

Către Agenția Medicamentului  
și Dispozitivelor Medicale

**NOTIFICARE**

pentru înregistrarea dispozitivelor medicale în Registrul de stat  
al dispozitivelor medicale

nr. 6 din 13.10.2023

Solicitantul **FCPC „DataControl” S.R.L.**, cu sediul **mun. Chișinău, str. N. Testemițanu, 17/6**, tel./fax: 022 27 37 12, e-mail: [contact@datacontrol.md](mailto:contact@datacontrol.md), solicit înregistrarea în Registrul de stat al dispozitivelor medicale a următoarelor categorii și tipuri de dispozitive medicale pentru introducerea și punerea la dispoziție pe piață a:

**Product: PTA Dilatation Catheter (OTW)**

**Model: Tercross PTA Dilatation Catheter (OTW)**

BD-T1220P4E	BD-T20150P4E	BD-T25150P4E	BD-T30150P4E	BD-T35150P4E
BD-T1520P4E	BD-T20200P4E	BD-T25200P4E	BD-T30200P4E	BD-T35200P4E
BD-T2040P4E	BD-T2540P4E	BD-T3040P4E	BD-T3540P4E	BD-T4040P4E
BD-T2080P4E	BD-T2580P4E	BD-T3080P4E	BD-T3580P4E	BD-T4080P4E
BD-T20120P4E	BD-T25120P4E	BD-T30120P4E	BD-T35120P4E	BD-T40120P4E
BD-T40150P4E	BD-T2080Q4E	BD-T2580Q4E	BD-T3080Q4E	BD-T3580Q4E
BD-T40200P4E	BD-T20120Q4E	BD-T25120Q4E	BD-T30120Q4E	BD-T35120Q4E
BD-T1220Q4E	BD-T20150Q4E	BD-T25150Q4E	BD-T30150Q4E	BD-T35150Q4E
BD-T1520Q4E	BD-T20200Q4E	BD-T25200Q4E	BD-T30200Q4E	BD-T35200Q4E
BD-T2040Q4E	BD-T2540Q4E	BD-T3040Q4E	BD-T3540Q4E	BD-T4040Q4E
BD-T4080Q4E				
BD-T40120Q4E				
BD-T40150Q4E				
BD-T40200Q4E				

Se anexează următoarele acte:

1. Declarație de Conformitate CE
2. Certificatul de Conformitate CE
3. Actul prin care producătorul își desemnează reprezentantul
4. Declarație pe propria răspundere.

Data **13.10.2023**

Semnătura \_\_\_\_\_

**Tabelul de recepționare a notificării**

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

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

Către  
Agenția Medicamentului  
și Dispozitivelor Medicale

## DECLARAȚIE PE PROPRIE RĂSPUNDERE

Solicitantul F.C.P.C. "DataControl" S.R.L., cu sediul în mun. Chișinău, str. N. Testemițanu 17/6, tel./fax: 022 27 37 12, e-mail: [contact@datacontrol.md](mailto:contact@datacontrol.md),

declar pe proprie răspundere, cunoscând prevederile art. 352<sup>1</sup>, Codul Penal al Republicii Moldova cu privire la falsul în declarații, că documentele și datele furnizate pentru notificarea dispozitivului medical:

