

EU Quality Management System Certificate

Certificate no.:
10000376655-PA-NoMA-DNK

Initial certification date:
26 August 2021

Valid Until:
26 August 2026

This is to certify that the quality system of

Coloplast A/S

Holtedam 1, 3050 Humlebaek, Denmark

SRN: DK-MF-000025526

For design, production, and final product inspection/testing of:

Foam wound dressings, Foam wound dressings with silver, Foam wound dressings with Ibuprofen, Antimicrobial Moisture Wicking Fabric with Silver, Superabsorbent wound dressing, Catheters for intermittent catheterization, Surgical accessories, Urological stents, Penile implants, Penile implant accessories, Drainage bags, Endourological instruments, Surgical Mesh, Single Incision Sling System, Urine bags, Penile prosthesis, Hydrocolloid wound dressings and Wound debridement pad, Gelling fiber dressing, Urethral dilation, Urological devices, urological accessories, Urinary/Suprapubic Indwelling Catheters and ostomy devices, Surgical Drainage.

Has been assessed and found to comply with respect to:

**The conformity assessment procedure described in Annex IX,
(Chapter I & III) of Regulation (EU) 2017/745 on Medical Devices**

Place and date:
Høvik, 13 November 2025

For the issuing office:
DNV Product Assurance AS – Notified Body 2460
Veritasveien 1, 1363 Høvik, Norway



Mariann Jeremiassen
Management Representative

Jurisdiction

Application of Regulation 2017/745 on medical devices, implemented in Norway by Act 7 May 2020 no. 37 on medical devices and Regulation 9 May 2021 no.1476 on medical devices by the Norwegian Ministry of Health and Care Services.

Certificate history:

Revision	Description	Report No.	Issue Date
0.0	Original Certificate	5254234	26 August 2021
1.0	Addition of Speedicath control	2499147	17 December 2021
2.0	Addition of Furlow	2499161	27 December 2021
3.0	Addition of Biosoft duo	2522909	03 February 2022
4.0	Blockchain data Changes	NA	03 February 2022
5.0	Addition of Titan and Titan Accessories	2522908	06 April 2022
6.0	Revision of wording and product name on certificate	2703041	06 May 2022
7.0	Addition of drainage bags	2707869	16 May 2022
8.0	Editorial change	NA	30 May 2022
9.0	Addition of Steerable Pusher and Hybrid Guidewire	2499146	17 June 2022
10.0	Addition of Biatain Ag Adhesive Biatain Ag Non-Adhesive and Biatain Silicone Ag	2522905	30 June 2022
11.0	Editing Furlow to be defined as reusable surgical instruments. Include Rossello and Brooks Dilator to reusable surgical instrument product list.	2706360	13 July 2022
12.0	Addition of Restorelle® Polypropylene Mesh and Altis® Single Incision Sling System	2499115	19 October 2022
13.0	Addition of Urine bags	2701701	25 October 2022
14.0	Addition of Genesis and adding sites relevant for production of devices included in this certificate. Adding EU technical documentation assessment certificate reference for Biosoft duo	2499117	22 November 2022
15.0	Move of location Coloplast Manufacturing France SAS from Champlan to Le Plessis-Pâté Revise EU technical documentation certificate number to Genesis	2793908	06 December 2022
16.0	To add Soft polyurethane (PU-S) double loop ureteral stents, Silicone double loop ureteral stents, Comfeel Plus, Vortek and Vortek hydro-coated double loop ureteral stents with hydrogel coating, Vortek Tumor stent - double loop ureteral stents, Biosoft duo multi-length hydro-coated double loop ureteral stents, Silicone hydro-coated double loop ureteral	2499125	03 February 2023



