

SOP#: MI-3

Non-NIH Site Participant Registration & Status Update

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NCI Clinical Director Signature:

POLICY

Research participants who sign an informed consent document for any Center for Cancer Research (CCR) protocol, including multi-site studies, are considered enrolled in a clinical research study at the time of consent.

All participants at the non-NIH site(s) that sign an informed consent document for a research protocol at the CCR are required to be registered via the CCR Central Registration Office (CRO). When the CCR is the Coordinating Center for a multi-site study, the CCR Study Coordinator is responsible for the registration of participants from non-NIH sites. A 2-step registration process will be used. Non-NIH sites will submit the registration form and notify the CCR Study Coordinator within **24 hours** of participant signing the consent document.

Once the non-NIH site notifies the CCR Study Coordinator (SC), the CCR SC has **24 hours** to complete the registration notification with the CCR CRO.

The CCR SC must be notified by the non-NIH site of any change in a registered participant status within **5 business days** of the non-NIH site becoming aware of the change. CCR SC must then notify the CRO within **5 business days** of becoming aware of the change. Examples of change in status are: screen failure, meets off treatment criteria, meets off study criteria, death.

For Experimental Therapeutics Clinical Trials Network (ETCTN) protocols where the CCR is the corresponding organization, non-NIH sites will not register with the CRO.

Important: Email communication between the CCR SC and non-NIH sites that contains Personally Identifiable Information (PII) must be sent encrypted via Outlook or the NIH Secure E-mail and File Transfer Service (SEFT) – see Resources.

PURPOSE

To describe the steps required to register a research participant from a participating non-NIH site with the CRO when the CCR is the Coordinating Center.

RESOURCES

- CCR Policies/Standard Operating Procedures [website](#)
 - ADCR-2: *CCR Participant Registration & Status Updates*
- CCR Guidelines [website](#) for Secure Email and File Transfer information

PROCEDURE: PARTICIPANT REGISTRATION

1: Provide the registration and status change forms to non-NIH site

- The CCR Study Coordinator (SC) is responsible for providing all non-NIH sites with the *NCI Center for Cancer Research Participant Registration Form* and *NCI Center for Cancer Research Participant Status Change Form*

2: Non-NIH site Registration Step 1

- Within 24 hours of the research participant signing the informed consent document, the non-NIH site will complete the *NCI Center for Cancer Research Participant Registration Form* and send to the CCR SC via **encrypted email**.
 - Signed page of the informed consent document must also be sent to verify enrollment.

Note: Information sent by the non-NIH site must be maintained by the CCR SC as a research record on the team's protocol shared network.

- If the site has included the patient's local medical record number (MRN), CCR SC will send an email confirmation to the non-NIH site that the form was received. The CCR SC will register the participant in PRES or with the Interim Registration Form (see ADCR-2).
- If the site has not included the patient's local MRN, the CCR SC will
 - send an email confirmation to the non-NIH site that the form was received, and
 - forward the registration form to the CRO who will register the participant.
- The question in PRES regarding eligibility may be answered "No" at this point.

3: Registration Step 2 - Verification of eligibility

- Once eligibility is confirmed by the non-NIH site, the site will send an email to the CCR SC indicating that the participant meets eligibility criteria.

Note: Information sent by the non-NIH site must be maintained by the CCR SC as a research record on the team's protocol shared network.

- Once the email confirming eligibility is received, the CCR SC will update PRES or the Interim Registration Form to indicate participant is eligible
- If participant is not eligible see procedure below to remove them from the study as a screen failure.

4: Notify non-NIH site of participant's registration

- The CCR SC is responsible for providing registration information to the non-NIH site. This includes the unique subject number for C3D, if applicable.
- The non-NIH site is responsible for notifying their local pharmacy of participant registration/randomization if applicable.

PROCEDURE: CHANGE IN PARTICIPANT STATUS

When there is a change in a registered participant's status (i.e.: screen failure, meets off treatment criteria, meets off study criteria, death), the non-NIH site must notify the CCR SC within 5 business days of becoming aware of the status change. The CCR SC must notify the CRO within 5 business days of becoming aware of the change.

1: Non-NIH site will complete *NCI Center for Cancer Research Participant Status Change Form*

- When there is a change in participant status (fails screening, meets off treatment criteria, meets off study criteria, death), the CCR Study Coordinator must provide *NCI Center for Cancer Research Participant Status Change Form* to the non-NIH site.
- Non-NIH site is responsible for completing the form and returning to the CCR SC via **encrypted email** within 5 business days.
- CCR SC is responsible for sending an email confirmation to the non-NIH site that the form was received
- The CCR SC will update the participant status in PRES or with the Interim Registration Form (see ADCR-2).

NCI Center for Cancer Research Participant Registration Form

This form MUST be sent via encrypted email

Required*

Protocol Information

1. Protocol number* _____
2. Registering Principal Investigator* _____
3. Outside registering institution* _____

Participant Information

4. Last name* _____
5. First name* _____
6. MRN* _____
7. Diagnosis* _____
8. Date of birth* _____
Please input date in format mm/dd/yyyy

9. Sex*

Male

Female

10. Gender*

Male

Female

Neither exclusively male nor female

Transgender male

Transgender female

Decline to answer

Other: _____

11. Ethnicity*

- Hispanic or Latino
- Not Hispanic or Latino
- Unknown

12. Race*

- White
- Black or African American
- Asian
- Native Hawaiian or other Pacific Islander
- American Indian or Alaska Native
- Multiracial
- Unknown

13. New Subject to NCI*

- Yes
- No

Consent & Study Information

14. Consent language used (if not English): _____

15. Phone consent? *

- Yes
- No

16. Eligible? *

- Yes
- No

17. Protocol consent date (mm/dd/yyyy): * _____

(The date the subject signed the consent. Not the version date of the consent)

18. Protocol cohort * _____

(Please use cohort names as specified in the protocol document)

19. Randomization registration? *

Yes *(Please continue to questions 21 & 22)*

No *(Please continue to question 20)*

20. Protocol arm* _____

(Please use arm names as specified in protocol document)

21. Stratified randomization? *

Yes
If yes, please answer question # 22

No

22. Stratification Information:* _____

NCI Center for Cancer Research Participant Status Change Form

This form MUST be sent via encrypted email

Required*

Protocol Information

1. Protocol number* _____

Participant Information

2. Last name* _____

3. First name* _____

4. MRN* _____

Protocol action

Note that taking a subject off-study also takes them off-treatment if they haven't previously been taken off-treatment

Off-treatment

Off-treatment date* _____

Off-treatment reason*:

- Adverse event
- Completed study
- Death
- Disease progression
- Lost to follow up
- Screen failure
- Withdrew consent

Off-Study

Off-study date*

Off-study reason*:

- Adverse event
- Death
- Disease progression
- Refused further treatment
- Subject non-compliance
- Treatment period completed