Submission Report

Section: Report

Introduction

ISO 13485 Voluntary Audit Report Program

Section 228 of the Food and Drug Administration Amendments Act of 2007 (FDAAA), which amends section 704(g)(7) of the Federal Food, Drug, and Cosmetic Act (the Act) (21 U.S.C. 374 (g)(7)) was amended to add the following provision:

"(F) For the purpose of setting risk-based inspectional priorities, the Secretary shall accept voluntary submissions of reports of audits assessing conformance with appropriate quality system standards set by the International Organization for Standardization (ISO) and identified by the Secretary in public notice. If the owner or operator of an establishment elects to submit audit reports under this subparagraph, the owner or operator shall submit all such audit reports with respect to the establishment during the preceding 2-year periods."

Effective October 1, 2011, FDA will begin a two year voluntary pilot program as described in the guidance document entitled "Medical Device ISO 13485:2003 Voluntary Audit Report Submission Pilot Program" found at [web address]. FDA will evaluate the outcomes of the pilot program at the end of this period. It is important to note that participation in the Medical Device ISO 13485:2003 Voluntary Audit Report Submission Pilot Program is entirely voluntary.

Complete the following question and answer form, as well as attach the necessary PDF files to submit your ISO 13485 Audit Reports to FDA's Center for Devices and Radiological Health via the FDA Electronic Submissions Gateway (ESG). To register with the FDA ESG, go to www.fda.gov/esg/.

Blue dots indicate required fields.

1.0 Inspection Date

Warning: Your submission must be sent in within the last 90 days of an audit.

Has an audit been performed in the last 90 calendar days?

Yes

2.0 Contact Information

Contact Information

Applicant Contact Information should be completed by the requester or submitter. The applicant will act as the primary point of contact for this ISO 13485 Audit Report Submission.

Definitions

Company - Refers to the organization as a whole for a specific contact reference point.

Establishment - Refers to the physical location of a facility.

Tips

The Address Book function can store re-usable contact information across multiple eSubmitter report types. Click the link below to view instruction on how to use the Address Book.

Address Book User Guide

2.1 Establishment Contact Information

Information:	Enter the facility contact information that was audited including the most responsible person.
Warring.	The FDA Establishment Identifier (FEI) will be validated upon FDA receipt. Any invalid FEI numbers and Establishment addresses will be rejected.

Establishment Contact Information		
Contact Information:		
Contact Name		
Occupation Title		
Email Address		
Address		

Submission Report				
Establishment Name				
Division Name				
Address				
Telephone Number				
FDA Establishment Identifier	(FEI)			
	Verify that your FEI and Facility Address are up to date. Any inconsistencies may cause the submission to be rejected. Establishment Registration & Device Listing			
Please select the associated	FDA Dis	strict Office *		
2.2 Auditing Organizatio	n Cont	act Information		
Please enter Auditing Organia	zation co	ontact information	*	
Contact Information:				
Contact Name				
Occupation Title				
Email Address				
Address				
Establishment Name				
Address				
Telephone Number				
Auditing Organization Members				
Team Member Name				
Is this team member an employee or a contractor to the auditing organization?				
Team Member Name				
Is this team member an employee or a contractor to the auditing organization?				
Team Member Name				
Is this team member an employee or a contractor to the auditing organization?				

Submission Report							
Team Member Name							
Is this team member an employee or a contractor to the auditing organization?							
Team Me	Team Member Name						
Is this tea	am memb	per an e	employee	or a contractor	to the auditing	organization?	
2.3 App	olicant C	Contac	ct Informa	ation			
Informati	ion:	Conta	ct name s	hould be the su	ubmitter of this	submission.	
Please e	nter your	contac	ct informati	ion			*
Contact	Informatio	on:					
Contact I	Name						
Occupati	ion Title						
Email Address							
Address	Address						
Establishment Name							
Address							
Telephor	Telephone Number						
3.0 Sub	3.0 Submission Content						
Enter your product for medical devices manufactured at this establishment. Product Codes can be entered manually or selected using filters. Click the link for additional instructions on how to use the Product Code Filters. Product Code Filter Guide							
Multiple Draduct Codes may be selected							
Multiple Product Codes may be selected * Device Classification C.E.R. Section							
Item	Product	Code			Class	Panel	C.F.R. Section

