

**DANA-FARBER / HARVARD CANCER CENTER  
STANDARD OPERATING PROCEDURES FOR CLINICAL RESEARCH**

<b>TITLE:</b> Documenting Study-Specific Training	
<b>SOP #:</b> ET-203	<b>Page:</b> 1 of 1

**Applicable Regulations  
& Guidelines:**

21 CFR 312.53    Selecting Investigators and Monitors  
May 1997        International Conference on Harmonization; Good Clinical Practice:  
Consolidated Guideline, section 4

**Other References:**

ET-202            Sample Delegation of Authority and Study Training Log

**Responsible Personnel:**

DF/HCC Principal Investigator (PI) or study staff designee

**Policy Statement:**

The PI or study staff designee is responsible for ensuring training on study-specific tasks and documentation of this training.

**Procedure:**

- 1) The PI or study staff designee will provide study-specific training for research protocols **as appropriate**.
  
- 2) Training includes, but is not limited to:
  - Initial training delivered at the site initiation visit
  - Consenting process
  - Ongoing training for modifications to study procedures
  - Training of new or replacement staff added after the site initiation visit
  - Study staff of all protocols utilizing electronic data capture (eDC) through the QACT must have training in Phase Forward “Inform” conducted by the QACT office.
  
- 3) The method to document study-specific training will vary according to the mechanism provided by the sponsor.
  - For industry sponsored studies, use the document(s) provided by the company.
  
  - For cooperative group or PI-initiated studies, there is no document provided by the sponsor. Use the *Sample Delegation of Authority and Study Training Log* attached to policy ET-202.

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