OMB N	o. 0910-0025; Exp. May 31, 2010
Section:	: eRadHealth Menu
Role	
What is your i	role?
Note:	If you are acting as an agent of the actual manufacturer, please select your role, for example, Importer or Consultant. Later in the report, under Manufacturer Data, you will be prompted to enter both manufacturer and submitter information.
Submissio	on Information
FDA or Sta	ate Inspector
Abbreviate	ed Report Applicability
OEM Lase	er Applicability
Section:	: Manufacturer Data
Introduction	

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, when you submit through it you will receive your acknowledgement email message with Accession Number within minutes!

Information about the FDA Electronic Submissions Gateway can be found at <a href="www.fda.gov/esg">www.fda.gov/esg</a>. 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.

You should be familiar with the regulatory requirements for radiological products at <a href="www.fda.gov/cdrh/radhealth/">www.fda.gov/cdrh/radhealth/</a> and medical devices available at <a href="www.fda.gov/cdrh/devadvice/">www.fda.gov/cdrh/devadvice/</a>. If you have specific questions about the regulations, please contact us at: <a href="DSMICA@fda.hhs.gov">DSMICA@fda.hhs.gov</a>.

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).

General Information

# General Information for Radiological Health Products

Manufacturers of products subject to performance standards under the Federal Food, Drug, and Cosmetic Act (FFDCA), Chapter V, Subchapter C - Electronic Product Radiation Control are required to furnish various reports to the Center for Devices and Radiological Health (CDRH).

The Radiological Health staff, CDRH developed this software application for the Product and Annual reports. This application will assist manufacturers of electronic products that emit radiation in providing adequate reporting of radiation safety testing and compliance with federal performance standards. Title 21 of the Code of Federal Regulations (CFR), Parts 1002 and 1003 specify Reporting and Notification requirements 1,2,3.

Reports submitted on radiation safety of electronic products must follow the appropriate form (21 CFR 1002.7). This software application serves the same report responsibility, so long as the submitter or manufacturer prints out the cover letter and sends it in along with the CD containing the report files. The submitter of the report will receive an acknowledgment letter (or email message) with the

accession number that CDRH assigns to the report. Please reference this accession number in the future when providing additional information about this model family in either a supplement or the annual report. If a report is incomplete or inadequate CDRH may reject it and return it for completion. CDRH will not enter a rejected report into our database.

#### CDRH DOES NOT APPROVE THESE REPORTS OR THE PRODUCTS BEING REPORTED.

It is the manufacturer's responsibility to certify that their products comply with all applicable standards (21 CFR 1010 - 1050), based on a testing program in accordance with good manufacturing practices. Prior to the shipment of products in interstate commerce, 21 CFR 1002 requires the manufacturer to submit the product and Annual Reports and to comply with all applicable importation requirements (21CFR 1005). If there are deficiencies, CDRH may disapprove the firm's quality control and testing program, determine that the product contains a radiation defect, or determine that the product fails to comply with a standard. CDRH will notify the manufacturer if we make such a determination. CDRH may require the manufacturer to cease introduction into U.S. commerce until deficiencies are corrected, and to initiate a corrective action program (21CFR 1003 - 1004) for products already introduced into commerce.

CDRH can now accept and process 'CeSub' electronic submissions at this time, if all attachments are PDF files only, and the cover letter is printed out and included with a real signature. Translate any text that appears in a language other than English into English in a complete and accurate manner. Keep a copy (save a copy to your hard drive) of the completed report in your records.

We are providing our new software applications for the old reporting forms upon request during this beta testing period of development in Spring, 2005. Other regulatory information is still available on the Internet under <a href="https://www.fda.gov/cdrh/radhealth/">www.fda.gov/cdrh/radhealth/</a>. No copyright exists for these forms.

Reproduce these forms as needed. If you would like to comment on the reporting forms, website, or future electronic submissions, you may direct the comments to **cdrhesub@cdrh.fda.gov**.

A complete Product Report is required for each product model or model family. Product Reports are now more generally referred to as Radiation Safety Reports to distinguish the Radiological Health submissions from medical device submissions. CDRH suggests that a complete report on one model of a family be submitted, with a separate Supplemental Report for each of the other models in the family. The Supplemental Report should respond in detail to the parts of the form where there are differences to report, referencing the number of the affected item. Items that are unchanged will still appear in the supplement from the original report.

When new models of a product are introduced, if the models satisfy the criteria for an established reporting exemption or if the new models do not involve changes in radiation emission or performance requirements, then the manufacturer need not report the models prior to introduction into commerce. Rather, the manufacturer is only required to identify them in the annual report, or in quarterly updates to the annual report. Quarterly updates to annual reports may be submitted using the Annual Report software included in this application. [See 21 CFR 1002.13(c).]