**Product: PTA Dilatation Catheter (OTW)**

**Model: Tercross PTA Dilatation Catheter (OTW)**

BD-T1220P4E	BD-T20150P4E	BD-T25150P4E	BD-T30150P4E	BD-T35150P4E
BD-T1520P4E	BD-T20200P4E	BD-T25200P4E	BD-T30200P4E	BD-T35200P4E
BD-T2040P4E	BD-T2540P4E	BD-T3040P4E	BD-T3540P4E	BD-T4040P4E
BD-T2080P4E	BD-T2580P4E	BD-T3080P4E	BD-T3580P4E	BD-T4080P4E
BD-T20120P4E	BD-T25120P4E	BD-T30120P4E	BD-T35120P4E	BD-T40120P4E
BD-T40150P4E	BD-T2080Q4E	BD-T2580Q4E	BD-T3080Q4E	BD-T3580Q4E
BD-T40200P4E	BD-T20120Q4E	BD-T25120Q4E	BD-T30120Q4E	BD-T35120Q4E
BD-T1220Q4E	BD-T20150Q4E	BD-T25150Q4E	BD-T30150Q4E	BD-T35150Q4E
BD-T1520Q4E	BD-T20200Q4E	BD-T25200Q4E	BD-T30200Q4E	BD-T35200Q4E
BD-T2040Q4E	BD-T2540Q4E	BD-T3040Q4E	BD-T3540Q4E	BD-T4040Q4E
BD-T4080Q4E				
BD-T40120Q4E				
BD-T40150Q4E				
BD-T40200Q4E				

Sunt autentice și corespund realității

Alexandru Grabazei, director

Semnătura \_\_\_\_\_

Data: 13.10.2023



**Terumo Europe NV  
Emerging Market Division**

Researchpark Haasrode 1520  
Interleuvenlaan 40  
3001 Leuven, Belgium  
Tel.: +32 16 38 13 08  
Fax: +32 16 38 16 01

[www.terumo-europe.com](http://www.terumo-europe.com)

**To: Whom It May Concern**

Ref: 2023/007/IS/MI

Leuven, January 18, 2023

**Letter of Authorization**

We, begin company-manufacturer **Terumo Europe N.V. (Belgium)**;  
and being the European Authorized representative of company-manufacturer **Terumo Corporation, Terumo Medical Corporation, Terumo Clinical Supply and Terumo Medical Products (Hangzhou)**;  
and being the appointed distributor for products from the company-manufacturer **PendraCare, MicroVention Europe, MicroVention Inc and Kaneka Corporation**;

hereby appoint following company (hereinafter - "Company"):

**FCPC "DataControl" SRL**  
20 Melestiu Street, MD-2001,  
Chisinau, Republic of Moldova,

to be our official representative at the responsible authorities of the Republic of Moldova for registration, renewal, variation of registration etc. of following medical products and devices manufactured and/or distributed by us:

**Accuforce PTCA dilatation catheter (RX)**  
**Angio-Seal VIP Vascular Closure Device**  
**Azur Detachment Controller**  
**Azur Peripheral Coil System**  
**Climber Guiding Catheter**  
**Croserio RX PTA Balloon Dilatation Catheter**  
**Crosstella OTW PTA Balloon Dilatation Catheter**  
**Destination Guiding Sheath (Terumo Corporation and Terumo Medical Corporation)**  
**Eliminate Aspiration catheter**  
**FemoSeal Vascular Closure System**  
**Finecross MG Coronary Micro-Guide catheter**  
**Glidesheath Slender Hydrophilic Coated Introducer Sheath**  
**Heartrail II Guiding Catheter**  
**HydroPearl Compressible Microspheres for Embolisation**  
**LifePearl Drug-elutable microspheres for embolisation**  
**Metacross® OTW PTA Balloon Dilatation Catheter**  
**Metacross® RX PTA Balloon Dilatation Catheter**  
**Navicross Support Catheter**  
**Occlusafe Temporary Occlusion Balloon Catheter**  
**Outlook Angiographic Catheter**  
**Progreat Micro Catheter System (Terumo Corporation and Terumo Clinical Supply)**  
**Radifocus Glidecath Angiographic Catheter (Terumo Corporation and Terumo Europe)**

