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en/de



EU Declaration of Conformity
EU-Konformitätserklärung

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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 / Produktkategorie	Device Class / Gerätekategorie	UMDNS Code / GMDN Code / EMDN Code
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

meets the following provisions:
mit den folgenden Bestimmungen übereinstimmt:

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:/ 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: TÜV Süd Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany, NB 0123
The quality management system also complies to EN ISO 9001 and EN ISO 13485./ Das Qualitätsmanagementsystem erfüllt weiterhin die Anforderungen gemäß EN ISO 9001 und EN ISO 13485.

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



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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



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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 (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 (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 (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 (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 (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 (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 (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

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EN ISO 5356-1:2015 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets
EN ISO 80369-1:2018 (ISO 80369-1:2018)	Small-bore connectors for liquids and gases in healthcare applications -- Part 1: General requirements
EN ISO 5367:2014 (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

Extend of conformity assessment / Umfang der Konformitätsbewertung		
Part Number / 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

**ЕС декларация за съответствие**

Документ №
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Drägerwerk AG & Co. KGaA
Moislinger Allee 53-55
23542 Lübeck
Germany

Упълномощен представител: N/A

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

Еднократен регистрационен номер (SRN): DE-MF-000005329

с настоящото декларира на своя отговорност, че

Име на продукта	Категория на уреда	Клас на уреда	Код UMDNS / Код GMDN / Код EMDN
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

отговаря на следните разпоредби:

ЕВРОПЕЙСКИ РЕГЛАМЕНТ (ЕС) 2017/745 относно медицински изделия. Извършено е проучване на системата за управление на качеството в съответствие с анекс IX (глави I и III и раздел 4) на регламента от нотифицирания орган:

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

Системата за управление на качеството също отговаря на EN ISO 9001 и EN ISO 13485.

За продукти, пуснати на пазара, тази декларация е в сила от датата на издаване. Всяка модификация на уреда, която не е разрешена от Dräger, обезсилва тази декларация.

Това е превод на оригиналния документ (en/de) и затова не е подписан.

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



ЕС декларация за съответствие

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

Име на продукта	Категория на уреда
ErgoStar	Catheter Mounts
Напълно или частично приложени стандарти:	
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 (IEC 62366-1: 2015 COR 1 2016 AMD1:2020)	Medical devices - Part 1: Application of usability engineering to medical devices
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ISO 18190:2016	Anaesthetic and respiratory equipment – General requirements for airways and related equipment
EN ISO 5356-1:2015 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



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EN ISO 80369-1:2018 (ISO 80369-1:2018)	Small-bore connectors for liquids and gases in healthcare applications -- Part 1: General requirements
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**ЕС декларация за съответствие**

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

Обхват на оценката на съответствие		
Номер на частта	Име на продукта	Основен идентификатор 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



Declaración UE de conformidad

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

Representante autorizado:

N/A

EC Certificate: G10 010578 0039

Valid until: 2025-03-17

Número de registro único (SRN):

DE-MF-000005329

por la presente declara bajo su exclusiva responsabilidad que

Nombre del producto	Categoría del dispositivo	Clase del dispositivo	Código UMDNS / Código GMDN / Código EMDN
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

cumple las siguientes disposiciones:

REGLAMENTO EUROPEO (UE) 2017/745 sobre los productos sanitarios. Se ha efectuado un examen del sistema de gestión de la calidad siguiendo el anexo IX (capítulos I y III y sección 4) del reglamento del organismo notificado: **TÜV Süd Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany, NB 0123**

El sistema de gestión de la calidad también cumple con EN ISO 9001 y EN ISO 13485.

Esta declaración será efectiva para los productos puestos en el mercado a partir de la fecha de publicación. Cualquier modificación del dispositivo no autorizada por Dräger invalidará esta declaración.

Esta es una traducción del documento original (en/de) y, por lo tanto, no lleva firma.

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

Nombre del producto	Categoría del dispositivo
ErgoStar	Catheter Mounts
Normas aplicadas total o parcialmente:	
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 (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 (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 (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 (ISO 80601-2-12:2020)	Medical electrical equipment - Part 2-12: Particular requirements for basic safety and essential performance of critical care ventilators
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EN ISO 10993-1:2020 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



Declaración UE de conformidad

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EN ISO 80369-1:2018 (ISO 80369-1:2018)	Small-bore connectors for liquids and gases in healthcare applications -- Part 1: General requirements
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Drägerwerk AG & Co. KGaA
Moislinger Allee 53-55
23542 Lübeck
Germany

Alcance de la evaluación de conformidad		
Número de referencia	Nombre del producto	UDI-DI básico
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



EU prohlášení o shodě

Č. dokumentu

Datum

Místo

Strana

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

Zplnomocněným zástupcem:

N/A

EC Certificate: G10 010578 0039

Valid until: 2025-03-17

Jednorázové registrační číslo (SRN):

DE-MF-000005329

tímto prohlašuje na svou výhradní zodpovědnost, že

Název produktu	Kategorie prostředku	Třída prostředku	Kód UMDNS / Kód GMDN / Kód EMDN
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

splňuje následující ustanovení:

NAŘÍZENÍ EVROPSKÉHO PARLAMENTU A RADY (EU) 2017/745 o zdravotnických prostředcích. Kontrola systému managementu kvality byla provedena oznámeným subjektem podle přílohy IX (kapitol I a III a oddílu 4) nařízení:

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

Systém managementu kvality splňuje rovněž požadavky norem EN ISO 9001 a EN ISO 13485.

