

July 22, 2020

InMode Ltd. % Amit Goren Regulatory Manager A. Stein - Regulatory Affairs Consulting Ltd. 20 Hata'as Str., Suite 102 Kfar Saba, 4442520 Israel

Re: K201150

Trade/Device Name: InMode RF Multi-System

Regulation Number: 21 CFR 878.4400

Regulation Name: Electrosurgical Cutting and Coagulation Device and Accessories

Regulatory Class: Class II

Product Code: GEI, PBX, ISA, NUV

Dated: April 22, 2020 Received: April 29, 2020

Dear Amit Goren:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. Although this letter refers to your product as a device, please be aware that some cleared products may instead be combination products. The 510(k) Premarket Notification Database located at https://www.accessdata.fda.gov/scripts/cdrh/efdocs/efpmn/pmn.cfm identifies combination product submissions. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

K201150 - Amit Goren Page 2

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803) for devices or postmarketing safety reporting (21 CFR 4, Subpart B) for combination products (see https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820) for devices or current good manufacturing practices (21 CFR 4, Subpart A) for combination products; and, if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to https://www.fda.gov/medical-device-problems.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance) and CDRH Learn (https://www.fda.gov/training-and-continuing-education/cdrh-learn). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice">https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice) for more information or contact DICE by email (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

Long Chen, Ph.D.
Assistant Director
DHT4A: Division of General Surgery Devices
OHT4: Office of Surgical
and Infection Control Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

K201150 Page 1 of 1

DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

Indications for Use

Form Approved: OMB No. 0910-0120

Expiration Date: 06/30/2020 See PRA Statement below.

510(k) Number <i>(if known)</i>
K201150
Device Name InMode RF Multi-System
Indications for Use (Describe) The InMode RF Multi-System with the Non-invasive Applicators employs RF energy for various applications:
•Forma (Plus), Plus (Plus Plus) and Plus90 for relief of minor muscle aches and pain, relief of muscle spasm, and temporary improvement of local blood circulation.
•WMface is intended for use in dermatologic procedures for non-invasive treatment of mild to moderate facial wrinkles and rhytids.
•BodyFX TM (WMBody)/MiniFX TM for Relief of minor muscle aches and pain, relief of muscle spasm, temporary improvement of local blood circulation and temporary reduction in the appearance of cellulite.
The InMode RF Multi-System with the Fractional Applicators employs RF energy for various applications:
•Fractora Applicator with 60 pins tip is designed for use in dermatological procedures requiring ablation and resurfacing of the skin.
•Fractora Applicator with 24 pins tip is intended for use in dermatological and general surgical procedures for electrocoagulation and hemostasis. At higher energy levels greater than 62mJ/pin, use of the applicator is limited to skin types I-IV
•Morpheus8 [™] for dermatological and general surgical procedures for electrocoagulation and hemostasis. At higher energy levels greater than 62mJ/pin, use of the applicator is limited to skin types I-IV
Type of Use (Select one or both, as applicable)
Prescription Use (Part 21 CFR 801 Subpart D) Over-The-Counter Use (21 CFR 801 Subpart C)
CONTINUE ON A SEPARATE PAGE IF NEEDED.

This section applies only to requirements of the Paperwork Reduction Act of 1995.

DO NOT SEND YOUR COMPLETED FORM TO THE PRA STAFF EMAIL ADDRESS BELOW.

The burden time for this collection of information is estimated to average 79 hours per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden, to:

Department of Health and Human Services Food and Drug Administration Office of Chief Information Officer Paperwork Reduction Act (PRA) Staff PRAStaff@fda.hhs.gov

"An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number."

510(K) SUMMARY INMODE RF MULTI-SYSTEM

510(k) Number $\underline{K201150}$

Applicant Name:

Company Name: InMode Ltd.

Address: Tabor Building, Shaar Yokneam

Yokneam 20692

Israel

Tel: +972-4-9097470 Fax: +972-4-9097471

E-mail: amit@asteinrac.com

Contact Person:

Official Correspondent: Amit Goren

Company Name: A. Stein – Regulatory Affairs Consulting Ltd.

Address: 20 Hata'as Str., Suite 102

Kfar Saba 4442500

Israel

Tel: +972-9-7670002 Fax: +972-9-7668534

E-mail: amit@asteinrac.com

Date Prepared: July 21, 2020

Trade Name: InMode RF Multi-System

Classification Regulation:

Classification Name	Regulation No.	Product Code
Electrosurgical cutting and	878.4400	GEI, PBX
coagulation device and accessories.		
Therapeutic massager.	890.5660	ISA
Laser surgical instrument for use in	878.4810	NUV
general and plastic surgery and in		
dermatology.		

