



WEBINAR INVITATION

WHO New Guidelines for Clinical Trials: Innovation and Best Practices



**3 December
2025**

REGISTER HERE



The registration form is available at this [link](#).

Please, fill it by 1st December 2025.

Accepted participants will receive a confirmation email.

The participation is free of charge.

GENERAL INFORMATION

The event will be held online via Microsoft Teams.

It is addressed to clinicians, biologists, principal investigators and researchers.

The official language is English.

ECM credits will not be provided.

Max n. of participants: 300.

ATTENDANCE CERTIFICATE

Upon request, participants who attend at least 75% of the event's duration online and complete the satisfaction survey will receive a certificate of participation.

AIMS & OBJECTIVES

The primary goal of this webinar is to disseminate and analyze the new WHO guidelines on clinical trials. It offers to researchers, physicians, regulators, and other key stakeholders a valuable opportunity for discussion, learning, and practical application of these standards.

In this light, the event aims to foster the implementation of the new guidelines within the European and International context, with a particular focus on best practices, ethics, and innovation in clinical research.

EVENT ORGANIZERS

SCIENTIFIC COORDINATOR

Maria Jose Ruiz Alvarez

ItaCRIN, Istituto Superiore di Sanità

SCIENTIFIC SECRETARIAT

Maria Buocervello

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Istituto Superiore di Sanità

ORGANIZING SECRETARIAT

Fabiola Giuliano

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CONTACT US

If you need more information or support, please write to the following email address: itacrin@iss.it

PRELIMINARY AGENDA

Moderator: *Maria José Ruiz Alvarez*

11:00 – 11:10 | Opening and Welcome

Welcome address by ItaCRIN

Elena Toschi

11:10 – 11:30 | Key Updates from the WHO Guidelines on Clinical Research

Presentation of the main developments and implications for implementation.

Rasmus Gjesing

11:30 – 13:25 | Roundtable Discussion

Challenges and Opportunities in Implementing the WHO Guidelines on Clinical Research Experts

Rasmus Gjesing, Ornella Gonzato, Carmine Iorio, Annalisa Landi, Claudia Louati, Federica Mantovani, Amélie Michon, Elena Petelos, Evelina Tacconelli

Discussion themes: Ethics and patient protection; Transparency and data management; Application of the guidelines in multicentre and international contexts; Practical examples from European clinical trials; Tools and strategies for integrating WHO guidelines into trial protocols and management; Interactive discussion with participant questions

13:25 – 13:30 | Conclusion and Next Steps

Summary of key discussion points

Acknowledgements and closing remarks

Maria José Ruiz Alvarez

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SPEAKERS & CHAIRS

Rasmus Gjesing

Regional Adviser, Access to Medicines and Health Products
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Carmine Iorio

Guest Researcher - Bioethic Unit
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Public Health Specialist and HTA expert
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