Toto prohlášení nabývá platnosti pro produkty uvedené na trh ke dni vydání. Jakákoli úprava prostředku, která není schválena společností Dräger, toto prohlášení zneplatní.

Toto je překlad původního dokumentu (en/de), a proto nenese podpis.

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

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 prohlášení o shodě

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Germany - Lübeck

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

Název produktu	Kategorie prostředku
ErgoStar	Catheter Mounts
Použité normy, v celku nebo z části:	
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 (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 (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
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ISO 15223-1:2021	Medical devices – Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General requirements.
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EN ISO 80601-2-13:2012+A2:2019 (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 (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 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



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Germany - Lübeck

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EN ISO 5367:2014 (ISO 5367:2014)	Anaesthetic and respiratory equipment - Breathing sets and connectors
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**EU prohlášení o shodě**

Č. dokumentu
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2022-12-02
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Drägerwerk AG & Co. KGaA
Moislinger Allee 53-55
23542 Lübeck
Germany

Rozsah posuzování shody		
Číslo dílu	Název produktu	Základní 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



EU-overensstemmelseserklæring

Dokumentnr.

Dato

Sted

Side

MDR108-007-2212-029-0

2022-12-02

Germany - Lübeck

1 / 4

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

Autoriseret repræsentant:

N/A

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

Individuelt registreringsnummer (SRN):

DE-MF-000005329

erklærer hermed på eget ansvar, at

Produktnavn	Apparatkategori	Apparatklasse	UMDNS-kode / GMDN-kode / EMDN-kode
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

opfylder følgende bestemmelser:

EUROPÆISK FORORDNING (EU) 2017/745 om medicinsk udstyr. En undersøgelse af kvalitetsstyringssystemet er blevet foretaget i overensstemmelse med bilag IX (kapitel I og III og afsnit 4) til forordningen af det bemyndigede organ:

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

Kvalitetsstyringssystemet overholder ligeledes EN ISO 9001 og EN ISO 13485.

Denne erklæring gælder for produkter, der markedsføres efter udstedelsesdatoen. Ved enhver ændring af udstyret, der ikke er godkendt af Dräger, mister denne erklæring sin gyldighed.

Dette er en oversættelse af det originale dokument (en/de) og er derfor ikke forsynet med en underskrift.

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



EU-overensstemmelseserklæring

Dokumentnr.

Dato

Sted

Side

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

Produktnavn	Apparatkategori
ErgoStar	Catheter Mounts
Standarder, der anvendes helt eller delvist:	
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 (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 (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 (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 (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 (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
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EN ISO 10993-1:2020 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets

da



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Germany - Lübeck

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

da



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Germany - Lübeck

4 / 4

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

Overensstemmelsesvurderingens omfang		
Varenummer	Produktnavn	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

et



ELi vastavusdeklaratsioon

Dokumendi nr
Kuupäev
Koht
Lk

MDR108-007-2212-029-0
2022-12-02
Germany - Lübeck
1 / 4

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

Volitatud esindaja:

N/A

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

Kordumatu registreerimisnumber (SRN): DE-MF-000005329

kinnitab käesolevaga oma ainuvastutusel, et

Toote nimi	Seadme kategooria	Seadme klass	UMDNS-kood / GMDN-kood / EMDN-kood
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

vastab järgmistele nõuetele:

EUROOPA MÄÄRUS (EL) 2017/745 meditsiiniseadmete kohta. Kvaliteedijuhtimise süsteemi on hinnatud teavitatud asutuses määruse IX lisa (peatükid I ja III ning jaotis 4) alusel:

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

Kvaliteedikontrolli süsteem vastab ka standarditele EN ISO 9001 ja EN ISO 13485.

Käesolev deklaratsioon kehtib toodete kohta, mis on turule toodud alates deklaratsiooni väljaandmise kuupäevast. Deklaratsioon kaotab kehtivuse, kui tootel tehakse muudatusi, mille kohta ei ole Drägerilt nõusolekut saadud.

Tegu on originaaldokumendi (en/de) tõlkega ja seetõttu ei ole sellel allkirja.