Audit Type

3.1 Auditing B	3.1 Auditing Bodies			
Select all that ap	[] Australia [] Canada [] European Union [] Japan			
Auditing Affilia	Auditing Affiliations			
Select an auditin	Select an auditing body affiliated with Australia.			
Select an auditin	g body affiliated with Canada.			
Select an auditin	g body affiliated with the European Union.			
Select an auditin	Select an auditing body affiliated with Japan.			
4.0 Audit Rep	orts			
Information:	Section 228 of the Food and Drug Administration Amendments Act of 2007 (FDAAA), which amends section 704(g)(7) of the Federal Food, Drug, and Cosmetic Act (the Act) (21 U.S.C. 374 (g)(7)). Section 228 was amended to add the following provision: (F) For the purpose of setting risk-based inspectional priorities, the Secretary shall accept voluntary submissions of reports of audits assessing conformance with appropriate quality system standards set by the International Organization for Standardization (ISO) and identified by the Secretary in public notice. If the owner or operator of an establishment elects to submit audit reports under this subparagraph, the owner or operator shall submit all such audit reports with respect to the establishment during the preceding 2-year periods.			
4.1 Most Rece	ent Audit Report			
4.1 Most Recent Audit Report				
Information:	mation: All documentation and data entered in this section should be associated with the most recent audit date within 90 days of this submission.			
Enter the most re	ecent Audit date within 90 days of this submission.	*		
Attach the original Audit Report.				

Attach the original ISO 13485 certificate *				
Attach any associated correspondence for this Audit Report period. *				
Are all of the above original documents in English?				
4.2 Preceding Audit Reports and Information				
Information: The following set of Audit Reports with a timeframe of 2 years, should precede the most recent Audit Report in Section 4.1.				
Were Audits conducted in the previous 2 years from the last day of the most recent Audit? * Yes				

Section: Previous Audit Reports			
1.0 Preceding	Audit Reports and Information		
Information:	The following set of Audit Reports with a timeframe of 2 years, should precede the most recent Audit Report in Section 4.1.		
How many Audits were conducted in the previous 2 years from the last day of the most * 5 recent Audit?			
1.1 First Prece	ding Audit Report		
Attach your Previ	ous Audit Reports	*	
If the ISO 13485	certificate is different for this Audit Report, please attach the relevant ISO 13485	certificate. *	
Attach any associated correspondence for this Audit Report. *			
Are all of the abo	ve original documents in English? *		
1.2 Second Preceding Audit Report			
Attach your Previous Audit Reports *			
If the ISO 13485 certificate is different for this Audit Report, please attach the relevant ISO 13485 certificate.			
Attach any associated correspondence for this Audit Report. *			
Are all of the above original documents in English?			
1.3 Third Preceding Audit Report			
Attach your Previous Audit Reports *			

Submission Report			
If the ISO 13485 certificate is different for this Audit Report, please attach the relevant ISO 13485 certificate.			
Attach any associated correspondence for this Audit Report period. *			
Are all of the abo	Are all of the above documents in English?		
1.4 Fourth Pro	eceding Audit Report		
Attach your Prev	rious Audit Reports. *		
If the ISO 13485	certificate is different for this Audit Report, please attach the relevant ISO 13485 certificate.		
Attach any asso	ciated correspondence for this Audit Report period. *		
Are all of the abo	Are all of the above documents in English?		
1.5 Fifth Prec	eding Audit Report documents		
Attach your Previous Audit Reports *			
If the ISO 13485 certificate is different for this Audit Report, please attach the relevant ISO 13485 certificate.			
Attach any associated correspondence for this Audit Report period. *			
Are all of the above documents in English?			
Attestation			
Stop:	Any submissions submitted after 90 days from the last day of the Audit will be rejected.		
Information:	To the best of my knowledge, the data and information submitted are truthful and accurate, and no material fact has been omitted.		
Full Name	*		
Date of Attestation *			