1 011011114					
Document No	Controlled Copy No.	Effective Date	Page 1 of 17		
			<b>Review: N/A</b>		
Prepared by	Date	Authorised By	Date		

Performance Qualification Protocol (PQP) for **Steam/Air Cycle** in the Production Steam Steriliser (Autoclave), located in .....

# PREPARED BY:

Validation Engineer	Date
PROTOCOL APPROVAL: This protocol has been approved by:	
Microbiologist	Date
Operations Manager	Date
QA Manager	Date

Document No	Controlled Copy No.	Effective Date	Page 2 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

# 1 Scope and Protocol Objective

The Production Steam Steriliser (autoclave), located in ...... shall be used for sterilising aseptically-filled vials of selected products.

To qualify the performance of the Fedegari Steam Steriliser (Autoclave) as part of a change control qualification study (refer to CR-14- xxxx "Recommence Manufacture of xxxx Acid Injection 15 mg in 1 mL) This Performance Qualification shall be limited to demonstrating consistency and efficacy of the steam/air sterilisation cycle, using 1mL water filled into 2 mL vials. Equivalence for xxx Injection 15 mg in 1 mL product, which has been aseptically filled into 2 mL vials, shall be demonstrated in the subsequent Process Validation Study (see PQ protocol kkk).

This protocol has been prepared with reference to the following regulatory guidelines:

The Performance Qualification study (PQP kkk ) for the autoclave equipment, included heat distribution studies for a porous load cycle only. This document shall include heat distribution studies for the steam/air sterilisation cycle, as part of the process development for the terminal sterilisation of filled vials.

The objective of this Performance Qualification, is to verify that the sterilising autoclave consistently provides the required sterility assurance, when operated under normal conditions, using standard minimum and maximum loading patterns and the specified settings:

The production cycle registered for Folic Acid product is 121.1  $^{\circ}$ C for fifteen (15) minutes, to provide a minimum F<sub>o</sub> = 8 min.

The standard loading patterns shall be as follows:

Minimum load Six (6) trays 2 mL vials, 340 vials per tray, across two autoclave shelves Maximum Load Nine (9) trays 2 mL vials, 340 vials per tray, across three autoclave shelves

A reduced cycle shall also be run, for the standard production load patterns, to demonstrate that sub-optimal conditions also yield an acceptable level of sterility assurance. This shall be achieved by changing the time-temperature combination for the standard production load patterns to 121.1 °C, for *ten* (10) minutes, which is 66% of the registered sterilising condition of 121.1°C for 15 minutes.

#### 2 Equipment Identification and Description

The Production Steam Steriliser is a multi-cycle, 210 L capacity, single-ended (*future:- double ended*) autoclave, manufactured by Fedegari, model FOB4S-TS (S/N aaaa). The installation and operation is described in the following supporting documents:

ED ED Commissioning – F3 Production Steam Steriliser (Fedegari FOB4S –TS Autoclave) IQP for F3 Steam Steriliser (Autoclave) IQR for F3 Steam Steriliser (Autoclave) OQP "OQP for F3 Steam Steriliser (Autoclave) OQR for Steam Steriliser (Autoclave) PQP for F3 Steam Steriliser (Autoclave) PQR for F3 Steam Steriliser (Autoclave) PVP Terminal Sterilisation of xxxx Product

#### 2.1 Definition of Terms - Engineering Specifications

The following terms have been used in this protocol and are explained below:

• SAL: Sterility Assurance Level

Document No	Controlled Copy No.	Effective Date	Page 3 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

- **PNSU:** Probability of a non-sterile unit
- D<sub>T</sub>: The time in minutes required for a one logarithm or 90% reduction of the population of microorganisms used a biological indicator under specified lethal conditions at a reference temperature T.
- F value: measure of the microbiological inactivation capability of a heat sterilisation process
- F<sub>o</sub> value: F value calculated using a reference temperature (T<sub>ref</sub>) of 121.1°C, with a z value of 10°C and a D value of 1 minute
- **F**<sub>phys</sub>: A term used to describe the delivered lethality, which is calculated using the actual physical parameters of the cycle against the reference temperature (T<sub>ref</sub>).
- Z value: The number of degrees of temperature change necessary to change the D-value by a factor of 10

## 2.2 Responsibilities

Validation Staff shall prepare the Performance Qualification Protocol (PQP) document and train the Production Staff in its execution.

