Manufacturer Incident Report (MIR) for Serious Incidents (MDR/IVDR) and Incidents (AIMDD/MDD/IVDD)

Reporting Template Version 7.2.1

European Union Medical Devices Vigilance System

Sec	Section 1: Administrative information						
1.1	Corresponding competent authority						
а	Name of receiving national competent authority (NCA) Ministero Della Salute						
b	EUDAMED number of NCA						
С	Reference number assigned by NCA for this incident Not Known						
d	Reference number assigned by EUDAMED for this incident						
1.2	Date, type, and classification of incident report						
а	Date of submission b Date of incident (e.g. 2012-10-23) c Manufacturer awareness date 2024-03-04 (e.g. 2012-10-23) 2023-11-01 to 2023-11-01 c Manufacturer awareness date						
d	Type of report ☑ Initial □ Follow up □ Combined initial and final □ Final (Reportable incident) □ Final (Non-reportable incident)						
е	In case of initial and follow-up reports, please indicate the expected date of the next report 2024-05-30 (e.g. 2012-10-23)						
f	Classification of incident Classification of incident Serious public health threat Death Unanticipated serious deterioration in state of health All other reportable incidents						
1.3	Submitter information						
1.3.1	Submitter of the report						
а	☑ Manufacturer ☐ Authorised representative ☐ Other, please specify						
b	Manufacturer's reference number for this incident 17907248						
С	If this incident involves multiple devices from the same manufacturer, please list the respective reference numbers of the other MIR forms you have submitted - NCA's local reference number						
	- EUDAMED's reference number						

	- Manufacturer's reference number		
	If this incident is covered under an FSCA, please		ide the relevant numbers:
d		5 pi 0 v	
	- NCA's local FSCA reference number		
	- EUDAMED's FSCA reference number		
	- Manufacturer's FSCA reference number		
е	Periodic Summary Report (PSR) ID		
f	If the incident occurred within a PMCF/PMPF inv	/estig	ation; please provide the Eudamed ID of that
	PMCF/PMPF investigation		
1.3.2	Manufacturer information		
а	Manufacturer organisation name		
	Boston Scientific Corporation – CRM		
b	Single registration number		
	N/A		
с	Contact's first name	d	Contact's last name
	Rossana	-	Perego
е	Email	f	Phone
Ŭ	MILBSCRegulatory@bsci.com, Giovanna.Pira@bsci.com		1-651-582-4000
g	Country		
	United States		
h	Street	i	Street number
	Hamline Avenue North		4100
i	Address complement	k	PO Box
1	City name	m	
	Saint Paul, MN		55112
1.3.3	Authorised representative information		
а	Authorised representative organisation name		
	Guidant Europe SA/NV- Boston Scientific		
b	Single Registration Number		
	N/A		
С	Contact's first name	d	Contact's last name
	Sophie		Vaillot
е	Email	f	Phone
	sophie.vaillot@bsci.com		N/A
g	Country		
	Belgium		
h	Street	i	Street number
	Green Square - Lambroekstraat 5D		N/A
i	Address complement	k	PO Box
1	City name	m	Postal code
	Diegem		1831

