Team-NB Position Paper list (update July 17th 2025)

Date	Name		Version	Status
17-07-25	Team-NB-PositionPaper-Ophan-IVDR-Medical- Devices-V1	The paper is addressing Orphan under the IVDR	V1	Active
27-06-25	Team-NB-PositionPaper-Software-Qualification under-the-IVDR-V2	The paper is addressing Software qualification under the IVDR.	V2	Active
09-04-25	Team-NB-PositionPaper-IVDR-Certification- Process-Consensus-Document-V1	The paper is describing in detail pre-application and application and Post application phases processes through which manufacturers may apply to Notified Bodies (NBs) for the certification of medical devices under the regulation (EU) 2017/746 (IVDR).	V1	Active
09-04-25	Team-NB-PositionPaper-EU-AI-Act-V2	The designation of notified bodies under the upcoming Artificial Intelligence Act	V2	Active
09-04-25	Team-NB-PositionPaper-BPG-TechnicalDocEU- MDR-2017-745-V3	Best Practice Guidance for the Submission of Technical Documentation under Annex II and III of Medical Device Regulation (EU) 2017/745	V3	Active
18-12-24	Team-NB-PositionPaper-MDR-Certification- Process-Consensus-Document	The paper is addressing to describe in detail the pre- application, application processes through which manufacturers may apply to Notified Bodies (NBs) for the certification of medical devices under the regulation (EU) 2017/745 (MDR).	V1	Active
10-12-24	Joint-Team-NB-IG-NB-PositionPaper-Al-in-MD- Questionnaire-V1.1	This questionnaire was prepared in accordance with MDGG 2022-14 requests.	V1.1	Active
11-09-24	Team-NB-PositionPaper-IVD-Transfer- Agreement	The paper is proposing a form for a contract between the 3 involved parties.	V1	Active
10-07-24	Team-NB-IVDConfirmationLetterTemplate-V2	The paper is proposing an updated version of the IVD Confirmation Letter template specific to IVD on the basis of the MD one.	V2	Active
02-07-24	Team-NB-PositionPaper- TransferAgreement_v02	Transfer Agreement for Surveillance of Legacy Devices specifying the terms of the transfer of the appropriate surveillance according to Regulation (EU) 2017/7451	V2	Active

19-02-24	Team-NB-PositionPaper-Classification-of-SARS-	The paper is addressing the reclassification of Covid-19	V1	Active
	Cov2	devices.		
15-12-23	Team-NB-PositionPaper-Lifetime-Medical-	MD Lifetime is addressing Lifetime with the objective to	V1	Active
	Device	consider different device types of a medical device		
		lifetime in terms of safe and effective use.		
10-08-23	Team-NB-PositionPaper-	Views on the amended timelines with regards to benefits	V1	Active
	MDRTransitionTimelines-NotifiedBodyCapacity	for the European patients and as the continuity in the		
	V1	availability of essential medical devices in the European		
		market		
10-07-23	Team-NB-PositionPaper-NB-	Template for notified body confirmation letter of the	V2	Active
	ConfirmationLetterEU2023-607-V2	status of a formal application in the framework of		
		Regulation EU 2023/607		
16-05-23	Team-NB PositionPaper HybridAudits V2	Notified bodies' paper on the application of hybrid audits	V2	Active
		to quality management system assessments under		
		MDR/IVDR		
25-02-23	Team-NB PositionPaper-BPG-technicalDoc EU-	Best Practice Guidance for the Submission of Technical	V1	Active
	IVDR-2017-746 V1	Documentation under Annex II and III of In Vitro		
		Diagnostic Medical Devices Regulation		
11-11-22	Team-NB PositionPaper Certificates under	Team NB Position in Response to MDCG 2022-14 Item	V1	Active
	conditions-V1	Number 17 – 'Certificates under Conditions'		
05-10-22	Team-NB-PositionPaper-VoluntaryTransfer-	Transfer Agreement specifying the terms of voluntary	V1	Active
	Agreement-V1-20221005	change of notified body under Regulation (EU) 2017/745		
		or Regulation (EU) 2017/746		
05-10-22	Team-NB PositionPaper	Class D measures in the absence of EU Reference	V1	Active
	ConformityAssessmentClassD V1	Laboratories- Points to consider for Notified Body		
		approach		
05-10-22	Team-NB PositionPaper CyberSecurity-V1	Recommendations to cybersecurity harmonised	V1	Active
		approach		
05-10-22	Team-NB PositionPaper Off Label Use-V1	Data generated from 'Off-Label' Use of a device under	V1	Active
		the EU Medical Device Regulation 2017/745.		
03-10-22	Team-NB-PositionPaper-Leveraging-evidence-	Leveraging directive conformity assessments to establish	V1	worked on
	from-Directives-DRAFT	compliance with the MDR requirements		

14-07-22	Team-NB-PositionPaper ConfAssessment-	Notified body approach for the Technical Documentation	V1	Active
	Multiplex IVD V1	assessment approach of multiplex in-vitro diagnostic		
		devices		
28-01-22	Team-NB-PositionPaper-	Team-NB Notified Bodies recommendations	V1	Active
	ModificationsSamplingPlan-V1	on the handling of modifications to the device sampling		
		plans		
01-12-21	Team-NB-PositionPaper-on-MDR_IVDR-	Notified Body position paper on MDR/IVDR	V3	Active
	Implementation-V3	Implementation		
09-11-21	Team-NB-PositionPaper-IVDR-Significant	Significant changes according to Article 110 (3) of	V1	Active
	changes-V1	Regulation EU 2017/746		
06-10-21	Team-NB-PositionPaper-Article117-NB-Opinion	Proposal for a Notified Body Opinion Template	V1	Active
	Template-V1			
06-10-21	Team-NB-PositionPaper-Artificial-Intelligence-	European Artificial Intelligence Regulation	V1	Active
	V1			
20-07-21	Team-NB-PositionPaper-ImplantCard-	Team-NB Position Paper on a risk-based approach for	V1	Active
	20210720	medical devices exempted from an implant card and		
		information to be supplied to the patient with an		
		implanted device per Article 18.3		
19-05-21	Team-NB-PositionPaper-ClassD-20210519-V4.4	Team-NB Notified Bodies considerations on conformity	V4.4	Active
		assessment for class D devices		
21-12-20	Team-NB PositionPaper Art117 Substantial	Position paper for the interpretation of device related	V1	Active
	Changes DrugDeviceCombination	changes in relation to a Notified Body Opinion as		
		required under Article 117 of Medical Device Regulation		
		(EU)2017/745		
18-11-20	Team-NB-PositionPaper-RemoteAudits-V1-	Position paper	V1	Active
	20201118	Remote Audit Survey : Analysis		
22-07-20	Team-NB Position Paper Technical Cooperation	Position Paper on the requirements for the EU	V1	Active
	Program TCP III -V1	MDR/IVDR Notified Body Partners under the Technical		
		Cooperation Program on Exchange of Medical Device		
		Quality Management System Regulation and ISO 13485		
		Audit Reports (TCP III)		

01-04-20	Team-NB Position-Paper Documentation Requirements Article117 V1	Position Paper on Documentation Requirements for Drug Device Combination Products falling in the Scope of Article 117 of MDR 2017/745.	V1	Active
11-03-20	Team-NB-Position paper on Dental Implants- 20200311-V1	Position Paper on Applicability of exemption rule to endosseous dental implants and dental implant abutments	V1	under revision

Legend Status: active / worked on /voting process /under revision