

QUALITY ASSURANCE DEPARTMENT

### VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name:	System ID:
System Name.	System ID.

## **VALIDATION PLAN**

## **FOR**

## **COMPUTER SYSTEM OF**

**UV (MICROBIOLOGY SECTION)** 

System Name	
System ID	
Location	
Effective Date	

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## VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

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1. PREPAR	ATION AND APPRO	VALS:	
			al of this Validation Plan. This approval is
	ibility of listed functions		01 v v v 1 upp10 v.u.
John Tespons	iomity of fished functions	ar arcas.	
	DOCUMENT DEVI	ELOPMENT	SIGN / DATE
Name	<b>:</b>	_	
Designation	•	_	
	DOCU	MENT REVIEW AND AI	PPROVAL
Sign / Date	;	_	
Name	<b>:</b>	_	
Designation	•		
	Engineering		
Sign / Date	;	_	
Name	<b>:</b>	_	
Designation	<b>:</b>	_	
	IT		
Sign / Date	:	_	
Name	<b>:</b>	_	
Designation	:	_	
	<b>Quality Control</b>		
		DOCUMENT APPROVA	AL
Sign / Date	<b>:</b>	_	
Name	:	_	
Designation	•	_	
-	<b>Quality Assurance</b>		

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VALIDA	ATION PLAN FOR	COMPUTER SY	STEM OF UV	
System Name:			Syste	em ID:
2. SIGNATURE OF EXECUTABLE All the executer involved in the M/s		to sign within pres	cribed format give	en below.
Name	Designation	Signature	Initial	Date
M/s				
Name	Designation	Signature	Initial	Date
3. REVISION HISTORY:				
Date	Supersedes	I	Reason for Revisi	on

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#### VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

#### 4. OBJECTIVE:

The objective of Validation Plan is to provide an organization approach towards the validation activities for the Computer System hardware and software of QC Department UV. This document will define the requirement and standards that must be followed for all the validation activities as apply to the Computer System of UV.

#### 5. SCOPE:

This document is applicable to validation of Hardware and Software system of Computer System (UV). This document shall define the test procedures, documentation, references and acceptance criteria in accordance with the guidelines laid down by the manufacturer of the system.

#### 6. SYSTEM DESCRIPTION:

Computer system of UV defines the controlling of analytical instrument connected to the system. It controlled the process of instrument connected to it by the software installed in the CS. The operator interface is carried out by CS screen. The CS is used to feed required parameters and set points in the system during operation. The system is connected to data server for printing and data backup. The system is secured by IT through password only within the system.

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## VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

## 7. 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 the necessary data for qualification activities.	
	> To prepare the Validation Plan, Risk Assessment, Installation Qualification,	
	Operational Qualification, Traceability Matrix and Validation Summary	
	Report.	
Validation Agency	> To execute the qualification in coordination with engineering, validation and	
	quality assurance team.	
	Comply with regulatory / Guidelines / Standards / validation plan requirements	
	throughout the validation life cycle.	
	> To submit qualification for approval.	
	> To provide the necessary data for system requirement specification activities.	
Engineering (M/s)	<ul> <li>To provide the necessary data for system requirement specification activities.</li> <li>To review Validation Plan.</li> </ul>	
(WI/S)	To review varidation Fian.	
	To provide the passessary date for system requirement specification activities	
IT (M/s)	<ul> <li>To provide the necessary data for system requirement specification activities.</li> <li>To review Validation Plan.</li> </ul>	
(W1/S)	To review varidation Flan.	
Quality Control	> To provide the necessary data for system requirement specification activities.	
(M/s)	> To review Validation Plan.	
Quality Assurance	> To approve and authorized the Validation Plan.	
(M/s)		

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#### VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

#### 8. REFERENCES:

The publication listed below form part of this reference documents. Each publication shall have latest revision in effect on the date of this document is approved for execution.

