

# **Certificate of Analysis**

(RE TYPED COPY OF ORIGINAL CERTIFICATE)

Date:	2015-07-30		
Product:	Chlamydia Rapid Test Card		

PRODUCT CODE/ ART. NO.	LOT NUMBER	EXPIRATION DATE	QUANTITY	PHYSICAL APPEARANCE
004A170	004A170-1579	2017-07	10.000	conforms

## **PRODUCT TEST RESULTS**

ANALYSIS	SPECIFICATIONS	TEST RESULTS	
Negative specimens	negative	negative	
1.0 x 10 <sup>7</sup> org/ml	positive	positive	

Further Test Criteria:
Chromatography
Reaction time
Visual testing
Discoloration at C & T Line





# **PRODUCT MANUFACTURING**

The above listed product has been manufactured in accordance with relevant standard (EN ISO 13485:2012/AC:2012). No radioactive material of any kind is utilized in the product, or in the manufacturing of the product.

The above listed product does not contain any virus, reactive by-product of the same, or metabolic by-product of hepatitis A, B, C, or D, or of the antibodies for the human acquired immune deficiency syndrome (AIDS).

If this product is shipped as bulk, ulti med Products (Deutschland) GmbH assumes no responsibility for the packing and label claims.

Management Management





# **EC DECLARATION OF CONFORMITY**

## For the purposes of Annex IV to directive 98/79/EC for In-vitro-Diagnostic medical devices

Manufacturer: ulti med Products (Deutschland) GmbH, Reeshoop 1, D-22926 Ahrensburg

Product: Chlamydia Rapid Test Card

 Product ID:
 004A170

 Lot:
 004A170-1579

 Expiry:
 2017-07

 Quantity:
 10.000

Classification: IVD Directive 98/79 Annex II List B

Notified Body: TÜV Süd Product Service GmbH, Ridlerstraße 65, D-80339 Munich

No. Of Notified Body: 0123

Certificate-No: V1 13 07 55233 036

 Date:
 2013-07-25

 Valid from:
 2013-09-19

 Valid until
 2018-09-18

We hereby declare that the above named In-vitro-Diagnostic medical device are manufactured by us meets the essential requirements of Directive 98/79/EC and is suitable for the intended application. Any accompanying documentation is subject to the manufacture's responsibility.

#### Laws, rules and standards, applied

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

devices

EN ISO 13485: 2012/ Medical devices – Quality management systems – Requirements for regulatory purposes

EN ISO 14971:2012 Medical devices – Application of risk management to medical devices

EN 980:2008 Symbols for use in the labelling of medical devices

**EN ISO 15223-1: 2012** Symbols to be used with medical device labels, labelling and information to be supplied – Part 1:

General requirements

EN ISO 18113-1:2013 Medical devices – In vitro diagnostic medical devices - Information supplied by the manufacturer

(labelling) - Part 1: Terms, definitions and general requirements

**EN ISO 18113-2:2013** In vitro diagnostic medical devices - Information supplied by the manufacturer (labelling) - Part 2: In

vitro diagnostic reagents for professional use

**EN 13612: 2002** Performance evaluation of in vitro diagnostic medical devices

EN 13641: 2002 Elimination or reduction of risk of infection related to in vitro diagnostic reagent

EN 1041: 2008/A1:2013 Information supplied by manufacturer of medical devices

Ahrensburg, 2015-07-30

Mulo Gugel

ulti med Products (Deutschland) GmbH

Matthias W. Engel Managing Director

Reeshoop 1, D-22926 Ahrensburg (Germany) ##49 (0) 4102 - 80090; Fax ##49 (0) 4102 - 50082

Tax-ID-No. DE 154 23 11 80, Handelsregister Ahrensburg: HRB 3562

Geschäftsführer: Matthias W. Engel.