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## Abstract

This research will focus on upgrading the Chromatography System 3 to reduce time and generate batch reports. In this way, system monitoring is more effective and documentation is more accurate. The system handles an excessive amount of equipment where each one is critical to the process and with the help of the upgrade the operator can be more effective with automation. The process should ensure the quality, safety, and efficiency of the manufactured chromatography system. Revalidation will be used as the only method since it is an upgrade of an existing and validated system.

**Key Terms** — Automation, Chromatography System, Installation Operation Performance Qualification (IOPQ), Validation.

## Problem Statement

Cost reduction, efficiency, and process improvement within biological and pharmaceuticals industries is a very well-known matter in question that business has been facing in today's day. This research in analyzing and upgrading the Chromatography System is focused not only on the reduction of data printing time but also on the lower paper consumption. The validation process in this research is very important because it reduces costs by reducing rejects, reworks, and downtime. The chromatography process is a complex process where flow, pH, conductivity, temperature, among others, are monitored all the time. The system currently has an operator interface terminal, which will be upgraded to a control and data acquisition system. The new system will monitor the process and collect the necessary data (pressure, flow rate, UV, date/time, process steps and other data), date/time, process steps and other information) to generate the necessary reports (pressure, flow, UV, date/time, process steps and other information) to generate the necessary reports including data trends, audit trail and others.

## Research Objectives

The objective of this research is to find a way to mitigate the time-wasting factors in handling reports, data, system verification. With the upgrade of the new software, the manufacturing area can achieve better management of reports, documentation, and monitoring in less time and paper.

## Research Contributions

This project contributes to maximum efficiency in the handling of documentation at the industrial level. With this Project.

- The Chromatography Skid # 3 operator interface terminal (OIT) will be upgraded to a Supervisory Control and Data Acquisition (SCADA) system.
- The SCADA systems will be connected to the Automation Integration Layer network. This capability will provide a Wonderware (System Platform) SCADA client along with Wonderware's InBatch solution to manage the Chromatography process using an S-88 Batch Management system.
- The system will be connected to the OSI PI System to support batch data storage.
- Rt Reports will be used to develop an electronic report to replace the paper.

The following are some of the advantages that the project will have Mitigate time loss: It will be possible to improve the workflow with manufacturing operators and quality experts. Cost reduction in using the same software, lowering the use of paper in the industry, and employee time in printing documentation.

## Background

This project will automate the Chromatography Manufacturing purification process. This software will be implemented on manufacturing equipment in the purification area in the biopharmaceutical industry, for this reason, it is important that this software complies with the regulations of the pharmaceutical industry. The Chromatography System 3 consist of a Semi-purified product tank, pump, filter, liquid control panel, buffer solution supply, solution supply flow control valves, air traps, manual actuator valves, operator interface, pressure sensors, level sensor, flow sensor, pH sensor, conductivity sensor, temperature sensors and Chromatography Column.

