

### Deutsche Akkreditierungsstelle

# Annex to the Partial Accreditation Certificate D-PL-11140-07-02 according to DIN EN ISO/IEC 17025:2018

**Valid from: 26.10.2022**Date of issue: 26.10.2022

This annex is a part of the accreditation certificate D-PL-11140-07-00.

Holder of partial accreditation certificate:

Fraunhofer Gesellschaft zur Förderung der angewandten Forschung eingetragener Verein Hansastraße 27 c, 80686 München

with its testing laboratory

## Fraunhofer Institut für Produktionstechnik und Automatisierung IPA Nobelstraße 12, 70569 Stuttgart

The testing laboratory meets the minimal requirements of DIN EN ISO/IEC 17025:2018 and, if applicable, additional legal and normative requirements, including those in relevant sectoral schemes, in order to carry out the conformity assessment activities listed below.

The management system requirements of DIN EN ISO/IEC 17025 are written in the language relevant to the operations of testing laboratories and confirm generally with the principles of DIN EN ISO 9001.

Field: Medical devices

**Testing fields/test items:** Biological tests of medical devices;

environmental monitoring

This certificate annex is only valid together with the written accreditation certificate and reflects the status as indicated by the date of issue. The current status of any given scope of accreditation can be found in the directory of accredited bodies maintained by Deutsche Akkreditierungsstelle GmbH at https://www.dakks.de.

Abbreviations used: see last page Page 1 of 3



#### Annex to the Partial Accreditation Certificate D-PL-11140-07-02

Testing field	Test item Device(category)	Type of testing Test	Regulation Testing method
Biological testing	Medical devices	Cytotoxicity test	DIN EN ISO 10993-5 SAA IVT 01-0
		Activity of metabolism after contact with extracts or direct contact (MTS-test)	applicable: DIN EN ISO 10993-1 DIN EN ISO 10993-12
Environmental monitoring in the process of production and testing of cleanliness of the products according to DIN EN ISO 13485:2021 <sup>1</sup> , Par. 6.4 and Par. 7.5			
Physical testing	Medical devices, Surfaces	Testing of particulate impurities (Light microscopy)	VDI 2083, Part 21

#### **Regulations:**

DIN EN ISO 10993-1 Biological evaluation of medical devices - Part 1: Evaluation and

2021-05 testing within a risk management process

DIN EN ISO 10993-5 Biological evaluation of medical devices - Part 5: Tests for in vitro

2009-10 cytotoxicity

DIN EN ISO 10993-12 Biological evaluation of medical devices - Part 12: Sample prepa-

2021-08 ration and reference materials

VDI 2083 Cleanroom technology - Cleanliness of medical devices in the

Part 21 manufacturing process

2019-10 B3.3, Particulate impurities, Light microscopy

(automated with image processing)

SAA IVT 01-0 In-vitro-Zytotoxizität nach DIN EN ISO 10993-5

Rev. 0

Valid from: 26.10.2022 Date of issue: 26.10.2022



#### Annex to the Partial Accreditation Certificate D-PL-11140-07-02

#### abbreviations used:

DIN German Institute for Standardization

EN European Standard

IEC International Electrotechnical Commission
ISO International Organization for Standardization
SAA IVT XXX Standard operating procedure of Fraunhofer IPA

VDI Association of German Engineers (German: Verein Deutscher

*Ingenieure)* 

<sup>1</sup> DIN EN ISO 13485:2021: Medical devices - Quality management systems - Requirements for regulatory purposes

Valid from: 26.10.2022 Date of issue: 26.10.2022