

## ***EC Declaration of Conformity***

**Manufacturer:** Beekley Corporation  
**Address:** One Prestige Lane  
Bristol, CT 06010 USA

**Devices:** Order Codes: 101, 102, 103, 104, 105, 106, 107, 108, 110, 111, 112, 113, 114, 115, 117, 118, 119, 120, 121, 122, 123, 128, 132, 150, 151, 153, 156, 187, 190, 191, 209, 210, 211, 215, 252, 300, 301, 302, 309, 310, 311, 315, 316, 318, 319, 322, 323, 418, 500, 502, 505, 601, 602, 603, 607, 608, 610, 632, 650, 651, 653, 654, 677, 690, 691, 609, 652, 703, 710, 711, 712, 713, 732, 750, 751, 752, 754, 790

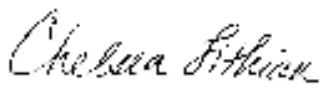
**Description:** Beekley Skin Markers, Beekley Specimen Radiography Devices, Protective Coverlets,  
**EC Product Class:** Class I (Annex IX, Rule 1)

Beekley Corporation declares that device/s Beekley Skin Markers listed on the attached Device Schedule conform to the relevant provisions of the EC Council Directive 93/42/EEC for Class I non-sterile (Annex IX, Rule 1), as implemented by the European Union's Medical Devices Regulations.

Beekley Corporation agrees to develop, implement and maintain a documented post-production experience monitoring process, including the notification of reportable events under the European Medical Device Vigilance System Guidelines.

Beekley Corporation confirms that no medicinal products/drugs are incorporated in any devices covered by the Device Schedule.

Beekley Corporation has appointed Emergo Europe as our EU Authorized Representative of: Molenstraat 15, 2513 BH The Hague, The Netherlands  
Tel: +31 70 345 8570 Fax: +31 70 346 7299 E-mail: [Europe@emergogroup.com](mailto:Europe@emergogroup.com)  
Signed by the Beekley Corporation designated representative:

Name:   
Chelsea Fithian

Title: International Sales & Marketing Manager

Date: January 14, 2010