INTERVENTIONAL ENROLLMENTS

Once informed consent has been obtained, the following steps should ideally be completed on the same day.

For studies involving bill review in Epic, the following steps must be completed before any study-related procedures are performed and no later than one business day following consent. For all other studies, subject reporting should occur on a monthly basis. Please reference SOP ADM-004 for additional details.

1. Enter the subject into the OnCore Pre-Screening Record
   a. Under the Subjects tab choose Pre-Screening
   b. Select the [New] button
   c. Enter the required information
      i. *Management Group > Choose from dropdown list. The management group should be the group that is handling the screening/coordination efforts for the particular subject.
      ii. *Contact Date = Date subject initially approached about the study
      iii. Referral Channel (if none apply, select “Internal Referral or Consult”)
      iv. Referring Physician = Treating Physician
      v. *Subject Identifier = MRN without leading zeroes
         1. Certain situations may allow for subjects who have a cancer diagnosis to be enrolled without a UF Health MRN. Contact the OnCore Support Team for more information and for assistance with protocol configuration in OnCore.
         2. If no UF Health MRN is found in the EMR and your protocol is configured to allow auto-generation of MRNs, you can auto-generate an OnCore-only MRN:
            a. In the Study Site block, select the blue arrow in the Study Site field and select “No UF Health MRN” from the drop down
            b. In New Subject Details block, click the “Generate” button to auto-generate the Subject MRN.
      vi. *Select the Disease Site or Diagnosis Group for the Subject
         1. For subjects with a diagnosed cancer use Disease Site and choose from dropdown list
         2. For subjects without a diagnosed cancer select: leave blank
      vii. *Race
      viii. *Ethnicity
      ix. *Subject Evaluated = ‘Yes’
      x. *Evaluator = Enter the name of the individual that screened the subject
      xi. Time Spent = Enter the time spent evaluating this subject for study
      xii. *Protocol Number = Enter the OnCore Protocol ID # (OCR Number)
      xiii. *Subject Consented = ‘Yes’
1. This will populate the [Add Subject] button, please complete the rest of the form before selecting
   xiv. Record Completed = ‘Yes’ if all eligibility has been confirmed; ‘No’ if screening procedures are still in process. Once all eligibility has been met, revise this response to ‘Yes’
   xv. Select [Add Subject] > This will re-direct you to the Subject Console

2. Register Subject into the OnCore Study Record under the Subject Console (* all fields/steps required)
   i. Study Site > Choose from dropdown list
   ii. Search for UF Health subjects by MRN and/or last name
      1. Select the appropriate patient from the “Subject Find Results”; this will auto-populate the demographic data from the EMR.
      2. Please confirm the populated race/ethnicity with the subject’s self-reported data. Reporting ‘unknown’ is not permitted. Please use the UFHCC demographics form to collect race and ethnicity data (Appendix A), subjects may select all that apply.
   iii. Select the “Add” button to continue into the study-specific registration section

   Manual creation of records in OnCore for subjects with a MRN is not allowed. If your subject cannot be found please contact the OnCore support team at oncore-support@ahc.ufl.edu or 352-273-5924.

3. Additional Information
   a. Documents/Info (* all fields/steps required)
      i. Click on the ‘Attachments’ tab
      ii. Select [Add]
         1. Document Type = Signed Informed Consent
         2. Version Date = Date of informed consent execution
         3. Attach PDF file of signed Informed Consent. File name should be: MRN_Study OCR#_Consent ID (example: 0123456_OCR1234_ICF-A)
         4. Description = Initial ICF
         5. Select [Add]
      iii. Open file to confirm completeness of upload
   b. Consent (* all fields/steps required)
      i. In the Subject Console, select the Consent tab
      ii. Select [Update] to add an available consent
      iii. In the Available Consents block, select the date the ICF was signed and then click [Select Consents]
      iv. Select the appropriate consent from the Available Consent pop up window
      v. Check the ‘Accepted’ radio button, and then select [Save]
c. **Eligibility (leave blank until you know the subject’s final eligibility status)**
   i. *Eligibility Status = Select the correct eligibility status (Eligible or Not Eligible) from the dropdown. As no waivers of eligibility are allowed, the ‘Eligible (O)’ should not be used.*
   ii. **Verified By = Study Coordinator that confirmed the subject’s eligibility**
   iii. **Version Date = Report the version date of the protocol used to determine eligibility**
   iv. *Status Date = the date that the subject is deemed fully eligible or a screen failure*
   v. **Select [Submit]**