**Radifocus Guide Wire GT with Gold Coil**  
**Radifocus Guide Wire M** (Terumo Corporation and Terumo Europe)  
**Radifocus Guide Wire M Non-Vascular**  
**RADIFOCUS® Glidewire Advantage™**  
**RADIFOCUS® Glidewire Advantage™ Track**  
**Radifocus Obturator**  
**Radifocus Torque Device** (Terumo Corporation and Terumo Medical Products (Hangzhou))  
**Radifocus Vessel Dilator**  
**Radifocus OPTITORQUE Angiographic Catheter** (Terumo Corporation and Terumo Europe)  
**Radifocus Introducer II (Transradial Kit)**  
**Radifocus Introducer II**  
**Roadsaver Carotid Artery Stent**  
**Runthrough® NS Extension Wire PTCA Guide Wire**  
**Runthrough® NS PTCA Guide Wire**  
**Ryujin Plus PTCA dilatation catheter (RX)**  
**Senri® PTA Balloon Dilatation catheter**  
**Tercross® PTA Dilatation Catheter (OTW)**  
**Ryurei PTCA Dilatation Catheter**  
**TR Band Radial Artery Haemostasis Band**  
**Ultimaster Sirolimus eluting coronary stent system**  
**Ultimaster Tansei Sirolimus eluting coronary stent system**  
**Ultimaster Nagomi Sirolimus eluting coronary stent system**

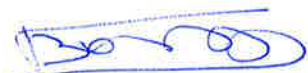
Hereby the Company is authorized to ensure that state registration (re-registration) of the abovementioned products is obtained and maintained in accordance with the legislation of Republic of Moldova.

For this purpose, the company can perform all acts, including but not limited: to submit, confirm, receive all necessary documents, including registration certificates, to reply to inquiries, questions or other communications from authorized institutions, after consultation with Regulatory department of Terumo Europe N.V, to conduct any field actions which may be necessary, in accordance with legislation of Republic of Moldova.

Registration certificates must be issued in the name of Terumo Europe N.V.

This authorization letter is valid for a period of 12 /twelve/ months from the date of issue, unless revoked earlier by Terumo Europe N.V.

For and behalf of Terumo Europe N.V.:



**Valérie Boydens**

Director Regulatory Affairs  
Terumo Europe NV



**TERUMO**  
TERUMO EUROPE N.V.  
Interleuvenlaan 40  
3001 LEUVEN, BELG

**EC Certificate**  
**Directive 93/42/EEC Annex II, excluding Section 4**  
**Full Quality Assurance System**  
**Medical Devices**

**Registration No.:** HD 60145252 0001

**Report No.:** 12031336 018

**Manufacturer:** Terumo Corporation  
44-1, 2-chome, Hatagaya  
Shibuya-Ku, Tokyo  
151-0072 Japan

**Products:** see attachement for products included

Replaces Approval, Registration No.: HD 60121893 0001

**Expiry Date:** 2024-05-26

The Notified Body hereby declares that the requirements of Annex II, excluding section 4 of the directive 93/42/EEC have been met for the listed products. The above named manufacturer has established and applies a quality assurance system, which is subject to periodic surveillance, defined by Annex II, section 5 of the aforementioned directive. For placing on the market of class III devices covered by this certificate an EC design-examination certificate according to Annex II, section 4 is required.

**Effective Date:** 2019-12-23

**Date:** 2019-12-23



Notified Body

*M. Aihara*  
M.Sc. M. Aihara

**TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg**  
TÜV Rheinland LGA Products GmbH is a Notified Body according to Directive 93/42/EEC concerning medical devices with the identification number 0197.

