



# ECHIPAMED

## P L U S

Moldova, MD - 2001, Chisinau, str. Valea Trandafirilor 24 "B", of. 2-7

tel. +373 (22) 234 349, 234 225; fax +373 (22) 234 225

e-mail: [office@echipamed.com](mailto:office@echipamed.com), [info@echipamed.com](mailto:info@echipamed.com)

Nr. F/N

din 17.09.2023

*Către Agenția Medicamentului  
și Dispozitivelor Medicale*

### Notificare

**pentru înregistrarea dispozitivelor medicale în Registrul de stat  
al dispozitivelor medicale**

Solicitantul **„Echipamed-Plus” SRL**, cu sediul str. Valea Trandafirilor 24B, of.80, MD-2001, mun. Chișinău, Republica Moldova, tel./fax: 022 23-42-25, e-mail: office@echipamed.com, solicit înregistrarea în Registrul de stat al dispozitivelor medicale a următoarelor categorii și tipuri de dispozitive medicale pentru introducerea și punerea la dispoziție pe piață a:

| Nr | Nr. Cat. | Denumire        | Denumire comercială | Model          |
|----|----------|-----------------|---------------------|----------------|
| 1  | MP01840  | Catheter Mounts | ErgoStar            | ErgoStar CM 40 |

Se anexează următoarele acte:

1. Declarații de conformitate CE.
2. Declarații de la producător privind desemnarea reprezentantului.
3. Declarație pe proprie răspundere.

Data 17.09.2023

Semnătura \_\_\_\_\_

### Tabelul de recepționare a notificării

(se completează de către Agenție în momentul depunerii notificării de către solicitant)

|                                                                                                 |  |
|-------------------------------------------------------------------------------------------------|--|
| Comentarii cu privire la acceptul/refuzul recepționării notificării, inclusiv motivul refuzului |  |
| Data/nr. de ordine atribuit notificării de către Agenție (în cazul acceptării recepționării)    |  |
| Numele, prenumele, funcția persoanei responsabile de recepționarea dosarului                    |  |
| Semnătura persoanei responsabile                                                                |  |



# ECHIPAMED

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Nr. F/N

din 17.09.2023

*Către Agenția Medicamentului  
și Dispozitivelor Medicale*

### Declarație pe proprie răspundere

Solicitant: **„Echipamed-Plus” SRL**, cu sediul str. Valea Trandafirilor 24B, of.80, MD-2001, mun. Chișinău, Republica Moldova, tel./fax: 022 23-42-25, e-mail: office@echipamed.com, declar pe proprie răspundere, cunoscând prevederile art. 352<sup>1</sup>, Codul Penal al Republicii Moldova cu privire la falsul în declarații, că documentele și datele furnizate pentru notificare dispozitivului medical:

| Nr | Nr. Cat. | Denumire        | Denumire comercială | Model          |
|----|----------|-----------------|---------------------|----------------|
| 1  | MP01840  | Catheter Mounts | ErgoStar            | ErgoStar CM 40 |

**Sunt autentice și corespund realității.**

Director Valeriu Iurchevici

Semnătura \_\_\_\_\_

| Nr. | Numărul de<br>catalog (referință) | Denumire generică (denumirea dispozitivului) | Denumire comercială<br>(brand) | Modelul        | Cod<br>GMDN | Clasa<br>dispozitivului |
|-----|-----------------------------------|----------------------------------------------|--------------------------------|----------------|-------------|-------------------------|
| 1   | MP01840                           | Catheter Mounts                              | ErgoStar                       | ErgoStar CM 40 | 61346       | Ila                     |



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Zentralstelle der Länder  
für Gesundheitsschutz  
bei Arzneimitteln und  
Medizinprodukten  
www.zls.de  
BS-MDR-099



Product Service

## EU Quality Management System Certificate (MDR)

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapters I and III  
(Class IIa and Class IIb Devices)

