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Submission Report

eRadHealth Menu

Introduction

Electronic Product Radiation Safety Reporting Form

This software application is intended to automate the hard copy product reporting forms in the effort of the Center for Devices and Radiological Health (CDRH) to become capable of accepting electronic submissions from industry and to improve our review process. This FDA Electronic Submission (eSub) software is the next version of the application developed to allow us to accept all Radiological Health reports and other submissions electronically and improve the ability of CDRH to accomplish its mandated product and industry evaluations in a timely and efficient manner.

All electronic reports and correspondence can either be transferred to CD and mailed to the address below, or can be sent via the FDA Electronic Submissions Gateway to CDRH. If you follow instructions to set up an account with the FDA Gateway, it currently may take several weeks,, but when you submit through it you will receive your acknowledgement email message with Accession Number within minutes! Or, in the interest of faster turn-around for a one-time urgent report of if you submit few reports, you may simply fill out this template creating the submission and then at 'Packaging' follow the instructions to transfer the files to a CD to mail in. This method of submitting your report will be acknowledged by an email with the Accession Number within several days.

Information about the FDA Electronic Submissions Gateway can be found at www.fda.gov/ForIndustry/ElectronicSubmissionsGateway/default.htm. Please contact the Gateway Helpdesk with your questions about that system.

Electronic submissions on CD should be mailed directly to the Document Control Center at:

U.S. Food and Drug Administration Center for Devices and Radiological Health Attn: eSubmitter Team Document Mail Center - WO66-0609 10903 New Hampshire Avenue Silver Spring, MD 20993-0002

Submissions received in the mail on CD will be processed within a few days of receipt.

Note about eSubmitter software:

Instructions provided in this software briefly summarize the requirements of the regulations under the Federal Food, Drug and Cosmetic Act (FFDCA), Chapter V, Subchapter C - Electronic Product

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Radiation Control, that applies to manufacturers of electronic products that emit radiation. The software provides questions relevant to requirements in the performance standards and may include explanations or clarification about the performance, labeling, and informational requirements of the standard. It does not replace the regulations, however, and if there is any conflict between the software and the regulations, the regulations must prevail. Throughout this application, pertinent sections of Title 21, Code of Federal Regulations, Chapter I, Subchapter J, are cited in parentheses. Please consult them before making design or procedural decisions.

Regulatory requirements for radiological products can be found at http://www.fda.gov/Radiation-EmittingProducts/default.htm and for medical devices are located at www.fda.gov/M/devaDvices/default.htm. If you have specific questions about the regulations, please contact us at: DSMICA@fda.hhs.gov.

If you have specific questions regarding this software, please contact the eSub team by email at: eSubmitter@fda.hhs.gov.

Thank you for using our electronic product reporting software. Please communicate your comments and suggestions to the eSub team as often as you like.

Thank you for your continued support of the FDA Electronic Submission Program (eSub).

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 control number. The OMB control number for this information collection is 0910-0025 (expires January 31, 2017).

Role

What is your role?

!* Manufacturer

Information:

The following screen provides several options for you to accurately define what type of eSubmission you intend to create for FDA. Below are explanations of your options. Please feel free to review this screen, advance to the next screen and view the picklists, but if you're confused, come back to read this screen again to be certain you are selecting the correct report or correspondence type you want to create.

Submission Information

Step 1	Use the radio buttons to identify the type of submission (Supplements should be prepared using the same does submission.)	
	f Submission is this? (Supplements should be submitted same document type as the original report.)	!* () Radiation Safety Report (Product) Report (21 CFR 1002.10) () Annual Report (21 CFR 1002.13) () Laser Light Show Documents (all relevant documents) (21 CFR 1040.11(c)) (•) Correspondence

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S (E N ()	(a) Variance Request (General, not Laser Light (Show) (21 CFR 1010.4) (b) Laser Original (Equipment/Component (Manufacturer Registration (21 CFR 1040.10(a)(3)(ii)) (c) Abbreviated Report (21 CFR 1002.12)
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Step 2	After answering the Submission Type question above, one of the questions below may become active and required (see the blue dot to the right of the question). If there is an active question, select the appropriate product area or document type from the question's pick list.		
What Type of P	roduct is this Radiation Safety Report about?		
What Type of P	roduct is this Annual Report about?		
What Laser Ligh	nt Show Document are you filing?		
What Type of C	What Type of Correspondence is this?		
Accidental Radi	ation Occurrence (21 CFR 1002.20)		
What Type of P	roduct is this Variance Request about?		

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Correspondence

Introduction

Information:

This section allows you to submit certain types of information or inquiries that are not part of a manufacturer's Product Report, Annual Report, or other reports as specified under 21 CFR 1002. However, some correspondence types would likely be submitted in conjunction with Product Reports. Examples of these would be Variance requests, Exemption requests, Laser Light Show notifications, follow-ups from FDA communications and audits, corrective actions, and notifications of product issues.

The following questions may seem a little too vague or not exactly appropriate to your situation but they are designed to be generic questions to suit many situations and issues. Please respond as well as possible and you have the opportunity to attach PDF letters or files if you like.

