# CERTIFICATE OF COMPLIANCE CERTIFICATE OF ANALYSIS

FOR

## INHALATION DEVICE

## **TORUS INHALER**

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### **1 FREIGABE FUNKTIONSSPEZIFIKATION / ÄNDERUNGSINDEX**

Version	Datum	Freigabe	ÄNDERUNGEN /CHANGES
0.1	14.02.2019	Markus Mezger	Erstausgabe/ First Issue

Version Datum

#### **ÄNDERUNGEN /CHANGES**

0.1	14.02.2019	Erstellung COA, COC für Torus Inhaler

### 2 REFERENZDOKUMENTE

Dokument /-herkunft	Version	Datum
Erstvorlage von Markus Mezger	0.1	21.04.2015
URS draft version	0.3	22.11.2018
FMEA draft version	0.1	16.12.2018

### **3 ABKÜRZUNGEN / ABBREVIATION**

Allgemeine Abkürzungen / General Abbrevation

Auftragnehmer
Actuator-Sensor-Interface (Feldbussystem)
Berufsgenossenschaft Vorschrift
Code of Federal Regulations
Deutsches Institut für Normung e. V.
Elektromagnetische Verträglichkeit
Europäische Norm
First In First Out (Speicherorganisationsprinzip)
Failure Mode and Effect Analysis
Gerätesicherheitsgesetz
Good Manufacturing Practice
Hochfrequenz
Human Machine Interface
International Electrotechnical Commission (Internationale Normungs Organisation
In Process Control, Industrie-PC
National Fire Protection Agency
Open Process Communication Standard
Supervisory Control And Data Acquisition (Linienleitsystem)
Standard Operation Procedure (Arbeitsanweisung)
Underwriters Laboratory (Amerikanische Sicherheits- und Normen Organisation)
Unterbrechungsfreie Stromversorgung
Verband der Elektrotechnik Elektronik Informationstechnik e.V.
Extended Markup Language (Communication format standard)
Title 21 Code of Federal Regulations Part 11 (US-amerikanisches Bundesgesetz zum Thema "Electronic Records, Elektronic Signatures")
Design Qualification (Designqualifizierung)
"wie gebaut" d.h. der tatsächlichen Installation entsprechend
Jede mögliche Kombination von Text, Graphiken, Daten, Bild- oder sonstigen Informationsdarstellungen in digitaler Form, die mit Hilfe eines Rechnersystems erstellt, modifiziert, gepflegt, archiviert, wiedergefunden oder verteilt werden.
Good Automated Manufacturing Practice (Leitfaden der ISPE zur guten Herstellungspraxis mit automatisierten Systemen)
Factory Acceptance Test (Prüfung beim Hersteller
Installation Qualification (Funktionsqualifizierung)
Operational Qualification
Performance Qualification (Leistungsqualifizierung)
Qualification Master Plan
Site Accetptance Test (Prüfung am Installationsort)
System Life Cycle (Lebenszyklus eines systems)
Open-Point-List
User Requirement Specification (Benutzeranforderung)
Proof of Principle

#### 3.1 Projekt Spezifische Abkürzungen / Project Specific Abbreviations

The 'definitions' section is used to define all terms and phrases which are used in the body of the specification but which may be ambiguous in their meaning. The definitions captured in this generic specification are likely to be required but care should be taken to capture any other words or phrases which require defining.

Active Ingredient	Active Ingredient The part of the formulation with therapeutic effect.		
Delivered Dose	The mass of active ingredient received by the patient during a single inhalation. This can also be known as the Emitted Dose.		
Device	The fully assembled inhaler (excluding all packaging).		
Device Pressure Drop	A pressure drop created through the device by an inspiratory flow rate.		
Dose Content Uniformity	The repeatability of the delivered dose during the lifetime of a device and between devices. Abbreviated to DCU.		
Fine Particle Dose	The mass of active ingredient with an aerodynamic diameter of 5 microns or less received by the patient during a single inhalation. Abbreviated to FPD.		
Fine Particle Fraction	The percentage of delivered dose with an aerodynamic particle size of 5 microns or less. Abbreviated to FPF.		
FMECA	Failure Mode, Effects and Criticality Analysis – a bottom up risk management technique / design tool that systematically identifies failures that lead to hazardous conditions.		
Formulation	The active ingredient stored in the inhaler and its associated lactose carrier.		
FTA	Fault Tree Analysis – a top down risk management technique / design tool that uses hazardous conditions to systematically identify device or system failures.		
Inhalation Path	The path of air into and out of the device and into the patient's mouth.		
Metered Dose	The mass of active ingredient removed from the bulk reservoir during normal operation or for discrete dose devices - the mass of active ingredient contained in a		