**No. G10 010578 0039 Rev. 09**

**Manufacturer:**

**Drägerwerk AG & Co. KGaA**

Moislinger Allee 53-55  
23542 Lübeck  
GERMANY

**SRN Manufacturer:**

DE-MF-000005329

The Certification Body of TÜV SÜD Product Service GmbH certifies that the manufacturer has established, documented and implemented a quality management system as described in Article 10 (9) of the Regulation (EU) 2017/745 on medical devices. Details on device categories covered by the quality management system are described on the following page(s). The Report referenced below summarises the result of the assessment and includes reference to relevant CS, harmonized standards and test reports. The conformity assessment has been carried out according to Annex IX Chapter I and III of this regulation with a positive result. The quality management system assessment was accompanied by the assessment of technical documentation for devices selected on a representative basis. The certified quality management system is subject to periodical surveillance by TÜV SÜD Product Service GmbH. The surveillance assessment shall also include an assessment of the technical documentation for the device or devices concerned on the basis of further representative samples. All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: [www.tuvsud.com/ps-cert?q=cert:G10\\_010578\\_0039\\_Rev.\\_09](http://www.tuvsud.com/ps-cert?q=cert:G10_010578_0039_Rev._09)

**Report No.:**

713253108\_CN

**Preceding Certificate No.:**

G10 010578 0039 Rev. 08

**Valid from:**

2023-03-14

**Valid until:**

2025-03-17

**Date of Initial Issuance:**

2020-03-18

**Issue date:**

2023-03-14

Christoph Dicks

Head of Certification/Notified  
Body





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Product Service

## EU Quality Management System Certificate (MDR)

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapters I and III  
(Class IIa and Class IIb Devices)

**No. G10 010578 0039 Rev. 09**

|                          |                                                                                                                                                                                                                                                                                                                                          |
|--------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <b>Classification:</b>   | Class IIa                                                                                                                                                                                                                                                                                                                                |
| <b>Device Group:</b>     | R02 - BREATHING CIRCUITS AND CATHETER MOUNTS<br>R0301 - RESPIRATORY MASKS<br>R030201 - VENTILATION BALLOONS<br>R0401 - VENTILATION FILTERS<br>R0402 - NATURAL BREATHING FILTERS<br>Z120301 - ANAESTHESIA AND PULMONARY VENTILATION<br>SUPPORT INSTRUMENTS<br>Z120309 - MEDICAL/MEDICINAL GAS PIPELINE SYSTEMS AND<br>RELATED ACCESSORIES |
| <b>Intended Purpose:</b> | -                                                                                                                                                                                                                                                                                                                                        |
| <b>Classification:</b>   | Class IIa                                                                                                                                                                                                                                                                                                                                |
| <b>Device Group:</b>     | Z12040192 - GENERAL MEDICINE DIAGNOSIS AND<br>MONITORING INSTRUMENTS - MEDICAL DEVICE SOFTWARE                                                                                                                                                                                                                                           |
| <b>Intended Purpose:</b> | -                                                                                                                                                                                                                                                                                                                                        |
| <b>Classification:</b>   | Class IIa                                                                                                                                                                                                                                                                                                                                |
| <b>Device Group:</b>     | Z120302 - VITAL SIGNS MONITORING INSTRUMENTS                                                                                                                                                                                                                                                                                             |
| <b>Intended Purpose:</b> | -                                                                                                                                                                                                                                                                                                                                        |
| <b>Classification:</b>   | Class IIa                                                                                                                                                                                                                                                                                                                                |
| <b>Device Group:</b>     | A060304 - INTRA-OPERATION FLUID COLLECTION DEVICES                                                                                                                                                                                                                                                                                       |
| <b>Intended Purpose:</b> | -                                                                                                                                                                                                                                                                                                                                        |
| <b>Classification:</b>   | Class IIb                                                                                                                                                                                                                                                                                                                                |
| <b>Device Group:</b>     | Z12040192 - GENERAL MEDICINE DIAGNOSIS AND<br>MONITORING INSTRUMENTS - MEDICAL DEVICE SOFTWARE                                                                                                                                                                                                                                           |
| <b>Intended Purpose:</b> | Software intended to provide clinical information for the purpose of<br>supporting patient management and the decision making process                                                                                                                                                                                                    |
| <b>Classification:</b>   | Class IIb                                                                                                                                                                                                                                                                                                                                |
| <b>Device Group:</b>     | Z120804 - NEONATOLOGY INSTRUMENTS                                                                                                                                                                                                                                                                                                        |



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Product Service

## EU Quality Management System Certificate (MDR)

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapters I and III  
(Class IIa and Class IIb Devices)

