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MEDICAL DEVICE SINGLE AUDIT PROGRAM Responsible Office/Division	Version Date: 2015-10-07	Effective Date: 2013-07-15
Title:	Project Manager:	
MDSAP QMS Training Review Checklist	Liliane Brown, USFDA	

Title of MDSAP Procedure/Policy/Guidance	MDSAP Procedure/Policy/ Guidance Number	Reviewed
MDSAP Functional Statement	MDSAP P0001	□Yes □ No
MDSAP Audit Model	MDSAP P0002	□Yes □ No
MDSAP Regulatory Authority Council and Lead Project Manager Authorities, Responsibilities, and Governing Rules	MDSAP P0003	□Yes □ No
MDSAP Assessment Program Procedure	MDSAP AS P0005	□Yes □ No
MDSAP Audit Time Calculation Procedure	MDSAP AU P0008	□Yes □ No
MDSAP Regulatory Authority Council (RAC) Appointment	MDSAP P0009	□Yes □ No
MDSAP AO Application for Recognition Procedure	MDSAP AS P0010	□Yes □ No
MDSAP Witnessed Audit Procedure	MDSAP AS P0012	□Yes □ No
MDSAP Conducting Stage 1 Assessment Procedure	MDSAP AS P0013	□Yes □ No
MDSAP Special Documentary Assessment Procedure	MDSAP AS P0014	□Yes □ No
MDSAP AO Nonconformities Procedure	MDSAP AS P0015	□Yes □ No
On Site Assessment /Audit Procedure(<i>Stage</i> 2, <i>Surveillance</i> , <i>Re-cognition</i> , <i>Critical Locations</i>)	MDSAP AS P0016	□Yes □ No
MDSAP Technical Reviews & Recognition Decision Making Procedure	MDSAP AS P0017	□Yes □ No
MDSAP QMS Audit Report Policy	MDSAP AU P0019	□Yes □ No
MDSAP Special Assessment Audit Procedure	MDSAP AS P0020	□Yes □ No
Appeals Procedure	MDSAP AS P0021	□Yes □ No
Implementing Suspension or Revocation of Recognition	MDSAP AS P0022	□Yes □ No
Flagging/Collection and Review of AO Audit Reports	MDSAP AS P0023	□Yes □ No
Type or Organization Eligible for Audit and Excluded from Participating	MDSAP AU P0024	□Yes □ No

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MDSAP Templates and FormsNumberMDSAP Project Team Work Item (PTWI) Proposal/Approval FormMDSAP F0003.1MDSAP Assessment Program FlowchartMDSAP AS F0005.1MDSAP AO Assessment ProgramMDSAP AS F0005.2	Reviewed □Yes □ No □Yes □ No □Yes □ No □Yes □ No
Proposal/Approval FormMDSAP Assessment Program FlowchartMDSAP AS F0005.1	□Yes □ No □Yes □ No
MDSAP Assessment Program Flowchart MDSAP AS F0005.1	□Yes □ No
	□Yes □ No
MDSAP AQ Assessment Program MDSAP AS F0005 2	
	□Yes □ No
Management File	□Yes □ No
MDSAP AO Assessment Program Approval MDSAP AS F0005.3 Form	
MDSAP Notice of Change MDSAP AS F0005.5	□Yes □ No
Auditor Training Working Instruction MDS/R MDS/R MO10006.1	
MDSAP AO Application for Recognition MDSAP AS F0010.1	
Form	
MDSAP AO Application for Recognition MDSAP AS F0010.2	□Yes □ No
Flowchart	
MDSAP AO Scorecard Form MDSAP AS F0010.3	□Yes □ No
MDSAP AO Participant Agreement MDSAP AS F0010.4	□Yes □ No
AO Recognition Application Additional MDSAP AS F0010.5 Information Sheet	□Yes □ No
AO Application Matrix MDSAP AS F0010.6	□Yes □ No
AO Critical Location Information Form MDSAP AS F0010.7	□Yes □ No
Auditor and Tech Expert Competency MDSAP AS F0010.8 Summary	□Yes □ No
