Declaration of Conformity Tensoplus®

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We,

BSN Medical SAS Rue du Millenaire / CS 90022 72320 VIBRAYE France (SRN: FR-MF-000000598)

hereby declare under our own responsibility, that this product family complies with the applicable regulations of the regulation (EU) 2017/745 of the European parliament and of the council on medical devices (MDR).

Prduct Name:

Tensoplus®

Basic UDI-DI:

4042809400364789S

Intended purpose:

Tensoplus is intended for fixation of wound dressings of all types and sizes as well as devices such as splints, tubes and paddings, for short term application of compression and to provide general support. The device is non-sterile and reusable. Intended Users are Health Care Professionals and patients. Use of product is not restricted to specific populations.

Conformity assessment route: Annex II+III

Classification rule:

Classification:

The Declaration of Conformity is performed in accordance with the quality management system according to EN ISO 13485, and fulfils the general safety and performance requirements of the regulation (EU) 2017/745.

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Compiled and released:

VIBRANE, 26.11.2021 Philippe Hatet Senior Project Manager BSN Medical SAS



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Article	Description	REF
72097-00006-03	TENSOPLUS COHESIVE 8CM X 3M WHITE 1 EN DE FR ES PT IT NL SV NO DA FI ZH AR HO	72097-06
72097-00007-03	TENSOPLUS COHESIVE 10CM X 3M WHITE 1 EN DE FR ES PT IT NL SV NO DA FI ZH AR HO	72097-07
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72097-00009-03	TENSOPLUS COHESIVE 10CM X 3M TAN 1 EN DE FR ES PT IT NL SV NO DA FI ZH AR HO	72097-09
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72097-00013-03	TENSOPLUS COHESIVE 8CM X 3M TAN 1 FR EN IT CS	72097-13
72097-00014-03	TENSOPLUS COHESIVE 10CM X 3M TAN 1 FR EN IT CS	72097-14