


Declaration of conformity				 ALBYN MEDICAL Smart Medical Group			
Description	EC Declaration of Conformity Hermes			Document	HE14.1_7		
				Revision	7		
Report Number		Date	05/01/2018	Sheet	1	OF	2

EC Declaration of Conformity

EC Declaration of Conformity Annex II (excluding section 4) to Medical Devices Directive 93/42/EEC

Manufacturer: ALBYN MEDICAL S.L.

Manufacturer's Address: POLIGONO INDUSTRIAL CORDOVILLA calle D, nº 1, 31191 – CORDOVILLA – NAVARRA, SPAIN

Device/s: HERMES (Ref: 98600000)

Description: Urine Urodynamic system

EC Product Class: IIa Rule 10 (Active Device for diagnosis)

CLASSIFICATION OF ME EQUIPMENT: CLASS I Medical equipment, externally powered, Type BF applied part.

Albyn Medical S.L. declares that device HERMES with reference 98600000 conforms to the relevant provisions of the EC Council Directive 93/42/EEC and the amendment 2007/47/EC and is in accordance with Annex II (excluding section 4), as implemented by the European Union's Medical Devices Regulations, as verified by our Notified Body:

**SGS United Kingdom Limited (Notified Body No. 0120)
202B Worle Parkway, Weston-super-Mare, BS22 6WA UK**

Standards:

Safety:

IEC 60601-1: 2005 + CORR. 1 (2006) + CORR. 2 (2007) third edition

EMC:

UNE-EN 60601-1-2:2008 + Corr:2010 (Medical Electrical Equipment)

Emission:

UNE-EN 55011:2011+A1:11 (conducted and radiated emission)

UNE-EN 61000-3-2:2006 + A1:10 + A2:10 (Harmonics).

UNE-EN 61000-3-3:2009 (Voltage fluctuations).

Immunity:

UNE-EN 61000-4-2:10

UNE-EN 61000-4-3:07 + A1:08 + A2:11

UNE-EN 61000-4-4:05+A1:10

UNE-EN 61000-4-5:07


UNE-EN 61000-4-6:09

UNE-EN 61000-4-8:11

UNE-EN 61000-4-11:05

Albyn Medical S.L. confirms that no other application has been lodged with another Notified Body for the same device related Quality Management System.

Albyn Medical S.L. agrees to develop, implement and maintain a formally-recognized Quality Management System to ensure continued adequacy and efficacy.

Declaration of conformity				 ALBYN MEDICAL Smart Medical Group			
Description	EC Declaration of Conformity Hermes			Document	HE14.1_7		
				Revision	7		
Report Number		Date	05/01/2018	Sheet	1	OF	2

Albyn Medical S.L. agrees to develop, implement and maintain a documented post-production experience monitoring process, including the notification of reportable events under the European Medical Device Vigilance System Guidelines.

Albyn Medical S.L. confirms that no medicinal products/drugs are incorporated in any devices covered by the Device Schedule.

Albyn Medical S.L. agrees to inform the appointed Notified Body of any planned or unplanned substantial change to the Quality Management System.

Albyn Medical S.L. agrees to inform the appointed Notified Body of any planned or unplanned significant change to the Device Schedule, including significant design change to devices.

Accessories included:

00004326 Puller for SmartDyn

00002785 Reusable pressure transducers

98800000/00004442 flowscale

Signed by the Albyn Medical S.L. designated representative:



Gonzalo Buil
Quality Manager
5th of January 2018