

EN

EU Declaration of Conformity

12-CEM-01-L

Manufacturer:	FUJIFILM Corporation
single registration number (SRN)	TBD
Address:	26-30, Nishiazabu 2-chome, Minato-ku, Tokyo 106-8620, JAPAN
Authorized Representative:	FUJIFILM Europe GmbH
single registration number (SRN)	TBD
Address:	Heesenstrasse 31 40549 Duesseldorf, GERMANY
Basic UDI-DI:	454741010100000000000353K
Trade name:	FUJI MEDICAL DRY LASER IMAGER DRYPIX Smart
Model Number:	DRYPIX6000
Applicable Products:	xxx51978 or later
Classification (MDR, Annex VIII):	Class I (Rule 13)

We, FUJIFILM Corporation, herewith declare in our sole responsibility that the product(s) identified in this declaration conforms to the provisions of the following Regulation(s) and Directive(s).

Regulation:
 Medical Device Regulation: REGULATION (EU) 2017/745 and their Annexes.

Directive:
 RoHS Directive: 2011/65/EU,(EU)2015/863

Common specifications ('CS'):
 No references to any CS

Conformity Assessment Procedure for Regulation (EU) 2017/745 :
Annex IV

Place and Date of issue

Kanagawa, JAPAN

2021-05-21

Signature :

Name :

Function :

Naotake Mitsumori

General Manager,

Quality Assurance and Regulatory Affairs Division,

Medical Systems Business Division

FUJIFILM Corporation