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

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



ELi vastavusdeklaratsioon

Dokumendi nr
Kuupäev
Koht
Lk

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

Toote nimi	Seadme kategooria
ErgoStar	Catheter Mounts
Osaliselt või täielikult kohaldatud standardid:	
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 (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 (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
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EN ISO 80601-2-13:2012+A2:2019 (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
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EN ISO 10993-1:2020 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets

et



ELi vastavusdeklaratsioon

Dokumendi nr
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2022-12-02
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EN ISO 80369-1:2018 (ISO 80369-1:2018)	Small-bore connectors for liquids and gases in healthcare applications -- Part 1: General requirements
EN ISO 5367:2014 (ISO 5367:2014)	Anaesthetic and respiratory equipment - Breathing sets and connectors
EN ISO 20417:2021	Medical devices - Information to be supplied by the manufacturer

et



ELi vastavusdeklaratsioon

Dokumendi nr
Kuupäev
Koht
Lk

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

Vastavushinnangu ulatus		
Osa number	Toote nimi	Peamine 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

**Δήλωση συμμόρφωσης ΕΕ**

Αρ. εγγράφου
Ημερομηνία
Τοποθεσία
Σελίδα

MDR108-007-2212-029-0
2022-12-02
Germany - Lübeck
1 / 4

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

Εξουσιοδοτημένος
αντιπρόσωπος:

N/A

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

Μεμονωμένος αριθμός εγγραφής (SRN):

DE-MF-000005329

δηλώνει με αποκλειστική ευθύνη ότι

Όνομα προϊόντος	Κατηγορία συσκευής	Κλάση συσκευής	Κωδικός UMDNS / Κωδικός GMDN / Κωδικός EMDN
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

συμμορφώνεται με τις ακόλουθες διατάξεις:

Ευρωπαϊκός κανονισμός (ΕΕ) 2017/745 περί ιατροτεχνολογικών προϊόντων. Πραγματοποιήθηκε έλεγχος του συστήματος διαχείρισης ποιότητας σύμφωνα με το παράρτημα IX (κεφάλαια I και III και ενότητα 4) του κανονισμού από τον κοινοποιημένο οργανισμό:

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

Το σύστημα διαχείρισης ποιότητας συμμορφώνεται επίσης με τα πρότυπα EN ISO 9001 και EN ISO 13485.

**Η παρούσα δήλωση ισχύει για προϊόντα που τίθενται στην αγορά από την ημερομηνία έκδοσης.
Οποιαδήποτε τροποποίηση στη συσκευή χωρίς την έγκριση της Dräger θα ακυρώσει την παρούσα δήλωση.**

Το παρόν αποτελεί μετάφραση του πρωτότυπου εγγράφου (από τα αγγλικά/γερμανικά) και γι' αυτό το λόγο δεν φέρει σφραγίδα.

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

Drägerwerk AG & Co. KGaA
Moislinger Allee 53-55
23558 Lübeck, Germany
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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



Δήλωση συμμόρφωσης ΕΕ

Αρ. εγγράφου
Ημερομηνία
Τοποθεσία
Σελίδα

MDR108-007-2212-029-0
2022-12-02
Germany - Lübeck
2 / 4

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

Όνομα προϊόντος	Κατηγορία συσκευής
ErgoStar	Catheter Mounts
Πρότυπα που εφαρμόζονται πλήρως ή εν μέρει:	
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 (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 (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
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EN ISO 10993-1:2020 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



Δήλωση συμμόρφωσης ΕΕ

Αρ. εγγράφου
Ημερομηνία
Τοποθεσία
Σελίδα

MDR108-007-2212-029-0
2022-12-02
Germany - Lübeck
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EN ISO 80369-1:2018 (ISO 80369-1:2018)	Small-bore connectors for liquids and gases in healthcare applications -- Part 1: General requirements
EN ISO 5367:2014 (ISO 5367:2014)	Anaesthetic and respiratory equipment - Breathing sets and connectors
EN ISO 20417:2021	Medical devices - Information to be supplied by the manufacturer

**Δήλωση συμμόρφωσης ΕΕ**

Αρ. εγγράφου
Ημερομηνία
Τοποθεσία
Σελίδα

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

Επέκταση αξιολόγησης της συμμόρφωσης		
Αριθμός εξαρτήματος	Όνομα προϊόντος	Βασικό 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



Déclaration de conformité UE

N° du document

Date

Ville

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2022-12-02

Germany - Lübeck

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

Mandataire:

N/A

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

Numéro d'enregistrement unique (SRN): DE-MF-000005329

déclare par la présente et sous sa seule responsabilité que le

Nom du produit	Catégorie de l'appareil	Classe de l'appareil	Code UMDNS / Code GMDN / Code EMDN
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

satisfait aux dispositions suivantes :

RÉGLEMENTATION EUROPÉENNE (UE) 2017/745 sur les dispositifs médicaux. Une vérification du système de gestion de la qualité a été réalisée conformément à l'annexe IX (chapitres I et III, ainsi que section 4) de la réglementation suivante par l'organisme notifié :

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

Le système de gestion de la qualité satisfait également aux normes EN ISO 9001 et EN ISO 13485.

La déclaration s'applique aux produits mis sur le marché à partir de la date de publication. Toute modification non autorisée par Dräger apportée sur l'appareil rend cette déclaration caduque.

Il s'agit d'une traduction du document original (en/de) et ne porte donc pas de signature.

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

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



Déclaration de conformité UE

N° du document

Date

Ville

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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

Nom du produit	Catégorie de l'appareil
ErgoStar	Catheter Mounts
Normes appliquées en totalité ou en partie :	
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 (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 (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 (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.
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2022-12-02

Germany - Lübeck

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EN ISO 80369-1:2018 (ISO 80369-1:2018)	Small-bore connectors for liquids and gases in healthcare applications -- Part 1: General requirements
EN ISO 5367:2014 (ISO 5367:2014)	Anaesthetic and respiratory equipment - Breathing sets and connectors
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Déclaration de conformité UE

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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

Étendue de l'évaluation de la conformité		
Référence de pièce	Nom du produit	IUD-ID de base
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



EU izjava o sukladnosti

Br. dokumenta

Datum

Mjesto

Stranica

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

Ovlašteni zastupnik:

N/A

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

Jedinstveni registracijski broj (SRN):

DE-MF-000005329

ovime izjavljuje pod vlastitom odgovornošću da je

Naziv proizvoda	Kategorija proizvoda	Razred proizvoda	UMDNS kod / GMDN kod / EMDN kod
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

sukladan sa sljedećim odredbama:

UREDBA (EU) 2017/745 o medicinskim proizvodima. Ocjena sustava upravljanja kvalitetom provedena je prema Prilogu IX. (poglavljju I. i III. i stavku 4.) uredbe od strane prijavljenog tijela:

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

Sustav upravljanja kvalitetom također je sukladan normama EN ISO 9001 i EN ISO 13485.

Ova izjava za proizvode stavljene na tržište stupa na snagu od datuma izdavanja. U slučaju bilo kakvih izmjena proizvoda koje nisu odobrene od strane tvrtke Dräger ova izjava gubi svoju valjanost.

Ovo je prijevod izvornog dokumenta (engl./njem.) i stoga ne sadrži potpis.

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



EU izjava o sukladnosti

Br. dokumenta

Datum

Mjesto

Stranica

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

Naziv proizvoda	Kategorija proizvoda
ErgoStar	Catheter Mounts
Norme primijenjene u cijelosti ili djelomično:	
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 (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 (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 (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 (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 (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 (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 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



EU izjava o sukladnosti

Br. dokumenta

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Stranica

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2022-12-02

Germany - Lübeck

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



EU izjava o sukladnosti

Br. dokumenta
Datum
Mjesto
Stranica

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

Opseg ocjene sukladnosti		
Broj dijela	Naziv proizvoda	Osnovni 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

**Dichiarazione di conformità UE**

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

Mandatario:

N/A

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

Numero di registrazione unico (SRN):

DE-MF-000005329

dichiara con la presente sotto la propria responsabilità che

Nome prodotto	Categoria dispositivo	Classe dispositivo	Codice UMDNS / Codice GMDN / Codice EMDN
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

è conforme alle seguenti disposizioni:

REGOLAMENTO EUROPEO (UE) 2017/745 relativo ai dispositivi medici. È stata effettuata una verifica del sistema di gestione della qualità ai sensi dell'allegato IX (capitoli I e III e sezione 4) del regolamento dell'organismo notificato:

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

Il sistema di gestione della qualità è altresì conforme alle norme EN ISO 9001 e EN ISO 13485.

La presente dichiarazione è valevole per i prodotti lanciati sul mercato a partire dalla data di pubblicazione. Qualsiasi modifica del dispositivo non autorizzata da Dräger invalida la presente dichiarazione.

Si tratta di una traduzione del documento originale (en/de) e non porta pertanto una firma.

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

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



Dichiarazione di conformità UE

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

Nome prodotto	Categoria dispositivo
ErgoStar	Catheter Mounts
Standard applicati integralmente o parzialmente:	
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 (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 (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 (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 (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 (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 (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 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



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EN ISO 80369-1:2018 (ISO 80369-1:2018)	Small-bore connectors for liquids and gases in healthcare applications -- Part 1: General requirements
EN ISO 5367:2014 (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

Estensione della valutazione di conformità		
Numero d'ordine	Nome prodotto	UDI-DI di base
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



ES atbilstības deklarācija

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

Pilnvarotais pārstāvis:

N/A

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

Vienotais reģistrācijas numurs (VRN):

DE-MF-000005329

pilnībā atbildot par to, apliecina, ka

Izstrādājuma nosaukums	Ierīces kategorija	Ierīces klase	UMDNS kods / GMDN kods / EMDN kods
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

atbilst šādiem noteikumiem:

EIROPAS REGULA (ES) 2017/745 par medicīnas ierīcēm. Kvalitātes vadības sistēmas pārbaudi veikusi pilnvarotā iestāde saskaņā ar regulas IX. pielikumu (nodaļas I un III, 4. sadaļa):

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

Kvalitātes vadības sistēma atbilst arī EN ISO 9001 un EN ISO 13485.