All symbols, units, and unusual terms in the report must be adequately defined and consistently used. Please use the terms as defined in Section 1040.10(b) and in the IEEE Standard Dictionary of Electrical and Electronic Terms (IEEE Std. 1001972 and ANSI C42.1001972).

**Definitions** 

## **Definitions for Rad Health Products**

#### **Manufacturers**

Manufacturer is any person or organization engaged in the business of manufacturing, assembling, or importing of electronic products (21 CFR1000.3(n)). Manufacturers of electronic products subject to 21 CFR1000-1050 must:

- Design and manufacture their products to be in compliance with applicable performance standards;
- Test their products to assure compliance;
- Certify compliance of their products;
- Maintain test and distribution records and a file of correspondence concerning radiation safety, safety complaints, and inquiries;
- Use the published reporting forms or electronic software application to submit reports to CDRH, including Product reports describing the manner of compliance of the product design and testing program and Annual Reports summarizing their compliance testing;
- Report accidental radiation occurrences (i.e., possible, suspected, or known exposures);
- Report any radiation defects or noncompliances; and
- Recall (i.e., repair, replace, or refund the purchase price of) defective or noncompliant products.

#### **Accidental Radiation Occurrences**

An accidental radiation occurrence means a single event or series of events that has/have resulted in injurious or potentially injurious exposure of any person to electronic product radiation as a result of the manufacturing, testing, or use of an electronic product.

## **Importers**

Importer is any person of organization engaged in the business of importing electronic products. An importer is considered to be a manufacturer. The requirements for Manufacturers given above also apply to importers if the requirements have not been done by the foreign manufacturer.

## **United States Agent for Foreign Manufacturers**

Every manufacturer of electronic products, prior to offering such product for importation into the United States, shall designate a permanent resident of the United States as the manufacturer's agent upon whom service of all processes, notices, orders, decisions, and requirements may be made for and on behalf of the manufacturer as provided in section 536(d) of the Radiation Control for Health and Safety Act of 1968 (21U.S.C. 360mm(d)) and this section. The agent maybe an individual, a firm, or a domestic corporation. For purposes of this section, any number of manufacturers may designate the same agent.

# From The Federal Food, Drug, and Cosmetic ActSec 536 [21 U.S.C. 360mm](d) Designation of agent for purposes of service

It shall be the duty of every manufacturer offering an electronic product for importation into the United

States to designate in writing an agent upon whom service of all administrative and judicial processes, notices, orders, decisions, and requirements may be made for and on behalf of said manufacturer, and to file such designation with the Secretary, which designation may from time to time be changed by like writing, similarly filed. Service of all administrative and judicial processes, notices, orders, decisions, and requirements may be made upon said manufacturer by service upon such designated agent at his office or usual place of residence with like effect as if made personally upon said manufacturer, and in default of such designation of such agent, service of process, notice, order, requirement, or decision in any proceeding before the Secretary or in any judicial proceeding for enforcement of this part or any standards prescribed pursuant to this part may be made by posting such process, notice, order, requirement, or decision in the Office of the Secretary or in a place designated by him by regulation.

Sec. 531 [21 U.S.C. 360hh] (1) the term **"electronic product radiation"** means:

- (A) any ionizing or non-ionizing electromagnetic or particulate radiation, or
- (B) any sonic, infrasonic, or ultrasonic wave, which is emitted from an electronic product as the result of the operation of an electronic circuit in such product.

Sec. 531 [21 U.S.C. 360hh](2) the term **''electronic product''**means:

- (A) any manufactured or assembled product which, when in operation,(i) contains or acts as part of an electronic circuit and (ii) emits (or in the absence of effective shielding or other controls would emit) electronic product radiation, or
- (B) any manufactured or assembled article which is intended for use as a component, part, or accessory of a product described in clause (A) and which when in operation emits (or in the absence of effective shielding or other controls would emit) such radiation.

Burden to Industry

# **Paperwork Reduction Act Statement**

Public reporting burden for this collection of information is estimated to average 26 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 Document Mail Center - WO66-0609 10903 New Hampshire Avenue Silver Spring, MD 20993-0002

"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."

#### Manufacturer and Report Information

#### Information:

This general report requests names, addresses, phone numbers, etc. for your firm, various officials of your firm, consultants who may assist in preparing the report, parent firm (if any), importer and designated agent (for foreign firms). Some of this information is mandatory and its absence will prevent you from completing the report submission. You can check for missing data using the "Missing Data" report from the "Output" menu.

If you are acting as an agent or consultant for another firm who is certifying the product (or laser light show), please enter the certifying manufacturer and list yourself as the report submitter, below.