**No. G10 010578 0039 Rev. 09**

**Intended Purpose:** Warming therapy devices intended to provide controlled ambient conditions for premature babies and neonates in closed and open care therapy

**Classification:** Class IIb

**Device Group:** Z120301 - ANAESTHESIA AND PULMONARY VENTILATION SUPPORT INSTRUMENTS

**Intended Purpose:** Devices for the purpose of ventilation and/or anesthesia

**Classification:** Class IIb

**Device Group:** Z120309 - MEDICAL/MEDICINAL GAS PIPELINE SYSTEMS AND RELATED ACCESSORIES

**Intended Purpose:** Devices intended to distribute or supply gases, vacuum, electricity or data to equipment in diagnostic, therapy or surgery

**Classification:** Class IIb

**Device Group:** R020107 - THERMOREGULATED BREATHING CIRCUITS

**Intended Purpose:** Inspiratory (and expiratory) heated disposable breathing circuit for conducting humidified breathing gas from humidifier to patient

**Classification:** Class IIb

**Device Group:** R020101 - STANDARD BREATHING CIRCUITS

**Intended Purpose:** Devices intended to administer gases for the purpose of ventilation

The validity of this certificate depends on conditions and/or is limited to the following: ./.

### Revision History:

| Rev. | Dated      | Report    | Description |
|------|------------|-----------|-------------|
| 00   | 2020-03-18 | 713169482 | -           |
| 01   | 2021-07-02 | 713184148 | -           |
| 02   | 2021-09-30 | 713215188 | -           |
| 03   | 2021-10-01 | 713215832 | -           |
| 04   | 2021-10-04 | 713215842 | -           |
| 05   | 2021-10-04 | 713219421 | -           |
| 06   | 2021-11-22 | 713229134 | -           |

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TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

TÜV SÜD Product Service GmbH • Certification Body • Ridlerstraße 65 • 80339 Munich • Germany



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Product Service

## EU Quality Management System Certificate (MDR)

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapters I and III  
(Class IIa and Class IIb Devices)

**No. G10 010578 0039 Rev. 09**

|    |            |              |   |
|----|------------|--------------|---|
| 07 | 2022-02-21 | 713213004    | - |
| 08 | 2022-10-06 | 713225304_CN | - |
| 09 | 2023-03-14 | 713253108_CN |   |

Supplemented: Device(s)/group of  
device(s) added

Drägerwerk AG & Co. KGaA, 23542 Lübeck, Germany

To whom it may concern

### Manufacturer's Authorization

May 25, 2023

We, Drägerwerk AG & Co. KGaA, Moislinger Allee 53-55, 23558 Lübeck, Germany, who is an established and reputable manufacturer of medical equipment, having factories at Lübeck (Germany), Telford (United States), Andover (United States) and Shanghai (China), do hereby declare that

**"Echipamed-Plus" SRL Valea Trandafirilor 24 "B", of. 2-7 MD-2001, Chisinau Republic of Moldova**

is our official distributor and local representative for Anesthesia Devices, Incubators, Ventilation Machines, Monitoring, Pendant and Operating Lights of Drägerwerk AG & Co. KGaA in the territory of the Republic of Moldova.

We declare that only above mentioned Company is authorized to do registration, quote, sell, subsequently negotiate and sign contracts, as well as to perform installation and after sales Service of Anesthesia Devices, Incubators, Ventilation Machines, Monitoring, Pendant and Operating Lights manufactured by us in their own name and on their own account.

This authorization letter will remain valid until 31.12.2024.

Martin Koch  
Managing Director Sub Region East Europe



Drägerwerk AG & Co. KGaA  
Moislinger Allee 53-55  
23558 Lübeck, Germany  
Postal address:  
23542 Lübeck, Germany  
Tel. +49 451 882-0  
Fax +49 451 882-2080  
info@draeger.com  
www.draeger.com  
UID-Nr. DE135082211

Bank details:  
Commerzbank AG, Lübeck  
IBAN: DE95 2304 0022 0014 6795 00  
Swift-Code: COBA DE FF 230  
Sparkasse zu Lübeck  
IBAN: DE15 2305 0101 0001 0711 17  
Swift-Code: NOLADE21SPL