Šī deklarācija ir spēkā izstrādājumiem, kas laisti tirgū no izdošanas datuma. Jebkādi ierīces pārveidojumi, kurus nav atļāvis Dräger, padarīs šo deklarāciju par spēkā neesošu.

Šis ir oriģinālā dokumenta (en/de) tulkojums, tādēļ uz tā nav paraksta.

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

Drägerwerk AG & Co. KGaA
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info@draeger.com
www.draeger.com
VAT no. DE135082211

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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

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Local court Lübeck HRB 7903 HL
General partner: Drägerwerk
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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



ES atbilstības deklarācija

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

Izstrādājuma nosaukums	Ierīces kategorija
ErgoStar	Catheter Mounts
Pilnībā vai daļēji piemērotie standarti:	
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 (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 (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
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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 (ISO 80601-2-12:2020)	Medical electrical equipment - Part 2-12: Particular requirements for basic safety and essential performance of critical care ventilators
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EN ISO 10993-1:2020 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



ES atbilstības deklarācija

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2022-12-02
Germany - Lübeck
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EN ISO 80369-1:2018 (ISO 80369-1:2018)	Small-bore connectors for liquids and gases in healthcare applications -- Part 1: General requirements
EN ISO 5367:2014 (ISO 5367:2014)	Anaesthetic and respiratory equipment - Breathing sets and connectors
EN ISO 20417:2021	Medical devices - Information to be supplied by the manufacturer



ES atbilstības deklarācija

Dokumenta Nr.
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Lappuse

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

Atbilstības novērtēšanas pagarinājums		
Daļas numurs	Izstrādājuma nosaukums	Pamata 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



ES atitikties deklaracija

Dokumento Nr.
Data
Vieta
Psl.

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

Igaliotasis atstovas:

N/A

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

Bendrasis registracijos numeris (BRN):

DE-MF-000005329

prisiimdami visą atsakomybę pareiškia, kad:

Prietaiso pavadinimas	Prietaiso kategorija	Prietaiso klasė	UMDNS kodas / GMDN kodas / EMDN kodas
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

atitinka šias nuostatas:

EUROPOS PARLAMENTO IR TARYBOS REGLAMENTA (ES) 2017/745 dėl medicinos prietaisų. Notifikuotoji įstaiga, atlikusi kokybės valdymo sistemos patikrinimą pagal reglamento IX priedą (I ir III skyrius bei 4 skirsnį):
TÜV Süd Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany, NB 0123

Kokybės valdymo sistema taip pat atitinka EN ISO 9001 ir EN ISO 13485 standartus.

Ši deklaracija taikoma prietaisams, pateiktiems į rinką jų išleidimo dieną. Atlikus neleistinus „Dräger“ prietaiso keitimus, ši deklaracija taps negaliojanti.

Tai yra originalaus dokumento vertimas (iš anglų / vokiečių k.), todėl nereikia parašo.

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



ES atitikties deklaracija

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

Prietaiso pavadinimas	Prietaiso kategorija
ErgoStar	Catheter Mounts
Iš dalies ar visa apimtimi taikyti standartai:	
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 (IEC 62366-1: 2015 COR 1 2016 AMD1:2020)	Medical devices - Part 1: Application of usability engineering to medical devices
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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 (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 (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 (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 (ISO 10993-1:2018)	Biological evaluation of medical devices -- Part 1: Evaluation and testing within a risk management process
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EN ISO 5356-1:2015 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



ES atitikties deklaracija

Dokumento Nr.
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MDR108-007-2212-029-0
2022-12-02
Germany - Lübeck
3 / 4

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



ES atitikties deklaracija

Dokumento Nr.
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MDR108-007-2212-029-0
2022-12-02
Germany - Lübeck
4 / 4

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

Išsami informacija apie atitikties vertinimą		
Prekės kodas	Prietaiso pavadinimas	Pagrindinis 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

**EU megfeleléségi nyilatkozat**

Dokumentum száma

Dátum

Hely

Oldal

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

Meghatalmazott képviselő:

N/A

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

Egyedi regisztrációs szám (SRN):

DE-MF-000005329

saját kizárólagos felelősségére kijelenti, hogy a

Termék neve	Készülékkategória	Készülékosztály	UMDNS-kód / GMDN-kód / EMDN-kód
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

megfelel a következő rendelkezéseknek:

AZ EURÓPAI PARLAMENT ÉS A TANÁCS (EU) 2017/745 RENDELETE az orvostechnikai eszközökről. A minőségirányítási rendszer vizsgálatát a bejelentett szervezet az irányelv IX. melléklete (az I. és III. fejezet és a 4. szakasz) szerint végezte:

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

A minőségirányítási rendszer megfelel továbbá az EN ISO 9001 és az EN ISO 13485 szabványoknak is.