Classification: Class II Medical Device

Predicate Device: The InMode RF Multi-System is substantially equivalent to the previously cleared predicate devices;

Manufacturer	Device	510(k) No.
	Main Predicate	
InMode Ltd.	InMode RF System	K182325
	Reference Predicates	
InMode Ltd.	InMode System MiniFX Applicator	K160329
InMode Ltd.	InMode WMBody Device	K131362
InMode Ltd.	InMode PLUS System	K172302
InMode Ltd.	WMFACE Applicator	K140926
InMode Ltd.	FRACTORA 60 pin	K102461
InMode Ltd.	InMode System with Fractora3D/3D-90	K180189
	Applicators	
InMode Ltd.	InMode FRF 24 pin Applicator	K151273
InMode Ltd.	InMode System with the Morpheus 12, 24, 40	K192695
	& T Applicators	
InMode Ltd.	InMode System with the Morpheus8 Applicators	K200947

Device Description:

The InMode RF Multi-System supports multiple radiofrequency (RF) applications and accessories. The InMode RF Multi-System is compatible with the Fractional RF Applicators and the Non-Invasive RF Applicators, and employs RF energy for various applications.

The InMode RF Multi-System consists of platform console with an AC/DC power supply unit, two applicator connectors, RF generator, RF measuring circuit, controller, footswitch and user interface including a touch screen. The RF Applicator is connected to the console via a cable and a footswitch activates the energy delivery to the applicator. The applicators are comprised of a handle and electrodes, and some of them are used with a single-use tip.

This 510(k)-file submission includes all the FDA-Cleared applicators under one submission with the RF-supporting console, as cleared in K182325, compatible with all applicators.

The below list comprises the set of applicators to be registered under the subject device:

- Non-Invasive RF Applicators:
 - o Forma (Plus)
 - o Plus (Plus Plus)
 - o Plus90
 - o WMface
 - o BodyFXTM (WMBody)
 - o MiniFXTM

- Fractional RF Applicators:
 - Fractora
 - 24 pins tip (FRF)
 - 60 pins tip
 - o Morpheus8TM
 - 12 pins tip (Prime Tip)
 - 24 pins tip (Fractora 3D)
 - 40 pins tip (Body Tip)
 - T tip

Following are the InMode RF Multi-System device specifications:

RF Max Output Power	65 Watt
RF Output Frequency	$1[MHz] \pm 2\%$
Dimensions	46cm W x 46cm D x 100cm H
	[18.2" W x 18.2" D x 40" H]
Console Weight	32.0Kg [70.548lb]
Main Line Frequency (nominal)	50-60 Hz
Input Voltage (nominal)	100-240 VAC

Intended Use/Indication for Use:

The InMode RF Multi-System with the Non-invasive RF Applicators employs RF energy for various applications:

- Forma (Plus), Plus (Plus Plus) and Plus90 for relief of minor muscle aches and pain, relief of muscle spasm, and temporary improvement of local blood circulation.
- WMface is intended for use in dermatologic procedures for non-invasive treatment of mild to moderate facial wrinkles and rhytids.
- BodyFXTM (WMBody)/MiniFXTM for relief of minor muscle aches and pain, relief of muscle spasm, temporary improvement of local blood circulation and temporary reduction in the appearance of cellulite.

The InMode RF Multi-System with the Fractional Applicators employs RF energy for various applications:

- Fractora Applicator with 60 pins tip is designed for use in dermatological procedures requiring ablation and resurfacing of the skin.
- Fractora Applicator with 24 pins tip is intended for use in dermatological and general surgical procedures for electrocoagulation and hemostasis.

 At higher energy levels greater than 62mJ/pin, use of the applicator is limited to skin types I-IV
- Morpheus8[™] for dermatological and general surgical procedures for electrocoagulation and hemostasis.

At higher energy levels greater than 62mJ/pin, use of the applicator is limited to skin types I-IV

Performance Standards:

The InMode RF Multi-System complies with the following recognized consensus standards:

- [Rec. Number 19-4] ANSI AAMI ES 60601-1:2005/(R)2012 and A1:2012, C1:2009/(R)2012 and A2:2010/(R)2012 (Consolidated Text) Medical electrical equipment Part 1: General requirements for basic safety and essential performance
- [Rec. Number 5-89] IEC 60601-1-6 Edition 3.1 2013-10 Medical electrical equipment Part 1-6: General requirements for basic safety and essential performance Collateral standard: Usability
- [Rec. Number 19-8] IEC 60601-1-2 Edition 4.0 2014-02 Medical electrical equipment Part 1-2: General requirements for basic safety and essential performance Collateral Standard: Electromagnetic disturbances Requirements and tests
- [Rec. Number 6-389] IEC 60601-2-2 Edition 6.0 2017-03 Medical electrical equipment Part 2-2: Particular requirements for the basic safety and essential performance of high frequency surgical equipment and high frequency surgical accessories

The test certificates and test reports documents conclude that the InMode RF Multi-System complies with the requirements of all the above-mentioned standards. The test certificates and reports for the InMode RF Multi System are provided in section 17 of this 510(k) submission.