#### Information:

Attention: Variance Applicants

If you are acting as an agent or consultant for, or on behalf of, or filing for, a company that will be manufacturing or producing a Class IIIb or IV projector or laser light show or both which require an approved variance, the following explanations may provide further clarification.

Manufacturer: This is the firm or company who is requesting the variance, will certify the product or show, and will be the holder and owner of the variance. This is not the agent or consultant who may be filing this report or Variance request for the manufacturer; that agent may be the submitter, identified in a later screen.

Responsible Individual: This person works for the Manufacturer and is responsible for compliance of the projector and/or show. In the case of laser light shows, he or she may be the company president, CEO, or the laser light show head operator or a manager who oversees the shows.

Reporting Official: This person works for the Manufacturer and is responsible for reports, recordkeeping, and submitting FDA required documents and correspondence.

#### Manufacturer Responsible for Product Compliance

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This is the firm that takes responsibility for certification that the product meets the performance standard. This firm develops and maintains the quality control and testing program that is the basis for the certification of this product. Additionally, this firm usually is the owner of the product design and manufacturing process design.

Select the Manufacturer's address from the Establishment Address book:		
Establishment Information:		
Establishment Name		
Division Name		
Home Page		
Physical Location:		
Address		
Telephone Number		
Fax Number		
Mailing Location:		
Address		

#### Responsible Individual

Note:

The responsible individual is the highest level and most responsible individual affiliated with this establishment.

Select the Responsible Individual from the Contact Address book:

Contact Information:		
Contact Name		
Occupation Title		
Email Address		
Establishment Informa	ation:	
Establishment Name		
Division Name		
Physical Location:		
Address		
Telephone Number		
Fax Number		
Mailing Location:		
Address		
Manufacturer's F	Report	ting Official
Note:	and qu	s the person at the manufacturing facility that is knowledgeable and responsible for addressing all aspects of the testing uality control procedures for certification as reported to FDA in the product report. Documentation of changes intesting uality control procedures submitted to FDA must be signed by this individual.
Select the Reporting 0	Official f	from Contact Address book:
Contact Information:		
Contact Name		
Occupation Title		
Email Address		
Establishment Informa	ation:	
Establishment Name		
Division Name		
Physical Location:		
Address		
Telephone Number		
Fax Number		
Mailing Location:		
Address		
Report Submitte	r	
Note:		ubmitter maybe a consulting individual or firm providing assistance in report preparation and maintenance. All nents prepared by the submitter must have the manufacturer's reporting official signature for authenticity of submitted

	docume	entation.	
Select the Submitter from	om the	Contact Address book:	
Contact Information:			
Contact Name			
Occupation Title			
Email Address			
Establishment Informat	tion:		
Establishment Name			
Division Name			
Physical Location:			
Address			
Telephone Number			
Fax Number			
Mailing Location:			
Address			
Comments:			
Internal Reference Nun	mber:		
Parent Establishn	nent		
Is there a parent establ	lishmer	12	[L]
is there a parent establish			
Select the Parent Estab	blishme	nt and Contact from the Contact Address book:	
Contact Information:			
Contact Name			
Occupation Title			
Email Address			
Establishment Informat	tion:		
Establishment Name			
Division Name			
Physical Location:			
Address			
Telephone Number			
Fax Number			
Mailing Location:			

Address					
Manufacturar Designated United States Agent					
Manufacturer Designated United States Agent					
Note:	Note: Manufacturers exporting to the U.S. must designate a U.S. agent, see 21 CFR 1005.25.				
Is there a United State	es agen	t that has been designated by the manufacturer?	[L]		
Written Agreeme	ent				
Item: 1 (could contain	in up to	10 items with none required)			
	1				
Note:		of the required responses below do not apply to your designated agent, enter 'NOT APF	PLICABLE' or 'NA.'		
	d Agent	from the Contact Address book:			
Contact Information:					
Contact Name					
Occupation Title					
Email Address					
Address					
Establishment Name  Division Name					
	Address				
Telephone Number Fax Number					
	Fax Number  Attach a copy of written agreement with the designated U.S. agent:				
[Multi-Line Plain Text]		sherit with the designated 0.5. agent.			
		[Single File Attachment ( ndf ing gif tif avi wmv ynt yml dtd sgml mol yls	csv zin)]		
The Attachment	File Attachment [Single File Attachment (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]				
Importer					
Item: 1 (could contain up to 10 items with none required)					
Select the Importer from the Contact Address book:					
Contact Information:					
Contact Name					
Occupation Title					
Email Address					
l					

