

PRODUCTION DEPARTMENT

## **USER REQUIREMENT SPECIFICATION**

NAME OF ITEM: VIAL WASHING MACHINE	PROTOCOL No
FUNCTIONAL AREA: PRODUCTION	Page No.: 1 of 12

## USER REQUIREMENT SPECIFICATION

NAME OF THE ITEM: VIAL WASHING MACHINE

FUNCTIONAL AREA: WASHING AREA

**PROTOCOL No. :** 

**CONTENT** 



PRODUCTION DEPARTMENT

## USER REQUIREMENT SPECIFICATION

NAME OF ITEM: VIAL WASHING MACHINE	PROTOCOL No
FUNCTIONAL AREA: PRODUCTION	Page No.: 2 of 12

## S.No. <u>Description</u>

URS Approval
Objective
Responsibilities
Equipment Description & Identification
User Requirements
Complementary aspects
Safety and environmental Protection
Cleaning maintenance and service
Rules and Regulation
Scope of Delivery
Installation ,Commissioning and Tests
Qualification/Validation
Guarantee/Warrantee



PRODUCTION DEPARTMENT

## USER REQUIREMENT SPECIFICATION

NAME OF ITEM: VIAL WASHING MACHINE	PROTOCOL No
FUNCTIONAL AREA: PRODUCTION	Page No.: 3 of 12

## 1.0 URS APPROVAL:

### **Protocol Prepared By:**

Functional area	Name	Signature	Date
Production			

### **URS Reviewed By:**

Functional area	Name	Signature	Date
Production			
Quality assurance			
Engineering			

### **URS Approved By:**

Functional area	Name	Signature	Date
Head Engineering			
Head Manufacturing			
Head Quality			



PRODUCTION DEPARTMENT

### **USER REQUIREMENT SPECIFICATION**

NAME OF ITEM: VIAL WASHING MACHINE	PROTOCOL No
FUNCTIONAL AREA: PRODUCTION	Page No.: 4 of 12

#### 2.0 Objective:

The purpose of this document is to ensure that all the critical aspects of the Equipment, cGMP & Safety features have been considered in designing the equipment/instrument and is properly documented.

#### 3.0 Responsibilities:

In accordance with the document, following functions shall be responsible for initiation and finalization of Equipment user requirement specification. When the work is carried by contract/ consulting staff, all the work is to be performed under the oversight of .....

#### 3.1 Preparation of Document

- User department to prepare the URS
- Ensures that the document is in compliance with current policies and procedures of cGMP regulations.
- Ensures that the content is sufficient, clearly defined, technically sound and accurate.
- It is a Guidance document to prepare the URS.

#### **3.2 Review of Document**

• To be reviewed by Head of the user department and functional department (Engineering & Quality assurance)

#### 3.3 Approval of Document

• Approval of document by Head Manufacturing/Head Engineering/Head Quality.

#### 4.0 Equipment Description & Identification:

#### 4.1 Scope:

This document covers all aspects of Users requirements for the Equipment along with all Attachment, Spare Parts, Change Parts and Accessories to be used in .....

Scope incorporates understanding and documentation of critical requirements such as system requirements, cGMP requirements, safety requirements, documentation requirements and operational requirements.

#### 4.2 Purpose:

For the washing of vials in washing area.



PRODUCTION DEPARTMENT

## USER REQUIREMENT SPECIFICATION

NAME OF ITEM: VIAL WASHING MACHINE	PROTOCOL No	
FUNCTIONAL AREA: PRODUCTION	Page No.: 5 of 12	

## 5.0 USER REQUIREMENTS

## 5.1 System Requirements:

S.No.	SYSTEM COMPONENTS	SYSTEM REQUIREMENTS	
1.	Identification (In case of Equipment	Details of Make, Name, Serial. No., Capacity, Model and Year of	
	/Instrument)	manufacture should be available	
2.	Model/Type	Vial washing machine with 20 track, feeding and loading	
		attachment confirming to current CGMP requirements.	
3.	Capacity	7.5 ml to 30 ml molded & tubular vials with Minimum	
		350 vials/minute	
4.	Potential Suppliers	1.Ambica	
5.	Contact parts (In case of Equipment)	SS316L with mirror finish	
6.	Non contact parts (In case of Equipment)	SS304 with mat finish	
0.	Non contact parts (in case of Equipment)	33304 With mat minish	
7.	Non metallic contact parts (In case of	• Any material with food grade quality having no potential impact	
	Equipment /Instrument)	on the products.	
		• Durable.	
		• Must be easily cleanable.	
8.	Motor & Electrical installations (In case of	Machine should be operated through PLC mounted on electrical	
	Equipment /Instrument)	control panel.	
9.	Machine assemblies (In case of Equipment /Instrument)	Must be covered with SS 304 with mirror finish.	
10.	Machine adjustments (In case of Equipment /Instrument)	Setting with Zero clearance with good accuracy.	
11.	Packaging & Transport	Should be packed and transported in such a way to avoid any	
		damage during transportation.	
12.	No. of requirements	01	
13.	Requirements for any power failure backup's (In case of Equipment /Instrument)	To be backed up by installed in-house DG set.	
14.	Gear box specifications(In case of Equipment /Instrument)	As per cGMP model	



