

This training workshop will take place in a virtual format via a digital platform

Live virtual Instructor-led Training Course

INTRODUCTION TO CLINICAL RESEARCH

Module 1: 18th November 2020 - Module 2: 25th November 2020



Virtual Classroom

Who Should Attend?

This course is designed for those with **no** or **little experience** in clinical research or those seeking **an overview of the** clinical research process, regulations, terminology, acronyms, roles and responsibilities within clinical research.

Course Objectives

The course offers a general overview of the drug development process and most relevant tasks and responsibilities within the clinical research setting. With this course you will:

- Understand how drugs are developed and who the key players are in the process
- Become knowledgeable about the terminology used in the clinical research environment
- Find out what rules, guidelines and laws are essential
- Learn how a clinical study is set up and performed
- Become familiar with and understand how to handle the documentation required in clinical research

Course Modules

08:30 - 12:00 CET (incl. breaks)

- Developing a drug in the pharmaceutical industry from the idea to the product you buy in the pharmacy and the contributions of clinical research
- What is a clinical study the four phases of clinical development
- Who are the key players and what are their roles in the process
- The language and abbreviations the clinical research team are using every day
- · What are the rules, guidelines and laws that need to be followed and why are they so important
- How is a clinical study performed from preparation over conduct to termination of a study
- A key to a successful study: the master study file. What documents are essential and how are they handled, filed, and archived?



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Course Leader

Tamara Schärer (MSc, ECPM), Managing Director, SRS Schärer Research Services, Basel, Switzerland Ms Schärer started working in the pharmaceutical industry as a Clinical Research Scientist in the field of CNS for Sandoz in 1996. After the merger to Novartis, she became the Therapeutic Area Coordinator for all Phase IV studies in Oncology. In 1999 Tamara moved to a small CRO as an International Project Manager and later trained as a GCP Auditor to become Quality Manager Clinical Operations. In 2002, she founded SRS Schärer Research Services.

Learning Methodology

The combination of live trainer interaction, information sharing and opportunities for conversations and discussions of the theory will help ensure you have a comprehensive understanding of the course content.

Technical Requirements

The training will be delivered via digital platform. In order get access you will need an internet connection, audio (from computer speakers or microphone), and a webcam (standalone or integrated with your monitor/laptop).

Continuing Education

This course is accredited by the Swiss Association of Pharmaceutical Professionals (SwAPP) and the Swiss Society for Pharmaceutical Medicine (SGPM).





INTRODUCTION TO CLINICAL RESEARCH

2 Modules

REGISTRATION FORM	REGISTRATION INFORMATION
First Name*:	Dates: Module 1 - 18 th November 2020 / Module 2 - 25 th November 2020
Last Name*:	Time: 09:20 42:00 CET (incl. breaks)
Company*:	Time: 08:30 – 12:00 CET (incl. breaks)
Job Title*:	Delivery method: Remotely via a digital platform. Delegates will receive login instructions within a week before the course starts.
Billing Address*: Telephone*:	Registration Fee: CHF 680.00 + 7.7% VAT 15 % reduction for SwAPP and SBA members Special fees for CTC employees Special rates available for PhD Students, please contact training@ctcresourcing.com
Telephone*: Email Address*:	Registration fees include: Course documentation. Upon
Background:	 Payment: After registration you will receive confirmation and an invoice with instructions on how to complete the bank transfer. Full prepayment is needed prior to the course start date.
Experience:	Cancellation Policy: • Cancellations received more than 14 days before course: 100% refund less bank charges & cancellation fee of CHF 50.00 • Cancellations received 7-14 days before the course: 50% refund • Cancellations received 7 days before the course: no refund
Are you a CTC employee? ☐ yes ☐ no	In case of no show, no refund will be made.
Are you a SwAPP member?	Please note that CTC reserves the right to cancel this course or to make minor alterations to the content and timing of the programme or to the identity of the speakers. In the unlikely event of cancellation, delegates will be offered a full refund. CTC will not, however, be held responsible for any related expense incurred by the participant. Transfer Policy Subject to agreement from CTC Clinical Trial Consulting AG (CTC) prior to the event, the registration may be transferred to another person from the same company or organization at no extra charge. Updated registration information will be required.
Do you wish to receive regular training updates?	Photography and Video Policy By attending this training, you give permission for pictures, images etc. of you, captured during the training through video, photo, and/or digital devices, to be used by CTC in promotional materials, publications, and website and waive any and all rights including but not limited to compensation or ownership. Privacy Policy At CTC, we're committed to protecting and respecting your privacy. To view our privacy policy, go to https://www.ctcresourcing.com/about-us/privacy- policy/
REGISTRATION: ◆ Email: training@ctcresourcing.com ◆ Internet: www.ctcresourcing.com ◆ Fax: +41 61 695 88 67 If you have NOT received an official confirmation 7 days	after registering, please call CTC at +41 61 695 88 66 or send
an email to training@ctcresourcing.com. THANK YOU!	