Witnessed Audit Flowchart MDSAP AS F0012.1	□Yes □ No
Witnessed Audit Manufacturer Profile Form MDSAP AS F0012.2	□Yes □ No
Witnessed Audit Report MDSAP AS F0012.3	□Yes □ No
Witnessed Audit Assessment MDSAP AS F0012.4	□Yes □ No
Checklist/Performance Checks	
Stage 1 Assessment Flowchart MDSAP AS F0013.1	□Yes □ No
Stage 1 Assessment Consolidated Report MDSAP AS F0013.2	□Yes □ No
Special Documentary Assessment Flowchart MDSAP AS F0014.1	□Yes □ No
Special Documentary Assessment Report MDSAP AS F0014.2 Form	□Yes □ No
AO Nonconformity Flowchart MDSAP AS F0015.1	□Yes □ No
AO Nonconformity Report Form MDSAP AS F0015.2	□Yes □ No
On-Site Assessment-Audit Process MDSAP AS F0016.1 Flowchart	□Yes □ No
Assessment-Audit Announcement Letter MDSAP AS F0016.2 Template	□Yes □ No
Assessment Audit Plan Form MDSAP AS F0016.3	□Yes □ No
On-Site Assessment-Audit Report Form MDSAP AS F0016.5	□Yes □ No

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Technical Reviews & Recognition Decision Flowchart	MDSAP AS F0017.1	□Yes □ No
AO Recognition Decision Form	MDSAP AS F0017.2	□Yes □ No
AO Recognition Decision Making Calculation Sheet	MDSAP AS F0017.3	□Yes □ No
Letter of Recognition Template	MDSAP AS F0017.4	□Yes □ No
Audit Report Letter to AOs	MDSAP AS F0017.5	□Yes □ No
Audit Report Template for AO Audit Manufacturer	MDSAP AU F0019.1	□Yes □ No
Special Assessment-Audit Flowchart	MDSAP AS F0020.1	□Yes □ No
Special Assessment-Audit Report	MDSAP AS F0020.2	□Yes □ No
Appeals Flowchart	MDSAP AS F0021.1	□Yes □ No
Appeal Request and Processing Form	MDSAP AS F0021.2	□Yes □ No
Implementing Suspension or Revocation of Recognition	MDSAP AS F0022.1	□Yes □ No
Collection and Review of AO Audit Reports Flowchart Overview	MDSAP AU F0023.1	□Yes □ No
Internal FDA Processing of AO Audit Reports Flowchart	MDSAP AU F0024.2	□Yes □ No
MDSAP QMS Title of Procedure	MDSAP QMS Procedure Number	Reviewed
MDSAP QMS Quality Manual	QMS P0001	□Yes □ No
MDSAP QMS Document Control and Approval Procedure		□Yes □ No □Yes □ No
MDSAP QMS Document Control and	QMS P0001	
MDSAP QMS Document Control and Approval Procedure MDSAP QMS Quality Policy (refer to the	QMS P0001 QMS P0002	□Yes □ No
MDSAP QMS Document Control and Approval Procedure MDSAP QMS Quality Policy (refer to the Quality Manual - Section 5) MDSAP QMS Quality Risk Management	QMS P0001 QMS P0002 N/A	□Yes □ No □Yes □ No
MDSAP QMS Document Control and Approval Procedure MDSAP QMS Quality Policy (refer to the Quality Manual - Section 5) MDSAP QMS Quality Risk Management Procedure MDSAP QMS Management Responsibility	QMS P0001 QMS P0002 N/A QMS P0004	□Yes □ No □Yes □ No □Yes □ No
MDSAP QMS Document Control and Approval Procedure MDSAP QMS Quality Policy (refer to the Quality Manual - Section 5) MDSAP QMS Quality Risk Management Procedure MDSAP QMS Management Responsibility and Management Review Procedure MDSAP QMS Control of Nonconforming	QMS P0001 QMS P0002 N/A QMS P0004 QMS P0005	□Yes □ No □Yes □ No □Yes □ No □Yes □ No