Establishment Inform	ation:				
Establishment Name					
Division Name	Division Name				
Physical Location:	Physical Location:				
Address					
Telephone Number					
Fax Number					
Mailing Location:					
Address					
Additional Manu	factur	ing Locations			
Item: 1 (could conta	in up to	100 items with none required)			
Note:	Produc codes proced	If any of the products certified in this report are manufactured at locations other than listed in the Manufacturer Responsible for Product Compliance section, then the names, addresses, and FDA registration numbers should be provided. In addition any codes used on labels to identify a manufacturing location must be provided. Each factory location must assure all production procedures are followed identically step by step as provided in this report. If the procedures are not the same then separate reports should be filed.			
Select the Manufactur	rer Addı	ess from the Establishment Address book:			
Establishment Inform	ation:				
Establishment Name	Establishment Name				
Division Name					
Home Page					
Physical Location:					
Address					
Telephone Number					
Fax Number					
Mailing Location:					
Address	Address				
Comments:	Comments:				
Code used on identification labels:					
•					
Section: Pro	Section: Product Data				
Product and Mod	Product and Model Identification				
. 75 dast and Wo					

At this time we are only accepting electronic versions of reporting guides contained within this software. Other reporting Note: guides that are not yet electronic are available for downloading from http://www.fda.gov/cdrh/comp/eprc.html. Product Type Reported Report Information Is this submission a supplement to an Annual Report submitted previously for the same reporting year? [L] Provide the Accession Number of the original report for which this is a supplement: (Note: Do not enter any Device Premarket Application or Notification document number here, such as PMAs, 510(k)s, IDEs, Please verify that your accession number matches the report type that is being filed. The third character of your accession number must correspond with its associated report type as shown in the table below: Third Character: **Report Type Description: Initial Product Report** Model Change Product Report 3 Annual Report 8 Abbreviated Report Variance Request Α R Laser OEM Registration and Listing Report [L] Are you requesting a new variance, a renewal, extension or amendment to a previous variance? If you are requesting a renewal, extension, or amendment, please provide the variance number that was issued by CDRH. If you are requesting a new variance, renewal, extension, or amendment, you must file a Variance Request separate from this Stop: report. To do this, open a new report (File > New) and select either "Laser Light Show Variance Request" or "Variance Request, Other" as your Type of Submission in the Submission Information Screen. If you select "Variance Request, Other" you must select the product for which you are requesting a variance at the end of the screen. Special Considerations Note: Check all items in this section that may apply to this submission. Noncompliances or Defects Does this document or any of its attachments contain: [L] A self-declaration or notification of noncompliance or defect? Provide an explanation:

[Multi-Line Plain Text]

## Responses to Noncompliances or Defects

Does this document or any of its attachments contain and of these responses concerning noncompliances?		
A refutation of noncompliances?	[L]	
A request for an exemption from notification?	[L]	
Corrective action plans you may be conducting?	[L]	
A description of any design changes that correct noncompliances for future production?	[L]	
Provide an explanation:		
[Multi-Line Plain Text]		

## **Exemption Requests**

Does this document of	r any of its attachments contain:	
Exemption of a product	for government use from a standard (1010.5)?	[L]
Exemption for products	for government use from reporting and recordkeeping (1002.51)?	[L]
Special exemption of pr	oducts from reporting and/or recordkeeping (1002.50)?	[L]
Request for approval of alternate labeling? [L]		
Application for alternate test procedures (1010.13)? [L]		
Provide an explanation:		
[Multi-Line Plain Text]		
Attach any necessary fi	les.	
[Multi-Line Plain Text]		
File Attachment	[Single File Attachment (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)	]

## Variance Requests

Message:	essage: Click the plus sign to list the requirements from which you are requesting a variance.				
This submission inclu	This submission includes an application for a variance from certain requirements.				
Item 1					
Item 2					
Item 3					
Provide an explanation and attach supporting files, if necessary. Click on the plus sign below to attach files.					
Details	[HTML Text]				

File Attachment		[Multiple File Attachments (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]
Stop:	For all The eletthe Me U.S. F. Center Attn: e Docum 10903 Silver Addition Food a Divisio 5630 F	Variance requests, two submissions must be made to the FDA.  ectronic version should be submitted following the Packaging Files for Submission instructions located under Output in enu bar, and explained in subsection 4.3 of the User Manual. If sending a CD & submittal letter, please mail to:  food and Drug Administration or for Devices and Radiological Health Submitter Team nent Mail Center - W066-0609 New Hampshire Avenue Spring, MD 20993-0002  ponally, a paper version (hard-copy) of the signed Variance request document should be submitted to:  and Drug Administration on of Dockets Management (HFA-305) Fishers Lane, Room 1061
	Rockvi	ille, MD 20857

