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| INTERNATIONAL GENERIC AND BIOSIMILAR MEDICINES ASSOCIATION | |
| 12.25 13.15 | OO Networking buffet lunch |
| 13.15 15.15 | Session 3 – International harmonisation of bioequivalence: current status and next steps (ICH M13A implementation and hot topics for M13C) Co-Chairs: Lei Zhang, FDA and Russ Rackley, Viatris |
| | M13 guideline series Lei Zhang, FDA Impact of M13A from a European perspective |
| | Jan Welink, MEB/EMA Challenges and opportunities of M13A guideline for generic industry Irmela Gabriel, Teva M13A implementation plan in the EU |
| | Kevin Blake, EMA Statistical challenges and opportunities in ICH M13C Helmut Schütz, University of Vienna M13C an opportunity to harmonise bioequivalence requirements for (NTI) drugs |
| | Paulo Paixão, University of Lisbon Faculty of Pharmacy Panel discussion |
| 15.15 15.30 | O Networking coffee break |
| 15.30 16.30 | Rapid session 4 – Compliance and oversight in BE |
| | Chair: Janja Luksa, Sandoz Regulatory expectations for data integrity Peter Twomey, EMA (invited) FDA's perspective Nilufer Tampal, FDA Data integrity in BE: sponsor's perspective Janja Luksa, Sandoz Panel discussion |
| 16.30 16.45 | Closure of the conference - Susana Almeida, IGBA |



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