ISMPP EU, 23–24 Jan, 2018 Day 1 Summary

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Disclosures

Co-owner/director of Aspire Scientific Ltd

Co-owner/director of The Publication Plan Ltd

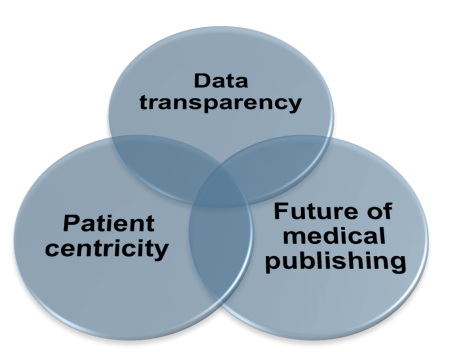


23-24 January 2018 • London, UK

2018 EUROPEAN MEETING OF ISMPP











ISMPP EU day 1 agenda

10.00	Welcome (Fiona Plunkett)
10.10	Year in review: a look back at 2017 (Jackie Marchington)
10.30	Medical publications and data disclosure – ensuring compliance, efficiency and effectiveness for publication and data disclosure teams (Moderator: Russell Traynor)
11.30	Speed research: part 1
13.00	Roundtable sessions
15.00	Panel discussion: Time to embrace change in medical publishing? (Moderator: Richard Smith)
16.30	Debate: Will we be replaced by robots? (Moderator: Martin Delahunty)
17.10	ISMPP & ISMPP CMPP TM update: the expanding role of ISMPP (Juliana Clark, Susan Scott)

Year in review: a look back at 2017 – Jackie Marchington



Open Pharma

• Encouraging pharmaceutical industry to participate in discussions on the future of publishing <u>and</u> drive change



New guidelines for abstracts/presentations

- Good Practice for Conference Abstracts and Presentations (GP-CAP)
- Currently available at PeerJ Preprints*



ICMJE Recommendations update

Updates on predatory journals, ethics and data sharing



Fake news?

- Predatory medical journals hit the New York Times
- TrialsTracker report that 45% of completed trials not reported on ClinicalTrials.gov or PubMed

Medical publications and data disclosure – ensuring compliance, efficiency and effectiveness for publication and data disclosure teams

Moderator: Russell Traynor

Panellists: Julie Ford, Karen Mittleman & Santosh Mysore



Challenges

- Ever-evolving regulatory and cultural environment
- Co-ordination of multiple local/international trials
- Maintaining anonymity of patient-level data
- Internal resource and communication



Solutions

- Robust internal procedures
- Training
- Personal responsibility/accountability
- Information management systems

Medical publications and data disclosure – ensuring compliance, efficiency and effectiveness for publication and data disclosure teams

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Integration across teams

- Collaboration between clinical trial teams responsible for 'non-interpretative' disclosure and publications teams managing 'interpretative' disclosure
- Moving from publication planning to integrative data disclosure planning



Educational Series on Data/Financial Transparency

- Launched by ISMPP Global Transparency Committee
- Developed to raise awareness of the numerous guidance documents and policies on these topics
- Available to ISMPP members

Time to embrace change in medical publishing?

Moderator: Richard Smith

Panellists: Chris Rains, Chris Winchester, Lise Baltzer, Martin Delahunty,

Rebecca Lawrence, Robert Kiley, Santosh Mysore, Theodora Bloom



Open Pharma

- Multi-stakeholder group
- Recognising the role that pharma has to play in helping to improve the publishing of science
- Helping to drive faster and increasingly transparent publication of industry-sponsored research
- Four workstreams established, including one on open access and one on pre-prints/post-publication peer review

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Open access

- Open access isn't only 'free to read'
- True open access also allows reuse
- Publishers often unwilling to issue suitable (CC-BY) licences for industry-sponsored research
- Multiple copyright licenses
- Can open access publication be mandated by pharma?

Time to embrace change in medical publishing?

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Pre-prints

- Allow open, pre-published peer review and may speed up dissemination of data
- MedArXiv pre-print server for medicine/heath sciences due to launch soon
- Journal attitudes to preprints?



Post-publication peer-review

- Another method of speeding up dissemination of results, including less impactful data
- Examples: F1000Research, Wellcome Open Research
- Articles indexed once passed peer review

Will we be replaced by robots?

Moderator: Martin Delahunty

Speakers: Russell Traynor, Harry Politis



Artificial intelligence (AI) is already here

- AI-driven drug discovery (e.g. collaborations between GlaxoSmithKline and Exscientia and Insilico Medicine)
- Numerous Al-focused start-ups focusing on the patient healthcare system interaction



AI in publishing

- Offers increased operational speed and efficiency
- Identifying peer reviewers (e.g. SubSift)
- Detecting plagiarism

Will we be replaced by robots?

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The future?

- Impact on clinicians
 - Augment the work of clinicians, at least initially
 - Some specialists (e.g. diagnostics) may be replaced
- Impact on content and delivery of education
 - Al can learn from user interaction
 - Intelligent e-learning programmes
- Impact on publications management
 - Moving from systems supporting, to those executing,
 SOPs
 - Publication professionals will not be replaced just yet

ISMPP and ISMPP CMPP™ update: the expanding role of ISMPP – Juliana Clark and Susan Scott



ISMPP

- Growing membership through education and collaboration
- Efforts include the AMWA-EMWA-ISMPP Joint Position
 Statement and MPIP Transparency Matters initiative
- Encourage member-driven research



ISMPP CMPPTM

- Changes to the four knowledge domains for the exam (developing and implementing a publication plan, fostering ethical/compliant behavior, monitoring trends)
- Next exam windows: March and September 2018
- Recertification now on a 3-year cycle
- Self-study recertification credits soon available

Thank you

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For written reports on the ISMPP EU meeting see:

<u>ThePublicationPlan.com</u>

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