

EU Declaration of Conformity

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

PRODUCT IDENTIFICATION	
Product Name	Code / Catalog Number
ImmuLex Pneumococcal antisera Pool, Group, Type and Factor	Multiple items - eg. 52390, 77322, 77334, 98925
Intended Purpose	Basic UDI-DI
ImmuLex™ pneumococcus Pool, Group, Type and Factor antisera products are intended for visual qualitative confirmation, serogrouping and serotyping of Streptococcus pneumoniae (pneumococcus) by use of a rapid agglutination test. The products are for testing identified and confirmed, pure cultured isolates and strains of pneumococcus	5713106Pneu_ImmuLexDL

RISK CLASS FOR DEVICES		
Device Classification		Rule:
Class:	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"> European In Vitro Diagnostic Regulation (2017/746) ISO 13485:2019 - Medical devices - Quality management systems - Requirements for regulatory purposes ISO 14971:2019 – Risk Management ISO 13612:2002 – Performance evaluation of in vitro diagnostic medical devices ISO 15223-1: 2021 - Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General requirements ISO 18113-1: 2011 - Information supplied by the manufacturer (labelling) – Part 1: Terms, definitions and general requirements ISO 18113-2: 2011 - Information supplied by the manufacturer (labelling) - In vitro diagnostic reagents for professional use 		

COMPANY REPRESENTATIVE:

Place/date: Hillerød, October 21th 2022

Signature: 
Ulla W. Berg

Title: QA/RA Manager