1.3.4	Submitter's details if not also manufacturer of	or au	thorised representative
а	Registered commercial name of company Boston Scientific Corporation – CRM		
h	Contact's first name		
b	Rossana	C	Contact's last name Perego
d	Email MILBSCRegulatory@bsci.com, Giovanna.Pira@bsci.com	l e	
			1-651-582-4000
f	Country United States		
		_	
g	Street	h	
	Hamline Avenue North		4100
i	Address complement	j	PO Box
k	City name	I.	Postal code
	Saint Paul, MN		55112
Se	ction 2: Medical device information		
2.1	Unique Device Identification (UDI)		
а	UDI device identifier/Eudamed ID	b	UDI production identifier
	Unknown		Unknown
с	Basic UDI-DI/Eudamed-DI	d	Unit of use UDI-DI
	Unknown		
2.2	Categorisation of device		
а	Medical device terminology		
	EMDN GMDN UMDNS(ECRI)	D/ED	MS LI Other, please specify
b	Medical device nomenclature code 36241		
		¢	
2.3	Description of device and commercial in	TOR	nation
а	Medical device name (brand/trade /proprietary c	or cor	nmon name)
	ENDOTAK RELIANCE		
b	Nomenclature text/Description of the device and	l ite i	ntended use
b	Defibrillator / Pacemaker Lead	1113 11	
с	Model		Catalogue/reference number
C	0148	d	
е	Serial number	f	Lot/batch number
g	Software version	h	Firmware version
I	Device manufacturing date (e.g. 2012-10-23)	j	Device expiry date (e.g. 2012-10-23) 2002-10-01
	2000-10-01		
k	Date when device was implanted (e.g. 2012-10-23)	I	Date when device was explanted (e.g. 2012-10-23)
	2000-12-01 to 2000-12-01		to

	Number of years Number of months	Number of days				
n	Implant facility o	Explant facility				
	Istituto Per La Sicurezza Sociale					
р	Notified body (NB) ID number(s) (if applicable) No	tified body (NB) certificate number(s) of device (if applicable)				
	1 2797 54	1612				
	2					
	Please indicate the date of <u>one</u> of the following:					
q	□ First declaration of conformity					
	I The device first CE marked					
	□ First placed on the market					
	□ First put into service					
	□ If software, date first made available					
	Year 2000 Month 5					
2.4	Risk class of device when placed on market	t				
а		efore the implementation of the MDD/AIMDD/IVDD				
b	MDD/AIMDD	IVDD				
	⊠ active implant □ class III	□ IVD Annex II List A				
		□ IVD Annex II List B				
		 IVD devices for self-testing IVD general 				
	□ class I					
	□ class Is					
	□ class Im					
	□ class Ism					
С						
Ŭ	MDR Type (Multiple choice) □ class III □ implantable	IVDR Type (Multiple choice) class D Image: state				
	□ class IIb □ active device	□ self-testing				
	□ class IIa □ intended to administer and/or	□ class C □ near-patient testing □ class B □ professional testing				
	class I remove a medicinal product sterile conditions	□ class A □ companion diagnostic				
	□ reusable surgical instruments	□ software □ instrument				
	□ software					
	□ systems					
	□ procedure packs □ custom-made					
	Market distribution of device (region/count	trv)				
2.5	(according to the best knowledge of the manu					
а	All EEA, Switzerland and Turkey					
	· · · · ·					
	Others:					

2.6	Use of accessories, associated devices or other devices
а	Relevant accessories used with the device being reported on (please list with corresponding Manufacturer if different from device being reported on)
b	Relevant associated devices used with the device being reported on (please list with corresponding Manufacturer if different from device reported on)

Section 3: Incident information derived from healthcare professional/facility/patient/lay user/other