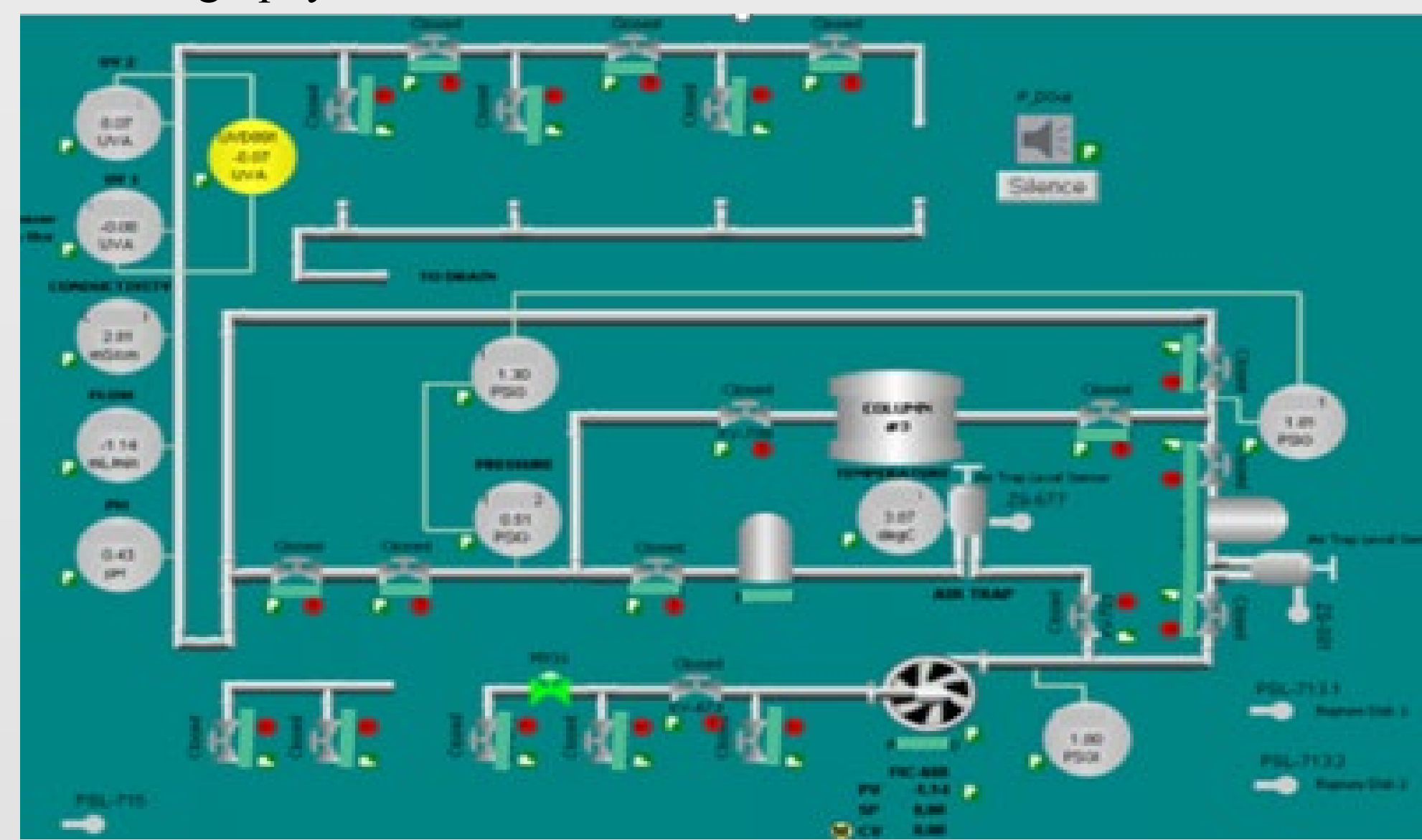


Figure 1: Chromatography System Skid #3

Validation is classified into four types. In this case, a revalidation of equipment and processes was carried out. Revalidation helps to ensure that modifications and upgrades to process environments, whether intentionally or unintentionally whether intentionally or unintentionally introduced, do not affect adversely affect the process characteristics of the product.

According to the US Food and Drug Administration (FDA) the assurance of product quality is derived from careful and systemic attention to a number of important factors, including selection of quality components and materials, adequate product and process design and statistical control of the process through in-process and end product testing [1].

