



Declaration of Conformity

As Legal Manufacturer
We, 3M Company, 3M Health Care,
3M Center, 2510 Conway Ave, Bldg. 275-5W-06
St. Paul, MN 55144 USA

hereby declare under our sole responsibility
that the CE marked products to which this declaration relates ,

3M™ Tegaderm™ Alginate, High Integrity Alginate Dressing
Product model numbers:
90110, 90112, 90114, 90120

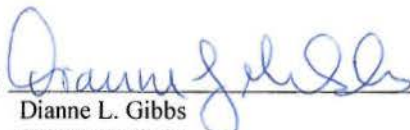
is classified,
per rule 4 of Annex IX of the Medical Device Directive 93/42/EEC, as amended per 2007/47/EC
as a Class IIb device
and

is in accordance with Annex II of Directive 93/42/EEC, as amended per 2007/47/EC
on the approximation of the laws of the European Union Member States concerning medical devices.

In addition, we declare that the above mentioned devices fulfill the applicable provisions of the Directive
93/42/EEC, as amended per 2007/47/EC

This declaration is made on the basis of the quality assurance certificate
CE Certificate 02242 delivered by BSI, 2797

EU Representative Address
3M Deutschland GmbH
Health Care Business
Carl-Schurz-Str. 1
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Signature: 
Dianne L. Gibbs
3M Health Care
Division Regulatory Affairs Manager
Medical Solutions Division

Date: 1 November 2019