The Performance Qualification Protocol shall be approved prior to execution by:

- The Microbiologist
- The Operations Manager
- The Quality Assurance Manager

The Production Supervisor shall be responsible for conducting the test protocols. Validation Staff shall assemble data, review results and draw conclusions from the test protocols in order to prepare the Performance Qualification Report (PQR).

The PQ report shall be reviewed by:

- The Microbiologist
- The Operations Manager
- The Quality Assurance Manager

#### 2.3 Other Relevant Documents

#### 3 **Process Description**

Sterilisation shall be by the moist heat process, using saturated steam, where  $F_0 > 8 D_T$  (see below). A steam-air mixture is used to control chamber pressure and assist in pressure equalisation between chamber and vials, particularly during the cool-down phase.

In-process controls for the sterilisation phase of the cycle shall be temperature TE1 in the liquid product and sterilisation phase hold time. Temperature (TE 8 on the auxilliary heating device) and chamber pressure (TP01) are also required to control heating and forced cooling phases of the cycle.

For the purposes of the PQ, sterilisation phase temperature and hold time data shall be processed to demonstrate that the physical characteristics of the cycle in terms of accumulated lethality ( $F_{phys}$ ) exceed the minimum cycle design criteria ( $F_o$ ), where:

 $F_o > 8 D_T$  for  $D_{Tref} = D_{121.1oC} = 1.0$  minutes and  $F_{phys} = \Delta t \sum 10^{(T-Tref)/z}$ 

Reference: BP Appendix XVIII "Methods of Sterilisation" and registered particulars, where  $F_0 > 8$  min is required.

Biological Indicators shall be selected, to demonstrate the survival probability of a non-sterile unit (PNSU)  $\leq 10^{-6}$  for both the normal production cycle ( $F_0 \geq 15$ ) and a reduced cycle ( $F_0 \geq 10$ ).

The selected cycle for normal operation is "P4 Steam Air Cycle", which has the following characteristics:

Heating phase to 121.1 °C

Document No	Controlled Copy No.	Effective Date	Page 4 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

- Sterilisation temperature 121.1 °C
- Sterilisation time 15 minutes
- Drying phase 10 minutes
- Forced cooling 30 minutes

The selected cycle for reduced cycle operation is a modified "P4 Steam Air Cycle", which has the following characteristics:

- Heating phase to 121.1 °C
- Sterilisation temperature 121.1 °C
- Sterilisation time 10 minutes
- Forced cooling 15 minutes

Refer to Appendices A and B for standard load components and loading patterns respectively. A standard load configuration shall be the combination of one set of load components, stacked in the specified loading pattern.

#### 4 Performance Qualification Tests

Tests to be conducted and acceptance criteria are defined in the attached Performance Qualification Test Sheets.

#### Tests to be performed:

- 1 Test Instrument Calibration
- 2 Vacuum Leak Rate Test
- **3** Heat Distribution Test (empty chamber temperature mapping)
- 4 Heat Distribution Test (loaded chamber temperature mapping)
- 5 Heat Penetration studies for:
  - Production cycle standard loads (121.1 °C for 15 minutes, two consecutive cycles) and for "Reduced" cycle standard loads (121.1 °C for 10 minutes, one cycle)
- 6 Biological challenge testing for standard and reduced cycle loads

All attachments of raw data in the form of thermal printer records, shall include photo or printed copies.

