



# EU Declaration of Conformity

**MANUFACTURER:**

**Bio-Rad Laboratories, Inc.**

**ADDRESS:**

9500 Jeronimo Rd  
Irvine, CA 92618  
UNITED STATES OF AMERICA

**EUROPEAN AUTHORIZED REPRESENTATIVE:**

**Bio-Rad**

**ADDRESS:**

3 boulevard Raymond Poincaré  
92430 Marnes-la Coquette, France

**PRODUCT(S) NAME(S)**

Liquichek Sedimentation Rate Control

**CATALOG NUMBER(S):**

514, 514X, 515

**GENERIC DEVICE GROUP CODE:**

GMDN Nomenclature: 55972

**GENERIC DEVICE GROUP TERM:**

GMDN Nomenclature: Erythrocyte Sedimentation Rate (ESR) IVD, Control

We hereby declare that the above mentioned product(s) meet(s) the provisions of the following Directives

Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on *in vitro* Diagnostic medical devices

**CLASSIFICATION:**

ANNEX II-A

DEVICE FOR SELF TESTING

ANNEX II-B

OTHER DEVICE

**CONFORMITY ROUTE**

ANNEX III

ANNEX IV.3 Full Quality System

ANNEX IV.4 Product EC Design Examination

ANNEX V EC Type Examination

ANNEX VII Production Quality System

**NEW PRODUCT(S)** (Notification according to article 10 point 4)

YES

NO

**APPLICABLE HARMONIZED STANDARDS:** *Listed in the Bio-Rad QSD Quality Manual Normative References*

Signature

Irvine, CA

Issued in

22-Feb-19

Date

Vindeep Kohli

Name

Regulatory Affairs Manager

Function

