NeuroNEXT Network

Standard Operating Procedure (SOP)

Site Selection and Qualification Version 4.0 SOP NN SS 401

Originators: NeuroNEXT CCC and DCC Personnel

Reviewed and Approved by:

Signature and Date:

Christopher S. Coffey Coffey Reason: I approve this document Date: Mar 8, 2024 08:20 CST

08-Mar-2024

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Merit Cudkowicz

Electronically signed by: Merit Cudkowicz Reason: I approve this document Date: Feb 22, 2024 17:01 CST

22-Feb-2024

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22-Feb-2024

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NN SS 401 Page 1 of 7

NEURONEXT NETWORK STANDARD OPERATING PROCEDURE FOR SITE SELECTION AND QUALIFICATION

SOP: NN SS 401 Version No.: 4.0 Issue Date: 01Mar2024

Effective Date: 15Apr2024

SITE SELECTION AND QUALIFICATION

Supersedes: Document Version 3.0

Effective Date: 08Apr2023

Signature and Date:

Dixie Ecklund

Electronically signed by: Dixie Ecklund Reason: I approve this document Date: Feb 24, 2024 17:20 CST

24-Feb-2024

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Electronically signed by: Stacey Grabert Reason: I approve this document Date: Feb 22, 2024 13:46 EST

22-Feb-2024

Name and Title: Stacey Grabert, Pharm.D, MS, (CCC Director of Quality Assurance)

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Joan Ohayon

Electronically signed by: Joan Ohayon Reason: I approve this document Date: Mar 11, 2024 11:20 EDT

11-Mar-2024

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NN SS 401 Page 2 of 7

NEURONEXT NETWORK STANDARD OPERATING PROCEDURE FOR SITE SELECTION AND QUALIFICATION

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1. POLICY

The NeuroNEXT Executive Committee (NEC) will select Clinical Study Sites (CSS) for participation in each Network study. The following criteria may be considered:

- Scientific expertise in the disease indication
- Projections of patient availability and diversity
- Geographic distribution
- History of productivity (when available)
- Site capacity

For the purposes of site selection, the funded NeuroNEXT Clinical Study Sites (CSS) are each considered as a single unit. NeuroNEXT CSS that do not have a fully executed Master Clinical Trial Agreement (MCTA) and Single Institutional Review Board (SIRB) Reliance Agreement (RA) on file at the Clinical Coordinating Center (CCC) will not be considered for participation in a NeuroNEXT study until such agreements are executed.

The Protocol Principal Investigator (PPI) may request the addition of non-NeuroNEXT CSS for participation to ensure adequate recruitment. If the NEC determines that there is a need for additional non-Network CSS, the NEC may make this recommendation to the NINDS. The addition of non-NeuroNEXT CSS is dependent upon NINDS approval. In the event that the PPI requests inclusion of non-US sites, this would require NINDS approval prior to NEC review of the request.

Any Non-NeuroNEXT CSS under consideration will be required to:

- confirm their interest in participating in the study;
- confirm the willingness of their Institution to sign a MCTA and SIRB RA with the CCC; and
- cede review for the study to the NeuroNEXT SIRB.

Each CSS selected to participate in a study will be qualified by the NeuroNEXT Data Coordinating Center (DCC) study monitor and/or Clinical Coordinating Center staff (CCC), unless prior qualification of the CSS is deemed to be sufficient. A qualification telephone screening process and/or pre-study site visit will be used to review the appropriateness of the investigator, his/her staff, facility, and resources, and to gauge the understanding of NeuroNEXT policies and applicable regulatory requirements by the investigator and his/her key research staff.

2. SCOPE

This SOP has been developed to be in alignment with federal regulations and Good Clinical Practices (GCP) as set forth in the 2016 Integrated Addendum to ICH E6(R1): Guideline for Good Clinical Practice E6(R2). The policies and procedures described in this SOP apply to the NeuroNEXT CCC and DCC within the context of their oversight and advisory roles for the NeuroNEXT Network, and to all NeuroNEXT investigators, staff, subcontractors, or other entities associated with the NeuroNEXT Network who manage, oversee, and conduct research regulated by FDA and/or applicable review committees.

3. ROLES AND RESPONSIBILITIES

Upon receipt of funding, the NEC is responsible for selecting CSS and communicating its final decision to all interested CSS.

NN SS 401 Page 3 of 7

NEURONEXT NETWORK STANDARD OPERATING PROCEDURE FOR SITE SELECTION AND QUALIFICATION

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The NEC is responsible for determining if there is a need for participation of additional non-NeuroNEXT CSS in a Network study, and if so, recommending those CSS to NINDS for approval according to the criteria described in the Policy section of this SOP.

NINDS is responsible for reviewing and approving/disapproving the addition of any non-NeuroNEXT CSS.

Prior to their consideration as a CSS, each CSS is responsible for: completing a study-specific questionnaire, confirming their interest in participating in the study, and confirming the willingness of their Institution to cede review for the study to the NeuroNEXT SIRB.

After site selection by the NEC, the DCC/CCC is responsible for conducting assessments of each CSS to determine if the investigator is appropriate and if the site is adequately prepared to conduct the study.

