

May 25, 2021

Diode Art Engineering doing business as Air Relax % Prithul Bom
Most Responsible Person
Regulatory Technology Services, LLC
1000 Westgate Drive, Suite 510k
Saint Paul, Minnesota 55114

Re: K211460

Trade/Device Name: Air Relax Plus Model AR-3.0

Regulation Number: 21 CFR 890.5650

Regulation Name: Powered inflatable tube massager

Regulatory Class: Class II

Product Code: IRP Dated: May 10, 2021 Received: May 11, 2021

#### Dear Prithul Bom:

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 <a href="https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm">https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm</a> 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>.

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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 <a href="https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products">https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products</a>); 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 <a href="https://www.fda.gov/medical-devices/medical-device-safety/medical-device-reporting-mdr-how-report-medical-device-problems">https://www.fda.gov/medical-device-problems</a>.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (<a href="https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance">https://www.fda.gov/training-and-continuing-education/cdrh-learn</a>) and CDRH Learn (<a href="https://www.fda.gov/training-and-continuing-education/cdrh-learn">https://www.fda.gov/training-and-continuing-education/cdrh-learn</a>). 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 (<a href="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">https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice</a>) for more information or contact DICE by email (<a href="DICE@fda.hhs.gov">DICE@fda.hhs.gov</a>) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

Vivek Pinto, PhD
Director
DHT5B: Division of Neuromodulation
and Physical Medicine Devices
OHT5: Office of Neurological
and Physical Medicine Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

# DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

## **Indications for Use**

510(k) Number (if known)

Form Approved: OMB No. 0910-0120

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

K211460			
Device Name Air Relax Plus Model AR-3.0			
dications for Use (Describe) he Air Relax Plus Model AR-3.0 is intended for the temporary relief of minor muscle aches and pains and for temporary crease in circulation to the treated areas in people who are in good health. The Air Relax Plus Model AR-3.0 simulates neading and stroking of tissues by using an inflatable garment.			
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.

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**Date Prepared:** April 29, 2021

**Applicant** Diode Art Engineering doing business as Air Relax

9535 Brasher St

Pico Rivera, CA 90660 Tel – 1.323.285.4231

Official Contact: Beomjoon Lee, General Manager

**Proprietary or Trade Name:** Air Relax Plus Model AR-3.0

**Common/Usual Name:** Powered Inflatable Tube Massager

Classification Name: IRP - Massager Powered Inflatable Tube (CFR 890.5650)

**Predicate Devices:** K191441 – Air Relax Model AR-1.0/AR-2.0

## **Device Description:**

This submission is for the Diode Art Engineering Air Relax Plus Model AR-3.0. The Air Relax Plus Model AR-3.0 is a powered inflatable tube massager. It is intended to temporarily relieve minor muscle aches and pains, and to temporarily increase circulation to the treated areas. It simulates manual kneading and stroking of tissues by use of an inflatable pressure cuff. The device is to be used by people who are in good health.

The device is a Class II, type BF applied part that receives power from a separately approved external IEC 60601-1 compliant power supply or optional battery pack.

The Air Relax Plus Model AR-3.0 consists of an air compressor unit with a control system, an inflatable "garment" (arms, legs and hips), plastic air tubing with a proprietary connector for connecting the device to the garment A description of each of these components is provided below. The hip garment is also referred to as "shorts".

The user interface is a front panel display and buttons.

The Air Relax Plus Model AR-3.0 contains an air compressor with a system control that allows the user to adjust the amount of air coming from the air compressorand going to the individual segments of the inflatable garment.

There is no electrical contact with the user and the device does not transfer or detect energy to or from the user.

The user interface of the Air Relax Plus Model AR-3.0 provides for starting and stopping the massage treatment, allows for adjusting time and intensity (pressure) of the treatment. The device also provides a proprietary keyed connector to the tubing

which connects to the garment. The tubing connector to the garment is also proprietary. The proprietary connectors ensure that neither the device, tubing nor garment can be misconnected to any other device or garment.

Pressure selection is performed by pressing pressure button multiple times. There are four pressure levels with approximate pressure levels as below:

Level 1: 60 mmHg Level 2: 100 mmHg Level 3: 140 mmHg Level 4: 170 mmHg

There are three modes (Progressive, Sequential, Full massage P, S and F) that determine the inflation sequence.

#### **Indications for Use:**

The Air Relax Plus Model AR-3.0 is intended for the temporary relief of minor muscle aches and pains and for temporary increase in circulation to the treated areas in people who are in good health. The Air Relax Plus Model AR-3.0 simulates kneading and stroking of tissues by using an inflatable garment.