## Responses to Communications from FDA

Does this document or any of its attachments contain:			
A response to an inspection?	[L]		
What was the date of the inspection?	[Date]		
A response to a warning letter from the Food and Drug Administration (FDA)?	[L]		
What was the date of the Warning Letter?	[Date]		
A response to a report review inquiry from the Center for Devices and Radiological Health (CDRH) (the inquiry may have been in the form of a letter, email, or phone call)?	[L]		
What was the date of the inquiry?	[Date]		
A response to any other communication from FDA?	[L]		
What was the date of the communication?	[Date]		
Provide an explanation:			
[Multi-Line Plain Text]			

## Additional Information

Is there any other relevant information or additional comments that would help expedite the review of this submission? Click the plus sign below to		
attach any supporting files.		
File Attachment [Multiple File Attachments (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]		
Details	[HTML Text]	

## Private Labeling

Is the product sold by other companies under different brand names?	[L]
---	-----

Private Labeling-Table				
Item: 1 (could contain up to 20 items with 1 required)				
	The state of the s			
Give the name and address	of the manufacturer:			
Establishment Information:				
Establishment Name				
Division Name				
Email Address				
Address				
Address				
Telephone Number				
Fax Number				
Give the firm establishment r	egistration number of the manufacturer listed above (if known):			
Enter brand names and/or m	odel designations in the following table by clicking on the Add button. If	you prefer to attach a file please click on the		
	"See File Attachment" as the first table entry.	you prefer to attach a file, please click on the		
Item 1				
Item 2				
Item 3				
List of Brand Names and/or N	Model Designations			
File Attachment	[Single File Attachment (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .	sgml, .mol, .xls, .csv, .zip)]		
Details	[HTML Text]			
The Original Equipment Man	ufacturer (OEM) accession number (if known):			
Explain how the brand name	s and model designations correspond with your own brand names and r	nodel designations:		
[Multi-Line Plain Text]				
Medical Devices				
Provide the premarket 510(k), IDE, HDE, PDP, or PMA filing numbers related to this medical product, if one of these numbers has been assigned by				
FDA yet.  [Multi-Line Plain Text]				
If it has not been assigned yet, provide an explanation and submit it as soon as you receive such a filing number.				
[Multi-Line Plain Text]				
Γ				

Note: See www.fda.gov/cdrh for more information onmedical device premarket clearance procedures.

Document Key: Specialized Response content is defined within straight brackets []; Special code: [L] List of Values.

OMB No. 0910-0025; Exp. May 31, 2010							
Section	Section: Ultrasonic Therapy Product						
000	JII. CIL.	4301110 1110.4	<u> </u>				
Model [	Designati	on					
Note:		•		•	_	•	oduct. If reporting a model name, leave the field blank.
Model De	signation (N	lames and/or Numbers	;):				
Item	Model Na	me		Family Name		rand Name	
Item 1							
Item 2	<u> </u>						
Item 3							
Produc	t Classific	cation					
Indicate I	halow the ti	me of product or fam	ally of product:	s covered by this repor	+		
		/pe or product or la	ily of product	5 COVERED BY MILE TOPO.		[L]	
	This report covers:  [L]  Is your system a continuous-wave (CW) unit or an amplitude-modulated (AM) unit?  [L]						
is your system a continuous-wave (Ovv) unit of an amplitude-modulated (Alvi) unit:							
Perform	Performance Specifications						
Note:  The information reported in this section will be used to determine whether the product complies with the requirements set forth in 21 CFR 1050.10(c). Several items must be reported in terms of definitions that are provided by the standard; please refer to 21 CFR Part 1050.10 for these definitions.							
Freque	ncy of Op	peration					
Note:		Provide the following	data for operat	ting frequency(ies) in MH:	z for either fixed, variab	le, or multipl	e frequency systems.
Fixed at _	MHz:						
MH	łz:						
Variable,	from	MHz to MHz:					r
Minimum	MHz:						
Maximum	MHz:						
Multiple, fixed at MHz, MHz, and MHz:							
MH	łz:						

MHz:			
MHz:			
Operating frequency (	Operating frequency (ies) are indicated to the user by:		
Output Paramete	ers for Continuous-Wave Units		
Temporal-average u	Itrasonic power:		
Message:	Variable, from Watts to Watts:		
Minimum Watts:			
Maximum Watts:			
Indicated to the user b	ру:		
Temporal-average ef	ffective intensity:		
Message:	Variable, from W/cm² to W/cm²:		
Minimum W/cm <sup>2</sup> :			
Maximum W/cm²:			
Indicated to the user b	ру:		
Output Parameters for Amplitude-Modulated Units			
Temporal-maximum ultrasonic power:			
Message: Variable, from Watts to Watts:			
Minimum Watts:			
Maximum Watts:			
Indicated to the user by:			
Temporal-maximum effective intensity:			
Message:	Message: Variable, from W/cm² to W/cm²:		
Minimum W/cm <sup>2</sup> :	Minimum W/cm²:		
Maximum W/cm²:			
Indicated to the user by:			
Output pulse width:			
Message: Fixed at milliseconds:			
milliseconds:			
Message:	Message: Variable from milliseconds to milliseconds:		
Minimum milliseconds	Minimum milliseconds:		
ı			

