

For office use only:

Clinical Investigation –Notification for Modification of Clinical Investigation under the Medical Device Regulation received on

Clinical Investigation – Notification for Modification of Clinical Investigation under the Medical Device Regulation Reference No.

MT-MDF17

Clinical Investigation – Notification for Modification of Clinical Investigation under the Medical Device Regulation (EU) 2017/745

The application is valid when submitted with the relevant documents and fees, where applicable.

Relevant correspondence should reach the Authority via Email address mdforms.medicinesauthority@gov.mt.

Refer to GL-MDF15 Guidance for Notification for Modification of Clinical Investigations under the Regulation (EU) 2017/745 and GL-MDF07 Guidance on fees in relation to Medical Devices. Guidance and Application Form are available on the Malta Medicines Authority website www.medicinesauthority.gov.mt.

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SECTION A: NOTIFICATION INTRODUCTION & APPLICANT DETAILS

A.1 Date of Notification (dd/mm/yyyy):		
Name & Surname:		
Email Address:		
Contact Number:		
A.2 Applicant (tick as applicable)		
Sponsor (within the Union): Fill in	n Section B	
Legal Representative (when Sponsor is outside the Union): Fill in Section C		
SECTION B: SPONSOR CONTACT DETAILS		
Organisation Name:	Telephone Number:	
Address:	Contact Name:	
	Job Title:	
	Email address:	
If Organisation is registered with the Auth	nority, quote reference number	



MDC004/01 Appendix 1 Version 1 **SECTION C: LEGAL REPRESENTATIVE CONTACT DETAILS**

Organisation Name:	Telephone Number:
Address:	Contact Name:
	Job Title:
	Email address:
If Organisation is registered with the A	Authority, quote reference number
SECTION D: INVESTIGATOR CO	NTACT DETAILS
Name:	Telephone Number:
Job Title:	
Email address:	
Healthcare Institution:	



MDC004/01 Appendix 1 Version 1 SECTION E: IDENTIFICATION OF THE CLINICAL INVESTIGATION

E.1 MMA Reference No.:
Clinical Investigation ID:
Clinical Investigation Title:
Date of previous authorisation/notification:
E.2 Clinical Investigation Plan (CIP) details:
CIP code:
CIP version no.:
CIP date:
SECTION F: INFORMATION ON THE DEVICE
F.1 Name of Device
F.2 Model of Device
F.3 Classification of Medical Device
F.4 Description of Device including its intended purpose
F.5 Are there any changes or modifications in relation to the device or its components since the previous application/notification to the Malta Medicines Authority?
Yes (If yes, provide a rationale and description of these changes)
\square No



SECTION G: NOTIFICATION FORM (EU)

Refer to the MDCG 2021-28 guidance in the Clinical Investigation and Evaluation section.

Docum	nentation in attachment:
	Filled in relevant forms Supporting documents
SECT	ION H: DETAILS OF PAYMENT
	Proof of Payment attached
SECT	ION I: DATA PROTECTION CONSENT STATEMENT
accord 2016/6 Protect Data I Medica initiall to the process	The applicant hereby consents to the processing of their personal data by the Medicines Authority and understands that this data shall be processed in lance with the General Data Protection Regulation (GDPR), Regulation (79/EU of the European Parliament and of the Council of 27 April 2016, the Data tion Act (Chapter 586 of the Laws of Malta) and the Malta Medicines Authority Protection Policy (P-MA05). The applicant also understands that the Malta ines Authority shall process this personal data in line with the purposes they are by collected for. Exceptions to the latter include when the data subject consents new purpose, when there is a legal provision requiring or allowing the new saing or when the new purpose is deemed compatible with the purposes the tall data were initially collected for.



Malta Medicines Authority Declaration for Form Submission

Date:

I, the applicant, declare that all information given in the application form is true, complete and correct. I also bind myself to inform immediately any change to details in the application form and annexes, where relevant, to the Malta Medicines Authority.

Company Name (if applicable):

Name & Surname:

Position: