

Audit readiness checklist



Sooner or later, every regulated company — from medical devices to cosmetics and pharmaceuticals — will face an audit. Use this general checklist to evaluate your organization and take proactive steps to prepare to pass an audit when the time comes.

Management responsibility and quality system	
<input type="checkbox"/>	Are your organization’s quality policy and quality objectives documented?
<input type="checkbox"/>	Does your organization have a quality manual documenting the quality system and its implementation?
<input type="checkbox"/>	Is the quality function separate from the production function?
<input type="checkbox"/>	Are quality procedures and responsibilities documented?
<input type="checkbox"/>	Does the quality department have designated authority for all quality matters?
<input type="checkbox"/>	Is the suitability and the effectiveness of the quality system reviewed by management at defined intervals?
<input type="checkbox"/>	Does your organization have a documented plan for recovery and return to operations following a disaster?
<input type="checkbox"/>	Does your organization have written procedures for handling customer complaints?
<input type="checkbox"/>	Do procedures ensure that root causes of customer complaints are investigated and resolved, and effectiveness of corrective and preventative actions is verified?
<input type="checkbox"/>	Are procedures in place to handle field alerts, recalls or market withdrawals?

Documentation and records	
<input type="checkbox"/>	Are procedures available for the control of documents and records?
<input type="checkbox"/>	Do these procedures address the handling of document changes?
<input type="checkbox"/>	Are there written manufacturing or service instructions?
<input type="checkbox"/>	Is there a process for the approval of procedures/service instructions?
<input type="checkbox"/>	Is there a system for record retention?
<input type="checkbox"/>	Are records retained in accordance with the applicable regulatory requirements?

Facilities	
<input type="checkbox"/>	Are requirements defined for the infrastructure needed to achieve conformity to product requirements?
<input type="checkbox"/>	To prevent product mix-up?
<input type="checkbox"/>	To ensure orderly handling of product?
<input type="checkbox"/>	Is there a security system to assure there is no entry by unauthorized persons?
<input type="checkbox"/>	Is there a written preventative maintenance program for all equipment and critical utilities?
<input type="checkbox"/>	Is there a pest control program including approved insecticides and areas applied?

Equipment	
<input type="checkbox"/>	Are manufacturing and lab equipment and critical utilities qualified according to the written protocols and industry standards?
<input type="checkbox"/>	Is equipment maintained and calibrated according to a preventive maintenance schedule?
<input type="checkbox"/>	Are the calibration maintenance intervals based on the manufacturer's specified frequencies?
<input type="checkbox"/>	Are records maintained for maintenance and calibration operations?
<input type="checkbox"/>	Are there written procedures for cleaning, specifying cleaning agents and methods?
<input type="checkbox"/>	Are changes authorized and approved through quality?
<input type="checkbox"/>	Is there an adequate system to assure that unclean instruments, equipment and machines are not used?
<input type="checkbox"/>	Is there proper storage of cleaned instruments and equipment so as to prevent contamination?
<input type="checkbox"/>	Are there written procedures for cleaning, specifying cleaning agents and methods?
<input type="checkbox"/>	Is there an adequate system for controlling changes to methods, documents related to equipment, machines?
<input type="checkbox"/>	Is re-qualification or re-validation performed post any major changes to the equipment and/or machine?

Purchasing controls and materials	
<input type="checkbox"/>	Is there a system in place to ensure that materials are only purchased from approved suppliers?
<input type="checkbox"/>	Has each supplier of material or component been inspected, evaluated or audited for proper manufacturing controls?
<input type="checkbox"/>	Are batch records used to document the material, equipment, machine and process(es) used in the production?
<input type="checkbox"/>	Are there written procedures for the receipt, testing and release for use of all materials?
<input type="checkbox"/>	Are incoming materials inspected?
<input type="checkbox"/>	Are incoming material/components quarantined until approved for use?
<input type="checkbox"/>	Are rejected components, material and containers quarantined and clearly marked to prevent their use?
<input type="checkbox"/>	Is there a segregated area for non-conforming product?
<input type="checkbox"/>	Is a final inspection performed on the completed product?
<input type="checkbox"/>	Is there a system for handling non-conforming product materials or test results to prevent re-occurrence?
<input type="checkbox"/>	Is each lot within each shipment of material or components assigned a distinctive code so material or components can be traced through manufacturing and distribution?

