



Benannt durch/Designated by
Zentralstelle der Länder
für Gesundheitsschutz
bei Arzneimitteln und
Medizinprodukten
www.zlg.de
ZLG-BS-245.10.07



Product Service

EC Certificate

Full Quality Assurance System

Directive 98/79/EC on In Vitro Diagnostic Medical Devices (IVDD), Annex IV excluding (4, 6)
(List A and B and devices for self-testing)

No. V1 095123 0008 Rev. 04

Manufacturer:

Hangzhou AllTest Biotech Co., Ltd.

550#, Yinhai Street
Hangzhou Economic and Technological Development Area
310018 Hangzhou
PEOPLE'S REPUBLIC OF CHINA

**Product Category(ies): Products for determination of infection markers
tumor markers and products for self testing**

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for design, manufacture and final inspection of the respective devices / device families in accordance with IVDD Annex IV. This quality assurance system conforms to the requirements of this Directive and is subject to periodical surveillance. For marketing of List A devices an additional Annex IV (4) certificate is mandatory. All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: [www.tuvsud.com/ps-cert?q=cert:V1 095123 0008 Rev. 04](http://www.tuvsud.com/ps-cert?q=cert:V1_095123_0008_Rev_04)

Report no.:

SH221064A02

Valid from:

2022-04-05

Valid until:

2025-05-26

Date,

2022-04-05

Christoph Dicks
Head of Certification/Notified Body



EC Certificate

Full Quality Assurance System

Directive 98/79/EC on In Vitro Diagnostic Medical Devices (IVDD), Annex IV excluding (4, 6)
(List A and B and devices for self-testing)

No. V1 095123 0008 Rev. 04

Model(s):

Toxo IgG/IgM Rapid Test,
Rubella IgM Rapid Test,
CMV IgM Rapid Test,
ToRCH IgM Combo Rapid Test,
PSA Rapid Test,
PSA Qualitative Rapid Test,
Chlamydia Rapid Test,
Sperm Concentration Rapid Test,
SP-10 Male Fertility Rapid Test,
hCG Rapid Test,
Digital hCG Pregnancy Test
LH Rapid Test,
FSH Rapid Test,
Vaginal pH Rapid Test,
Ferritin Rapid Test,
TSH Rapid Test,
H.pylori Rapid Test,
Urinary Tract Infections Test,
FOB Rapid Test,
Vitamin D Rapid Test

Facility(ies):

Hangzhou AllTest Biotech Co., Ltd.
550#, Yin Hai Street, Hangzhou Economic and Technological
Development Area, 310018 Hangzhou, PEOPLE'S REPUBLIC OF
CHINA

EC Declaration of Conformity

Manufacturer:

Name: HANGZHOU ALLTEST BIOTECH CO., LTD.

Address: #550, Yinhai Street, Hangzhou Economic & Technological Development Area,
Hangzhou -310018, P.R. China

European Representative:

Name: MedNet EC-REP GmbH

Address: Borkstrasse 10, 48163 Muenster, Germany

Product Name: HCG (Human Chorionic Gonadotropin) Pregnancy Rapid Test (Urine)

Cat. No.: See Attachment 1

Analyte: Human Chorionic Gonadotropin in human Urine

Model: See Attachment 1

Classification: Self-testing of IVDD 98/79/EC

Conformity Assessment Route: IVDD 98/79/EC Annex IV

EDMA Code: 12 70 05 02 00

We, HANGZHOU ALLTEST BIOTECH CO., LTD., herewith declare that we are exclusively responsible for this declaration of conformity. We herewith declare that the above mentioned products meet the transposition into national law, the provisions of the following EC Council Directives and Standards. All supporting documentations are retained under the premises of the manufacturer.

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

Standard Applied: EN ISO 13485:2016, EN ISO 14971:2012, EN 13975:2003, EN ISO 18113-1:2011, EN ISO 18113-4:2011, EN 13612:2002/AC: 2002, EN ISO 17511:2003,
EN ISO 23640:2015, EN 13641:2002, EN 13532:2002, EN ISO 15223-1:2016

Notified body: TUV SUD Product service GmbH, Ridlerstrasse 65, 80339 Munich, Germany
(0123)

(EC) Certificate(s): V1 095123 0008 Rev.04

Expire date of the Certificate: 2025-05-26

Place, Date of First issue of DOC: in Hangzhou on 2016-09-14

The Date of Issue of DOC on 2022-04-05

Signature: 

Name: GAO FEI (Position: General Manager)