Registered office: Lübeck  
Commercial register:  
Local court Lübeck HRB 7903 HL  
General partner:  
Drägerwerk Verwaltungs AG  
Registered office: Lübeck  
Commercial register:  
Local court Lübeck HRB 7395 HL

Chairman of the Supervisory Board  
for Drägerwerk AG & Co. KGaA  
and Drägerwerk Verwaltungs AG:  
Stefan Lauer  
Executive Board:  
Stefan Dräger (chairman)  
Rainer Klug  
Gert-Hartwig Lescow  
Dr. Reiner Piske  
Anton Schrofner

CERTIFICAT

CERTIFICADO

CERTИФИКАТ

認證證書

CERTIFICATE

ZERTIFIKAT



Management Service

# CERTIFICATE

The Certification Body  
of TÜV SÜD Management Service GmbH  
certifies that

## Dräger

**Drägerwerk AG & Co. KGaA**  
Moislinger Allee 53-55, 23542 Lübeck  
Germany

for the Scope of application

Design and development, production and distribution of  
diagnostic and therapeutic medical devices and installations  
as well as consulting and services  
in the field of medical technology

**Revalstraße 1, 23560 Lübeck**  
Germany

for the Scope of application

**Production and distribution of diagnostic  
and therapeutic medical devices and installations**

has established and applies  
a Quality Management System.

An audit was performed, Order No. **707037695**.

Proof has been furnished that the requirements according to

### ISO 9001:2015

are fulfilled.

The certificate is valid from **2021-01-15** until **2024-01-14**.

Certificate Registration No.: **12 100 49423 TMS**.

Head of Certification Body  
Munich, 2021-01-13





# Certificate

**No. Q5 010578 0031 Rev. 01**

**Holder of Certificate:** Drägerwerk AG & Co. KGaA

Moislinger Allee 53-55  
23542 Lübeck  
GERMANY

**Certification Mark:**



**Scope of Certificate:** Design, Development, Manufacture and Distribution of Diagnostic and Therapeutic Medical Devices and Installations as well as Consulting and Services in the Field of Medical Technology. Diagnostic and Therapeutic Medical Devices and Installations: Anaesthetic Equipment, Infusion Equipment, Pediatric Equipment for Warming- and Photo-Therapy, Lung Ventilator Equipment, Monitoring Equipment, Clinical Decision Support Software, Patient Data Management Software, Equipment for Suction, Breathing-, Inhalation-, O2- and Aerosol-Therapy, Medical Gas Management and Supply Systems as well as Medical Lights

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: [www.tuvsud.com/ps-cert?q=cert:Q5 010578 0031 Rev. 01](http://www.tuvsud.com/ps-cert?q=cert:Q5 010578 0031 Rev. 01)

**Report No.:** 713193628  
**Valid from:** 2021-01-18  
**Valid until:** 2024-01-13

Date, 2021-01-18

C.D.M

Christoph Dicks  
Head of Certification/Notified Body



# Certificate

No. Q5 010578 0031 Rev. 01

**Applied Standard(s):** EN ISO 13485:2016  
Medical devices - Quality management systems -  
Requirements for regulatory purposes  
(ISO 13485:2016)  
DIN EN ISO 13485:2016

**Facility(ies):** Drägerwerk AG & Co. KGaA  
Mölsinger Allee 53-55, 23542 Lübeck, GERMANY

Design, Development, Manufacture and Distribution of  
Diagnostic and Therapeutic Medical Devices and Installations  
as well as Consulting and Services in the Field of Medical  
Technology. Diagnostic and Therapeutic Medical Devices and  
Installations: Anaesthetic Equipment, Infusion Equipment,  
Pediatric Equipment for Warming- and Photo-Therapy, Lung  
Ventilator Equipment, Monitoring Equipment, Clinical  
Decision Support Software, Patient Data Management  
Software, Equipment for Suction, Breathing-, Inhalation-,  
O2- and Aerosol-Therapy, Medical Gas Management  
and Supply Systems as well as Medical Lights

Drägerwerk AG & Co. KGaA  
Revalstraße 1, 23560 Lübeck, GERMANY

Manufacture and Distribution of Diagnostic and  
Therapeutic Medical Devices and Installations as well  
as Consulting and Services in the Field of Medical  
Technology. Diagnostic and Therapeutic Medical Devices and  
Installations: Anaesthetic Equipment, Infusion Equipment,  
Pediatric Equipment for Warming- and Photo-Therapy, Lung  
Ventilator Equipment, Monitoring Equipment, Clinical  
Decision Support Software, Patient Data Management  
Software, Equipment for Suction, Breathing-, Inhalation-,  
O2- and Aerosol-Therapy, Medical Gas Management  
and Supply Systems as well as Medical Lights

./.





