

EU DECLARATION OF CONFORMITY – BIOPAD BIOPSY PADS

Date: 21/07/2021

This Declaration of Conformity has been issued solely by CellPath Limited (Company Number 01831261) in accordance with the definitions and regulations as described in the **EU Regulation 2017/746 of The European Parliament and of the Council on In-Vitro Diagnostic Medical Devices**.

General Information

Manufacturer CellPath Limited

Address of Production Facility CellPath Ltd
Unit 80, Mochdre Enterprise Park
Newtown, Powys
Mid Wales, UK
SY16 4LE

EU Authorised Representative EMERGO EUROPE (NL-AR-000000116)
Prinsessegracht 20
2514 AP The Hague
The Netherlands

Product Name Histological tissue cassette biopsy insert IVD

Product Description A flexible and porous material designed to securely hold a biopsy specimen within a tissue cassette, to prevent loss of the specimen during specimen processing (e.g., fixation, dehydration, infiltration) in preparation for subsequent cytological or histological examination. The flexible and porous material (e.g., paper, synthetic polymer) is intended to allow infiltration of reagents and maximize fluid-exchange during tissue processing. It is available in various forms (e.g., sheet, bag, foam pad) and is placed in a tissue processing cassette, with the biopsy specimen wrapped or contained inside it. This is a single-use device.

Product Classification CLASS A
(Classified in accordance with EU
Regulation 2017/746)

Registered Office:

CellPath Ltd, 80 Mochdre Enterprise Park,
Newtown, Powys, SY16 4LE Mid Wales, UK
Registration Number: 01831261

DEVICE(S) INFORMATION

PART NUMBER	GTIN	UDI-DI	DESCRIPTION	YEAR OF CE MARKING
EBA-0101-03A	05060470564438	05060470EBA001RC	BIOPAD BIOPSY PAD – BLACK	2011
EBA-0102-03A	05060470564445	05060470EBA001RC	BIOPAD BIOPSY PAD – WHITE	2009
EBA-0104-03A	05060470564452	05060470EBA001RC	BIOPAD BIOPSY PAD – GREEN	2014
EBA-0106-03A	05060470564469	05060470EBA001RC	BIOPAD BIOPSY PAD – BLUE	2003

Conformity to the aforementioned regulation is demonstrated by the company's compliance to the following international manufacturing standard:

- **EN ISO 13485:2016** - Medical Devices, Quality Management Systems - Requirements for Regulatory Purposes


Signed on behalf of the Manufacturer CellPath Limited

Address of Issue CellPath Ltd
 Unit 80, Mochdre Enterprise Park
 Newtown, Powys
 Mid Wales, UK
 SY16 4LE

Date of Issue 21/07/2021

Name of Signatory Paul Webber

Position of Signatory Joint Managing Director

Signature 
Paul Webber (Jul 21, 2021 10:23 GMT+1)