EU-Declaration of Conformity

Manufacturer	Chromsystems Instruments & Chemicals GmbH		
Address	Am Haag 12 82166 Gräfelfing, Germany		
SRN (single registration number)	DE-MF-000010089		

Order No.	Device Description	EMDN Code		
Basic UDI-DI: 42503176000B5Q				
6000-B	HPLC Reagent Kit Catecholamines in urine	W01010499		
6000-BK	HPLC Basic Kit Catecholamines in urine	W01010499		
5001	Mobile Phase	W01019099		
5002	Mobile Phase	W01019099		
6033	Analytes Mix	W0101050399		
6009	Urine Calibration Standard	W0101050302		
6004	Internal Standard	W0101050399		
6055	Neutralisation Buffer	W01019099		
6006	Elution Buffer	W01019099		
6007	Sample Clean Up Columns	W01019099		
6100	HPLC column	W01019099		
Basic UDI-DI: 4250317102040502B				
0040	Endocrine Urine Control, Normal Range	W0101050207		
0050	Endocrine Urine Control, Pathological Range	W0101050207		

Device Intended Purpose	The Chromsystems assay "Catecholamines in urine" is an in vitro diagnostic medical device designed for professional users in clinical laboratories for the quantitative determination of noradrenaline (syn. norepinephrine), adrenaline (syn. epinephrine), and dopamine in human urine samples. Sample preparation is carried out manually, and analytic separation is done via high performance liquid chromatography with electrochemical detection (HPLC-ECD). The assay is intended to be used for patients in whom the urinary levels of noradrenaline, adrenaline and dopamine are of clinical importance, primarily as an aid to diagnosis and monitoring of suspected pheochromocytoma and paraganglioma.			
Risk Class	C, as per EU Regulation 2017/746, Annex VIII, Rule 3h			
GMDN Code	63823 "Multiple biogenic amine neurotransmitter IVD, kit, liquid chromatography"			
Notified Body	TÜV Süd Product Service GmbH Ridlerstraße 65, 80339 Munich, Germany	Identification No. 0123		

Conformity Assessment	Conformity assessment based on a quality management system and on assessment of technical documentation - Annex IX Chapters I and III						
Certificates issued	EU Quality management System Certificate (IVDR) No. V12 057136 0015 Rev01						
Declarations							
This EU declaration of conformity is issued under the sole responsibility of the manufacturer. The devices that are covered by the present declaration are in conformity with the In-Vitro Diagnostic Medical Devices Regulation (2017/746/EU) (IVDR).							
Following Common Specifications were considered as part of determining device conformity with the IVDR:							
Not applicable as no Commo	n Specifications exi	st for the concerned device.					
Additional information	Additional information n/a						
This EU declaration of conformity is issued by							
		Rall find					
Gräfelfing, December 18 th , 20 Michael Meier, Managing Dir		Gräfelfing, December 18 th , 20 Dr. Ralf Fischer, PRRC	24				
EU declaration of conformity	valid until:	December 17 th , 2029	Version: 1.0				