#### **Tests 4-6 Summary Table:**

Load Description	Empty Chamber	Water Heat Distribution		Water Heat Penetration	
		(probes outside vials, no Bls)		(probes inside	vials, with Bls)
		Minimum load	Maximum load	Minimum load	Maximum load
		(6 trays)	(9 trays)	(6 trays)	(9 trays)
Production Cycle	3	2	2	2	2
(121.1°C, 15 min)					
Reduced Cycle	0	0	0	1	1
(121.1°C, 10 min)					
Biological Indicators	No	No	No	Yes	Yes

Acceptance criteria for the overall program shall be based on the following:

- 1. Successful calibration data for all measuring instruments
- 2. Successful completion of all test functions listed in Section 4.

#### **Protocol Report Reference:**

The results of all test studies shall be included in the Performance Qualification Report (PQR): .... "PQR for Steam/Air Cycle"

Document No	Controlled Copy No.	Effective Date	Page 5 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

The PQR shall include a summary report, with a list of all attachments to the protocol. Deviations from the Performance Qualification Protocol shall be listed in the report and filed electronically on copies of Form "Validation Deficiency Report" (VDR).

#### 4.1 Rationale for Testing Schedule

Two consecutive standard production cycles shall be tested for each load configuration to demonstrate consistency of autoclave performance. The first cycle shall be run under "cold start" conditions, where it is the first heated run of the day (note Vacuum Leak Rate Testing may be run first, because it does not require the chamber to be heated).

One "reduced" cycle shall be tested for each load configuration, to demonstrate lethality >  $10^{-10}$  under sub-optimal conditions. A reduced cycle of 121.1°C for 10 minutes has been selected, as this temperature-time combination delivers approximately half the lethality of the production cycle in terms of F<sub>o</sub> design value:

121.1 °C for 15 minutes  $F_o$  = 15 minutes 121.1 °C for 10 minutes  $F_o$  = 10 minutes

In order to preserve raw data, the PQ report shall include copies of all raw test data, which has been recorded on thermal paper. All qualification test cycles shall be logged in the summary report.

As a minimum, biological indicators shall be placed in the first, "cold start" heat penetration standard load cycle and in the "reduced" qualification cycle. Biological indicators shall be presented in a liquid form, inside sealed glass ampoules, which are of a similar size to the filled product vials.

#### The Biological indicator shall have:

Population > 5 x  $10^5$  Geobacillus stearothermophilus (ATCC 7953) per ampoule, with a "D" value of not less than 1.5 minutes and not more than 2.0 minutes.

A Vacuum Leak Rate Test shall be conducted prior to performance testing, with the temperature probes in place and thereafter on a daily basis, to confirm that the leakage rate remains within specification and that the autoclave chamber constitutes a sealed unit, during operation.

Document No	Controlled Copy No.	Effective Date	Page 6 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

Test 1	st 1 Title of Test: Test Instrument Calibration		
<b>Objective</b> To Verify that the F3 oven successfully calibrated.	and autoclave data recorder and temperature probe	s used for the qualification are	
	d to 11 temperature probes, shall be temporarily ion study, to provide and record data for thermal n studies.	<b>Reference:</b> TR vvvv "Extended Calibration Schedule"	
Sampling and Recording P	lan:	· ·	
	a recorder is calibrated at a minimum frequency of one udy or within one month of execution of this protocol.	e per year (as recorded in TR	
Test Equipment and Materi	als		
Dry heat block and other test (NATA) standards.	equipment supplied by calibration service contractor	referenced to recognized	
Acceptance Criteria The temperature monitoring	device shall be:		
	ver the scale range 50 °C to 150 °C C at the sterilisation temperature (121.1 °C)		
The temperature monitoring or within one month of the qu	equipment shall meet the above calibration requirement alification activity.	ents before and after PQ study,	
Test Results: Copies of calibration certifica	tion for protocol test equipment shall be included in the	Date:	
Summary of results (analys	· · · ·		
Acceptance Criteria met: F	Pass/Fail		
Analysed By: Validation S	taff		
Reviewed By: Microbiolog	ist		
Approved By: Operations	Manager		

