Submission Report Page 1 of 20

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

Submission Report Page 2 of 20

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 you are preparing. (Supplements should be prepared using the same document type as the original submission.)				
	of Submission is this? (Supplements should be submitted e 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			

Submission Report Page 3 of 20

	() Variance Request (General, not Laser Light Show) (21 CFR 1010.4) () Laser Original Equipment/Component Manufacturer Registration (21 CFR 1040.10(a)(3)(ii)) () Abbreviated Report (21 CFR 1002.12)
--	--

Step 2	After answering the Submission Type question above, one o may become active and required (see the blue dot to the rights an active question, select the appropriate product area or question's pick list.	nt of the question). If there
What Type of P	oduct is this Radiation Safety Report about?	ļ*
Laser Light Sho	ws	
What Type of P	oduct is this Annual Report about?	
What Laser Ligh	t Show Document are you filing?	
What Type of C	orrespondence is this?	
What Type of P	oduct is this Variance Request about?	

Submission Report Page 4 of 20

Manufacturer Data					
Manufacturer	Manufacturer Responsible for Product Compliance				
Note:	perfo progi the o	is the firm that takes responsibility for certification that the product meets the ormance standard. This firm develops and maintains the quality control and testing ram that is the basis for the certification of this product. Additionally, this firm usually is owner of the product design and manufacturing process design. The product design and manufacturing process design.			
Select the Manu	ufactur	rer's address from the Establishment Address book: *			
Establishment I	nforma	ation:			
Establishment N					
Division Name					
Home Page					
Physical Location	on:				
Address					
Telephone Num	nber				
Fax Number					
Mailing Location	า:				
Address					
Telephone Num	nber				
Fax Number					
Responsible	Indiv	idual			
Note:		responsible individual is the highest level and most responsible individual affiliated with establishment.			
Select the Resp	onsibl	le Individual from the Contact Address book:			
Contact Informa					
Contact Name	1011.				
Occupation Title					
Email Address					
Establishment I	nforma	ı			
Establishment Name					
Division Name					
Physical Location	on:				
Address					
Telephone Num	nber				
Fax Number					

Submission Report Page 5 of 20

Mailing Location	ı.		
Address	<i>'.</i>		
Telephone Num	her		
Fax Number	ibei		
I ax Number			
Manufacturer	's De	eporting Official	
iviariulaciulei	3116	sporting Official	
Note:	addre repoi	is the person at the manufacturing facility that is knowledgeable and responsible for essing all aspects of the testing and quality control procedures for certification as rted to FDA in the product report. Documentation of changes intesting and quality rol procedures submitted to FDA must be signed by this individual.	
Select the Repo	rting (Official from Contact Address book: *	
Contact Informa	tion:		
Contact Name			
Occupation Title)		
Email Address			
Establishment li	nforma	ation:	
Establishment N	lame		
Division Name			
Physical Location	on:		
Address			
Telephone Num	ber		
Fax Number			
Mailing Location	ı:		
Address			
Telephone Num	ber		
Fax Number			
Report Subm	itter		
Note:	prepa by th	submitter may be a consulting individual or firm providing assistance in report aration and maintenance. Documents or submissions such as this one that are prepared e submitter must have an accompanying authorization letter from the manufacturer's rting official for authenticity.	
Select the Subn	Select the Submitter from the Contact Address book:		
Contact Informa	Contact Information:		
Contact Name			
Occupation Title)		
Email Address			
Establishment l	nforma	ation:	
Establishment N	Establishment Name		
Division Name			

Submission Report Page 6 of 20

Physical Location:	
Address	
Telephone Number	
Fax Number	
Mailing Location:	,
Address	
Telephone Number	
Fax Number	
Comments:	
Internal Reference N	umber:
Parent Establishn	nent
Is there a parent esta	blishment? *
Select the Parent Est	ablishment and Contact from the Contact Address book:
Contact Information:	
Contact Name	
Occupation Title	
Email Address	
Establishment Inform	ation:
Establishment Name	
Division Name	
Physical Location:	
Address	
Telephone Number	
Fax Number	
Mailing Location:	
Address	
Telephone Number	
Fax Number	
Manufacturer Des	signated United States Agent
Note: Man	ufacturers exporting to the U.S. must designate a U.S. agent, see 21 CFR 1005.25.
Is there a United Stat	es agent that has been designated by the manufacturer?
Importer	