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	stents, Silicone Pyelostent double loop ureteral stents, Silicone Stenostent double loop ureteral stents, Rigid polyurethane (PU-R) double loop ureteral stents, Alprep Pad. Editorial change of name for Biosoft® duo double loop ureteral stents		
17	Include Wound debridement pad in scope to cover Alprep pad, to add Connectors, Catheter valve and Urethral dilation devices.	2797428	21 February 2023
18	Include Endourological instrument- Non steerable pusher	2807829	02 May 2023
19	Include Band-Aid hydrocolloid gel plaster	2727928	16 June 2023
20	Include Single loop ureteral stent, Elephant Suction-Irrigation Devices, Retrace ureteral access sheath, Ureteral dilators, Comfeel Plus Contour, Comfeel Plus Transparent, Guidewires – Stainless Steel	2738064	20 July 2023
21	Include Aris introducers and Dormia PCNL. Place urological accessories under same group. Move Biosoft® duo double loop ureteral stents to the Urological stents.	2712975	27 September 2023
22	Include Dormia No-Tip, Dormia N.Stone and Percutaneous Nephrostomy Dilators	2997369	07 November 2023
23	Include Biatain Fiber and Freudenberg Introducer	2499173	01 December 2023
24	Include Ureterostomy catheter and Short term enterocystoplasty catheter	2679416	07 December 2023
25	Include Biatain Silicone Sacral and Multishape. Move Biatain Fiber to other wound dressings and add intended purpose. Add Costa Rica site.	2499172	21 December 2023
26	Include Short term uretero-sigmoidostomy catheter, Single loop ureterostomy catheter, Percutaneous puncture needles, and ureteric catheters	3051268	19 January 2024
27	Include Folsil Catheters and SenSura Mio Baby	2499151	5 February 2024
28	Include Biatain Silicone Fit	3051268	04 March 2024
29	Include X-Flow prostatectomy catheter, Hydro X-Flow prostatectomy catheter, PA/PU double loop ureteral stent sets, Detour and Bonee	2499142	16 April 2024
30	Include Neoplex catheters without balloon. Corrected address for site 2 Espergærde.	2499113	27 May 2024
31	Include Floppy tip hydro-coated ureteric catheter	3123417	19 June 2024

Lack of fulfilment of conditions as set out in the Certification Agreement may render this Certificate invalid.

NOTIFIED BODY 2460: DNV Product Assurance AS, Veritasveien 1, 1363 Høvik, Norway, Tel +47 67 57 88 00, www.dnv.com

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32	Include Virtue Male Sling System	2499113	10 July 2024
33	Include Biatain Superabsorber dressing	2930666	28 August 2024
34	Include Biatain Ibu Non-Adhesive and Biatain Ibu Soft-Hold dressings	2499129, 2499130	25 November 2024
35	Include Assura/Alterna Post Op Ostomy bag	3071290	20 January 2025
36	Admin correction: change product name Assura/Alterna Post Op Ostomy bag to Sterile ostomy bags	3071290	22 January 2025
37	Admin correction: change of product description from Ostomy baseplates and bags (refeeding) to Ostomy bags	3071290	23 January 2025
38	Include Supraflow supra-pubic drainage set with silicone balloon catheter and Cystodrain integral set for supra-pubic drainage	2499143	07 February 2025
39	Include Silicone nephrostomy balloon catheter	2499157	02 May 2025
40	Include InterDry	2499116	21 May 2025
41	Include Percutaneous nephrostomy set with simple loop vortek catheter	2954648	04 June 2025
42	Include Vortek J and malecot nephrostomy catheters	3339904	10 June 2025
43	Include Escat transcystic drains in PVC, Multitubular drain silicone, Tubular drains silicone, Delbet drains, Multi-perforated tubular drain silicone radiopaque	2979526	13 November 2025

Products covered by this Certificate:

Product Description (and intended purpose for class IIb)	Product Name	Class
Silicone dressings Moist wound healing and exudate management	Biatain Silicone	IIb
	Biatain Silicone Lite	IIb
	Biatain Silicone Non-Border	IIb
Silicone dressings Moist wound healing and exudate management and may also be used as part of pressure injury prevention therapy	Biatain Silicone Sacral and Multishape	IIb
	Biatain Silicone Fit	
Gelling fiber dressing Moist wound healing and exudate management of moderate to high exuding wounds, including cavity wounds	Biatain Fiber	IIb
Superabsorbent wound dressing The product is intended for moist wound healing and exudate management	Biatain Superabsorber	IIb

