



EQulP's Mission

EQulP's mission is to improve policies and practices in clinical research at Boston Children's Hospital with the aim to continually maximize the protection of human subjects, and to promote good clinical practices regarding research conduct and documentation. EQulP aims and goals are created to educate and support the research community to safely conduct compliant clinical research. Please feel free to contact us with questions or comments

EQulP Services

Study Audit (required or optional)

A one-time, confidential study audit to ensure compliance with applicable regulations and policies, and to evaluate study conduct, organization and documentation. The EQulP office aims to help research teams implement tools and strategies to improve identified 'problem' areas.

Study audits are required for routine, not-for-cause audits (random selection) and directed, for-cause audits (IRB requested). PI/staff can request an optional full or partial study audit (e.g. ensure compliance, during staff changes, to prepare for outside audit).

New/Transfer Investigator Training (required)

New Investigators (conducting clinical research for the first time) and Transfer Investigators (conducting clinical research at BCH for first time) must meet with the EQulP office prior to IRB approval to review applicable resources, regulations and BCH policies.

One-on-One Assistance (optional)

EQulP office can provide one-on-one assistance regarding best practices for study documentation to investigators and their research teams.

Talks/Presentations (optional)

Upon request, the EQulP staff is available to present various topics about research compliance and good clinical practices.

Educational Initiatives and Materials

Educational initiatives and materials are continuously developed and disseminated to assist the research community. Guidelines and templates are available on the EQulP website.

Study Tools & Templates

The **EQulP website** offers many study tools and templates to download and customize for your study. Guidance documents are available for most templates.

- **Subject Screening Log**
- **Subject Enrollment Log**
- **Subject Eligibility Checklist**
- **Study Visit Checklist**
- **Informed Consent Checklist**
- **Memo-to-File**
- **Monitoring Log**
- **Recruitment Activity Log**
- **Remuneration Log**
- **Minor Deviation Log**
- **Drug & Device Accountability Logs**
- **IRB Tracking Log**
- **Consent Revision Log**
- **Clinical Trial Study Documents**
- **CRF-Source Document Log**
- **Roles and Responsibilities Log**
- **Staff Training Log**
- **Staff Signature Log**
- **Clinical Trial Tasks**
- **Subject Case History**
- **Electronic Document Storage**
- **Storing Consent in Medical**

Contact Information

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