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Product Service

## EC Certificate

Full Quality Assurance System

Directive 93/42/EEC on Medical Devices (MDD), Annex II excluding (4)  
(Devices in Class IIa, IIb or III)

**No. G1 010578 0037 Rev. 01**

### Manufacturer:

**Drägerwerk AG & Co. KGaA**

Moislinger Allee 53-55  
23542 Lübeck  
GERMANY

### Facility(ies):

Drägerwerk AG & Co. KGaA  
Revalstraße 1, 23560 Lübeck, GERMANY

Drägerwerk AG & Co. KGaA  
Moislinger Allee 53-55, 23542 Lübeck, GERMANY

### Product Category(ies):

Anaesthetic equipment with standard accessories,  
Infusion equipment with standard accessories,  
Pediatric equipment with standard accessories,  
Lung ventilator equipment with standard accessories,  
Monitoring equipment with standard accessories,  
Equipment for suction, breathing-, inhalation-, oxygen-  
and aerosol-therapy with standard accessories,  
Medical supply units and terminal units for pressurized  
medical gases and vacuum,  
Pipelines for compressed medical gases and vacuum,  
Anaesthetic gas scavenging systems, Components for  
medical gas management systems, Software for diagnosis based on clinical  
data Incl. patient data, monitoring and device parameter, Visualization,  
diagnostic and therapeutic software for anesthesia and respiratory devices

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for design, manufacture and final inspection of the respective devices / device categories in accordance with MDD Annex II. This quality assurance system conforms to the requirements of this Directive and is subject to periodical surveillance. For marketing of class III devices an additional Annex II (4) certificate is mandatory. See also notes overleaf.

**Report No.:** 713162398

**Valid from:** 2020-01-15  
**Valid until:** 2024-05-26

**Date,** 2019-12-09

Christoph Dicks  
Head of Certification/Notified Body

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|                                        |                        |
|----------------------------------------|------------------------|
| Released                               | Document Status        |
| 2022-12-02T12:03:13                    | Released Date          |
| Please see associated Document Release | Author(s) and Approval |
| Reference: DR-00073321                 |                        |

en/de



EU Declaration of Conformity  
EU-Konformitätserklärung

Document No. / Dokument Nr.  
Date / Datum  
Place / Ort  
Page / Seite

MDR108-007-2212-029-0  
2022-12-02  
Germany - Lübeck  
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Drägerwerk AG & Co. KGaA  
Moislinger Allee 53-55  
23542 Lübeck  
Germany

Authorised representative /  
Europäischer Bevollmächtigter: N/A

EC Certificate: G10 010578 0039  
Valid until: 2025-03-17

Single registration number (SRN)/  
einmalige Registrierungsnummer: DE-MF-000005329

hereby declares under its sole responsibility that the /  
erklärt hiermit in alleiniger Verantwortung, dass

| Product Name / Produktbezeichnung | Device Category /<br>Produktkategorie | Device Class /<br>Gerätekategorie | UMDNS Code /<br>GMDN Code /<br>EMDN Code   |
|-----------------------------------|---------------------------------------|-----------------------------------|--------------------------------------------|
| ErgoStar                          | Catheter Mounts                       | Ila                               | UMDNS 14-080/<br>GMDN 61346/<br>EMDN R0202 |

meets the following provisions:  
mit den folgenden Bestimmungen übereinstimmt:

|                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |
|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <p>European regulation (EU) 2017/745 on medical devices. An examination of the quality management System has been carried out following Annex IX (Chapters I and III and section 4) of the regulation by the Notified Body:/<br/>Verordnung (EU) 2017/745 über Medizinprodukte. Eine Überprüfung des Qualitätsmanagementsystems, nach den Regeln wie in Anhang IX (Kapitel I und III und Abschnitt 4) der Verordnung beschrieben, wurde durch die Benannte Stelle vorgenommen:<br/><b>TÜV Süd Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany, NB 0123</b></p> |
| <p>The quality management system also complies to EN ISO 9001 and EN ISO 13485./<br/>Das Qualitätsmanagementsystem erfüllt weiterhin die Anforderungen gemäß EN ISO 9001 und EN ISO 13485.</p>                                                                                                                                                                                                                                                                                                                                                                              |

