

EC Declaration of Conformity

PRODUCT IDENTIFICATION		Platelet Storage Products		
Product name		Model/number		
Platelet Incubator		PC100i, PC100h, PC900i, PC900h, PC1200i, PC1200h, PC2200i, PC2200h, PC3200i, PC3200h, PC4200i, PC4200h, PC100-Pro, PC900-Pro, PC1200-Pro		
MANUFACTURER				
Name of company Address			Representative	
Helmer Inc. 14400 Be Boulevard Noblesvil 46060 USA			Renee Schultz	
REGISTRATION INFORMATION				
Notified Body ID#	Certificate Number			
BSI 2797	CE 544457	CE 544457		
AUTHORIZED REPRESENTATIVE				
Name of company Address			Telephone/email	
Emergo Europe CONFORMITY ASSE	Prinsessegr 2514 AP Th The Netherl	e Hague	+31.70.345.8570 - phone +31.70.346.7299 - fax europe@emergogroup.com	
Device classification	.OCIVILIY I	Poute to co	mpliance	
Class IIa Rule 2		Annex V of MDD 93/42/EEC Council Directive as amended by 2007/47/EEC		

Helmer Inc. declares that the above-mentioned products meet the provision of the Council Directive 93/42/EEC for Medical Devices as amended by 2007/47/EEC and Directive 93/42/EEC, RoHS Recast Directive 2011/65/EU including the amendment to Annex II described in Commission Delegated Directive (EU) 2015/863.

COMPANY REPRESENTATIVE: Renee Schultz

TITLE: Director of Regulatory Affairs SIGNATURE:

DATE: 01/01/2021