Burden to Industry

Paperwork Reduction Act Statement

Public reporting burden for this collection of information is estimated to average 2 hours per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, completing, and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to:

U.S. Food and Drug Administration Center for Devices and Radiological Health Attn: eSubmitter Team Document Mail Center - WO66-0609 10903 New Hampshire Avenue Silver Spring, MD 20993-0002

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Submitter Information

Message:	Please provide the following information regarding the submitter of this report. If you are not associated with a manufacturing establishment, enter N/A for Establishment Name on the Establishment Identification Tab. If you are associated with a Government Agency, please complete the Establishment Identification information.					
Copy from conta	Copy from contact address list					
Contact Informa	ation:					
Contact Name	Contact Name					
Occupation Title	9					
			_			

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Email Address				
Establishment Information:				
Establishment Name				
Division Name				
FDA Establishment Identifier (FEI)				
Central File Number (CFN)				
Registration Number				
Owner/Operator Number				
D&B D-U-N-S Number				
Physical Location:				
Address				
Telephone Number				
Fax Number				
Mailing Location:				
Address				
Telephone Number				
Fax Number				

Manufacturer Information

Message:	Please provide any information known regarding the manufacturer of the product being reported.			
Copy from contact address list				
Contact Informa	Contact Information:			
Contact Name	Contact Name			
Occupation Title	;			
Email Address	Email Address			
Establishment li	nformation:			
Establishment N	lame			
Division Name	Division Name			
FDA Establishm	FDA Establishment Identifier (FEI)			
Central File Nun	Central File Number (CFN)			
Registration Nu	Registration Number			
Owner/Operator Number				
D&B D-U-N-S N	D&B D-U-N-S Number			
Physical Location	Physical Location:			
Address	Address			
Telephone Num	Telephone Number			
Fax Number	Fax Number			
Mailing Location	Mailing Location:			
Address				

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Telephone N	lumber				
Fax Number					
			,		
Product In	formatio	on			
Note:	Each product that CDRH regulates is assigned a product code by CDRH. Please provide the following information regarding the product being reported.				
	•				
-Enter a key choose. (If you are no - Select the l - The remain	ter search word to so to finding best mate ning fields to find the	h icon (next to the earch the databa the correct produc the to your produc will be filled in for e code that you a	ise. You w uct, try oth it. or you whe	n). You will see a product coo vill be provided a list of produc- ner words and/or variations of en you select your product co- for, use RZZ (Other)	the keywords.)
Item	Category			Product Code	Performance Standard
Details					
Describe the	product	and its intended	use. Attac	ch any supporting documents	if necessary.
Details					

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Accid	Accidental Radiation Occurrence				
A DO 1	- (('				
ARO II	ntroduction				
Note:	Note: Accidental radiation occurrences (AROs) must be reported to CDRH by manufacturers (pursuant to 21 CFR 1002.20), regardless of whether injury occurred, or it was a situation presenting the potential for injury, or involving a product malfunction. The report must include identification of the product involved, the circumstances and details of the incident, and actions taken to prevent recurrence. If your product is a medical device, an adverse event must be reported using the MedWatch form and following the Medical Device Reporting regulations explained on the following website: http://www.fda.gov/cdrh/mdr/.				
Model	Designation	un.			
IVIOGEI	Designation	·11			
Note:	Note: Report the model name and/or number, model family, brand name, or other designation of the product. If reporting a model family, provide the model designation of each model. If you do not have a model family or brand name, leave the field blank.				
		Names and/or Numbers	1	*	
Item	Model Nam	e	Family Name	Brand Name	
Descri	ption of the	Radiation Occurre	nce		
		ntal Radiation Occurren vious ARO report?	ice (ARO) report or a	*	
What wa	as the date o	f the previous ARO rep	ort?		
person(s)' actions wh	nen exposed to the radi	ding the accidental radiation occu ation, and causes of the occurrer king on the Add button below.		
Details					
	If this involves a medical device, has a Medical Device Report (MDR) been submitted to FDA?				
1 0	(O				
Location	on of Occu	rence			
residend address zeroes i	ce, a factory, s, provide res if unknown.	a tanning salon, schoo ponses to the best of y	the Accidental Radiation Occurre I, restaurant, airport, etc). If you do our ability, or enter Unknown. For	lo not know the exact	
Establis	Establishment Name				
Address	3				
1		I			

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Telephone Nur	mber
Please give sta same.)	art and end dates of event. (They may be the *
Persons Invo	olved
Please list the of Occurrence.	number of people exposed in the Accidental Radiation *
Please list the	number of people adversely affected. *
	number of potentially exposed people who have not * adverse reactions.
Please list the	number of unexposed people who were involved. *
Type of reporta	* [] Death [] Serious Injury [] Malfunction [] Other
If Other, spec	cify the the type of event.
Please list the	nature and magnitude of exposure and/or injuries.
Details	
Are the affected	d person(s) employees of the product manufacturer? *
Did the affected	d person(s) have any responsibility toward the operation of the equipment?
Remarks	
A (: -)	
Actions Take	en e
	to control, correct, or eliminate the causes and to prevent reoccurrence. If unknown, e as such in your response.
Please list the a Occurrence.	actions, to date, taken by the manufacturer in response to the Accidental Radiation
Details	
Please list futur Occurrence.	re actions to be taken by the manufacturer in response to the Accidental Radiation
Details	
Other Import	tant Information
Please list any	other pertinent information and/or attach a file.
Details	
Stop:	You have reached the end of this report. Please verify that all PDFs that are to be included in this submission are correctly attached to a specific file attachment question. Otherwise, they will not be packaged with your report. Check to make sure you have no missing data (select

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Missing Data Report from the Output menu). Once you have confirmed that there is no missing data and all your files are attached, click on the Package Submission icon on the tool bar.