Inhaler	Certificate of Analysis	Marrata a Calutian
27.08.2018	Certificate of Compliance	ivientatec Solution
Nominal Dose	discrete dose chamber. The metered dose for the capsule device is the capsule itself. The label claimed delivered dose per	
	normal operation, which is equal to the nominal, delivered dose.	
Normal Operation	The operation of the device performing all the operating steps in the correct operating sequence and generating a device pressure drop of 4kPa. A 4kPa pressure drop represents a medium resistance inhaler as defined by the EP and USP. Where a low or high resistance inhaler is being specified the pressure drop values will need to be altered accordingly.	
Nominal Actuation	[TBD]. This will be the orientation of the device during	
Orientation normal actuation.		
Nominal Inhalation	[TBD]. This will be the orientation of the device during	
Onentation	normal initialation.	
Patient	The person who uses the device to receive the formulation.	
PIF	Median Peak Inspiratory Flow (PIF)	
Particle Size Distri	bution The distribution of particle sizes in the delivered dose.	
RSD	Relative Standard Deviation = 100 x (Standard deviation / Mean).	
Secondary Packag	ing All external packaging surrounding the device during pre-sale storage.	
Shot weight The mass of formulation delivered from the device on a single actuation during normal operation.		
S.G.	Specific Gravity = density of a substance / density of water at 4°C (1000kg/m <sup>3</sup> ).	
Specification	A dose is defined as being delivered to specification if its fine particle dose is within $\pm 20\%$ of the fine particle dose during normal operation.	

### 4 EINFÜHRUNG / INTRODUCTION

This document has been written as a detailed description for a dry powder inhaler (DPI) Torus inhaler device.

This document contains regulatory requirements and industry standards present at the time of writing. A careful check of all current requirements should be made prior to issuing the specification for a product. Where no regulatory or industry standards exist, Mematec Solutions has used other source materials and its experience of DPI design to set requirements.

Blue text is text that has been added for instructions on how to use the generic specification template. This does not form part of the generic requirements specification and should be removed from the tailored specification

#### 4.1 Zielvorgabe des Dokuments / Objective of the document

The objective of this document is to define the requirements for the [TBD] dry powder inhaler. It provides the full set of requirements against which the system will be validated and, as such, forms the primary reference document for use during design and development.

### **5 RICHTLINIEN / REFERENCED DOCUMENTS**

Where appropriate, sources or fuller explanations of requirements should be given. This section defines the abbreviations used for these references to reduce the amount of text required in the body of the specification.

These documents only form requirements to the extent explicitly stated wherever they are referenced in this document.

[ACPS]	Parametric tolerance Interval test for Delivered Dose Uniformity" Advisory Committee for Pharmaceutical Science October 2005	
[ADULTDATA]	Adultdata: The handbook of adult anthropometric and strength measurements (Published by Product Safety and Testing Group (PSTG), 1998).	
[BATT]	Directive 2006/66/EC of the European Parliament and of the Council on batteries and accumulators and waste batteries and accumulators	
[BIO]	Draft Guidance for Industry - Bioavailability and Bioequivalence Studies for Nasal Aerosols and Nasal Sprays for Local Action: FDA, 2003	
[CHILDDATA]	Childdata: The handbook of child measurements and capabilities (Published by PSTG, 1995).	
[CHOKING]	BS EN 71-7:2002 - Safety of toys. Finger paints. Requirements and test methods.	

