Working with WCG-WIRB – Yeshiva University
Agenda

- Overview of WCG and WIRB
- Submission Process
- Submission Form Assistance
  - Initial Review
  - Change in Research (Modifications)
  - Promptly Reportable Information (PRI)
  - Continuing Review Report Form (CRRF)
- MyConnexus Submission Overview
- Note on the New Common Rule
Overview of WCG-WIRB
Several Companies, One United Mission:
Provide the people who perform clinical trials with the highest quality of services to accelerate the scientific advancement of human health, while ensuring that the risks of progress never outweigh the value of human life.
Institutions – Part of WCG-WIRB’s DNA

- >200 Academic Medical Centers
- 2,800 hospital facilities under contract
- Dedicated Account Manager
- WCG-WIRB learns how you operate
- WCG-WIRB flexes to your requirements
Ethics and Compliance History and Experience

Unmatched in the Industry

- 50+ Years of Regulatory Experience – First Commercial IRB 1968
- Senior Advisors to: FDA, OHRP, AAHRPP
- Longest AAHRPP Accreditation History – First IRB Accredited
- Re-Accredited with No Findings
- 20 Successful FDA Audits - Dating Back to 1983
- Over 500 Successful Sponsor/CRO Audits
- 40+ Certified IRB (CIPs) Professionals on Staff
- WCG-WIRB: 5 US Panels (10 meetings per week)
Submission Assistance and MyConnexus Submission Overview
Preparing Your **Initial Review Submission:**
**Additional PI Site**

- Contact WCG-WIRB Client Services (or Account Manager) to ask whether WCG-WIRB has reviewed your protocol (provide # or title)

- If yes –
  - Request WCG-WIRB approved ICF Templates for the study
  - Ask whether the study falls under the Single Review Solution (SRS)

- If no –
  - Compile all protocol/site documents and submit in Connexus as a new study.
Preparing Your **Initial Review Submission**

Smart PDF forms are available in two locations:

- **WIRB.com** > Download Forms
- **MyConnexus** > Quick Access Links > IRB Forms and Guides
Preparing Your **Initial Review Submission**

- Complete Initial Review Submission Form
  - Select WIRB as Destination IRB

### Destination Institutional Review Board (IRB)

*To which WCG IRB is this application being submitted?*

*If you have questions, please call or email the selected IRB*

- Aspire IRB (Aspire)  
  - (877) 366-5414  
  - email@aspire-irb.com
- Copernicus Group IRB (CGIRB)  
  - (888) 303-2224, (919) 465-4310  
  - irb@cgirb.com
- Hummingbird IRB (HIRB)  
  - (855) 447-2123  
  - info@HummingbirdIRB.com
- Midlands IRB (MLIRB)  
  - (800) 636-4445, (913) 385-1414  
  - info@mlirb.com
- New England IRB (NEIRB)  
  - (800) 232-9570, (617) 243-3924  
  - info@neirb.com
- Western IRB (WIRB)  
  - (800) 562-4789  
  - clientservices@wirb.com
Preparing Your **Initial Review Submission**

- Complete Initial Review Submission Form
  - Select Submission Type (2 Common Options):

  **Submission Type**
  
  *Indicate the type of submission:
  - New protocol with no Principal Investigator (PI) or site information
  - Site being added to existing protocol, or change of Principal Investigator (PI)
  - New protocol and Principal Investigator (PI) (combined submission)

  *For clinical use of a Humanitarian Use Device (HUD), Expanded Access, Compassionate Use, and Emergency Use see separate application forms on the IRB Web site.*
Preparing Your **Initial Review Submission**

- Complete Initial Review Submission Form
- Exempt from IRB oversight:

**IRB Determinations**

*If the IRB determines that the submission does not represent human research or represents research that is exempt from regulation, do you want the IRB to issue an exempt or not human research determination instead of conducting IRB review? For research to be exempt from regulation, the research must be limited to:*

- Evaluation of educational methods
- Surveys/interviews/focus groups
- Benign behavioral interventions
- Use/review of specimens/information collected for purposes other than the proposed research
- Evaluation of taste and food quality
- Creation of a biobank
- Use of data and specimens from a biobank

