

Certificate of Completion

Introduction to the Clinical Drug Development Process: ICH Good Clinical Practice for Clinical Trial Sites

**Module 1: Course Overview & Overview of ICH Good
Clinical Practice**

Congratulations to

Hilde Bollen

Membership Affiliation and #:
Employee Number:

For Successfully Completing Training
on

June 28, 2018

Provided by: **Quintiles Global Learning**



This document must be retained by the licensee for a period of four years after the course concludes

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".

Certificate of Completion

Introduction to the Clinical Drug Development Process: ICH Good Clinical Practice for Clinical Trial Sites

**Module 2: Clinical Investigator Obligations &
Qualifications, Resources, IRBs/IECs**

Congratulations to

HILDE BOLLEN

Membership Affiliation and #:
Employee Number:

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on

July 1, 2018

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**Module 3: Subject Informed Consent & Protocol
Compliance**

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**Module 4: Investigational Product, Randomization, and
Unblinding & Source Documents and Case Report Form**

Completion
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**Module 5: Safety Reporting, Financial Disclosure, & Study
Closeout, Trial Termination, and Record Retention**

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**Module 6: Non-compliance, Scientific Misconduct and
Fraud & Monitoring and Preparing for Audits and
Inspections**
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Module 7: The European Clinical Trial Directives

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