Document No	Controlled Copy No.	Effective Date	Page 7 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

Test 2	Title of Test: Vacuum Leak Rate Test	
<b>Objective</b> To verify integrity of the cham	ber and door seals via pressure hold test under vacuum.	
port cover and insert tempera	ak Rate Test cycle. a "Validation Port" in the autoclave chamber. Remove the ture probes via the stainless steel adapter provided. Festing after mounting and on removal of test equipment.	Reference: SOP xxxx
-	ng days of operation, and also including: ng qualification probes lification runs	
over a 16 minute pressure ho where $P_1$ = approximately 15 kPa (al therefore where absolute press $P_2$ = equilibration pressure with	m. The controller measures and calculates the leak rate para	
Acceptance Criteria The rate of pressure change s	shall not exceed 0.13 kPa/min over a 16 minute pressure hole $(P_3-P_2) \le 2.08 \text{ kPa}$	l phase, i.e.
Test Results:		Date:
Attach copies of autoclave pri	nter records to the PQR.	
Summary of results (analys	is and conclusion)	
Acceptance Criteria met: P	ass/Fail	
Analysed By: Validation Sta	aff	
Reviewed By: Microbiologi	st	
Approved By: Operations M	Manager	
Approved By: QA Manager		

Document No	Controlled Copy No.	Effective Date	Page 8 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

Test 3	Title of Test: Heat Distribution Study (Empty Chamber Mapping)

#### Objective

Physical qualification of the Production sterilisation cycle to demonstrate that heat distribution within the empty chamber is consistent. Program 4 (Steam/air) shall be used as the standard sterilisation cycle.

#### Test Method

Fit the 11 thermocouple probes through the validation port in the autoclave door as described in Test Sheet 2 and check connections to the oven and autoclave data recorder. Close the door and perform a leak rate test cycle, as for Test Sheet 2.

Probe placement inside the chamber shall reflect the results from previous heat distribution studies (see Operational Qualification) for determination of hot and cold spots within the chamber. Record the position of each probe on a loading pattern diagram (see Appendix B).

The F3 oven and autoclave data recorder drain probe (AC1-B) shall be used to reference the temperature in the drain.

Select and run "Program 4" (steam/air cycle) whilst synchronising the F3 oven and autoclave data recorder start and finish times to collect all relevant data at 1 minute intervals. At least one cycle must be the first hot cycle run in the autoclave for that day ("cold start").

Copy, sign and date, all thermal printer records and record the test method and results on TR 019H068. Process the raw data according to the procedure outlined in Appendix C.

#### Sampling and Recording Plan

Three runs for this first, heat distribution study (empty chamber mapping) on the standard Production cycle (Program 4).

Test Equipment, Materials and Calibrations	References
Refer to OI 003171 'Use of oven/autoclave data recorder"	OI xxxxx OI xxxxx

Document No	Controlled Copy No.	Effective Date	Page 9 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

