

EC Declaration of Conformity

Name	Device Description	Basic UDI-DI
Pipette (FS1028)	Single-use Mouthpiece (link	376025345FF1023T9
	between the mouth and the	
	TABATABA®V2 device)	

Medical devices conform to the following standards:

NF EN ISO 13485 :2016 : Medical Device – Quality Management System

NF ISO 2859-1 :2000 : Sampling procedures for inspection by attributes — Part 1: Sampling schemes indexed by acceptance quality limit (AQL) for lot-by-lot inspection

EN ISO 10993-1:2009/AC: 2010: Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process. EN ISO 10993-5:2009: Biological evaluation of medical devices - Partie 5 : Tests for in vitro cytotoxicity

EN ISO 10993-10:2010: Biological evaluation of medical devices - Partie 10 : Tests for irritation and skin sensitization

NF EN ISO 14971:2019: Medical devices - Application of risk management to medical devices

NF EN 1041 + A1: 2013: Information supplied by the manufacturer of medical devices

EN ISO 15223-1:2016: Medical devices - Symbols to be used with medical device labels, labelling and information to be supplied - Part 1: General

I the undersigned, Dounia Benbachir, Quality Manager of the FIM MEDICAL company located at 51 rue Antoine Primat Villeurbanne - FRANCE assure and declare that the medical devices listed above belong to class I (Rule 5) and satisfy the provisions of annex I (Essential Requirements), annex VII (Evaluation of Tabataba® V2 FF1006DTR100 technical file) of the 93/42/EEC directive and its local adaptation (Book II of the public health code).

Villeurbanne, 12/07/2021, D.BENBACHIR

Quality Manager

CE