This declaration is effective for products placed on the market as of the date of issue. Any modifications of the device not authorized by Dräger will invalidate this declaration./  
Diese Erklärung ist gültig für ab dem Ausstellungsdatum in Verkehr gebrachte Produkte. Jede nicht durch Dräger autorisierte Modifikation an dem Produkt führt zur Ungültigkeit dieser Erklärung.

Drägerwerk AG & Co. KGaA  
Moislinger Allee 53-55  
23558 Lübeck, Germany  
Postal address:  
23542 Lübeck, Germany  
Tel +49 451 882-0  
Fax +49 451 882-2080  
info@draeger.com  
www.draeger.com  
VAT no. DE135082211

Bank details:  
Commerzbank AG, Lübeck  
IBAN: DE95 2304 0022 0014 6795 00  
Swift-Code: COBA DE FF 230  
Sparkasse zu Lübeck  
IBAN: DE15 2305 0101 0001 0711 17  
Swift-Code: NOLADE21SPL

Registered office: Lübeck  
Commercial register:  
Local court Lübeck HRB 7903 HL  
General partner: Drägerwerk  
Verwaltungs AG  
Registered office: Lübeck  
Commercial register:  
Local court Lübeck HRB 7395 HL

Chairman of the Supervisory Board for  
Drägerwerk AG & Co. KGaA and  
Drägerwerk Verwaltungs AG:  
Stefan Lauer  
Executive Board:  
Stefan Dräger (chairman)  
Rainer Klug  
Gert-Hartwig Lescow  
Dr. Reiner Piske  
Anton Schrofner



## EU Declaration of Conformity EU-Konformitätserklärung

Document No. / Dokument Nr.

Date / Datum

Place / Ort

Page / Seite

**MDR108-007-2212-029-0**

**2022-12-02**

**Germany - Lübeck**

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For the signature on behalf of Dräger see upper left corner of page 1./  
Für die Unterschrift im Namen von Dräger siehe linke obere Ecke der Seite 1.

Director  
Quality & Regulatory Affairs  
Business Unit Hospital Consumables & Accessories  
Medical Division

Timo Harms

Director  
Research & Development  
Business Unit Hospital Consumables & Accessories  
Medical Division

Kalle Heckmann



## EU Declaration of Conformity EU-Konformitätserklärung

Document No. / Dokument Nr.  
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2022-12-02  
Germany - Lübeck  
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Drägerwerk AG & Co. KGaA  
Moislinger Allee 53-55  
23542 Lübeck  
Germany

| Product Name / Produktbezeichnung                                                     | Device Category / Produktkategorie                                                                                                                 |
|---------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------|
| ErgoStar                                                                              | Catheter Mounts                                                                                                                                    |
| Applied Standards in full or in part / Vollständig oder teilweise angewendete Normen: |                                                                                                                                                    |
| IEC 60601-1:2005 +A1:2012+A2:2020                                                     | Medical electrical equipment – Part 1 General requirements for basic safety and essential performance                                              |
| EN 62366-1:2015+A1:2020<br>(IEC 62366-1: 2015 COR 1 2016 AMD1:2020)                   | Medical devices - Part 1: Application of usability engineering to medical devices                                                                  |
| EN 60601-1-6: 2010+A1:2015+A2:2021<br>(IEC 60601-1-6: 2010 AMD 1 2013 A2:2020)        | Medical electrical equipment - Part 1-6: General requirements for basic safety and essential performance - Collateral standard: Usability          |
| EN ISO 14971:2019<br>(ISO 14971 :2019)                                                | Medical devices – Application of risk management to medical devices                                                                                |
| ISO 15223-1:2021                                                                      | Medical devices – Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General requirements.          |
| EN ISO 80601-2-12:2020<br>(ISO 80601-2-12:2020)                                       | Medical electrical equipment - Part 2-12: Particular requirements for basic safety and essential performance of critical care ventilators          |
| EN ISO 80601-2-13:2012+A2:2019<br>(ISO 80601-2-13:2011+AMD1:2015 +AMD2:2018)          | Medical Electrical Equipment – Part 2-13: Particular Requirements for Basic Safety and Essential Performance of an Anaesthetic Workstation         |
| EN ISO 18562-1:2020<br>(ISO 18562-1:2017)                                             | Biocompatibility evaluation of breathing gas pathways in healthcare applications - Part 1: Evaluation and testing within a risk management Process |
| EN ISO 10993-1:2020<br>(ISO 10993-1:2018)                                             | Biological evaluation of medical devices -- Part 1: Evaluation and testing within a risk management process                                        |
| ISO 18190:2016                                                                        | Anaesthetic and respiratory equipment – General requirements for airways and related equipment                                                     |

