



Pharmaceutical Vendor Qualification Checklist

 Show only Checklist

Display Style
Default 

Vendor Information & Initial Assessment

Collection of basic vendor details and a preliminary risk assessment.

Vendor Legal Name

Write something...

Vendor Contact Person

Write something...



Vendor Address

Write something...

Years in Business

Enter a number...

Primary Business Type

- ☐ Raw Materials
- ☐ Packaging
- ☐ Equipment
- ☐ Services
- ☐ Other

Initial Risk Assessment Level

- ☐ Low
- ☐ Medium
- ☐ High

Initial Assessment Date

Enter date...

Financial Stability & Business Practices

Evaluation of vendor's financial health and ethical business conduct.

Annual Revenue (USD)

Enter a number...

Debt-to-Equity Ratio

Enter a number...

Credit Rating Agency

☐ Moody's

☐ Standard & Poor's

☐ Fitch

☐ Not Rated

Summary of Financial Stability Assessment

Write something...

Business Ethics Program

- ☐ Yes, documented program
- ☐ Yes, informal policy
- ☐ No program in place

Date of Last Financial Review

Enter date...

Quality Management System (QMS)

Assessment of the vendor's QMS, including policies, procedures, and documentation.

QMS Documentation Availability

- ☐ Complete & Current
- ☐ Partially Available
- ☐ Not Available

Summary of QMS Documentation Reviewed

Write something...

Number of Documented Procedures

Enter a number...

Date of Last QMS Audit

Enter date...

QMS Elements Assessed (Select all that apply)

- ☐ Document Control
- ☐ CAPA
- ☐ Change Management
- ☐ Training
- ☐ Internal Audits
- ☐ Management Review

Copy of QMS Manual (if available)

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Evidence of Management Review

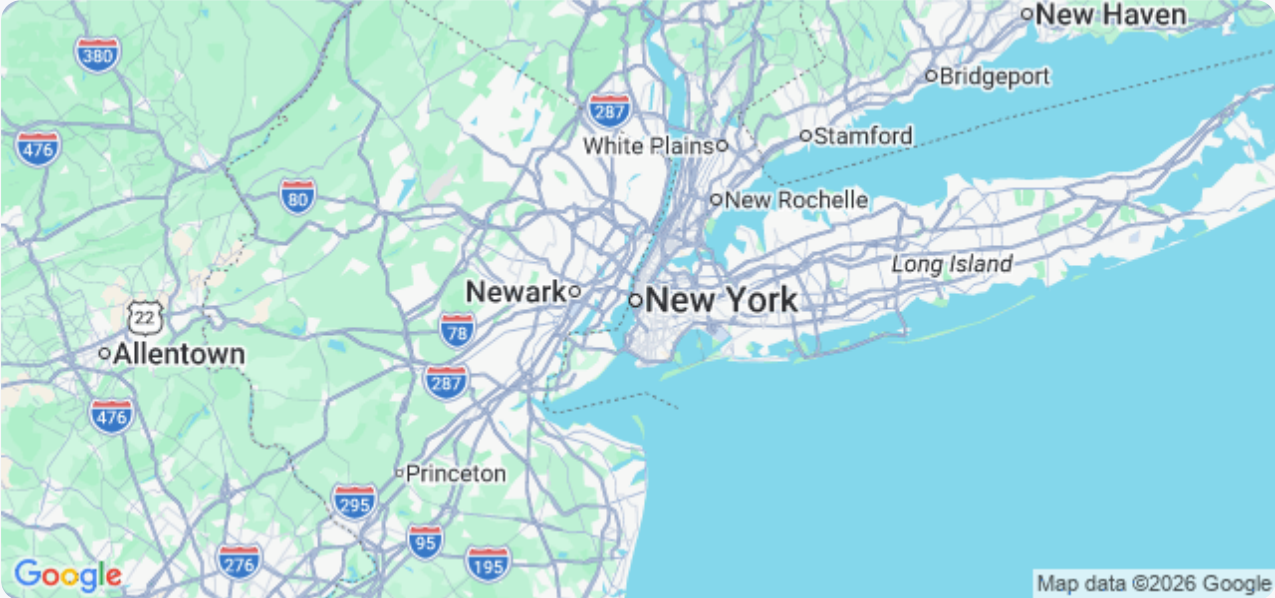
- ☐ Yes
- ☐ No
- ☐ Not Applicable

Facility & Equipment

Verification of the vendor's facilities and equipment suitability for pharmaceutical materials or services.