Non-Clinical (Bench) Performance Data:

The following performance data were provided in support of the substantial equivalence determination:

Electrical safety and electromagnetic compatibility (EMC)

Electrical safety and EMC testing were conducted on the InMode RF Multi-system. The system complies with the IEC 60601-1, IEC 60601-2-2 standards for safety and the IEC 60601-1-2 standard for EMC.

Software Verification and Validation Testing

Software verification and validation testing were conducted and documentation was provided as recommended by FDA's Guidance for Industry and FDA Staff, "Guidance for the Content of Premarket Submissions for Software Contained in Medical Devices." The software for this device was considered as a "major" level of concern, since a failure or latent flaw in the software could directly result in serious injury or death to the patient or operator.

Animal Performance Data / Histology Data:

Non-Applicable

Clinical Performance Data:

Non-Applicable

Substantial Equivalence:

The following tables provide a comparison information for the InMode RF Multisystem and its main predicate, the InMode RF System and for the subject device applicators and their compatible predicate reference devices:

	Subject Device InMode RF Multi-System K201150	Main Predicate Device InMode RF System K182325
General		
Product Code	GEI	GEI
Class	Class II	Class II
Manufacturer	InMode Ltd.	InMode Ltd.
RF platform	InMode RF Multi-System	InMode RF System
Prescription or OTC	Prescription use only	Idem
Target Population	Adults requiring treatment as specified in the indication for use	Idem
Anatomical sites	Body parts requiring treatment as specified in the indication for use	Idem
Environment Used	Hospital or Clinic setting	Idem
Device description /	The InMode RF Multi-System	The InMode RF System is a
Design	is a line power, electro	line power, electro thermal,
	thermal, radio frequency	radio frequency system with
	system with integral	integral temperature and
	temperature and impedance	impedance feedback. The
	feedback. The system is	system is capable of

	Subject Device	Main Predicate Device
	InMode RF Multi-System	InMode RF System
	K201150	K182325
	capable of delivering up to 65	delivering up to 40 watts of
	watts of power.	power.
Device components	The InMode RF Multi-System	The InMode RF System
1	consists of the same following	consists of the following
	components as the InMode RF	components:
	System, with additional	• Electrosurgical Unit which
	applicators as described in next	includes the power supply, RF
	table:	generator, controller and LCD
	 Electrosurgical Unit which 	touch screen control and
	includes the power supply,	display panel
	RF generator, controller and	• RF measuring circuit
	LCD touch screen control	• AC/DC power supply
	and display panel	• Applicators
	• RF measuring circuit	• Applicators connector
	• AC/DC power supply	• Footswitch
	• Designated applicators	
	• 2 Applicator connectors	
	• Footswitch	
Input Power		
Main Line Frequency	60 - 50Hz	idem
(nominal)	00 2012	Idelli
Input Voltage (nominal)	100-240VAC	
Input Current (rms)	2A	
Dimensions		
Difficusions		
System	46cm W x 46cm D x 100cm H	idem
	[18.2" W x 18.2" D x 40" H]	
Weight		
System	32.0Kg [70.5lb]	idem
RF Output Parameters		
Maximum Output Power	65[W]	40[W]
Frequency	1MHz	idem
Crest Factor (Rated Load)	1.4± 2%	idem
Safety & adherence to consensus standards		

	Subject Device InMode RF Multi-System K201150	Main Predicate Device InMode RF System K182325
Standards Met	System adheres to:	idem
	• IEC 60601-1	
	• IEC 60601-1-2	
	• IEC 60601-2-2	
Biocompatibility	Materials are biocompatible.	idem
Compatibility with	InMode RF System is	idem
Environment and Other	compliant with the IEC 60601-	
Devices	1-2 (EMC Safety) standard	
Electrical Safety	Power Requirements:	idem
	100-240 VAC 50-60 Hz	
	The InMode RF System is	
	compliant with the IEC 60601-	
	1 standard.	
Mechanical Safety	The InMode RF System is	idem
	compliant with the IEC 60601-	
61 1.0.6	1 standard.	
Chemical Safety	Not Applicable	idem
Thermal Safety	The InMode RF System is	idem
	compliant with the IEC 60601-	
	1 standard.	
Radiation Safety	The InMode RF System is	idem
	compliant with the IEC 60601-	
	1-2 (EMC Safety) standard	

