

Guidelines

National Institute of Neurological Disorders and Stroke

Clinical Study Quality Control / Quality Assurance Checklist

**PROCESS CHECKLIST FOR NINDS CLINICAL STUDIES
(AND PREPARING FOR A SITE VISIT)**

This checklist outlines a review of study organization and processes, with a focus on data management.

Note: NINDS has established these guidelines as a resource for items that NINDS or its contractor may review during a site visit. Definitions of underlined terms are available in the NINDS [Glossary](#).

| | | YES | NO | N/A |
|---|---|--------------------------|--------------------------|--------------------------|
| Overview - Study Administration and Procedures | | | | |
| 1. | Are all study documents, including protocol , manual of procedures (MOP) , data collection forms, statistical analysis plan (SAP) , etc. consistent with data management procedures? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. | Are the MOP , protocol, data collection forms, informed consent , etc., easily accessible, in a centrally located binder (electronic or paper), to assist study investigators? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. | Are there accessible participant files that contain source documentation of clinical observations such as lab results, medical record, progress notes, etc.? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. | Is there a study regulatory binder* that contains key study documents such as Institutional Review Board (IRB) approval , protocol versions, informed consent form , C.V.s, forms, financial disclosures , site monitoring reports ? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. | Does the training plan describe how and when procedures for quality assurance (QA) are implemented? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. | Does the training plan include procedures on how to train new staff? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 7. | Does the Drug / Device Distribution Plan specify procedures for the storage of, preparation of, dispensing of and handling unused intervention as well as procedures for completing treatment accountability logs? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 8. | Are there written plans for obtaining, handling, storing, and sending participant samples/materials? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 9. | Are there written procedures for obtaining and transmitting laboratory data? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 10. | Are there procedures in place for following participants from screening and enrollment through completion of the study? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 11. | Is there documentation of pre-screening and screening procedures so that data on eligible and ineligible individuals are captured in an appropriate format? Is a screening log provided? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

| | | YES | NO | N/A |
|-----|---|--------------------------|--------------------------|-------------------------------------|
| 12. | Does the informed consent include statements about the use of the data and specimen sharing for future research? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 13. | Is there a written procedure to insure that the current copy of the IRB approved informed consent form is signed before each participant is enrolled? | | | |
| 14. | Has the manual of procedures (MOP) , which includes the protocol, CRFs, informed consent, study staff roster, screening log, and standard operating procedures (SOPs) , been distributed to all clinical sites and updated as needed? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 15. | <i>Have the following study operation procedures or plans been created for the MOP:</i> | | | |
| | a. Organizational Plan | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | b. Safety Monitoring Plan | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | c. Training Plan | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | d. Study Communications Plan | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | e. Maintaining MOP | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | f. Site Signature Log/Delegation of Authority (Description of Responsibility) | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | g. Recruitment Plan | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | h. Screening and Informed Consent | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | i. Enrollment and Randomization | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | j. Retention Plan | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | k. Study Timelines/Study Visits | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | l. Drug/Device Plan | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | m. Laboratory Specimen Plan | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | n. Blinding/Unblinding | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | o. Concomitant Medications | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | p. Data Management | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | q. Source Documentation | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | r. Case Report Form completion | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | s. Adverse Events (AEs)/Serious Adverse Events (SAEs) | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | t. Participant Withdrawals from study and Lost-to-Follow-ups | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | u. Protocol Deviations and violations | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | v. Quality Assurance (QA)/Quality Control (QC) procedures | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | w. Monitoring Plan | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| | x. Study Completion | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |

| | | YES | NO | N/A |
|--------------------------------------|---|--------------------------|--------------------------|--------------------------|
| | y. Website (if applicable) | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| Randomization | | | | |
| 16. | Are there written procedures to assure that participants are randomized according to the randomization plan ? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 17. | Are there written procedures for maintaining the confidentiality of the randomization code ? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 18. | Is there a procedure that verifies the correct randomization number was assigned? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 19. | Are there written procedures to ensure that the randomization assignment stays with the participant through the entire data collection process? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 20. | Are masking/blinding and unmasking/unblinding procedures in place? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| Data Collection (Data system) | | | | |
| 21. | Is there a schedule of participant contacts (i.e. study visits)? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 22. | Are there written procedures that guide data collection at each participant contact? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 23. | Is there a complete description and definition of how each data item is to be collected on each study form for each participant contact? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 24. | Do the forms and data collected at each participant contact correspond to and reflect the statistical analysis plan ? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 25. | Are there adverse event (AE) forms and do they include the necessary data to generate safety reports? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 26. | Are there automated range and logic checks built into the system? | | | |
| Data Management | | | | |
| 27. | Is there a detailed description of how forms are sent or transmitted to the data-coordinating center? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 28. | Is there a Data Management Plan or do written procedures document data handling from collection through analysis? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 29. | Are there tracking procedures that document and confirm participant enrollment, data collected, forms completed, and forms received at the data collection/coordinating center? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 30. | Are there written procedures that describe how data are transformed from paper into a computer system, edited, and transferred to an analysis data base, as relevant? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

| | | YES | NO | N/A |
|----------------------------------|---|--------------------------|--------------------------|-------------------------------------|
| 31. | Are there procedures for correcting data so that changes can be identified for accuracy and completeness in a systematic way? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 32. | Are there procedures in place that identify and track the status of each participant throughout the study? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 33. | Are there procedures in place for data cleaning? | | | |
| 34. | Are there automated range and logic checks? | | | |
| Safety Plan | | | | |
| 35. | Is a Safety Monitoring Plan in place that outlines independent oversight in the form of a DSMB / Safety Monitoring Body (SMB) /Medical Safety Monitor ? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 36. | Are there procedures in place for documenting and reporting AEs, serious AEs and unexpected AEs , according to NIH Guidelines (http://grants.nih.gov/grants/guide/notice-files/not99-107.html)? See also OHRP Guidelines and NIH Policy regarding unanticipated problems involving research subjects or others (UPIRTSOs). OHRP: http://www.hhs.gov/ohrp/policy/advevntguid.html NIH: http://grants.nih.gov/grants/policy/hs/data_safety.htm | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| Compliance and Monitoring | | | | |
| 37. | Are screening, recruitment, enrollment, and retention reports reviewed regularly and action plans documented? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 38. | Are protocol deviation reports reviewed regularly and violations documented systematically? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 39. | Are there data quality reports that describe missing or erroneous data reviewed regularly to detect and correct problems? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 40. | Are site-monitoring reports generated to provide feedback regarding problems and issues discovered during site visits and to report on the quality of data reviewed? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| Quality Standards | | | | |
| 41. | Have quality standards been established for enrollment and accrual deviations, drop-outs, and data entry and analysis? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 42. | Are procedures in place for correcting inaccurate data and documenting the changes systematically? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 43. | Are procedures in place for amending the protocol and the MOP and documenting the changes? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |

| | | YES | NO | N/A |
|-----|---|--------------------------|--------------------------|-------------------------------------|
| 44. | Are procedures in place to modify quality control reports, if necessary, to capture correct data? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 45. | Are procedures in place to modify training, if necessary, so clinical study site personnel accurately collect data according to the procedures specified in the protocol? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |

**Even if the study is not under IND, the expectation is that there is a binder that holds all study related documents (IRB submissions and approvals, CVs for key study staff, etc. Study binders may be electronic and/or paper.)*