Inhaler	Certificate of Analysis		
27.08.2018	Certificate of Compliance	Mematec Solutior	
	ASTM F963-07e1- Standard Consumer Safety Spe Toy Safety.	ecification on	
[CYTO]	ISO10993-parts 1 to 18: 1997-2008 - Cytotoxicity.		
[DOSE COUNTER]	Guidance for Industry - Integration of Dose-Countin Mechanisms into MDI Drug Products: FDA, 2003.	ng	
[DROP]	IEC 60068-2-32:1975 Environmental Testing- Test Free fall	s.Test Ed:	
[ELEC SAFETY]	ELEC SAFETY] IEC 60601-1 Medical electrical equipment.General requirements for safety.		
	IEC 60601-1-1 Medical electrical equipment, collat standard. Safety requirements for medical electricated	eral al systems.	
	IEC 60601-1-4 Medical electrical equipment, collat standard. General requirements for programmable medical systems	eral electrical	
[EMC]	IEC 60601-1-2 Medical electrical equipment, collateral standard. Electromagnetic compatibility:requirements and tests		
	IEC 61000-4-2 2001 Electromagnetic compatibility measurement techniques – electrostatic discharge tests	: Testing and immunity	
	IEC 61000-4-2 2001 Electromagnetic compatibility measurement techniques – Radiated , radi electromagnetic field immunity test.	<ul> <li>Testing and</li> <li>frequency,</li> </ul>	
[EMEA1]	Note for Guidance on Dry Powder Inhalers: CPMP	, 1998.	
[EMEA2]	Guideline on the pharmaceutical quality of inhalation products: CHMP, 2006	on and nasal	
[EP]	European Pharmacopoeia, Sixth Edition, 2008		
[EPAG]	Summary of test methods for use and misuse of in EPAG. Available from <a href="https://www.epag.co.uk/Inuse.htm">www.epag.co.uk/Inuse.htm</a>	halers:	
[GUIDANCE]	Draft Guidance for Industry - Metered Dose Inhaler (MDI) and Dry Powder Inhaler (DPI) Drug Products: FDA,1998.		
[IPAC-RS]	A Parametric Tolerance Interval Test for Improved Control of Delivered Dose Uniformity of Orally Inhaled and nasal Drug Products: IPAC-RS, Nov. 2001.		
[ISO13485] BS EN ISO 13485: 2003 - Quality Systems - Medical Devices - particular requirements for the application of EN ISO 9001.			

Inhaler	Certificate of Analysis		
27.08.2018	Certificate of Compliance		
[ISO20072]	Draft BS ISO 20072 2007 Aerosol drug delivery devices design verification – Requirements and test methods		
[ISO9001]	Quality Systems, BS EN ISO 9001: 2000 (will be updated 2008		
[ודו]	ISO10993-1-17: 1992-2007 – Intracutaneous Toxicity or Irritation.		
[LABEL]	Guidance on Medical Device Patient Labelling; Final Guidance for Industry and FDA reviewers: FDA, 2001.		
[LIGHT] BS 2782-5, Method 540G:1995: Methods of testing p Optical and colour properties, weathering. Methods of ex to laboratory light sources. Open flame carbon-arc lamps			
[MDD]	Medical Device Directive, 93/42/EEC, 1993.		
	Essential requirements Annex 1/93/42/EEC		
[PARTICULATES]	"Best Practises for Managing Quality and Safety of Foreign Particles" J Blanchard et al Pharmaceutical Research 24.3 2007 471-479		
	"Measurement and Identification of Foreign Particles in a QbD Environment Streamlining with Efiicient Analytical Methods" O.Valet and M. Lankers Respiratory Drug Delivery 2008		
	"Proposal for Foreign Particles Testing in Orally Inhaled and Nasal Drug Products" J. Hart Respitory Drug Delivery 2004		
[OLDERADULT]	Older Adultdata: The handbook of Measurements and Capabilities of the Older Adult (Published by PSTG,2000).		
[QSR]	Current Good Manufacturing Practice (cGMP) for Medical Devices & In Vitro Diagnostic Products - Quality Systems Regulation, 21 CFR Part 820, FDA, April 2008.		
[RECYCL]	ISO 14021:1999 Environmental Labels and Declarations		
[ROHS]	Directive 2002/95/EC of the European Parliament and of the Council of 27 January 2003 on the "Restriction of the Use of certain Hazardous Substances in electrical and electronic equipment"		
[Control]	Directive 21CFR 820.30 for US required Design controls		
[SENS]	ISO10993-1-17: 1992-2007 – Sensitization.		
[USP]	United States Pharmacopoeia 31, 2008.		
[VENTILATION]	BS 7272-1:2000 - Writing and marking instruments. Specification for safety caps.		

