

MEDICAL GRADE TREADMILLS

.==

3017 Full Vision Drive Newton, KS 67114

Phone: 316-283-3344 Fax: 316-283-9522 www.trackmastertreadmills.com

Declaration of Conformity

MANUFACTURER		
Name of company	Address	Representative
Full Vision, Inc	3017 Full Vision Drive Newton, KS 67114	Doug Pauls, Compliance Manager
SRN: US-MF-000002038	USA	

AUTHORIZED REPRE	JTHORIZED REPRESENTATIVE	
Name of company	Address	
Emergo Europe	Prinsessegracht 20 2514 AP The Hague	
SRN: NL-AR-000000116	The Netherlands	

Product name	Model/number	Basic UDI
Trackmaster Treadmill	TMX428 110V / 317-07926	
	TMX428 220V / 317-07927	08601760006FVITMR4
	TMX428CP 110V / 317-07928	
	TMX428CP 220V / 317-07929	
	T2100-ST1 / 317-07926GE / 2097357-001	
	T2100-ST2/317-07927GE/2097357-002	
Intended Purpose	Photo	
The medical tree desille and inte	11	

The medical treadmills are intended as stressing devices, by providing motion to patient, to be interfaced with a variety of cardiac and pulmonary stress testing systems. The treadmill is intended to be operated by the physician, therapist, or operator acting under authorization of the physician with training per IFU under the supervision of a physician and / or therapist, with sufficient knowledge of the indications and contraindications. The medical treadmills are intended to be used in a medical facility or wellness center.

Certain models have a control panel to operate the treadmill.

Caution: Treadmill does not provide any kind of medical treatment diagnostic or assessment.

	7/1	
/-		
	6	

CONFORMITY ASSESSMENT		
Device classification	Route to compliance	Standards applied
Class 1 Rule 13	Annex II and III of MDR 2017/745	See List below

Standard Number	Standards Organisation	Standard Title	Version
13485	ISO	Medical Devices Quality Management Systems Requirements	2016
14971	ISO	Medical Devices – Application of risk management to medical devices	2019
15223-1	ISO	Medical devices – Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General requirements	2016
17664	ISO	Processing of health care products – Information to be provided by the medical device manufacturer for the processing of medical devices	2017
62304	IEC	Medical device software – Software life-cycle processes	2006/AMD 1:2015
62366	IEC	Medical devices – Application of usability engineering to medical devices	2016
10993-1	ISO	Biological evaluation of medical devices – Part 1: Evaluation and testing within a risk management process	2018
60601-1	IEC	Medical electrical equipment – Part 1: General requirements for basic safety and essential performance	2005+A1:2012
60601-1-1	IEC	Medical electrical equipment – Part 1-1: General requirements for safety – Collateral standard: Safety requirements for medical electrical systems	2000
60601-1-2	IEC	Medical electrical equipment – Part 1-2: General requirements for basic safety and essential performance – Collateral standard: Electromagnetic disturbances – Requirements and tests	2014
60601-1-6	IEC	Medical electrical equipment – Part 1-6: General requirements for basic safety and essential performance – Collateral standard: Usability	2010+A1:2013
61000-3-2	IEC	Electromagnetic compatibility (EMC) – Part 3-2: Limits – Limits for harmonic current emissions (equipment input current ≤ 16 A per phase)	(Ed:4.0): 2014
61000-3-3	IEC	Electromagnetic compatibility (EMC) – Part 3-3: Limits – Limitation of voltage changes, voltage fluctuations and flicker in public low-voltage supply systems, for equipment with rated current ≤ 16 A per phase and not subject to conditional connection	(Ed:3.0): 2013
D4169-16	ASTM	Standard Practice for Performance Testing of Shipping Containers and Systems	16

RoHS Compliance

The previously listed products, manufactured by Full Vision Inc, as well as the accessories and options available, are in compliance with the RoHS Directive (EU Directive 2002/95/EC and subsequent amendments). RoHS Compliant means that the substances restricted by the EU Directive 2011/65/EU (RoHS 2) and EU Directive 2015/863/EU (RoHS 3) and subsequent amendments of the European Parliament are not contained in a finished product above threshold limits stated below.

Restricted Substance	Maximum Threshold Limit
Lead	0,1 %
Mercury	0,1 %
Cadmium	0,01 %
Hexavalent chromium	0,1 %
Polybrominated biphenyls (PBB)	0,1 %
Polybrominated diphenyl ethers (PBDE)	0,1 %
Bis (2-ethylhexyl) phthalate (DEHP)	0,1 %
Butyl benzyl phthalate (BBP)	0,1 %
Dibutyl phthalate (DBP)	0,1 %
Diisobutyl phthalate (DIBP)	0,1 %

REACH Compliance

Full Vision, Inc.

The EU REACH Regulation (EC) No 1907/2006 of the European Community, concerning the Registration, the Evaluation and the Authorization of the Chemical substances (REACH), was released in June 2007. The instruction of the regulation establishes specific duties and obligations for Companies in the European Union (EU) that manufacture or import substances on their own, in preparations, as well as in articles. Under the structure of the REACH regulation, **Full Vision Inc**, is a manufacturer and supplier of "articles" to our EU Customers, and is therefore not obliged to register or pre-register the products we supply. Full Vision Inc products (articles) do not contain Substances of Very High Concern (SVHC) above the threshold value declared as per ECHA. Customers are encouraged to visit the site https://echa.europa.eu/candidate-list-table for the most up to date information on the current list of SVHCs under REACH.

This declaration of conformity is issued under the sole responsibility of **Full Vision, Inc**. **Full Vision, Inc** declares the device(s) that is covered by the present declaration is in conformity with this Regulation (EU) MDR 2017/745 and, if applicable, with any other relevant Union legislation that provides for the issuing of an EU declaration of conformity.

Authorized Signatory Bran Clayer	4-27-21	ORPOR 1
Signature	Date of issue	SEAL
Brian Hayes Name		TANSAS
Medical Device Business Manager	Newton, KS	Ę
Title	Place of Issue	Company Stamp

MDR DoC Trackmaster Treadmills Rev 3

Revision	Description	Author	Date
1	Initial Release to EU MDR 2017/745	B. Tucker	2021/02/02
2	Added Full Vision SRN number, corrected Route to compliance to Annex II and III instead of Annex IV, added RoHS directive (EU) 2015/863, Added REACH regulation 1907/2006,	B. Tucker	2021/03/15
3	Added additional model numbers to product identification for the T2100-ST1 and ST2	B. Tucker	2021/04/27