



Read this manual prior to performing any task!

miha bodytec Inc.
2171 Executive Drive Suite 200
Addison, Illinois 60101
USA
Telephone: +1 833 367 6442
E-mail: usa@miha-bodytec.com
Internet: www.miha-bodytec.us

TD150168 – Rev. 0

This manual was created by:
kothes GmbH
Internet: www.kothes.com
© miha bodytec Inc. 2021



Conscientious handling of the miha bodytec m.ove

Conscientious handling is the prerequisite for successful and safe use. The trainer must take part in a training session held by the manufacturer to be able to safely operate the EMS training device, hereinafter referred to as “device”.

Before starting with the workout, the trainer instructs the athlete/patient in the basic operating functions and displays on this device for the workout. The trainer points out consequences caused by misuse.

Information about this manual

This manual enables safe and efficient handling of the device. This manual is an integral part of the device and must be kept in the immediate vicinity of the device and accessible to the trainer at any time.

The trainer must be able to read and understand this manual. The trainer must have carefully read and understood this manual before handling the device. The prerequisite for safe training is compliance with all safety instructions and procedural instructions specified in this manual. The local regulations for the prevention of accidents and the general safety regulations for the location in which the device is used also apply.

The illustrations in this manual provide a basic understanding and may vary from the actual design.

Safety instructions

Safety instructions in this manual are identified by symbols. The safety instructions are introduced by signal words which express the extent of the hazard.

 **DANGER**

DANGER indicates a hazardous situation which, if not avoided, will result in death or serious injury.

 **WARNING**

WARNING indicates a hazardous situation which, if not avoided, could result in death or serious injury.

 **CAUTION**

CAUTION indicates a hazardous situation which, if not avoided, could result in minor or moderate injury.

NOTICE

NOTICE indicates important, non-safety-related information, such as property and environmental damage.

Safety instructions in procedural instructions

Safety instructions can refer to specific, individual procedural instructions. Such safety instructions are embedded in the procedural instructions so that they do not interrupt the reading flow when performing the activity. The signal words described above are used.

Example:

➔ **⚠ WARNING! Risk of injury if detaching the electrodes while the training program is running!**

Make sure that no training program is active. To do so, switch to the main menu.

Tips and recommendations



This symbol highlights useful tips and recommendations, as well as useful information for efficient and fault-free operation.

Identifiers in this manual

To highlight procedural instructions, results, lists, references, and other elements, the following identifiers are used in this manual:

Identifier	Explanation
➔	Step-by-step procedural instructions
⇒	Results of procedural steps
↗	References to sections of this manual and to other applicable documents
■	Lists without a defined sequence
[Button]	Controls (e.g., buttons, switches), display elements (e.g., signal lamps)
"Display"	Screen elements (e.g., buttons, assignment of function keys)

Other applicable documents

The following document applies in addition to this manual:

- User manual for i-body® connect wireless
- Installation manual for travel station m.ove
- Installation manual for work station m.ove

Copyright

The contents of this manual are protected by copyright. Use of these contents is permissible within the framework of use of the device. Any other use is not permitted without the written permission of the manufacturer.

Table of contents

1	Overview and scope of delivery	7
1.1	Scope of delivery	7
1.2	Equipment	10
2	Safety	17
2.1	Intended use	17
2.2	Misuse	18
2.3	Risks due to physical condition: Contraindications	19
2.4	A team for your safety	20
2.5	Warnings	21
2.6	Reporting of adverse events	29
2.7	Symbols on the miha bodytec m.ove	30
2.7.1	Symbols on device	30
2.7.2	Rating plates	31
2.7.3	Cleaning symbols	34
2.8	Environmental protection	35
3	Technical data	36
4	Electromagnetic compatibility	42
4.1	Compliance details	42
4.2	EMC Warnings	46
5	Basic information on EMS training	48
5.1	Safety first	48
5.2	EMS training	48
5.3	Training frequency and regeneration	49
6	Understanding the device	52
6.1	Explanation of terms	52
6.2	Opening menu tabs	55
6.3	Opening menus	56
6.4	Entering text	59
7	Customization and setup	61
7.1	Adjusting the device setup	61
7.2	Selecting the language	62
7.3	Program memory settings	63
7.4	Setting up the training plan memory	63
7.5	Setting up synchronized start	64
7.6	Adjusting the favorites menu	66
7.7	Backing up data	67
7.8	Restoring/importing data	68
7.9	Installing updates	69
7.10	Using networks	70
7.11	Resetting to factory settings	71
7.12	Displaying statistics	71