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Urinary Catheters	Sterile intermittent catheters	Is
Surgical instruments	Reusable surgical instruments	Ir
Penile implants System surgically implanted for the management of erectile dysfunction	Titan	Ilb*
Penile implant accessories To facilitate assembly and implant of the Titan IPP	Titan	Ilb*
Drainage bags	Sterile drainage bags	Is
Endourological instrument	Steerable Pusher	Ila
	Non steerable pusher	Ila
	Hybrid Guidewire	Ila
	Guidewires – Stainless Steel	
Foam dressing with silver	Biatain Ag Adhesive	III*
Foam dressing with silver	Biatain Ag Non-Adhesive	III*
Foam dressing with ibuprofen	Biatain Ibu Non-Adhesive	III*
Foam dressing with ibuprofen	Biatain Ibu Soft-Hold	III*
Silicone foam dressing with silver	Biatain Silicone Ag	III*
Surgical Meshes	Restorelle® Polypropylene Mesh	III*
	Virtue Male Sling System	III*
Single Incision Sling System	Altis® Single Incision Sling System	III*
Antimicrobial Moisture Wicking Fabric with Silver	InterDry	III*
Urine bags	Sterile urine collection bags	Is
Penile prosthesis	Genesis® Malleable Penile Prosthesis	Ilb*
Urological stents Drainage of the upper urinary tract over fistulas or ureteral obstacles and healing of the ureter	Soft polyurethane (PU-S) double loop ureteral stents	Ilb*
	Silicone double loop ureteral stents	Ilb*
	Vortek® double loop ureteral stents	Ilb*
	Vortek® hydro-coated double loop ureteral stents	Ilb*
	Vortek® Tumor stent - double loop ureteral stents	Ilb*
	Biosoft duo multi-length hydro-coated double loop ureteral stents	Ilb*

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	Rigid polyurethane (PU-R) double loop ureteral stents	IIb*
	Silicone hydro-coated double loop ureteral stents	IIb*
	Biosoft® duo double loop ureteral stents	IIb*
	Double loop ureteral stent set in polyamide	IIb*
	Double loop ureteral stent set in polyurethane	IIb*
Urological stents Drainage of the upper urinary tract and ureter healing during management of localized stenosis of ureteropelvic junction.	Silicone Pyelostent double loop ureteral stents	IIb*
Urological stents Drainage of the upper urinary tract and ureter healing during management of ureteral stenosis.	Silicone Stenostent double loop ureteral stents	IIb*
Urological Implants Long-term palliative treatment of ureteral obstruction.	Detour subcutaneous ureteral bypass	IIb*
Hydrocolloid wound dressings	Band-Aid hydrocolloid gel plaster for minor wounds Band-Aid hydrocolloid gel plaster for blisters	IIa
Hydrocolloid wound dressing Moist wound healing and exudate management	Comfeel Plus	IIb
	Comfeel Plus Contour	IIb
	Comfeel Plus Transparent	IIb
Wound debridement pad	Alprep Pad	Is
Urological accessories	Sterile Connectors	Is
	Sterile Tuohy Borst Adapter	Is
	Sterile Catheter valve	Is
	Sterile Urethral dilations	Is
	Sterile Ureteral dilators	Is
	Freudenberg Introducer	Is
	Sterile Percutaneous nephrostomy dilators	IIa
	Ureteric catheter for retrograde uretero-pyelography	Is
	Bonee needle for bladder injection	IIa

Urological devices	Percutaneous puncture needles	Ila
Urinary/Percutaneous Indwelling Catheters	Single loop ureteral stent	Ilb
	Ureterostomy catheter	Ilb
	Folysil Silicone Catheter	Ilb
	Folysil Silicone Catheter - Long-term	Ilb*
	Short term enterocystoplasty catheter	Ila
	Short term uretero-sigmoidostomy catheter	Ila
	Single loop ureterostomy catheter	Ilb
	Ureteric drainage catheter in neoplex	Ila
	Ureteric drainage catheter in polyamide	Ila
	Ureteric interventional catheter	Ila
	X-Flow prostatectomy catheter	Ila
	Hydro X-Flow prostatectomy catheter	Ila
	Neoplex catheters without balloon	Ila
	Floppy tip hydro-coated ureteric catheter	Ila
	Silicone nephrostomy balloon catheter	Ilb
	Percutaneous nephrostomy set with simple loop vortek catheter	Ilb
	Vortek J percutaneous nephrostomy catheter	Ilb
	Vortek malecot percutaneous nephrostomy catheter	Ilb
Urinary/Suprapubic Indwelling Catheters The devices are intended to be used for: Supra-pubic drainage of urine from the bladder and bladder instillation of physiological saline solution by the supra-pubic rout	Supraflow supra-pubic drainage set with silicone balloon catheter	Ilb
Urinary/Suprapubic Indwelling Catheters The devices are intended to be used for: - Supra-pubic drainage of the urinary bladder. - Bladder instillation of physiological saline solution by the supra-pubic route. - Replacement of supra-pubic drainage catheter.	Cystodrain integral set for supra-pubic drainage	Ilb
Surgical Accessories	Elefant Suction-Irrigation Devices	Ila
Ureteral access sheath, Ureteral access sheath with ureteral dilator	Retrace ureteral access sheath	Ila
Introducer needles	Aris introducers	Ila



