Clinical investigation – application form under Medical Device Regulation.

Application form version 1.1

Section 1: Clinical investigation identification			
1.1 Sponsor identification			
Name:			
	Street name:	Street number:	
Address	Postal code:	City:	
	Country:	<u> </u>	
Telephone numb	per:		
Email:			
Contact	person of the sponsor		
First name:			
Last name:			
Telephone numbe	r:		
Email:			

Sponsor's legal representative identification

Do you have a leg	gal representative?		
Yes	No		
		legal representative (section 1.2)	
1.2 Legal represent	tative identification		
Organisation nan	ne:		
	Street name:	Street number:	
Address	Postal code:	City:	
	Country:		
Telephone numb	er:		
Email:			
Contact per	rson of the legal representa	<u>tive</u>	
First name:			
Last name:			
Telephone number	•		
Email:			

Contact person for the clinical investigation

Same as contact person of sponsor Same as contact

clinical investigation.

person of legal representative

Other con	tact person for the clinical inves	<u>tigation</u>		
First name:				
T .				
Last name:				
	Street name:	Street number:		
Address	Postal code:	City:		
	Country:			

If you selected other, please fill in the section below related to the other contact person for this

1.3 Clinical investigation type

Select the appropriate regulatory pathway for the application :
Clinical investigation application (MDR Art. 62(1)/IVDR Art. 58(1))
PMCF investigation notification (MDR Art. 74(1))
Other clinical investigation application/notification - national application (MDR Art. 82(1))
1.4 Submission type
First submission in the EEA
First submission at the national level (clinical investigation has been already submitted in EEA)
In this case, please provide the clinical investigation ID (CIV-ID) provided
Resubmission Please provide the CIV-ID if already available
1.5 Participating countries within the EU/EEA/UK (Northern Ireland), Turkey and Switzerland
Select the participating countries for the clinical investigation

1.6 Participating countries outside EU/EEA, Northern Ireland, Turkey and Switzerland

If this study is part of a multi-site clinical investigation outside the EU/EEA, Northern Ireland, Turkey and Switzerland please provide a list of all the non EU/EEA countries the study plans to be carried out in.
1.7 Clinical investigation plan (CIP)
CIP code:
CIP version:
CIP date:
1.8 Clinical investigation title
Full title :
Short title :
Title for lay people:

Section 2: Clinical investigation description

2.1 Scientific opinion

Has the manufacturer consulted with an expert panel as outlined in Art. 61(2) of Regulation (EU) 2017/745.

Y Yes

No

2.2 Design of the clinical investigation

Exploratory investigation Confirmatory investigation

Observational investigation

First in human investigation

Not first in human

2.3 Design methodology

Case Control Controlled Cross-sectional Double blind

Parallel Randomised Open

Other:

2.4 Development stage

Pilot stage Pivotal stage Post-market stage

2.5 Objectives and endpoints

Drive any objective (c).	
Primary objective(s):	
Secondary objective(c).	
Secondary objective(s):	
Other objective(s):	
Primary endpoint(s):	

Secondary endpoint(s):	
Other endpoint(s):	
2.6 Synopsis of the clinical investigation	
Overall synopsis:	

2.7 Planned number of subjects

In Europe:	
In Asia:	
In Africa:	
In North America:	
In South America:	
In Oceania:	
Total planned number of subjects:	
2.8 Duration of clinical investigation	
Estimated start date:	
Estimated end date:	
2.9 Population	
2.9.1 Medical condition	
Is there an associated medical condition?	
Yes No	
Is the medical condition considered to be rare?	
Yes No	
2.9.2 Therapeutic area	
Select the therapeutic area that the clinical investigation fal	ls under
	If "other" is selected, please define the
	therapeutic area:
2.9.3 Gender of subjects	
Female Male Other	

0 0 5 EL	
2.9.5 Exclusion criteria	

2.9.4 Inclusion criteria

${\bf 2.9.6\,Type\,\,of\,subjects\,\,that\,\,the\,\,clinical\,\,investigation\,\,plans\,\,to\,\,recruit}$

Healthy	Patients	Vulnerable population	Incapacited subjects
Minors	Pregnant women	Breastfeeding women	Patients in emergency situations
Other (plea	ase specify)		

2.9.7 Age range of the participants that the clinical investigation plans to include

In utero	Adults (from 18 to 84 years)
Newborns (from 0 to 27 days)	Elderly (from 85 years)
Infants and toddlers (from 28 days to 23 months)	
Children (from 2 to 12 years)	
Adolescents (from 12 to 17 years)	

2.10 Scope of the investigational device

2.10.1 Combined investigation Medical Device/In Vitro Diagnostic?

Yes No		
If yes, please provid	de the related IVD performance st	tudy identification number
	he application submitted in paralle	el with an application for a clinical trial on medicinal
products?		
Yes No		
If yes, please provide	the EU Clinical Trial Number:	
9 11 Coordinat	ling at increase the section of	
<u>2.11 Coorumat</u>	ting investigator	
First name:		
Last name:		
Last name:		
	Street name:	Street number:
Address	Postal code:	City:
Address		
	Country:	<u>_</u>
	J	
Telephone number	•	·
Telephone number	•	
Email:		

Section 3: Investigational device(s)

3.1 Investigational medical device

3.1.1 Device purposes

3.1.2 Device type

Implantable	System

Active device Non-medical purpose

Measuring function Sterile

Reusable surgical instrument Software

Intended to administer or remove medicinal

substance

3.1.3 Invasiness

Is it an invasive medical device?