Facility Address

 Set My Current Location



Map data ©2026 Google

Square Footage of Manufacturing Area

Enter a number...

Equipment Types Present (Select all that apply)

- ☐ Reactors
- ☐ Dryers
- ☐ Mills
- ☐ Filters
- ☐ Packaging Equipment
- ☐ Analytical Instruments

Last Facility Inspection Date

Enter date...

Facility Layout Diagram

 Upload File

HVAC System Temperature Control Range

Enter a number...

Description of Cleaning and Sanitation Procedures

Write something...

Personnel & Training

Review of vendor's personnel qualifications and training programs.

Number of Qualified Personnel

Key Personnel Qualifications (e.g., GMP, Degree)

- ☐ GMP Training
- ☐ Relevant Degree
- ☐ Other

Last Training Completion Date (Key Personnel)

Brief Description of Key Personnel Training Program

Training Records (Example)

 Upload File

Verification of Personnel Background Checks

- ☐ Yes
- ☐ No
- ☐ N/A

Details on Background Check Procedures (if applicable)

Write something...

Regulatory Compliance & Audits

Confirmation of adherence to relevant regulatory requirements and history of audits.

Last Audit Score

Enter a number...

Date of Last Regulatory Inspection

Enter date...

Summary of Findings from Last Regulatory Inspection

Write something...

Compliance with GMP Guidelines?

☐ Yes

☐ No

☐ N/A

Relevant Regulatory Frameworks (Select all that apply)

☐ FDA

☐ EMA

☐ WHO

☐ PIC/S

☐ Other

Copy of Latest Regulatory Audit Report

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Product/Service Specifications & Testing

Validation of product/service specifications and testing protocols.

Detailed Product/Service Specifications

Write something...

Testing Methodology Alignment (e.g., USP, EP, JP)

- ☐ USP
- ☐ EP
- ☐ JP
- ☐ Other (Specify)

Acceptance Criteria Limit (e.g., Purity %)

Certificate of Analysis (CoA)

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CoA Issue Date

Testing Parameters Verified (Select all that apply)

- ☐ Identity
- ☐ Purity
- ☐ Assay
- ☐ Impurities
- ☐ Water Content
- ☐ Other (Specify)

Deviations & Resolutions (if any)

Write something...

Change Control & Corrective Actions

Review of vendor's change control process and management of corrective and preventive actions.

Change Request Originated From:

- ☐ Quality Assurance
- ☐ Manufacturing
- ☐ Engineering
- ☐ Regulatory Affairs
- ☐ Other

Description of Change/Deviation

Write something...

Risk Score (assigned)

Enter a number...

Date of Deviation/Change Initiation

Enter date...

Root Cause Analysis Findings

Write something...

Potential Impact Areas (select all that apply)

- ☐ Manufacturing Process
- ☐ Product Quality
- ☐ Equipment
- ☐ Documentation
- ☐ Regulatory Compliance

Corrective Actions Planned

Write something...

Planned Completion Date of Corrective Actions

Enter date...

Signature of Responsible Person

Contractual Agreements & Performance Monitoring

Assessment of contractual obligations and ongoing performance monitoring processes.

Contract Start Date

Enter date...

Contract Expiration Date

Enter date...

Agreed Upon Price/Rate

Enter a number...

Payment Terms

- ☐ Net 30
- ☐ Net 60
- ☐ Other (Specify)

Key Performance Indicators (KPIs)

Write something...

Performance Rating (Scale 1-5)

Enter a number...

Performance Review Comments

Write something...

Contract Renewed?

- ☐ Yes
- ☐ No

Requalification & Periodic Review

Establishing a schedule for requalification and periodic reviews of vendor status.

Last Requalification Date

Enter date...

Review Frequency (in months)

Enter a number...

Review Type

☐ Document Review

☐ On-site Audit

☐ Combination

Summary of Review Findings

Write something...

Reviewer Signature

Next Review Date

Enter date...