GAMP 5	Good Automated Manufacturing Practices, Version 5, Guideline	
	Document for Automated Systems from International Society of	
	Pharmaceutical Engineering	
21 Code of Federal	Current Good Manufacturing Practice in Manufacturing, Processing,	
Regulations (CFR), Part	Packing, or Holding off Drugs; General	
210		
21 Code of Federal		
Regulations (CFR), Part	Current Good Manufacturing Practice for finished Pharmaceuticals	
211		
21 Code of Federal	21 Code of Federal Regulations (CFR), Part 11 Electronic Records,	
Regulations (CFR), Part	Electronic Signatures, Final Rule Electronic Submissions;	
11	Establishment of Public Docket, Notice	
ICH Q9	International Conference of Harmonization (ICH) quality risk	
	assessment Q9	
EU GMP	Laying down the principles and guidelines of GMP in respect of	
	medicinal products for human use.	
WHO	Appendix 5, validation of computerized systems.	

#### 9. DOCUMENTATION PROCEDURE:

- Qualification activities will be performed as defined in the approved document.
- All documentation will be completed during the execution of the qualification.
- Recording of information will be made in permanent ink.
- Fill out complete information in the verification table provided.
- Do not keep any space blank. Mark blank space with a single line throughout the appropriate space with mentioning NA (Not Applicable) and put initial and date.
- Correct the mistakes by drawing a single line through the incorrect data, recording the correct information and then initial sign and date the change.

### 10. QUALIFICATION COMPLETION AND APPROVAL:

- Verify that all tests required by qualification are completed and attached.
- Verify that all amendments and discrepancies are documented, approved and attached.
- If all items in the qualification for the Computer System of UV have been reviewed and found to be acceptable, sign the corresponding block in the qualification completion and approval form.

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## VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

#### 11. ACCEPTANCE CRITERIA:

- Installation completion as per manufacturer's recommendations & cGMP requirements.
- The supply of all necessary documentation from manufacturer/Installer.
- The system is operating as intended and is under state of control.
- Operational features meet system requirements and system specifications.

#### 12. DOCUMENTATION MANAGEMENT:

All quality and project relevant documents delivered by ..... are handled through ......'s document management system. Each document has a unique ID and is version. The identification number of a document has the following structure:

<.....>-<Project No.>-<Document Name>-Version

The author's name, the file name, the document number (document code and Revision No.) and the total pages number are included in the document footer in order to clearly assign each page to a certain document.

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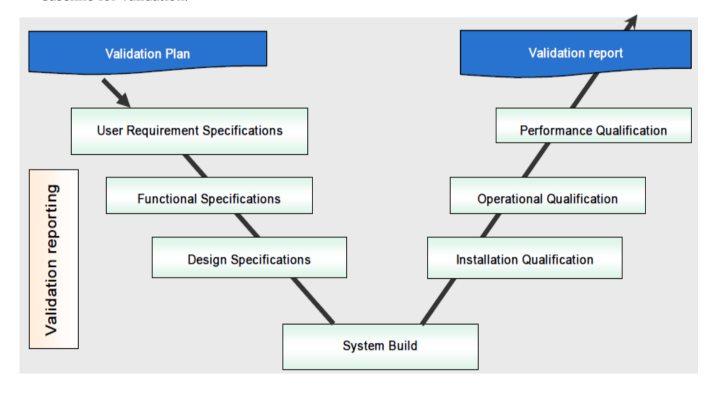
System Name: System ID:

#### 13. REFERENCE DOCUMENTS:

- a. Standard operating procedures
- b. System requirement specifications
  - System operation manual
  - System bill of material

#### 14. V-MODEL OF GAMP:

The system development life cycle is based on the GAMP-5 development life cycle and the ISPE baseline for validation.



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### VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

#### 15. VALIDATION APPROACH:

For the validation, GAMP 5 guidelines have been considered. As per GAMP Software Life Cycle approach is considered for all automated control systems. The following table depicts categorization of various software systems as per GAMP 5.

The system categorization is intended to evaluate and determine appropriate validation activities and deliverables. Once a system is evaluated as a whole, the functionality of individual components can be assessed for potential risk to data integrity and tested accordingly.

In determining the system categorization, functionality and intended use of the system are to be considered.