PRODUCTION DEPARTMENT

## USER REQUIREMENT SPECIFICATION

NAME OF ITEM: VIAL WASHING MACHINE	PROTOCOL No
FUNCTIONAL AREA: PRODUCTION	Page No.: 6 of 12

## 5.2 Technical Description

S.No.	Specification	SYSTEM REQUIREMENTS
1.	Machine should be operated by manual and auto mode on PLC.	YES
2.	21 CFR part 11 compliant software	YES
3.	Machine should be PLC based control .	YES
4.	• Out feed lifter, pusher and out feed platform for vials provide in machine.	YES
5.	• Pump, pipe fittings (ss316with silicon pipe), pipe fitting for drain, tanks (ss316, minimum 50 ltrs).	YES
6.	Lifter device, Extra infeed conveyor, pre-inspection table , infeed conveyor, Infeed lifter, loosening device, should be provide with machine.	YES
7.	Washing sequence- 1 <sup>st</sup> wash Compressed Air 2 <sup>nd</sup> wash Re-circulated Water 3 <sup>rd</sup> wash Re-circulated Water 4 <sup>th</sup> wash Compressed Air 5 <sup>th</sup> wash Purified Water 6 <sup>th</sup> wash Compressed Air 7 <sup>th</sup> wash Purified Water 8 <sup>th</sup> wash Compressed Air 9 <sup>th</sup> wash WFI Water 10 <sup>th</sup> wash Compressed Air	YES



PRODUCTION DEPARTMENT

## USER REQUIREMENT SPECIFICATION

NAME OF ITEM: VIAL WASHING MACHINE	PROTOCOL No
FUNCTIONAL AREA: PRODUCTION	Page No.: 7 of 12

## 6.0 COMPLEMENTARY ASPECTS

## 6.1 Training

S.No.	SPECIFICATION	SYSTEM REQUIREMENTS
6.1.1	The vendor Shall supply all available information for the	YES
	adequate exploitation of equipment. For the Compliance of	
	this purpose at the Job site and/ or at the Vendors Shop.	
	Vendor's technical staff shall train customer's personnel. The	
	scope of the	
	Training will be agreed during the contract signature.	
6.1.2	The supplier is to include the personnel training activities.	YES
	The contractor is to specify the foreseen time for:	
	Operator/Supervisor training	
	Manager Training	
	Electrical maintenance training	
	Mechanical Maintenance training	
6.1.3	The contractor is to specify the personnel background needed	YES
	for each of the operators maintenance.	

## S.No. SPECIFICATION SYSTEM REQUIREMENTS 6.2.1 The System or its parts as provided for in the scope of supply shall be pre-installed at the vendors shop prior to delivery to customer site. Installation will be completed and documented including mechanical parts as well as electrical connections of all parts to facilitate taking over tests at Vendors shop prior to delivery. YES

### 6.3 Supplier Technical Documentation Requirements:

<ul> <li>Equipment/Systems electrical drawing</li> </ul>	Pre Installation Requirements will be supplied by Vendor
Equipment/Systems electrical drawing.	Vendor
Point to point wiring diagram	
<ul> <li>6.3.2 LIST.</li> <li>Equipment and instrument list with Component description.</li> <li>Electrical component parts list with Description.</li> <li>Function check list.</li> <li>Documentation list.</li> <li>Spare part list</li> </ul>	YES YES YES List of spares required for smooth operation will be provided by the Vendor at the time of ordering.



