TANZANIA MEDICINES AND MEDICAL DEVICES AUTHORITY



APPLICATION FORM FOR REGISTRATION OF MEDICAL DEVICES

Please read this section carefully before completing the form

- 1. Please check the corresponding boxes in the "Encl." column if any document is enclosed and indicate the respective indexes in the submission folder
- 2. Please check the boxes as appropriate

Note	Part A: Particulars of Applie	cant	Ere al
A1	Applicant's name		Encl.
	Address of Head Office		-
	Post Code:	Country:	-
	Contact Person:	Telephone:	
	Fax:	E-mail:	-
	Website:		-
	Part B: Particulars of the Mar	nufacturing Site(s)	
	Name		
B1	Physical address of the site		
	Post Code:	Country:	
	Contact Person:	Telephone:	-
	Fax:	E-mail:	-
	Website:	1	-
			-

Particulars of the Manufactu manufacturing site)	are (if different from	
Name		
Physical address of the site		
Post Code:	Country:	
Contact Person:	Telephone:	

	Quality Management System Established by the Manufacturing Site(s):	
B2	Mention current Standards with which the system complies :	
	System certified by	
	and a certified copy of the certificate is enclosed.	
	Indicate areas covered by Quality Management System Device	
	design	
	Production	
	Post-production processes	
	Others (<i>please specify</i>)	
	Part C: Particulars of Authorized Representative (AR)	
C1	LRP's name	
	Address of the registered business premise	

Contact person:	Telephone:	
Fax:	E-mail:	
Contact telephone for public e <i>given above</i>):	enquiries (<i>if different from the number</i>	
	enquiries (<i>if aifferent from the number</i>	

	Certified copy of business registration certificate with business registration number: is enclosed	
C2	Certified copy of Power of attorney or formal agreement or any other official authorization of the LRP is enclosed	
СЗ	The AR is also an importer of the device named in Part D	
	Part D: Particulars of the Device	
D1	Generic name of the Device	
D2	Brand name of the device	
D3	Model/Series/System (<i>if applicable</i>)	
D4	Family (<i>if applicable</i>)	
D5	Country of origin	

	Select GMDN (Global Medical Device Nomenclature) Categories:
D6	 01 - Active implantable device 02 - Anaesthetic and respiratory devices 03 - Dental devices 04 - Electro mechanical devices 05 - Hospital hardware 06 - In vitro diagnostic devices 07 - Non-active implantable devices 08 - Ophthalmic and optical devices 09 - Reusable instruments 10 - Single use devices 11 - Technical aids for disabled persons 12 - Diagnostic and therapeutic radiation devices 13 - Complimentary therapy devices 14 - Biologically -derived devices 15 - Healthcare facility products and adaptations 16 - Laboratory equipment 17 - Others
D7	Description of the device (<i>Please enter appropriate GMDN description</i> . If none of the descriptions in GMDN appear
	appropriate, enter a short description of the device)
D8	GMDN Code: (Please enter if known)
D9	Other common descriptions of the device:
D10	Intended use of device
D11	Class of the medical device:
	Class A
	Class B
	Class C
	Class D

D12	Reasons for classifying the device as Class A, B, C or D device:	
D13	 History No previous recalls, reportable adverse incidents, banning in other countries or post-market surveillance studies. Yes (<i>Please tick the appropriate boxes and provide details</i>): 	
	Recalls completed or in progress	
	Any reportable adverse incidents bearing implications to the Device	
	The device banned previously in other countries Pro-	
	active post-market surveillance studies	
	Performance and safety:	
D13	International or national standards with which the device complies	
	(Please enclose copy of the standard)	
	Part E: Marketing Approvals in Foreign countries	
E1	Mention the countries where the device has obtained marketing approvals	
	(Please enclose certified copy of validmarketing authorization)	
E2	Mention the countries where the device approval is still pending	
	Part F: Declaration of conformity (DoC)	
F1	Submit a written declaration of conformity. The DoC should contain the following:-	

(i)	
	has been classified accordingly and has met applicable
	conformity assessment element.
(ii	Information is sufficient to identify the device including its
	nomenclature.
(ii	i) The risk class allocated to the device.
(iv	y Which of the conformity assessment elements have been
	applied
(v	The date from which the DoC is valid.
(v	i) The name and address of the device manufacturer.
(v	ii) The name, position and signature of the responsible person
	who has been authorized to complete the DoC.

Declaration by applicant

I, the undersigned certify that all the information in this form and accompanying documentation is correct and true to the best of my knowledge.

Name:

Position:______
Signature: ______
Official stamp:

Date: _____