

**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 das die Produkte**Contiplex® C,
Contiplex® C NRFit®**Katheterset für kontinuierliche periphere
NervenblockadenBasis UDI-DI: 40392390000085632
(Artikelnummern siehe Anlage I)mit den Anforderungen der Medizinprodukte
Verordnung (EU) 2017/745 übereinstimmen**Konformitätsbewertungsverfahren**
nach Anhang IX
der oben genannten Verordnung**Klassifizierung**
gemäß Anhang VIII der oben genannten
Verordnung
Klasse IIa**Benannte Stelle**
TÜV SÜD Product Service GmbH
Kennnummer 0123**Gültig bis**
gemäß gültigem EU Zertifikat
(G10 012974 0611)hereby declare in our own responsibility
that the products**Contiplex® C,
Contiplex® C NRFit®**

Continuous peripheral nerve block catheter set

Basic UDI-DI: 40392390000085632
(article numbers see attachment I)are in conformity with the requirements of the
Medical Device Regulation (EU) 2017/745**Conformity Assessment Procedure**
according to annex IX
of the Regulation named above**Classification**
according to annex VIII of the Regulation named
above
Class IIa**Notified Body**
TÜV SÜD Product Service GmbH
Identification number 0123**Valid until**
according to our valid EU Certificate
(G10 012974 0611)

Anlage I / Attachment I**Basic UDI-DI 403923900000085632**

Art.-Nr. / Art. No.	Produktname / Product name	Klasse / Class
4898115	Contiplex® C	Ila
4898115NR	Contiplex® C NRFit®	Ila
4898130	Contiplex® C	Ila
4898130NR	Contiplex® C NRFit®	Ila

Document amendment information

Version	Description of the changes
1.0	Initial Version under 2017/745 MDR

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