## **Patient Population:**

Adults

#### **Environments of Use:**

Clinics, hospital, athlete training, and home environments

## Table of the Similarities and Differences of Predicate vs. Subject Device

<b>Model Name</b>	Subject Device	Predicate Device	Comment
510(k)	Air Relax Plus Model AR-3.0	Air Relax Model AR-1.0/AR-	
Number	510(k) K211460	2.0	
		510(k) K191441	
Classification	Class II Device, IRP (21	Class II Device, IRP (21	Identical
	CFR890.5650)	CFR890.5650)	
Indications	The Air Relax Plus Model AR-	The Air Relax /Compressible Identical	
for use	3.0 is intended for the	limb Sleeve System (AR-1.0,	
	temporary relief of minor	AR-2.0) is intended for the	
	muscle aches and pains and for	temporary relief of minor muscle	
	temporary increase in	aches and pains and for the	
	circulation to the treated areas	temporary increase in	
	in people who are in good	circulation to the treated areas in	
	health. The Air Relax Plus	people who are in good health.	
	Model AR-3.0 simulates	The Air Relax/ Compressible	
	kneading and stroking of	Limb Sleeve System (Model:	
	tissues by using an inflatable	AR-1.0, AR-2.0) simulates	
	garment	kneading and stroking of tissues by using an inflatable garment	
OTC or	OTC	OTC	Identical
Prescription	ore	ore	lucifical
Environment	Clinics, hospital, athlete	Clinics, hospital, athlete	Identical
of Use	training, and home	training, and home	Identical
or ese	environments	environments	
Compliance	ES 60601-1, IEC 60601-1-2,	ES 60601-1, IEC 60601-1-2,	Identical
with	IEC 60601-1-11	IEC 60601-1-11	
standards			
Mode of	Sequential/Peristaltic	Sequential/Peristaltic	Identical
Operation	•	•	
Power	100~240V, 50/60Hz	120 V, 60Hz	Although the
			power rating of
			the subject device
			is different
			both devices
			comply with
			ANSI/AAMI
			ES60601-1, so the
			difference does
			not affect safety
D :	60 170	0.220	and effectiveness.
Device	60 - 170 mmHg	0-230 mmHg	Lower
Pressure			range than
range Garments	Nylon with a Polyurethane	Nylon with a Polyurethane	K191441 Identical with
material	Laminate	Laminate	K191441
material	Lammate	Lammate	K171441

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Model Name	Subject Device	Predicate Device	Comment
510(k)	Air Relax Plus Model AR-3.0	Air Relax Model AR-1.0/AR-	
Number	510(k) K211460	2.0	
		510(k) K191441	
Leg Garment	Yes size in inches:	Yes size in inches:	Identical in size,
	L 13.4 x 8.25 x 35.6	L 13.4 x 8.25 x 35.6	construction and
	XL 15.4 x 8.25 x 43.1	XL 15.4 x 8.25 x 43.1	materials
	XXL 15.4 x 8.25 x 47.0	XXL 15.4 x 8.25 x 47.0	
Arm Garment	Yes	No	See reference
			device comparison
			below
Hip Garment	Yes	No	See reference
			device comparison
NI I C	4	4	below
Number of	4	4	Identical
Inflatable			
garment			
segments Weight	1.67kg (3.7 pounds)	1.67kg (3.7 pounds)	Identical
Dimensions	9.5" x 4.7" x 7.5"	9.5" x 4.7" x 7.5"	Identical
(W x H x D)	9.3 x 4.7 x 7.3	9.3 × 4.7 × 7.3	Identical
Housing	Molded ABS enclosure	Molded ABS enclosure	Identical
Materials and	Worded ABS enclosure	Worded ABS enclosure	Identical
Constructions			
Patient	Non-conductive garments	Non-conductive garments	Identical
contact	Tron conductive garments	Tion conductive garments	racinical
Safety	Button at control unit allows	Button at control unit allows	Identical
Features	user to stop or pause	user to stop or pause	
	therapy session at any time.	therapy session at any time.	
Modes	4 Modes :	4 Modes :	Identical to
	"P" mode inflates	"A" mode inflates	predicate except
	and deflates chambers from	and deflates chambers from	mode names differ
	bottom up, one at a time	bottom up, one at a time	
	"S" mode also inflates	"B" mode also inflates	
	chambers from bottom up, but	chambers from bottom up, but	
	maintains pressure in lower	maintains pressure in lower	
	chambers as works its way to	chambers as works its way to	
	top.	top.	
	"F" mode is inflates all	"C" mode is inflates all	
	chambers and maintained	chambers and maintained	
	pressure at same time and release pressure all chambers.	pressure at same time and	
	release pressure an chambers.	release pressure all chambers.	
	At "Target" mode, user can	At "Manual" mode, user can	
	select specific chamber to	select specific chamber to	
	inflates	inflates	