3.1	Nature of incident							
а	Provide a comprehensive description of the incident, including (1) what went wrong with the device (if applicable) and (2) a description of the health effects (if applicable), i.e. clinical signs, symptoms, conditions as well as the overall health impact (i.e. Death; life-threatening; hospitalization - initial or prolonged; required intervention to prevent permanent damage; disability or permanent damage; congenital anomaly/Birth defects; indirect harm; no serious outcome)							
	It was reported that the right ventricular (RV) lead exhibited high out of range pacing impedance measurements of greater than 2000 Ohms along with increasing threshold measurements. The patient underwent provocation maneuvers and no oversensing or noise was observed. It was believed that the change in values is due to the age of the lead as it has been implanted for 24 years. The output was increased to maximum value. Data was sent into Technical Services (TS) for review. The RV lead remains in service and no adverse effects were reported. No additional information is available. If additional information becomes available the report will be updated at that time.							
3.2	Medical device problem information							
а	IMDRF Medical device problem codes (Annex A) Coding with IMDRF terms is a mandatory requirement.							
	Choice 1 (most relevant) Choice 2 Choice 3 Choice 4 Choice 5 Choice 6							
	IMDRF 'Medical device problem codes' Code Code Code Code Code Code							
	If you think the incident is unique and a suitable IMDRF term is missing, briefly explain:							
b	Number of patients involved							
С	What is the current location of the device? Healthcare facility/carer Distributor Patient/user Discarded In transit to manufacturer Image: Remains implanted							
	□ Manufacturer □ Unknown □Other:							
d	Operator of device at the time of the incident							
	□ Healthcare professional I Patient/lay user □ Other, please describ∈							
е	Usage of device (as intended) Initial use Reuse of a single use medical device Reuse of a reusable medical device Reuse of a reusable medical device Orbolem noted prior use Orbolem noted prior use							
f	Remedial actions taken by healthcare facility, patient or user subsequent to the incident							
	See Incident Narrative.							

3.3	Patient information	า						
а	IMDRF 'Health Effect' Coding with IMDRF te				nt.			
		Choice 1 (most relevant)	Choice	2	Choice 3	Choice 4	Choice 5	Choice 6
	IMDRF 'Clinical signs, symptoms, and conditions	Code	Code		Code	Code	Code	Code
	codes' (Annex E)	E2403		<u> </u>			Code	
	IMDRF 'Health impact' codes (Annex F)	Code F26	Code		Code	Code	Code	Code
	If you think the incide	nt is unique an	nd a suitab	le IMD	ORF term is r	nissing, briefly	explain:	·
b	Age of patient at the ti	ime of the incion months	dent	days				
с	Gender 🛛 Fema	ale □ Ma	le	□Un	known E	Not applicable	9	
d	Body Weight (kg)]						
е	List any of the patient	's prior health	condition of	or med	dication that	may be releva	nt to this incide	ent
2 1	Initial reporter (car	ho hoalthe	aro profo		aal of facili	ty patient la	av usor)	
3.4	Initial reporter (car		are profe	essio	nal of facili	ty, patient, la	ay user)	
3.4	Initial reporter (car Role of initial reporter ⊠ Healthcare profess		-				ay user)	
а	Role of initial reporter ⊠ Healthcare profess	ional 🛛 Patiei	nt □ Lay	user			ay user)	
	Role of initial reporter	ional	nt □ Lay	user			ay user)	
а	Role of initial reporter Healthcare profess Name of healthcare fa	ional □ Patien acility where in zZZA SOCIALE	nt □ Lay	user			ay user)	
a b c	Role of initial reporter Healthcare profess Name of healthcare fa ISTITUTO PER LA SICURE Healthcare facility rep	ional □ Patien acility where in zZZA SOCIALE	nt □ Lay	user	□ Other, ple	ease specify	ay user)	
a	Role of initial reporter Healthcare profess Name of healthcare fa ISTITUTO PER LA SICURE	ional □ Patien acility where in zZZA SOCIALE	nt □ Lay	user	□ Other, ple		ay user)	
a b c	Role of initial reporter Healthcare profess Name of healthcare fa ISTITUTO PER LA SICURE Healthcare facility rep Contact's first name Roberto Email	ional □ Patien acility where in zZZA SOCIALE	nt □ Lay	user	Contact's	ease specify	ay user)	
a b c d	Role of initial reporter Healthcare profess Name of healthcare fa ISTITUTO PER LA SICURE Healthcare facility rep Contact's first name Roberto Email rtomassoni@omniway.sm	ional □ Patien acility where in zZZA SOCIALE	nt □ Lay	user curred	Contact's	ease specify	ay user)	
a b c	Role of initial reporter Healthcare profess Name of healthcare fa ISTITUTO PER LA SICURE Healthcare facility rep Contact's first name Roberto Email	ional □ Patien acility where in zZZA SOCIALE	nt □ Lay	user curred	Contact's Tomassoni Phone	ease specify	ay user)	
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a b c d f i	Role of initial reporter Healthcare profess Name of healthcare fa ISTITUTO PER LA SICURE Healthcare facility rep Contact's first name Roberto Email rtomassoni@omniway.sm Country San Marino Street VIA SCIALOIA, 20	ional D Patier	nt □ Lay	user curred e) g	Contact's Contact's Tomassoni Phone +011(378)05	last name	ay user)	
a b c d f	Role of initial reporter Healthcare profess Name of healthcare fa ISTITUTO PER LA SICURE Healthcare facility rep Contact's first name Roberto Email rtomassoni@omniway.sm Country San Marino Street	ional D Patier	nt □ Lay	user curred e) g	Contact's Tomassoni Phone +011(378)05	last name	ay user)	
a b c d f i	Role of initial reporter Healthcare profess Name of healthcare fa ISTITUTO PER LA SICURE Healthcare facility rep Contact's first name Roberto Email rtomassoni@omniway.sm Country San Marino Street VIA SCIALOIA, 20	ional D Patier	nt □ Lay	user curred e) g	Contact's Contact's Tomassoni Phone +011(378)05	ease specify last name	ay user)	

