

PROTOCOL No.:

# OPERATIONAL QUALIFICATION PROTOCOL CUM REPORT FOR STIRRER

| EQUIPMENT ID No.        |             |
|-------------------------|-------------|
| LOCATION                | LIQUID LINE |
| DATE OF QUALIFICATION   |             |
| SUPERSEDES PROTOCOL No. | NIL         |



PROTOCOL No.:

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| DESIGNATION          | NAME | SIGNATURE | DATE |
|----------------------|------|-----------|------|
| HEAD<br>(PRODUCTION) |      |           |      |



**STIRRER** 

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### **2.0 OBJECTIVE:**

- To verify that the equipment operates in accordance with the design and user requirements as defined by set Acceptance Criteria and complies with relevant cGMP Requirements.
- To verify the Operational features of Stirrer and to ensure that it produces desired Quality & rated output according to manufactures specifications.
- To verify all the Operational features from user point of view of the Equipment, Cleaning Procedure, Start up & Shut down Procedure and Safety Features

### 3.0 SCOPE:

- The scope of this operational qualification protocol cum report is limited to qualification of Stirrer installed in Liquid Line.
- The Stirrer is a standalone unit with plug in type electrical connections for operation and is on castor wheel. Hence, may be moved as per requirement to other area of operation which shall not change the performance of equipment.
- This Protocol will define the methods and documentation used to perform OQ activity the Stirrer for OQ. Successful completion of this Protocol will verify that Stirrer meet all acceptance criteria and ready for Routine Use.



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### 4.0 **RESPONSIBILITY:**

The Validation Group, comprising of a representative from each of the following departments, shall be responsible for the overall compliance of this Protocol cum Report:

| DEPARTMENTS       | NTS RESPONSIBILITIES   |  |  |
|-------------------|--|--|--|
| Quality Assurance | <ul> <li>Preparation, Review, Authorization and Compilation of the Operational Qualification Protocol cum Report.</li> <li>Co-ordination with Production and Engineering to carryout Operation Qualification.</li> <li>Monitoring of Operation Process.</li> <li>Post Approval of Operational Qualification Protocol cum Report after Execution</li> </ul>             |  |  |
| Production        | <ul> <li>Review &amp; Pre Approval of Operational Qualification Protocol cum Report.</li> <li>To Co-ordinate and support for execution of Operation Qualification study as per Protocol.</li> <li>Post Approval of Operational Qualification Protocol cum Report after Execution</li> </ul>  |  |  |
| Engineering       | <ul> <li>Review &amp; Pre Approval of Operational Qualification Protocol cum Report.</li> <li>Co-ordination, Execution and technical support in Stirrer Operational Qualification Activity.</li> <li>Responsible for Trouble Shooting (if occurs during execution).</li> <li>Post Approval of Operational Qualification Protocol cum Report after Execution</li> </ul> |  |  |



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### **5.0 EQUIPMENT DETAILS:**

| Equipment Name                  | Stirrer     |
|---------------------------------|-------------|
|                                 |             |
| Equipment ID.                   |             |
| Manufacturer's Name             |             |
| Model                           | GMP         |
| Supplier's Name                 |             |
| <b>Location of Installation</b> | Liquid Line |

### **6.0 EQUIPEMENT DESCRIPTION:**

Stirrer is suitable for emulsifying, dispersing, mixing and comminuting of liquid to Liquid products. It is based on rotor- stator principle. It is available in plain as well as water jacketed model which are suitable for heat sensitive products.

Special design facilitates adjustment of the grinding gap by an exterior screw by means of handle even during operation.

Stirring is an important step in pharmaceutical manufacturing process.

### **Operation:**

Product is fed to the operating area of a rotor, having a speed of 2800 RPM by specially designed feed device. The product is processed by high shear, pressure & friction between two Phase, and also, which exerts their force on it by means of pressing & releasing action.

### 7.0 PRE – QUALIFICATION REQUIREMENTS:

### **7.1** Verification of Documents:

- Executed and approved design qualification document
- Technical specification of equipment
- Certificate of material of construction of components.