**EU Declaration of Conformity**  
**EU-Konformitätserklärung****Document No. / Dokument Nr.****Date / Datum****Place / Ort****Page / Seite****MDR108-007-2212-029-0****2022-12-02****Germany - Lübeck****4 / 5**

|                                           |                                                                                                           |
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| EN ISO 5356-1:2015<br>(ISO 5356-1:2015)   | Anaesthetic and respiratory equipment; Conical connectors – Part 1:<br>Cones and sockets                  |
| EN ISO 80369-1:2018<br>(ISO 80369-1:2018) | Small-bore connectors for liquids and gases in healthcare<br>applications -- Part 1: General requirements |
| EN ISO 5367:2014<br>(ISO 5367:2014)       | Anaesthetic and respiratory equipment - Breathing sets and<br>connectors                                  |
| EN ISO 20417:2021                         | Medical devices - Information to be supplied by the manufacturer                                          |

**EU Declaration of Conformity**  
**EU-Konformitätserklärung**

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**Drägerwerk AG & Co. KGaA**  
**Moislinger Allee 53-55**  
**23542 Lübeck**  
**Germany**

| Extend of conformity assessment / Umfang der Konformitätsbewertung |                                   |                           |
|--------------------------------------------------------------------|-----------------------------------|---------------------------|
| Part Number /<br>Sachnummer                                        | Product Name / Produktbezeichnung | Basic UDI-DI              |
| MP01840                                                            | ErgoStar CM 40                    | 0404867512080076K19Z000ST |
| MP01845                                                            | ErgoStar CM 45                    | 0404867512080076K19Z000ST |
| MP01850                                                            | ErgoStar CM 50                    | 0404867512080076K19Z000ST |
| MP01855                                                            | ErgoStar CM 55                    | 0404867512080076K19T010RL |
| MP01860                                                            | ErgoStar CM 60                    | 0404867512080076K19Z000ST |
| MP01890                                                            | ErgoStar AC 90                    | 0404867512080076K19T020RP |
| MP01895                                                            | ErgoStar AC 95                    | 0404867512080076K19T020RP |



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**Drägerwerk AG & Co. KGaA**  
**Moislinger Allee 53-55**  
**23542 Lübeck**  
**Germany**

Reprezentant autorizat:

N/A

EC Certificate: G10 010578 0039

Valid until: 2025-03-17

Număr unic de înregistrare (SRN):

DE-MF-000005329

### declară prin prezenta pe proprie răspundere că

| Numele produsului | Categoria dispozitivului | Clasa dispozitivului | Codul UMDNS /<br>Codul GMDN /<br>Codul EMDN |
|-------------------|--------------------------|----------------------|---------------------------------------------|
| ErgoStar          | Catheter Mounts          | Ila                  | UMDNS 14-080/<br>GMDN 61346/<br>EMDN R0202  |

### îndeplinește următoarele cerințe:

REGULAMENTUL (UE) 2017/745 privind dispozitivele medicale. O analiză a sistemului de management al calității a fost efectuată conform Anexei IX (capitolele I și III și secțiunea 4) a reglementării Organismului notificat:

**TÜV Süd Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany, NB 0123**

Sistemul de management al calității îndeplinește de asemenea cerințele standardelor EN ISO 9001 și EN ISO 13485.