*For more information see 45 CFR §46.104(d)*

![Yes or No poll](image)

Note: WCG-WIRB does not require status updates if the research is found exempt from IRB oversight. However, you must provide substantive changes in your research for re-evaluation.
Preparing Your **Initial Review Submission**

- **Complete Initial Review Submission Form**
  - **Site Contacts**

  ![Image of Research Contact Form]

  **Contacts**
  Are there any designated contacts for this research (e.g., Sponsor contact, Contract Research Organization (CRO) contact, Site Management Organization (SMO) contact, study coordinator contact)?
  - Yes
  - No

  **Research Contact**
  - *Contact Type*
  - Prefix *First*
  - Middle
  - *Last*
  - Suffix
  - *Email*
  - *Phone*

  **Degrees**
  - *Company/Institution/Organization*

  **Mailing address for the above individual:**
  - *Address Line 1*
  - Address Line 2

  **City**
  - *State*
  - *Postal Code*

  **Country**

  - Copy this person on IRB correspondence
  - Send continuing review forms to this person to be filled out and returned to the IRB

  **Add another contact**

Note: WCG-WIRB will allow the Student to be listed as the PI as long as they are capable of the oversight of the research. The Faculty Advisor should be listed as a coordinator.
Preparing Your **Initial Review Submission**

- Complete Initial Review Submission Form
  - Research Locations—List all where research activity conducted

![Research Location Form](image)
Preparing Your **Initial Review Submission**

- Complete Initial Review Submission Form
  - Select YES to Institutional Services and Indicate Institution Name/Number:

![Institutional Services form]

**Yeshiva University** Institution ID: **127265**
Preparing Your **Initial Review Submission**

- Complete Initial Review Submission Form
- Human Subjects Protection Training Requirement

**Research Team Training**

The Principal Investigator (PI) must ensure that all investigators and research staff undergo training on the ethics and regulations of human subject protections before being involved in the conduct of this research. For clinical research, the Principal Investigator (PI) must ensure that all investigators and research staff undergo training on Good Clinical Practice (GCP).

*Have all investigators and research staff involved with the conduct of this research taken one or more of the following programs and all applicable training programs noted as required?*

- ACRP Certified Clinical Investigator Training
- CenterWatch: Protecting Study Volunteers in Research
- Collaborative IRB Training Initiative (CITI)
- DIA Certified Investigator (CCI)
- SOCRA Clinical Research Professional (CRP)
- Tri-Council Policy Statement online training (TCPS)
- WCG Academy

**Note:** WCG-WIRB does not require Certificates of Training.
Preparing Your **Initial Review Submission**

- Complete Initial Review Submission Form
  - Conflict of Interest / Financial Interest Disclosure

**Financial Interest Disclosure**

*Does the Principal Investigator (PI), the PI's immediate family, or any other research personnel or their immediate families, have any of the following financial interests in an entity that is sponsoring the research, or an entity that is manufacturing the product or service being tested?*

- Any remuneration from the entity in the previous twelve months that exceeds $5,000, when aggregated for the individual and their immediate family. *(Remuneration includes salary and any payment for services not otherwise identified as salary, such as consulting fees, honoraria, or paid authorship)*
- Any equity interest in the entity. *(Equity interest includes any stock, stock option, or other ownership interest)*
- Any intellectual property rights and interests *(e.g., patents, copyrights)*
- Any governance or executive relationship with the entity *(e.g., board of director, CEO)*

- [ ] Yes
- [x] No
Submission Documents

- The form will guide you as to what documents are required for submission
  - There will be ⚠️ signs in the form if supplemental forms are required
  - Reference the end of the form for a list of required documents

- The form includes a tool to check for missing entries
  - Missing entries will be highlighted
Preparing Your **Initial Review Submission**

- Ensure that you have complied with your local Yeshiva University processes
Preparing Your **Initial Review Submission**

- **Log into Connexus**

  **Create a New Account**
  If you are a new user, you must create a new account to access the system. Fill in the form with the required information and click the Register button to continue. If you need help, click the Registration Help link. You can also Request Support via the provided link.