Test 3	Title of Test: Heat Distribution Study (Empty Chamber Ma	apping)		
Acceptance Criteria: 1. Verify consistency of o • Heating phase to 1 • Sterilisation tempe • Sterilisation time 1 • Drying phase 10 m • Forced cooling 15	erature 121.1 °C 5 minutes ninutes			
2. Record the overall time	e for cycle and verify that heating time is consistent.			
3. Leak rate tests remain	within specification.			
<ol> <li>Stratification within the other.</li> </ol>	chamber is acceptable i.e. thermocouple probes measure within +/- 2	2ºC of each		
5. All measured thermoco during the sterilisation	ouple temperatures show 121.1°C +3/-0°C and do not fluctuate by mo hold phase.	ore than 1 °C		
6. At least 10 thermocoup	ple probes remain active during the study			
	mperature and pressure readings, during the sterilisation hold time, ag s and record the results.	gainst		
8. For information, calcula	8. For information, calculate and record $F_{phys}$ for each thermocouple probe, using: $F_{phys} = \Delta t \sum 10^{(To-Tref)/z}$			
Where: $\Delta t = 1 \text{ minute}$ $T_o = \text{thermocouple te}$ $T_{ref} = 121.1 \text{ K (°C)}$ Z = value specific to	emperature K (°C) biological indicator used or 10 °C if not specified			
	at loads for each probe, as indicated by $F_0$ , are consistent.			
Test Results: Attach copies of marked-up proven and autoclave charts and	robe location (loading diagrams), printer records, TR 019H068, F3 I spreadsheet data.	Date:		
Summary of results: (analysi	is and conclusion)			
Acceptance Criteria Met: Pas	ss /Fail			
Analysed By: Validation Sta	ff			
Reviewed By: Microbiologis	Reviewed By: Microbiologist			
	Approved by: Operations Manager			
Approved by: QA Manager				

Document No	Controlled Copy No.	Effective Date	Page 10 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

Test 4	Title of Test:	Heat Distribution Study	(Loaded Chamber Mapping)	
<b>Objective</b> Physical qualification of the Production sterilisation cycle to demonstrate that heat distribution within the loaded chamber is consistent and to locate the "cold spot" within the chamber.				
	<b>Test Method</b> Load 6 trays of vials into the autoclave chamber, distributed across two shelves. Temperature probes shall be placed adjacent to vials and held in place with autoclave tape if necessary.			
			ous heat distribution study (Test 3) for n of each probe on a loading pattern	
Seal the liquid load probe TE1 spot" (Test 3). Similarly, for a			ation previously identified as the "cold	
The F3 oven and autoclave da drain.	The F3 oven and autoclave data recorder drain probe (AC1-B) shall be used to reference the temperature in the drain.			
Select and run "Program 4", whilst synchronising the F3 oven and autoclave data recorder start and finish times to collect all relevant data at 1 minute intervals. The cycle must be the first hot cycle run in the autoclave for that day ("cold start").				
Repeat the above cycle, leaving the coldest probes in place and rearranging the "hottest" probes, so that they are re-located, to new, previously unmapped and therefore potentially cold, locations.				
Repeat for 9 trays of vials.				
Copy, sign and date, all thermal printer records and record the test method and results on TR 019H068. Process the raw data according to the procedure outlined in Appendix C. <b>Sampling and Recording Plan</b>				
Two runs each for minimum and maximum loads, (Program 4).				
Test Equipment, Materials ar	nd Calibrations		References OI 003121.1	
Refer to OI 003171 'Use of ove	n/autoclave data	a recorder"	OI 003171	

Document No	Controlled Copy No.	Effective Date	Page 11 of 17	
			<b>Review: N/A</b>	
Prepared by	Date	Authorised By	Date	

Test 4	Title of Test: Heat Distribution Study (Loaded Chamber N	lapping)
<ul> <li>Heating phase to</li> </ul>	perature 121.1 °C 15 minutes minutes	
	me for cycle and verify that heating time is consistent. Use the data to a up time during the heat penetration study.	set upper and
3. Leak rate tests rema	in within specification.	
4. Stratification within th other.	ne chamber is acceptable i.e. thermocouple probes measure within +/-	2°C of each
<ol><li>All measured thermo during the sterilisatio</li></ol>	couple temperatures show 121.1°C +3/-0°C and do not fluctuate by mo n hold phase.	ore than 1 °C
6. At least 10 thermoco	uple probes remain active during the study	
	temperature and pressure readings, during the sterilisation hold time, a es and record the results.	gainst
Where: $\Delta t = 1 \text{ minute}$ $T_o = \text{thermocouple}$ $T_{ref} = 121.1 \text{ K} (^{\circ}\text{C})$ Z = value specific	ulate and record $F_{phys}$ for each thermocouple probe, using: $F_{phys} = \Delta t \sum 10^{(To-Tref)/z}$ temperature K (°C) to biological indicator used or 10 °C if not specified mulated heat loads for each probe, as indicated by F <sub>0</sub> , are consistent.	
<ol> <li>Verify that the "cold-s loads.</li> </ol>	spot" location is constant, in two consecutive cycles, for minimum and r	naximum
Test Results: Attach copies of marked-up oven and autoclave charts ar	probe location (loading diagrams), printer records, TR 019H068, F3 nd spreadsheet data.	Date:
Summary of results: (analy	rsis and conclusion)	
Acceptance Criteria Met: P	ass /Fail	
Analysed By: Validation S		
Reviewed By: Microbiolog		
Approved by: Operations N	-	
Approved by: QA Manager		

