

MICROBIOLOGY DEPARTMENT

INSTALLATION QUALIFICATION PROTOCOL FOR VERTICAL LAMINAR FLOW UNIT

Pre - Execution Approval

	Name	Designation	Signature	Date
Prepared By				
Reviewed By				
Reviewed By				
Reviewed By				
Approved By				

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INSTALLATION QUALIFICATION PROTOCOL FOR VERTICAL LAMINAR FLOW UNIT

1.0 Objective:

The purpose of installation qualification is as follows

- To provide documented evidence that the mentioned Vertical Laminar Flow Unit is installed as per design.
- To ensure that the Horizontal Vertical Flow Unit installed confirms to purchase specifications and manufacturer literature, and to document the information that the Vertical Laminar Flow Unit meets the specification.

2.0 Scope:

Scope is limited to the following

Equipment / System Name	Vertical Laminar Flow Unit	
ID Number		
Location	Cooling Zone	

3.0 Equipment / System Description:

Klenz FloTM Vertical Laminar flow clean air workstation provides a contained environment to protect the product. It provide clean zone and particle free conditions by unidirectional down ward airflow from the HEPA filters situated above the working area. This shall produce clean zone with air quality, which is equivalent or better than class 5 ISO 144644 - 1 / specifications.

The Vertical Laminar flow clean air workstation is required to provide local class 5 (ISO 144644 - 1) grade environment for aseptic processing. The equipment should be designed for continuous run and shall not experience any major breakdown because of workmanship. The equipment is intended to be in operation in all the three shift 7 days a week.

The Vertical Laminar flow clean air workstation shall be used for maintaining class 5 (ISO 14644 - 1) environment during unloading of sterilized equipments, accessories, materials, etc.



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4.0 Checklist for Preinstallation verification:

The purpose of the checklist is to confirm the availability of required documents for installation and to verify the availability of components and parts as per the approved purchase order in presence of the technical personnel of the vendor.

Preinstallation verification checklist is enclosed as Annexure - I.

5.0 Checklist for Installation verification:

Installation of Vertical Laminar Flow Unit shall be verified for the compliance with the critical parameters mentioned in the Functional Design Specification.

The purpose of this checklist is to check and document the received material and installation at site with respect to drawings, details, and data sheets approved during DQ and GEP. GEP considers the basic engineering requirements for skilful maneuvering, means those methods and practices, which the administrator determines to be consistent with scientific and engineering principles.

Installation verification checklist is enclosed as Annexure - II.

6.0 Any Changes identified towards equipment design / lay out.

Refer Annexure - III.

7.0 Recommendations and Conclusions:

8.0 References:

Purchase order (already available with Design Qualification)

Packing list supplied by vendor (Not applicable).

List of spares (Not applicable).

Installation Qualification submitted by vendor.

Impact Assessment analysis.

9.0 Annexure

Annexure - I : Check list for Preinstallation Verification.

Annexure - II : Check list for Installation Verification.

Annexure - III : List of Changes / Deviation.

Annexure - IV: Installation Qualification Submitted by the vendor.

Annexure - V : Impact Assessment Analysis.

Annexure - VI: Summary Report of Installation Qualification



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10.0 Abbreviations:

IQ : Installation Qualification

DQ : Design Qualification

Post execution approval:

	Name	Designation	Signature	Date
Compiled By				
Reviewed By				
Reviewed By				
Reviewed By				
Approved By				



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Annexure - I

Checklist for Preinstallation Verification

S.No.	Main Components Accessories / Documents	Code / Doc No.	Actual	Remarks
1.	Purchase Order No.			
2.	Vendor's Name	Klenzaids Bioclean		
3.	Instrument Make	Klenzaids		
4.	Instrument Model No.	Klenz Flo TM		
5.	Design qualification Reference	Design qualification submitted by the vendor		
6.	Factory Acceptance Test (FAT) Certificate	Should be provided		
7.	Instrument Manual	Instrument Manual submitted by the vendor		



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Annexure - II

Checklist for Installation Verification

S.No.	System Data	Acceptance Criteria	Actual	Remarks
Α.		Equipment /Instrument specific details		
1.	Material of Construction	Should be SS 304		
2.	Curtain	Should be Antistatic PVC curtain		
3.	HEPA filter Screen (MOC)	Should be SS 304		
4.	HEPA filters	Should be EU - 13, 610 x 915 x 75 mm (Antimicrobial)		
		Should be EU - 6, 765x298x45 mm (Biocidal)		
5.	Pre filters	Should be EU - 6, 728 x 298 x 45 mm (Biocidal)		
		Should be EU - 6, 840 x 298 x 45 mm (Biocidal)		
6.	DOP Port	100 % DOP test port Should be provided		



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S.No.	System Data	Acceptance Criteria	Actual	Remarks
Α.	Equipment /Instrument specific details			
7.	Electrical construction	a) Fluorescent light:4'L; 40-Wattsb) Independent controls for lighting, blowers& Sockets.		
8.	Relay / Switchgear	Should be Provided		
9.	Indicators	Power ONBlower ONLight switchShould be provided		
10.	Differential pressure gauge	Should be 0 to 25 mm WC		
11.	Special features	Soft touch key pad switch Should be provided		
12.	Motor	Should be 0.5 HP; 1440 RPM;		
13	Blower (Impeller)	Should be 210 x 85 mm		
14.	Overall Dimensions (W x D x H)	Should be 1065 x 2746 x 610 mm		



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S.No.	System Data	Acceptance Criteria	Actual	Remarks
В.	Location suitability			
1.	Location	Should be place Cooling Zone		
C.		Utiliti	es	
1.	Electrical power supply	Should be 230 V; AC; 50 Hz		
D.		Safet	y	
1.	Alarm	Audiovisual alarm for motor blower trip Should be provided		
Е.	MOC Certificates	Should be provided		
F.	Calibration Certificates	Should be provided		
G.	Testing Certificates	Should be provided		
Н.	Drawing Details	Drawing No.		
1.	Drawing of Vertical Laminar Air Flow Unit			



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Annexure - III

List of Changes / Deviations

S.No.	Description of Change / Deviations	Justification based on impact analysis

V	erifi	ied	By:
			•

Approved By:



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Annexure - VI

Summary Report of Installation Qualification

Checks	Observations (Yes / No)	Reviewed By Sign / Date
All test procedures executed and verified as per the protocol.		
All criteria set forth in the installation qualification were met.		
Deviation if any		
Summary:		

Summary Report Prepared By:

Date & Sign