



## EU Declaration of Conformity

Manufacturer Name: BiopSafe ApS

Manufacturers Address: Bygstubben 4  
DK-2950 Vedbæk, Denmark  
Email address: info@biopsafe.com

Authorised Representative Name: Not Applicable due to manufacturing location inside the EU.

Name of the Device: BiopSafe® Biopsy Container

Product code: 3178-200-xx size 20 ml., and  
3178-600-xx size 60 ml.

Intended purpose: The BiopSafe® Biopsy Container is intended to preserve tissue in a closed formalin system.

Basic UDI-DI: 5714678BiopSafeNG

Registration at  
The Danish Medicines Agency: 2023113470

EUDAMED registration: Not applicable until June 2029 (Pending)

Device Classification: Medical Device In Vitro Diagnostic, Class A non-sterile.

Conformity assessment route: The Conformity assessment route for the CE marking of the BiopSafe is according to the Medical Device IVD EU Regulation 2017/746:  
Classification MD IVD Class A non-sterile: EU Declaration of Conformity according to Annex II + Annex III. Self-certified.

Notified Body information: Not applicable due to BiopSafe® Classification MD IVD Class A non-sterile.

This EU Declaration of Conformity is issued under the sole responsibility of BiopSafe ApS.  
At this moment, BiopSafe ApS declare that the BiopSafe® specified above meets the requirements of Medical Device EU 2017/746 IVD Regulation.  
All supporting documentation is retained at the premises of BiopSafe ApS.

Location and date of issue:

Vedbæk, 4<sup>th</sup> of December 2023

Signature:

A handwritten signature in blue ink, appearing to be "Søren Christensen", written over a horizontal dotted line.

Mr. Søren Christensen  
CEO