

Start-up Resource – NINDS Spinal Muscular Atrophy CDE Recommendations

The National Institute of Neurological Disorders and Stroke (NINDS) and other Federal agencies and international organizations have the common mission of developing data standards for clinical research. Through the efforts of subject-specific working groups, topic-driven data elements have been created. The first set of Common Data Elements (CDEs) for Spinal Muscular Atrophy was developed in 2011. The Core data elements to be used by an investigator when beginning a research study in this disease/disorder are listed in this resource document. All other recommendations are listed on the website and should be considered based on study type.

Each CDE or instrument could be classified according to the definitions below:

General Core: A data element that is required for all NINDS funded studies.

Disease Core: A data element that collects essential information applicable to any disease-specific study, including all therapeutic areas. The NINDS and its appointed working groups assign the disease "Core" classification based on the current clinical research best practices. In each case, the disease Core CDEs are a small subset of the available CDEs, where it is anticipated that investigators will need to collect the disease Core CDEs on any type of study. These are required for all disease-specific studies.

Disease Supplemental - Highly Recommended: A data element which is essential based on certain conditions or study types in clinical research studies. In most cases, these have been used and validated in the disease area. These data elements are strongly recommended for the specified disease condition, study type or design.

Disease Supplemental: A data element which is commonly collected in clinical research studies. Use depends upon the study design, protocol or type of research involved. These are recommended, but not required, for studies.

Disease Exploratory: A data element that requires further validation but may fill current gaps in the CDEs and/or substitute for an existing CDE once validation is complete. Such data elements show great promise but require further validation before they are ready for prime-time use in clinical research studies. They are reasonable to use with the understanding that it has limited validation in the target group.



Summary of Core Recommendations: Spinal Muscular Atrophy CDEs

National Institute of Health (NIH) Resources:

The NINDS also strongly encourages researchers to use these NIH developed materials for NINDSsponsored research, when appropriate. Utilization of these resources will enable greater consistency for NINDS-sponsored research studies. These tools are free of charge.

Core CDEs for all NINDS Studies:¹

- NIH Toolbox
- Quality of Life in Neurological Disorders (Neuro-QOL)
- Patient-Reported Outcomes Measurement Information System (PROMIS)

Domain/Sub-Domain	CDE Name	CDE ID	Study Type
Participant Characteristics; Demographics	Birth date	C00007	All studies
Participant Characteristics; Demographics	Ethnicity USA category	C00020	All studies
Participant Characteristics; Demographics	Race USA category	C00030	All studies
Participant Characteristics; Demographics	Birth sex assigned type	C58676	All studies
Participant Characteristics; Demographics	Gender identity type	C58677	All studies
Participant History and Family History; General Health History	Medical history condition text	C00322	All studies
Participant History and Family History; General Health History	Medical history condition SNOMED CT code	C00313	All studies

¹ Note: Education year count C00015 is no longer a general Core CDE



Domain; Sub-domain	CDE Name	CDE ID
Participant/Subject Characteristics; Social Status	Education level USA type	C00012
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Trial number	C10171
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function test date and time	C11098
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function test position type	C11100
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function test mouth apparatus type	C11101
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function test not done reason	
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary Function Test seat position type	
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	tion Testing/Respiratory Pulmonary function test equipment manufacturer name	
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function test equipment model name	C12304



Domain; Sub-domain	CDE Name	CDE ID
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function test equipment software program name	C12305
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Ulna length measurement	C12306
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Ulna length measure tool name	C12307
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function complete exhalation indicator	C12310
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function index of lung function type	C12311
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function index of lung function best trial measurement	C12312
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function maximal pressure trial difference indicator	C12313
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function forced vital capacity cough exhalation indicator	C12314
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function forced vital capacity leak indicator	
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function forced vital capacity large value range indicator	C12316



Domain; Sub-domain	CDE Name	CDE ID
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function forced vital capacity early termination indicator	C12317
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function forced vital capacity peak flow indicator	C12318
Outcomes and Endpoints; Pulmonary Function Testing/Respiratory Status	Pulmonary function lung function measurement	C12516
Assessments and Examinations; Physical/Neurological Examination	Hand preference type	C00023
Assessments and Examinations; Vital Signs and Other Body Measures	Vital signs date and time	C01519
Assessments and Examinations; Vital Signs and Other Body Measures	Heart rate	C01521
Assessments and Examinations; Vital Signs and Other Body Measures	Weight measurement	C01541
Assessments and Examinations; Vital Signs and Other Body Measures	Body mass index value	C11131
Assessments and Examinations Laboratory Tests and Biospecimens/Biomarkers	Lab specimen collection date and time	C01701
Assessments and Examinations Laboratory Tests and Biospecimens/Biomarkers	Pregnancy test date and time	C01702
Assessments and Examinations Laboratory Tests and Biospecimens/Biomarkers	Pregnancy test specimen type	C01704



Domain; Sub-domain CDE Name		CDE ID	
Assessments and Examinations Laboratory Tests and	Lab test name	C01705	
Biospecimens/Biomarkers			
Assessments and Examinations Laboratory Tests and	Lab test result value	C01706	
Biospecimens/Biomarkers			
Assessments and Examinations Laboratory Tests and	Lab test abnormality significance type	C01707	
Biospecimens/Biomarkers			
Assessments and Examinations Laboratory Tests and	Lab test result status	C01709	
Biospecimens/Biomarkers			
Assessments and Examinations Laboratory Tests and	Pregnancy test qualitative result value	C01710	
Biospecimens/Biomarkers			
Assessments and Examinations Laboratory Tests and	Lab test result unit of measure	C01711	
Biospecimens/Biomarkers			
Assessments and Examinations Laboratory Tests and	Pregnancy test not applicable reason	C01714	
Biospecimens/Biomarkers			
Assessments and Examinations Laboratory Tests and	Pregnancy test performed indicator	C10896	
Biospecimens/Biomarkers			
Assessments and Examinations Laboratory Tests and	Mutational analysis results available indicator	C12783	
Biospecimens/Biomarkers			
Assessments and Examinations Laboratory Tests and	Mutational analysis performed family member indicator	C12784	
Biospecimens/Biomarkers			



