Manufacturer's Periodic Summary Report (PSR) Medical Devices Vigilance System (MEDDEV 2.12/1 rev 7)

v.12/11

1. Administration Information				
To which NCA(s) is this report being sent?				
Date of this report				
Reference number assigned by the manufacturer				
Reference number assigned by NCA				
Type of report				
Initial report				
Follow up report Follow up Number s				
☐ Final report				
2. Information on submitter of the report				
Status of submitter				
Manufacturer				
Authorised Representative within EEA, Switzerland and T	urkey			
☐ Others: (identify the role) :				
3. Manufacturer information				
Name				
Contact name				
Address				
Postcode	City			
Phone	Fax			
E-mail	Country			
4. Authorised Representative information				
Name				
Contact name				
Address				
Postcode	City			
Phone	Fax			
E-mail	Country			
5. Submitter's information (if different from section 3 or 4)				

Submitter's name					
Contact name					
Address					
Postcode	City				
Phone	Fax				
E-mail	Country				
6. Medical Device Information					
Class AIMD Active Implants MDD Class III MDD Class IIb MDD Class IIa MDD Class I	 IVD Annex II List A IVD Annex II List B IVD Devices for self-testing IVD General 				
Nomenclature system (preferable GMDN)	Nomenclature code				
Nomenclature text	·				
Notified Body (NB) ID – Number					

Model number(s) or Family Name			Catalogue number(s)			
7. PSR Information						
PSR Type:						
☐ Incidents described in a Field Safety Notice			Common and well documented incidents			
If Incidents described in a Field Safety Notice, Manufacturers reference number for FSN/FSCA						
Stage of PSR reporting based on:						
Observed Failure mode Root cause						
Nature of problem agreed for PSR reporting						
Summary period agreed:						
Every month Every 2 months Every 3 months Every 6 months Every 12 months						
The figures in the table below relate to:	□ EEA + CH+ TR		R recipients NCA's in Section 1	Single Member State Please name:-		
Date of PSR New incidents this period	Total num incidents		Total number resolved	Total number in progress		

8. Manufacturer's comments / investigation results Investigation update for this period

Initial corrective actions / preventive actions implemented by the manufacturer

Recommended actions for this period, if any

Expected date of next PSR report

9. Distribution

The medical device has been distributed to the following Countries

Within EEA, Switzerland and Turkey:

🗌 AT	🗌 BE	🗌 BG	□сн	CY	□cz	🗌 DE	🗌 DK	🗌 EE	🗌 ES
🗌 FI	🗌 FR	🗌 GB	🗌 GR	🗌 HU	🗌 IE	🗌 IS	🗌 IT	🗌 LI	🗌 LT
🗌 LU	🗌 LV	🗌 MT	🗌 NL	🗌 NO	🗌 PL	🗌 PT	🗌 RO	🗌 SE	🗌 SI
🗆 SK	🗌 TR								

Candidate Countries:

🗌 HR

All EEA, Candidate Countries, Switzerland and Turkey

Others:

10. Comments

Submission of this report does not, in itself, represent a conclusion by the manufacturer and / or authorized 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.

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Name City date