

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	
ImmuView S. pneumoniae and L . pneumophila	2451 & 10557	
Urinary Antigen Test		
Intended Purpose	Basic UDI-DI	
The ImmuView® S. pneumoniae and L. pneumophila Urinary Antigen Test is an in vitro rapid lateral flow test, also known as a lateral flow immunochromatographic assay. The assay is intended for qualitative detection of Streptococcus (S.) pneumoniae and Legionella (L.) pneumophila antigens in urine specimens from patients with symptoms of pneumonia. The ImmuView® S. pneumoniae and L. pneumophila Urinary Antigen Test can be read visually or used in conjunction with the ImmuView® reader. The assay is intended to aid in diagnosis of S. pneumoniae and of L. pneumophila serogroup 1 infections. The assay is further intended to aid in the diagnosis of S. pneumoniae infections by detection of S. pneumoniae antigen in cerebrospinal fluid (CSF). Results from the ImmuView® S. pneumoniae and L. pneumophila Urinary Antigen Test should be interpreted in conjunction with the patient's clinical evaluation and other diagnostic methods.	5713106ImmuView_PutLutZH	



RISK CLASS	FOR DEV	'ICES
Device Clas	sification	Rule:
Class:	С	IVDR rule 3 (b)
C		no / Chandanda

Common Specifications / Standards

SSI Diagnostica A/S declares that the above-mentioned products meet the provision of the following EU legislation:

- 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:

Hilleroed, February 21th 2023

Signature:

Ulla W. Berg

Title:

QA/RA Manager