Inhaler	Certificate of Analysis		
27.08.2018	Certificate of Compliance	Mematec Solution	
[VIBRATION1]	IEC 60068-2-64: 1993- Environmental testing. Test methods. Test Fh. Vibration, broad band random (digital control) and guidance		
	IEC 60068-2-27: 2008 Environmental testing.T guidance: Shock	est Ea and	
[VIBRATION2]	IEC 60068-2-6: 2008- Environmental testing. To Test Fc. Vibration (sinusoidal).	est methods.	
	IEC 60068-2-7: 1993 - 60068-2-7: 1983 - Environm Test methods. Test Ga and guidance. Accelera state.	ental testing. ation, steady	
	IEC 60068-2-29: 1993 - Environmental testing. T Environmental testing procedures. Tests. Te guidance. Bump.	est methods. st Eb and	
	US Standards:		
	ASTM D3332-99 (2004)– Standard Test I Mechanical-Shock Fragility of Products, Using Sho	Vethods for ck Machines.	
	ASTM D3580-95 (2004)– Standard Test Methods (Vertical Linear Motion).	for Vibration	
	ASTM D5112-98 (2003)– Standard Test method (Horizontal Linear Sinusoidal Motion) Test of Produ	for Vibration icts.	
	Electrical Standards:		
	IEC 60601-1: 2005 Medical Electrical Equipm General Requirements for Safety.	ent- Part 1:	
[WEEE]	Directive 2002/96/EC of the European Parliament a Council of 27 January 2003 on "Waste Electrical ar Equipment"	and of the Id Electronic	
[FDA]	Draft Guidance from 22. June 2011. Applying Human and usability engineering to optimize medical device	an Factors e design	

### 6 EINLEITUNG/ GENERAL DESCIPTION

#### 6.1 Anwendungsbereich / Area of Use:

Blister based device for Seritide.

### 7 PATENTE:

Torus Patente which are applied.

"EU- Geschmacksmuster" 003003086-0001 "Designschutz in D" 402015201982.3

### 8 MODULARER AUFBAU, MODULAR STRUCTURE



Standard:

Varianten/ Variants:

Blister strip 61 cavity and later 31 cavities. 31 Cavity is not part of initial design order

**Optionen/ Options:** 

Different strength of Seritide

#### 9 PACKAGING

The Device Torus will be available full assembled and delivered in bulk. The device is packed in a PE Bag. The number of device per bag is 500x.

### **10 SHIPMENT OF THE DEVICE**

- Shipment procedure

The Device will be shipped in 1x cartons with the amount of 1000 Devices. The Dimension of the carton is: LxBxH in cm (dimension can be varied from Customer to customer)

### **11 STABILITY OF THE DEVICE**

The Device and is selected materials are designed for a minimum time of storage in ambient temperature of minimum 2 years.

### **12 MATERIAL SELECTION AND CERTIFICATES**

Description of the materials which	ale useu.	
Description of the materials	Device, blister based, called	Desciption:
and components:	TorusHaler	
		<ul> <li>Dimensions, Sizes,</li> </ul>
	ROFW	<ul> <li>ø:approx. 86mm, W:approx.30mm</li> </ul>
	14	Weight:
Torus Inhaler		- Material: plastic housing with Alu -Alu
		Blister
		<ul> <li>Ambient conditions, RH and T 15-</li> </ul>
		30°C at 35-65% RH
		<ul> <li>Tolerances are defined in mematec parts</li> </ul>
		drawings
		- Colour Body Base: White
Description of the materials	Pulver, Powder:	Description:
and components		
	Seritide in different strenght.	Description
		- Mass: 12,5mg
		- Density: tbd
		<ul> <li>Moisture sensitivity: needs to be protected</li> </ul>
		against moisture
		- Melting point: tbd
		- Segregation: yes due to blend
		- Particle size: 2-100µ
		- Particle size distribution:
Description of the materials	Part 1	Description:
and components	Mouth Piece	
		- Dimensions: 24 70mm x 45 6mm x 13 42mm
		- Material: EDA Confirm PP
		- Colour : white
		- Environmental conditions: 15-85% RH and T: -
Mouthniece		
Modulpiece		- Surface: electro- polished
		$= 0.001 \text{ g/mm}^3$
		- Mass $= 1.2  g$
		- 1.2  g
	•	
	Part in contact with patient	
	requires FDA confirmation	

