

## Ongoing guidance development and deliverables of MDCG Subgroups – May 2022\*

*\*This is not an exhaustive list of ongoing work performed by MDCG Subgroups*

Scope	Group Deliverables	Consult prior to MDCG**	Planned MDCG Endorsement	Additional Comments
<i>** Stakeholders are observers in 13 MDCG subgroups and are consulted on a regular basis; further to that other MDCG subgroups are consulted as indicated</i>				
<b>1. Notified Bodies Oversight (NBO)<sup>1</sup></b>				
MDR + IVDR	<i>Q&amp;A on requirements notified bodies – update of MDCG 2019-6</i>	Notified bodies	N/A	Permanent NBO Work Item
MDR+IVDR	<i>Updates of guidance documents and templates on the designation and re-assessment process</i>	Notified bodies	2022	Q2 2022: NBOG PBG 2017-1 Revision 1 to include re-assessment process
MDR + IVDR	<i>Updates of guidance documents and templates on qualification and authorisation of personnel</i>	Notified bodies	TBD	Work starting in 2022
MDR + IVDR	<i>Template List of standard fees</i>	Notified bodies and MDCG Stakeholders	2022	
IVDR	<i>Guidance on appropriate surveillance according to Article 110(3)</i>	IVD WG and MDCG Stakeholders	2022	
MDR	<i>Notified Body Technical Documentation Assessment Report</i>	Notified bodies and relevant MDCG Subgroups	2022	
MDR	<i>Revision of MDCG 2020-3 Guidance on significant changes regarding the transitional</i>	MDCG Stakeholders	2022	

<sup>1</sup> Stakeholders are not part of this group as it covers requirements set out by designating authorities specifically for notified bodies; stakeholders are consulted on mature and final drafts.

	<i>provision under Article 120 of the MDR with regard to devices covered by certificates according to MDD or AIMDD</i>	and relevant MDCG subgroups		
MDR + IVDR	<i>Position on the concept of “Hybrid audits” carried out by notified bodies, including definition</i>	Notified bodies	2022	
<b>2. Standards</b>				
MDR + IVDR	<i>Updates of guidance document MDCG 2021-5 on standardisation for medical devices</i>	NBO, IVD	Q3 2022	
MDR + IVDR	<i>“Cookbook” for harmonised standards</i>		Q2 2022	Proposed by CLC TC 62, not intended to become a MDCG-endorsed document
<b>3. Clinical Investigations and Evaluation (CIE)</b>				
MDR	<i>Clinical Investigation Report Summary Template</i>		2022	
<b>4. Post-Market Surveillance and Vigilance (PMSV)</b>				
MDR + IVDR	<i>Guidance on Periodic Safety Update Report requirements</i>		Q2 2022	PSUR for MDR to be later adapted for IVDR
MDR + IVDR	<i>Guidance on Post-Market Surveillance requirements</i>	MS	Q3 2022	Work to be coordinated with the Market Surveillance WG
MDR + IVDR	<i>Q&amp;A document on Vigilance terms and concepts Q&amp;A document on Art 87 to 90 on Vigilance requirements</i>		Q3 2022 Q3 2022	Task force work has been divided in 2 groups respectively on definitions and on Art 86-90 interpretation
MDR + IVDR	<i>Development of harmonised reporting forms for incidents</i>		Q2 2022	Several Task Forces on-going on the updating of the MIR form and the Trend report form

5. Market Surveillance (MS) <sup>2</sup>				
MDR + IVDR	<i>Authorised Representatives</i>	IVD	2022	
MDR + IVDR	<i>In-house devices</i>	IVD	2022	
MDR + IVDR	<i>Update MDCG 2021-27 Q&amp;A on Importers &amp; Distributors</i>	IVD	Q.4 2022	
MDR + IVDR	<i>Update MDCG 2021-26 Q&amp;A on repackaging &amp; relabelling activities under Article 16</i>	IVD	Q.4 2022	
MDR + IVDR	<i>Update MDCG 2019-7 of PRRC Guidance</i>	TBD	Q.4 2022	
6. Borderline & Classification (B&C)				
7. New Technologies				
MDR + IVDR	<i>Legal status of app providers</i>		Q4 2022	
MDR + IVDR	<i>Guidance on MDSW - Hardware combination systems</i>	B&C	Q2 2022	
8. Eudamed				
IVDR	<i>Guidance on harmonised administrative practices and alternative technical solutions until EUDAMED is fully functional (IVDR)</i>	IVD	2022	IVDR Implementation

<sup>2</sup> Stakeholders are not part of this group as it covers requirements set out by competent authorities; stakeholders are consulted on mature and final drafts.

9. Unique Device Identification (UDI)				
10. International Matters				
	N/A			
11. <i>In Vitro</i> Diagnostic Medical Devices (IVD)				
IVDR	<i>In-house devices</i>	MS	2022	Joint with Market Surveillance MDCG sub-group, draft in preparation
IVDR	<i>Analysis of IVDR in context of hypothetical scenarios of an urgent response to a health crisis</i>	N/A	2022	In progress
IVDR	<i>Performance study application/notification form</i>	CIE	2022	Template in development
IVDR	<i>Minor revision of MDCG 2021-22 – Clarification on “first certification for that type of device” and corresponding procedures to be followed by notified bodies</i>	N/A	2022	Addition of notes, based on experience collected so far
IVDR	<i>Minor revision of MDCG 2020-16 – Classification of IVDs</i>	B&C	2022	Addition of specific points

12. Nomenclature				
MDR + IVDR	<i>Procedures for the annual and ad-hoc updates of the EMDN</i>	N/A	Q2-Q3 2022	
MDR + IVDR	<i>FAQ on EMDN</i>	N/A	Q3 2022	
MDR + IVDR	<i>Mapping EMDN-GMDN package</i>	N/A	N/A	The outcome of this exercise is highly dependent on level of cooperation ensured by GMDN.
13. Annex XVI				
MDR	<i>Guidance document on the use of equivalence criteria for Annex XVI products</i>	CIE, NBO	Q4 2022	
MDR	<i>Guidance document on the classification of Annex XVI products</i>	B&C	Q4 2022	