Ez a nyilatkozat a kiállítását követően forgalomba hozott termékekre érvényes. A készüléken végzett bármilyen, a Dräger által nem engedélyezett módosítás érvényteleníti a nyilatkozatot.

Ez az eredeti dokumentum (en/de) fordítása, és ezért nem szerepel rajta aláírás.

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



EU megfeleléségi nyilatkozat

Dokumentum száma

Dátum

Hely

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2022-12-02

Germany - Lübeck

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

Termék neve	Készülékkategória
ErgoStar	Catheter Mounts
Teljesen vagy részben alkalmazott szabványok:	
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 (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 (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 (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 (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 (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 (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 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



EU megfeleléségi nyilatkozat

Dokumentum száma

Dátum

Hely

Oldal

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2022-12-02

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

**EU megfeleléségi nyilatkozat**

Dokumentum száma

Dátum

Hely

Oldal

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

A megfeleléséértékelés meghosszabbítása		
Cikkszám	Termék neve	Alapvető 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



EU-verklaring van overeenstemming

Documentnr.

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2022-12-02

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

Gemachtigde:

N/A

EC Certificate: G10 010578 0039

Valid until: 2025-03-17

Enkelvoudig registratienummer (SRN):

DE-MF-000005329

verklaart hierbij onder haar volledige eigen verantwoordelijkheid dat

Productnaam	Apparaatcategorie	Apparaatklasse	UMDNS-code / GMDN-code / EMDN-code
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

voldoet aan de volgende bepalingen:

EUROPESE VERORDENING (EU) 2017/745 voor medische hulpmiddelen. De aangemelde instantie heeft het kwaliteitsborgingssysteem onderzocht overeenkomstig Bijlage IX (hoofdstukken I en III en deel 4) van de verordening:

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

Het kwaliteitsmanagementsysteem voldoet ook aan EN ISO 9001 en EN ISO 13485.

Deze verklaring geldt voor producten die op de markt zijn gebracht vanaf de datum van afgifte. Elke modificatie van het product waarvoor Dräger geen toestemming heeft gegeven, maakt deze verklaring ongeldig.

Dit is een vertaling van het originele document en benodigt derhalve geen ondertekening.

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

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



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

Productnaam	Apparaatcategorie
ErgoStar	Catheter Mounts
Volledig of gedeeltelijk toegepaste 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 (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 (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 (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 (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 (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 (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 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



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



EU-verklaring van overeenstemming

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

Reikwijdte van conformiteitsbeoordeling		
Onderdeelnummer	Productnaam	Basis 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

**Deklaracja zgodności UE**

Nr dokumentu

Data

Miejsce

Strona

MDR108-007-2212-029-0

2022-12-02

Germany - Lübeck

1 / 4

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

Upoważniony przedstawiciel:

N/A

EC Certificate: G10 010578 0039

Valid until: 2025-03-17

Pojedynczy numer rejestracyjny (SRN):

DE-MF-000005329

deklaruje niniejszym na swoją wyłączną odpowiedzialność, że

Nazwa produktu	Kategoria urządzenia	Klasa urządzenia	Kod UMDNS / Kod GMDN / Kod EMDN
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

spełnia wymogi następujących przepisów:

ROZPORZĄDZENIE PARLAMENTU EUROPEJSKIEGO I RADY (EU) 2017/745 w sprawie wyrobów medycznych. Zostało przeprowadzone badanie systemu zarządzania jakością zgodnie z Załącznikiem IX (rozdziały I i III, sekcja 4) Rozporządzenia przez jednostkę notyfikowaną:

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

System zarządzania jakością spełnia też normy EN ISO 9001 i EN ISO 13485.

Niniejsza deklaracja dotyczy produktów wprowadzonych na rynek wg daty wydania. Wszelkie modyfikacje urządzenia niezatwierdzone przez Dräger spowodują utratę ważności niniejszej deklaracji.

Jest to tłumaczenie oryginalnego dokumentu i dlatego nie jest opatrzone podpisem.