Maximum millisecond	Maximum milliseconds:			
Indicated to the user by:				
Output pulse repetit	ion rate:			
Message:	lessage: Fixed at pulses/second:			
pulses/second				
Message:	Variable, from to pulses/second:			
Minimum pulses/seco	nd:			
Maximum pulses/seco	ond:			
Message:	User selected from the available settings (list available settings):			
Setting 1:				
Setting 2:				
Setting 3:				
Indicated to the user b	ру:			
Timer Specificat	ions			
Timer accuracy for s	settings of:			
Less than 5 minutes:				
Provide the number of minutes: (+/ minutes)				
Between 5 and 10 minutes:				
Provide the number of minutes as a percent: (+/ percent)				
Greater than 10 minutes:				
Provide the number of minutes: (+/ minutes)				
Maximum timer setting: ( minutes)				
How does ultrasonic emission automatically terminate at the end of preset time?  [HTML Text]				
How can ultrasonic emission be terminated prior to the end of the preset time?				
[HTML Text]				
How is radiation emission routinely terminated?				
[HTML Text]				
Applicators				
Type of applicators:				
Type of applicators:				

Collimating, with an e	ffective i	radiating area (ERA) of cm²:		
Diverging, with an effe	ective ra	diating area (ERA) of cm²:		
Focusing, with focal a	area of _	cm² and a focal length of cm:		
Transducer Configu	ration:			
Single Crystal (specify	y materia	al):		
For multiple elements	s, describ	be each element, the manner in which connected, and the resulting effe	ect on the radiated field.	
[HTML Text]				
Cables				
	electrica	I power to the transducer indicated to the user?		
[HTML Text]				
	e or oper	n connection indicated to the user?		
[HTML Text]				
Labeling Require	ement			
Labeling Require	CITICITA			
Note:	in 21 C	information reported in this section will be used to determine whether the product complies with the requirements set forth 1 CFR Parts 801, 1010.2, 1010.3 and 1050.10(d). Most of the items below require that a copy of the label be attached; if els are unavailable at the time of reporting, please provide a specification control drawing.		
	idbolo (	are unavaliable at the time of reporting, preade provide a openioanon	onto diamig.	
Certification				
Note:	label ce	ort 1010.2 of 21 CFR requires that the product (generator and applicator, if detachable) bear a permanently affixed tag or one certifying that it complies with the provisions of Part 1050.10. Provide the following information concerning the rtification label:		
The manner in which	the labe	l is attached:		
[HTML Text]				
The location of the lab	bel:			
[HTML Text]				
Attach a sample of the	e label.			
File Attachment		[Single File Attachment (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .	sgml, .mol, .xls, .csv, .zip)]	
Details	Details [HTML Text]			
Identification				
Note:	label gi other th may be	110.3 of 21 CFR requires that the product (generator and applicator, if of ving the following information:(a) The name and address of the manufator that of the manufacturer, the name and address of the individual or given on the label, provided that such individual or company has preves of the manufacturer.)(b) The place, month, and year of manufacture.	acturer. (Where the product is sold under a name company under whose name the product is sold iously supplied the CDRH with the name and	

	orm if the manufacturer has previously supplied the CDRH with the codes and their meaning). The month and year of nanufacture must be given without abbreviation and with the year as a four-digit number (for example: Manufactured: September 1978.)		
Message:	Provide the following information concerning the identification label:		
The manner in which the	e label is attached:		
[HTML Text]	[HTML Text]		
The location of the label:			
[HTML Text]			
Attach a sample of the label.			
Attach a sample of the	abel.		
Attach a sample of the File Attachment	abel.  [Single File Attachment (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]		

## Generator Labels

Note:	Part 1050.10(d)(3) of 21 CFR requires that each ultrasonic therapy generator bear a label giving the following information:		
	(a) The brand name, model designation, and serial number of the generator.		
	(b) The ultrasonic frequency (unless variable, and indicated on the controls).		
	(c) The type of waveform (continuous wave or amplitude modulated).		
	In addition to the above, generators employing amplitude modulated waveforms are required to bear additional labeling giving the following information:		
	(a) Pulse duration and repetition rate (unless variable, and indicated on the controls).		
	(b) An illustration of the waveform.		
	(c) The ratio of the temporal-maximum effective intensity to the temporal-average effective intensity. If this ratio is a function of any operation control setting, then the range of the ratio shall be given, and the waveform illustration shall be for the maximum value of this ratio.		
Message:	Provide the following information concerning the generator label:		
The manner in which the label is attached:			
[HTML Text]			
The location of the label:			
The location of the lab	el:		
The location of the lab	el:		
[HTML Text]			

