

## DECLARATION OF CONFORMITY

**Manufacturer:** Research and Development Department  
Of Salmonella Center IMMUNOLAB Ltd.  
Kładki 24, 80-822 Gdansk, Poland

This declaration of conformity is issued under the sole responsibility of the manufacturer.

**Name of the *in vitro* diagnostic medical device:** SIT OH-Sal  
Quick Identification Test  
for Salmonella spp.  
**catalog number PS006, code UDI-DI 5905753743806**

**Single Registration number (SRN):** PL-MF-000042563

**Basic UDI-DI number:** 590575374PS006FA

**Risk classification:** Class A defined for *in vitro* diagnostic medical device  
(IVDR, annex VIII)

**We declare** that product covered by this declaration is in conformity with the relevant Union harmonisation legislation: REGULATION OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL (EU) 2017/746 of 5 April 2017 on *in vitro* diagnostic medical devices and repealing Directive 98/79/EC and Commission Decision 2010/227/EU and have been marked with the sign:



Harmonised specifications and standards were used for conformity assessment in order to demonstrate the safety and effectiveness of an *in vitro* diagnostic medical device:

SPEC/PS006	Specification of SIT OH-Sal Quick Identification Test for Salmonella spp.
PN-EN ISO 14971:2019	Medical devices - Application of risk management to medical devices
PN-EN ISO 15223-1:2022-01	Medical devices -- Symbols to be used with information to be supplied by the manufacturer -- Part 1: General requirement
PN-EN ISO 20417:2021-10	Medical devices -- Information to be supplied by the manufacturer
PN-EN ISO 13485:2016-04	Medical devices – Quality management systems – Requirements for regulatory purposes

We declare the implementation, maintenance and improvement of the Quality Management System in accordance with the standard PN-EN ISO 9001:2015-10 in the scope of design and development, production and sales of the aforementioned medical device for *in vitro* use.

Gdańsk, 02.09.2024

(place, date)

  
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Chairman of the Board, Dorota Lieder