

Juristische Fakultät



## **EU Clinical Trial Regulation New Legal Framework for Multicenter Clinical Trials**

## Friday, December 16, 2016, 1 pm - 5 pm

## **WWZ Auditorium**

University of Basel, Law Faculty, Peter Merian-Weg 8, CH-4002 Basel

## **Program**:

12:30	Registration
13.00 – 13.05	Welcome Prof. Dr. DiplBiol. Herbert Zech, University Basel, Faculty of Law
13:05 – 13.15	Introduction Prof. Dr. Claudia Seitz, M.A., University Basel, Faculty of Law Dr. Alexander Meier, Novartis Pharma AG
13.15 – 13.45	<b>EU Clinical Trail Regulation and the Swiss Human Research Act</b> Prof. Dr. Claudia Seitz, M.A., University Basel, Faculty of Law
13.45 – 14.15	<b>EU Clinical Trial Regulation – A View from Legal and Regulatory</b> Dr. Alexander Meier, Novartis Pharma AG
14.15 – 14.45	Coffee Break
14.45 – 15.30	EU Clinical Trial Regulation – A View from the Regulator loana Ratescu, Legal Advisor, European Medicines Agency (EMA)
15.30 – 16.00	EU Clinical Trial Regulation – A View from the Industry Dr. Judith Creba, Executive Director EU Regulatory Strategy, Novartis Pharma AG
16.00 – 16.30	EU Clinical Trial Regulation – A View from Data Privacy Noemi Alonso Calvo, Data Privacy Officer, Sandoz International GmbH
16.30 – 17.00	Discussion
17.00	Apéro

No registration fee, please register by email to zlsr-ius@unibas.ch by December 12, 2016

Organized by: Dr. Alexander Meier, Novartis Pharma AG, Head Legal Global & Pharma Drug

Development, Basel

Prof. Dr. Claudia Seitz, M.A., Center for Life Sciences Law (CLSL), University Basel,

Faculty of Law

Apéro provided by Novartis Pharma AG