



EC Design-Examination Certificate

Directive 93/42/EEC on Medical Devices, Annex II Section 4

No. Issued To: CE 664636 Terumo Medical Corporation 2101 Cottontail Lane Somerset New Jersey 08873 USA

In respect of:

Angio-Seal Vascular Closure Devices

BSI has performed a design examination on the above devices in accordance with the Council Directive 93/42/EEC Annex II Section 4 and Regulation 722/2012. The design conforms to the requirements of this directive and regulation. For marketing of these products an additional Annex II excluding Section 4 certificate is required.

For and on behalf of BSI, a Notified Body for the above Directive (Notified Body Number 0086):

Frank Lee, EMEA Compliance & Risk Director

First Issued: 20 January 2017

Date: 20 January 2017

Expiry Date: 19 January 2022

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Validity of this certificate is conditional on the quality system being maintained to the requirements of the Directive as demonstrated through the required surveillance activities of the Notified Body.

This certificate was issued electronically and is bound by the conditions of the contract.

Information and Contact: BSI, Kitemark Court, Davy Avenue, Knowlhill, Milton Keynes MK5 8PP. Tel: + 44 345 080 9000 BSI Assurance UK Limited, registered in England under number 7805321 at 389 Chiswick High Road, London W4 4AL, UK. A member of BSI Group of Companies.





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Supplementary Information to CE 664636

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Angio-Seal Vascular Closure Device

Model Number	Description		
610120	6F Angio-Seal STS-Plus	2	
610122	8F Angio-Seal STS-Plus		
610132	6F Angio-Seal VIP		
610133	8F Angio-Seal VIP		
C610136	6F Angio-Seal Evolution		
C610137	8F Angio-Seal Evolution		

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Certificate History

Date	Reference Number		Action	
20 January 2017	10167452	First Issue.		A Col

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