standards for Collecting Data		
Describe the data collection/generation methods		
Describe measures to ensure data quality		
Providing Access to Data		
Describe what level of access (free public access, restricted access, no		
access) of data will be provided and when it will be made available (must be		
within 30 months of a one-off collection; 12 months for close of a cycle for		
an ongoing collection)		
Describe when, where, and how the data will be available (e.g., URL for		
downloading public access dataset, discoverability and procedures for		
gaining access to restricted dataset)		
If free public access to the data will not be provided, give a justification		
For data that will be released, describe procedures for data security,		
privacy / confidentiality (removal of PII, data use agreements, website		
security, etc.)		
Description of Standards Accompanying Release of Data		
Describe the established standards to be used to ensure usability and		
interoperability of data (e.g., ICD codes, CSV files, etc.)		
Describe the documentation that will be available regarding data source		
(e.g., population studied, response rate, etc.)		
Describe the documentation that will be available for analysis (e.g., data		
dictionary, sample code)		
Archival and Long-Term Data Preservation		
Describe the planned long-term preservation (how long the data will be		
stored, when the data can be accessed, who has access) or the justification		
for no long-term preservation		
If applicable, name the planned final location of the data (publicly accessible		
repository, institutional or governmental repository, etc.) and describe how		
it can be accessed (including link or contact information for archived data)		