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

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



Deklaracja zgodności UE

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

Nazwa produktu	Kategoria urządzenia
ErgoStar	Catheter Mounts
Zastosowane normy (w całości lub w części):	
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 (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 (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 (ISO 14971 :2019)	Medical devices – Application of risk management to medical devices
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EN ISO 10993-1:2020 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



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

**Deklaracja zgodności UE**

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

Zakres oceny zgodności		
Numer części	Nazwa produktu	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

**Declaração de conformidade da UE**

Nº. do documento

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

Mandatário:

N/A

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

O número de registro único (SRN):

DE-MF-000005329

declara, sob exclusiva responsabilidade, que

Nome do produto	Categoria do equipamento	Classe do equipamento	Código UMDNS / Código GMDN / Código EMDN
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

está em conformidade com as seguintes disposições:

REGULAMENTO (UE) 2017/745 relativo aos dispositivos médicos. Um exame do sistema de gerenciamento de qualidade foi realizado seguindo o Anexo IX (Capítulos I e III e a seção 4) do regulamento pelo Órgão notificado: **TÜV Süd Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany, NB 0123**

O sistema de gerenciamento de qualidade também está em conformidade com a EN ISO 9001 e a EN ISO 13485.

Esta declaração é válida para produtos colocados no mercado a partir da data de emissão. Quaisquer modificações no equipamento não autorizadas pela Dräger invalidarão esta declaração.

Este documento é uma tradução do documento original (en/de) e, portanto, não precisa ser assinado.

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



Declaração de conformidade da UE

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

Nome do produto	Categoria do equipamento
ErgoStar	Catheter Mounts
Normas aplicadas total ou parcialmente:	
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 (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 (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 (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 (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 (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 (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 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



Declaração de conformidade da UE

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Data

2022-12-02

Local

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



Declaração de conformidade da UE

Nº. do documento

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Data

2022-12-02

Local

Germany - Lübeck

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

Extensão da avaliação de conformidade		
Número da peça	Nome do produto	UDI-DI básico
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



Declarație de conformitate UE

Nr. document

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2022-12-02

Germany - Lübeck

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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 / Codul GMDN / Codul EMDN
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ 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



Declarație de conformitate UE

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2022-12-02
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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 (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 (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 (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 (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 (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 (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 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



Declarație de conformitate UE

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Germany - Lübeck
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EN ISO 80369-1:2018 (ISO 80369-1:2018)	Small-bore connectors for liquids and gases in healthcare applications -- Part 1: General requirements
EN ISO 5367:2014 (ISO 5367:2014)	Anaesthetic and respiratory equipment - Breathing sets and connectors
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**Declarație de conformitate UE**

Nr. document

Data

Localitatea

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2022-12-02

Germany - Lübeck

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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

**EÚ vyhlásenie o zhode**

Dokument č.
Dátum
Miesto
Strana

MDR108-007-2212-029-0
2022-12-02
Germany - Lübeck
1 / 4

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

Splnomocnený zástupca:

N/A

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

Jedinečné registračné číslo (SRN):

DE-MF-000005329

týmto na vlastnú zodpovednosť vyhlasuje, že

Názov výrobku	Kategória zariadenia	Trieda zariadenia	Kód UMDNS / Kód GMDN / Kód EMDN
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

spĺňa nasledujúce nariadenia:

EURÓPSKE NARIADENIE (EÚ) 2017/745 o zdravotníckych pomôckach. Preskúmanie systému riadenia kvality bolo vykonané notifikovaným orgánom podľa prílohy IX (kapitoly I a III a oddielu 4) nariadenia:
TÜV Süd Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany, NB 0123

Systém riadenia kvality tiež spĺňa normy STN EN ISO 9001 a STN EN ISO 13485.

Toto vyhlásenie pre výrobky uvedené na trh nadobúda platnosť dňom vydania. Akékoľvek zmeny zariadenia, ktoré neschválila spoločnosť Dräger, vedú k strate platnosti tohto vyhlásenia.

Toto je preklad pôvodného dokumentu (en/de) a preto na ňom nie je uvedený podpis.

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



EÚ vyhlásenie o zhode

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Strana

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2022-12-02
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Drägerwerk AG & Co. KGaA
Moislinger Allee 53-55
23542 Lübeck
Germany

Názov výrobku	Kategória zariadenia
ErgoStar	Catheter Mounts
Použité normy v úplnom alebo v čiastočnom znení:	
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 (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 (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 (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 (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 (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 (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 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



EÚ vyhlásenie o zhode

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



EÚ vyhlásenie o zhode

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

Rozsah posúdenia zhody		
Objednávacie číslo	Názov výrobku	Základné 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

Pooblašчени predstavnik:

N/A

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

Enotna registrska številka (SRN):

DE-MF-000005329

izjavlja z vso odgovornostjo, da

Ime izdelka	Kategorija naprave	Razred naprave	Koda UMDNS / Koda GMDN / Koda EMDN
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

izpolnjuje naslednje določbe:

EVROPSKA UREDBA (EU) 2017/745 o medicinskih pripomočkih. Sistem upravljanja kakovosti je na podlagi Priloge IX (poglavji I in III in razdelek 4) uredbe preveril priglašeni organ:

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

Sistem upravljanja kakovosti je skladen tudi z EN ISO 9001 in EN ISO 13485.

