



**Terumo Europe NV  
Emerging Market Division**

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**To: Whom It May Concern**

Ref: 2021/038/IS/MI

Leuven, 02 April 2021

**Letter of Authorization**

We, company-manufacturer **Terumo Europe N.V. (Belgium)**, with a manufacturing facility located at Interleuvenlaan 40, 3001, Leuven, Belgium, being a truly official representative of company-manufacturer Terumo Corporation (Japan) with manufacturing facilities located worldwide, 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 Evolution Vascular Closure Device**  
**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**  
**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**  
**Radifocus Glidecath Angiographic Catheter**  
**Radifocus Guide Wire GT with Gold Coil**  
**Radifocus Guide Wire M**  
**Radifocus Guide Wire M Non-Vascular**

**RADIFOCUS® Glidewire Advantage™**  
**RADIFOCUS® Glidewire Advantage™ Track**  
**Radifocus Obturator**  
**Radifocus Torque Device**  
**Radifocus Vessel Dilator**  
**Radifocus OPTITORQUE Angiographic Catheter**  
**Radifocus Introducer II (Transradial Kit)**  
**Radifocus Introducer II**  
**Roadsaver Carotid Artery Stent**  
**Runthrough® NS Extension Wire PTCA Guide Wire**  
**Runthrough® NS PTCA Guide Wire**  
**Ryuji Plus OTW PTCA dilatation catheter (OTW)**  
**Ryuji 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**

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 on 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.:

**Fien Aerts**

VP Regulatory & Vigilance  
Terumo Europe NV

  
**TERUMO**  
TERUMO EUROPE N.V.  
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3001 LEUVEN, BELGIUM