**TÜV Rheinland**  
**LGA Products GmbH**  
**Tillystraße 2, 90431 Nürnberg**

**Attachment to  
Certificate**


**Registration No.:** HD 60145252 0001  
**Report No.:** 12031336 023

**Manufacturer:** Terumo Corporation  
44-1, 2-chome, Hatagaya  
Shibuya-ku, Tokyo  
151-0072 Japan

**Products included:**

- Blood Bags
- Blood Donor Set with/without Blood Transfusion Filter
- Blood Transfusion Filter
- Intravenous Catheter
- Intravenous Administration Set
- Hypodermic Syringe
- Winged Needle
- Dental Needle
- Other Medical Needle
- Blood Administration Set
- Lancet

**Notified Body**

  
**M.Sc. M. Aihara**



**Date: 2020-10-23**

**TÜV Rheinland**  
**LGA Products GmbH**  
**Tillystraße 2, 90431 Nürnberg**

**Attachment to  
Certificate**

**Registration No.:** HD 60145252 0001  
**Report No.:** 12031336 023

**Manufacturer:** Terumo Corporation  
44-1, 2-chome, Hatagaya  
Shibuya-ku, Tokyo  
151-0072 Japan

**Products included:**

- Extra-corporeal Membrane Oxygenator
- Cardiopulmonary Bypass Arterial Line Blood Filter
- Heart-Lung Bypass Defoamer
- Cardiotomy Reservoir
- Cardiopulmonary Bypass Blood Reservoir
- Haemoconcentration Filter
- Centrifugal Pump
- Angiographic Catheter
- Balloon Dilatation Catheter
- Catheter Guide Wire
- Guiding Catheter
- Catheter Introducer
- Stents
- Extension Tube
- Temperature Control Unit for Heart-Lung Bypass System Module
- Infusion Pump
- Syringe Infusion Pump
- Clinical Electronic Thermometer
- Portable insulin infusion pump
- Portable insulin infusion administration set

**Notified Body**



**Date:** 2020-10-23

*M. Aihara*  
**M.Sc. M. Aihara**



## DECLARATION OF CONFORMITY

We, **TERUMO CORPORATION**

**44-1, 2-chome, Hatagaya, Shibuya-ku, Tokyo 151-0072, Japan**

being the manufacturer of:

**Tercross**

**PTA Dilatation Catheter (OTW)**

**Product : PTA Dilatation Catheter (OTW)**

declare that the above products of **Class IIa** are in conformity with the provisions of the EC Council Directive 93/42/EEC of 14 June 1993, as amended, concerning medical devices, and have been subject to the conformity assessment procedure laid down in Article 11, 2 and 11, 3(a) of the Directive, relating to the "Full quality assurance" set out in Annex II, and by certification of Annex II, excluding Section 4 under the supervision of TÜV Rheinland LGA Products GmbH (Registration No.: HD 60145252 0001), Tillystraße 2, 90431 Nürnberg Germany, as Notified Body authorized by the German Competent Authority and carrying the Notified Body No. 0197.

Authorized European Representative :

TERUMO EUROPE N.V.

Interleuvenlaan 40, 3001 Leuven, Belgium

Object of the declaration: see appendix A

Tokyo, February 10, 2020

(place and date of issue)



Toshio Nakashima

General Manager

Quality Assurance Department

TERUMO CORPORATION

Appendix A - List of Code Number Structure

B D - T □ □ □ □ □ □ □ □  
 1 2 3 4 5 6 7 8 9 10 11 12

Character number	Characters	Denotation
1-2	Product name	BD: PTA CATHETER
3	Destination	-: for export/ domestic use
4	Product name	T: Tercross (OTW)
5-6	Balloon diameter	12: 1.25 mm 15: 1.5 mm 20: 2.0 mm 25: 2.5 mm 30: 3.0 mm 35: 3.5 mm 40: 4.0 mm
7-(8), 9*	Balloon length	20: 20 mm 40: 40 mm 80: 80 mm 120: 120 mm 150: 150 mm 200: 200 mm
(9), 10*	Catheter length	P: 100 cm Q: 148 cm
(10), 11*	Adaptation wire	4: Wire adaptation of 0.014' '
(11), 12*	Place of destination	E: for domestic market/ export

\*:When balloon length is 2 digits, digit numbers are adapted (8)~(11)

When balloon length is 3 digits, digit numbers are adapted 9\*~12\*