Production	
<input type="checkbox"/>	Have any of your manufacturing processes been validated?
<input type="checkbox"/>	Does your facility have design control for any products manufactured?
<input type="checkbox"/>	Are planned and unplanned deviations documented?
<input type="checkbox"/>	Is there a procedure for the documentation and investigation of non-conformances?
<input type="checkbox"/>	Is production data reviewed for potential trends in non-conforming product?
<input type="checkbox"/>	Are adverse trends addressed and is appropriate management notified?
<input type="checkbox"/>	If your firm subcontracts the assembly and/or packaging of the product — are contractors frequently audited?

Laboratory controls	
<input type="checkbox"/>	Do you have an on-site laboratory to test incoming materials and finished product?
<input type="checkbox"/>	Are non-conformances tracked and trended?
<input type="checkbox"/>	Are all your contract labs/manufacturers qualified?
<input type="checkbox"/>	Is there a procedure for investigation of out-of-specification (OOS) test results to assure that a uniform procedure is followed to determine why the OOS result occurred and that corrective actions are implemented?
<input type="checkbox"/>	Do your contract labs/suppliers follow current good manufacturing practice /good laboratory practice requirements?
<input type="checkbox"/>	Is there a written specification on how to qualify contract labs/manufacturers?

Change control / notification	
<input type="checkbox"/>	Is there a system in place to manage process changes? If yes, is this process detailed in a written procedure?
<input type="checkbox"/>	Do you routinely notify customers of changes that potentially affect the quality of the product or service provided?
<input type="checkbox"/>	Do you routinely notify customers of key business changes (e.g. acquisitions, manufacturing location changes, etc.)?
<input type="checkbox"/>	Do you notify customers in advance of significant changes to processes/materials? What is the notification process?

Corrective and preventive action	
<input type="checkbox"/>	Are procedures in place for corrective and preventive action?
<input type="checkbox"/>	Are corrective and preventive actions assigned and tracked to closure?
<input type="checkbox"/>	Are records maintained for corrective and preventive actions?
<input type="checkbox"/>	Are corrective and preventive actions reviewed for effectiveness?
<input type="checkbox"/>	Are corrective actions periodically reviewed by top management?

Internal audits / supplier management	
<input type="checkbox"/>	Is there a documented internal audit program in place at your organization?
<input type="checkbox"/>	Are internal audits performed on a schedule and corrective actions taken as appropriate?
<input type="checkbox"/>	Are internal audits periodically reviewed by top management?
<input type="checkbox"/>	Does your organization maintain an approved supplier list?
<input type="checkbox"/>	Does your organization have a procedure/process outlining the selection and monitoring of suppliers providing critical products and/or services?

Training	
<input type="checkbox"/>	Internal QMS training
<input type="checkbox"/>	Job-specific training
<input type="checkbox"/>	Are employee training records maintained in compliance with control of records?
<input type="checkbox"/>	Are personnel qualifications/skills in accordance with the job performed and job descriptions available?



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Sarah Raux

Frameworks

Frameworks provide structured requirements to help achieve and demonstrate regulatory standards.

Active

MDSAP

Implementing

Requirement status

Met (1%)

Attention required (77%)

In progress (6%)

Not started (17%)

ISO 9001:2015

Implementing

Requirement status

Met (4%)

Attention required (71%)

In progress (21%)

Not started (4%)

Inactive

Enable a framework to view requirement status

BETA | ISO 42001:2023

Enable