PRODUCTION DEPARTMENT

## **USER REQUIREMENT SPECIFICATION**

NAME OF ITEM: VIAL WASHING MACHINE	PROTOCOL No
FUNCTIONAL AREA: PRODUCTION	Page No.: 8 of 12

### 6.4 Technical Manuals

S.No.	Specification	Requirements
6.4.1	Operating handbook	YES
6.4.2	Trouble Shooting Guide	YES
6.4.3	Equipment Description	YES
6.4.4	Equipment specification	YES
6.4.5	Calibration Instruction	YES
6.4.6	Maintenance Instruction	YES
6.4.7	Maintenance Handbook	YES

### 7.0 SAFETY AND ENVIRONMENTAL PROTECTION

S.No.	Specification	Requirements
7.1	All motors have to be thermally Protected.	YES
7.2	All the Installation must be in accordance with the cGMP.	YES
7.3	The cGMP concerning safety must be applied.	YES
7.4	Safety interlock-	YES
	<ul> <li>Emergency stop button should be provided in the control panel.</li> <li>washing machine stop Maximum vial accumulation at tunnel infeed .</li> <li>washing machine start Minimum vial accumulation at tunnel infeed</li> <li>Main switch, in feed vial jam, lifter vial jam, pusher vial jam, discharge vial jam, provide in machine.</li> </ul>	

#### 8.0 CLEANING MAINTENANCE AND SERVICE

S.No.	Specification
8.1	In accordance with cGMP guidelines the units must be easy to clean, to disinfect, and where necessary, to sterilize.
8.2	The Supplier should guarantee that, if required, a service team can be on site within one working day.
8.3	The design should be such as to allow mechanical cleaning of the surface and that the cleanliness of the surface can be checked easily.
8.4	All machine parts, in particular instrumentation, should be constructed so that they can be easily removed and calibrated.
8.5	All special tools required for running and maintenance should be best.
8.6	A spare parts delivery guarantee with in time.

#### 9.0 RULES AND REGULATION

These standards, recommendation and requirements are considered the minimum. Specifications that are more stringent or expansive take the precedence. In case of conflict between published requirements, final determination is the responsibility of the Owners Representative.



PRODUCTION DEPARTMENT

## USER REQUIREMENT SPECIFICATION

NAME OF ITEM: VIAL WASHING MACHINE	PROTOCOL No
FUNCTIONAL AREA: PRODUCTION	<b>Page No.:</b> 9 of 12

### **10. SCOPE OF DELIVERY**

S.No.	Specification	Requirements
10.1	Units described in the specific system requirements including all necessary controls and instrumentation.	YES
10.2	The complete mechanical and electrical installation.	YES
10.3	The Connections to all the necessary utilities, exhaust, and waste lines necessary for its operation.	Yes
10.4	All piping and cabling of the units itself.	YES
10.5	Wiring and cable run: all wiring and cable run is part of the supply will supply the main power switches to be located in correspondence to the electrical and control cabinets delivered by the equipment supplier.	YES
10.6	All internal contacts of the supplied equipment for the required utilities.	YES
10.7	Unload on site of the equipment: the supplier is required to define all the necessary handling devices required to the unloading operation. The supplier will inform at least 4 weeks in advance the day of delivery and the list of required handling devices.	YES
10.8	Assembling operation: the required consumable, the internal transportation, the assembling tools and the required personal are part of the supply.	YES
10.9	A complete set of commissioning spare parts.	YES
10.10	All special tools necessary for use and maintenance of the supplied equipment.	YES
10.11	A complete set of two years spare parts should be listed quoted and offered as option.	YES
10.12	All test activities as specified in this document.	YES
10.13	Training in the use and maintenance of the equipment.	YES
10.14	A complete set of documentation as specified in this document.	YES



PRODUCTION DEPARTMENT

## USER REQUIREMENT SPECIFICATION

NAME OF ITEM: VIAL WASHING MACHINE	PROTOCOL No
FUNCTIONAL AREA: PRODUCTION	Page No.: 10 of 12

### 11.0 INSTALLATION, COMMISSIONING AND TESTS

### 11.1 General

Sr. No.	Specification	Requirements
11.1.1	The Contractor must specify for each piece of equipment the Guaranteed	YES
	performance and the guaranteed system performance. These values will be tested during the acceptance tests.	
11.1.2	In addition the functionality described in the user requirements and detailed in the system specifications will be tested.	YES

### **11.2 INSTALLATION, COMMISSION**

Sr. No.	Specification	Requirements
11.2.1	The commissioning tests will be carried out in accordance with a written test plan	YES
	developed by the supplier with clearly stated test procedures and acceptance	
	criteria.	
11.2.2	The contractor will approve successfully completed tests and will specify items	YES
	requiring additional work. Representatives fromWill attend and	
	participate in the commissioning tests as required.	
11.2.3	The installation and commissioning of the system will be performed at the	YES
	Facility by the contractor.	
11.2.4	The commissioning can only start once all the foreseen documents have been	YES
	delivered by the supplier to	
11.2.5	All equipment should be properly installed, adjusted, leveled, tagged, and	YES
	connected with utilities.	
11.2.6	Point to point checks on wiring and pneumatic should be performed.	YES
11.2.7	All instruments should be properly calibrated.	YES
11.2.8	A equipment (instrument) used for qualification must be listed and approved by	YES
	······	
11.2.9	The calibration equipment must have all the necessary documents to demonstrate	YES
	their maintenance & use.	
11.2.10	The last calibration of all this equipment must be less than 6 months old, and	YES
	evidenced by certificate.	
11.2.11	Verification that the interior surfaces of equipment are free of practices and dirt	YES
	and all points of product contact meet the specified material requirements.	
11.2.12	All the clearances and tolerances specified in the drawing or recommended by	YES
	component manufacturers are correct.	