## **Applicator Labels**

Part 1050.10(d)(4) of 21 CFR requires that each ultrasonic therapy applicator bear a label giving the following information:(a) The brand name, model designation, and serial number of the applicator.(b) Thedesignation of the generator for which the applicator is intended.(c) The ultrasonic frequency, effective radiating area, maximum beam nonuniformity ratio, type of applicator (focusing, collimating, diverging), and, for focusing applicators, the focal length and focal area.

Message:	Provide the following information concerning the applicator label:	
The manner in which the label is attached:		
[HTML Text]		
Attach a sample of the label.		
File Attachment	[Single File Attachment (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]	
Details	[HTML Text]	

## **Operation Controls**

Note:	Part 1050.10(d)(1) of 21 CFR requires that each operation control be clearly labeled, identifying the function controlled and, where appropriate, the units of measure of that function. If a separate control and indicator are associated with the same function, labeling the units of measure of that function is required for the indicator but not for the control.		
9 / 1	Provide drawings, photographs, or other documents, which show clearly the location and labeling of all such controls. Click on the Add button below to attach and select the files.		
File Attachment	[Multiple File Attachments (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]		
Details	[HTML Text]		

## Service Controls

Note:	Part 1050.10(d)(2) of 21 CFR requires that each service control that is accessible without displacement or removal of any part of the product be clearly labeled, identifying the function controlled and including the phrase for service adjustment only.		
Provide drawings, photographs, or other documents, which show clearly the location and labeling of all such controls. Click on the Add button below to attach and select the files.			
File Attachment [Multiple File Attachments (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]			
Details [HTML Text]			

## Information Requirements

Note: Pro	ovide the following information regarding servicing information, user information, and product description.
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## Servicing Information

Note:  Part 1050.10(f) (1) of 21 CFR requires a manufacturer to provide to servicing dealers and distributors adequate instru for operation, service, and calibration of the product. This must include:(a) A description of those controls and proceed could be used to increase radiation emission levels.(b) A schedule of maintenance necessary to keep the product in compliance with 21 CFR 1050.10.(c) Any safety precautions that may be necessary regarding ultrasonic exposure.			
Attach a copy of the servicing information clearly identified above. Click on the Add button below to attach and select the files.			
File Attachment		[Multiple File Attachments (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]	
Details		[HTML Text]	

## User Information

Part 1050.10(f)(2) of 21 CFR requires a manufacturer to provide users with adequate instructions for assembly, operation, and Note: safe use of the product. This must include: (a) A discussion of all operation controls and a description of the effect of each control. (b) A schedule of maintenance necessary to keep the product in compliance with 21 CFR 1050.10. (c) Any safety precautions that may be necessary regarding ultrasound exposure. (d) A description (including textual discussion and diagrams, plots or photographs) of the spatial distribution of the radiated field. The description must include the statement that it applies for the radiation emitted into the equivalent of an infinite medium of distilled, degassed water at 30°C and with line voltage variations in the range of ± 10%, or the rated value. (e) The uncertainties in magnitude, expressed in percentage error, of the ultrasonic frequency, effective radiating area, and (when applicable) the ratio of the temporal-maximum to temporal-average effective intensity, pulse duration, pulse repetition rate, focal area, and focal length. (f) The error in indication of radiated power and intensity. (g) The error in indication of present treatment time. (h) A listing of all controls, adjustments, and procedures for operation and maintenance, including the warning "Caution -- use of controls or adjustments or performance of procedures other than those specified herein may result in hazardous exposure to ultrasonic energy." Attach a copy of the user information to the preceding sections, clearly identified above. Click on the Add... button below to attach and select the files.

#### **Product Description**

[HTML Text]

File Attachment

Details

Note:	Note:  In order to adequately review a manufacturer's product, CDRH requires that a product report provide a thorough physical description of the product. Such a description must include:(a) Photographs or drawings or the generator and applicator.(b) complete schematic diagram of the product.		
Provide the information listed above as attachments. Click on the Add button below to attach and select the files.			
File Attachment	[Multiple File Attachments (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]		
Details	[HTML Text]		

[Multiple File Attachments (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]

#### **Testing Programs**

The information reported in this section will be used to determine whether the manufacturer's testing programs are adequate for certification (21 CFR 1010.2) and that the products are in compliance with the performance standard. Each item in this
section must be addressed individually and in detail.

