

## Section 4

### Declaration of Conformity

Manufacturer Bio-Rad Laboratories  
Address 9500 Jeronimo Road  
Irvine, CA 92618

European Bio-Rad  
Representative 3, Boulevard Raymond Poincare  
Marnes-La-Coquette FRANCE 92430

Products/  
Model Code Liquichek™ Urinalysis Control

Product	Catalog Number
Liquichek™ Urinalysis Control Bilevel	435
Liquichek™ Urinalysis Control Level 1	436
Liquichek™ Urinalysis Control Level 2	437

Classification Non-listed (In-Vitro Diagnostic Directive, Annex II)

We herewith declare that the above-mentioned products meet the provisions of the following EC Council Directives and Standards. All supplying documentation is retained under the premises of the manufacturer and the notified body.

### DIRECTIVES

#### General applicable directives:

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

#### Standards:

Harmonized Standards (published in the Official Journal of the European Communities) applicable to this product are: 98/79/EC: 1998, EN ISO 13485:2000, EN 980: 2003, EN 1441:1997, EN 928:1995, ISO 14971:2001, EN 13641:2002, EN 13612:2002, EN 13640:2002, 88/379/EEC: 2003, 67/548/EEC: 1967.

Notified Body: TÜV Rheinland Product Safety, GmbH  
World Headquarters  
Am Grauen Stein  
Konstantin-Wille-Straße 1  
D - 51105 Cologne

Signature   
Name Fri  
Date 4/20/03  
Position Regulatory Affairs Specialist