

EG-KONFORMITÄTSERKLÄRUNG gemäß Richtlinie 93/42/EWG
EC Declaration of Conformity acc. to Directive 93/42/EEC

Wir, P. J. Dahlhausen & Co. GmbH, Emil-Hoffmann-Str. 53, 50996 Köln, erklären hiermit eigenverantwortlich, dass unsere nachfolgend genannten Medizinprodukte den Anforderungen der Richtlinie 93/42/EWG über Medizinprodukte entsprechen:

We, P. J. Dahlhausen & Co. GmbH, Emil-Hoffmann-Str. 53, 50996 Cologne, hereby declare under our sole responsibility that the medical devices described hereafter are in conformity with the provisions of the Directive 93/42/EEC on medical devices:

Produkt: <i>Product:</i>	Sauerstoffmaske mit Reservoirbeutel und Sicherheitsverbindungs-schlauch <i>Oxygen mask with reservoir bag and Sure Flow Tubing</i>
REF:	01.000.08.130 – 01.000.08.132
Verfahren gemäß RL 93/42/EWG: <i>Procedure acc. to MDD 93/42/EEC:</i>	Anhang II ohne (4) <i>Annex II excluding (4)</i>

Die Erklärung basiert auf dem EG-Zertifikat, das von der TÜV SÜD Product Service GmbH, Ridlerstr. 65, 80339 München (Benannte Stelle Nr. 0123), in Übereinstimmung mit der Richtlinie 93/42/EWG über Medizinprodukte ausgestellt wurde (Zertifikat-Nr. G1 015692 0502 Rev. 01).

The declaration is based on the EC-Certificate issued by the TÜV Süd Product Service GmbH, Ridlerstr. 65, 80339 Munich (notified body no. 0123), in accordance with the Council Directive 93/42/EEC concerning medical devices (Certificate No. G1 015692 0502 Rev. 01).

Unser Qualitätsmanagementsystem entspricht den Anforderungen der DIN EN ISO 13485:2016 und ist von einer akkreditierten Zertifizierstelle (TÜV SÜD Product Service GmbH) zertifiziert.

Our Quality Management System meets the requirements of the DIN EN ISO 13485:2016 and is certified by an accredited certification body (TÜV SÜD Product Service GmbH).

Hiermit erklären wir, dass wir für die Erstellung dieser EG-Konformitätserklärung die alleinige Verantwortung tragen.

We hereby declare that we are solely responsible for the preparation of this EC Declaration of Conformity.

Diese Erklärung ist gültig bis zum 26.05.2024. / *This declaration is valid until 26.05.2024.*

Köln / Cologne, 19.04.2021

P. J. Dahlhausen & Co. GmbH



Petra Hardt
Leitung QM/RA
Manager QM/RA



**Add value.
Inspire trust.**

TÜV SÜD Product Service GmbH · Ridlerstr. 65 · 80339 Munich · Germany

P. J. Dahlhausen & Co. GmbH
Adam-Riese-Straße 4
50996 Köln

Your reference/letter of	Our reference/name	Tel. extension/Email	Fax extension	Date	Page
15692	713347895, 713347896, 713347898	medical_devices@tuvsud.com	/	2024-09-25	1 of 8

**TÜV SÜD Product Service GmbH
Confirmation Letter
CL 015692 0508 Rev. 00**

Reference: 713347895 | 713347896 | 713347898

To whom it may concern,

Confirmation of the status of a formal application, written agreement, and appropriate surveillance in the framework of Regulation EU 2023/607 amending Regulations (EU) 2017/745 (in the following referenced as MDR) as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices.

With this letter TÜV SÜD Product Service GmbH, designated under MDR and identified by the number 0123 on NANDO, confirms that we have received a formal application in accordance with Section 4.3, first subparagraph of Annex VII of MDR and has signed a written agreement in accordance with Section 4.3, second subparagraph of Annex VII of MDR with the above stated manufacturer with the following SRN Number:

SRN Number: DE-MF-000006357

The devices covered by the formal application and the written agreement mentioned above are identified in the Tables below.

- Table 1 identifies the devices for which an MDR application has been received, written agreement concluded and for which TÜV SÜD Product Service GmbH is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive.
- Table 2 identifies the devices for which an MDR application has been received and a written agreement concluded, but TÜV SÜD Product Service GmbH has not yet taken the responsibility for appropriate surveillance of the corresponding devices under the applicable Directive.

Registered Office: Munich
Trade Register Munich HRB 85742
UniCredit Bank GmbH · BIC HYVEDEMMXXX
IBAN DE13 7002 0270 0048 8522 11
VAT ID No. DE129484267
Information pursuant to § 2 [1] DL-InfoV
(Germany) at tuvsud.com/imprint

Supervisory Board:
Holger Lindner (Chairman)
Board of Management:
Walter Reithmaier (CEO)
Patrick van Welij

TÜV SÜD Product Service GmbH
Zertifizierstelle für Medizinprodukte /
Certification Body for Medical Products
Ridlerstr. 65
80339 Munich
Germany

tuvsud.com/ps
Hotline: +49 89 50084-747





If devices covered by certificates issued under Directive 90/385/EEC (AIMDD) or Directive 93/42/EEC (MDD) that expired after 26 May 2021 and before 20 March 2023, without having been withdrawn, this letter also confirms that

- the manufacturer signed the written agreement under MDR by the date of MDD/AIMDD certificate expiry; or
- provided evidence that a competent authority of a Member State had granted a derogation or exemption from the applicable conformity assessment procedure in accordance with Article 59(1) of MDR or Article 97(1) of the MDR respectively.

