

EU Declaration of Conformity

Name of manufacturer: UAB "Medicinos linija"
Single registration number: LT-MF-000002719
Basic UDI-DI 4772178ML11PMTQ
Registered place of business and address of manufacturer: Aviacijos str. 28, LT-77103 Šiauliai, Lithuania
Date of issue of the declaration: 26th of May 2021
Place of issue: Šiauliai
Expiry date: 26th of May 2026

Medical device family: Prophylaxis materials
Medical device name: Teeth surface polishing paste
Intended use: Teeth surface polishing paste is intended for cleaning and polishing the surface of the tooth during restorative and prophylactic dental treatment.

Medical device trade names and specification:

<i>Medical device trade name</i>	<i>Catalogue number (REF)</i>	<i>Specification</i>
i-FASTE	IFCSTP	10g paste coarse, sample
	IFCTP	30g paste coarse
	IFC100	100g paste coarse
	IFCD1	100x2g paste coarse
	IFMSTP	10g paste medium, sample
	IFMTP	30g paste medium
	IFM100	100g paste medium
	IFMD1	100x2g paste medium
	IFFSTP	10g paste fine, sample
	IFFTP	30g paste fine
	IFF100	100g paste fine
	IFFD1	100x2g paste fine
	IFMB30	30g paste medium, bubble mint
	IFMB100	100g paste medium, bubble mint
	IFMB250	250g paste medium, bubble mint
	IFMM30	30g paste medium, mangosteen

	IFMM100	100g paste medium, mangosteen
	IFMM250	250g paste medium, mangosteen
	IFMC30	30g paste medium, cherry
	IFMC100	100g paste medium, cherry
	IFMC250	250g, paste medium, cherry
	IFMCSP	2g paste medium, sample, cherry
	IFMCSTP	10 g paste, sample, cherry
	IFMMSP	2g, paste medium, sample, mangosteen
	IFMMSTP	10g paste medium, sample, mangosteen
	IFMBSP	2g paste medium, sample, bubble mint
	IFMBSTP	10g paste medium, sample, bubble mint
	IFMSK1	3x10g paste medium, sample (mangosteen, cherry, bubble mint)
	IFMCD1	100x2g paste medium, cherry
	IFMMD1	100x2g paste medium, mangosteen
	IFMBD1	100x2g paste medium, bubble mint
dline Prophy Paste	22000	30g paste coarse
	22001	100g paste coarse
	22004	100x2g paste coarse
	22010	30g paste medium
	22011	100g paste medium
	22014	100x2g paste medium
	22020	30g paste fine
	22024	100x2g paste fine
PROFEX	C12530	100g paste coarse, peppermint flavour, green colour
	C12620	100g paste medium, peppermint flavour, green colour

- This EU declaration of conformity is issued under the sole responsibility of the manufacturer.
- Risk class of the device:
 - Class I, according regulation (EU) 2017/745 of the European Parliament and of the Council Annex VIII, rule 5.
- Medical device complies with the fundamental requirements for medical devices listed in (EU) 2017/745 of the European Parliament and of the Council Annex I.
- Standards and common specifications applied:

EN ISO 13485:2016 Medical devices. Quality management systems. Requirements for regulatory purposes (ISO 13485:2016); EN ISO 13485:2016/AC:2018 Medical devices - Quality management systems - Requirements for regulatory purposes (ISO 13485:2016); EN ISO 14971: 2019 Medical devices - Application of risk management to medical devices (ISO 14971:2019); EN ISO 7405:2018 Dentistry - Evaluation of biocompatibility of medical devices used in dentistry (ISO 7405:2018, Corrected version 2018-12); EN ISO 10993-1:2018 Biological evaluation of medical devices – Part 1: Evaluation and testing within a risk management process (ISO 10993-1:2018); EN ISO 10993-5:2009 Biological evaluation of medical devices – Part 5: Tests for in vitro cytotoxicity (ISO 10993-5:2009); EN ISO 10993-10:2010 Biological evaluation of medical devices – Part 10: Tests for irritation and skin sensitization (ISO 10993-10:2010); EN ISO 10993-11:2017 Biological evaluation of medical devices – Part 11: Tests for systemic toxicity (ISO 10993-11:2017); EN ISO 10993-18:2009 Biological evaluation of medical devices – Part 18: Chemical characterization of materials (ISO 10993-18:2009); EN ISO 10993-19:200 Biological evaluation of medical devices – Part 19: Physico-chemical, morphological and topographical characterization of materials (ISO 10993-19:2006); EN ISO 15223-1: 2016 Medical devices - Symbols to be used with medical device labels, labelling and information to be supplied - Part 1: General requirements (ISO 15223-1:2016, Corrected version 2017-03); EN 1041: 2008+A1:2013 Information supplied by the manufacturer of medical devices; EN 1641:2009 Dentistry - Medical devices for dentistry – Materials; MEDDEV 2.7/1 rev.4, 2016 Clinical evaluation: a guide for manufacturers and notified bodies under Directives 93/42/EEC and 90/385/EEC; MEDDEV 2.12/1 rev.8, 2013 Guidelines on a medical devices vigilance system; MEDDEV 2.12/2 rev.2, 2012 Post market clinical follow-up studies a guide for manufacturers and notified bodies; MEDDEV 2.2/3 rev.3, 1998 "USE-BY" date.

- Referenced technical documentation is retained under the premises of the manufacturer.

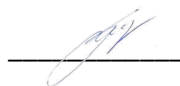
On behalf of UAB "Medicinos linija",

General Manager



Rima Žurauskaitė-Storace

Quality Executive Manager



Simona Daunienė

Declaration history

Date	Action
2021-05-26	First issue
2021-10-14	Included REF No. : IFMB100; IFMB250; IFMM250; FMM100; IFMC100; IFMC250; IFMCSP; IFMCSTP; IFMMSP; IFMMSTP; IFMBSP; IFMBSTP; IFMSK1; IFMC30; IFMM30; IFMB30; IFMCD1; IFMMD1; IFMBD1