

**Konformitätserklärung
Declaration of Conformity**

Wir

We

**B. Braun Melsungen AG
Carl-Braun-Str. 1
34212 Melsungen
Deutschland/Germany
SRN DE-MF-000000201**erklären in eigener Verantwortung,
dass die Produkte**Certodyn® Universaladapter
Certodyn® Universaladapter Paed**EKG Positionskontrolle für Zentrale Venenkatheter
(Artikelnummern und Basic UDI-DI siehe Anlage I)mit den Anforderungen der Medizinprodukte
Verordnung (EU) 2017/745 übereinstimmen**Konformitätsbewertungsverfahren**
nach Artikel 52 Absatz 7 der oben genannten
Verordnung**Klassifizierung**
gemäß Anhang VIII der oben genannten
Verordnung
Klasse I**Gültig bis 2024-10-21**
gemäß gültigem ISO Zertifikat
Q5 012974 0590hereby declare in our own responsibility
that the products**Certodyn® Universal Adapter
Certodyn® Universal Adapter Paed**Central Venous Catheter ECG Positioning Control
(article numbers and Basic UDI-DI see attachment I)are in conformity with the requirements of the
Medical Device Regulation (EU) 2017/745**Conformity Assessment Procedure**
according to article 52 section 7
of the Regulation named above**Classification**
according to annex VIII of the Regulation named
above
Class I**Valid until 2024-10-21**
according to our valid ISO Certificate
Q5 012974 0590

Anlage I / Attachment I**Basic UDI-DI 403923900000219ZT**

Art.-Nr. / Art. No.	Produktname / Product name	Klasse / Class
4150228	Certodyn® Universal Adapter	I
4150724	Certodyn® Universal Adapter Paed	I

Document amendment information

Version	Description of the changes
4.0	Addition of Certificate Number Q5 012974 0590
3.0	Adaption of Validity
2.0	Adaption of Validity
1.0	First issue of DoC acc. to MDR replacing DoC acc. to MDD, document no.: 130-003CE, version 3.0

Title: Declaration of Conformity - 130-003-MDR - Certodyn Universal Adapter Initiator: Sandra ? Staufenberg

This document is signed electronically in compliance with the B. Braun electronic signature policies and procedures by following persons:

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