

## PHARMA DEVILS

#### VALIDATION SUMMARY REPORT FOR COMPUTER SYSTEM

System Name: HPLC	System ID:
System Manie: III De	Dystem ID.

## **VALIDATION SUMMARY REPORT**

### **FOR**

### **COMPUTER SYSTEM**

**OF** 

## **HPLC**

System Name	HPLC
System ID	
Location	Instrument Lab
Effective Date	

**Document No.:** Page 1 of 10



System Name: HPLC System ID:

#### TABLE OF CONTENTS

1.	PREPARATION AND APPROVALS	.3
2.	REVISION HISTORY	.3
3.	OBJECTIVE AND SCOPE	.4
4.	REFERENCES	.4
5.	ROLE AND RESPONSIBILITY	.5
6.	DELIVERABLE DOCUMENTS	.6
7.	INSTALLATION QUALIFICATION TEST STATUS	.7
	OPERATION QUALIFICATION TEST STATUS	
	PERFORMANCE QUALIFICATION TEST STATUS	
	ABBREVIATION	
11.	SUMMARY & CONCLUSION	10



**Document No.:** 

# PHARMA DEVILS QUALITY ASSURANCE DEPARTMENT

#### VALIDATION SUMMARY REPORT FOR COMPUTER SYSTEM

System Name:	HPLC			System ID:
The signatur	e listed below	TD APPROVALS:  v indicates the preparation are ability of listed functional area.		his Validation summary report. This
	REPOR	T DEVELOPMENT		SIGN / DATE
Name Designation	:			
	REPOR	T REVIEW AND APPRO	VAL (M/S	)
Sign / Date Name Designation	<b>:</b>	g		
Sign / Date Name Designation	<b>:</b>			
Sign / Date Name Designation	<b>:</b>	ntrol		
	RI	CPORT AUTHORIZATIO	N (M/S	)
Sign / Date Name Designation	:			
2. REVISIO	ON HISTOR	RY:		
Da	nte	Supersedes		Reason for Revision

Page 3 of 10



System Name: HPLC System ID:

#### 3. OBJECTIVE AND SCOPE:

The objective of this summary report is to collect sufficient data and the qualification executed pertaining to the Computer system of HPLC at ......

Successful completion of this document will provide the successfully validated of the computer system of HPLC.

#### 4. REFERENCES:

The publication listed below form part of this report's reference documents. Each publication shall be the latest revision in effect on the date this report is approved for execution unless noted otherwise. Except as modified by the requirements specified herein or the details of the drawings, work included in this report shall conform to the applicable provisions of these publications.

GAMP 5	Good Automated Manufacturing Practices, Version 5, Guideline document for Automated Systems from International Society of Pharmaceutical Engineering
21 Code of Federal Regulations (CFR), Part 210	Current Good Manufacturing Practice in Manufacturing, Processing, Packing, or Holding off Drugs; General
21 Code of Federal Regulations (CFR), Part 211	Current Good Manufacturing Practice for finished Pharmaceuticals
WHO	Appendix 5, validation of computerized systems.
VP	-
SRS	-
RA	-
IQ	-
OQ	-
PQ	-
TM	-

**Document No.:** Page 4 of 10



# PHARMA DEVILS GUALITY ASSURANCE DEPARTMENT

#### VALIDATION SUMMARY REPORT FOR COMPUTER SYSTEM

System Name: HPLC System ID:

#### 5. ROLE AND RESPONSIBILITY:

The Validation team comprising of representative from each of the following departments should be responsible for overall compliance with this validation plan.

Department	Responsibilities	
	> To collect data necessary for the generation, execution of this report from	
	M/s	
Validation Agency	> To prepare the validation summary report.	
()	> Comply with regulatory / Guidelines / Standards / validation plan	
	requirements throughout the validation life cycle.	
	> To submit validation documents for approval.	
	Project Management and Planning.	
Engineering	> To provide the necessary data for qualification activities.	
(M/s)	> To co-ordinate during execution of qualification activities.	
	> To review and approve the validation documents.	
	> To provide the necessary data for qualification activities.	
IT	> To co-ordinate during execution of qualification activities.	
(M/s)	> To review and approve the validation documents.	
	> To provide the necessary data for qualification activities.	
Quality Control	> To co-ordinate during execution of qualification activities.	
(M/s)	> To review the validation documents.	
<b>Quality Assurance</b>	> To approve and authorize the validation documents.	
(M/s)		

**Document No.:** Page 5 of 10



System Name: HPLC System ID:

#### 6. DELIVERABLE DOCUMENTS:

- Validation Plan
- > System Requirement Specification
- > Gap and Risk Assessment
- > Installation Qualification
- > Operational Qualification
- > Performance Qualification
- > Traceability Matrix
- > Validation Summary Report

**Document No.:** Page 6 of 10





System Name: HPLC System ID:

### 7. INSTALLATION QUALIFICATION TEST STATUS:

S.No.	Critical Feature	Pass/Fail	Discrepancy (Y/N)	Checked By / Date
1.	Identification of System Details			
2.	Verification of Master Documents			
3.	Verification of Capacity Requirement			
4.	Verification of Hardware Components			
5.	Verification of Software Components			
6.	Verification of Physical and Logical Security Control			
7.	Verification of Test Instruments Calibration and Traceability			
8.	Verification of Power Utility			
9.	Verification of Environmental Condition			
10.	Verification Of Communication Link Between Server to Computer System			
11.	Verification of General System Installation			
12.	Verification of Standard Operating Procedures			

**Document No.:** Page 7 of 10



# PHARMA DEVILS QUALITY ASSURANCE DEPARTMENT

#### VALIDATION SUMMARY REPORT FOR COMPUTER SYSTEM

System Name: HPLC System ID:

### **8. OPERATION QUALIFICATION TEST STATUS:**

S.No.	Critical Feature	Pass / Fail	Discrepancy? (Y/N)	Checked By / Date
1.	Verification of Field Instruments Calibration			
2.	Verification of Windows Security			
3.	Verification of System Start-up & Shutdown.			
4.	Verification of Password Security			
5.	Verification of User Level and Rights			
6.	Verification of Application software Screens.			
7.	Verification of System Response Failure.			
8.	Verification of Electronic Data Security.			
9.	Verification of Audit Trail .			
10.	Verification of Report Generation.			
11.	Verification of User Prevented from Alternating			
11.	Date and Time			
12.	Verification of Data Back Up			
13.	Verification of system software as per 21 CFR part			
13.	11 Clauses			

Document No.: Page 8 of 10



## PHARMA DEVILS

#### VALIDATION SUMMARY REPORT FOR COMPUTER SYSTEM

System Name: HPLC System ID:

### 9. PERFORMANCE QUALIFICATION TEST STATUS:

S.No.	Critical Feature	Pass / Fail	Discrepancy? (Y/N)	Checked By / Date
1.	Verification of Control Loop Test			

**Document No.:** Page 9 of 10





System Name: HPLC	System ID:
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#### 10. ABBREVIATION:

Abbreviations	Description
GMP	Good Manufacturing Practices
IS	Information Services
IQ	Installation Qualification
OQ	Operation Qualification
PQ	Performance Qualification
QA	Quality Assurance
TM	Traceability Matrix
SOP	Standard Operating Procedure
SRS	System Requirement and Specification
TS	Technical Services
WHO	World Health Organization

TS	Technical Services
WHO	World Health Organization
11. SUMMARY	& CONCLUSION:
Compiled by:	Date:

Document No.: Page 10 of 10