



# DECLARATION OF CONFORMITY

ACCORDING TO In Vitro Diagnostic Medical Devices Regulation (EU) 2017/746

## EU Representative

**SUNGO Cert Gmbh**  
**Harffstr. 47, 40591 Düsseldorf, Germany**  
**SRN: DE-AR-000010869**

## Device Classification

**Classification:** Class A.

**Rule:** According to Rule 5, Annex VIII, of  
In Vitro Diagnostic Medical Devices  
Regulation (EU) 2017/746.

## Applicable Standards

EN ISO 20417: 2021,  
EN ISO 15223-1:2016,  
EN ISO 18113-1:2011,  
EN ISO 14971:2019

## Remark

*The declaration of conformity is valid in connection  
with the release technical document  
CE/IVDR-UIP-05.*

*All the supporting documentation is retained at the  
premises of the manufacturer.*

*The Declaration of Conformity is exclusively under  
the sole responsibility of the manufacturer.*

## Manufacturer

**Name:** Shaoxing Shangyu Mingji Plastic Co., Ltd.  
**Address:** Shengqiao, Xiaoyue, Shangyu, Shaoxing,  
Zhejiang, China 312367

## Product Information

**Name:** SEROLOGICAL PIPETTE  
**Model:** MSP0101, MSP0201, MSP0501, MSP1001,  
MSP2501, MSP5001  
**GMDN:** 43375  
**Basic UDI-DI:**  
**Classification:** Class A

## Conformity Assessment

Compliance of the designated product with the In Vitro  
Diagnostic Medical Devices Regulation (EU) 2017/746  
has been assessed by issuing the EU declaration of  
conformity referred to in Article 17 after drawing up the  
technical documentation set out in Annexes II and III.

## Declaration

We herewith declare that the above-mentioned  
products meet the requirements of In Vitro Diagnostic  
Medical Devices Regulation (EU) 2017/746 and the  
applicable standards above.

Signature: *Song ying* Date: 2021.8.19

Position: GM Place: Shaoxing/China





