

# Document Title Declaration of Conformity for Diagnostic Medical Devices (IVDD)

In Vitro Diagnostic Medical Devices (IVDD)— Clearblue Digital Ultra Early Pregnancy Test - CB15

> Owner Regulatory Affairs

Document no. DC-0277

## **Declaration of Conformity Certificate**

Manufacturer:

SPD Swiss Precision Diagnostics GmbH

47, Route de Saint-Georges 1213 Petit-Lancy, Geneva

**SWITZERLAND** 

Product:

**Clearblue Digital Ultra Early Pregnancy Test** 

(Variants listed in Appendix 1)

Product Code:

**CB15** 

Device Classification:

Self - Test

Conformity

Annex IV (excluding sections 4 and 6) of the Directive:

Assessment Procedure:

98/79/EC on In Vitro Diagnostic Medical Devices

EC Certificate No:

98/79/EC IVDD 099371 0009

EC Certificate Expiry Date:

2025-05-26

Notified Body:

TÜV SÜD Product Service GmbH, Ridlerstraße 65, D-80339

München, Germany. Notified Body Number: 0123

We hereby declare that the above-mentioned product meets the provisions of the Council Directive 98/79/EC for In Vitro Diagnostic Medical Devices as transposed into National Legislation.

This declaration is issued under the sole responsibility of the manufacturer stated above.

For and on behalf of SPD Swiss Precision Diagnostics GmbH

Kim French

Position: Regulatory Director

Place of Issue: SPD Development Company Limited

Clearblue Innovation Centre, Stannard Way, Priory Business Park,

Bedford, MK44 3UP, UNITED KINGDOM

Date: 25may 2022



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### **Appendix 1** Variant Codes

Pouched Device Code	Description
CB15001	Novus Pouched Device-Symbolic
CB15002	Novus Pouched Device-English
CB15003	Novus Pouched Device-German
CB15004	Novus Pouched Device-Spanish
CB15005	Novus Pouched Device-French
CB15006	Novus Pouched Device-Dutch
CB15007	Novus Pouched Device-Italian
CB15010	Novus Pouched Device-Swe /Fin



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Pre Design Freeze	Post Design Freeze□	Non Project ⊠

### 1 Summary

File note to accompany DC-0277 - Clearblue Digital Ultra Early Pregnancy Test - CB15.

#### 2 Detailed Information

#### 2.1 Details of Change

To provide the statement that the EU Authorised Representative for the device named above with the European Union is:

Medical Device Safety Service GmbH (MDSS) has been appointed SPD's Authorised Representative for products placed on the market in EEA. Details for MDSS are:

Medical Device Safety Service GmbH Schiffgraben 41 Hannover 30175 Germany

#### 2.2 Declaration

We hereby declare that DC-0277 - Clearblue Digital Ultra Early Pregnancy Test - CB15 complies with all the provisions of the In Vitro Diagnostic Medical Device Directive 98/79/EC (IVDD).

Accordingly, the mark with Notified Body registration number 0123 will be placed on all production batches placed on the market in the European Union and EEA member states as defined in Section 3 of this document. The manufacturer retains all supporting documentation. The declaration of conformity is issued under the sole responsibility of the manufacturer.

This is not a significant change according to MDCG-20022-6 - Guidance on significant changes regarding the transitional provision under Article 110(3) of the IVDR.

### 3 Approval

For and on behalf of SPD Swiss Precision Diagnostics GmbH

Name: Kim FRENCH

Signature. Vinanch

Position: REGULATORY DIRECTOR

Date: 30NOV2022



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### 4 Revision History

Revision	Author	Reason for issue
03	Tosin Akisanya	Changes to make cover notes specific to DoC
02	Kate Sharman	Update to include the 'Declaration' section and definition of EEA
01	Kate Sharman	New Release