

Good Regulatory Review Practices (GRRP) Working Group Update

Working Group Chair(s):

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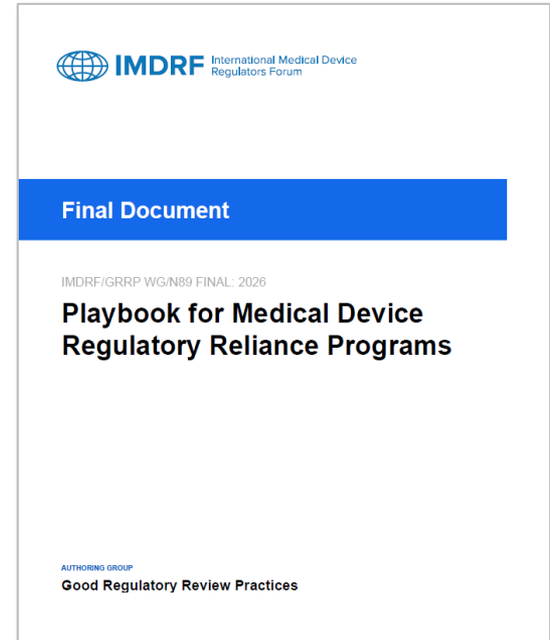


Reliance Playbook published in February 2026

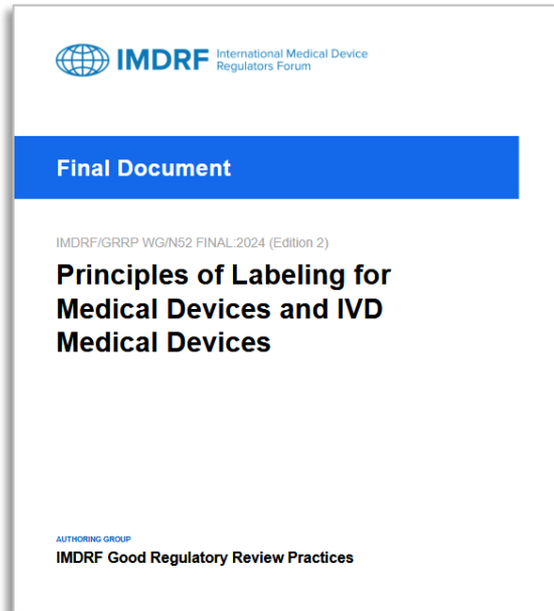
- General strategies and specific considerations for developing and implementing regulatory reliance programs within and across regulatory jurisdictions
 - GRRP WG appreciates the valuable feedback on the draft document (over 400 comments received)
- Real-world experience with reliance programs, including the discussion during the March 9 IMDRF Industry Workshop, will inform future IMDRF efforts to promote and enhance regulatory reliance activities



N89



Upcoming Work: Updating Principles of Labeling for Medical Devices and IVD Medical Device (N52)



- Work item approved in September 2025 to update labeling document to include additional guidance on electronic labeling practices and terminology
 - Clarify appropriate use cases for e-labeling within the IMDRF labeling framework
 - Promote timely availability of updated labeling
 - Maintain equitable access for users and patients
- Expected completion date of mid-2027