Part 2 Base Wheel	Description: - Dimensions: L= 21.2mm , Ø= 18.35mm - Material: No FDA confirmed POM - Color: white - mass = 1.7 g - Environmental conditions: 15-85% RH and - Temperature: 10°C - 45°C - Surface: Line polish RZ 1 - Hole for mounting position
Part 3 Outer case	Description of the - Dimensions: 66.78mm x 49.62mm x27.80mm - Material: FDA Confirm ABS - Colour : white - Environmental conditions: 15-85%RH - Temperature: -10°C - 45°C - Surface: polished - Density = 0.001 g/mm3 - mass = 10.1 g - Volume = 10100mm3
Part 4 Manifold	Description - Colour : white - Environment condition: 15-85%RH - Temperature: 10°C – 45°C - Surface: polished Puffer for empty blister strip and puffer for blister strip with 61 filled powder cavities. - Density = 0.001 g/mm3 - Mass = 9.1 g - Volume = 9100 mm3
Part 5 Counting Wheel	Description -60 figures in total, 55figures in black printing. - Figures 5-1 in red printing. - Printing in tamp print technology - No- Stop function end of strip. -Key hole for assembly position - Counting wheel counts remaining shots left - Material in ABS - Letter Size 2,5mm -mass = 1.2 g
	Part 2 Base Wheel

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Description of the materials and components Body Base	Part 6 Body Base	Description of the - Material: FDA Confirm ABS - Colour : White - Environment: 15-85%RH - Temperature: -10°C – 45°C - Surface: polished - Density = 0.001 g/mm3 - Mass = 12.5 g - Volume = 12500 mm3
Description of the materials and components Body Base half top	Part 7 Body Base half top	<ul> <li>Material: FDA Confirm ABS</li> <li>Colour : white</li> <li>Environment condition: 15-85%RH</li> <li>Temperature: -10°C - 45°C</li> <li>Surface: polished</li> <li>Density = 0.001 g/mm3</li> <li>Mass = 14.5 g</li> <li>Volume = 14500 mm3</li> </ul>
Description of the materials and components Index Wheel ratchet	Part 8 Index Wheel ratchet	Description - Dimensions: L= 19.3 mm , Ø= 18.195mm - Material: None FDA Confirm POM - Colour : White -mass = 1.3 g - Environment condition: 15-85% RH - Temperature: -10°C – 45°C - Surface: polished
Description of the materials and components Index Wheel	Part 9 Index Wheel	Description - Dimensions: L= 16.50mm , Ø= 22.74mm - Material: None FDA Confirm POM - Colour : White -mass = 2.3 g - Environment condition: -15-85% RH - Temperature: -10°C – 45°C - Surface: polished

Contracting wheel arm	Part 10 Contracting wheel arm	<ul> <li>Description</li> <li>Dimension: L= 15.3mm , Ø= 20.35mm</li> <li>Material: None FDA Confirm POM</li> <li>Colour : White</li> <li>mass = 1.5 g</li> <li>Environment condition: 15-85% RH</li> <li>Temperature: -10°C – 45°C</li> <li>Surface: polished</li> </ul>
Description of the materials and components Contractiong wheel ratchet	Part 11 Contractiong wheel ratchet	Description - Dimensions: L= 19mm , Ø= 17mm - Material: None FDA Confirm POM - Colour : White -mass = 0.7 g - Environment condition: 15-85% RH - Temperature: -10°C – 45°C - Surface: polished
		-
Description of the materials	Part 12	Description
and components Contracting wheel gear	Contracting wheel gear	<ul> <li>Dimension: L= 2mm, ø= 22.4mm</li> <li>Material: None FDA Confirm POM</li> <li>Colour : White</li> <li>mass = 0.4 g</li> <li>Environment condition: 15-85% RH</li> <li>Temperature: -10°C – 45°C</li> <li>Surface: polished</li> </ul>
Description of the materials and components	Part 13 Idler Gear	Description - Dimension: L= 2mm, ø= 10.52mm - Material: FDA Confirm ABS - Colour : white - mass = 0.6 g
Idler Gear		<ul> <li>Environment condition: 15-85%RH</li> <li>Temperature: -10°C – 45°C</li> <li>Surface: polished</li> <li>Assembly position will be fixed by pin. The fixation will keep all gear box parts in the starting position. Will hold the counting wheel in position in pre assembly, avoids falling out of housing</li> </ul>