Document No	Controlled Copy No.	Effective Date	Page 12 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

Test 5	Title of Test: Heat Penetration Study	
<b>Objective</b> Physical qualification of the effectively for the standard lo temporarily modified for the "r	e sterilisation cycle to demonstrate that heat penetrates o oad configurations set out in this protocol. Program 4 (Stea reduced" qualification cycle.	equipment load items am/Air Cycle) shall be
condensate can drain freely fr Seal temperature probes ins liquid, without touching the si in Appendix D. Allow 24 hour Position 11 temperature prob to reflect the results from pre	terilising chamber, so that steam and air can flow around and rom each item. side the product vials so that the probe is immersed in the des. Refer to the photograph and sealing method described rs for the sealant to cure before proceeding. es, sealed inside product vials, within the autoclave chamber evious heat distribution studies for determination of hot and	Reference AS ISO 11134-2003 7.4.2 And SOP 003112
cold spots within the chamber Seal the liquid load probe T previously identified as the "c and TE5.		
The F3 oven and autoclave of the temperature in the drain.		
Place a Biological Indicator "Biological challenge testing f		
Record the position of each diagram (see Appendix B).		
For the first two runs, select a the data recorder start and f Select and run a reduced cyc for <b>10</b> minutes.		
All validation cycles, for each Test 6.		
The first validation cycle run autoclave for that day, to ensu		

Document No	Controlled Copy No.	Effective Date	Page 13 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

Test 5	Title of Test: Heat Penetration Study		
<b>Test Method</b> <i>(continued)</i> Open the door and prepare for the next test cycle by removing the used Biological Indicator adjacent to each temperature probe (See test schedule "Biological challenge testing for standard loads"). On completion of the third ("reduced") test cycle, remove the load and repeat the vacuum leak rate test cycle as for test schedule 2.			
Re-instate production cycl	e conditions to the Steam//	Air cycle.	
Repeat for 9 trays of vials	, laid across 3 autoclave sh	ielves.	
Copy all thermal printer records and record the test method and results on TR 019H068. Graphical presentation of data is by using secure software as described in OI 003173 "Operator Instructions for Data Review – F3 oven and autoclave data recorder". Use the "Review" software to download the raw data and process it on electronic spreadsheets, according to the procedure outlined in Appendix C.			ctions for Data Review – F3 oven
Sampling and Recording			
	ors (see Test 6). Record Penetration Results Shee		R 019H068 "F3 Production Steam
6 trays (minimum load)	2 x Production cycle (at least 1 "cold start")	1 x "reduced" cycle ("cold start")	1 x Vacuum Leak Rate Test
9 trays (maximum load)	2 x Production cycle (at least 1 "cold start")	1 x "reduced" cycle ("cold start")	1 x Vacuum Leak Rate Test
Test Equipment and Materials			
			and Operation of Temperature PQ report as described in Test

#### Acceptance Criteria:

For minimum and maximum load patterns:

1. Verify consistency of operation of sterilisation production cycle parameters:

- Heating phase to 121.1 °C
- Sterilisation temperature 121.1 °C
- Sterilisation time 15 minutes
- Drying phase 10 minutes
- Forced cooling 30 minutes

Record the overall time for the cycle and verify that the load conditioning phase (heating), prior to sterilisation, operates consistently.