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Ostomy baseplates and bags (refeeding)	SenSura Mio Baby	Ila
Ostomy bags	Sterile ostomy bags	Is
Stone extractors	Dormia PCNL	Ila
	Dormia No-Tip / Dormia N.Stone	Ila
Surgical Drainage	Escat transcystic drains in PVC	Ila
	Multitubular drain silicone	
	Tubular drains silicone	
	Delbet drains	
	Multi-perforated tubular drain silicone radiopaque	

* Class III and class IIb devices referred to in the second subparagraph of Article 52(4): Technical documentation assessment is covered by a separate EU Technical Documentation Assessment Certificate No.: C536740, C545160, C566662 NoMa DNK, C569918-NoMa-DNK, C581486 NoMA DNK, C582792 NoMa DNK, C591315 NoMa DNK, C582794 NoMa DNK, C591820 NoMa DNK, C592004 NoMa DNK, C594157 NoMa DNK, C687110, C672963 NoMa DNK, C683022, C677114 NoMA DNK, C677116.

The complete list of devices is filed with the Notified Body

Sites covered by this certificate

Site Name	Address
Coloplast A/S	Holtedam 1, 3050 Humlebaek, Denmark
Coloplast A/S	Aa. Louis-Hansens Allé 15, 3060 Espergærde, Denmark
Coloplast Hungary KFT	Búzavirág út 15, 2800 Tatabánya, Hungary
Coloplast Hungary KFT	Coloplast utca 2, 4300 Nyírbátor, Hungary
Coloplast Hungary KFT	Kerék utca 3, 2800 Tatabánya, Hungary
Coloplast (China) Ltd.	No. 202, Baocheng Rd, Xiangzhou District, Zhuhai 519030, China
Coloplast Corporation	1601 West River Road North, Minneapolis, MN 55411, USA
Coloplast Manufacturing US, LLC	1601 West River Road North, Minneapolis, MN 55411, USA
Coloplast Manufacturing France SAS	9 Avenue Edmond Rostand, CS 70218, 24206 Sarlat-la-Canéda Cedex, France
Coloplast Manufacturing France SAS	20 rue Blaise Pascal, 24200 Sarlat La Canéda, France
Coloplast Manufacturing France SAS	2 Rue Jacqueline Auriol, 91220 Le Plessis-Pâté, France
Coloplast Manufacturing France SAS	Lieudit La Boursidière, Centre d'Affaires, 92350 Le Plessis Robinson, France
Coloplast Volume Manufacturing Costa Rica SA	Calle 58, Zona Franca La Lima, La Lima, 30106, Cartago, Costa Rica

Terms and conditions

The certificate is subject to the following terms and conditions:

- Any producer (see 2001/95/EC for a precise definition) is liable for damage caused by a defect in his product(s), in accordance with directive 85/374/EEC, as amended, concerning liability of defective products.
- The certificate is only valid for the products and/or manufacturing premises listed above.
- The Manufacturer shall fulfil the obligations arising out of the quality system as approved and uphold it so that it remains adequate and efficient.
- The Manufacturer shall inform the Notified Body of any intended updating of the quality system and the Notified Body will assess the changes and decide if the certificate remains valid.
- Periodical audits will be held, in order to verify that the Manufacturer maintains and applies the quality system. Notified Body reserves the right, on a spot basis or based on suspicion, to pay unannounced visits.
- For the class III devices and IIb devices falling under Article 52 (4) covered this certificate is dependent on the continued validity of the EU Technical Documentation Assessment Certificate, covering the devices.

The following may render this Certificate invalid:

- Changes in the quality system affecting production.
- Periodical audits not held within the allowed time window

Specific conditions - Class I devices, Systems and Procedure Packs:

- For class I device being placed on the market in a sterile condition, Class I devices with a measurement function and class I devices being reusable surgical instruments covered by this certificate the audit by the notified body of the quality management system was limited to the aspects required under article 52(7) of the regulation.
- For system and procedure packs being placed on the market in a sterile condition, covered by this certificate the audit by the notified body of the quality management system was limited to the aspects required under article 22(3) of the regulation.
- For Custom Made Class III implantable device the certification only relates to the Quality management system. Technical documentation assessment and issuance of EU Technical Documentation Assessment Certificate does not apply.