Submission Report Page 7 of 20

Additional Manufacturing Locations

Submission Report Page 8 of 20

P	r۸	d	П	ct		a	ta
	··	ч	ч	UL.	_	ч	LU

Product and Model Identification

Attention - Information about this section

In this section you'll be asked to identify several required or optional things which will help FDA/CDRH staff to prioritize their reviews. You'll be asked to consider the following aspects:

- (1) Identify your product's radiation type and the CDRH Product Code.
- (2) Enter an Accession number if this will be a report supplment. If you are preparing a supplement, you'll see that after entering a valid 7-digit Accession number many questions will no longer be required (they will either be disabled or will be optional, meaning they will no longer have the blue dot).
- (3) You will also have several questions that are of high significance for FDA/CDRH why you might be submitting this report or correspondence. Please read these questions carefully, referring to the 21 CFR regulations on the website www.FDA.gov if you are unsure if the question is relevant to your firm's situation.
- (4) If you find that you have more information that you want the FDA/CDRH to read but it doesn't seem to fit under other questions, we have a final "*Additional Information*" question in this section which invites you to add comments and/or attach a file that provides further information from your firm about this submission. This is the place to add that extra information.

Product Type Reported

What is the product co	ode?
To select the three let	ter product code,
Select the appropria from which to choose.Select the best mate	
Category	
Product Code	
Performance Standard	
If Other, provide a cat	tegory name for this specific product.

Submission Report Page 9 of 20

Report Information

Is this the first time you've submitted a report on the particular type of product selected in the Product Type Reported section?

.

Since this is not the first time you've reported on this type of product, then is this a report supplement to a previously reported model family?

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, etc. See Accession number description below.)

Are you requesting a new variance, a renewal, extension or amendment to a previous variance?

*

Stop:

If you are requesting a new variance, renewal, extension, or amendment, you must file a Variance Request separate from this report. To do this, open a new report (File > New) and select either "Laser Light Show Variance Request" or "Variance Request (General, not Laser Light Show)" as your Type of Submission in the Submission Information Screen. If you select "Variance Request (General, not Laser Light Show)r" you must select the product for which you are requesting a variance with the pick list in the bottom section of the screen.

Special Considerations

Information:

If this product will require a formally approved Variance from a certain performance requirement, you will need to complete two Reports for FDA, both (1) this Radiation Safety Report (RSR) on this product, and (2) a Variance Request report. This eSubmitter software application package includes a general Variance Request form as well as the specific Laser Light Show Variance Request form. Both the Product RSR file and the appropriate Variance Request Correspondence file must be submitted to CDRH following the regular files packaging procedures in this application. Both may be transferred to the same CD or submitted via the FDA ESG to submit to the FDA/CDRH.

In addition, any Variance Request form must be printed out and the signed hard-copy sent to FDA's Division of Dockets Management at:

Division of Dockets Management (HFA-305) Food and Drug Administration 5630 Fishers Lane Rockville, MD 20852

NOTE: There is no need to send a copy of the CD to Division of Dockets Management.

