



Biomedical Alliance in Europe

Training day: Understanding & impacting the EU Health Agenda

Learn how EU policies & regulations impact your daily work as a clinician or researcher and how you can contribute to better decision-making

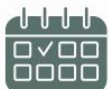


One-day training session for BioMed Alliance members, with key speakers from the EU health policy field focussing on:

- How does the EU work?
- Impact of EU policies on health systems
- How to influence the EU health agenda?
- How can experts contribute?



Date



27 November

Location



**BioMed Alliance
office, Brussels**

Register now



Funded by the European Union. Views and opinions expressed are however those of the author(s) only and do not necessarily reflect those of the European Union or HaDEA. Neither the European Union nor the granting authority can be held responsible for them. This workshop is supported by the EU4Health Operating Grant EXHALE, funded under grant agreement nr. 101176496



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Draft Programme

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- 10.00-10.10 **Welcome & introduction**
- Alan Fraser, Outgoing chair Regulatory Affairs Committee
- 10.10-12.00 **Opening discussion: involving clinical experts in EU health policy**
Chair: Tom Melvin
- European Health Union & the role of the Commission in working with experts to advance health
 - o Rainer Becker, Director for Medical Products & Innovation, European Commission
 - Involving experts in evidence based regulatory affairs
 - o Ivana Silva, Healthcare Professionals and Learned Societies Liaison, European Medicines Agency (EMA)
 - Do policy makers need medical specialists at the table?
 - o Elizabeth Macintyre, BioMed Alliance President
- 12.00-13.00 **Expert Perspectives on Health Policy: From clinical practice, to advocacy and regulation**
Chair: Elizabeth Macintyre
- How to navigate an increasingly complex health policy field
 - o Marieke Meijer (BioMed Alliance)
 - The view from a national regulator turned regulatory science academic – what is effective?
 - o Tom Melvin (BioMed Alliance & Trinity College)
 - Representing the academic and clinical community at EU institutions
 - o Alan Fraser (BioMed Alliance)
- 13.00-14.00 **Lunch**
- 14.00-15.00 **Implementing the EU regulation on health technology assessment**
Bela Dajka (European Commission) and Robin Doeswijk (EHA)
- Introduction to the legislation, how it impacts clinical care and how experts can contribute to the system
- 15.00-15.30 **Coffee break**
- 15.30-17.00 **Medical devices & diagnostics: IVDR/MDR**
Paul Piscoi & Olga Tkachenko (European Commission), Tom Melvin
- Introduction to the legislations, how they impact clinical care and how experts can contribute to the system
- 17.00-18.00 **Digital health: The European Health Data Space (EHDS)**
Jerome de Barros (European Commission)
- Introduction to the legislation, how it impacts clinical care and how experts can contribute to the system
- 18.00-19.00 **Conclusion & Networking Cocktail**
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