



EU DECLARATION OF CONFORMITY (DoC)

Manufacturer:	Becton, Dickinson and Company 7 Loveton Circle Sparks, Maryland 21152 USA.														
Manufacturer SRN:	US-MF-000018910														
Authorised Representative:	Becton Dickinson Ireland Ltd. Donore Road, Drogheda Co. Louth, A92 YW26 Ireland														
Authorised Representative SRN:	IE-AR-000007610														
Product:	<table border="1"><thead><tr><th>Catalogue No.</th><th colspan="2">Product Trade Name</th></tr></thead><tbody><tr><td>443624</td><td colspan="2">BD Phoenix™ M50 Automated Microbiology System Instrument</td></tr><tr><td>441107</td><td colspan="2">BD Phoenix™ Update Data</td></tr><tr><td>443866</td><td colspan="2">BD Phoenix™ M50 Automated Microbiology System Application Software</td></tr></tbody></table>			Catalogue No.	Product Trade Name		443624	BD Phoenix™ M50 Automated Microbiology System Instrument		441107	BD Phoenix™ Update Data		443866	BD Phoenix™ M50 Automated Microbiology System Application Software	
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	441107	BD Phoenix Update Data									
443866	Phoenix M50 Automated Microbiology System Application Software										
Intended Purpose:	<p>The BD Phoenix™ Automated Microbiology System is intended for the rapid identification (ID) and antimicrobial susceptibility testing (AST) of clinically significant bacteria. The BD Phoenix System provides rapid results for most aerobic and facultative anaerobic Gram-positive bacteria as well as most aerobic and facultative anaerobic Gram-negative bacteria of human origin. The BD Phoenix System is also intended for the rapid identification of yeast and yeast-like organisms.</p>										
Notified Body:	Not applicable, device(s) self-certified										
<p>We, as the manufacturer of the device(s) take sole responsibility for and hereby declare that the above-mentioned product(s) meet(s) the provisions of the following Directives/ Regulation(s):</p> <ul style="list-style-type: none"> Regulation (EU) 2017/746 on In vitro Diagnostic Medical Devices. Directive 2011/65/EU as amended (EU) 2015/863 on Restriction of the use of certain hazardous substances in electrical and electronic equipment as amended by Commission Delegated Directive 2015/863 (RoHS) 											

Conformity Assessment Route:

<input type="checkbox"/> ANNEX IX Technical File Examination	EC CERTIFICATE No.: N/A EC Certificate Expiration Date: N/A
<input type="checkbox"/> ANNEX IX Full Quality System	EC CERTIFICATE No.: N/A EC Certificate Expiration Date: N/A
<input type="checkbox"/> ANNEX X Type Examination	EC CERTIFICATE No.: N/A EC Certificate Expiration Date:
<input type="checkbox"/> ANNEX XI Production Quality System	EC CERTIFICATE No.: N/A EC Certificate Expiration Date: N/A
<input checked="" type="checkbox"/> ANNEX I & II+III	N/A

**Common Specifications (CS):**


Number:	Title:	Full or Partial Application:
Not Available	Not Available	Not Available

Common Specifications have not been issued for this product.

Devices Covered by this DoC:

SKU#	Device Name	Device Class
443624	BD Phoenix™ M50 Automated Microbiology System Instrument	Class A
441107	BD Phoenix™ Update Data ^a	
443866	Phoenix M50™ Automated Microbiology System Application Software ^a	

^a Product is not in scope for RoHS Directive 2011/65/EU

Authorised Signatory:	
Name & Title:	Anne Zavertnik, Vice President, Regulatory Affairs
Place of Issue:	Sparks, MD, USA
Date of Issue:	23-Mar-2022
Signature:	<p>DocuSigned by:</p> <p><i>Anne Zavertnik</i></p> <p> Signer Name: Anne Zavertnik Signing Reason: I approve this document Signing Time: 23-Mar-2022 6:47:19 PM GMT DC6A638A32E64A8A91F9D8DE330F0415</p>

DECLARATION OF CONFORMITY Revision History:

Version:	Detailed Change Description:
01	Initial release.
02	Updated to add Manufacturer SRN.
03	Removed WEEE Directive as it is not a CE-marking regulation. Added footnote to clarify that the RoHS directive applies to instruments only. Minor formatting changes.