

DECLARATION OF CONFORMITY

Date of issue	Not applicable	Valid until	Not applicable
CE Assessment certificate	This product has been assessed on conformity according to the essential requirements as listed in Annex I of the European Directive 98/79/EC on in vitro diagnostic medical devices (IVDD). Certification by a Notified Body is not required.		
Certificate no.	Not applicable		

Product code	IQP-396	Product name	HITA/ert™ Kit
EDMA code	130204.90	EDMA term	Other Primary Hemostasis/Platelet Factors
EMDN code Level 4	W010302	EMDN term	Haemostasis Reagents (Coagulation)
GMDN code	61512	GMDN term	Platelet factor 4-heparin complex total antibody IVD, kit, agglutination, rapid
Basic UDI - DI	87179530223IQP-396NX		
Intended use	Immune heparin induced thrombocytopenia (HIT) is a distinct syndrome in which laboratory detection of the pathological HIT antibodies is diagnostically useful. The HITA/ert™ Kit, a qualitative, rapid, non-automated and flow cytometric platelet activation assay, which detects antibodies based on their characteristic platelet-activating properties in venous blood samples of patients suspected of HIT. This HITA/ert™ Kit should be used as a screening test and only be executed and interpreted by well-trained and authorized laboratory technicians. The results should always be used in conjunction with clinical findings and other serological tests.		

Manufacturer name and Address	IQ Products BV, Rozenburglaan 13a, 9727 DL Groningen, The Netherlands		
Single Registration number (SRN)	NL-MF-000010812		
Competent Authority	Dutch Health Care Inspectorate CIBG, registration no. NL-CA002-24457		
Harmonized Standards Referenced	<ul style="list-style-type: none"> • EN ISO 13485:2016 • EN 13641:2002 • EN 13612:2002 • EN 13975:2003 • EN ISO 14971:2012 	<ul style="list-style-type: none"> • EN ISO 18113-1:2011 • EN ISO 18113-2:2011 • EN ISO 23640:2015 • EN ISO 15223-1:2021 	
Other regulations/standards by which the product is regulated	Regulation (EC) No. 1272/2008; Regulation (EU) No. 453/2010; Regulation (EU) No. 2015/830.		

This declaration of conformity is issued under the sole responsibility of IQ Products BV.

- (1) The product meet the provisions of the Directive 98/79/EC on in vitro diagnostic medical devices which apply to the products;
- (2) The HITAlert™ kit is placed on the market for the first time in 2012;
- (3) The product concerned is produced at the manufacturing facility of IQ Products B.V. in conformance with the design control procedure requirements as specified in Directive 98/79/EC and the records are available for review;
- (4) The product comply with all essential requirements of the Directive 98/79/EC on in vitro diagnostic medical devices;
- (5) IQ Products BV has a Quality System in place based on ISO 13485:2016. The Quality System has been certified and the certificate has been issued by Notified Body DEKRA Certification B.V. (Identification number 0344) on February 01, 2022 and expires on February 01, 2025;

We hereby certify that as of the date of this declaration, the product described above conform with the provisions of the European Directive 98/79/EC on in vitro diagnostic medical devices (IVDD). All supporting documentation is retained at IQ Products BV.

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Date of issue: 2 February 2022

Place: Groningen, the Netherlands

Signature: 

Name: Christine M. Kebbedies

Function: Manager QA & RA

Company: IQ Products BV.