MDSAP QMS Document Control and Approval Procedure MDSAP QMS Quality Policy (refer to the Quality Manual - Section 5) MDSAP QMS Quality Risk Management Procedure MDSAP QMS Management Responsibility and Management Review Procedure MDSAP QMS Control of Nonconforming Processes or Services Procedure MDSAP QMS Control of Quality Record	QMS P0001 QMS P0002 N/A QMS P0004 QMS P0005 QMS P0006	 □Yes □ No
MDSAP QMS Document Control and Approval Procedure MDSAP QMS Quality Policy (refer to the Quality Manual - Section 5) MDSAP QMS Quality Risk Management Procedure MDSAP QMS Management Responsibility and Management Review Procedure MDSAP QMS Control of Nonconforming Processes or Services Procedure MDSAP QMS Control of Quality Record Procedure MDSAP QMS Internal Assessment	QMS P0001 QMS P0002 N/A QMS P0004 QMS P0005 QMS P0006 QMS P0007	 □Yes □ No
MDSAP QMS Document Control and Approval Procedure MDSAP QMS Quality Policy (refer to the Quality Manual - Section 5) MDSAP QMS Quality Risk Management Procedure MDSAP QMS Management Responsibility and Management Review Procedure MDSAP QMS Control of Nonconforming Processes or Services Procedure MDSAP QMS Control of Quality Record Procedure MDSAP QMS Internal Assessment Procedure	QMS P0001 QMS P0002 N/A QMS P0004 QMS P0005 QMS P0006 QMS P0007 QMS P0008	 □Yes □ No
MDSAP QMS Document Control and Approval Procedure MDSAP QMS Quality Policy (refer to the Quality Manual - Section 5) MDSAP QMS Quality Risk Management Procedure MDSAP QMS Management Responsibility and Management Review Procedure MDSAP QMS Control of Nonconforming Processes or Services Procedure MDSAP QMS Control of Quality Record Procedure MDSAP QMS Internal Assessment Procedure MDSAP QMS Corrective Action Procedure	QMS P0001 QMS P0002 N/A QMS P0004 QMS P0005 QMS P0006 QMS P0007 QMS P0008 QMS P0009	 □Yes □ No

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MDSAP QMS Continual Improvement Procedure	QMS P0013	□Yes □ No
MDSAP QMS Training Procedure	QMS P0014	□Yes □ No
MDSAP QMS Templates and Forms	MDSAP QMS Document Number	Reviewed
MDSAP QMS New Document Proposal/Change Request (NDP/CR) Form	QMS F0002.1	□Yes □ No
MDSAP QMS New Document Proposal (NDP) Form	QMS F0002.2	□Yes □ No
MDSAP QMS Risk Management Governance Structure	QMS F0004.1	□Yes □ No
MDSAP QMS Risk Management Process Steps	QMS G0004.2	□Yes □ No
MDSAP QMS Risk Management Flowchart	QMS F0004.3	□Yes □ No
MDSAP QMS RM Identifying and Analyzing Risks Form	QMS F0004.4	□Yes □ No
MDSAP QMS Risk Treatment Action Plan	QMS F0004.5	□Yes □ No
MDSAP QMS Management Review Report Form	QMS F0005.1	□Yes □ No
MDSAP QMS Management Review Agenda Form	QMS F0005.2	□Yes □ No
MDSAP QMS Nonconformity Report (NCR) Form	QMS F0006.1	□Yes □ No
MDSAP QMS Control of Quality Records Flowchart	QMS F0007.1	□Yes □ No
MDSAP QMS Internal Assessment Summary Report Form	QMS F0008.1	□Yes □ No
MDSAP QMS Internal Assessment Checklist	QMS F0008.2	□Yes □ No
MDSAP QMS Internal Assessment Qualification/Training Form	QMS F0008.3	⊡Yes ⊡No
MDSAP QMS Internal Schedule- Instruction Form	QMS F0008.4	⊡Yes ⊡No