The transition timelines in accordance Article 120 (3) of MDR that apply to the devices covered by this letter, subject to the manufacturer's continued compliance to the other conditions specified in Article 120 (3c) of MDR, are shown below:

- 26 May 2026 for Class III custom-made implantable devices
- 31 December 2027 for Class III devices and Class IIb implantable devices (except sutures, staples, dental fillings, dental braces, tooth crowns, screws, wedges, plates, wires, pins, clips and connectors)
- 31 December 2028 for other Class IIb devices, Class IIa, Class I devices placed on the market in sterile condition, measuring function
- 31 December 2028 for devices not requiring the involvement of a notified body under MDD but requiring it under MDR (e.g., class I devices that qualify as re-usable surgical instruments)

We reserve the right to invoice any issuance, copies, amendments and / or changes of the confirmation letter according to effort.

For confirmation letter validity see www.tuvsud.com/ps-cert?q=cert:CL 015692 0508 Rev. 00

In case of inquiries please contact medical_devices@tuvsud.com.

On behalf of the Notified Body TÜV SÜD Product Service GmbH,
2024-09-25

TÜV SÜD Product Service GmbH
Medical and Health Services

A handwritten signature in blue ink, appearing to be 'T. Alt'.

Torsten Alt
Conformity Assessment Responsible (CARE)

TÜV SÜD Product Service GmbH
Medical and Health Services

A handwritten signature in blue ink, appearing to be 'A. Fazlija'.

Arianit Fazlija
Application Reviewer



Table 1: Devices covered by this letter and for which TÜV SÜD Product Service GmbH is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
34342Neutralelektrode3D	☒ Class IIb / Class IIb implantable (exempted)	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342SillikonballonND	☒ Class IIb implantable (non-exempted)	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342AbsaugkatheterWU	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G2 015692 0503 Rev. 00 NB# 0123
34342AbsaugsystemG5	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev.01 NB# 0123
34342ASchlauchSQV	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342BeatmungsbGF	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342BeatmungsbH9	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342BezugLampeEZ	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342CSchuheSG3	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342DarmrohrXL	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342DrainageUD	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342Einnahmeglas7T	☒ Class I devices with measuring function	☒ N/A	☒ Certification as follows: Certificate # G3M 015692 0506 Rev. 00 NB# 0123
34342ElektrodenreinigerLF	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
34342EmbolekkVQ	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342Endotrachealtubus34	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342Endotrachfuehrung89	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342FadenMS4V	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342FadenziehSA3	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342Filter79	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342FrazierHY	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342GasinsufflationWF	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342GasprobensQA	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342GefaessA8	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342GuedeltubeN4	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342HautklammergeraetCK	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G2 015692 0503 Rev. 00 NB# 0123
34342HautstanzeY3	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342InfusionsWX	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
34342IrrigationsSTB	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342KatheterapliHT	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342KlammerentMM	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342KuenstlicheNase9P	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342KVerschlussPF	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342LaryngealmaskeMT	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342LatexballonkHE	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342Nabelk3F	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342OrthosaugerMG	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342PEEPVentilYB	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342RDrainage7M	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342Redon6P	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342Reservoir2J	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342SauerstoffkF7	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
34342Sauerstoffver3G	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342SaugAE	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342SBeutel92	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342SetK9H	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342SetM9M	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342SetS9Z	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342SilikonMaskeMY	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342SkalpelleSL	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342SkalpellklingeMZ	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342SMaskeZQ	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342SondeD9Y	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342SondeMAJ	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342SpuelsFA	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342Stuhldraining73	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
34342SUltraschallK7	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342SVerschlussX7	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342TKatheterD7	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342TrachealsQ8	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342TSbeutelGV	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342TSystemNY	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342UBbeutelA4	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev. 01 NB# 0123
34342UBbeutelMNL	☒ Class I devices in sterile condition ☒ Class I devices with measuring function	☒ N/A	☒ Certification as follows: Certificate #; G2MS 015692 0505 Rev. 00 NB# 0123
34342UrokatheterKD	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342VBezugSAT	☒ Class I devices in sterile condition	☒ N/A	☒ Certification as follows: Certificate # G2S 015692 0504 Rev.01 NB# 0123
34342VerneblerSW8	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123
34342Wenditubus42	☒ Class IIa	☒ N/A	☒ Certification as follows: Certificate # G1 015692 0502 Rev. 01 NB# 0123



Table 2: Devices covered by this letter and for which TÜV SÜD Product Service GmbH is NOT responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
34342LaryngoskopC8	☒ Class I reusable surgical instruments	☒ N/A	☒ N/A - Device did not require a Notified Body certificate under Directives

Confirmation Letter Version History

Date	TÜV SÜD Product Service GmbH internal reference traceable to each version of the letter	Action
2024/09/25	713347895, 713347896, 713347898	Initial issue