

Declaration of Conformity

This European Declaration of Conformity is issued under the sole responsibility of the manufacturer.

MANUFACTURER								
Name of Company	Address	Address						
Mirion Technologies (Capintec), Inc.		7 Vreeland Road,	7 Vreeland Road,)			
		Florham Park, NJ	Florham Park, NJ 09732					
		USA	USA					
AUTHORIZED REPRESENTATIVE								
Name of Company	Address	SRN	Telephone/fax/email					
Atlantico Systems,	34 Oldfield, Galway,	IE-AR-000000208	+35391443609/					
Ltd.	Ireland		info@AtlanticoSystems.com					
PRODUCT IDENTIFICA	ATION							
Product Name		Product Code / C	Product Code / Catalog		Basic UDI-DI/EMDN			
		Number						
C CRC® -PC Smart Chamber HL Chamber		5130-30253	5130-30253		08599420061004Q/			
CRC® -PC Smart Chamber RpH Chamber		5130-33024	5130-33024		Z11029005			
Intended Purpose								
The CRC Dose Calibrators are intended to measure radiopharmaceuticals or brachytherapy								
sources prior to administration to patients.								

RISK CLASS FOR MEDICAL DEVICES					
Device	Standards				
Classification					
Class: Im Rule: 13, Annex VIII Conformity Assessment per Annex IX	IEC 60601-1-2 Ed. 4.0, (2014): Medical Electrical Equipment – Part 1-2: General requirements for basic safety and essential performance - Collateral Standard: Electromagnetic disturbances - Requirements and tests ANSI/AAMI ES60601-1:2005/A1:2012 Issued: 2012/08/20 Medical electrical equipment – Part 1: General requirements for basic safety and essential performance CAN/CSA-C22.2 No. 60601-1:14: Third Edition Issued: 2014/03/01 – Medical electrical equipment – Part 1: General requirements for basic safety and essential performance IEC 60601-1-6 Issued: 2013/10/29 Ed: 3.1 Medical electrical equipment – Part 1-6: General requirements for basic safety and essential performance – Collateral Standard: Usability				



IEC 62366:2014, Edition 1.1 Issued 2014/01/28 – Medical devices – Application of usability engineering to medical devices

IEC 62304:2006 + A1:2015 issued Medical device software – Software life cycle processes

NOTIFIED BODY			
Name of Company	ID	Address	Certificate Reference(s)
	Number		
Intertek Medical	2862	Torshamnsgatan 43	286201185244
Notified Body AB		Box 1103, SE-164 22	
		Kista, Sweden	

Autograph declares that the above-mentioned products meet the provision of the following EU legislation:

• MDR (EU) 2017/745 (LVFS 2003:11 as amended by LVFS 2009:07)

COMPANY REPRESENTATIVE:

Mary anne Yusko

Mary Anne Yusko

TITLE: Director, Regulatory Affairs

SIGNATURE:

DATE September 9, 2024

PLACE: Mirion Technologies (Capintec), Inc.

7 Vreeland Road, Florham Park, NJ 07932 USA