PRODUCTION DEPARTMENT

## USER REQUIREMENT SPECIFICATION

NAME OF ITEM: VIAL WASHING MACHINE	PROTOCOL No
FUNCTIONAL AREA: PRODUCTION	Page No.: 11 of 12

Sr. No.	Specification	Requirements
11.2.13	On site verification that valves and other equipment with moving parts are in their	YES
	normal position if in a power down condition and move in the correct direction	
	with the correct speed and precision.	
11.2.14	Verification that all the Input and Output points are connected and labeled	YES
	according to the documentation and that all the along the input values have been	
	scaled in accordance with the system specification and process requirements. That	
	all equipment components requiring configuration	
11.2.15	The commissioning should demonstrate that the system supplied by the contractor	YES
	has been properly installed and that the functions are in accordance with	
	User Requirements specifications, Vendors System specifications	
	Manuals and other Documentation.	

### 11.3 Site Acceptance Test (SAT)

Sr. No.	Specification	Requirements
11.3.1	This test will be carried out once the commissioning will be completed. The scope	YES
	will be to verify the performance and the functionality of the system integrated	
	with the other factory systems (Including sterility testing of at least 02 days).	
11.3.2	The test will be carried out to verify the system response with the expected	YES
	productivity of the system.	
11.3.3	Details on the test realization will be defined during the project Phase. The supplier	YES
	is asked to specify the proposed duration for SAT and the standard procedure	
	proposed.	
11.3.4	During SAT the required functionality, performances and system reliability are	YES
	met.	
11.3.5	The Functionality described in the User Requirements Specification and in the	YES
	System Specifications are verified and met.	
113.6	All the documentation agreed has been delivered.	YES



PRODUCTION DEPARTMENT

## USER REQUIREMENT SPECIFICATION

NAME OF ITEM: VIAL WASHING MACHINE	PROTOCOL No
FUNCTIONAL AREA: PRODUCTION	Page No.: 12 of 12

### 12.0 QUALIFICATION/VALIDATION

Sr. No.	Specification	Requirements
12.1	The maintenance Qualification is responsibility of the customer. However, the	YES
	supplier is responsible for delivering the basic documents for maintenance	
	qualification.	
12.2	This includes all side costs such as : calibration measuring equipment and	YES
	instruments: manpower (IQ and OQ will take place completely on)	
12.3	Time Schedule for IQ/OQ execution will be developed by With the	YES
	supplier.	
12.4	Suppliers personnel used for IQ/OQ must be well trained and experienced. This	YES
	should be documented.	
12.5	The onsite test run performed by the supplier might become part of the IQ.	YES
12.6	Main IQ/OQ steps such as calibration must be performed and documented in	YES
	accordance to a SOP approved by	
12.7	All equipment used for qualification must be listed and approved by	YES
	The calibration equipment should be well documented.	
12.8	The last Recalibration of all this equipment should be less than 06 month old.	YES
	Proofed by Certificate.	
12.9	OQ can only start after IQ approved by	YES
12.10	IQ will be carried out by During Installation phase. IQ will include	YES
	the tests performed by the contractor.	
12.11	Part of the OQ will be carried out by During commissioning and	YES
	SAT phase. OQ will include the tests performed by the contractor.	
12.12	After installation of the equipment at customers site. Complementary IQ & OQ	YES
	tests will be performed by the Customer and may be supervised by a member of	
	Technical staff.	
12.13	Qualification documents (In case of equipments/Instruments)	DQ, IQ, OQ & PQ

#### **13.0 GAURANTEE/WARRANTEE**

Sr. No.	Specification	Requirements
13.1	The System must be guaranteed including all the sub- system and components for	YES
	a period of 12 months from the date of the system acceptance for a 03- shift	
	operation.	
13.2	The servicing companies involved for the Sub- systems maintenance must be	YES
	declared and the maintenance group organization described. Furthermore, the	
	contractor will be directly responsible of the system assistance and the required	
	operation will be co- ordinate by him.	
13.3	In case of failures, the intervention will be guaranteed by the contractor within a	YES
	maximum time limit. The contractor is asked to specify the maximum time limit.	
13.4	The supplier is asked to propose as option maintenance and assistance contract	YES
	after the guarantee expiration.	