Form Approved: OMB Number 0910-0025 Expiration Date: January 31, 2017

ACCIDENTAL RADIATION OCCURRENCE REPORT

See Burden Statement on page 5.

Note: Items with an asterisk (*) require a response.

		SUBMITTER	NFOR	MATION				
If you are not submitting this the problem, you may enter y and provide your home or oth	our own compa							
Contact Information								
Contact Name (Title, first name, last name)*				pation Title				
Email Address*								
Establishment Identificatior	n (Manufacturer	of the radiation-en	nitting p	product beii	ng reported, if known)			
Establishment Name								
Division Name								
Submitter Address								
Address					Telephone Number*			
Street*								
City*		State*	Zip Co	de*	Fax Number			
	INFORMATI	ON REGARDING	PRO					
Product Manufacturer Name (If	known)							
Product Manufacturer Address	(If known)							
Street (Line 1)			Stree	t (Line 2)				
City	Te	erritory, Province, or	State	ate Country Zip or Postal Code				
Product Model Designation (If k	known)	Model Name or Nu	mber		lel Family Designation	Brand Name		
Please provide any other inforradiation exposure incident.	ormation known	regarding the mai	nufactu	irer of the p	product that was involved	l in the accidental		

If you are aware that the manufacturer was informed about the incident, please provide the contact information below.

Contact Information (Including whom you contacted and address)

PRODUCT INFORMATION					
Product Types (Please select the best match (only one). Note that product types are grouped into radiation categories.)					
Acoustic Radiation	Microwave EMF Radiation (Continued)				
Therapeutic Ultrasonic Devices (Including diathermy and stimulators)	Microwave Identification, Safety, Security, and Surveillance Products				
Ultrasonic Medical Devices (Miscellaneous) (Including lithotriptors)					
Diagnostic Ultrasound Devices	Microwave Medical Products				
Sonic Medical Products (Including hearing aids and	Microwave Heating and Drying Products				
vibrators)	Microwave Communication, Data Transmit, and Measurement Products (Including CB radios, cell phones,				
 Ultrasound Non-Medical Products (Including jewelry cleaners and intrusion security systems) 	walkie-talkies, household remote controllers) Nuclear Magnetic Resonance Devices				
Sonic Non-Medical Products	 Household ELF Products (Including electric blankets) 				
Veterinary Diagnostic Ultrasonic Products	☐ Other Microwave Product				
Veterinary Therapy Ultrasonic Products					
Other Sonic or Ultrasonic Product	Optical Radiation				
Ionizing Radiation	Medical Laser Products (Including surgical devices and laser therapy)				
Personnel Security Systems (Including backscatter and transmission x-ray systems)	Surveying, Leveling, Alignment Laser Products (Including laser pointers, laser levels)				
Cargo Non-Intrusive Security Systems	Laser Light Show/Display Products				
Cabinet X-Ray Systems, Non-Medical (Including baggage	Toy, Novelty, Play Laser Products				
x-ray systems) Industrial X-Ray Systems (Excluding Cabinet)	Safety, Security, Surveillance Laser Products (Including night vision systems, traffic speed systems and intrusion				
Analytical X-Ray Systems, Non-Medical	detection systems)				
High Voltage Vacuum Switches	Research, Scientific, Laboratory Laser Products				
Industrial Particle Beam Systems	Material Processing Laser Products (Including welders,				
TVs and video monitors (<i>Not</i> including flat-screen TVs)	cutters, engravers) Data Measurement, Transmit, Control Laser Products				
Medical Diagnostic X-Ray Equipment	(Including fiber optic communication systems, laser vision				
Dental Diagnostic X-Ray Equipment	systems and process control systems)				
Therapeutic X-Ray Systems	 Utility/Peripheral Laser Products (Including laser printers, bar code scanners, CD and DVD systems) 				
Veterinary X-Ray Systems	☐ In Vitro and Other Medical Laser Products (Including				
X-Ray Bone Densitometers	Veterinary devices)				
X-Ray Film and Film Processing Materials	Patient Positioning Medical Laser Products				
Cabinet X-Ray Systems, Medical Medical Accelerators	Other Laser Products				
Non-Medical Accelerators	Sunlamp Products (Including sunlamps and tanning beds)				
High Voltage Vacuum Tubes	Mercury Vapor Lamps				
Cathode Ray Tube (Without Electronics Chassis)					
Cold-Cathode Gas Discharge Tubes	Ultraviolet Commercial/Consumer Products				
Other X-Ray Product	Ultraviolet Surveillance & Detection Products				
	Ultraviolet Hygiene Products (Including UV sanitizers)				
Microwave EMF Radiation	General Optical Products, Medical (Including surgical lamps)				
Microwave Ovens (Food Prep)	General Optical Products, Non-Medical (Including LEDs				
Microwave Hyperthermia Therapy Devices	and fluorescent lamps)				
Microwave Diathermy Machines					

