# **Submission Report**

Section: Main Menu

Welcome

# Welcome to the CDRH Electronic Submissions Software (CeSub)

This software application is intended to automate the current paper submission process. This software contains a number of data capturing tools and helpful dialog boxes to reduce redundant responses for you, and to allow us to capture data in a more useful, structured format. These benefits will enable CDRH to improve our review process and reduce lengthy review times.

For your convenience, an email account has been established to support any questions that you may have regarding the use of this software. Please email any questions or comments to the CeSub team at: **cdrhesub@cdrh.fda.gov.** Please be sure to include your name, company name and contact information in the email.

Thank you again for using our electronic product reporting software. We look forward to hearing from you soon.

What type of product is this submission referring to?

Radiation Emitting Product (OMB No. 0910-0025; Expiration Date: December 31, 2006)

#### Welcome (Cont.)

Department of Health and Human Services Food and Drug Administration

Form Approved:
OMB Number 0910-0025
Expiration Date: December 31, 2006

Section: eRadHealth Menu

Role

What is your role?

\* Manufacturer

#### Submission Information

What Type of Submission is this? (Supplements should be submitted selecting the same document type as the original report.)

Correspondence

What Type of Product is this Annual Report about?

What Type of Correspondence is this?

Government Agency's Exemption Request

What Type of Product is this Radiation Safety Report about?

What Type of Product is this Variance Request about?

What Laser Light Show Documents are you filing?

#### **Section: 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 1 hour 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:

Food and Drug Administration CDRH (HFZ-240) 1350 Piccard Drive Rockville, MD 20850

Please DO NOT RETURN this application to this address.

"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."

## 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 contact address list				
Contact Information:				
Contact Name				
Occupation Title				
Email Address				
Establishment Information:				
Establishment Name				
Division Name				
FDA Establishment Identifier (FEI)				
Central File Number (CFN)				
Registration Number				
Owner/Operator Number				
Physical Location:				
Address				
Telephone Number				
Fax Number				
Mailing Location:				
Address				

## Manufacturer Information

Message:	Please provide any information known regarding the manufacturer of the product being reported.			
Copy from contact address list				
Contact Information:				
Contact Name				
Occupation Title				
Email Address				
Establishment Information:				
Establishment Name				
Division Name				
FDA Establishment Identifier (FEI)				
Central File Number (CFN)				
Registration Number				
Owner/Operator Number				
Physical Location:				

Address						
Telephone Number						
Fax Number						
Mailing Location:						
Address	Address					
Product Infor	matior	า				
-						
Message:	Pleas	se provide the following information regarding the product being reported.				
What product typ	e is bein	g reported?	*			
Note:	Each	product that CDRH	regulates is assigned a product code by CDRH.			
What is the produ	ıct code	?				
If you know the ti	ree lette	er code, enter it in th	e space provided.			
If you do not,						
	aarch icc	on (next to the trach	can). You will see a product code filter dialog box.			
-Enter a keyword	to searc	ch the database. You	will be provided a list of product codes from which to	choose.		
- Select the best	match to	your product.	other words and/or variations of the keywords.)			
			vhen you select your product code. ng for, use RZZ (Other)			
Product Code						
Device Class						
Classification Panel						
C.F.R. Section						
Describe the pro-	duct and	its intended use. At	tach any supporting documents if necessary.			
File Attachment						
Details						
Section: C	orre	spondence	Details			
Model Desig	nation					
Item: 1						
Model Family De	signatior	ո:				
Model Designation	n (Name	es and/or Numbers)				
ltem						
Brand or Trade Names, if different:						
Item						

#### Additional Information

Please provide an explanation of your concerns, your questions, and/or your requests below.

File Attachment			
Details			
Error:	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 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.		
Message:	Form	m FDA 3642 General Correspondence (03/06)	