### 7.1.1 Procedure:

- Verify the above mentioned documents for availability, completeness and approval status
- If any deviation is observed the same has to be recorded giving reasons for deviation and approved. Deviation should be approved by Authorized person.
- Approved Drawings and supporting documents would form a part of the IQ Protocol cum report.



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### 7.1.2 Acceptance Criteria:

• All the documents should be available, complete and approved by respective authorities.

### 8.0 CRITICAL VARIABLES TO BE MET:

### **8.1** Verification of documents:

The results of any tests should meet the limits and acceptance criteria specified in the test documents. Any deviations or issues should be rectified and documented prior to OQ commencing

| S.No. | Document Name                                      | Completed<br>(Yes/No) | Checked By<br>(Engineering)<br>Sign/Date | Verified By<br>QA<br>Sign / Date |
|-------|--|-----------------------|--|----------------------------------|
| 1.    | DQ Protocol Cum Report                             |                       |  |                                  |
| 2.    | IQ Protocol Cum Report                             |                       |  |                                  |
| 3.    | Draft SOP for operation & Cleaning of Stirrer      |                       |  |                                  |
| 4.    | Draft SOP for Preventive<br>Maintenance of Stirrer |                       |  |                                  |

| Checked By Production Sign/Date: | Verified By Quality Assurance Sign/Date: |
|----------------------------------|--|
| Inference:                       |  |
|                                  |  |
|                                  |  |
|                                  |  |
|                                  |  |
|                                  |  |
|                                  | Reviewed By                              |
|                                  | Manager QA                               |
|                                  | Sign/Date:                               |
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### **8.2** Test Equipment Calibration:

| Equipment /<br>Instruments<br>Name | Equipment /<br>Instrument I.D. | Calibration On | Due On | Observed By<br>Sign/Date |
|------------------------------------|--------------------------------|----------------|--------|--------------------------|
|                                    |                                |                |        |                          |
|                                    |                                |                |        |                          |
|                                    |                                |                |        |                          |

| Checked By Production Sign/Date: | Verified By Quality Assurance Sign/Date: |
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| Inference:                       |  |
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|                                  | Reviewed By Manager QA Sign/Date:        |



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### **8.3** Operational And Functional Checks:

Operate the Stirrer as per Manufacturer's Manual / SOP and Check for the following functions of the Equipment. The Equipment should function as desired.

| Item               | Acceptance criteria             | Observation | Observed By (Engineering) Sign/Date |
|--------------------|---------------------------------|-------------|-------------------------------------|
| Power              | Connect 3Ph, 415V, AC           |             |                                     |
| supply             | supply to the control panel     |             |                                     |
|                    | through proper isolator         |             |                                     |
| Motor &            | Check the direction of motor    |             |                                     |
| drive              | shows on machine by direct      |             |                                     |
|                    | arrow.                          |             |                                     |
| ON-OFF             | Green Button Operation Starts   |             |                                     |
| Operation          | & Red Operation Stops as        |             |                                     |
| <b>Push Button</b> | Required                        |             |                                     |
| Blades             | Check that blades should be     |             |                                     |
|                    | properly tightened.             |             |                                     |
|                    |                                 |             |                                     |
| Application        | Stirrer is Suitable for, Mixing |             |                                     |
|                    | Comminuting of Liquids to       |             |                                     |
|                    | Liquid.                         |             |                                     |

| Checked By (Production) | Verified By<br>(Quality Assurance) |
|-------------------------|------------------------------------|
| Sign/Date:              | Sign/Date:                         |
| Inference:              |                                    |
|                         |                                    |
|                         |                                    |
|                         |                                    |
|                         |                                    |
|                         |                                    |
|                         | Reviewed By                        |
|                         | (Manager QA)                       |
|                         | Sign/Date:                         |



