

3200 Lakeside Drive Santa Clara, CA 95054

Tel 408-845-3000 Fax 408-845-3333

DECLARATION OF CONFORMITY

Manufacturer:

Abbott Vascular

Address:

3200 Lakeside Drive

Santa Clara, California 95054, USA

Additional

Manufacturing Sites:

26531 Ynez Road

Temecula, California 92591, USA

Building PR-17, Road #2 km. 58.0

Cruce Davilla

Barceloneta 00617, Puerto Rico

Device Name:

HI-TOROUE PROGRESS PTCA Guide Wires

Device Classification:

Class III

GMDN Code:

35094 – Cardiac catheter guidewire, single-use

Classification Rationale:

The following Annex IX definition(s) apply to the HI-TORQUE PROGRESS PTCA Guide Wires for purposes of classifications: Per Rule 6, Annex IX, all surgically invasive devices intended for transient use are in Class IIa unless they are: Intended specifically to control, diagnose, monitor, or correct a defect of the heart or of the central circulatory system through direct contact with these parts of the heavy in which page they are in Class III.

parts of the body, in which case they are in Class III.

Authorized European Representative:

Abbott Vascular International BVBA

Park Lane

Culliganlaan 2B

1831 Diegem, Belgium



Model Numbers:

HI-TORQUE PROGRESS PTCA Guide Wires

Catalog Number	Tip Radiopacity	Wire Diameter	Wire Length
1011836	3	.014"	190
1011837	3	.014"	300
1011838	3	.014"	190
1011839	3	.014"	300
1011840	3	.014"	190
1011841	3	.014"	300
1011842	3	.014"	190
1011843	3	.014"	300
1011844	3	.014"	190
1011845	3	.014"	300

I, the undersigned, hereby declare that the medical devices specified above conform with the applicable *Essential Requirements* listed in Annex I and Annex II of EC Council Directive 93/42/EEC.

Directive 2006/42/EC on Machinery and directive 1989/686/EC on Personal Protective Equipment do not apply.

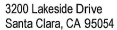
This declaration is supported by an EC quality system and design examination approval listed below.

Quality Management System - ISO 13485:2003, Certificate: FM72377, FM510125

EC Design Examination Certificate: CE 553292 Full Quality Assurance Certificate: CE 510108

Notified Body: British Standards Institution, a notified body authorized by the United Kingdom Competent Authority, Notified Body Identification Number 0086.

British Standards Institution Kitemark Court Davy Avenue Knowlhill Milton Keynes MK5 8PP United Kingdom





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This Declaration of Conformity expires upon revision or with the revision or expiration of any of the supporting certificates listed above.

This Declaration of Conformity is issued under the sole responsibility of the manufacturer.

Ву:	Slave	Date: _	8 /22/14
Susan Slane Divisional VP, Glo Abbott Vascular	bal Quality and Compliance		
Place of issue:	Temecula 8/72/14	Date of issue:_	8/72/14