



## CE Declaration of Conformity

In accordance with Regulation (EU) 2017/745 of the European Parliament and of the Council of 05 April 2017 concerning medical devices

Medical Device	Family: Wound Care non-sterile
<p><b>Gauze Compresses</b></p> <ul style="list-style-type: none"> <li>- BeeSana® Mullkompressen (REF: 1000, 1002, 1004, 1006, 1008, 1010, 1012, 1014, 1016, 1018, 1023, 1025, 1038, 1052, 1053, 1235, 2708, 2712, 2714, 2723)</li> <li>- BeeSana® Mullschlitzkompressen (REF: 4291, 4192, 4122, 4160, 4159)</li> <li>- BeeSana® Mullzuschnitt (REF: 1330, 1335)</li> <li>- BeeSana® Verbandmull (REF: 1301)</li> </ul>	
<p><b>Non-woven Compresses</b></p> <ul style="list-style-type: none"> <li>- BeeSana® Vlieskompressen (REF: 4291, 4192, 4122, 4160, 4159, 1124, 1125, 1126, 1127, 1789, 1790, 1791, 1792, 1796, 1797, 1798, 1799, 2790, 2791, 2798, 2799, 11261)</li> <li>- BeeSana® Vliesschlitzkompressen (REF: 1793, 1794, 1795)</li> </ul>	
<p><b>Dressing pad</b></p> <ul style="list-style-type: none"> <li>- BeeSana® Saugkompresse (REF: 1224, 1225, 1227, 1228, 1229, 1230, 1231, 1244)</li> </ul>	
<p><b>Gauze Sponges</b></p> <ul style="list-style-type: none"> <li>- BeeSana® Mulltupfer (REF: 1102, 1106, 1109, 1113, 1116, 1348, 4460, 4461, 4462)</li> </ul>	
<p><b>Cellulose Sponges</b></p> <ul style="list-style-type: none"> <li>- BeeSana® Zellstofftupfer (REF: 1177; 1190; 1278)</li> </ul>	
<p><b>Adhesive Strips</b></p> <ul style="list-style-type: none"> <li>- ABE® Injektionspflaster (REF: 9525)</li> </ul>	
Intended use	Non-sterile products for general wound care. The devices are intended for the treatment of wounds with function such as absorption of blood or exudates, wound covering or swabs.
Basis UDI-DI according to Annex VI, Part C	GMN42500164W001classicwcnsGZ
Medical device class according to Annex VIII	I
Chosen conformity assessment procedure	The technical documentation according to Annex II and Annex III of Regulation (EU) 2017/745 is available.



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CE-mark since	Since 1998 according to 93/42/EEC and since 05.2021 according to Regulation (EU) 2017/745.
Validity of this CE Declaration of Conformity	07.12.2025

Manufacturer	Meditrade GmbH Medipark 1 83088 Kiefersfelden
Single Registration Number according to Article 31	DE-MF-000008937

We hereby declare in our sole responsibility the conformity of the above medical device with Regulation (EU) 2017/745 of the European Parliament and of the Council of 05 April 2017 on medical devices.

Meditrade hereby declares that medical devices covered by this declaration comply with this Regulation and, where applicable, with other relevant Union provisions which require the issuance of an EU declaration of conformity.

Common specifications applied:

There are no common specifications for these devices according to Article 9 of Regulation (EU) 2017/745.

Kiefersfelden, 08.12.2022

Martin Unterberg, PRRC

Person responsible for regulatory compliance under Article 15 of Regulation (EU) 2017/745, Meditrade GmbH