Description of the materials and components Drive Wheel gear	Part 14 Drive Wheel gear	Description - Material: None FDA Confirm POM - Colour : White -mass = 0.9 g - Environment condition: 15-85% RH - Temperature: -10°C – 45°C - Surface: polished
Description of the materials and components Cover	Part 15 Cover	Description - Dimension : L= 3.1mm , Ø= 21.3mm - Material: None FDA Confirm POM - Colour : White -mass = 2 g - Environment condition: -15-85% RH - Temperature: -10°C – 45°C Surface: Polished for laser printing of variable and fixed data. Eroded surface for better grip while handling the device
Decemination of the metanicle	Dert 40	
Description of the materials	Part 16 Blistor Strip	Material Lid foil:
Blister Strip		Lid foil Paper 50 g/m² PET 12 μm Aluminum 20 μm HSL-peelable 7 g/m²
	Paper: stiffness, avoids sticking to sealing tool	Locating mandrel in HH machine: 150mm Film width: 117 mm Blister strip length: 710 mm
	<ul> <li>Paper: stiffness, avoids sticking to sealing tool</li> <li>PET: stability</li> <li>Aluminium: barrier</li> <li>HSL: wide sealing window, easy peel</li> <li>PVC: rigidity, sealing layer</li> <li>Aluminium: barrier, form ability</li> <li>oPA: supporting the forming</li> </ul>	Locating mandrel in HH machine: 150mm Film width: 117 mm Blister strip length: 710 mm <b>Material Base Foil:</b> mandrel in HH machine: 150mm Film width: 115 mm Blister strip length: 710 mm Blister strip length: 710 mm
	<ul> <li>Paper: stiffness, avoids sticking to sealing tool</li> <li>PET: stability</li> <li>Aluminium: barrier</li> <li>HSL: wide sealing window, easy peel</li> <li>PVC: rigidity, sealing layer</li> <li>Aluminium: barrier, form ability</li> <li>oPA: supporting the forming</li> </ul>	Locating mandrel in HH machine: 150mm Film width: 117 mm Blister strip length: 710 mm <b>Material Base Foil:</b> mandrel in HH machine: 150mm Film width: 115 mm Blister strip length: 710 mm Base foil PVC 100 µm Aluminium 45 µm

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### **13 RESPIRATOY DEVICE RESISTANT DATA**

#### **Device Resistent Data**

Flow Rate in l/min	Pressure Drop P1 in kPa	Pressure Drop P2 in kPa	Pressure Drop P1 in kPa	Pressure Drop P2 in kPa
	Mematec MDPI		GSK Originator	r Device
20	1,65	0,88	1,60	0,36
40	2,10	0,88	2,40	0,36
60	4,05	0,88	3,7	0,36
80	6,2	0,88	5,4	0,36
90	7,35	0,88	6,5	0,36

#### Salmeterol

Flow Rate in l/min at 4 kPa	Delivered dose in µg	Fine particle dose in µg	Fine particle fraction in %	MMAD	GSD
74,6	49,13	8,322	16,939	2,668	2,433
74,55	48,925	7,725	15,790	2,570	2,471
72,62	50,463	9,407	18,642	2,523	2,445
74,76	46,405	8,890	19,157	2,371	2,511

#### Fluticasone

Flow Rate in l/min at 4 kPa	Delivered dose in µg	Fine particle dose in µg	Fine particle fraction in %	MMAD	GSD
74,6	213,066	44,675	20,968	2,629	2,346
74,55	215,271	48,675	22,611	2,546	2,313
72,62	241,113	59,638	24,734	2,569	2,267
74,76	211,853	55,958	25,223	2,393	2,398

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### **14 USER DESCRIPTION:**



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### **15 ENVIROMENTAL REQUIREMENTS**

Description

Device will deliver doses to specification and have no visible defects when operated in an environment of 18°C to 28°C at 25%R.H. to 75%R.H. [ISO20072] (This simulates 'in use' conditions only.)

Device will deliver doses to specification and have no visible defects when operated in an environment of 5+/- 3°C and 40+/- 2°C at 50% R.H. [ISO20072]

Device will operate to specification when stored below 0°C for at least 3 hours and then immediately used [EMEA2 4.2.1.15].