Similarly, verify reduced cycle parameters:

- Heating phase to 121.1 °C
- Sterilisation temperature 121.1 °C
- Sterilisation time 10 minutes
- Drying phase 10 minutes
- Forced cooling 30 minutes
- 2. Vacuum Leak Rate Tests remain within specification.
- 3. Stratification within the chamber is acceptable i.e. temperature probes measure within +/- 2 °C of each other.

Document No	Controlled Copy No.	Effective Date	Page 14 of 17 Review: N/A
Prepared by	Date	Authorised By	Date
Test 5	Title of Test: Heat Pen	atration Study	
	Title of Test: Heat Pen	-	
<ol> <li>For all cycles, all measur more than 1 °C during th</li> </ol>	ed temperature probe tempe e sterilisation hold phase.	ratures show 121.1 °C +3/-0	°C and do not fluctuate by
5. At least 10 thermocouple	es stay active during the study	1.	
6. Chemical indicators show	v a positive indication for thre	shold sterilisation temperatur	e.
7. Correlate autoclave tem steam tables and record	• •	gs, during the sterilisation ho	ld time, against saturated
8. Calculate and record $F_{phy}$		e on at least one cycle per loa \t ∑10(T <sub>o</sub> -T <sub>ref</sub> )/z	d configuration, using:
Where:	- phys –		
$\Delta t = 1 \text{ minute}$	robe temperature K (°C)		
$T_o = temperature p$ $T_{ref} = 121.1 \text{ K} (^{\circ}\text{C})$	Tobe temperature K ( C)		
z = value specific	to biological indicator used or	10 °C if not specified	
Test Results:			Date:
Attach copies of loading diagrams, TR 019H068, raw data including printer records, F3 oven and autoclave data recorder charts, spreadsheets with formula checks to the PQR.			
Summary of results (analys	sis and conclusion)		
Acceptance Criteria met: F	Pass/Fail		
Analysed By: Validation St	taff		
Reviewed By: Microbiolog	ist		
Approved By: Operations	Manager		

Document No	Controlled Copy No.	Effective Date	Page 15 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

Test 6	Title of Test: Biological challenge testing for standard loads			
	sterilisation cycle to demonstrate that the de Biological Indicator system results in a PNSU			
<b>Test Method</b> This test is run concurrently v Indicators placed adjacently i	vith Test 5 (i.e. temperature probes and Biolo n the same loads).	gical <b>Reference</b> AS ISO 11134-2003 7.4.2 d)		
Guidelines for handling Bio	logical Indicators (BIs):			
<ul> <li>Collect the required numl to the facility.</li> </ul>	per of BIs from the Microbiology Laboratory.	Transfer the ampoules in a sealed bag		
ampoule is damaged, the Production, so that the ar 10% H <sub>2</sub> O <sub>2</sub> in 70% ethano	es carefully. Keep spare gloves and sample on transfer immediately to the sample jar and rea can be cleaned down accordingly, using a I solution and leave to soak for 60 minutes. C Microbiology Laboratory for disposal.	seal. Change gloves and notify sporicidal sanitant; e.g. spray with		
<ul> <li>Treat all autoclaved BIs a transfer back to the Micro</li> </ul>	is potentially viable and handle as described a biology Laboratory.	above. Place in a sealed bag for		
	Bls issued and the number returned from/to th ty). Inform the Production Manager on duty i			
For each standard load con	figuration:			
Place a Biological Indicator (I study").	BI) adjacent to each temperature probe locat	ion (see test schedule "heat penetration		
Record the position and tag of	of each BI.			
Complete the sterilisation cyc	le (See test schedule "heat penetration study	")		
penetration study"). Open th	study after the first sterilisation cycle has ne door and prepare for the next test cycle i mperature probe. Transfer the used BIs to	by removing the used BI and placing a		
Sampling and Recording P	an:			
6 trays (minimum load)	2 x Production cycle (at least 1 "cold start")	1 x "reduced" cycle ("cold start")		
9 trays (maximum load)	2 x Production cycle (at least 1 "cold start")	1 x "reduced" cycle ("cold start")		