Product Description

Description of product and its intended use

ACCIDENTAL RADIATION OCCURRENCE INFORMATION

Location of Occurrence

Please provide the physical location where the Accidental Radiation Occurrence took place (e.g., at a residence, a factory, a tanning salon, school, restaurant, airport, etc.). If you do not know the exact address, provide responses to the best of your ability, or enter "Unknown."

Location or Establishment Name

Specific	Section of	Location o	r Establishment	(If applicable)
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Address				Telepho	Telephone Number		
Street							
City State		State	State Zip Code		Fax Number		
Date of Event* From	То	To Web Address					
Persons Involved							
Number of people exposed in the Accidental Radiation Occurrence		of people y affected*	Number of unex people who wer		Number of potentially exposed people who have not exhibited any adverse reactions*		
Type of reportable event	Death	Serious Inju	ry 🗌 Malfunc	tion] Other		

Description of the nature and magnitude of exposure and/or injuries

ACCIDENTAL RADIATION OCCURRENCE INFORMATION (Continued)

Description of the Radiation Occurrence

Is this a new Accidental Radiation	Occurrence (ARO) r	eport or a supplement	to a previous ARO r	report filed by you or yo	ur organization?
(Please select one.)*					

New ARO report

Supplement to previous ARO report (Enter date of previous report below.)

Date of previous ARO report, if applicable (mm/dd/yyyy) (Required entry* only if "Supplement to previous ARO report" is selected.)

Description of circumstances surrounding the accidental radiation occurrence (Please include a description of the activities leading up to the event and actions that occurred during the event, as well as any suspected causes of the occurrence.)*

Actions Taken

The actions described below are those taken to control, correct, or eliminate the causes and to prevent reoccurrence. If unknown, you may state "Unknown" below.

Description of specific actions, to date, taken by the manufacturer in response to the Accidental Radiation Occurrence*

ACCIDENTAL RADIATION OCCURRENCE INFORMATION (Continued)

Actions Taken (Continued)

Description of future actions to be taken by the manufacturer, if known, in response to the Accidental Radiation Occurrence (If this is a preliminary ARO report from the manufacturer, please indicate that further investigation is ongoing.)*

If this involved a	medical device.	has a Medical Devi	ice Report (MDR)	been submitted to FDA?*

Yes No N/A Unknown

Other Important Information (Please enter below)

Feel free to send in medical documentation regarding the incident and injuries.

Please mail this completed FORM FDA 3649 to the address to the right:

U.S. Food and Drug Administration Center for Devices and Radiological Health Document Mail Center – WO66-G609 10903 New Hampshire Avenue Silver Spring, MD 20993-0002

This section applies only to requirements of the Paperwork Reduction Act of 1995.

DO NOT SEND YOUR COMPLETED FORM TO THE PRA STAFF EMAIL ADDRESS BELOW.

The burden time for this collection of information is estimated to average 2 hours per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden, to:

Department of Health and Human Services Food and Drug Administration Office of Chief Information Officer Paperwork Reduction Act (PRA) Staff *PRAStaff@fda.hhs.gov* "An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number."