

Procedure:

Declaration of Conformity

Becton Dickinson and Company

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Authorized Benex Limited

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Conformity
Assessment
Directive 98/79/EC of the European Parliament and of the Council. Annex III of Directive 98/79/EC

Product: REF Product Name

291311 BD BBL™ Sensi-Disc™ Teicoplanin 30 µg

We hereby declare that the above mentioned products comply with the European In Vitro Diagnostic Medical Devices Directive and its relevant transposition into national laws of the member states into which we place the devices. This declaration is issued under the sole responsibility of Becton, Dickinson and Company.

Date: 12-Jan-2022

Name and Authority: Anne Zavertnik

Vice President Regulatory Affairs, IDS

Signature: One Swift

Technical File Number: BALTER291311

RECORD REVISION HISTORY TABLE	
Revision	Description of Changes
A	Initial Release