

MEDICAL DEVICE ADVERSE EVENT REPORTING FORM

Materiovigilance Programme of India (MvPI)

This form is intended to collect information on Medical Devices Adverse Event in India. The form is designed to be used voluntarily by Manufacturer/Importer/Distributor of Medical Devices, Healthcare Professionals and anyone with direct/indirect knowledge of Medical Devices Adverse Event.

General Information					
 Date of Report : Type of Report : Initial □ F Reporter Reference for MDMC on 		·		Month-Year • Case No.	
Reporter Details					
1. Type of Reporter : (a) Manufact (e) Patient 2. In case, where the reporter is not (a) Has the reporter informed the Yes No (b) Is the reporter also submitting Yes No (c) No (c) Name (c) Name (c) Address (c) Tel. /Mobile (c) Email (c) Patient (e) Patient	t manu ne incid	\square (f) Others \square specuracturer, fill the following deta	ails:-	or (d) Healthcare Professional	
Device Category					
Device Category Medical Device		In Vitro Diagnostics (I\	/D)	Medical Equipments / Machin	es
		In Vitro Diagnostics (IV I. Kits II. Reagents III. Calibrator IV. Control Material V. Others VI. IVD electronic reader/ Analyzer	/D)	Medical Equipments / Machin I. Therapeutic □ Diagnostic III. Therapeutic & Diagnostic III. Preventive IV. Assistive V. Imaging VI. Invasive □ Non-Invasive VII. Others	es

- If Medical Devices/Equipments/Machines : Please fill all the sections i.e. A, B, C, D, E & F
- If in Vitro Diagnostics (IVD): Please fill sections i.e. A (except 6, 7, 8, 13, 14 & 16), B (except 1, 2, 6 & 8), D, E, & F

(A)	Device Details											
Dev	rice Name / Trade Nam	e / Brand Name:										
	Details	Name					A	ddres	s			
Ma	anufacturer											
Im	porter											
Dis	stributor											
1.	a) Is the device notified,	regulated in India/	:	Yes		No						
	b) Device Risk Classificat	tion as per India MDR 2017	:	Α		В		С		D		
2.	License No. (Manufacture	e/Import)	:									
3.	Catalogue No.		:									
4.	Model No.		:									
5.	Lot / Batch No.		:									
6.	Serial No.		:									
7.	Software Version		:									
8.	Associated Devices / Acc		:									
9.	Nomenclature Code if ap	plicable; GMDN/UMDNS	:									
	UDI No. (If applicable)		:									
	Installation Date		:									
	Expiration Date		:									
	Last preventive maintena		:									
	Last calibration date (dd,	/mm/yyyy)	:									
	Year of manufacturing		:									
	How long was device/Equ		:									
17.	Availability of device for	<u></u>	: Yes □ retu	⊔ ırn to	No	iactur	or or	impor	ter/dis	tribut	or F	1
	If no, was the device des	stroyed in Still III use	rell	1111 LU	manul	actui	Ci UI	mpol	cei/uis	ינוזטענ(JI	_
18.		s per manufacturer claim /Ir	nstruction	for us	se/usei	r man	ual:	Yes		No		
	If no specify usage											
19.	For devices not regulated	d / notified in India	: Regu	lator ,	/ Regu	latory	statı	ıs in c	ountry	of or	igin	

(B) Event Description					
 Date of Event / Near miss incident: Date of Implant/Explant (If applicated) Location of Event: Hospital Premise ☐ Manufacture/Ded) Home ☐ Others ☐ Device Operator:- Healthcare Professional ☐ Patient Problem noted prior to use/near miss. Device disposition / Current location a) Returned to company ☐ If yest Remains implanted in patient Within the healthcare facility At patient home Destroyed Others (specify) Is device in use after incidence: Yes 	ole): istributor pre nt	ners 🗆	If serice a) Dea b) Life c) Disc d) Hos e) Cor f) Any g) Rec Imp 8. Non se 9. Whether	Threatening ability or permanent despitalization agenital anomaly /birth other serious (Imp. repuired intervention to pairment / damage descrious event er other medical devices	continue
10. Detail description of Event:-	es 🗆 No				
For manufacturer/authorized representations of a similar Adverse Event in India in past 3 years 12. Frequency of occurrence of similar Adverse Event in globally	esentative Year Year	No. of Advers	Similar e Events Similar e Events	Total No. Supplied Total No. Supplied	Frequency of Occurrence (%) Frequency of Occurrence (%)
in past 3 years					
(C) Patient Information, His	tory & Oı	ıtcome			
1. Patient Hospital ID : 2. Patient Initial : 3. Age : 4. Gender : Male □ Fe 5. Weight : 6. Other relevant history, including pre conditions			a) Rec b) Not c) Dea d) Oth		YY)

(D) Healthcare Facility Information (if available)	
1. Name:2. Address:3. Contact Person Name at the site of event:4. Tel. No.:	
(E) Causality Assessment	
1. Investigation action taken:	
2. Post spuss of problem (Applicable for follow up / final reports):	
2. Root cause of problem (Applicable for follow up / final reports):	
(F) Manufacturer/Authorized Representative Investigation & Action taken	
1. Manufacturer/Authorized Representative device risk analysis report:	
2. Corrective / preventive action taken:	
3. Device history review:	

B) Event Description (Continued)
). Detail description of Event:-
E) Causality Assessment (Continued)
Investigation action taken:
Root cause of problem (Applicable for follow up / final reports):
F) Manufacturer/Authorized Representative Investigation & Action taken (Continued)
F) Manufacturer/Authorized Representative Investigation & Action taken (Continued) Manufacturer/Authorized Representative device risk analysis reports
F) Manufacturer/Authorized Representative Investigation & Action taken (Continued) . Manufacturer/Authorized Representative device risk analysis report:
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Where to report?

Duly filled Medical Device Adverse Event Reporting Form can be sent to Indian Pharmacopoeia Commission, Ministry of Health and Family Welfare, Government of India, Sector-23, Rajnagar, Ghaziabad-20002, Tel-0120-2783400, 2783401 and 2783392, FAX:0120-2783311 or email to mvpi.ipcindia@gmail.com Or Call on Helpline no. 1800 180 3024 to report Adverse event.

Partnering Organizations







Disclaimer

Confidentiality: The patient's identity is held in strict confidence and protected to the fullest extent. Programme staff is not expected to and will not disclose the reporter's identity in response to a request from the public. Submission of a report does not constitute an admission that medical personnel or manufacturer or the product caused or contributed to the adverse event.