Noncompliances or Defects

Does this document	t or any of its attachments contain:	
A notification of nonco	ompliance or defect?	*
You may provide an e	explanation and/or attach a document here:	
Details		

Submission Report Page 10 of 20

Responses to Noncompliances or Defects

You may add an explanation and/or attach a document here:

Does this document or any of its attachments contain any of these responses concerning noncompliances or defects? A refutation of noncompliances or defects identified to your firm? * A request for an exemption from notification to purchasers (see 21 CFR 1003.21 and 1003.30)? Corrective action plans you intend to implement to correct noncompliances or defects discovered in past or current production? Note: If you are submitting a Corrective Action Plan (CAP) following 21 CFR 1004 and information on design changes for future production, the design change information must be submitted in a Radiation Safety (Product) Report or supplemental report. Both the proposed CAP and the design changes may be submitted in one document if you prepare a product report and choose to include the CAP in it as a file attachment. Alternatively, you may create a separate eSubmission for the CAP using the "Correspondence" type template and selecting "Followup correspondence to FDA." A description of any design changes that correct noncompliances for future production? Note: If you are submitting information on product design changes for future production due to a discovery of noncompliances or defects in current production, you must use the Radiation Safety (Product) Report template to create the report. Correspondence templates may be used to submit other information such as a proposed corrective action plan pertaining to a noncompliance or defect.

Exemption Requests

Details

Does this document or any of its attachments contain:		
Exemption of a product for government use from a standard (21 CFR 1010.5)?	*	
Exemption for products for government use from reporting and recordkeeping (21 CFR 1002.51)?	*	
Special exemption of products from reporting and/or recordkeeping (21 CFR 1002.50)?	*	
Request for approval of alternate labeling?	*	
Application for alternate test procedures (21 CFR 1010.13)?	*	
You may provide an explanation and/or attach any relevant documents here:		

Variance Requests

Information:

Please note: in addition to responding to these questions below, a separate General Variance Request or Laser Light Show Variance Request form must be completed and submitted to CDRH, with a hard copy sent to FDA's Division of Dockets Management as instructed below for any variance request. The information requested on this screen does not constitute the full structured content of the variance request. The 2 types of Variance forms can be created in eSubmitter by selecting the appropriate Variance submission type under the eRad Health Menu section of this application.

Submission Report Page 11 of 20

Messag	ie:	lick the plus sign to list the requirements from which you are requesting a variance.
This sul	This submission includes an application for a variance from certain requirements.	
Item	No Info	nation Provided.
Provide	an expl	ation and attach supporting files, if necessary. Click on the plus sign below to attach files.
Details		
Provide an explana Details Stop: Fo Th ins Us U. Ce Att Do 10 Sil Ac su Fo Dir 56		or all Variance requests, two submissions must be made to the FDA. the electronic version should be submitted following the Packaging Files for Submission structions located under Output in the Menu bar, and explained in subsection 4.3 of the ser Manual. If sending a CD & submittal letter, please mail to: S. Food and Drug Administration enter for Devices and Radiological Health ttn: eSubmitter Team ocument Mail Center - WO66-0609 O903 New Hampshire Avenue ilver Spring, MD 20993-0002 dditionally, a paper version (hard-copy) of the signed Variance request document should be ubmitted to: ood and Drug Administration ivision of Dockets Management (HFA-305) 630 Fishers Lane, Room 1061 ockville, MD 20857

Responses to Communications from FDA

Does this document or any of its attachments contain:			
A response to an FDA inspection?	*		
What was the date of the inspection?			
A response to a Warning letter or a Notification of Noncompliance or Defect from the FDA?	*		
What was the date of the Warning Letter or other notification letter?			
A response to a report review inquiry from the CDRH (the inquiry may have been in the form of a letter, email, or phone call)?	*		
What was the date of the inquiry?			
A response to any other communication from FDA?	*		
What was the date of the communication?			
Provide an explanation:			

Additional Information

Here's your opportunity to add anything else to this submission that you want to tell the FDA!

Submission Report Page 12 of 20

	evant information or additional comments that would help expedite the review of this e plus sign below to attach any supporting files.	
Details		
Private Labeling		
Is the product sold by	other companies under different brand names?	*
Medical Devices		

If it has not been submitted yet, or if your device is exempt from premarket clearance or approval, please provide an explanation. The device regulations can be found in 21 CFR 807 - device manufacturer

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.

registration and device listing.

Submission Report Page 13 of 20

Report Identification

1.0 General Definitions

General Definitions

This software application should be followed for all lasers and products containing, incorporating, or intended to incorporate, a laser or laser system [see the definition of "laser product" in section 21 CFR1040.10(b)(21)].

