

A program of the National Cancer Institute of the National Institutes of Health

Steps for Coordinating Centers to Open ETCTN Trials

This checklist pertains to steps taken by the Coordinating Center and Study Principal Investigator (PI) leading up to ETCTN study activation. For a checklist of steps involved in actual site participation in an ETCTN trial (e.g., obtaining regulatory approval, enrolling patients, entering data), refer to <u>Steps for Participating Sites in ETCTN Trials</u>. During the lead-up to study activation described in this document, the Study PI's site should prepare for study participation according to the site participation document.

<u>Opon Cancer Evaluation Therapy Program (CTEP) Approval on Hold pending</u> Food and Drug Administration	
	A) review, Central Institutional Review Board (CIRB) approval, and study build in Oncology Patient
Enrollment Network (OPEN)/Interactive Web Response System (IWRS)/Rave:	
	The CTSU will announce Approval on Hold from CTEP in the CTSU Bi-Monthly Broadcast.
	For studies being filed to a new Investigational New Drug (IND), CTEP will submit the protocol to the FDA.
	You may be asked to respond to FDA comments and/or submit a revision while simultaneously
	undergoing CIRB review.
	Note: FDA review and approval will take four to five weeks. This activity must be included in study
	activation timelines (i.e., related to the Operational Efficiency Working Group [OEWG] timelines).
	CTEP will submit the protocol and consent to the CIRB concurrently with FDA review.
	The CIRB will contact you to submit the CIRB application. If a protocol revision is requested, it should be
	sent directly to the CIRB.
	Theradex will build OPEN eligibility forms and Medidata Rave website and forms. Once drafts are
	available, Theradex will contact the PI/coordinating center for approval of the forms and to clarify any
	other study-related issues necessary for study build and activation.
	Note: CIRB review/approval and study build in OPEN and Rave will take four to six weeks and will be done
	concurrently. These activities must be included in study activation timelines (i.e., related to OEWG
	deadlines).
	The Protocol and Information Office (PIO) will forward the protocol to the NCI Clinical Trial Reporting
	Office (CTRO), who will register the trial at ClinicalTrials.gov and obtain a National Clinical Trial (NCT)
	number for the study.
	Once CIRB approval is granted, CTEP will review any modifications made during the CIRB review.
	Once CIRB and CTEP have approved the protocol documents, CTEP will submit the protocol to the FDA (if
_	it was not previously submitted as part of the IND filing process).
	As the coordinating center or the Lead Protocol Organization (LPO), create and submit a study specific
_	template Delegation of Tasks Log (DTL) to CTEP via the DTL application on the <u>CTSU members' website</u> .
	For more information, refer to the <u>DTL Template Browser Help (for LPOs)</u> . An approved template DTL is
	required in order to obtain CTEP approval for the study.
П	The PIO will issue a CTEP approval letter upon receipt of FDA approval of a new IND, or submission of
_	protocol to existing IND and completion of Rave build and resolution of any remaining hold issue such as
	drug supply or PI registration.
	ding supply of Firegistration.
Upon CTEP approval:	
	CTSU Operations staff will contact the coordinating center to prepare for study activation and web
	posting. This communication will include an explanation of the remaining steps.
	CTSU Regulatory staff will assist in setting up the protocol in the Regulatory Support System (RSS) to
	allow for the proper collection of regulatory approvals and other study-related documentation.
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Version Date: June 2021 Page 1 of 2

timelines. Leading up to study activation: ☐ The Study PI should send a brief email to the CIRB at earlyphasecirb@emmes.com to include the following (or similar) language: "NCI Protocol #123 is CTEP approved and ready for distribution for IRB review at the participating organizations listed on the protocol cover page." Please copy the CTSU on this email at ctsuprotocol@westat.com and ctsuregsetup@coccg.org. ☐ CTSU Operations and Regulatory staff will communicate with the study team to finalize the Study Activation Memo and to confirm/finalize the study setup in RSS. ☐ CTSU Operations staff will request that the study team review, all requirements for study activation via email. ☐ When ready to open to enrollment, and in close collaboration with the study team, the Lead Academic Organization's (LAO) Grant Administrator will access the Change Request to PIO feature in RSS/Protocols Application to change the study status to Active. ☐ CTSU Operations staff will complete the posting of supporting documentation on the CTSU website protocol page. ☐ The CIRB documents will post to the protocol page on the CTSU website using an automated process based on the activation/release date conveyed by the study team and recorded in RSS/Protocols Application by the CTSU Regulatory Office. Upon study activation: ☐ All study documentation, including CIRB documentation, will be available on the CTSU website protocol ☐ The CIRB's Study Specific Worksheet for Local Context will be available via IRBManager. ☐ The study will be released in OPEN/Theradex IWRS and ready to enroll patients (active studies). ☐ Medidata Rave study invitations will be sent to those persons holding Rave roles at approved sites. ☐ The CTSU will send a notification of study activation to all appropriate Grant/Contract PIs and Grant Administrators, and will include notice of the activation on the CTSU website and in the next CTSU Bimonthly Broadcast. Upon CTEP/CIRB approval of study amendments and study status changes (ongoing basis): ☐ CTSU Operations staff will contact the study team to prepare for amendment releases. ☐ CTSU Regulatory staff will work with the study team to set up study amendments in RSS. ☐ CTSU Operations staff will complete the posting of protocol and supporting documentation on the CTSU website protocol page, and will include notice of the amendment activation on the CTSU website and in the next CTSU Bi-monthly Broadcast. ☐ The Coordinating Center and Study PI or designee (i.e. study team) should notify the CTSU Protocol Team ctsuprotocol@westat.com of trial status changes after initial activation as needed and inform the OPEN (CTSUOPENFORMS@westat.com), Rave (CTMSSupport@theradex.com), and Regulatory (CTSURegSetup@coccg.org) teams. ☐ CTSU Operations staff will communicate with the study team to obtain a status change memo. CTSU Operations will also update the Group Protocol Status in RSS on behalf of the LAO, send a notification of trial status change to all appropriate Grant/Contract PIs and Grant Administrators in case of immediate temporary or permanent closure, and include notice of the status change in the next CTSU Bi-monthly Broadcast.

Note: These activities can take several days to complete and should be included in protocol activation

Version Date: June 2021 Page 2 of 2