

## EC Declaration of Conformity (Directive 98/79/EC)

**Manufacturer:**

Chengdu Polytech Biological Technology  
Co.,Ltd

**Address:**

No.1 Tianhe Road, Western Zone of Chengdu  
High and New Tech, Sichuan, China.

**European Representative:**

Renault-Petersen Limited

**Address:**

5 Bankside, Hanborough Business  
Park, Witney OX29 8LJ UK

**Product Details:** SMT100 Chemistry Reagent

**Classification:** Other/General

**Conformity**

**Assessment Route:** Annex III without section 6

We herewith declare that above mentioned products meet the provisions of the Council Directive 98/79/EC for in vitro medical devices. All supporting documentation is retained under these premises and/or the premises of manufacturer's subcontractors.

### DIRECTIVE

**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 (IVDD 98/79/EC)

**Standard Applied:**

Standards list for which documented evidence of compliance can be provided as attachment.

**Quality assurance system standards:** EN ISO 13485: 2012; EN ISO 13485: 2012/AC: 2012(Certification number: SX 60079107 0001)

**Place, Date of Issue:** Chengdu, China, February 08, 2017

**Signature:****Name/Position:**

  
Peng Ran/ Managing Director



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### Applied Standard List

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Standard	Title
EN ISO 13485:2012	Medical devices-Quality management
EN ISO 13485:2012/AC:2012	systems-Requirements for regulatory purposes
EN ISO 14971:2013-04	Medical devices-Application of risk management to medical devices
EN ISO 15223-1:2016-11	Medical devices-Symbols to be used with medical device labels, labelling and information to be supplied-Part 1: General requirements
EN ISO 18113-1:2013-01	In vitro diagnostic medical devices-Information supplied by the manufacturer(labelling)-Part 1: Terms, definitions and general requirements
EN ISO 18113-2:2013-01	In vitro diagnostic medical devices-Information supplied by the manufacturer(labelling)-Part 2: In vitro diagnostic reagents for professional use
EN 13612:2002-08	Performance evaluation of in vitro diagnostic medical devices
EN 13640:2002	Stability testing of in vitro diagnostic reagents
EN 62366:2016-05; VDE 0750-241:2016-05	Medical devices-Application of usability engineering to medical devices

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**SMT-100 Chemistry Reagent types list:**

<b>The Types of Biochemistry Reagent Disc</b>	
<b>14 Conventional Parameters</b>	TC,ALT,AMY,ALB,ALP,GLU,GGT,UA,AST,TBIL,UREA,CREA, TG,TP
<b>13 Health Check Parameters</b>	ALT,ALB, AST, AMY, CA, CREA, GLU, TBIL, CK,TG,UREA, PHOS, TP, GLOB*
<b>10 liver Function</b>	ALT,TBA,AMY,ALB,ALP.GGT,AST,TBIL,CHE,TP
<b>8 Kidney Function</b>	Co2, CA,ALB,GLU,UA,P,UREA,CREA
<b>7 Electrolyte</b>	K+, Na+, Cl, CA, CO2, Mg, P
<b>6 Lipid Parameters</b>	TC,ALT,DHL,LDL*,GLU,AST,TG