#### Incoming Component Testing

Note:	Fully describe all tests that are performed on components whose perfomance can affect compliance with this standard. This description should include but is not limited to:(a) Identify the component tested and its function.(b) State whether the component is tested on a 100 percent or sampling basis. If tested on a sampling basis, provide all sampling parameters and the basis for selecting the Acceptable Quality Level.(c) Describe the corrective action taken following unit or lot rejection (i.e., return component to manufacturer, test 100 percent of components, increasesampling level). If the sampling level is increased, provide the complete rationale for this procedure, and any revised acceptance criteria.		
Provide the above information as an attachment for each tested component. For example, if transducer crystals and timers are among the components tested, attach the description of the testing of crystals as one file attachment and the description of the testing of timers as the second file attachment, and so forth. Click on the Add button to add and select the files to be attached.			
,			

File Attachment	[Multiple File Attachments (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]
Details	[HTML Text]

#### Calibration of Test Instruments

note: ni in di aa aa	y describe the instruments used in any test conducted to ensure compliance with this standard. This should include, but is limited to, the following:(a) The manufacturer, model number, type (e.g., radiation force), accuracy, and resolution of the rument used to measure ultrasonic power.(b) The procedure by which the above instrument is calibrated. Include a cription of any calibrated source used, stating the accuracy and by whom calibrated.(c) The manufacturer, model number, complete specifications of the hydrophone used to measure ultrasonic intensity.(d) A description of the scanning aratus used to measure the spatial distribution of the radiated field.(e) A description of, and calibration procedures for, any are instrument used for compliance testing.			
Provide the above inform	Provide the above information as attachments below. Click on the Add button to add and select the files to be attached.			
File Attachment	[Multiple File Attachments (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]			
Details	[HTML Text]			

## **Production Testing**

Note:	Fully describe all tests that are performed on the product during or after production to ensure compliance with this standard. The description of each test must include, but is not limited to, items (a) through (e) below. Note that part 1050.10(e) of 21 CFR requires that measurements of ultrasonic radiation be made with the radiation emitted into the equivalent of an infinite medium of distilled, degassed water at 30°C, and with line voltage variations in the range of ±10% of the rated value.
	(a) Identify all instruments reported in the Calibration of Test Instruments section above that are used for the test.
	(b) State the sources and magnitudes of uncertainty in the test.
	(c) State whether the component or parameter is tested 100 percent or sampling basis. If tested on a sampling basis, include lot size, proportion of total production tested, method of sample selection to ensure randomness, and the rationale for sampling rather than testing on a 100 percent basis. It must be clearly demonstrated that such a sampling program will ensure compliance of all certified products.
	(d) Decribe the test procedure in detail, including any assumptions that are taken from the results. For example, in the description of the test for accuracy of indicated power, state the specific power levels at which the measurement is made, the error in indicated power at each point, and the range over which the average error is assumed to hold.
	(e) Describe the corrective action taken following unit or lot rejection (i.e., Increase sampling, test 100 percent).

Provide the above information as an attachment for each parameter tested. For example, present the description of the test for error in indicated power as a file attachment. Click on the Add... button to add and select the files to be attached. The parameters tested during production should include, but are not limited to:

- (a) Error in indication of temporal-average ultrasonic power (CW units).
- (b) Error in indication of temporal-maximum ultrasonic power (pulsed units).
- (c) Error in measured value of effective radiating area.
- (d) Error in the determination of the ratio of temporal-maximum effective intensity to temporal-average effective intensity.
- (e) Error in indication of preset treatment time.
- (f) Proper operation of manual and automatic treatment termination devices.(g) Proper operation of visual "ultrasound on" indicator.
- (h) Proper operation of indicators of pulse duration, pulse repetition rate, and ultrasonic frequency (where applicable).

File Attachment	[Multiple File Attachments (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]
Details	[HTML Text]

#### Life Testing

Note:	Fully describe all tests that are performed on the product to ensure that it is capable of complying with the standard throughout its life. This should include, but is not limited to:(a) Sample size, frequency of sampling, and selection criteria.(b) Description of the test, including the sources and magnitudes of error, parameters measured or monitored, instruments used, and length of test or equivalent length of test.			
Provide the above information as an attachment. Click on the Add button below to add and select the files to be attached.				
File Attachment		[Multiple File Attachments (.pdf, .jpg, .gif, .tif, .avi, .wmv, .xpt, .xml, .dtd, .sgml, .mol, .xls, .csv, .zip)]		
Details		[HTML Text]		
Provide an estimate of	Provide an estimate of the useful life of the product (in years):			
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 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 FDA 3644 Guide for Preparing Product Reports for Ultrasonic Therapy Products (03/06)			

Document Key: Specialized Response content is defined within straight brackets []; Special code: [L] List of Values.