

## EU Declaration of Conformity

<b>Manufacturer:</b>	SSI Diagnostica A/S Herredsvejen 2 DK-3400 Hillerød Denmark
<b>SRN no.</b>	DK-MF-000023922


PRODUCT IDENTIFICATION	
<b>Product Name</b>	<b>Code / Catalog Number</b>
<i>Salmonella</i> Sero-Quick ID kit	18350
<b>Intended Purpose</b>	<b>Basic UDI-DI</b>
The <i>Salmonella</i> Sero-Quick ID Kit is intended for complete serotyping of <i>S. Enteritidis</i> (1,9,12:g,m:-) and <i>S. Typhimurium</i> (1,4,[5],12:i:1,2). Sero-Quick ID Kit is used as an in vitro diagnostic aid for qualitative manual complete or partial bacterial serotyping by slide agglutination. It is important to use pure culture isolates for determination of bacterial antigens.	5713106S_Sero-Q_ID_KitVL

RISK CLASS FOR DEVICES		
<b>Device Classification</b>	<b>Rule:</b>	
<b>Class:</b>	<b>B</b>	IVDR rule 6
Common Specifications / Standards		
SSI Diagnostica A/S declares that the above-mentioned products meet the provision of the following EU legislation: <ul style="list-style-type: none"> <li>• European In Vitro Diagnostic Regulation (2017/746)</li> <li>• ISO 13485:2019 - Medical devices - Quality management systems - Requirements for regulatory purposes</li> <li>• ISO 14971:2019 – Risk Management</li> <li>• ISO 13612:2002 – Performance evaluation of in vitro diagnostic medical devices</li> <li>• ISO 15223-1: 2021 - Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General requirements</li> <li>• ISO 18113-1: 2011 - Information supplied by the manufacturer (labelling) – Part 1: Terms, definitions and general requirements</li> <li>• ISO 18113-2: 2011 - Information supplied by the manufacturer (labelling) - In vitro diagnostic reagents for professional use</li> </ul>		

### COMPANY REPRESENTATIVE:

Place/date: Hillerød, October 21<sup>th</sup> 2022

Signature:



Ulla W. Berg

Title:

QA/RA Manager