d. **On Study (REQUIRED, even if the subject does not go on study)***Make sure the patient is closed out of EPIC***
   i. **Sequence number can be entered later (once the subject has been formally registered/randomized or is otherwise considered ‘On Study’ per protocol – see subject status definitions below) AND/OR automatically assigned by OnCore (required if subject goes on study)**
   ii. **On Study date = The date the subject is registered/randomized for studies that have a formal process or the date that Eligibility is confirmed for studies that do not have a formal registration/randomization process. Leave date blank until confirmed. (*required if subject goes on study; please see subject status definitions below)**
   iii. *Disease site > select from dropdown
      1. If subject does not have a cancer diagnosis (i.e. is a healthy volunteer or control subject) select: C999 – Healthy Volunteer*
   iv. **Histology > select from dropdown (if the exact histology is not listed, confirm with treating investigator)**
      1. If subject does not have a cancer diagnosis (i.e. is a healthy volunteer or control subject) – leave Histology blank
   v. **Diagnosis Date > select “not available” if not known otherwise enter the actual or approximate date**
   vi. *Disease Site Group - This field will be defaulted to the primary DSG that was designated when the protocol was initially entered into the system. Some protocols serve more than one DSG. It is important to credit the correct DSG with the enrollment as this information feeds into the Cancer Center’s DSG Progress Reports. The group selected should reflect the subject’s disease site. **However, all subjects under the age of 18 must be assigned to the Pediatric DSG.**
   vii. *Study Site > Enter the site where the subject will be seen/treated if it not already defaulted to the correct site*
   viii. **Zip at registration (NOTE: this is the zip code field that is pulled for data reporting). Please enter the Country of Residence into the subject’s Demographic page if the subject is not from the United States**
   ix. Under the Subject Staff box, assign an individual to the following required roles (select [Add] after assigning an individual to each role):
      1. Consenting individual = Person who co-signed the ICF
2. **Enrolling Faculty** = An Enrolling Faculty must be listed for every subject consented to an interventional protocol. This may be the same person listed as the consenting individual. The individual identified as the enrolling faculty will receive the Academic Points associated with the accrual. Only clinical faculty with RVU targets are eligible for Academic Points credit.

3. **Primary Study Coordinator** = primary study coordinator for this subject. If multiple coordinators work on the same study, assign the coordinator who will be primarily responsible for this subject. Only one primary coordinator should be assigned per subject.

x. Select [Submit]

### NON-INTERVENTIONAL ENROLLMENTS

Enrollments to non-interventional studies may be logged into OnCore using the interventional steps outlined above or via the summary accrual steps described below.

Summary accrual for each study should be updated on a **quarterly** basis (1/1-3/31, 4/1-6/30, 7/1-9/30, and 10/1-12/31) at a **minimum**:

- **From Date** = 1st of the month
- **Thru Date** = End of the month
- **Accrual** = Enter number of patient(s) enrolled during the month

  1. Enter the subject into the OnCore Accrual Record

     **Note: Study must have been “Open to Accrual” to allow subject accrual entry**

a. Under the Protocols PC Console tab choose Accrual
b. Select the [Update] button
c. Enter the required information
   i. *From Date = 1st of the month*
   ii. *Thru Date = End of the month*
   iii. *Accrual = Enter number of patient(s) enrolled during the month*

   1. i, ii, and iii can either be individual subject and date entry or can be a cumulative range if the demographic information is **the same**. These dates should reference when the subject(s) was formally enrolled in the study (i.e. registered/randomized)

   a. For example, if 5 subjects were enrolled over a one-month period and their demographic information was similar, then all 5 accruals could be entered at the same time. Any enrollments added using a cumulative range must be entered within the same calendar year (i.e. enrollments must be broken down by year).

