

COLLABORATIVE INSTITUTIONAL TRAINING INITIATIVE (CITI PROGRAM)

COMPLETION REPORT - PART 1 OF 2 COURSEWORK REQUIREMENTS*

* NOTE: Scores on this Requirements Report reflect quiz completions at the time all requirements for the course were met. See list below for details. See separate Transcript Report for more recent quiz scores, including those on optional (supplemental) course elements.

- **Name:** roland moskowitz (ID: 37561)
- **Institution Affiliation:** Schulman IRB (ID: 1920)
- **Institution Email:** rollie.moskowitz@rapidmedicalresearch.com
- **Institution Unit:** investigator

- **Curriculum Group:** CITI Good Clinical Practice
- **Course Learner Group:** GCP Course that Expires in 2 Years
- **Stage:** Stage 1 - Basic
- **Description:** This ICH E6 GCP Investigator Site Training meets the Minimum Criteria for ICH GCP Investigator Site Personnel Training identified by TransCelerate BioPharma as necessary to enable mutual recognition of GCP training among trial sponsors.

- **Record ID:** 23322064
- **Completion Date:** 30-May-2017
- **Expiration Date:** 30-May-2019
- **Minimum Passing:** 80
- **Reported Score*:** 91

REQUIRED AND ELECTIVE MODULES ONLY

	DATE COMPLETED	SCORE
Overview of ICH GCP (ID: 1352)	30-May-2017	4/4 (100%)
ICH - Comparison Between ICH GCP E6 and U.S. FDA Regulations (ID: 1354)	30-May-2017	4/5 (80%)
Conducting Investigator-Initiated Studies According to FDA Regulations and GCP (ID: 1355)	30-May-2017	3/3 (100%)
Investigator Obligations in FDA-Regulated Research (ID: 1356)	30-May-2017	5/5 (100%)
Managing Investigational Agents According to GCP Requirements (ID: 1357)	30-May-2017	4/5 (80%)
Overview of U.S. FDA Regulations for Medical Devices (ID: 1358)	30-May-2017	3/3 (100%)
Informed Consent in Clinical Trials of Drugs, Biologics, and Devices (ID: 1359)	30-May-2017	4/4 (100%)
Detecting and Evaluating Adverse Events (ID: 1360)	30-May-2017	4/4 (100%)
Reporting Serious Adverse Events (ID: 1361)	30-May-2017	4/4 (100%)
Monitoring of Clinical Trials by Industry Sponsors (ID: 1362)	30-May-2017	4/5 (80%)
Audits and Inspections of Clinical Trials (ID: 1363)	30-May-2017	5/5 (100%)
Completing the CITI GCP Course (ID: 1364)	30-May-2017	No Quiz
Overview of New Drug Development (ID: 1351)	30-May-2017	4/5 (80%)
The CITI Good Clinical Practice Course for Clinical Trials Involving Drugs and Devices (ID: 1350)	30-May-2017	2/3 (67%)

For this Report to be valid, the learner identified above must have had a valid affiliation with the CITI Program subscribing institution identified above or have been a paid Independent Learner.

Verify at: www.citiprogram.org/verify/?k777e109e-bbab-4cdb-8322-d1bc59022c12-23322064

Collaborative Institutional Training Initiative (CITI Program)

Email: support@citiprogram.org

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COLLABORATIVE INSTITUTIONAL TRAINING INITIATIVE (CITI PROGRAM)

COMPLETION REPORT - PART 2 OF 2

COURSEWORK TRANSCRIPT**

** NOTE: Scores on this [Transcript Report](#) reflect the most current quiz completions, including quizzes on optional (supplemental) elements of the course. See list below for details. See separate Requirements Report for the reported scores at the time all requirements for the course were met.

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- **Record ID:** 23322064
- **Report Date:** 30-May-2017
- **Current Score**:** 91

REQUIRED, ELECTIVE, AND SUPPLEMENTAL MODULES

	MOST RECENT	SCORE
The CITI Good Clinical Practice Course for Clinical Trials Involving Drugs and Devices (ID: 1350)	30-May-2017	2/3 (67%)
Overview of New Drug Development (ID: 1351)	30-May-2017	4/5 (80%)
Overview of ICH GCP (ID: 1352)	30-May-2017	4/4 (100%)
ICH - Comparison Between ICH GCP E6 and U.S. FDA Regulations (ID: 1354)	30-May-2017	4/5 (80%)
Conducting Investigator-Initiated Studies According to FDA Regulations and GCP (ID: 1355)	30-May-2017	3/3 (100%)
Investigator Obligations in FDA-Regulated Research (ID: 1356)	30-May-2017	5/5 (100%)
Managing Investigational Agents According to GCP Requirements (ID: 1357)	30-May-2017	4/5 (80%)
Overview of U.S. FDA Regulations for Medical Devices (ID: 1358)	30-May-2017	3/3 (100%)
Informed Consent in Clinical Trials of Drugs, Biologics, and Devices (ID: 1359)	30-May-2017	4/4 (100%)
Detecting and Evaluating Adverse Events (ID: 1360)	30-May-2017	4/4 (100%)
Reporting Serious Adverse Events (ID: 1361)	30-May-2017	4/4 (100%)
Audits and Inspections of Clinical Trials (ID: 1363)	30-May-2017	5/5 (100%)
Monitoring of Clinical Trials by Industry Sponsors (ID: 1362)	30-May-2017	4/5 (80%)
Completing the CITI GCP Course (ID: 1364)	30-May-2017	No Quiz

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