Laser Reporting and Recordkeeping (21 CFR 1002)

Applicability of reporting and recordkeeping requirements for laser products:

Class I, IIa, II, and IIIa laser products and laser products containing such lasers will require: Product Report, Annual Report, test records, manufacturer's distribution records, and dealer/distributor distribution records.

Note that for Class I laser products containing lasers of Class I, IIa, II, or IIIa no Supplemental Reports are required. Furthermore, some Class I laser products have already been exempted from the requirement for distribution records (see Notice to Industry dated August 9, 1988, Laser Notice # 41).

Class IIIb and IV laser products require all of the above plus Supplemental Reports when the criteria requiring submission of Supplemental Reports are met.

The laser standard applies to all laser products manufactured after August 1, 1976 (1040.10(a)), unless the products are either: sold to a manufacturer for use as components (or replacements) in products that will be certified (1040.10(a)(l)); sold by or for a manufacturer as repair or replacement components if they are properly labeled as such and have installation instructions (1040.10)(a)(2)); or intended for export only, are labeled as such, and comply with the requirements of the importing country (1010.20). Manufacturers of laser products that are sold to other manufacturers for use as components in their products are required to register and list such products.

Laser Definitions from 21 CFR 1040.10(b)

Laser means any product that can be made to produce or amplify electromagnetic radiation at wavelengths greater than 250 nm but less than or equal to 13,000 nm or, after August 20, 1986, at wavelengths equal to or greater than 180 nm but less than or equal to 1.0X106 nmprimarily by the process of controlled stimulated emission.

Laser energy source means any product intended for use inconjunction with a laser to supply energy for the operation of the laser. General energy sources such as electrical supply mains or batteries shall not be considered to constitute laser energy sources.

Laser product means any manufactured product or assemblage of components which constitutes, incorporates, or is intended to incorporate a laser or laser system. A laser product is any device that

Submission Report Page 14 of 20

constitutes, incorporates, or is intended to incorporate a laser or laser system (1040.10(b)(21)). A laser or laser system that is intended for use as a component of an electronic product shall itself be considered a laser product.

Laser radiation means all electromagnetic radiation emitted by a laser product within the spectral range specified in paragraph1040.10(b)(19) that is produced as a result of controlled stimulated emission or that is detectable with radiation so produced through the appropriate aperture stop and within the appropriate solid angle of acceptance, as specified in 1040.10(e).

Laser system means a laser in combination with an appropriate laser energy source with or without additional incorporated components. See paragraph 1040.10(c)(2) of the laser product performance standard for an explanation of the term "removable laser system."

2.0 Specific-Purpose Product Definitions

Specific-Purpose Product Definitions

Demonstration laser product means any laser product manufactured, designed, intended, or promoted for purposes of demonstration, entertainment, advertising display, or artistic composition. The term "demonstration laser product" does not apply to laser products which are not manufactured, designed, intended, or promoted for such purposes, even though they may be used for those purposes or are intended to demonstrate other applications. Demonstration laser products (1040.10(b)(13)) include:

laser products promoted for classroom demonstration of optical phenomena;

artistic displays and their associated apparatus;

laser light show projectors; and

laser light shows and displays themselves.

A general-purpose, scientific, medical or industrial laser product is not considered to be a demonstration laser product when it is demonstrated to a prospective purchaser. Demonstration laser products are restricted in their outputs to Class IIIa with its accompanying restrictions to Class I for short pulses and invisiblewavelengths(1040.11(c)). Because these levels are too low for effective use incommercial theatrical lighting effects, CDRH may grant variances (1010.4) to manufacturers of laser light shows and display devices. As a condition of the variance, the manufacturer must agree to adhere to several safety conditions to provide a level of safety to the publice quivalent to a fully compliant product. Consult the Compliance Guide for Laser Products, September 1985, Appendix B, Clarification of Certain Laser Light Show Requirements, for more information.