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### 8.4 Safety Testing / Interlocking:

| Item              | Acceptance Criteria                    | Observation | Observed By<br>(Engineering)<br>Sign/Date |
|-------------------|--|-------------|---|
| To deactivate the | The machine should stop immediately    |             |   |
| equipment in      | and should not start when started till |             |   |
| event             | emergency stop switch is released      |             |   |
| Off an emergency  | The machine should be made to turn off |             |   |
| Stop              | during any emergency.                  |             |   |
| Noise Level       | Below 80 db                            |             |   |

| Checked By (Production) Sign/Date: | Verified By (Quality Assurance) Sign/Date: |
|------------------------------------|--|
| Inference:                         |  |
|                                    |  |
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|                                    | Reviewed By (Manager QA) Sign/Date:        |



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| 8.5 | Power | <b>Failure</b> | Verification  |
|-----|-------|----------------|---------------|
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| Item                | Acceptance Criteria  | Observation | Observed By<br>(Engineering)<br>Sign/Date |
|---------------------|--|-------------|---|
| Main Power Shut     | Equipment stops in a safe and                                      |             |   |
| Down                | secure condition.  |             |   |
| Main Power Restored | Equipment can be restarted with no problems or adverse conditions. |             |   |

| Checked By (Production) Sign/Date: | Verified By (Quality Assurance) Sign/Date: |
|------------------------------------|--|
| Inference:                         |  |
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|                                    | Reviewed By (Manager QA) Sign/Date:        |



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### 9.0 **REFERENCES**:

- Design Qualification Protocol cum Report
- Installation Qualification Protocol cum Report
- GA Drawing
- Operating manual
- Wiring Diagram

### **10.0 DOCUMENTS TO BE ATTACHED:**

- Operation And Maintenance Manual
- Copy Of Draft SOP's
- Any Other Relevant Documents

| 11.0 | DEVIATION FROM PRE-DEFINED SPECIFICATION, IF ANY:             |
|------|---|
|      |   |
|      |   |
|      |   |
| 12.0 | CHANGE CONTROL, IF ANY:                                       |
|      |   |
|      |   |
|      |   |
|      |   |
| 12.0 | DEVIEW (INCLUSIVE OF FOLLOWID ACTION IF ANY).                 |
| 13.0 | REVIEW (INCLUSIVE OF FOLLOW UP ACTION, IF ANY ):              |
| 13.0 | REVIEW (INCLUSIVE OF FOLLOW UP ACTION, IF ANY ):              |
| 13.0 | REVIEW (INCLUSIVE OF FOLLOW UP ACTION, IF ANY ):              |
|      | REVIEW (INCLUSIVE OF FOLLOW UP ACTION, IF ANY ):  CONCLUSION: |
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PROTOCOL No.:

STIRRER

| 15.0 | RECOMMENDATION: |
|------|-----------------|
|      |                 |
|      |                 |
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|      |                 |

### **16.0 ABBREVIATIONS:**

Amp. : Ampere

cGMP : Current Good Manufacturing Practices

HP : Horse power

ID. : Identification

IQ : Installation Qualification

KW : Kilo watt

MCB : Miniature circuit break

mm : Millimetre

MOC : Material of construction

NLT : Not less than

No. : Number

OQ : Operational Qualification

QA : Quality Assurance

SS : Stainless steel

WHO : World Health Organization



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### 17.0 POST- APPROVAL:

### PREPARED BY:

| DESIGNATION                           | NAME | SIGNATURE | DATE |
|---------------------------------------|------|-----------|------|
| OFFICER/EXECUTIVE (QUALITY ASSURANCE) |      |           |      |

### **REVIEWED BY:**

| DESIGNATION                              | NAME | SIGNATURE | DATE |
|--|------|-----------|------|
| OPERATING MANAGER<br>(QUALITY ASSURANCE) |      |           |      |
| HEAD<br>(ENGINEERING)                    |      |           |      |

### **APPROVED BY:**

| DESIGNATION          | NAME | SIGNATURE | DATE |
|----------------------|------|-----------|------|
| HEAD<br>(PRODUCTION) |      |           |      |