## Team-NB Position Paper list (update December 18th 2023)

Date	Name	Scope/description	Version	Status
19-02-24	Team-NB-PositionPaper-Classification-of-SARS-	The paper is addressing the reclassification of Covid-19	V1	Active
	Cov2-20240202	devices. It is proposed that the Covid-19 devices are		
		initially reclassified as C (from class D); once more data is		
		available on the post-pandemic phase, if no new issue		
		concerning patient health is identified, a further		
		reclassification to class B should be considered.		
15-12-23	Team-NB-PositionPaper-Lifetime-Medical-	MD Lifetime is addressing Lifetime with the objective to	V1	Active
	Device-20231127	consider different device types of a medical device		
		lifetime in terms of safe and effective use.		
11-08-23	Team-NB-PositionPaper-LegacyMD-	Transfer Agrement for Surveillance of Legacy Devices	V1	Active
	Surveillance-Transfer-Agreement-V1-20230811	specifying the terms of the transfer of the appropriate		
		surveillance according to Regulation (EU) 2017/7451		
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		
19-04-23	Team-NB-PositionPaper BPG TechnicalDoc EU-	Best Practice Guidance for the Submission of Technical	V2	under
	MDR-2017-745-V2	Documentation under Annex II and III of Medical Device		revision
		Regulation (EU) 2017/745		
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		

16-12-22	Team-NB PositionPaper AI Designation-V1	The designation of notified bodies under the upcoming	V1	Active
		Artificial Intelligence Act		
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			

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		
Team-NB-PositionPaper-ClassD-20210519-V4.4	Team-NB Notified Bodies considerations on conformity	V4.4	Active
	assessment for class D devices		
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		
Team-NB-PositionPaper-RemoteAudits-V1-	Position paper	V1	Active
20201118	Remote Audit Survey : Analysis		
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)		
Team-NB Position-Paper Documentation	Position Paper on Documentation Requirements for Drug	V1	Active
Requirements Article117 V1	Device Combination Products falling in the Scope of		
	Article 117 of MDR 2017/745.		
Team-NB-Position paper on Dental Implants-	Position Paper on Applicability of exemption rule to	V1	under
20200311-V1	endosseous dental implants and dental implant abutments		revision
	Team-NB-PositionPaper-ClassD-20210519-V4.4  Team-NB PositionPaper Art117 Substantial Changes DrugDeviceCombination  Team-NB-PositionPaper-RemoteAudits-V1-20201118  Team-NB Position Paper Technical Cooperation Program TCP III -V1  Team-NB Position-Paper Documentation Requirements Article117 V1  Team-NB-Position paper on Dental Implants-	medical devices exempted from an implant card and information to be supplied to the patient with an implanted device per Article 18.3  Team-NB-PositionPaper-ClassD-20210519-V4.4  Team-NB Notified Bodies considerations on conformity assessment for class D devices  Team-NB PositionPaper Art117 Substantial Changes DrugDeviceCombination Changes in relation to a Notified Body Opinion as required under Article 117 of Medical Device Regulation (EU)2017/745  Team-NB-PositionPaper-RemoteAudits-V1- Position paper Remote Audit Survey : Analysis  Team-NB Position Paper Technical Cooperation Program TCP III -V1  Position Paper on the requirements for the EU 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)  Team-NB Position-Paper Documentation Requirements for Drug Device Combination Products falling in the Scope of Article 117 of MDR 2017/745.  Team-NB-Position paper on Dental Implants- Position Paper on Applicability of exemption rule to endosseous dental implants and dental implant	medical devices exempted from an implant card and information to be supplied to the patient with an implanted device per Article 18.3  Team-NB-PositionPaper-ClassD-20210519-V4.4 Team-NB Notified Bodies considerations on conformity assessment for class D devices  Team-NB PositionPaper Art117 Substantial Changes DrugDeviceCombination Position paper for the interpretation of device related changes in relation to a Notified Body Opinion as required under Article 117 of Medical Device Regulation (EU)2017/745  Team-NB-PositionPaper-RemoteAudits-V1- Position paper Remote Audit Survey: Analysis  Team-NB Position Paper Technical Cooperation Position Paper on the requirements for the EU 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)  Team-NB Position-Paper Documentation Requirements for Drug Device Combination Products falling in the Scope of Article 117 of MDR 2017/745.  Team-NB-Position paper on Dental Implants- Position Paper on Applicability of exemption rule to endosseous dental implants and dental implant

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