   iv. Select Institution from the drop down

   v. *Select the Internal Accrual Reporting Group (this may populate automatically)*

   vi. *Gender*

   vii. *Age Group*
1. Age ranges in OnCore are presented in 10 year increments.

viii. *Ethnicity
   1. If available, please confirm the populated race/ethnicity with the subject’s self-reported data. Reporting ‘unknown’ is not permitted.

ix. *Race
   1. If available, please confirm the populated race/ethnicity with the subject’s self-reported data. Reporting ‘unknown’ is not permitted.

x. *Select the Disease Site or Diagnosis Group for the Subject
   1. For subjects with a diagnosed cancer, use Disease Site and choose from dropdown list
   2. For subjects without a diagnosed cancer, leave this blank

xi. Recruited by either the Investigator or treating physician (if they are the same; if the name on the signed Consent form is not a physician or faculty member, enter the Investigator name]. If not known, leave blank.

xii. Zip code = subject’s zip at registration

xiii. Select [Add] > This will update the below listing of accruals and update the Accrual table.

xiv. Select [Close]

**If the study is not set up for Summary Accrual, you will not have the [update] button to select on the Accrual tab. If you believe you should have the option to report accruals via the Summary mechanism please contact Ashley Anderson at adanderson@ufl.edu.

UNDERSTANDING SUBJECT STATUS FOR INDIVIDUALLY REGISTERED SUBJECTS

The Cancer Center requires that all subject status updates for interventional enrollments be made within one day of adjustment. The primary study coordinator is responsible for ensuring OnCore accurately reflects the current status of all subjects. The “normal” sequence of statuses for a subject is as follows:

1. **Consented:** Subject has at least one consent signed date entered and linked to a consent as added in the PC Console to the subject’s study site. This is the initial status for all subjects who consent to a study.

2. **Eligibility:**
   (1) **Eligible:** Subject met all eligibility criteria and will be moved to On Study
   (2) **Ineligible:** Subject failed one or more eligibility criteria, withdrew consent prior to registration/randomization, or will otherwise not be moved to On Study
   (3) **Eligible(O):** Eligible with override: Online questionnaire complete with at least one criteria not met, however, explanation and PI approval to override, or eligible(O) entered with associated date. This status should no longer be utilized. Status described for informational purposes only.
3. **On Study:** Subject is considered *accrued* when On Study date is entered in OnCore. Subjects may be moved to the On Study status once they have been formally registered/randomized or are otherwise considered evaluable per the protocol. This step typically occurs following verification of eligibility. For studies that do not include a “treatment”, subjects will not be moved to the On or Off Treatment statuses. Instead, those subjects will remain On Study during the study participation/intervention period and then will move to an On Follow-up or Off Study status as appropriate.

4. **On Treatment:** Subject is considered On Treatment when associated with a Treatment Arm and an On Treatment date is entered within the subject console.

5. **Off Treatment:** An Off Treatment Date is entered for the subject. This date should reflect either the day after the last dose of therapy for subjects that complete treatment per protocol or the day that the decision is made to discontinue therapy otherwise. All subjects should be immediately moved into follow-up status or off study depending on the study.

6. **On Follow-up:** A Follow-up Start Date is entered for the subject. All subjects in long term follow-up should be in this status. Typically, the On Follow-up date is the date after the subject’s Off Treatment date.

7. **Off Study:** An Off Study date is entered for the subject. This status is used when it is determined that no further interaction with the subject is required. You should confirm how long each subject is to be followed per protocol and their consent. If they are followed indefinitely, this date is only entered upon death (will populate automatically if you enter an expiration date), the subject withdraws from further follow-up, or the study is closed at the IRB (you will have to manually enter this date for all subjects).

**SERIOUS ADVERSE EVENT REPORTING**

The Cancer Center requires that all SAEs occurring in subjects enrolled on UF interventional IITs be logged into OnCore.

In the Subject Console > SAEs tab, click **New**.

