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
Over	view - Study Administration and Procedures			
1.	Are all study documents, including protocol , <u>manual of</u>			
	procedures (MOP), data collection forms, statistical analysis			
	plan (SAP), etc. consistent with data management procedures?			
	Are the MOP , protocol, data collection forms, informed			
2.	consent , etc., easily accessible, in a centrally located binder			
	(electronic or paper), to assist study investigators?			
	Are there accessible participant files that contain source			'
3.	documentation of clinical observations such as lab results,			
	medical record, progress notes, etc.?			
	Is there a study regulatory binder * that contains key study			
4.	documents such as Institutional Review Board (IRB)			
т.	<u>approval</u> , protocol versions, <u>informed consent form</u> , C.V.s,			
	forms, financial disclosures, site monitoring reports?			
5.	Does the <u>training plan</u> describe how and when procedures for			
J.	quality assurance (QA) are implemented?			
6.	Does the training plan include procedures on how to train			
0.	new staff?			
	Does the <u>Drug / Device Distribution Plan</u> specify procedures			
7.	for the storage of, preparation of, dispensing of and handling			
7.	unused intervention as well as procedures for completing			
	<u>treatment accountability</u> logs?			
8.	Are there written plans for obtaining, handling, storing, and			
0.	sending participant samples/materials?			
9.	Are there written procedures for obtaining and transmitting			
	laboratory data?			
10.	Are there procedures in place for following participants from			
	screening and enrollment through completion of the study?			
	Is there documentation of pre-screening and screening			
11.	procedures so that data on eligible and ineligible individuals			
	are captured in an appropriate format? Is a screening log			
	provided?			

		YES	NO	N/A
12.	Does the <u>informed consent</u> include statements about the use of the <u>data and specimen sharing</u> for future research?			
	Is there a written procedure to insure that the current copy of			
13.	the IRB approved informed consent form is signed before each			
	participant is enrolled?			
	Has the <u>manual of procedures (MOP)</u> , which includes the protocol, CRFs, informed consent, study staff roster, screening			
14.	log, and <u>standard operating procedures</u> (SOPs), been			
	distributed to all clinical sites and updated as needed?			
	Have the following study operation procedures or plans been created			
15.	for the MOP:			
	a. Organizational Plan			
	b. Safety Monitoring Plan			
	c. <u>Training Plan</u>			
	d. Study Communications Plan			
	e. Maintaining MOP			
	f. Site Signature Log/Delegation of Authority			
	(Description of Responsibility)			
	g. Recruitment Plan			
	h. Screening and Informed Consent			
	i. Enrollment and Randomization			
	j. <u>Retention Plan</u>			
	k. Study Timelines/Study Visits			
	l. <u>Drug/Device Plan</u>			
	m. Laboratory Specimen Plan			
	n. Blinding/Unblinding			
	o. Concomitant Medications			
	p. Data Management			
	q. Source Documentation			
	r. Case Report Form completion			
	s. Adverse Events (AEs)/Serious Adverse Events (SAEs)			
	t. Participant <u>Withdrawals</u> from study and <u>Lost-to-</u>			
	Follow-ups			
	u. <u>Protocol Deviations</u> and violations			
	v. Quality Assurance (QA)/Quality Control (QC)			
	procedures Maritaria Plan			
	w. Monitoring Plan			
	x. Study Completion	1 1 1		

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

		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?				
32.	Are there procedures in place that identify and track the status of each participant throughout the study?				
33.	Are there procedures in place for data cleaning?				
34.	Are there automated range and logic checks?				
Safet	y Plan				
35.	Is a <u>Safety Monitoring Plan</u> in place that outlines independent oversight in the form of a <u>DSMB / Safety Monitoring Body</u> (<u>SMB) / Medical Safety Monitor</u> ?				
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				
Com	oliance and Monitoring				
37.	Are screening, recruitment, enrollment, and retention reports reviewed regularly and action plans documented?				
38.	Are <u>protocol deviation reports</u> reviewed regularly and violations documented systematically?				
39.	Are there data quality reports that describe missing or erroneous data reviewed regularly to detect and correct problems?				
40.	Are <u>site-monitoring reports</u> generated to provide feedback regarding problems and issues discovered during site visits and to report on the quality of data reviewed?				
Quali	Quality Standards				
41.	Have quality standards been established for enrollment and accrual deviations, drop-outs, and data entry and analysis?				
42.	Are procedures in place for correcting inaccurate data and documenting the changes systematically?				
43.	Are procedures in place for amending the <u>protocol</u> and the <u>MOP</u> and documenting the changes?				

		YES	NO	N/A
44.	Are procedures in place to modify quality control reports, if necessary, to capture correct data?			
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?			

^{*}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.)