3.0 Show Name

What is (are) the name(s) of the light show or display?			*
Item	Show Name	Projector Model	Brand or Trade Name

Submission Report Page 15 of 20

4.0 Va	ria	nce		
Attach a copy of your variance application (FDA Form 3147) and/or, if approved, attach your variance * approval letter (or variance number). Click on the Add button below to attach any supporting files.				
»	Is variance application attached? *			*
»	Is a copy of your variance approval letter attached?			*
	»	Or provide current variance number:		

Submission Report Page 16 of 20

Equipment Information

5.0 Projection Equipment

6.0 Show Venue

The laser light show or display takes place in:				
Item	No	No Information Provided.		
	»	If "o	ther" has been selected, please specify:	
Note:			Be sure to provide beam path diagrams/floor plans for ea above, unless certain drawings are general enough to co shall be attached following Part 9.0 Diagrams and Drawin	ver more than one type. Drawings

The laser light show or display takes place:				
Item	No	No Information Provided.		
	»	If "other" has been selected, please specify:		

7.0 Show Locations, Dates, and Times

Note:

Give specific location(s), date(s), and time(s) for the show if this information is known at the time this report is submitted. If not, advanced written notification must be made as early as possible to appropriate Federal, State, and local authorities. To be considered timely, this written notice must be submitted 30 days prior to the opening of the show. When the show dates become known to the manufacturer less than 30 days prior to the show date, the required information must be provided verbally by phone or by FAX to CDRH. A confirming formal written notice, including the date of the phone notification and the name of the CDRH individual to whom the information was given must be submitted within 14 days. Written confirmation would not be needed following a FAXed notification. CDRH must be notified of every show that your firm intends to produce. If notifications are not routinely received in a timely manner your variance may be revoked.

Click on the Add... button below to attach any supporting files.

Details

8.0 Light Show Effects Produced

The lase	The laser light show uses the following laser effects:		
Item	No Information Provided.		
If "other	If "other" has been selected, please specify:		
Note:	Be sure that the beam path diagrams included in your response to Part 9.0 are sufficient to illustrate all of the effects indicated above. Several effects may be included in a single diagram.		

Submission Report Page 17 of 20

9.0 Diagrams and Drawings of Show Venue

9.1 Provide both plan and elevation drawings with dimensions of the show or display. If the setup varies from show to show, then provide this information for a typical show. * If no drawings are attached. please add an explanation in the text box. Click on the Add... button below to attach any supporting

*Be sure to include in the drawings:

- (1) the location of the projector(s) and control panel(s), audience, performer(s), operator
- (s), mirrors, mirror balls, display screens (or other targets), and beam termination points;
- (2) the direct and reflected laser radiation beam path;
- (3) the laser radiation levels in each beam including the wavelength, maximum power, and scan parameters (if scanned) for the worst case from a human access point of view;
- (4) the minimum separations of the laser radiation fields (or beams) from reference locations in audience and performer areas in both vertical and horizontal directions; and

(5) any direct or reflected beams into audience or performer locations.

Are drawings attached?

10.0 Laser Radiation Levels

10.1 Describe how each of the laser radiation levels, indicated on the drawings attached in 9.0, were determined. If any levels were derived from calculations rather than directly measured, provide the actual calculations that were made. Click on the Add... button below to attach any supporting files.

11.0 Scanning Safeguards

11.1 Will there be audience scanning* from any of the planned effects?

Note:

* Audience scanning is considered to be any scanning, projection, or reflection of laser or collateral radiation into audience or other accessible, uncontrolled area. Scattered radiation coming from diffuse reflectors such as fog, smoke, mist or similar diffusing media is not considered audience scanning. However, all radiation must be below Class I levels if it reaches into audience or other uncontrolled areas. A scanning safeguard is required whenever a laser light show includes audience scanning to assure that the laser radiation levels in audience areas will not exceed Class I limits if there is a scan failure. See the companion publication, "Compliance Guide for Laser Products," for further discussion.

11.2 Do any of the planned effects require laser radiation (direct or scanned beams) to be viewed by operators, performers, or employees?