Table of contents

8	Before the training	72
8.1	Eating and drinking.....	72
8.2	Selecting undergarments and shoes.....	72
8.3	Moistening and applying electrodes.....	73
9	Training with the device	83
9.1	Safety during training.....	83
9.2	Connecting the device.....	84
9.3	Operation with the rechargeable battery.....	85
9.4	Individual training settings.....	85
10	Overview of programs/training plans	87
10.1	Selecting individual training settings.....	87
10.2	Stress standards.....	87
10.3	Training programs.....	88
10.4	Training plans.....	89
10.5	Training.....	94
10.6	Transponder system.....	98
10.7	Training with synchronized start.....	102
11	After the training	103
11.1	Taking off electrodes.....	103
11.2	Cleaning and storage.....	104
11.2.1	Cleaning the i-body® and the individual electrodes.....	104
11.2.2	Cleaning the external cables.....	106
11.2.3	Cleaning the control unit.....	107
12	Packaging and storage	108
12.1	Symbols on the packaging.....	108
12.2	Transport and storage.....	109
12.3	Environmental protection.....	110
13	Ensuring optimum function	111
13.1	Maintaining the device.....	111
13.2	Maintaining the electrodes.....	112
13.3	Handling error messages.....	112
14	Index	113

1 Overview and scope of delivery

1.1 Scope of delivery

miha bodytec m.ove

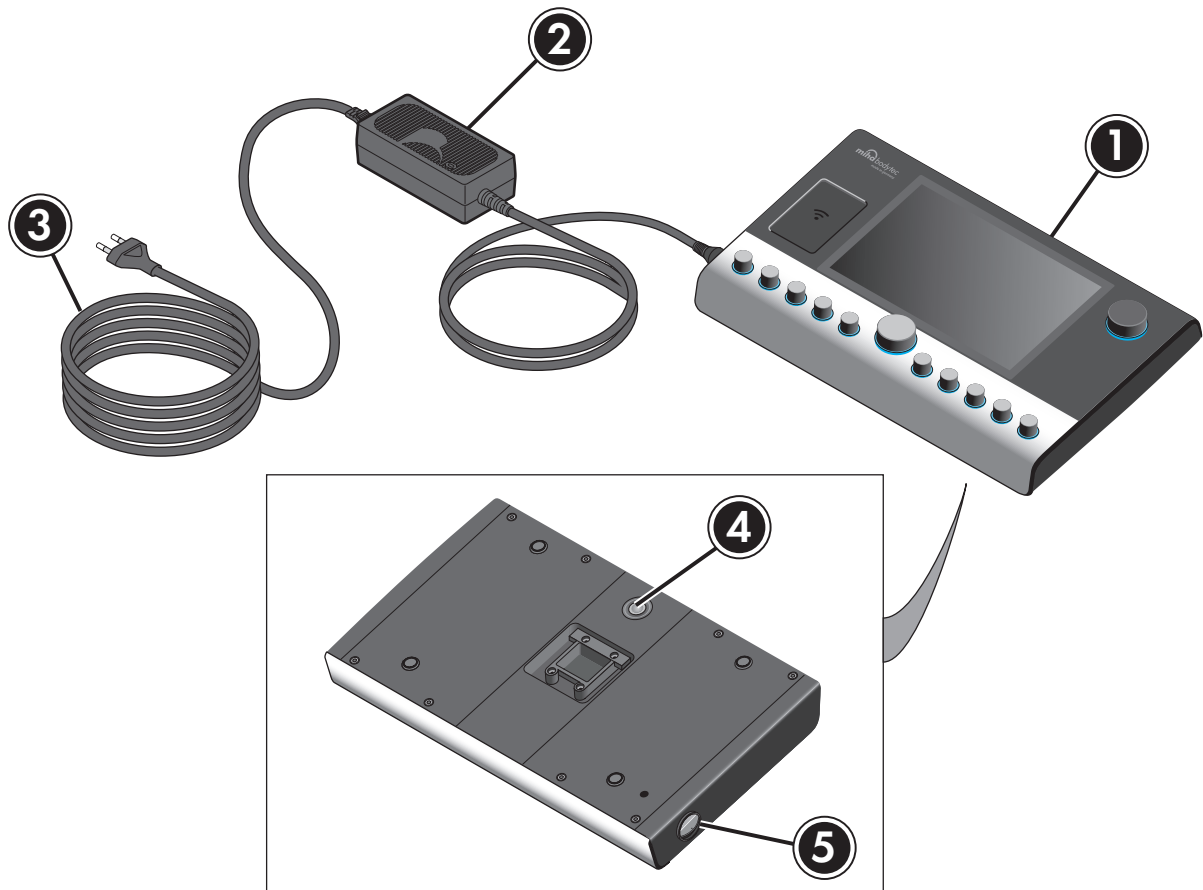


Fig. 1: Overview of miha bodytec m.ove

- 1 Control unit (🔗 “Control unit (top side)” on page 8)
- 2 Power supply

- 3 Power cable
- 4 On/off switch
- 5 Connection for power supply unit incl. cover

What is EMS training?