4. APPLICABLE REGULATIONS AND GUIDELINES

21 CFR 312.50	General Responsibilities of Sponsors
21 CFR 312.53	Selecting Investigators and Monitors
21 CFR 312.70	Disqualification of a Clinical Investigator
ICH E6, 2.0	The Principles of ICH GCP
ICH E6, 5.1	Quality Assurance and Quality Control
ICH E6, 5.6	Investigator Selection
ICH E6, 5.7	Allocation of Responsibilities
FDA	Information Sheet Guidance for IRBs, Clinical Investigators, and Sponsors: FDA Inspections of Clinical Investigators (June 2010)
FDA	Information Sheet Guidance for IRBs, Clinical Investigators, and Sponsors: Clinical Investigator Administrative Actions – Disqualification (May 2010)

5. REFERENCES TO OTHER APPLICABLE SOPS

NN GA 104	Conflict of Interest and Financial Disclosure Requirements for Clinical Study Sites
NN GA 106	Publication Policy Development
NN GA 107	Data Sharing
NN GA 109	Sharing Data with Industry Collaborators
NN SS 402	Site Initiation Visits and Site Training
NN SM 601	Single Institutional Review Board (SIRB) Reliance Process
NN SM 602	Single Institutional Review Board Reporting

6. ATTACHMENTS AND REFERENCES

NN SS 401 – A Document History

NN SS 401 Page 4 of 7

NEURONEXT NETWORK STANDARD OPERATING PROCEDURE FOR SITE SELECTION AND QUALIFICATION

SOP: NN SS 401 Version No.: 4.0 Issue Date: 01Mar2024 Effective Date: 15Apr2024

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7. TERMS AND ABBREVIATIONS

The following terms and abbreviations are used in this document:

CCC Clinical Coordinating Center at Massachusetts General Hospital

CSS Clinical Study Site(s)

DCC Data Coordinating Center at The University of Iowa

FDA U.S. Food and Drug Administration

ICH International Council for Harmonisation

MCTA Master Clinical Trial Agreement
NEC NeuroNEXT Executive Committee

PPI Protocol Principal Investigator

RA Reliance Agreement

SIRB Single Institutional Review Board

8. SPECIFIC PROCEDURES

A. Site Selection

#	Who	Task	Attachment / References	Related SOP
1.	CSS	Prior to site selection, the CSS must confirm its interest in participation and willingness to cede review of the study to the SIRB.		NN SM 601 NN SM 602
2.	CSS	Must have executed MCTA and RA on file at CCC prior to consideration for site selection. Non-NN sites must provide in writing that they would be willing to enter into study specific CTA and RA prior to being considered for site selection. The CTA and RA for non-NN sites must be fully executed prior to site activation.		NN GA 104 NN GA 106 NN GA 107 NN GA 109 NN SM 601 NN SM 602
3.	NEC	Select sites for each Network study based on criteria stated in the Policy section of this SOP, and communicate final site selection to all interested CSS.		

NN SS 401 Page 5 of 7

NEURONEXT NETWORK STANDARD OPERATING PROCEDURE FOR SITE SELECTION AND QUALIFICATION

SOP: NN SS 401 Version No.: 4.0 Issue Date: 01Mar2024

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B. Site Qualification

#	Who	Task	Attachment / References	Related SOP
1	NINDS	General site qualification is part of the NN grant review process. Sites awarded NN site grants have provided documentation of their training and qualification to participate is clinical trials across a spectrum of neurological disorders and diseases.		
2	PPI/NEC/DCC/CCC	During the site selection process sites provide information via study specific study site survey regarding training, qualifications and feasibility of performing as a participating site.		
3	DCC/CCC	A Site Initiation Webinar/Visit (SIW/SIV) is conducted for each CSS selected for each trial during which regulatory, SIRB, NeuroNEXT and protocol requirements are reviewed with CSS staff.		NN SS 402

NN SS 401 Page 6 of 7

NEURONEXT NETWORK STANDARD OPERATING PROCEDURE FOR SITE SELECTION AND QUALIFICATION

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4.0

Minor edits for clarity

SITE SELECTION AND QUALIFICATION

Supersedes : Document Version 3.0

Effective Date :08Apr2023

Attachment NN SS 401 - A. Document History

NeuroNEXT Network Standard Operating Procedure (SOP) Site Selection and Qualification **SOP NN SS 401** Reason or Justification Effective Version **Description of Modification** Issue Date Reviewer(s) for Modification Date N/A 06Apr2012 06May2012 1.0 New N/A Clarified that the CCC participates in assessment and qualification of 2.0 Updates for version 2.0 21Sep2016 21Oct2016 CSS. Minor edits and formatting N/A corrections. Updated "1996 ICH E6 Consolidated Guidance" to "2016 Integrated Addendum to ICH E6(R1): Guideline for Good Clinical Practice E6(R2)". 3.0 Changed CIRB to SIRB. Updated Updates for version 3.0 22Feb2023 08Apr2023 Catherine signature block to accommodate Gladden for electronic signatures. Additional minor updates throughout.

Periodic review

01Mar2024

15Apr2024

Preeti Paul

NN SS 401 Page 7 of 7

NN SS 401 Site Selection and Qualification v4.0 clean

Final Audit Report 2024-03-11

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- Stacey Grabert (SGrabert@mgh.harvard.edu) authenticated with Adobe Acrobat Sign.

Challenge: The user opened the agreement.

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Tania Leeder (tleeder@mgb.org) added alternate signer ecklundd@uiowa.edu. The original signer dixie-ecklund@uiowa.edu can still sign.

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2024-03-08 - 9:19:54 AM GMT- IP address: 172.226.137.0

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Signature Date: 2024-03-08 - 2:20:36 PM GMT - Time Source: server- IP address: 128.255.113.139

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ohayonj@ninds.nih.gov authenticated with Adobe Acrobat Sign.

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Signer ohayonj@ninds.nih.gov entered name at signing as Joan Ohayon

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