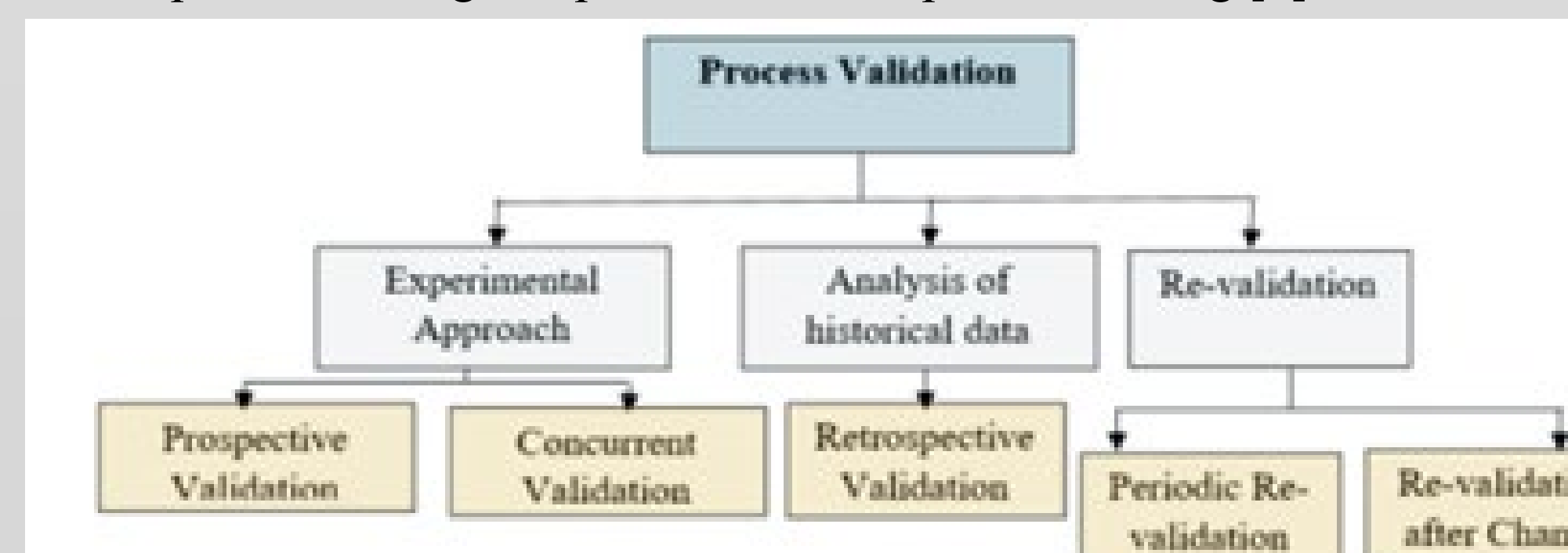


Figure 2: Classification of Validation

The equipment qualification test must be passed before starting the validation. The main stages of qualification include design qualification, Installation Qualification (IQ), Operational Qualification (OQ), and Performance Qualification (PQ). These four major documents are commonly used to validate every piece of manufacturing machinery [4].

The System monitors the process, gathering required data (i.e. Pressure, flow, UV, date/time, process steps and other information) to generate the required electronic reports including data trending, audit trail, HETP calculations and others. Process automation manages business processes for uniformity and transparency and typically handled by dedicated software and business software. Automation is desirable if the highest productivity is anticipated and to eliminate most human influence.

## Methodolgy

With the information obtained, we will be able to analyze if our project will be effective. In addition, we ensure the fulfillment of one of the objectives, where the RtReports system can be used to generate electronic reports.

Once the proposal has been analyzed and approved, the upgraded system and resources for the project will be acquired. With the acquisition, we proceed to perform qualification exercises. In this process, it is necessary to generate the necessary documentation, such as the System design, Validation Plan, IOQ protocol, Summary Report, SOP. This step will test compliance with 21 CFR Part 11 of the federal regulations. The computerized system will be tested for compliance with data integrity.

The initial plan is to implement this new system on Chromatography Systems 3 at Purification. In this phase of the project, the Chromatography System used is evaluated:

- Installation of the upgraded System
- Activation of the alarms
- Creation of the recipe in InBatch System
- Creation of electronic reports.

The documents are developed to help define the requirements of the business. The documents are developed to help define the requirements of the business. The User Requirements Specification (URS) defines the business user needs of System. The purpose of the System Design Specification is to provide a description of the Chromatography Skid system in terms of the functions that it will perform, and the facilities required to satisfy the URS. The Installation and Operational Qualification Protocol (IOQP) define the objectives, methodology, acceptance criteria, documentation verification and test activities required to provide evidence that the Chromatography Skid #3 was installed per requirements.

## Results and Discussion

As part of the system testing, full sequence of process steps according to the applicable approved operational procedure and recipe was challenged with the following conditions: The buffers solutions currently used for equilibration and purification of the column were used to test flow and volume controls for each step in its approved process sequence. Information shall include Test, Acceptance Criteria, and Pass/Fail. Actual results must confirm a passing result or acceptable test case. Upon completion of the qualification, the final releases were backed up to show that the SCADA is included in the plant backup system.