EMS stands for **E**lectro**M**yo**S**timulation. This means that the muscles are stimulated by electrical pulses. Complete body training which addresses all muscle groups is possible with up to 10 pairs of electrodes.

During conventional training, electrical signals from the brain initiate contractions and thus movement of the muscles. With EMS, electrical pulses from outside activate the muscles. It makes no difference to the muscle whether electrical stimuli are sent from the brain or from electrodes: It reacts by contracting.

Overview and scope of delivery

Special features and benefits:

- All muscle groups can be activated using an electrode system with up to 10 pairs of electrodes.
- Static and dynamic training with interaction of deliberate muscle contraction and EMS.
- Exercise postures increase the contraction of the deliberately stimulated muscles.
- Positive and negative electrodes are not on the same muscle.
- Agonist and antagonist are stimulated simultaneously.
- EMS training can cause more intense muscle contractions than classic strength training. At the same time, there is comparatively very little stress on the joints.
- Deeper muscle groups can be easily reached.
- The training is very intense and thus short (10 – 20 minutes).

Control unit (top side)

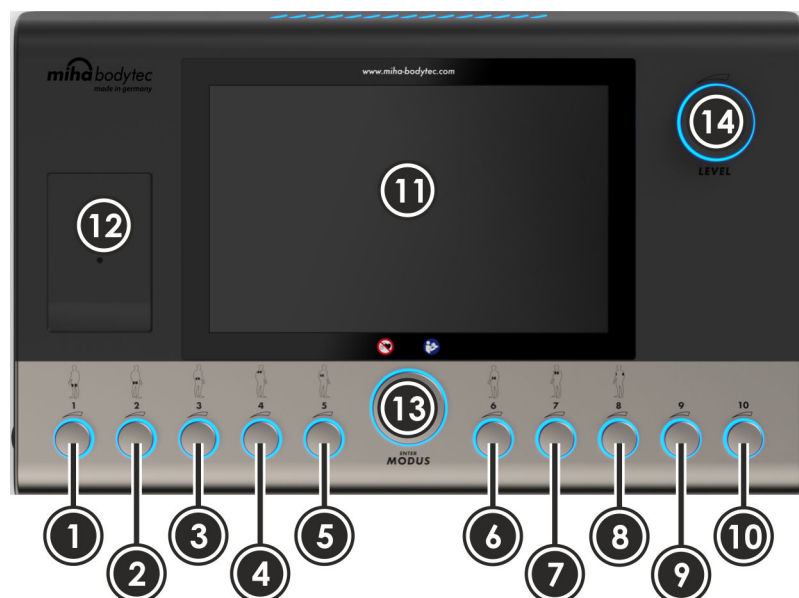


Fig. 2: Control unit (top side)

- 1 Level controller for legs
- 2 Level controller for buttocks
- 3 Level controller for lower back
- 4 Level controller for upper back
- 5 Level controller for side back
- 6 Level controller for abdomen
- 7 Level controller for chest
- 8 Level controller for arms
- 9 No function assigned
- 10 No function assigned
- 11 LC display
- 12 Transponder card contact surface
- 13 Multi-function button for start/stop and for setting the programs and the program parameters
- 14 Main controller for setting the pulse strength

All controls with the exception of the on/off switch, which is located on the underside (Fig. 3), are located on the top side of the control unit.

The main controller (Fig. 2/14) enables the athlete to regulate pulse strength of all outputs simultaneously. A connected pair of electrodes is an output in this context.

The pulse strength of individual outputs can be regulated using the relevant level controllers (Fig. 2/1 – 8). The image above each of these rotary switches shows the relevant part of the body for which the pulse strength is regulated. For example, using the level controller for legs (Fig. 2/1), the pulse strength sent to the legs is regulated using the i-body® straps flex.

The two level controllers without any assigned body parts (Fig. 2/9, 10) do not have any function.

