

DECLARATION OF CONFORMITY

We, **TERUMO CORPORATION**
44-1, 2-chome, Hatagaya, Shibuya-ku, Tokyo 151-0072, Japan

being the manufacturer of:

RADIFOCUS Vessel Dilator

Product : Catheter Introducer

declare that the above products of **Class IIa** are in conformity with the provisions of Annex I and Annex II of the EC Council Directive 93/42/EEC of 14 June 1993, as amended, concerning medical devices, and have been subject to the conformity assessment procedure laid down in Article 11, 3(a) of the Directive, relating to the "Full quality assurance" set out in Annex II, under the supervision of TÜV Rheinland Product Safety GmbH (Registration No.: HD 60026344 0001), as Notified Body authorized by the German Competent Authority and carrying the Notified Body No. 0197.

Authorized European Representative :
TERUMO EUROPE N.V.
Interleuvenlaan 40, 3001 Leuven, Belgium

Tokyo, October 13, 2009
(place and date of issue)



Hiroshi Nakagomi
General Manager
Quality Assurance Department
TERUMO CORPORATION

 **TERUMO®**