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	
** Stakeho	lders are observers in 13 MDCG subgroups and are consulted o	n a regular basis; furth	er to that other MD	CG subgroups are consulted as indicated	
1. Not	1. Notified Bodies Oversight (NBO) ¹				
MDR + IVDR	Q&A on requirements notified bodies – update of MDCG 2019-6	Notified bodies	N/A	Permanent NBO Work Item	
MDR+IVDR	Updates of guidance documents and templates on the designation and re-assessment process	Notified bodies	2022	Q2 2022: NBOG PBG 2017-1 Revision 1 to include re-assessment process	
MDR + IVDR	Updates of guidance documents and templates on qualification and authorisation of personnel	Notified bodies	TBD	Work starting in 2022	
MDR + IVDR	Template List of standard fees	Notified bodies and MDCG Stakeholders	2022		
IVDR	Guidance on appropriate surveillance according to Article 110(3)	IVD WG and MDCG Stakeholders	2022		
MDR	Notified Body Technical Documentation Assessment Report	Notified bodies and relevant MDCG Subgroups	2022		
MDR	Revision of MDCG 2020-3 Guidance on significant changes regarding the transitional	MDCG Stakeholders	2022		

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.

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

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

² 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.	Jnique Device Identification (UDI)					
10.	10. International Matters					
	N/A					
11.	In Vitro Diagnostic Medical Devices (IVD)					
IVDR	In-house devices	MS	2022	Joint with Market Surveillance MDCG sub-group, draft in preparation		
IVDR	Analysis of IVDR in context of hypothetical scenarios of an urgent response to a health crisis	N/A	2022	In progress		
IVDR	Performance study application/notification form	CIE	2022	Template in development		
IVDR	Minor revision of MDCG 2021-22 – Clarification on "first certification for that type of device" and corresponding procedures to be followed by notified bodies	N/A	2022	Addition of notes, based on experience collected so far		
IVDR	Minor revision of MDCG 2020-16 – Classification of IVDs	B&C	2022	Addition of specific points		

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