Sec	Section 4: Manufacturer analysis								
4.1	Manufacturer's preliminary comments								
а	For initial and follow-up reports: preliminary results and conclusions of manufacturer's investigation								
	N/A								
b	Initial actions (corrective and/or preventive) implemented by the manufacturer								
	N/A								
с	What further investigations do you intend in view of reaching final conclusions?								
	N/A								
4.2	Cause investigation and conclusion								
- 	For Final (Reportable incident): Description of the manufacturer's evaluation concerning possible ro								
a	causes/causative factors and conclusion								
b	For Final (Non-reportable incident): Fill out rationale for why this is considered not reportable								
с	Is root cause confirmed?								
d	Has the risk assessment been reviewed?								
	No If 'No', rationale for no review required:								
	If the risk assessment has been reviewed, is it still adequate?								
	□ Yes								
	□ No Results of the assessment:								
е	IMDRF 'Cause Investigation' terms and codes (Annex B, C, D)								
	Coding with IMDRF Choice 1 Choice 2 Choice 3 Choice 4 Choice 5 Choice 6 Choice 7 Choice 8								
	terms is a mandatory (most relevant) requirement.								
	IMDRF Cause Code Code Code Code Code Code Code Code investigation: Type of Image: Code Image: Co								
	investigation (Annex B)								

	IMDRF Cause investigation: Investigation findings (Annex C)	Code	Code Co	ode Code	Code	Code		
	IMDRF Cause investigation: Investigation conclusion (Annex D)	Code		ode Code	Code	Code		
	If you think the incider	nt is unique an	d a suitable IN	IDRF term is n	nissing, briefly	explain:		
f	IMDRF Component co Coding with IMDRF te	erms is a mano	datory requirer					
		Choice 1 (most relevant)	Choice 2	Choice 3	Choice 4	Choice 5	Choice 6	
	IMDRF 'Component' codes (Annex G)	Code	Code	Code	Code	Code	Code	
	If you think the incider	nt is unique an	nd a suitable IN	/DRF term is r	nissing, briefly	explain:		
g	Description of remedia (For a FSCA, fill in the FSC/		ctive action/pre	eventive action	/field safety co	rrective action	(FSCA)	
h	Time schedule for the implementation of the identified actions							
i	Final comments from t	he manufactur	rer on cause ir	nvestigation an	d conclusion			