Yes No

3.1.4 Device Identifiers

Generic denomination:
Device trade name: Model:
Device name:
European Medical Device nomenclature (https://webgate.ec.europa.eu/dyna2/emdn/)
Medical device classification:
Medical device classification:
Classification rule:
Device description:
Intended (clinical) purpose:
Does the device contain or incorporate medicinal substance(s)?
Yes No
Tes No
If yes, please provide the medicinal substance(s) name(s):
The device incorporates, as an integral part, or it is manufactured using:
Non-viable tissues of human origin or their derivatives with an ancillary action
Non-viable cells of human origin or their derivatives with an ancillary action Non- viable tissues of animal origin or their derivatives with an ancillary action Non-
viable cells of animal origin or their derivatives with an ancillary action
Non-viable biological substance other than those referred to in the previous points None of these proposals/Not applicable
1020 of these proposition for applicable

Is the Investigational Device CE marked?
Yes No
If yes, please provide the information in the box below.
To what extent is the intended purpose of the device in the clinical investigation covered by the CE-mark?
CE marked device will be used outside the scope of its CE mark
CE marked device will be used within the scope of its CE mark and no additional procedures are foreseen in the clinical investigation
CE marked device will used within the scope of its CE mark, but additional procedures are foreseen in the clinical investigation
Are those additional procedures considered to be burdensome and/or invasive?
Yes No
Please, comment why do you consider as such?
Information related to the Notified body involved, if applicable:
Notified body number:
Notified body name:
3.2 Previous clinical investigation
Has this device been investigated in a clinical investigation within the EEA previously?
Yes No
If yes, please provide the relevant reference number(s) (such as SIN, CIV-ID, other reference(s)) of the previo clinical investigations
3.3 Scientific opinion/view
Has the investigational/study device been subject to a national scientific view/opinion from an Expert Panel
Yes No
3.4 Manufacturer of the investigational device
Is the manufacturer the same as the sponsor?
Yes No
If no, please fill in the requested information in section 3.4.1 and 3.4.2.

3.4.1 Manufacturer information

Organisation nar	me:	
_	Street name:	Street number:
Address	Postal code:	City:
	Country:	I.
Telephone numb	per:	
Email:		
Contact p	person of the manufacturer	
First name:		
Last name:		
Telephone number	::	
Email:		
3.4.2 Auth	norised representative	
Organisation nar	ne:	
	Street name:	Street number:
Address	Postal code:	City:
	Country:	L.
Telephone numb	oer:	
Email:		
Contact p	person of the authorised repr	<u>esentative</u>
First name:		
Last name:		
Telephone number	:	
Email:		

Additional devices could be added by using a duplicated section 3, in appendix to this application form.

Section 4: Comparator

4.1 Applicability of section 4

Is there a comparator included in the clinical investigation	ation?.		
Yes No			
If yes, the section from 4.2 needs to be completed.			
4.2 Type of comparator			
Therapy			
Placebo			
No treatment			
Medical device			
4.2.1 Medical device as comparator			
Is the comparator medical device CE marked?	Yes	No	
If yes, will the CE marked comparator medical device	be used in t	the clinical invest	gation within the
scope of its CE mark? Yes No			
Generic denomination:			
Device trade name:	Mod	lel:	
Device name:			
European Medical Device Nomenclature :			
Medical device classification:			
Device description:			

Intended (clinical) purpose:
Does the comparator device contain or incorporate medicinal substance(s)?
Yes No
If yes, please provide the medicinal substance(s) name(s):
The comparator device incorporates, as an integral part, or it is manufactured using:
Non-viable tissues of human origin or their derivatives with an ancillary action Non-
viable cells of human origin or their derivatives with an ancillary action Non- viable tissues of animal origin or their derivatives with an ancillary action Non-viable cells
of animal origin or their derivatives with an ancillary action
Non-viable biological substance other than those referred to in the previous points
None of these proposals/Not applicable

Additional comparators could be added by using a duplicated section 4, in appendix to this application form.

Section 5: National information

5.1 Study site information

Please provide the list of sites taking part in the clinical investigation

Name of institution	Site address	Investigator attached to this site	Contact information of investigators
,			
			<u> </u>

Additional sites could be added by using a duplicated section 5.1, in appendix to this application form

5.2 Ethics committee information

Select the applicab	Select the applicable option:		
Ethics com	Ethics committee opinion available		
Ethics com	Ethics committee opinion under review		
Ethics com	Ethics committee opinion is not mandatory before submission to the competent authority		
If an ethics committee has to be selected by the sponsor before submission, please provide the ethics committee information's below.			
Organisation name:			
_	Street name: Street number:		
Address	Postal code:	City:	
	Country:	<u> </u>	
Telephone number	Telephone number:		
Email:			

5.3 Status of the clinical investigation

Is the sponsor considered as commercial according to national legislation?
Yes No
5.4 Expected number of subjects recruited within the Member State
How many subjects are expected to be recruited into the study in the Member State you are applying to?
I hereby certify that the information and documentation submitted with this notification is correct in detail and all the information requested has been supplied. The investigated (medical) device
complies with the applicable general safety and performance requirements, apart from those covered by he investigation and that every precaution has been taken to protect the health and safety of the patient and/or user.
I confirm that all the clinical investigations information collected for this application, has been done in compliance with the European data protection legislation (GDPR).
Date:
Name:
Position: