

Wir, die Firma

Nous, la maison

We, the company

**Cendres+Métaux SA**  
**Rue de Boujean 122**  
**CH-2501 Biel/Bienne**

Bevollmächtigter

Mandataire

Authorized Representative

**Cendres+Métaux France SAS**  
**Les petites Buffeteries**  
**F-49124 St-Barthélémy d'Anjou**erklären hiermit in alleiniger  
Verantwortung, dass die folgen-  
den Produktedéclarons sous notre seule  
responsabilité que les produits  
suivantsdeclare under our sole responsibility that  
the following products**Stege****Barres****Bars**

gemäß beiliegender Liste

selon liste annexée

according to attached list

[List\\_CM-Bars LDP-KE3 Version 5](#)konform sind zu den folgenden  
normativen Dokumenten:sont conformes aux documents  
normatifs suivants:are in conformity with the  
following normative documents:**Grundlegende Anforderungen**  
gemäß Anhang I, 93/42/EWG,  
2007/47/EG**Exigences essentielles**  
selon l'annexe I, 93/42/CEE,  
2007/47/CE**Essential Requirements**  
according to annex I, 93/42/EEC,  
2007/47/EC**Normen:** vgl. beiliegende Liste**Normes:** cf. liste annexée**Standards:** see attached list[List of laws, standards and norms Product Group Bars \(KE 3\) Edition 05](#)**Klassifizierung** gemäß  
Anhang IX, 93/42/EWG:  
vergl. beiliegende Liste**Classification** selon  
l'annexe IX, 93/42/CEE:  
Cf. liste annexée des produits**Classification** according to  
annex IX, 93/42/EEC:  
see attached list of products[List\\_CM-Bars LDP-KE3 Version 5](#)**Konformitätsbewertungs-  
verfahren:****Procédure d'évaluation de la  
conformité:****Conformity assessment procedure:**

93/42/EWG

93/42/CEE

93/42/EEC

Klasse I: Anhang VII

Classe I: Annexe VII

Class I: Annex VII

Übrige Klassen: Anhang II  
(ohne Abschnitt 4)Autres classes: Annexe II  
(sans section 4)All other classes: Annex II  
(Without section 4)**Benannte Stelle:****Organisme notifié:****Notified Body:**Klasse I: Benannte Stelle nicht  
erforderlich.Classe I: Pas d'organisme notifié  
nécessaire.

Class I: No notified body needed.

Übrige Klassen:

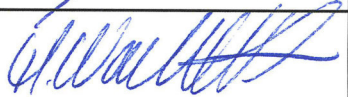
Autres classes: MNC

All other classes:

mdc medical device certification GmbH, Kriegerstraße 6, 70191 Stuttgart / Germany (CE 0483)

**Gültigkeit****Validité****Validity****Änderungen:****Modifications:****Changes:**[Änderungsantrag MD5-3252](#)[demande modification MD5-3252](#)[change request MD5-3252](#)Diese Konformitätserklärung wird  
mit den folgenden Unterschriften  
beglaubigt:Cette déclaration de conformité  
est certifiée par les signatures  
suivantes:This declaration is certified by the  
following signatures:Biel/Bienne, **2021-07-05**  
Bibiana Gamber  
Director of Quality & Regulatory  
Raymond Cuany  
Product Safety Officer

| #  | Catalog Number | Product Name                                          | Device group         | Family | Class MDD | Rule MDD      |
|----|----------------|-------------------------------------------------------|----------------------|--------|-----------|---------------|
| 1  | 05050010       | Ackermann-Bar A Female part E                         | Ackermann-Bar        | Bars   | Ila       | 5, 3rd indent |
| 2  | 05050011       | Ackermann-Bar B Female part E                         | Ackermann-Bar        | Bars   | Ila       | 5, 3rd indent |
| 3  | 052043         | Dolder® System Female part E micro L50                | Dolder® System       | Bars   | Ila       | 5, 3rd indent |
| 4  | 052046         | Dolder® System Female part E macro L50                | Dolder® System       | Bars   | Ila       | 5, 3rd indent |
| 5  | 054746         | Dolder® System Female part E micro L25                | Dolder® System       | Bars   | Ila       | 5, 3rd indent |
| 6  | 054747         | Dolder® System Female part E macro L25                | Dolder® System       | Bars   | Ila       | 5, 3rd indent |
| 7  | 05000680       | Dolder® System Female part T micro L47.5              | Dolder® System       | Bars   | Ila       | 5, 3rd indent |
| 8  | 05000681       | Dolder® System Female part T macro L47.5              | Dolder® System       | Bars   | Ila       | 5, 3rd indent |
| 9  | 05001125       | Dolder® System Female part D macro L50                | Dolder® System       | Bars   | Ila       | 5, 3rd indent |
| 10 | 05001201       | Dolder® System Female part D micro L50                | Dolder® System       | Bars   | Ila       | 5, 3rd indent |
| 11 | 050527         | Round bar with rider Female part E                    | Round Bar with Rider | Bars   | Ila       | 5, 3rd indent |
| 12 | 055801         | Round bar with rider Female part E                    | Round Bar with Rider | Bars   | Ila       | 5, 3rd indent |
| 13 | 05000679       | Round bar with rider Female part E L50                | Round Bar with Rider | Bars   | Ila       | 5, 3rd indent |
| 14 | 05050014       | Ackermann-Bar Male part P3 L60                        | Ackermann-Bar        | Bars   | Ilb       | 8, general    |
| 15 | 052053         | Dolder® System Male part E macro L50 (Bar attachment) | Dolder® System       | Bars   | Ilb       | 8, general    |
| 16 | 052057         | Dolder® System Male part E micro L50 (Resilient bar)  | Dolder® System       | Bars   | Ilb       | 8, general    |
| 17 | 052061         | Dolder® System Male part E macro L50 (Resilient bar)  | Dolder® System       | Bars   | Ilb       | 8, general    |
| 18 | 05000289       | Dolder® System Male part E micro L50 (Bar attachment) | Dolder® System       | Bars   | Ilb       | 8, general    |
| 19 | 052028         | Round bar with rider Male part P3 L200                | Round Bar with Rider | Bars   | Ilb       | 8, general    |
| 20 | 052030         | Round bar with rider Male part P3 L50                 | Round Bar with Rider | Bars   | Ilb       | 8, general    |

|           | First Name / Name | Function                          | Date       | Signature                                                                             |
|-----------|-------------------|-----------------------------------|------------|---------------------------------------------------------------------------------------|
| Creation: | Dylan Moretti     | Junior Regulatory Affairs Manager | 01.07.2021 |  |
| Review:   | Markus Blümli     | Product Manager                   | 1.7.2021   |  |
| Release:  | Matthias Walther  | Director of Development           | 01.07.2021 |  |

**Change Control:**

| Version   | Date       | Description of the Change                                                                                                                                                  | Reason of the Change                                                       |
|-----------|------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------|
| Version 2 | 01.02.2016 | Integration of all spare parts in the lists (no spare part articles on the list of Bars)                                                                                   | spare parts are in al points of veiw equivalent to the rest of articles    |
| Version 2 | 11.02.2016 | Integration of all products on demand in this list (17 classe IIa-rule 8 articles)                                                                                         | products on demand are in all points of view equal to the rest of articles |
| Version 3 | 12.06.2020 | Basis ist die "Device Master List". Produte die im SAP Status Desinvestiert haben wurden von der Liste entfernt. Klassifikation der Produkte überprüft und ggf. angepasst. | Im Rahmen des NB Wechsels auf mdc werden die Produktlisten überprüft.      |

|           |                |                                                                                                                                                                                                                                                                                                                     |                                                                                                                                                                                                      |
|-----------|----------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Version 4 | 30.11.2020/dym | 1) Change of class and MDD rules for class IIa/IIb articles that are now in class I.<br>2) Change of classes and MDD rules for the articles that are now in NoMed (Spare Part) class.<br>3) Deletion of the NoMed and NoMed (Spare Part) articles.<br>4) Grouping of Class I articles in a single list of articles. | 1) Änderungsantrag MD5-2943<br>2) Änderungsantrag MD5-3008<br>3) NoMed class articles must not be in the Declarations of Conformity.<br>4) Provides to have a single list with all class I articles. |
|           | 08.12.2020/dym | Deletion of the disinvested products                                                                                                                                                                                                                                                                                | After verification of the list by mbl and after checking in SAP.                                                                                                                                     |
| Version 5 | 30.06.2021/dym | New classification rule (according MDD) for the Female Parts.                                                                                                                                                                                                                                                       | AEA MD5-3252                                                                                                                                                                                         |

**Valid for the project and/or the product(s): Bars (KE3)**

## 1. Laws for medical devices

The actual laws for medical devices are on CL 4.101 ("Gesetzessammlung Teil 2")

## 2. Standards and norms for medical devices

Based on SNV Service

| No | Code          | No       | Title                                                                                                                                    | Issue date | Valid                               |
|----|---------------|----------|------------------------------------------------------------------------------------------------------------------------------------------|------------|-------------------------------------|
| 1  | SN EN ISO     | 9001     | Quality management systems – Requirements                                                                                                | 2015-09    | <input type="checkbox"/>            |
| 2  | SN EN ISO     | 14001    | Environmental Management System                                                                                                          | 2015-09    | <input type="checkbox"/>            |
| 3  | SN EN ISO     | 13485    | Medical devices, Quality management systems, Requirements for Regulatory purposes                                                        | 2016-03    | <input checked="" type="checkbox"/> |
| 4  | SN EN ISO     | 14971    | Medical devices - Application of risk management to medical devices                                                                      | 2012-09    | <input checked="" type="checkbox"/> |
| 5  | SN EN ISO     | 15223-1  | Medical devices - Symbols to be used with medical device labels, labelling and information to be supplied - Part 1: General requirements | 2017-02    | <input checked="" type="checkbox"/> |
| 6  | SN EN         | 1041+A1  | Information supplied by the manufacturer of medical devices                                                                              | 2013-10    | <input checked="" type="checkbox"/> |
| 7  | SN EN ISO/IEC | 17050-1  | Conformity assessment - Supplier's declaration of conformity - Part 1: General requirements                                              | 2010-08    | <input checked="" type="checkbox"/> |
| 8  | SN EN ISO/IEC | 17050-2  | Conformity assessment - Supplier's declaration of conformity - Part 2: Supporting documentation                                          | 2004-11    | <input checked="" type="checkbox"/> |
| 9  | IEC           | 62366-1  | Medical devices - Part 1: Application of usability engineering to medical devices                                                        | 2015-02    | <input checked="" type="checkbox"/> |
|    |               |          | Technical Corrigendum                                                                                                                    | 2016-07    |                                     |
|    | SN EN         | 62366+A1 | Medical devices - Part 1: Application of usability engineering to medical devices                                                        | 2016-05    |                                     |
| 10 | SN EN ISO     | 17664    | Sterilization of medical devices - Information to be provided by the manufacturer for the processing of resterilizable devices           | 2018-04    | <input checked="" type="checkbox"/> |
| 11 | SN EN         | 1639     | Dentistry - Medical devices for dentistry - Instruments                                                                                  | 2010-03    | <input type="checkbox"/>            |
| 12 | SN EN         | 1640     | Dentistry - Medical devices for dentistry - Equipment                                                                                    | 2010-03    | <input type="checkbox"/>            |
| 13 | SN EN         | 1641     | Dentistry - Medical devices for dentistry - Materials                                                                                    | 2010-03    | <input type="checkbox"/>            |
| 14 | SN EN ISO     | 12836    | Dentistry - Digitizing devices for CAD/CAM systems for indirect dental restorations - Test methods for assessing accuracy                | 2015-12    | <input type="checkbox"/>            |
| 15 | SN EN         | 10204    | Metallic products - Types of inspection documents                                                                                        | 2004-12    | <input type="checkbox"/>            |
| 16 | ISO           | 22674    | Dentistry - Metallic materials for fixed and removable restorations and appliances                                                       | 2016-01    | <input checked="" type="checkbox"/> |
| 17 | SN EN ISO     | 4049     | Dentistry -- Polymer-based restorative materials                                                                                         | 2010-03    | <input type="checkbox"/>            |
| 18 | SN EN ISO     | 10477    | Dentistry -- Polymer-based crown and bridge materials                                                                                    | 2005-03    | <input type="checkbox"/>            |
| 19 | ISO           | 9693-1   | Dentistry - Compatibility testing - Part 1: Metal-ceramic systems                                                                        | 2012-02    | <input type="checkbox"/>            |
| 20 | SN EN ISO     | 9333     | Dentistry – Brazing materials                                                                                                            | 2006-10    | <input type="checkbox"/>            |
| 21 | SN EN ISO     | 28319    | Dentistry - Laser welding                                                                                                                | 2010-08    | <input type="checkbox"/>            |

| No | Code      | No        | Title                                                                                                                                                                                                                                                                 | Issue date | Valid                               |
|----|-----------|-----------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------|-------------------------------------|
| 22 | ASTM      | F 67      | Standard Specification for Unalloyed Titanium, for Surgical Implant Applications (UNS R50250, UNS R50400, UNS R50550, UNS R50700)<br><br>Titan Grad 1(1083)/2 KV(780)/3(720)/4(721)/4 KV(725)/4 KV Dolder(1082)/4 CP4 low iron(1273)/4B ISO 5832-2(977)/Hannibal(976) | 2013       | <input checked="" type="checkbox"/> |
| 23 | ISO       | 5832-2    | Implants for surgery - Metallic materials - Part 2: Unalloyed titanium<br><br>Titan Grad 2(556)/2 KV(780)/3(720)/4(721)                                                                                                                                               | 2018-03    | <input type="checkbox"/>            |
| 24 | ASTM      | F 136     | Standard Specification for Wrought Titanium-6Aluminum-4Vanadium ELI (Extra Low Interstitial) Alloy for Surgical Implant Applications (UNS R56401)<br><br>Syntax(555)/TiAl6V4 ELI(555)/Titan64(555)/Titan Grade 5(555)/Grade 23(555)/Certain(995)                      | 2013       | <input checked="" type="checkbox"/> |
| 25 | ASTM      | B 348     | Standard Specification for Titanium and Titanium Alloy Bars and Billets<br><br>Titan Grade 36/Ti45Nb (1228)                                                                                                                                                           | 2013       | <input type="checkbox"/>            |
| 26 | ASTM      | F 899-12b | Standard Specification for Wrought Stainless Steels for Surgical Instruments                                                                                                                                                                                          | 2012       | <input checked="" type="checkbox"/> |
| 27 | ASTM      | F2820     | Standard Specification for Polyetherketoneketone (PEKK) Polymers for Surgical Implant Applications                                                                                                                                                                    | 2012       | <input type="checkbox"/>            |
| 28 | SN EN ISO | 7153-1    | Surgical instruments - Materials - Part 1: Metals                                                                                                                                                                                                                     | 2016-12    | <input checked="" type="checkbox"/> |
| 29 | SN EN     | 10088-3   | Stainless steels - Part 3: Technical delivery conditions for semi-finished products, bars, rods, wire, sections and bright products of corrosion resisting steels for general purposes 1.4435 (stainless steel: Medstahl, Chr Spezial 35)(1028)                       | 2014-12    | <input checked="" type="checkbox"/> |
| 30 | ASTM      | F75       | Standard Specification for Cobalt-28 Chromium-6 Molybdenum Alloy Castings and Casting Alloy for Surgical Implants (UNS R30075)                                                                                                                                        | 2012       | <input type="checkbox"/>            |
| 31 | ASTM      | F799      | Standard Specification for Cobalt-28Chromium-6Molybdenum Alloy Forgings for Surgical Implants (UNS R31537, R31538, R31539)                                                                                                                                            | 2011       | <input type="checkbox"/>            |
| 32 | ASTM      | F1537     | Standard Specification for Wrought Cobalt-28Chromium-6Molybdenum Alloys for Surgical Implants (UNS R31537, UNS R31538, and UNS R31539)                                                                                                                                | 2011       | <input type="checkbox"/>            |
| 33 | ISO       | 5832-4    | Implants for surgery - Metallic materials - Part 4: Cobalt-chromium-molybdenum casting alloy                                                                                                                                                                          | 2014-09    | <input type="checkbox"/>            |
| 34 | ISO       | 5832-12   | Implants for surgery - Metallic materials - Part 12: Wrought cobalt-chromium-molybdenum alloy                                                                                                                                                                         | 2007-05    | <input type="checkbox"/>            |
| 35 | ISO       | 13356     | Implants for surgery -- Ceramic materials based on yttria-stabilized tetragonal zirconia (Y-TZP)                                                                                                                                                                      | 2015-09    | <input type="checkbox"/>            |
| 36 | SN EN ISO | 18064     | Thermoplastic elastomers - Nomenclature and abbreviated terms                                                                                                                                                                                                         | 2015-02    | <input type="checkbox"/>            |
| 37 | SN EN ISO | 1797-1    | Dentistry - Shanks for rotary instruments - Part 1: Shanks made of metals                                                                                                                                                                                             | 2017-09    | <input checked="" type="checkbox"/> |
| 38 | SN EN ISO | 6872      | Dentistry - Ceramic materials                                                                                                                                                                                                                                         | 2015-09    | <input type="checkbox"/>            |
| 39 | ISO       | 5832-3    | Implants for surgery - Metallic materials - Part 3: Wrought titanium 6-aluminium 4-vanadium alloy                                                                                                                                                                     | 2016-10    | <input checked="" type="checkbox"/> |

## 3. Standards and norms for testing medical devices

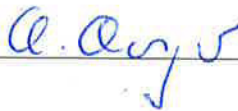
| No | Code      | No       | Title                                                                                                                                | Issue date         | Valid                               |
|----|-----------|----------|--------------------------------------------------------------------------------------------------------------------------------------|--------------------|-------------------------------------|
| 1  | ASTM      | D 638    | Standard Test Method for Tensile Properties of Plastics                                                                              | 2014               | <input type="checkbox"/>            |
| 2  | ASTM      | D 790-17 | Standard Test Methods for Flexural Properties of Unreinforced and Reinforced Plastics and Electrical Insulating Materials            | 2017               | <input type="checkbox"/>            |
| 3  | ASTM      | B 265-15 | Standard Specification for Titanium and Titanium Alloy Strip, Sheet, and Plate                                                       | 2015               | <input type="checkbox"/>            |
| 4  | DIN       | 51004    | Thermal analysis; determination of melting temperatures of crystalline materials by differential thermal analysis                    | 1994-06            | <input type="checkbox"/>            |
| 5  | DIN       | 51045-1  | Determination of the thermal expansion of solids - Part 1: Basic rules (WAK)                                                         | 2005-08            | <input type="checkbox"/>            |
| 6  | ISO       | 6892-1   | Metallic materials - Tensile testing - Part 1: Method of test at room temperature                                                    | 2016-06            | <input type="checkbox"/>            |
| 7  | ISO       | 10271    | Dentistry - Corrosion test methods for metallic materials (edition 2)                                                                | 2011-08            | <input type="checkbox"/>            |
| 8  | ISO       | 6507-1   | Metallic materials - Vickers hardness test - Part 1: Test method (ISO 6507-1:2005)                                                   | 2018-01            | <input type="checkbox"/>            |
| 9  | ISO       | 7405     | Dentistry - Evaluation of biocompatibility of medical devices used in dentistry<br>AMD 1: Positive control material                  | 2008-12<br>2013-07 | <input checked="" type="checkbox"/> |
| 10 | ISO       | 7491     | Dental materials -- Determination of colour stability                                                                                | 2000-09            | <input type="checkbox"/>            |
| 11 | SN EN ISO | 10993-1  | Biological evaluation of medical devices - Part 1: Evaluation and testing<br>AC                                                      | 2010-03<br>2010-09 | <input checked="" type="checkbox"/> |
| 12 | SN EN ISO | 10993-3  | Biological evaluation of medical devices - Part 3: Tests for genotoxicity, carcinogenicity and reproductive toxicity                 | 2014-12            | <input type="checkbox"/>            |
| 13 | SN EN ISO | 10993-5  | Biological evaluation of medical devices -Part 5: Tests for in vitro cytotoxicity                                                    | 2009-11            | <input checked="" type="checkbox"/> |
| 14 | SN EN ISO | 10993-6  | Biological evaluation of medical devices. Tests for local effects after implantation                                                 | 2017-05            | <input type="checkbox"/>            |
| 15 | SN EN ISO | 10993-10 | Biological evaluation of medical devices - Part 10: Tests for irritation and delayed-type hypersensitivity                           | 2013-12            | <input checked="" type="checkbox"/> |
| 16 | SN EN ISO | 10993-11 | Biological evaluation of medical devices – Part 11: Tests for systemic toxicity                                                      | 2009-08            | <input type="checkbox"/>            |
| 17 | SN EN ISO | 10993-12 | Biological evaluation of medical devices - Part 12: Sample preparation and reference materials                                       | 2012-09            | <input type="checkbox"/>            |
| 18 | SN EN ISO | 10993-14 | Biological evaluation of medical devices - Part 14: Identification and quantification of degradation products from ceramics          | 2009-08            | <input type="checkbox"/>            |
| 19 | SN EN ISO | 10993-15 | Biological evaluation of medical devices - Part 15: Identification and quantification of degradation products from metals and alloys | 2009-11            | <input type="checkbox"/>            |
| 20 | SN EN ISO | 10993-18 | Biological evaluation of medical devices - Part 18: Chemical characterization of materials                                           | 2009-08            | <input type="checkbox"/>            |
| 21 | ISO       | 14801    | Dentistry - Implants - Dynamic loading test for endosseous dental implants                                                           | 2016-11            | <input type="checkbox"/>            |
| 22 | SN EN ISO | 20795-1  | Dentistry - Base polymers, Part 1: Denture base polymers                                                                             | 2013-05            | <input type="checkbox"/>            |
| 23 | SN EN ISO | 20795-2  | Dentistry - Base polymers - Part 2: Orthodontic base polymers                                                                        | 2013-05            | <input type="checkbox"/>            |

Validity for the project and/or the product is confirmed by x

**Approval**

Alain Denzer  
Clinical Affairs Manager

Datum: 6.3.2019

Visum: 

Remi Meier  
Head of Quality Management

Datum: 6.3.2019.

Visum: 

#### 4. Änderungskontrolle

Ist eine Schulung erforderlich, hat der Prozesseigner die betroffenen Personen zu Schulung. Die Schulung ist mittels FO 6.301 „Nachweis interne Schulung“ zu dokumentieren. Der Nachweis ist QMA zur Archivierung abzugeben.

Die Schulungen sind rasch möglichst durchzuführen da sonst die Änderung nicht durchgesetzt werden kann.

#### 5. Änderungsprotokoll

| Änderungs-Datum | Seite | Pos./ Punkt | Beschreibung der Änderung                                                                                                              | Begründung der Änderung.                                                                                                                                        | Antragsteller | Schulung notwendig <sup>1)</sup><br>Schulung nicht notwendig <sup>2)</sup> |
|-----------------|-------|-------------|----------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------|----------------------------------------------------------------------------|
| 13.10.2017      | alle  | Alle        | Neugestaltung des Formulars im Excel für einfachere Verwaltung und Sortierung                                                          | Neugestaltung des Formulars im Excel für einfachere Verwaltung und Sortierung<br>Änderung ohne Schulungsbedarf, da Anwender an der Verbesserung beteiligt waren | ble           | <input type="checkbox"/> <input checked="" type="checkbox"/>               |
| 13.10.2017      | Alle  | alle        | Einfügen der neuen Versionen von:<br>- ISO 22674:2016-01<br>- ISO 1797-1:2017-05<br>- SN EN ISO 10993-6:2017-05<br>- ISO 14801:2016-11 | Generelle Aktualisierung ohne Schulungsbedarf                                                                                                                   | ble           | <input type="checkbox"/> <input checked="" type="checkbox"/>               |
| 13.10.2018      | N/A   | N/A         | Versionsänderung von 03 auf 04 wurde nicht dokumentiert                                                                                | N/A                                                                                                                                                             | N/A           | <input type="checkbox"/> <input checked="" type="checkbox"/>               |
| 13.12.2018      | alle  | alle        | Normenaktualisierung<br>Layoutwechsel auf Word                                                                                         | Generelle Aktualisierung ohne Schulungsbedarf                                                                                                                   | jka           | <input type="checkbox"/> <input checked="" type="checkbox"/>               |

<sup>1)</sup> Datum der durchgeführten Schulung:

<sup>2)</sup> Falls keine Schulung notwendig ist, bitte im Änderungsprotokoll begründen.

## Non-significant changes (MDCG 2020-3) affecting the content of the Declarations of Conformity (DoC) issued under EU MDD 93/42/EEC

| Change number – Description | Affected content                                                                                                                                                                                                                |
|-----------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| MD5-3639<br>EU-REP change   | <b>Previous EU-REP:</b> Cendres+Métaux France SAS, Les petites Buffeteries,<br>49124 St-Barthélémy d'Anjou, France<br><b>New EU-REP:</b> QualRep Services B.V., Utrechtseweg 310 – Bldg B42, 6812<br>AR Arnhem, The Netherlands |

Biel/Bienne, 11.10.2023

A blue ink signature of Dylan Moretti.

Dylan Moretti  
Regulatory Affairs Manager \*  
Cendres+Métaux SA

A blue ink signature of Matthias Walther.

Matthias Walther  
Director of Quality & Regulatory  
Cendres+Métaux SA

*\* Person responsible for compliance with the regulatory requirements MDR (EU) 2017/745 in accordance with Article 15.*