## **Comparison to reference device:**

Model Name 510(k) Number	Subject Device Air Relax Plus ModelAR-3.0 510(k) K211460	Reference Device Rapid Reboot 510(k) K182668	Comment
Indications for use	The Air Relax Plus Model AR- 3.0 is intended for the temporary relief of minor muscle aches and pains and for temporary increase in circulation to the treated areas in people who are in good health. The Air Relax Plus Model AR-3.0 simulates kneading and stroking of tissues by using an inflatable garment	The Rapid Reboot Compression Therapy System is indicated for the temporary relief of minor muscle aches and pains and for temporary increase in circulation to the treated areas in people who are in good health. The Rapid Reboot Compression Therapy System simulates kneading and stroking of tissues by using an inflatable garment.	Identical
Treatment Time	User determines Therapy time. Choosefrom 15, 30	User determines Therapy time. Choose from 10, 20, or 30 minute session time, with option to add additional 10 minutes to any therapy time.	Similar to K182668 except time is limited to 30 minutes in subject device
Arm Garment	Yes Long: 13.8" x 33.5"	Yes Regular: 18" x 38" Long: 18" x 44"	Similar in size and construction. Materials are identical to K191441
Hip Garment	Yes Large: 24.5" x 31.5"	Yes Regular : 26" x 32" Large : 26" x 35"	Similar in size and construction. Materials are identical to K191441

## **Substantial Equivalence Discussion**

In the above detailed tables we have compared the Air Relax Plus Model AR-3.0 to the predicate for equivalence of:

#### Indications -

The Air Relax Plus Model AR-3.0 is intended for the temporary relief of minor muscle aches and pains and for temporary increase in circulation to the treated areas in people who are in good health. The Air Relax Plus Model AR-3.0 simulates kneading and stroking of tissues by using an inflatable garment. **These indications are identical to the predicate.** 

**Prescriptive** – The Air Relax Plus Model AR-3.0 is OTC as is the predicate.

**Design, Technology and Principle of Operation** – The Air Relax Plus Model AR-3.0 has equivalent design and features when compared to the predicate and has identical technology to the predicate.

**Performance and Specifications** – The Air Relax Plus Model AR-3.0 has equivalent specifications of performance when compared to the predicate.

**Compliance with standards** – The subject device declares compliance with ES 60601-1, IEC 60601-1-11 and IEC 60601-1-2 which is identical to the predicate.

#### Materials -

The patient contacting materials of the Air Relax Plus Model AR-3.0 are the inflatable garments which are identical to the predicate device 510(K) K191441.

## **Patient Population –**

The Air Relax Plus Model AR-3.0 and predicates are indicated for adults.

## **Environment of Use –**

The Air Relax Plus Model AR-3.0 and predicates are for use in clinics, hospital, athlete training, and home environments.

#### Differences -

There are no differences between the proposed device and the predicate device that raise any new safety and effectiveness concerns.

### **Performance Testing**

#### Bench:

The device has been tested to ensure that all requirements have been met, this includes:

- Testing of all controls
- Testing of all indicators
- Testing of performance

The device has also been tested to the requirements of the following standards:

- AAMI / ANSI ES60601-1:2005 + A1: 2012 Medical electrical equipment part 1: general requirements for basic safety and essential performance
- IEC 60601-1-2: 2014 Collateral standard: Electromagnetic Disturbances Requirements and Tests
- IEC 60601-1-11: 2015, Collateral standard: requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment

#### **Animal:**

No animal testing was performed

#### Clinical:

No clinical testing was performed

#### Differences –

There are no differences between the proposed device and the predicate device that raise any new safety and effectiveness concerns.

## **Substantial Equivalence Rationale**

The Air Relax Plus Model AR-3.0 is viewed as substantially equivalent to the predicate device because:

**Indications** – are identical to the predicate

**Prescriptive** – The Air Relax Plus Model AR-3.0 and predicate are OTC.

**Design, Technology and Principle of Operation** – The Air Relax Plus Model AR-3.0 has equivalent design and features when compared to the predicate and has identical technology to the predicate.

**Performance and Specifications** – The Air Relax Plus Model AR-3.0 has equivalent specifications of performance when compared to the predicate.

**Compliance with standards** – The subject device declares compliance with ES 60601-1, IEC 60601-1-11 and IEC 60601-1-2 which is identical to the predicate.

**Materials** – The patient contacting materials of the Air Relax Plus Model AR-3.0 are the inflatable garments they are identical to the predicate.

**Environment of Use** – Clinics, hospital, athlete training, and home environments, not specified for predicate but predicate is OTC.

**Features -** The Air Relax Plus Model AR-3.0 has equivalent features when compared to the predicate.

#### Conclusion

The Air Relax Plus Model AR-3.0 is substantially equivalent to the predicate in indications for use, patient population, environment of use, technology characteristics, materials, specifications / performance and compliance with international standards. Minor differences as detailed in the substantial equivalence table above do not raise questions of safety and effectiveness.