Device will deliver doses to specification when operated in an environment of between 10% and 90% R.H., at 25°C. (This requirement is recommended due to variation of surface charge and brittleness of capsules with ambient humidity.) [EMEA1, 2.4.5], [EMEA2 4.2.1.17].

#### 15.1 GMP – Requirements / Hygiene

Funktions- Beschreibung

Device will not require cleaning during normal use. Cleaning of the device is possible using regular clean water

Device will have a Mouth Piece cover to protect the mouth Piece of dust and dirt

Device will deliver a dose to specification immediately after the patient cleans the device as per the cleaning instructions for the device [EPAG]. For example, cleaning the device with a cloth will not leave a static charge on it that would prevent the device from subsequently delivering a dose to specification.

#### **15.2 Environment Protection of the Device**

Description of fulfilled requirements:

Device will be RoHS confirm

Device will be Reach Declaration confirm

Device will be Packaging and Packaging Waste confirm

Device will be WEE- Declaration confirm

#### 15.3 Moisture

Will deliver doses to specification if there is a delay of up to 60 seconds between actuation and inhalation.

#### 15.4 Partikel / Particulates

There are currently no clear guidelines on the particulate requirements for inhalation products.

#### Funktions- Beschreibung

Should not contain any removable parts that may be classified as a choking hazard as defined by the Small Parts Cylinder criterion detailed in [CHOKING/ Verstopfen]. There are currently no guidelines specifically addressing choking hazards for medical devices. The choking hazard presented by a removable part can be mitigated by ventilation i.e. allowing airflow through the part to delay asphyxiation pending medical intervention [VENTILATION]. However, it is recommended that removable parts be designed to be large enough to not present a choking hazard as opposed to seeking to mitigate this risk through ventilation.

Shall allow no more than  $50\mu$ g/day of foreign particles having aerodynamic diameter less than or equal to 10  $\mu$ m to become detached from the device (equivalent to 5% of the National Ambient Air Quality Standard) [PARTICULATES].

Shall allow no more than [TBD] particles with longest axis greater than [TBD] µm in length to become detached from the device and reach the inhalation path during [TBD] normal operations. (The dimension and number used is not stipulated by regulatory bodies - it is a patient confidence issue).

Shall allow no more than [TBD] particles with longest axis greater than 100µm in length to become detached from the device and reach the inhalation path during [TBD] normal operations following an impact as defined in section 6.1. (The dimension and number used is not stipulated by regulatory bodies - it is a patient confidence issue).

#### 15.5 Qualitätsmanagement / Quality Systems

Description

Device is, manufactured and distributed in accordance with the following reference documents: [QSR], [ISO9001], [ISO13585], [MDD].

Device is designed accordance with ISO 13485 (Design and manufacturing of medical devices

Device is designed accordance with MDD 93/42/EEC from 14.06.1993 (Medizinprodukte Richtlinie/ Medical Device Directive)

#### 15.6 Recycling Information

Description

Device has appropriate recycling information marked on product as defined in URS Device has appropriate colours according to the convention defined in URS

#### 15.7 Herstellung / Manufacturing

#### Molding/ Components & Subassembly

Description

Plastic parts to be molded in a room with the classification ISO 8, 100.000

Device is assembled in a room with the classification ISO 8, 100.000

Plastic parts will be final checked in a room with the classification ISO 8, 100.000

Should, where necessary, have fully functional major subassemblies self- contained for transportation without the need for jigs and fixtures.

The Device Torus have all non-metallic parts & subassemblies made under grade C EU GGMP cleanroom conditions (compares to US Federal standard 209E class 10,000).

Molding tool material is done in stainless steel. The air path is done in stainless steel. Alternative standard steel in high quality.

No lubrication and mold release agent are used for the molding process

#### **15.8 Documents to be Delivered**

Allgemein / General

#### Description

The following documents are available with the device Torus Ellipse. Layout Drawing of the device with the dimension

Plastic Parts of the device with Material Certificates

Metal parts with Material specification

CE Confirmation Declaration

#### 15.8.1.1 Bedienungsanleitung / Operation Manual

Description The handling of the device is shown in the handling instruction. A handling Video can be down loaded on the Mematec solution homepage