Ta izjava velja za izdelke, na trg dane z datumom izdaje. Vsaka sprememba naprave brez soglasja družbe Dräger razveljavi to izjavo.

To je prevod originalnega dokumenta (en/de) in zato ni podpisan.

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



Izjava EU o skladnosti

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

Ime izdelka	Kategorija naprave
ErgoStar	Catheter Mounts
V celoti ali deloma uporabljeni standardi:	
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 (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 (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 (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 (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 (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 (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 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



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

**Izjava EU o skladnosti**

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

Obseg ugotavljanja skladnosti		
Kataloška številka	Ime izdelka	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

**EU-vaatimustenmukaisuusvakuutus**

Asiakirjan nro
Päivämäärä
Paikka
Sivu

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

Valtuutetulla edustajalla:

N/A

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

Rekisterinumero:

DE-MF-000005329

vakuuttaa täten yksinomaisella vastuullaan, että

Tuotenimi	Laitteen luokitus	Laiteluokka	UMDNS-koodi / GMDN-koodi / EMDN-koodi
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

täyttää seuraavat vaatimukset:

EUROOPAN PARLAMENTIN JA NEUVOSTON ASETUS (EU) 2017/745 lääkinnällisistä laitteista. Laatujärjestelmän on tarkastanut asetuksen liitettä IX (I ja III luvun 4 kohtaa) noudattaen seuraava ilmoitettu laitos:
TÜV Süd Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany, NB 0123

Laatujärjestelmä täyttää lisäksi standardien EN ISO 9001 ja EN ISO 13485 vaatimukset.

Tätä vakuutusta sovelletaan tuotteisiin, jotka on saatettu markkinoille antamispäivästä alkaen. Laitteeseen tehtävät muutokset, joita Dräger ei ole hyväksynyt, mitätöivät tämän vakuutuksen.

Tämä on alkuperäisen (saksan-/englanninkielisen) asiakirjan käännös, eikä siinä siksi ole allekirjoitusta.

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



EU-vaatimustenmukaisuusvakuutus

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

Tuotenimi	Laitteen luokitus
ErgoStar	Catheter Mounts
Kokonaan tai osittain sovellettavat standardit:	
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 (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 (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 (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 (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 (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
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EN ISO 10993-1:2020 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets

**EU-vaatimustenmukaisuusvakuutus**

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

**EU-vaatimustenmukaisuusvakuutus**

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

Vaatimustenmukaisuuden arvioinnin laajuus		
Osanumero	Tuotenimi	Yksilöllinen 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

**EU-försäkran om överensstämmelse**

Dokument nr.

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2022-12-02

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

Auktoriserad representant:

N/A

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

Enkelt registreringsnummer (SRN):

DE-MF-000005329

förklarar härmed under sitt eget ansvar att

Produktnamn	Enhetskategori	Enhetsklass	UMDNS-kod / GMDN-kod / EMDN-kod
ErgoStar	Catheter Mounts	Ila	UMDNS 14-080/ GMDN 61346/ EMDN R0202

uppfyller följande bestämmelser:

EUROPEISKA FÖRORDNINGEN (EU) 2017/745 om medicintekniska produkter. En undersökning av kvalitetshanteringssystemet har utförts enligt bilaga IX (kapitel I och III och avsnitt 4) till förordningen av det anmälda organet:

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

Kvalitetshanteringssystemet uppfyller även EN ISO 9001 och EN ISO 13485.

Denna försäkran gäller för produkter som släpps ut på marknaden från och med utgivningsdatum. Alla ändringar av enheten som inte godkänts av Dräger ogiltiggör denna försäkran.

Detta är en översättning av originaldokument (en/de) och därför har det inte någon signatur.

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



EU-försäkran om överensstämmelse

Dokument nr.
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2022-12-02
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Drägerwerk AG & Co. KGaA
Moislinger Allee 53-55
23542 Lübeck
Germany

Produktnamn	Enhetskategori
ErgoStar	Catheter Mounts
Helt eller delvis tillämpade standarder:	
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 (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 (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
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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 (ISO 80601-2-12:2020)	Medical electrical equipment - Part 2-12: Particular requirements for basic safety and essential performance of critical care ventilators
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EN ISO 10993-1:2020 (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 (ISO 5356-1:2015)	Anaesthetic and respiratory equipment; Conical connectors – Part 1: Cones and sockets



EU-försäkran om överensstämmelse

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EN ISO 80369-1:2018 (ISO 80369-1:2018)	Small-bore connectors for liquids and gases in healthcare applications -- Part 1: General requirements
EN ISO 5367:2014 (ISO 5367:2014)	Anaesthetic and respiratory equipment - Breathing sets and connectors
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Drägerwerk AG & Co. KGaA
Moislinger Allee 53-55
23542 Lübeck
Germany

Omfattning bedömning av överensstämmelse		
Artikelnummer	Produktnamn	Bas 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