



EC Declaration of Conformity

We

CIVCO Medical Instruments Co., Inc. dba CIVCO and CIVCO Medical Solutions
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 Kalona, Iowa 52247 United States
 Phone: 319.248.6757
 Fax: 319.656.4451
 Single Registration Number (SRN): US-MF-000008229

declare under our sole responsibility that the medical imaging accessory product described hereafter...

TD 301B Manual Steppers
 as specified in master list ML 301B Manual Steppers
 EU UDI-DI: 0841436101TD301BLK

is in conformity with the general safety and performance requirements and provisions of Medical Device Regulation 2017/745 for Medical Devices,

as a Class I product following Rule 1 from Annex VIII of Medical Device Regulation 2017/745

and is subject to the procedure set out in Annex IX of Medical Device Regulation 2017/745.

Relevant Common Specifications:
 None

Kalona, Iowa USA, 26 May 2020

European Regulatory Representative:
 SRN: DE-AR-000005009
 MPS
 Medical Product Service GmbH
 Borngasse 20
 35619 Braunfels, Germany
 Phone: 49 6442 32370
 Fax: 49 6442 32578

Person Responsible for Regulatory Compliance			
Name: <u>Laurence Marshall Jr</u>			
Signature: <u><i>Laurence Marshall Jr</i></u> Date: <u>3/20/2023</u>			
Date	Rev	Description	Issued By
26 May 2020	1	Initial MDR Release	Kg
19 July 2021	2	Add Single Registration Number (SRN)	Kg
14 March 2023	3	Add EAR SRN	kg



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TD 301B Manual Steppers Intended Use Statement:

The equipment is intended to hold and manipulate ultrasound imaging probes and report position during prostate brachytherapy, cryotherapy, transperineal template-guided biopsy and/or fiducial marker placement procedures (including volume determination of the prostate gland) and/or the application of radionuclide source(s) into the body during brachytherapy.