  **Existing User Login**
  If you already have an existing account, log in to the right.

  **Live Support**
  Click the Live Support ONLINE button to chat with a representative. If there are no representatives available, you can leave a message.

  **Forgot Password**
  Click the Forgot Password link to reset your password.

  **Quick links**
  Access helpful links without having to login.
Preparing Your **Initial Review Submission: Additional PI Site**

- Find Study (if **additional site** to existing study)
- Click the “**My Studies**” tab
- Find the study to be submitted and click on the **blue** “IRB Tracking” number to select
- Under “Submissions for this Study” select “Submit New Investigator” at the top right of your screen
Preparing Your **Initial Review** Submission: Additional PI Site

- Complete Wizard, Upload Documents, and Submit!

![Submission Form](image)
Preparing Your Initial Review Submission: New Study

Make a Submission to the IRB

Is your submission:

- Initial Review Submission
- Submission for an Already Approved Study or Investigator

This Initial Review Submission is for:

- Review of a New Research Protocol
  Will be redirected to the new protocol submission wizard
- Specific Investigator/ Site's Participation in a Study
  Please select your Study from the list on the next page to make your submission or Request Access if your Study is not shown
Preparing Your **Initial Review** Submission: New Study

Typically used by a CRO or Sponsor; used when needing to submit a new study protocol (and associated documents) that has not yet been reviewed by the IRB.

- **Submission Name:**
- **Sponsor Protocol ID:**
- Is this a Single Review Solution submission?:
- Where would you like to route this submission?:
- Who are you requesting this New Research submission to be reviewed by?:
- Is this research minimal risk and being conducted at a single site?:
- Comments or notes concerning the submission:

[Submit]  [Cancel]
After You Submit...

- You receive a **Submission Tracking Number**
- WCG-WIRB staff prepares the submission
- A WCG-WIRB panel or expedited reviewer reviews the research for your site
- WCG-WIRB staff assembles and finalizes documents
- You receive an e-mail with a link to all WCG-WIRB outcome documents
  - If the research is approved, you receive your approval documents
  - If the decision is to Defer or Disapprove, the link contains a regulatory letter with rationale for the decision
- Outcome documents are posted to MyConnexus for reference
After You Submit…

- You will receive a Certificate of Action “COA” with your Outcome Documents. It will list the following:
  - WCG-WIRB Board Action Date “Approval Date”
  - Expiration Date
  - Approved Research Location(s) and PI
  - The documents that were reviewed
  - List of study personnel on the email distribution list.

- You can add others to the MyConnexus workspace for your PI.
- Use the Contact Information Update Form for changes to study contacts or continuing review contacts.
Contact Information Update Form

- Necessary to update the contacts who receive CRRFs
Modifications and Amendments

Amendments and modifications can be submitted using our “Change in Research” form

- Add additional research locations
- Indicate if there is a change in financial disclosure
- Sponsors can submit changes on your behalf
The “Promptly Reportable Information” form is used to report any adverse events or unanticipated problems:

- The form will guide you as to what problems to report
- WCG-WIRB will review the report and if significant, communicate with appropriate parties
- If we find that the event does not constitute an increased risk to subjects, we will file it without action.
- If you need an email stating the event was filed, contact your Account Manager.
The “Continuing Review Report Form” will be sent automatically to the individual(s) listed on the Initial Review form

- Forms are sent out approximately 86 days before the study expiration date, and are due approximately 56 days before the study expiration date
- All sites are brought on to a single protocol-level expiration date
- If you receive initial approval within 90 days of the protocol-level expiration date, you will automatically be brought on to the next continuing review period
- Work order is reviewed 10-14 days before expiration date
Note on the New Common Rule

- WCG-WIRB will only review federally-funded studies under the new Common Rule unless requested otherwise.
- Consent Form Template with a summary is available on our Download Forms page.
We Are Here to Partner With You – Contact Us!

General Questions
WCG-WIRB Client Services
+1 360.252-2500 | clientservices@wirb.com

Escalated/Urgent Questions
Jon Gellert
+1 360.570.1309 | jgellert@wirb.com

Presentation by
Christopher Gennai, CIP
+1 360.252.2460 | cgennai@wirb.com
Thank You