

Declaration of Conformity

File: F_Dec._IVD_08
Page: 1 / 1
Date: February-2023

Manufacturer : FEATHER SAFETY RAZOR CO., LTD.
3-70, Ohyodo-Minami 3-chome, Kita-ku, Osaka, 531-0075 JAPAN

Facility : FEATHER SAFETY RAZOR CO., LTD. MINO SITE
600-1, Matsumori, Mino-city, Gifu, 501-3753 JAPAN

SRN Number JP-MF-000018505

Authorized Representative : pfm medical ag
Wankelstr. 60, 50996 Köln, GERMANY

SRN Number DE-AR-000005340

We, FEATHER SAFETY RAZOR CO., LTD. located at 3-70, Ohyodo-Minami 3-chome, Kita-ku, Osaka, 531-0075 JAPAN herewith declare under our sole responsibility that the device(s) covered by this declaration is in conformity with Regulation (EU) 2017/746, General Safety and Performance requirements, and relevant Union legislation. Any alterations made without our consent shall render this declaration null and void.

Product category : Pathological Instruments

Product group : Microtome Blades

Classification : Class A (acc. to Annex VIII of IVDR 2017/746)

Product name : **FEATHER MICROTOME BLADE**

Model No. (Ref. No.) : S35 (02.075.00.000), S22 (02.075.00.001), A22 (02.075.00.002),
C35 (02.075.00.003), S35L (02.075.00.004), R35 (02.075.00.005),
N35 (02.075.00.006), S35LL (02.075.00.009), FHP-01 (02.075.00.010),
A35 (02.075.00.011), N35HR (02.075.00.014), UNIVERSAL (02.074.00.000),
FHP-02 (02.075.00.016)

Basic UDI-DI 490247001MIBF5


Intended purpose of use : The devices are intended for thin-sectioning of tissue that has been processed for pathological examination.

EC-Directive : Regulation (EU) 2017/746 (In-Vitro Diagnostic Medical Device Regulation)

Conformity Assessment Route : Annex II and III, EU DECLARATION OF CONFORMITY

Common specifications N/A

Place, Date : GIFU JAPAN, February 1, 2023

Signature :  Satoshi Mitsuishi

Title : Executive Director