

## **Declaration of Conformity**

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according to Directive 98/79/EC, on in vitro diagnostic medical devices						
Maker	Getein Biotech, Inc.					
(Name, Address)	No. 9 Bofu Road, Luhe District, Nanjing, 211505, China					
Authorized Representative (Name, Address)	Lotus NL B.V. Koningin Julianaplein 10, 1e Verd, 2595AA, The Hague, Netherlands.					
Medical device	Description		FIA8000 Quantitative Immunoassay Analyzer FIA8600 Quantitative Immunoassay Analyzer Cardiac Troponin I Fast Test Kit One Step Test for cTnl (Colloidal Gold) cTnl Rapid Test (Colloidal Gold Assay) One Step Test for NT-proBNP (Colloidal Gold) One Step Test for NT-proBNP/cTnl (Colloidal Gold) One Step Test for CK-MB/cTnl/Myo (Colloidal Gold) One Step Test for hs-CRP+CRP (Colloidal Gold) One Step Test for D-Dimer (Colloidal Gold) One Step Test for D-Dimer (Colloidal Gold) One Step Test for PCT (Colloidal Gold) One Step Test for β2-MG (Colloidal Gold) One Step Test for MAIb (Colloidal Gold) One Step Test for MAIb (Colloidal Gold) One Step Test for MAIb (Colloidal Gold) One Step Test for Cysc (Colloidal Gold) One Step Test for HCG+β (Colloidal Gold) One Step Test for PCT/CRP (Colloidal Gold) One Step Test for PCT/CRP (Colloidal Gold) One Step Test for CK-MB/cTnl/H-FABP (Colloidal Gold) One Step Test for CK-MB/cTnl/H-FABP (Colloidal Gold) One Step Test for K-MB/cTnl (Colloidal Gold) One Step Test for TA/T3 (Colloidal Gold) One Step Test for TSH (Colloidal Gold) One Step Test for TSH (Colloidal Gold) One Step Test for TSH (Colloidal Gold) One Step Test for T3 (Colloidal Gold) One Step Test for T4/T3 (Colloidal Gold) One Step Test for T6OB (Colloidal Gold) One Step Test for T6OB (Colloidal Gold) One Step Test for FOB (Colloidal Gold) One Step Test for FOB (Colloidal Gold) One Step Test for SAA (Colloidal Gold) One Step Test for FA pylori (Colloidal Gold) One Step Test for SAA (Colloidal Gold)			

PCT Fast Test Kit (Immunofluorescence Assay) β2-MG Fast Test Kit (Immunofluorescence Assay) mAlb Fast Test Kit (Immunofluorescence Assay) NGAL Fast Test Kit (Immunofluorescence Assay) CysC Fast Test Kit (Immunofluorescence Assay) CK-MB Fast Test Kit (Immunofluorescence Assay) CK-MB/cTnl Fast Test Kit (Immunofluorescence Assay) HCG+β Fast Test Kit (Immunofluorescence Assay) HbA1c Fast Test Kit (Immunofluorescence Assay) PCT/CRP Fast Test Kit (Immunofluorescence Assay) CK-MB/cTnl/H-FABP Fast Test Kit (Immunofluorescence Assay) H-FABP Fast Test Kit (Immunofluorescence Assay) 25-OH-VD Fast Test Kit (Immunofluorescence Assay) TSH Fast Test Kit (Immunofluorescence Assay) T3 Fast Test Kit (Immunofluorescence Assay) T4 Fast Test Kit (Immunofluorescence Assay 25-OH-VD Fast Test Kit (Immunofluorescence Assay) FOB Fast Test Kit (Immunofluorescence Assay) H. pylori Fast Test Kit (Immunofluorescence Assay) SAA Fast Test Kit (Immunofluorescence Assay) LH Fast Test Kit (Immunofluorescence Assay) FSH Fast Test Kit (Immunofluorescence Assay) AMH Fast Test Kit (Immunofluorescence Assay) PRL Fast Test Kit (Immunofluorescence Assay) **CK-MB Control** cTnl Control Myo Control NT-proBNP Control **D-Dimer Control CRP Control PCT Control** β2-MG Control mAlb Control NGAL Control CysC Control H-FABP Control HbA1c Control HCG+B Control CK-MB/cTnl/Myo Control CK-MB/cTnl Control NT-proBNP/cTnl Control **TSH Control** T4/T3 Control T3 Control T4 Control Others Classification of products according to directive Batch/serial No. Type, production term (if applicable)

	EN ISO 14971:2012	EN ISO 23640:2015	EN ISO 13485:2016
Applicable	EN 13612:2002	EN ISO15223-1:2012	EN ISO 18113-2:2011
coordination	EN 1041:2008	EN ISO 18113-1:2011	EN ISO 18113-3:2011
standards:	IEC 61010-1:2010	IEC 61010-2-081:2015	IEC 61010-2-101:2015
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Signatory representative declares herein the above mentioned device meets the basic requirements of the European Parliament and the Council's in vitro diagnostic medical devices directive: 98/79/EC Annex III. This declaration of conformity is based on European Parliament and the Council's 98/79/EC directive Annex III. The compiled technical file and quality system document according to 98/79/EC directive Annex III are testified and the quality system certificate has issued by TÜV Rheinland (Shanghai) Co., Ltd.

General Manager: Enben Su

(place and date of issue)

(name and signature or equivalent

marking of authorized person)

