

# **Guidance Document:**

Clinical Evaluation Pathways MDR 2017/745



**Note:** Applicable devices given for each clinical evaluation route are listed as per the MDR and MDCG 2020-6. There may be other classes of devices not listed that can be applied to **some** of the clinical evaluation routes. The onus is on the manufacturer to review the MDR and the guidance documents and choose which route can be correctly applied to their device type. NSAI will review in accordance with the requirements of the MDR and MDCG guidance documents.

Clinical Evaluation Route options	Requirement	Applicable devices
Article 61(3)	a) Equivalence route	
	b) Clinical Investigations	All devices -Non-Legacy or some legacy devices that do not
	AND	have sufficient clinical evidence, brand new devices under MDR
	c) Alternative treatment options (cannot claim only option c)	
Article 61(4)	Exception for Implantable and class III devices who do not want to perform a clinical investigation:	
	-Manufacturer has made modifications to a device, which they have already marketed (under the directives or regulation)	
	-Can claim equivalence to a device marketed by the same Manufacturer	
	-Notified Body agrees with equivalence claim	Implantable and class III devices
	-Clinical evaluation of the marketed device is sufficient to demonstrate conformity to the GSPRs (CER of the marketed MDR compliant)	
	-Manufacturers must perform a PMCF study and show NSAI the plan (which should include a study) to demonstrate safety and performance of the device to be CE marked	



Article 61(5)	Exception for Implantable and class III devices who do not want to perform a clinical investigation:  -Manufacturers can claim equivalence to a different device that they don't manufacture themselves, however it must be CE- Marked under the MDR  - Provide a contract in place that explicitly allows the Manufacturers of the 2nd device full access to the technical documentation of the equivalent device on an ongoing basis  - The original clinical evaluation has to be performed in accordance with the requirements of the MDR (CER of the equivalent device must be MDR compliant)  - Manufacturer needs to provide clear evidence of this to NSAI	Implantable and class III devices
Article 61(6a)	Exception for manufacturers of legacy implantable and class III devices who do not want to perform a clinical investigation:  -Need to base the clinical evaluation on sufficient clinical data (as per MDCG 2020-6)  -Compliant to the relevant product specific common specification (CS) where such a CS is available (In the absence of CS, Manufacturers will need to prove sufficient clinical evidence)  Note: If a Manufacturers claims article 61 (6a & 6b) and no CS exists at the time of CE marking, and the relevant CS becomes available or released post CE marking, the manufacturer must update their technical documentation to comply with the relevant common specifications or run the risk of losing the CE mark	Legacy Implantable and class III devices



Article 61(6b)	If your device is a suture, staple, dental filling, dental braces, tooth crowns, screws, wedges, plates, wires, pins, clips, connectors,  -Manufacturers must base their clinical evaluation on sufficient clinical data (as per MDCG 2020-6)  -Manufacturers must be compliant with the relevant CS  -Compliant to the relevant product specific CS where such a CS is available (In the absence of CS, Manufacturers will need to prove sufficient clinical evidence)	WET devices
Article 61(9)	- the requirement to demonstrate a clinical benefit in accordance with this Chapter and Annexes XIV and XV shall be understood as a requirement to demonstrate the performance of the device.  -Clinical evaluations of those products shall be based on relevant data concerning safety, including data from post-market surveillance, PMCF, and, where applicable, specific clinical investigation.  -Clinical investigations shall be performed for those products unless reliance on existing clinical data from an analogous medical device is duly justified.  NOTE: As per the regulation (Article 61(9)), a manufacturer may either perform a clinical investigation for these Annex XVI devices or claim reliance on an analogous medical device.	Annex XVI devices



Article 61(10)	Where the demonstration of conformity with general safety and performance requirements based on clinical data is not deemed appropriate, the Manufacturers shall provide:  -Adequate justification which is based on the results of the manufacturer's risk management and on consideration of the specifics of the interaction between the device and the human body, the clinical performance intended and the claims of the manufacturer.  -The manufacturer shall duly substantiate in the technical documentation referred to in Annex II why it considers a demonstration of conformity with general safety and performance requirements that is based on the results of non-clinical testing methods alone, including performance evaluation, bench testing and pre-clinical evaluation, to be adequate.  - This will be considered only for low-risk devices, with no clinical benefit,	Only for low-risk devices where there is no clinical benefit. Not for class III & Implantable devices
	hence the device does not have a positive impact on the health of an individual, expressed in terms of a meaningful, measurable, patient-relevant clinical outcome(s), including outcome(s) related to diagnosis, or a positive impact on patient management or public health, Examples of devices that may be considered under this article are a lab fridge, a lab scale for weighing or measuring blood products, etc.	
MDCG 2020-6 Section 1.2	Legacy devices claiming WET must fulfil the following criteria below, by providing detailed rationale why the device fulfils these criteria and must provide supporting documents to justify the rationale given for each criteria- The common features of the devices which are well-established technologies are that they all have:  Relatively simple, common and stable designs with little evolution Their generic device group has well-known safety and has not been associated with safety issues in the past	Legacy devices claiming WET



- Well-known clinical performance characteristics and their generic device group are standard of care devices where there is little evolution in indications and the state of the art
- A long history on the market.

**NOTE 2:** A manufacturer that claims that a device qualifies as a WET device must specify what level of evidence has been provided based on MDCG 2020-6, appendix III table.

Reliance solely on complaints and vigilance is not sufficient.



## Important Information To Be Considered Prior To Submission Of Application

As per MDCG 2020-6, legacy devices which have been placed on the market have been subjected to conformity assessment and therefore are presumed to have been supported by clinical data. Post market clinical data together with the clinical data generated for the conformity assessment under the MDD/AIMDD will be the basis of the clinical evaluation process for legacy devices under the MDR, hence manufacturer's must state what clinical evaluation route (equivalence and/or clinical investigation) was used during the initial conformity assessment.

For legacy devices, please state what clinical evaluation route was used during your initial conformity assessment (when the device was first CE marked): Equivalence, clinical investigation, or both equivalence and clinical investigation.

#### **Applicable devices**

All Legacy devices.

If the clinical evaluation route during the initial conformity assessment (when the device was first CE marked) was based on equivalence, and you have not presented an equivalent device/argument to meet the MDR requirements, or no clinical investigation(s) have been performed during this MDR submission, the below statements shall apply during the review of your file:

As per MDCG 2020-6 Section 5, page 9 of 22, the European Commission guidance MEDDEV 2.12/2 regarding PMCF also notes that in the case that clinical evaluation was based exclusively on clinical data from equivalent devices for initial conformity assessment, the certifying notified body shall verify that **PMCF studies** have been conducted.

#### **Applicable devices**

All Legacy devices which previously claimed equivalence during their initial assessment (when the device was first CE marked).

As per MEDDEV 2.12/2 all MDR new application (Not Legacy devices) claiming equivalence need to provide a PMCF study plan. Please ensure to include PMCF clinical study plan in your submission.

### **Applicable devices**

All MDR new application (Not Legacy devices) claiming equivalence.