

EU Declaration of Conformity

Manufacturer:

ndd Medizintechnik AG

Address:

Technoparkstrasse 1

CH-8005 Zürich, Switzerland

Herewith we declare our sole responsibility for the declaration of conformity.

We declare under our sole responsibility that the medical devices:

Product name:

Spirette

EasyOne FlowTube

Product

Breathing mouthpieces

designation:

Product type:

Pulmonary Function Testing Devices

Model number:

Spirette:

2050-0 (basic unit)

2050-1, 2050-5, 2050-6, 2050-10, 2050-1GE, 2050-5GE, 2050-1HS, 2050-5HS

(packaging configurations) EasyOne FlowTube:

5050-0 (basic model)

5050-50, 5050-200, 5050-500, 5050-50MCK, 5050-200MCK (packaging

configurations)

Classified as:

Class IIa

according to annex IX of directive 93/42/EEC

meets all provisions of the directive 93/42/EEC which apply to it.

Applied standards:

See List of Applied Standards

Authorised

Johner Medical GmbH

Representative:

Office Frankfurt Speicherstrasse 16

60327 Frankfurt am Main, Germany

ndd Medizintechnik AG follows the procedure related to the EC declaration of conformity set out in Annex II of Directive 93/42/EEC which involves the intervention of the Notified Body:

TÜV SÜD Product Service GmbH, Notified Body 0123 Ridlerstrasse 65, 80339 Munich, Germany

Validity of the Declaration of Conformity corresponds to the validity of the EC Certificate G1 005204 0002, Rev. 01.

This declaration of conformity covers the products that have been released for production from the date of issuance of this Declaration of Conformity onward.

Andreas Senn,

Director Quality, Regulatory

Affairs & Clinical Affairs

Georg Harnoncourt,

Zurich, 25.May.2021