4.3	Similar incidents (for Final (Reportable inciden	t))						
4.3.1	Use of IMDRF terms and codes for identifying simila	ar incidents						
а	Identification of similar incidents using IMDRF Adverse Tick-mark which code or combination of codes were use							
	IMDRF code relating to most relevant 'Medical device problem' (Annex A) □ IMDRF code relating to most relevant 'Investigation finding' (Annex C, 'Cause investigation) □ □ Other - enter description of what similar incidents are based on and the rationale why the above IMDRF codes were not used							
4.3.2	Use of in-house terms/codes for identifying similar i	incidents (only for transition period)						
а	If similar incident were not identified by IMDRF codes b and terms below.	ut by in-house codes, please provide the codes						
	Code/term for most relevant medical device problem	Choice 1 Code						
	Code/term for most relevant root cause evaluation	Term Code Term						
	Other - enter description of what similar incidents ar were not used	re based on and the rationale why the above codes						
4.3.3	Number of similar incidents and devices on the mar	ket						
а	Indicate on which basis similar incidents were identified Model Software Lot/Batch Details of the selection made above							
b	 Indicate to what criteria the number of devices on the m (tick the most appropriate): Devices placed on the market or put into service Units distributed within each time period Number of tests performed Number of episodes of use (for reusable devices) Active installed base Units distributed from the date of declaration of conf time period Number of devices implanted Other -describe 							

С	Enter the number of similar incidents and devices on the market for the indicated time You must use yearly time periods unless: A: a different time period has been specified by the European vigilance Working Grou B: the device has not been on the European market for more than three years								
		Time pe	eriod (N)	Time per	iod (N-1)	Time per	iod (N-2)	Time per	iod (N-3)
			e = incident ar 2-10-23)	calendar year one year before incident (e.g. 2012-10-23)		calendar year two years before incident (e.g. 2012-10-23)		,	
	Start Date						,		
	End Date	End Date							
		Number of similar incidents	Number of devices on market	Number of similar incidents	Number of devices on market	Number of similar incidents	Number of devices on market	Number of similar incidents	Number of devices on market
	Country of incident								
	EEA + CH + TR								
	World								
d	Comments on h	now similar	incidents a	nd associate	ed number o	of devices c	on the mark	et were det	erminec

Section 5: General Comments

Local Affiliate Contact Information: Name: Boston Scientific S.p.A. Contact Name: Rossana Perego Address: Viale Forlanini 23 City/Postal Code: Milano, 20134 Country: Italy Phone Number: +39(02)26983225 Fax Number: +39(02)26983230 Email: MILBSCRegulatory@bsci.com, Giovanna.Pira@bsci.com

If the device is received for analysis an investigation which involves altering the device will take place. Boston Scientific Corporation will assume destructive analysis can begin unless contacted immediately following submission of this report opposing the analysis.

Coded summary of report (will be auto populated from previous selections)									
	Medical device name ENDOTAK RELIANCE								
	Basic UDI-DI Unknow	n							
	UDI device Unknow	n			UDI produ identifier	uctior Unkn	iown		
	IMDRF adverse event reporting terms and codes IMDRF=International Medical Device Regulators Forum. Coding with IMDRF terms is a mandatory requirement.								
	IMDRF clinical signs, symptoms, conditions codes	E2403							
	IMDRF health impact codes	F26							
	IMDRF Medical device problem codes	A072201							
	IMDRF Component codes								
	IMDRF Cause investigation: Type of investigation								
	IMDRF Cause investigation: Investigation findings.								
	IMDRF Cause investigation: Investigation conclusion.								

Submission of this report does not represent a conclusion by the manufacturer and / or authorised representative or the national competent authority that the content of this report is complete or accurate, that the medical device(s) listed failed in any manner and/or that the medical device(s) caused or contributed to the alleged death or deterioration in the state of the health of any person.

I affirm that the information given above is correct to the best of my knowledge.

Before signing and submitting	
Check the form	Save as PDF
Date 2024-03-04	
Signature/Digital Signature	
Send as XML file	Submit XML by Email
Send as PDF file	Submit PDF by Email

Before signing and submitting