Hangzhou AllTest Biotech Co., Ltd.
#550, Yinhai Street,
Hangzhou Economic & Technological
Development Area,
Hangzhou -310018, P.R. China

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杭州奥泰生物技术股份有限公司 Hangzhou AllTest Biotech Co.,Ltd	文件号 Document No.: ZTC-QC-005-R-001
妇女健康类 COA The Women's Health COA	生效日期 Effective Date: 2019 年 06 月 21 日

Certificate of Analysis

Product Name: Pregnancy (hCG) Rapid Test Midstream (Urine)

Catalog No.: FHC-103H

Batch No.: HCG25020112

Quantity:2000PCS

Expiry Date:2028-01

Date of Sampling:2025-03-03

Date of Analysis:2025-03-03

Specification:25mIU/ml

Other information:/

QC Item		QC Criterion	QC Result	Conclusion
Physical	Appearance	Good	Good	Pass
Functional Performance	Positive Sample	Positive	100% Positive	Pass
	Negative Sample	Negative	100% Negative	Pass

Others	/
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Final QC Conclusion:	This batch of product met the QC Criteria.
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QC supervisor: Freeman.zhong

Date:2025.03.03





[Stampa](#) | [Scarica il dataset](#)

Elenco dei dispositivi medici

Criteri di ricerca:
Denominazione fabbricante:
Codice fiscale fabbricante:
Partita IVA / VAT number fabbricante:
Codice nazione fabbricante:
Denominazione mandatario:
Codice fiscale mandatario:
Partita IVA / VAT number mandatario:
Codice nazione mandatario:
Tipologia dispositivo:
Identificativo di registrazione attribuito dal sistema BD/RDM:
Codice attribuito dal fabbricante: **verify FHC-103H**
Nome commerciale e modello:
Classificazione CND:
Descrizione CND:
Normativa:
Classe CE (valida solo per dispositivi medici di classe, impiantabili attivi e IVD):

Elenco dispositivi individuati

Dati aggiornati al:02/03/2025

DISPOSITIVO MEDICO/ASSEMBLATO										FABBRICANTE/ASSEMBLATORE					
TIPOLOGIA	IDENTIFICATIVO			NOME COMMERCIALE	CND	NORMATIVA	CLASSE CE	DATA PRIMA PUBBLICAZIONE	DATA FINE		RUOLO AZIENDA	DENOMINAZIONE	CODICE FISCALE	PARTITA IVA/VAT NUMBER	NAZI
	DI	ISCRITTO AL	CODICE ATTRIBUITO DAL						IMMISSIONE	IN					
DISPOSITIVO	REGISTRAZIONE	REPERTORIO	FABBRICANTE/ASSEMBLATORE	E MODELLO						COMMERIO					
	BD/RDM														
					W0102160302							HANGZHOU			
				VERIFY SELF	- HCG - TEST		ST - Test				FABBRICANTE	ALLTEST			CI
Dispositivo	2750489	N	VERIFY FHC-103H	TEST	RAPIDI E	D.L.vo	autodiagnostici					BIOTECH CO.,			
				GRAVIDANZA	"POINT OF	332/2000	(non inclusi			25/02/2025		LTD.			
				CARE"			nell'al. II)								
											MANDATARIO	MEDNET EC-REP		DE126042714	D
											GMBH				

FARMADATI ITALIA Srl

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parafarmaco@farmadati.it

L'Ufficio PARAFARMACO è a Vs disposizione dal lunedì al venerdì (8:30 -18:30) per: attribuzione codici paraf, aggiornamento anagrafica prodotti e prezzi e consulenza e informazioni.

Nella Tabella sono riportati i codici base 10 e base 32 attribuiti ai prodotti per l'elaborazione del Barcode tipo 39 in base 32.

N.B. nel codice base 32 non sono utilizzabili le lettere: A,E,I,O. I codici paraf notificati con il presente modulo sono univoci e validi per tutto il territorio nazionale. La variazione della grammatura o della descrizione del prodotto, comporta l'attribuzione di un nuovo paraf.

Codice base 10	Codice base 32	Codice a barre	Codice EAN	Descrizione prodotto	Ditta	Codice articolo (ditta)	Iva	Prezzo al pubblico indicativo	Data prezzo al pubblico	Data inizio commercio
950086367	WB2BQZ		6936983170451	VERIFY HCG GRAVIDANZA SELFTEST	VERIFY Srls	FHC-103H	22	9,90	01/12/2024	01/12/2024