**Această declarație are efect pentru produsele puse pe piață începând cu data emiterii. Orice modificare a dispozitivului neautorizată de Dräger va anula această declarație.**

Aceasta este o traducere a documentului original (en/de) și din această cauză nu necesită o semnătură.

Director  
 Quality & Regulatory Affairs  
 Business Unit Hospital Consumables & Accessories  
 Medical Division

Director  
 Research & Development  
 Business Unit Hospital Consumables & Accessories  
 Medical Division

Timo Harms

Kalle Heckmann



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Drägerwerk AG & Co. KGaA  
Moislinger Allee 53-55  
23542 Lübeck  
Germany

| Numele produsului                                                              | Categoria dispozitivului                                                                                                                           |
|--------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------|
| ErgoStar                                                                       | Catheter Mounts                                                                                                                                    |
| Standarde aplicate în totalitate sau parțial:                                  |                                                                                                                                                    |
| IEC 60601-1:2005 +A1:2012+A2:2020                                              | Medical electrical equipment – Part 1 General requirements for basic safety and essential performance                                              |
| EN 62366-1:2015+A1:2020<br>(IEC 62366-1: 2015 COR 1 2016 AMD1:2020)            | Medical devices - Part 1: Application of usability engineering to medical devices                                                                  |
| EN 60601-1-6: 2010+A1:2015+A2:2021<br>(IEC 60601-1-6: 2010 AMD 1 2013 A2:2020) | Medical electrical equipment - Part 1-6: General requirements for basic safety and essential performance - Collateral standard: Usability          |
| EN ISO 14971:2019<br>(ISO 14971 :2019)                                         | Medical devices – Application of risk management to medical devices                                                                                |
| ISO 15223-1:2021                                                               | Medical devices – Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General requirements.          |
| EN ISO 80601-2-12:2020<br>(ISO 80601-2-12:2020)                                | Medical electrical equipment - Part 2-12: Particular requirements for basic safety and essential performance of critical care ventilators          |
| EN ISO 80601-2-13:2012+A2:2019<br>(ISO 80601-2-13:2011+AMD1:2015 +AMD2:2018)   | Medical Electrical Equipment – Part 2-13: Particular Requirements for Basic Safety and Essential Performance of an Anaesthetic Workstation         |
| EN ISO 18562-1:2020<br>(ISO 18562-1:2017)                                      | Biocompatibility evaluation of breathing gas pathways in healthcare applications - Part 1: Evaluation and testing within a risk management Process |
| EN ISO 10993-1:2020<br>(ISO 10993-1:2018)                                      | Biological evaluation of medical devices -- Part 1: Evaluation and testing within a risk management process                                        |
| ISO 18190:2016                                                                 | Anaesthetic and respiratory equipment – General requirements for airways and related equipment                                                     |
| EN ISO 5356-1:2015<br>(ISO 5356-1:2015)                                        | Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets                                                              |



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|                                           |                                                                                                        |
|-------------------------------------------|--------------------------------------------------------------------------------------------------------|
| EN ISO 80369-1:2018<br>(ISO 80369-1:2018) | Small-bore connectors for liquids and gases in healthcare applications -- Part 1: General requirements |
| EN ISO 5367:2014<br>(ISO 5367:2014)       | Anaesthetic and respiratory equipment - Breathing sets and connectors                                  |
| EN ISO 20417:2021                         | Medical devices - Information to be supplied by the manufacturer                                       |



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Drägerwerk AG & Co. KGaA  
Moislinger Allee 53-55  
23542 Lübeck  
Germany

| Evaluarea extinsă a conformității |                   |                           |
|-----------------------------------|-------------------|---------------------------|
| Cod articol                       | Numele produsului | UDI-DI de bază            |
| MP01840                           | ErgoStar CM 40    | 0404867512080076K19Z000ST |
| MP01845                           | ErgoStar CM 45    | 0404867512080076K19Z000ST |
| MP01850                           | ErgoStar CM 50    | 0404867512080076K19Z000ST |
| MP01855                           | ErgoStar CM 55    | 0404867512080076K19T010RL |
| MP01860                           | ErgoStar CM 60    | 0404867512080076K19Z000ST |
| MP01890                           | ErgoStar AC 90    | 0404867512080076K19T020RP |
| MP01895                           | ErgoStar AC 95    | 0404867512080076K19T020RP |