Enter the following information in the Subject SAE Update section (at the top):

- **Event Date:** enter the start date of the event
- **Event End Date:** enter the end date of the event
- **Reported Date:** enter the date you initially reported the event (to sponsor, IRB, DISC, etc.)
- **Reported By:** enter your name
OnCore Subject Registration and Reporting Guidance
UFHCC Requirements
Version 4.0 (02/20/2019)

• **Event Narrative**: enter the symptoms/diagnosis, relevant testing, required interventions, etc. as applicable
• **Protocol Attribution**: select attribution from the drop down list
• **Outcome**: select outcome from the drop down list
• **Consent Form Change Required**: yes/no
• **SAE Classification**: select from list

In the Adverse Event Details section in the middle of the page, enter the following information:

• **Category**: select the appropriate CTCAE category from list
• **AE Detail**: select the appropriate CTCAE term from list
• **Grade/Severity**: select grade from drop down list
• **Unexpected**: yes/no
• **DLT**: yes/no
• **Action**: select action from drop down list
• **Attribution**: select the attribution for each component of the trial from the drop down lists

Click **Add** (in the middle of the page). The details you enter appear in the table.

In the Adverse Events Details section in the middle of the page, document a second category using the **Select Detail** link if necessary.

Click **Submit** to save the basic SAE information.

**DEVIATION REPORTING**

The Cancer Center requires that, at a minimum, all major deviations/violations related to subjects enrolled on UF interventional IITs be logged into OnCore.

A deviation/violation is a variance from the approved protocol procedures. Deviations specific to an individual subject are entered via the Subject Console > Deviations tab.

Navigate to the **Subject Console > Deviations** tab.

The table at the top of the page shows the subject’s deviations. The bottom table shows any visits with a visit date outside of the planned date’s tolerance. This table is for informational purposes, and you can use it as a guide for entering deviations if needed.

Click **New** to create a deviation.
The **Date Discovered** and **Reported By** fields default to the current date and user, but they can be changed. When entering Deviation data, the required fields are marked with an asterisk.

Enter the following:

- **Date Discovered**: enter the date of discovery
- **Reported By**: enter your name
- **Deviation Date**: enter the date the deviation occurred
- **Category**: select category from drop down list
- **Description of Deviation**: please describe the deviation including the root cause and whether or not the deviation occurred to prevent an immediate risk to the subject
- **Effect on Patient Safety**: please describe whether or not the event affected the safety or clinical care of the subject
- **Action Taken**: please describe any corrective and preventative actions taken
- **Report to IRB?**: select from drop down list

Click **Submit** to create the deviation.

Other information can be entered to record when it should be reported to the IRB, the date it was reported, and the date it was reported to the sponsor.

Click the **Deviations** vertical tab to see the updated page.

The **Delete** link allows you to delete a deviation record entered in error.

Publication Date:
Effective Date:
Scope: This process applies to all subjects consented to cancer-relevant interventional, observational and ancillary/correlative studies.

Approved By:

Alison Ivey
Administrative Director
UFHCC Clinical Research Office
Appendix A

Research Subject Information Sheet

The following demographic information is required by one or more research studies for which you have given consent to participate. This information may be reported to the study sponsor(s) for statistics on the demographics of research subjects. Study sponsors can be an industry sponsor, academic institution, national health organization or other type of cancer group. This information may also be used for internal quality improvement activities within the University of Florida Health Cancer Center. Unless otherwise specified in your research study’s informed consent, your name will not be reported to the study sponsor(s).

Subject’s Name: ____________________________________________________________

Date of Birth: ______________

Primary Race: Select all that apply

- American Indian or Alaska Native □
- Asian □
- Black or African American □
- Native Hawaiian or Other Pacific Islander □
- White □
- Decline to report □

Ethnicity: Choose One

- Hispanic or Latino □
- Not Hispanic or Latino □
- Decline to report □

The information below is optional and will only be used by University of Florida Health Cancer Center staff in order to perform follow up activities or to schedule/re-schedule appointments.

Home Address: ____________________________________________________________

Home Phone #: ____________________________________________________________

Cell Phone # (if applicable): ________________________________________________

If you do not have a cell phone, please provide a local contact phone number: ______________________________________________

Local address (if applicable): ________________________________________________

Email Address (only if you prefer to be contacted by email*): ________________________

Spouse or Caregiver’s Name: ________________________________________________

Name of Person Completing Form: ____________________________________________