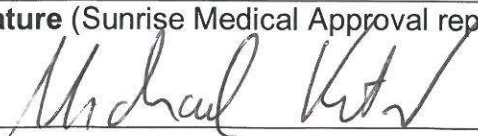


EC DECLARATION OF CONFORMITY

Declaración de Conformidad CE

Company: <i>Campania:</i>	Sunrise Medical S.L. Polígono Bakiola, 41 48498 Arrankudiaga Vizcaya / ESPAÑA
Product: <i>Producto:</i> (May include accessories) <i>(Incluye accesorios)</i>	SUNLIFT MICRO (MICRO) SUNLIFT MINI (G130R) SUNLIFT MIDI (G150) SUNLIFT MAJOR (G175)
We, Sunrise Medical declare under our sole responsibility that the product(s) to which this declaration relates, is a class 1 device, and is in conformity with the requirements of EC Council Directive for Medical Devices 93/42/EEC of 14 June 1993 amended by 2007/47/EEC of 21 March 2010. <i>Por la presente declaramos que los productos arriba mencionados, son clase I, y que son conformes a los requisitos de la Directiva de dispositivos Médicos 93/42/CEE, modificada por la Directiva 2007/47/CE de 21 de marzo de 2010</i> This was verified with conformity evaluation procedures according to Medical Device Directive Annex VII. <i>It ha sido verificado con conformidad a los procedimientos de evaluación acorde al anexo VII de la directiva de productos sanitarios.</i> Sunlift hoist are in compliance with the standard EN ISO 10535:2007. Hoists for the transfer of disabled persons – Requirements and test methods	

Michael Kutzer- Director European PDM and R&D	B	18/12/2015
Approval Name and Function <i>Nombre y Función</i>	Revision <i>Revision</i>	Approval Date <i>Fecha de aprobación</i>

Signature (Sunrise Medical Approval representative) <i>Firma</i> 	 SUNRISE MEDICAL. www.SunriseMedical.de	Sunrise Medical GmbH & Co. KG Kahlbachring 2-4 D-69254 Malsch/Heidelberg Telefon +49 7253 980 0 Telefax +49 7253 980 111
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