Technological Characteristic	Subject Device InMode RF Multi-System K201150	Reference Predicates K102461 - FRACTORA 60 pin Applicator K151273 - InMode FRF 24 pin Applicator K192695 & K200947- Morpheus8 (Fractora 3D) Applicator with the 12, 24, 40 & T tip heads
Product Code	GEI	idem
Class	Class II	
Manufacturer	InMode Ltd.	Idem
Device Technological Characteristics		
Energy Used	Fractional RF	idem
Specifications		
Max RF Energy output	65W	idem

Technological Characteristic	Subject Device InMode RF Multi-System K201150	Reference Predicates K102461 - FRACTORA 60 pin Applicator K151273 - InMode FRF 24 pin Applicator K192695 & K200947- Morpheus8 (Fractora 3D) Applicator with the 12, 24, 40 & T tip heads
RF Frequency	1 MHz	idem
Max RF Energy levels Tip needles penetration depth	FRACTORA 60 pin: 60W InMode FRF 24 pin: 60W Morpheus8 12, 24, 40: 60W Morpheus8 T: 30W FRACTORA 60 pin: 0.2mm InMode FRF 24 pin: 2.5mm Morpheus8 12, 24 pins: 4.0mm Morpheus8 40 pin: 7.0mm Morpheus8 T: 0.5mm	FRACTORA 60 pin: 62W InMode FRF 24 pin: 62W Morpheus8 12, 24 & 40: 60W Morpheus8 T: 30W idem
General Aspects		
Biocompatibility	Materials are biocompatible.	idem
Sterility	All tips are Gamma-sterilized	idem
Reprocessing	Handle to be reprocessed in accordance with user manual instructions. Tips are for single-use	idem

Technological Characteristic	Subject Device InMode RF Multi-System	Reference Predicates K131362- BodyFX (WMBody) K140926-WMFace K160329- MiniFX K172302- Forma (Plus), Plus90 and Plus (Plus-Plus)
Product Code	GEI, PBX, ISA/NUV	idem
Class	Class II	
Manufacturer	InMode Ltd.	idem
Device Technological Characteristics		
Energy Used	Bipolar RF	idem
Specifications		
Maximal RF output power	BodyFX (WMBody): 50 [W] WMFace: 65 [W] MiniFX: 25 [W]	idem

Technological Characteristic	Subject Device InMode RF Multi-System	Reference Predicates K131362- BodyFX (WMBody) K140926-WMFace K160329- MiniFX K172302- Forma (Plus), Plus90 and Plus (Plus-Plus)
	Forma (Plus), Plus (Plus- Plus): 50 [W] Plus90: 50 [W]	
RF Frequency	1 MHz	idem
General Aspects		
Biocompatibility	Materials are biocompatible.	idem
Sterility	NA	NA
Reprocessing	Applicators to be reprocessed in accordance with user manual instructions.	idem

The indications for use and technological characteristics of the InMode RF Multi-System are substantially equivalent to the indications for use and technological characteristics of the InMode RF System (predicate device, subject of K182325) and applicators (reference devices).

The design of and components included in the InMode RF Multi-System, including the main system platform, with power supply, RF generator, controller and display panel are similar to the design and components found in the main predicate device except for slight design modifications to support the utilization of all of the system applicators, mainly in software design. The subject device applicators possess similar technological principals to the predicate devices' applicators. Minor changes were done for some of the applicators, still the device performance and safety are maintained due to the exact functionality and mechanism of operation. Both the subject and predicate devices present similar performance specifications (for the specified indications for use) and similar monitoring features (where applicable) in order to maintain the desired performance specifications. The safety features and compliance with safety standards in the InMode RF Multi-System are similar to the safety features and compliance with safety standards found in the predicate device. Patient contacting materials are also similar. Any minor differences in the technological characteristics do not raise new safety or effectiveness concerns. Furthermore, the subject device underwent performance testing including software validation testing, electrical and mechanical safety testing according to IEC 60601-1, electromagnetic compatibility testing according to IEC 60601-1-2 and high frequency of surgical equipment testing according to IEC 60601-2-2. User interface was slightly changed in order to support all applicators, and labeling was updated to allow appropriate use.

Consequently, it can be concluded that the InMode RF Multi-System is substantially equivalent to the predicate InMode RF System, cleared under 510(k) K182325 and the

Page 5-10

InMode RF Multi-System 510(k) file Section 5 – 510(k) Summary

reference device cleared under 510(k) K1024615, K131362, K140926, K1512735, K160329, K172302, K180189, K192695 and K200947 and therefore, may be legally marketed in the USA.

Therefore, the InMode RF Multi-System may be legally marketed in the USA.

Conclusions:

Based on the performance testing and comparison to predicate device and reference devices, the InMode RF Multi-System is substantially equivalent to the InMode RF System (predicate device) and Applicators (reference devices).