

Signature:

CU Medical Systems, Inc.

No. of Document: DOC-EU-HD(Rev.1)

Declaration of Conformity

This declaration of conformity	is issued under the sole responsibility of the manufacturer.
Manufacturer:	CU Medical Systems, Inc.
(Address based on land lot numbers)	Donghwa Medical Instrument Complex 1647-1 Dongwha-ri, Munmak-eup, Wonju-si, Gangwon-do, 220-801 Republic of Korea
(Address based on road names and building numbers)	130-1, Donghwagongdan-ro, Munmak-eup, Wonju-si, Gangwon-do Republic of Korea
	Tel: +82 33 747 7657 Fax: +82 33 747 7659
EU Authorized	Medical Device Safety Service, GmbH
Representative:	Schiffgraben 41, 30175 Hannover, Germany
Notified Body:	Nemko AS
	Gaustadalléen 30, N-0373 Oslo, Norway
	Identification no. 0470
Type of Product:	Defibrillators
Model No.:	CU-HD1
Classification:	Class IIb, according to Rule 9 of Annex IX of Directive 93/42/EEC
EU Directive(s):	93/42/EEC concerning medical devices, as amended by 2007/47/EC
	pove mentioned medical device(s) is(are) in conformity with applicable DIRECTIVE 93/42/EEC concerning medical devices as amended by
Date of Issue:	August 30, 2016
Signature:	HaRok Na, Chief Executive Officer