

QUALITY ASSURANCE DEPARTMENT

STANDARD OPERATING PROCEDURE		
Department: Quality Assurance	SOP No.:	
Title: Retention and Disposal of Documents	Effective Date:	
Supersedes: Nil	Review Date:	
Issue Date:	Page No.:	

1.0 OBJECTIVE:

To establish and implement a procedure for retention and disposal of documents used in various departments.

2.0 SCOPE:

This SOP is applicable to the master document and related data available in company.

3.0 RESPONSIBILITY:

Executive - Quality Assurance

Head - Quality Assurance

4.0 **DEFINITION(S)**:

NA

5.0 PROCEDURE:

- 5.1 Retain the master documents and related data for the retention time mentioned as per Annexure-II.
- 5.2 Ensure that retention documents are readily available throughout the period of retention to trace the information. All raw materials, both actives and excepients shall be procured only from approved suppliers.
- 5.3 Ensure that the integrity of the documents are not affected by storage condition throughout the period of retention.
- 5.4 Raise document disposal proposal after the retention period.
- 5.5 Get approval from Manager -QA for disposal of these documents.
- 5.6 Enter the details of the documents to be disposed in 'Document Disposal Record' (enclosed Annexure-I).
- 5.7 Send the documents along with copy of approved document disposal record to housekeeping department for destruction either by Shredding.
- 5.8 Ensure that documents are destroyed.



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5.9 Retain the original copy of the document disposal record for reference.

6.0 ABBREVIATION(S):

BMR – Batch Manufacturing Record

BPR – Batch Packing Record

QA – Quality Assurance

7.0 REFERENCE(S):

NA

8.0 ANNEXURE (S):

ANNEXURE I: Document disposal details

ANNEXURE II: Retention time of documents



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9.0 REVISION CARD:

S.No.	REVISION No.	REVISION DATE	DETAILS OF REVISION	REASON (S) FOR REVISION



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	Annex	ture I	
DOCUMENT DISPOSAL RECORD			
	Name of the Department:		
Type of document (Tick (!) the ap	ppropriate)		11
☐ Standard Operating Procedure	l	☐ Usage Log Books (Equipment and Instruments)	
☐ Qualification and Validation		☐ Analytical Reports, CO	As and supporting data.
(Protocol & Reports)			
☐ Calibration Reports	[☐ Other QC Records (Specifications, STP, Method	
	•	of analysis, RM/PM/FG I	nward records, Stability
	1	records)	
□ BMR / BPR	[☐ Validation Master Plan	
☐ Site Master File	[☐ Complaint files / recall	
☐ Change control proposals	[☐ Destruction records	
☐ Control sample records	[☐ Self-Inspections reports	
☐ Training records	1	☐ Vendor audit reports	
□ OOS / NCR/Deviation records	[☐ Annual Product Review	
☐ Pest control records	[☐ MMF/ Master BMR, BPR	
☐ Maintenance related records	[☐ Warehouse Records (GRN, Inward/Stock register)	
☐ Other Records			
Brief details of the documents to be	destroyed		
Approved By(Manager-QA):			
Disposal	Name	Signature	Date
Done by			
Checked by (QA)			



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Annexure II RETENTION TIME OF DOCUMENTS

S.No.	Document Details	Retention Time
1	Standard Operating Procedure	Permanent
2	Qualification and Validation (Protocol & Reports)	Permanent
3	Calibration Reports	6 Years
4	BMR/BPR	Expiry + 1 Year
5	Usage Log Books (Equipment and Instruments)	6 Years
6	Analytical Reports, COAs and supporting data.	Expiry + 1 Year
7	Other QC Records (Specifications, Method of analysis, RM/PM/FG	6 Years
	Inward records, Stability records)	
8	Site Master File	Permanent
9	Validation Master Plan	Permanent
10	Complaint files / recall	6 Years
11	Change control proposals	Permanent
12	Destruction records	Permanent
13	Control sample records	6 Years
14	Self-Inspections reports	6 Years
15	Vendor audit reports	Permanent
16	Training records	6 Years
17	OOS / NCR/Deviation records	6 Years
18	Annual Product Review	Permanent
19	Pest control records	6 Years
20	MMF/ Master BMR, BPR	Permanent
21	Warehouse Records (GRN, Inward/Stock register)	6 Years
22	Maintenance related records	6 Years
23	Other Records	6 Years
24	Dossiers filed	Permanent