

BIO-RAD**BIO-RAD LABORATORIES
CLINICAL DIAGNOSTICS GROUP
EC DECLARATION OF CONFORMITY****MANUFACTURER:**

Bio-Rad Laboratories, QSD

ADDRESS:Bio-Rad Laboratories, QSD
9500 Jeronimo Rd,
Irvine CA 92618
Bio-Rad
3, Boulevard Raymond Poincare
Marnes-la-Coquette, France 92430**EUROPEAN AUTHORIZED
REPRESENTATIVE:****PRODUCT(S) NAME(S) and CATALOG NUMBER(S):**
Lyphochek® Immunoassay Plus Control

Catalog Number: 370, 371, 372, 373, 370X

CLASSIFICATION:

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- ANNEX II-A
-
-
- ANNEX II-B

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- DEVICE FOR SELF TESTING
-
-
- OTHER DEVICE

CONFORMITY ROUTE

-
- ANNEX III
-
-
- ANNEX IV.3 Full Quality System
-
-
- ANNEX IV.4 Product Design Examination

EC CERTIFICATE No.: 19347-1
Name of Notified Body : LNE/G-MED
Notified Body Identification No.: 0459
Expiration Date : 27.11.2013

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- ANNEX V Type Examination

EC CERTIFICATE No.:
Name of Notified Body :
Notified Body Identification No.:
Expiration Date:

-
- ANNEX VII Production Quality System

NEW PRODUCT(S) (Notification according to article 10 point 4) YES NO**GENERIC DEVICE GROUP CODE:**EDMS Nomenclature: 12-50-01-30
GMDN Nomenclature: None**GENERIC DEVICE GROUP TERM (EDMS Nomenclature): Multi Constituents Immunochemistry Controls**

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

APPLICABLE DIRECTIVE:Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on *in vitro* Diagnostic medical devices**APPLICABLE HARMONIZED STANDARDS:**EN 13641:2002
EN ISO 14971: 2007
EN ISO 15225:2000
EN 375:2001EN 980: 2008
EN 13485:2003
EN 13612:2002
EN 13640:2002

Signature

IRVINE CA USA

Issued in

12/2/10

Date

Vasif Vora

Regulatory Affairs Representative

Name

Function

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IBR-002-01, Ref 11, Effective Date 12-21-09

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