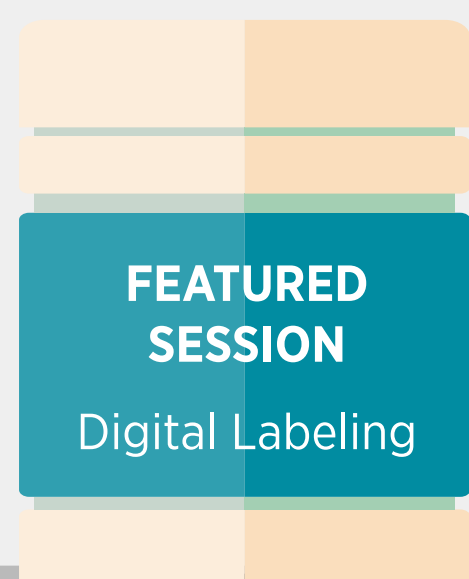
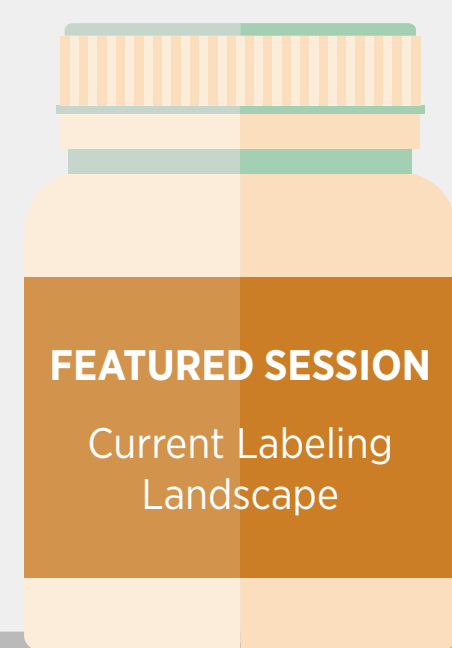




Hot Topics and Featured Sessions at DIA's Global Labeling Conference

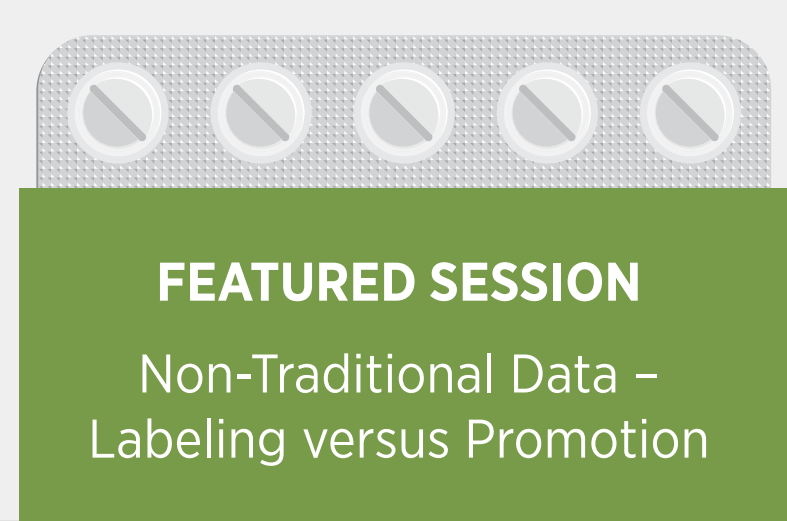
DIA's *Global Labeling Conference* is the premier forum for regulators and industry to update their knowledge of local and global labeling-related policies and to examine the impact of changes on regulatory compliance.

- Key labeling requirements in major markets
- Implications of recent regulations on labeling texts in major markets
- Evolving landscape of labeling in the future world of personalized medicine



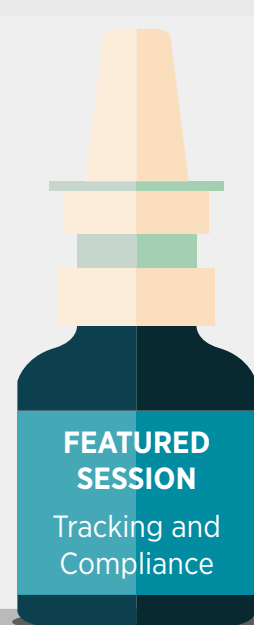
Explore the benefits, status, and current developments surrounding digital and electronic labeling across regions

Changes in regulatory strategy in areas beyond US, EU and Japan including the evolution of drug prescribing information regulations in Latin America and the Caribbean, and the changes in the MHRA and the UK post-Brexit



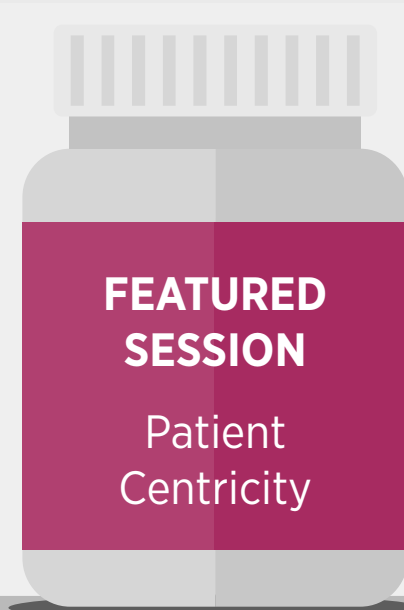
- Explore the different types of non-traditional data and the pros/cons and possibility for inclusion in labeling or promotional materials
- Current guidelines and practices of inclusion of the non-traditional data in promotional materials

Explore labeling throughout drug development from a global and local perspective and identify a strategy for creating and advocating for resources in your department



End-to-end tracking on a global scale, contemporary approaches for labeling operations governance groups, and opportunities to apply human error analysis techniques to labeling development

- Patient-centric drug labeling for all stakeholders
- Value of health literacy in effective patient-centered labeling



- Current guidance and regulations for combination product and medical device labeling development from conceptualization to agency approval,
- Industry and regulator perspectives

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