Document No	Controlled Copy No.	Effective Date	Page 16 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

$F_{BIO} = D_{121.1} (Log N_0 - Log N_F)$ from heat penetration study)				
	Calculate the actual log reduction (N <sub>F</sub> = actual PNSU) of micro-organisms using: $F_{BIO} = D_{121.1} (Log N_0 - Log N_F)$ where: $F_{BIO} = F_{phys}$ (as found from heat penetration study) $D_{121.1}$ and N <sub>0</sub> (from BI data)			
Test Equipment and Materials         See TR gggg "RMTR – M.org – Biological Indicator Ampoules (RM 6119)         QC shall quality-test and incubate a non-sterilised spore strip as a positive control, according to:         SOP 007J026 "Procedure for Determination of the Concentration of <i>Geobacillus stearothermophilus</i> (ATCC 7953)         Spore Suspension and Strips used for Autoclave Validation".         Results shall be recorded on:				
<ul> <li>Acceptance Criteria         There shall be no growth of BIs, which have undergone the sterilising cycles.     </li> <li>There shall be positive growth of the control BIs, which have <i>not</i> undergone the sterilising cycles.</li> <li>Bio-indicators shall comply with specification before use: Glass ampoules <i>G.stearothermophilus</i>, The Biological indicator shall have:         Population &gt; 5 x 10<sup>5</sup> Geobacillus stearothermophilus (ATCC 7953) per ampoule, with a "D" value of not less than 1.5 minutes and not more than 2.0 minutes.     </li> <li>Initial population shall be confirmed by QC.</li> <li>PNSU calculations are for information only and shall be reviewed with reference to original background bioburden data, where available.</li> </ul>				
TR 010X066 and Certificate of Analysis for the BI ampoules s and conclusion)	Date:			
nss/Fail ff et anager				
	Is – Biological Indicator Ampoules (RM 6119) ate a non-sterilised spore strip as a positive control, according Determination of the Concentration of <i>Geobacillus stearotherr</i> used for Autoclave Validation". In steriliser (Autoclave) Biological Indicator Result Sheet". Is, which have undergone the sterilising cycles. of the control BIs, which have <i>not</i> undergone the sterilising cy th specification before use: Glass ampoules <i>G.stearothermo</i> <i>trillus stearothermophilus (ATCC 7953)</i> per ampoule, than 1.5 minutes and not more than 2.0 minutes. irmed by QC. formation only and shall be reviewed with reference to origination only and shall be reviewed with reference to origination only and shall be reviewed with reference to origination only and shall be reviewed with reference to origination only and shall be reviewed with reference to origination only and shall be reviewed with reference to origination only and shall be reviewed with reference to origination only and shall be reviewed with reference to origination only and shall be reviewed with reference to origination only and shall be reviewed with reference to origination only and shall be reviewed with reference to origination. Ss/Fail ff t			

Document No	Controlled Copy No.	Effective Date	Page 17 of 17 Review: N/A
Prepared by	Date	Authorised By	Date

# Appendix A: Standard Load Components for Qualification

## Minimum Load

No. off	Description	Material	Mass single item (g)	Total mass (kg)
Total mass (kg)				

## Maximum Load

No. off	Description	Material	Mass single item (kg)	Total mass (kg)
	Total mass (	ka)		
Total mass (kg)				

• Designed to simulate Product xxx 15 mg in 1 mL product, aseptically filled into 2 mL glass vials, stoppered and capped.

#### Appendix B: Loading Pattern Diagrams

Appendix C: Procedure for processing raw data from heat distribution/penetration studies

Appendix D: Procedure for sealing probes inside vials for heat penetration studies