If the answer to 11.1 or 11.2 is yes, describe how the radiation levels that reach into audience areas are maintained at Class I levels by scanning. Your attached description must include details of the required scan failure safeguard, including a discussion of the means of detection of the scanning, the theory of the operation of the scanning safeguard, and its speed of response in order to show that it will prevent the scanned radiation from exceeding the Class I limits.

Submission Report Page 18 of 20

11.3	Will any laser radiation greater than Class I STRIKE BUT NOT BE VIEWED by operators, performers or other employees?		
»	Describe, in detail, the operation of the scan failure safeguard or other means which will prevent exposure to beams exceeding Class II. If a scan safeguard is used, include a discussion of the detection of scanning, the operation, and the speed of response of the safeguard to show that it will prevent the scanned radiation from exceeding the limits of Class II. If other means are used, such as pressure pads or infared beams, describe in detail as well.		
	Details		
12.0 C	perator Controls		
12.1	Is the show under th	e continuous control of an operator? *	
12.2	Does the laser opera	ator perform tasks in addition to operation of the laser projector?	
	Describe those task		
»	Decembe these tack	·	
12.3	Can the operator see all of the propagating beam paths, their terminations and the audience at times during the performance?		
»	» Explain how adequate surveillance is provided:		
	I		
12.4		nel assist in providing surveillance of the laser display?	
»	State the number of persons, their identification (job titles), their duties, and how they assist in providing surveillance. Describe how they are in constant communication with the operator.		
What ou	ualifications* are requi	ired of laser operators for your show?	
vviiat qt	daillications are requ	- Ined of laser operators for your snow!	
that the ope projection s training film from the Las		variances are required by the variance to employ trained operators or to assure rators receive adequate training to qualify them for the safe use of the laser extem and presentation of the light show effects. Useful information including sometimes, reference books, and programs on the safe use of lasers may be obtained the ser Institute of America (LIA) and from the American National Standards Institute SI standard Z136.1).	
12.6	If your show is not under the continuous control of an operator, is a person designated to be responsible for the immediate termination of the laser radiation in the event of equipment malfunction, audience unruliness, or other unsafe conditions?		
»	Explain alternate control:		
12.7	How is this person of	esignated? What are his or her duties?	
12.8	What qualifications a	are required for this person?	

Submission Report Page 19 of 20

13.0	Proj	ection Equipment Controls	
13.1			
13.1	Ai	Are one or more readily accessible controls provided to immediately terminate laser radiation?	
	»	Number of controls:	
13.2	De	Describe the location of these controls and their operation relative to your show.	

Submission Report Page 20 of 20

Test and Notification Procedures

14.0 Test Procedures

- Attach a copy of the written setup, alignment, and test procedures to be followed prior to the operation of the laser light show at each location. If setup procedures are not a seperate form, provide a detailed description below of procedures that are followed. Click on the Add... button below to attach any supporting files.
- 14.2 When are these setup, alignment, and test procedures performed?
- 14.3 What laser radiation levels are used during setup, alignment, and checkout? (in milliwatts)
- 14.4 Is a written record of the results of the setup, alignment, and test procedures maintained?
 - Explain how adequate quality assurance is maintained:
 - You may attach records such as Standard Operating Procedures (SOPs), Quality Safety Checklists, and Daily Logs.

Details

Note:

Adequate recordkeeping would include, but not limited to: (1) sketches showing the location of the laser projector(s), operator(s), performer(s), audience, beam paths, viewing screens, wall mirrors, mirror balls, and other surfaces that may be struck by the laser beams; (2) information on scanning patterns, velocity, and frequency; and/or (3) laser radiation levels used in each effect.

15.0 Notification Procedures

What procedures are followed for notification of appropriate Federal (CDRH, FAA), State and local agencies? Either attach a file, form letters, or describe procedures in the text box below. Click on the Add... button to attach any supporting files.

What Federal, State and local agencies are notified or would be notified?

Item No Information Provided.

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.