MDSAP QMS Internal Audit/Self Resolution of Findings Form	QMS F0008.5	⊡Yes □No
MDSAP QMS Corrective Action/Problem (CAPR) Form	QMS F0009.1	⊡Yes □No
MDSAP QMS Corrective Action/Problem (CAPR) Flowchart	QMS F0009.2	⊡Yes ⊡No
MDSAP QMS Complaint and Customer Feedback (CF) Form	QMS F0011.1	⊡Yes ⊡No
MDSAP QMS Training Review Checklist Form	QMS F0014.1	□Yes □No
MDSAP QMS Training Evaluation Form	QMS F0014.2	□Yes □No

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International Standards	Standard Numbe	r Reviewed		
Quality Management Systems – Fundamentals and Vocabulary	ISO 9000	□Yes □No		
Conformity Assessment – Vocabulary and General Principles	ISO/IEC 17000	□Yes □No		
Medical Devices – Quality Management Systems – Requirements for regulatory purposes	ANSI/AAMI/ISO 1348	5 🗆 Yes 🗆 No		
Quality Management Systems – Requirements	ISO 9001	□Yes □No		
Conformity Assessment – General Requirements for Accreditation Bodies Accrediting Conformity Assessment Bodies	ISO/IEC 17011	□Yes □No		
Conformity Assessment – Requirements for Bodies Providing Audit and Certification of Management Systems	ISO/IEC 17021	⊡Yes ⊡No		
Medical Devices – Application of Risk Management to Medical Devices	BS EN ISO 14971	□Yes □No		
Regulatory Authority Guidance				
Documents, Regulations and Other	Guidance Doc. No	b. Reviewed		
Materials				
AUST				
Australian Regulatory Guidelines for Medical Devices (ARGMD)	N/A	□Yes □No		
Uniform Recall Procedure for Therapeutic Goods (Australia)	N/A	□Yes □No		
BRA	ZIL			
Brazilian Medical Devices Good Manufacturing Practices Resolution	RDC 16/2013	□Yes □No		
CANADA				
Quality Management System Audits Performed by Health Canada Recognized Registrars	GD210: ISO 13485	⊡Yes ⊡No		
JAF	AN			
Japanese QMS Ordinance	MHLW MO169	□Yes □No		
UNITED STATES				
Guidance Document - Medical Device Reporting for Manufacturers	N/A	□Yes □No		
Compliance Program Guidance Manual – Inspection of Medical Device Manufacturers	7382.845	□Yes □No		
Guidance Document – Design Control Guidance for Medical Device Manufacturers	N/A	⊡Yes □No		
Investigations Operations Manual (IOM)	N/A	□Yes □No		

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(OPTIONAL)		
Regulatory Procedures Manual (RPM)	N/A	□Yes □No
Guide To Inspections of Quality Systems – QSIT (OPTIONAL)	N/A	□Yes □No
IMDRF DO	CUMENTS	1
Requirements for Medical Device Auditing Organizations for Regulatory Authority Recognition	IMDRF/MDSAP/ WG/N3	⊡Yes ⊡No
Competence and Training Requirements for Auditing Organizations	IMDRF/MDSAP/ WG/N4	⊡Yes □No
Regulatory Authority Assessment Method for the Recognition and Monitoring of Medical Device Auditing Organizations	IMDRF/MDSAP/ WG/N5	⊡Yes ⊡No
Regulatory Authority Assessor Competence and Training Requirements	IMDRF/MDSAP/ WG/N6	⊡Yes □No
Grading Nonconformities Issued to Auditing Organizations by Recognizing Regulatory Authorities; and Decision-Making Principles and Criteria for the Recognition of Auditing Organization	IMDRF/MDSAP/ WG/N11	⊡Yes ⊡No