



# EU Declaration of Conformity

for In Vitro Diagnostic Medical Devices  
according to Annex IV of Regulation (EU) 2017/746

**Manufacturer:** BioMaxima S.A., Vetterów 5, 20-277 Lublin, Poland

**SRN:** Not Available

**Product Name:** Mueller Hinton 2 LAB-AGAR™ + 5% KB

**Basic UDI-DI:** Not Available

**Classification (IVDR, Annex VIII):** Class A, Rule 5

We herewith under our sole responsibility declare that the above mentioned products meet the provisions of the Regulation (EU) 2017/746

The manufacturer is exclusively responsible for the declaration of conformity.

Place and date of issue:

Lublin, 27.05.2022

Signed on behalf of BioMaxima S.A.:

Henryk Lewczuk  
VicePresident

A blue ink signature of Henryk Lewczuk.

Patrycja Paniak -Sankowska  
Proxy

A blue ink signature of Patrycja Paniak -Sankowska.