



# **Declaration of Conformity**

We herewith declare, in exclusive responsibility, that the instruments

Tissue Processor MTP, MTM, MTM II
Paraffin Wax Embedding Centre MPS/P1, MPS/P2, MPS/P, MPS/W, MPS/C, MPS/CX
Rotary Microtome CUT4062, CUT5062, CUT6062
aquatec
ergotec workplaces
Sledge Microtome cuttec S
Cryostat MTC, MEV, MEV plus, MNT
Staining Machine MSM, MAS
Linear Stainer cromatec I & II
Cytocentrifuge CS I / II
Cover Slipper MCS I / II
printtec s & c, printtec s high pressure, autoloader
Floating Bath / Slide Warmer: slidetec WATER, slidetec WATER, slidetec WATER s

are developed, designed and manufactured in compliance with the applicable directives

- Council Directive 2014/35/EU, (Low Voltage) and
- Council Directive 2014/30/EU (Electromagnetic Compatibility)
- Directive 2011/65/EU (restriction of the use of certain hazardous substances in electrical and electronic equipment)

including their amendments up to the date mentioned below.

The following harmonized standards were applied:

#### • EN 61010-1:2010

Safety requirements for electrical equipment for measurement, control and laboratory use - Part 1: General requirements

### • EN 61326-1:2013

Electrical equipment for measurement, control and laboratory use - EMC requirements -

Part 1: General requirements

## • EN 61000-3-2:2014

Electromagnetic compatibility (EMC)
Part 3-2: Limits - Limits for harmonic current emissions

#### • EN 61000-3-3:2013

Electromagnetic compatibility (EMC)
Part 3: Limits Section 3: Limitation of voltage fluctuations and flicker in low-voltage
supply systems for equipment with rated current ≤ 16 A

In addition, the following in-house standards were applied: DIN EN ISO 9001:2015; EN ISO 13485:2016

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Stefan Schock Managing Director

Tel.: +49 6131 95871-0 Geschäftsführer: Dr. W. Berg, S. Schock HRB 8504, Mainz WAT.-No.: DE232749604

Commerzbank AG Mainz BIC: COBADEFF550 IBAN: DE27 5504 0022 0200 043800