Category	Software Type	Validation Approach
3	Non-configured Software e.g. Firmware based application COTS software Instruments	<ul> <li>Abbreviated Lifecycle approach</li> <li>Risk based approach to supplier assessment</li> <li>Record version (and configuration of environment) and verify correct installation.</li> <li>Risk based tests against requirements. (calibrations for instruments)</li> <li>Procedures in place for maintaining compliance and fitness for intended use.</li> </ul>
4	Configurable Software Packages, e.g. DAS IPC ERP DCS BMS LIMS HMI	<ul> <li>Life Cycle approach</li> <li>Risk based approach to supplier assessment.</li> <li>Record version number, verify correct installation</li> <li>Risk based testing to demonstrate applicable works as designed in a test environment and within the business process.</li> <li>Procedure in place for maintaining compliance and fitness for intended use.</li> <li>Procedures in place for managing data.</li> </ul>
5	Custom Software e.g. internally or externally developed IT applications. Custom ladder logic Spreadsheets (macro)	Same as configurable, plus:  More rigorous supplier assessment.  Possession of full life cycle documentation  Design and source code review

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#### VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

#### 16. CATEGORIZATION OF THE CONTROL SYSTEM:

#### 16.1 Computer System

The Computer control system falls under the **Category-3** Non-configurable software package as defined by **GAMP-5** guidlines. hence, verification & configuration and testing of operation against user requirement will be performed.

## 16.2 Hardware Category And Software Category

- Hardware Category 1 Standard hardware component
- Hardware Category 2 Custom built hardware component

Category	GAMP-4	GAMP-5
1	Operating Software	Infrastructure Software
2	Firmware	No longer used
3	Standard Software	Non configured Software
4	Configurable Software	Configured Software
5	Custom Software	Custom Software

#### 17. DOCUMENT SCOPE:

The documents scope of this validation plan is to establish the project framework for carrying out quality assurance and project management measures impacting ...... and M/s ...... for the project as described. The documents scope should define the activities to be performed, which will perform them, the control mechanisms to be used and the deliverables

- Validation Plan
- System Requirement Specification
- Risk Assessment Protocol
- Installation Qualification
- Operation Qualification
- Performance Qualification
- Traceability Matrix
- Validation Summary Report

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## VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

## 17.1 Risk Assessment

	This document is to provide the analyses the risk of utilization of the Computer
	Systemof UV as per the cGMP and GxP and to identify the possible areas of risk,
Definition	where the existing laid down appropriate controls or measures requires further
	strengthening. To suggest suitable solutions (action plan) to mitigate or minimize the
	risk.
Phase	Designing
Control	Review
Executor	-
Prerequisites	SRS is approved
Acceptance	-
Outcomes	Risk Assessment

## 17.2 Installation Qualification

	The objective of the installation qualification test is to verify the Computer System of
	UV installed at the M/s
	This includes the following tests:
	Identification of System Details
	Verification of Master Documents for computer system
	Verification of capacity Requirement of computer system
Definition	Verification of Hardware Components
	Verification of Software Components
	Verification of Physical and Logical Security Control
	Verification of Test Instruments Calibration and it's Traceability
	Verification of Power Supply
	Verification of Environmental Condition

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## VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

	<ul> <li>Verification of Communication Link Between Server To Computer System</li> <li>Verification of General System Installation</li> <li>Verification of Standard Operating Procedures</li> </ul>
Phase	Commissioning
Control	Review
Executor	-
Prerequisites	Risk Assessment is Pre approved
Acceptance	M/s
Outcomes	Installation Qualification

## 17.3 Operational Qualification

	The objective of the operational qualification test is to verify the function of Computer			
	System of UV installed at the M/s			
	This includes the following tests:			
	Verification of Field Instrument Calibration			
	Verification of Windows Security			
Definition	Verification of System Start-up & Shutdown			
	Verification of Password Security			
	Verification of User access and security features of the system			
	Verification of Application software Screens.			
	Verification of System Response Failure.			
	Verification of Electronic Data Security.			
	Verification of Audit Trail.			
	Verification of Report Generation.			
	Verification of User Prevented From Alternating Date and Time.			

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## VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

	Verification of Data Back Up.
	Verification of system software as per 21 CFR part 11 Clauses.
Phase	Commissioning
Control	Review
Executor	
Prerequisites	Risk Assessment is Pre approved
Acceptance	M/s
Outcomes	Installation Qualification

## 17.4 Performance Qualification

Definition	The objective of the performance qualification test is to verify the function of
	Computer System of UV installed at the M/s
	This includes the following tests:
	Verification of Control Loops Test
Phase	Commissioning
Control	Review
Executor	
Prerequisites	Operational Qualification is approved
Acceptance	M/s
Outcomes	Performance Qualification

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## VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

