

## EC Declaration of Conformity

We, the Manufacturer:

**Weaver and Company**  
**565 Nucla Way, Unit B**  
**Aurora, Colorado 80011**  
**United States of America**  
**1-800-525-2130**

declare under sole responsibility that the topical medical devices described below:

**Nuprep<sup>®</sup> Skin Prep Gel**  
**Ten20<sup>®</sup> Conductive Paste**

are medical devices subject to Council Directive 93/42/EEC of June 14, 1993, amended by Directive 2007/47/EC of September 5, 2007, concerning medical devices, are Class I devices according to the classification criteria of Annex IX of the directive, Rule 1;

are non-invasive medical devices not intended as sterile devices;

do not perform a measuring function;

are therefore eligible for and conform with the conformity assessment procedure described in Annex VII of the directive; and

meet the essential requirements of Annex I of the Council Directive.

In addition, the medical devices described above conform with Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances.

The initial date of the CE mark on Nuprep Skin Prep Gel and Ten20 Conductive Paste was June 1, 1998.

Authorized Representative:

**Emergo Europe**  
**Molenstraat 15**  
**2513 BH The Hague**  
**The Netherlands**  
**+31 70 345 8570**



Chris Cooper, CEO  
Management with Executive Responsibility  
Weaver and Company

20 May 2014

Date