Table 1: Installation Qualification Test Cases

Test Case#1	Pre-Requisites Verification	Acceptance Criteria	Pass/Fail
	Change Control	Approved	Pass
	User Requirements Specification for Skid #3	Approved	Pass
	System Design Specification for Skid #3	Approved	Pass
Test Case #2	Calibration Verification	Acceptance Criteria	Pass/Fail
	Instrumentation associated to the Controls System upgrade project.	Not past due	Pass
Test Case #3	Wiring and Drawings Verification	Acceptance Criteria	Pass/Fail
	New electrical wiring for the Control System upgrade was verified against approved drawings.	Verified	Pass
	All monitored system conditions have the correct associated graphic color/icons and all measured variables have the correct associated values.	Verified	Pass
Test Case #4	Electrical Utilities Verification	Acceptance Criteria	Pass/Fail
	Voltage (VAC)	115 ± 10%	Pass
	Chassis Ground Connected	Yes	Pass
	UPS Service	Yes	Pass
	Frequency	60 Hz ± 2%	Pass
Test Case #5	Environmental Condition Verification	Acceptance Criteria	Pass/Fail
	Temperature	0° C to 50° C	Pass
	Relative humidity	10% to 80%	Pass
Test Case #6	Screen Test Verification	Acceptance Criteria	Pass/Fail
	All phase, Control Modules, Equipment Module, and process trending screen modules function per specifications	Displayed	Pass
Test Case #7	Interlocks / Manual Operation Verification	Acceptance Criteria	Pass/Fail
	Local interlocks associated to the Skid Y-1730 control system operate in accordance to design specifications. System components can be operated manually.	Message displayed	Pass
Test Case #8	Warning and Alarms Verification	Acceptance Criteria	Pass/Fail
	Warnings and Alarms associated to the Skid Y-1730 control system operate in accordance to design specifications.	Message displayed and verified	Pass

Table 2: Operational Qualification Test Cases

Operational Qualification			
Test Case#9	Operational Procedures Verification	Acceptance Criteria	Pass/Fail
	Operational procedure(s) challenged and updated as applicable. Redlined procedures are required to be approved before system release.	Verified	Pass
Test Case #10	Flow Control System Verification	Acceptance Criteria	Pass/Fail
	Set flow to 1,000 ml/min	Process value maintained	Pass
	Set flow to 2,000 ml/min	Process value maintained	Pass
Test Case #11	Phase sequence Test Verification	Acceptance Criteria	Pass/Fail
	Follow the test procedures during this test execution. The Equipment Phases can be executed independently in semiautomatic mode per specifications.	Displayed	Pass
Test Case #12	Recipe Sequence/ Automatic Test Verification	Acceptance Criteria	Pass/Fail
	Follow the test procedures during this test execution. The certified InBatch recipe can be executed in automatic mode per specifications. The system has Unit Procedures and Operational Procedures per S88 standard architecture.	Displayed	Pass
Test Case #13	Report Data Verification	Acceptance Criteria	Pass/Fail
	The data storage system stores the batch data as electronic records.	Displayed	Pass
	The system produces Batch reports accurately retaining the meaning and content of the electronic records.	Properly generated	Pass
	Batch Report data is accurate and complies with user requirements (User ID, Batch, Batch status, Start and End times, Operational Unit, Recipe, Bath Summary [Alarms, CPPs, Duration, Material Addition, Reconciliation Trends]).	Information is contained	Pass
	Verify that the batch report is available in electronic format (e.g PDF).	Exported	Pass
Test Case #14	Preventive Maintenance Verification	Acceptance Criteria	Pass/Fail
	Verify SAP Preventive Maintenance related to the project was updated.	Updated	Pass
Test Case #15	Electronic Record Protection verification	Acceptance Criteria	Pass/Fail
	Access to the system drive files and folders where Electronic Records and Electronic Signatures are stored was not allowed to Process Operator, Engineer, and Supervisor account levels.	Allowed	Pass
Test Case #16	Audit Trail Verification	Acceptance Criteria	Pass/Fail
	Audit trail that records user entries and actions to create or modify electronic records is electronically available for review.	Performed	Pass

## Conclusion

The objective of this project was to simplify the monitoring of the purification process of System Skid#3 in the Pharmaceutical Industry. All Test Cases included as part of the Process Qualification (IOPQ) have been successfully documented and closed. Based on the comparison of the results presented in this report against the acceptance criteria, it can be concluded that the Skid #3 System Upgrade sequence challenged during the installation and operational validation activities was completed successfully. Reports generated with Batch Data were verified and approved by Quality Personnel. Consequently, by upgrading to the new software, the manufacturing area can achieve better reporting, documentation and monitoring in less time and paper. This system upgrade will allow for more accurate and automated monitoring, where the need for operators per shift can be reduced. In addition, the advantages of implementing this system upgrade are more cost effective which prevents unauthorized access to the system to comply with 21 CFR Part 11 of the Code of Federal Regulations, ensuring a standard of quality and compliance with data integrity (Refer to Tables 1 and 2).

## References

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