## 17.5 Traceability Matrix

Definition	The traceability matrix is to provide the assurance that mapped between IQ and OQ.  The traceability matrix contains all the traceability mentioned in system requirement specifications.
Phase	Commissioning
Control	Review
Executor	
Prerequisites	SRS, IQ and OQ is approved
Acceptance	M/s
Outcomes	Traceability Matrix

## 17.6 Validation Summary Report

	This validation summary report is to collect sufficient data and the qualification
	executed pertaining to the Computer Systemof UVSuccessful completion of
Definition	this document will provide the successfully validated of the Computer Systemof
	UVThis report describes the successful validation qualification for the
	Computer Systemof UV
Phase	Commissioning
Control	Review
Executor	
Prerequisites	IQ and OQ is approved
Acceptance	M/s
Outcomes	Validation Summary Report

## 18. CHANGE CONTROL

All changes in the control system during the validation activities shall be handled as per the change control SOP.

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#### VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

#### 19. MAINTENANCE AND SUPPORT:

#### 20. STANDARD OPERATING PROCEDURE:

A number of SOP's will be developed for the operations that support the control systems during this validation exercise. Each SOP is listed below.

- System Security
- Desktop Policy for Computer Operated
- Data Backup, Archiving and Retrieval
- Change Control
- Software in Laboratory and GMP System
- System operation

#### 21. DISCREPANCIES HANDLING DURING COMPUTER SYSTEM QUALIFICATION:

- In case of discrepancy observed during qualification, document in the defined column in each table and document the details of the observation in the discrepancy log sheet.
- Inform to engineering IT and quality assurance about discrepancy.
- Investigate the discrepancy and ensure the possible impact.
- If discrepancy does not have potential to impact on operation as well as performance of the system, close the discrepancy with proper justification.
- The engineering, IT and QA will decide whether discrepancy is acceptable or not.
- If discrepancy is acceptable, provide conclusion and recommendation if any into respective column.

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# PHARMA DEVILS

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ystem Name:	System ID:	
DISCREPANCY AND CORREC	TIVE ACTION FORM:	
Protocol Reference		
Discrepancy Number		
DISCREPANCY:		
Describe the Discrepancy		
Reported by	Date	
CORRECTIVE ACTION:		
Describe corrective action taken	(Attach additional sheets if necessary)	
Describe corrective action taken	(Attach additional sheets if necessary)	
Describe corrective action taken	(Attach additional sheets if necessary)	
Describe corrective action taken	(Attach additional sheets if necessary)	
Reported by	(Attach additional sheets if necessary)  Date	
Reported by		
Reported by		
Reported by DISPOSITION ACTION:	Date	
Reported by DISPOSITION ACTION: Acceptable? Yes	Date	
Reported by DISPOSITION ACTION: Acceptable? Yes	Date	
Reported by DISPOSITION ACTION: Acceptable? Yes	Date	
Reported by DISPOSITION ACTION: Acceptable? Yes	Date	
Reported by DISPOSITION ACTION: Acceptable? Yes Discussion	No	

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## VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

### 22. APPROVAL AND DELIVERABLES:

The complete validation is governed by a series of quality assurance measures. The following table lists the validation deliverables. It is assumed that all documents are submitted by ..... as required.

Deliverable	Original Location	Validation Agency (Developer)	M/S (Reviewer)	M/S (Reviewer)	M/S (Reviewer)	M/S (Approver)
Validation Plan System Requirement Specfication Risk Assessment Protocol Installation Qualification Operational Qualification Traceability Matrix Validation	Validation Dept.	(Developer)	Engineering	IT	Quality Control	Quality Assurance
Summary Report						

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## VALIDATION PLAN FOR COMPUTER SYSTEM OF UV

System Name: System ID:

## 23. ABBREVIATION:

Abbreviations	Description		
GMP	Good Manufacturing Practices		
SRS	SystemRequirement Specification		
RA	Risk Assessment		
IQ	Installation Qualification		
OQ	Operation Qualification		
PQ	Performance Qualification		
QA	Quality Assurance		
TM	Traceability Matrix		
VSR	Validation Summary Report		
SOP	Standard Operating Procedure		
NA	Not Applicable		
IO	Input Output		
ICH	International Conference of Harmonization		
CS	Computer System		
ID	Identification		
WHO	World Health Organization		

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