1) OHRPP’s “Learn at Lunch” – *upcoming presentations*

**Topic: “All About Adverse Events”**

Presenter: **Anya Rosensteel**, Quality Improvement Unit, OHRPP

Summary: A review of OHRPP guidance and tools to support adverse event assessment, documentation, and reporting (when appropriate)

Date: **October 26, 2021** (today!)

Time: **Noon-1pm**

Location: **Zoom** ([Register](#) for this meeting)

**Topic: “How to Respond to IRB Requests”**

Presenter: **Tiffany Rose**, Quality Improvement Unit, OHRPP

Summary: A review of types of IRB Requests and how to address them

Date: **November 16, 2021**

Time: **Noon-1pm**

Location: **Zoom** ([Register](#) for this meeting)

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2) BruinIRB Expanded Access Submissions

- All Expanded Access submissions should be submitted through the [BruinIRB](#) system. As of October 8th, webIRB has been updated to no longer allow Expanded Access as a clinical trial option in section 8.3.

- Please refer to [OHRPP Guidance and Procedures: Use of Drugs and Biologics in Clinical Research and Treatment](#) or [OHRPP Guidance and Procedures: Use of Devices in Clinical Research and Treatment](#) for information about FDA Expanded Access programs.

3) BruinIRB Training Guides

- [BruinIRB Training Guides](#) are available for the following topics:
  - [Creating a New Study](#)
  - [Create and Submit a Continuing Review or Modification](#)
  - [Updating Your Contact Information and Profile](#)
  - [How to Respond to IRB Requests](#)
  - [Navigating BruinIRB for Researchers](#)
  - [PI Assurances for a New Study](#)

4) Updated OHRPP Guidance

- [Funding Considerations for Federally-Funded and Industry-Sponsored Human Research](#)
5) OHRPP’s Office Hours

- Register once to join any session:

Upcoming Office Hours:
- November 9, 2021 8:30am
- November 23, 2021 8:30am

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