Domain; Sub-domain	CDE Name	
Assessments and Examinations Laboratory Tests and	Gene mutation detected result type	C12785
Biospecimens/Biomarkers		
Assessments and Examinations Laboratory Tests and	Mutational analysis performed indicator	C12944
Biospecimens/Biomarkers		
Assessments and Examinations Laboratory Tests and	Specimen source type	C12226
Biospecimens/Biomarkers		
Assessments and Examinations Laboratory Tests and	Tissue specimen size measurement	C12229
Biospecimens/Biomarkers		
Assessments and Examinations Laboratory Tests and	Tissue specimen collection date and time	C12230
Biospecimens/Biomarkers		
Assessments and Examinations Laboratory Tests and	Preservation technique used type	C12233
Biospecimens/Biomarkers		
Assessments and Examinations Laboratory Tests and	Histochemical stains used type	C12235
Biospecimens/Biomarkers		
Assessments and Examinations Laboratory Tests and	Biopsy amyloid present on Congo red stain indicator	C12236
Biospecimens/Biomarkers		
Assessments and Examinations Laboratory Tests and	Biopsy and autopsy histochemical stains diagnostic	C12249
Biospecimens/Biomarkers	abnormalities present type	
Assessments and Examinations Laboratory Tests and	Biopsy and autopsy fiber abnormality type	C12525
Biospecimens/Biomarkers		



Domain; Sub-domain	CDE Name	
Assessments and Examinations Laboratory Tests and	Biopsy and autopsy type 1 predominance fiber percentage	C1252
Biospecimens/Biomarkers	value	
Assessments and Examinations Laboratory Tests and	Biopsy and autopsy type 2 predominance fiber percentage	C1252
Biospecimens/Biomarkers	value	
Assessments and Examinations Laboratory Tests and	Biospecimen fragment collect count	C1227
Biospecimens/Biomarkers		
Assessments and Examinations Laboratory Tests and	Specimen histochemical stain diagnostic abnormality type	C1227
Biospecimens/Biomarkers		
Assessments and Examinations Laboratory Tests and	Tissue specimen collection anatomic site	C1228
Biospecimens/Biomarkers		
Assessments and Examinations Laboratory Tests and	Tissue specimen section thickness measurement	C1228
Biospecimens/Biomarkers		
Assessments and Examinations Laboratory Tests and	Tissue specimen section count	C1228
Biospecimens/Biomarkers		
Assessments and Examinations; Non-Imaging Diagnostics	ECG atrial arrhythmia type	
Assessments and Examinations; Non-Imaging Diagnostics	ECG assessment date and time	
Assessments and Examinations; Non-Imaging Diagnostics	ECG global result type	
Assessments and Examinations; Non-Imaging Diagnostics	ECG heart rate	C0450



Domain; Sub-domain	CDE Name	CDE ID	
Assessments and Examinations; Non-Imaging Diagnostics	ECG heart rhythm result type	C04504	
Assessments and Examinations; Non-Imaging Diagnostics	ECG left ventricular hypertrophy indicator	C04505	
Assessments and Examinations; Non-Imaging Diagnostics	ECG right ventricular hypertrophy indicator	C04506	
Assessments and Examinations; Non-Imaging Diagnostics	ECG previous myocardial infarction indicator	C04508	
Assessments and Examinations; Non-Imaging Diagnostics	ECG QT interval	C04512	
Assessments and Examinations; Non-Imaging Diagnostics	ECG QTc interval	C04512	
Assessments and Examinations; Non-Imaging Diagnostics	ECG ST segment abnormality indicator	C0451	
Assessments and Examinations; Non-Imaging Diagnostics	ECG T wave abnormality indicator	C04514	
Assessments and Examinations; Non-Imaging Diagnostics	ECG ventricular arrhythmia type	C0451	
Disease/Injury Related Events; History of Disease/Injury Event	Clinical event or milestone type	C1061	
Disease/Injury Related Events; History of Disease/Injury Event	Clinical event or milestone achieved indicator	C1061	
Disease/Injury Related Events; History of Disease/Injury Event Clinical event or milestone achieved age value		C1265	
Disease/Injury Related Events; History of Disease/Injury Event Milestone lost age value		C12659	
Disease/Injury Related Events; History of Disease/Injury Event	Milestone currently able indicator	C12660	



General Core for all Studies:

Investigators should review the FDA's <u>"Guidance for Industry: Suicidal Ideation and Behavior: Prospective Assessment of Occurrence in</u> <u>Clinical Trials</u> for the most up-to-date information about suicidal ideation and behavior. One scale that FDA suggests is the Columbia Suicide Severity Rating Scale (C-SSRS) (available at <u>Columbia Suicide Severity Rating Scale Website</u>).

Core Instrument for Spinal Muscular Atrophy Studies:

Manual Muscle Testing

For the complete list of NINDS CDE recommendations for SMA, please see the <u>NINDS CDE website</u>.