If a transponder card is placed on the transponder card contact surface (Fig. 2/12), the settings stored on this card can be edited or loaded automatically (↪ *Chapter 10.6 “Transponder system” on page 98*).

The multi-function button (Fig. 2/13) is used for navigating through the menus and for activating programs. The multi-function button is also used for starting and stopping workouts.

The LC display (Fig. 2/11) shows menus and provides status information on the device. A virtual person – an avatar – is shown on the LC display during training. The athlete can use the movements of the avatar, in addition to the instructions of the trainer, as an orientation.

Control unit (underside)

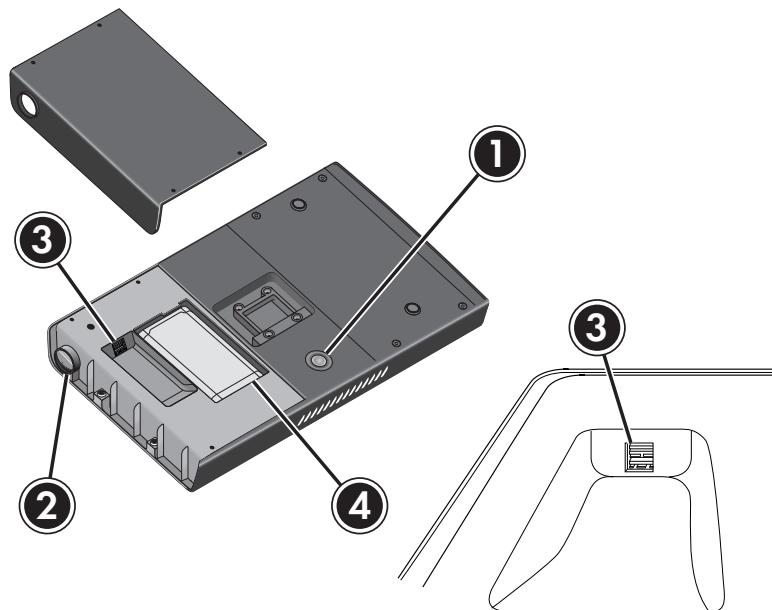


Fig. 3: Control unit (underside)

- 1 On/off switch
- 2 Power supply connection incl. cover
- 3 USB port
- 4 Rechargeable battery

The on/off switch (Fig. 3/1) is embedded in the case on the underside of the control unit. A built-in terminal strip is located in a depression on the underside, which is protected by a service cover.

The supply cable of the power supply unit is connected to the power supply connection (Fig. 3/2) by means of a magnetic plug connection.

The miha bodytec USB flash drive can be inserted into the USB port (Fig. 3/3) for performing updates and for saving device settings.

The rechargeable battery (Fig. 3/4) located in a designated rechargeable battery compartment incl. connector is used to supply power without a power supply during mobile use.

1.2 Equipment

Safety thanks to genuine miha bodytec equipment



Using equipment not obtained from miha bodytec or authorized dealers represents an increased safety risk. Only use genuine miha bodytec equipment.



It is not allowed to connect additional parts not mentioned in this manual to the device, the electrodes or the cables.

Rechargeable battery for m.ove



Rechargeable battery for miha bodytec m.ove

Fig. 4: Rechargeable battery for m.ove

travel station m.ove



Mobile floor stand for storing the equipment, with a holder for the control unit when in mobile use.

For further information, please refer to [TD 15303 – Installation instructions for travel station m.ove](#).

Fig. 5: travel station m.ove

Overview and scope of delivery

work station m.ove



Medical cart for storing the equipment, with a holder for the control unit when in mobile use in specific facilities.


For further information, please refer to  TD 15304 – Installation instructions for work station m.ove.

Fig. 6: work station m.ove

i-body® connect wireless



Wireless stimulation unit that can be attached to the body of the athlete for wireless training via a Bluetooth® connection.

See the “i-body® connect wireless” manual for further information.

Fig. 7: i-body® connect wireless

i-body® connect charger



Fig. 8: i-body® connect charger

Charger for i-body® connect wireless.

See the “i-body® connect wireless” manual for further information.

Transponder card



Fig. 9: Transponder card

Transponder card for storing the last values, individual time specification, and individually adjusted program.

i-body®



Fig. 10: i-body®

The i-body® electrode vest for applying electrodes to the upper body.

Table 1: Electrode sizes for each vest size

Vest size	Size 1 and size V1	Size 2, size 3 and size V2 in in ²
2 electrodes for abdomen	22.85 in ²	27.56 in ²
2 electrodes for chest	10.31 in ²	12.81 in ²

Overview and scope of delivery

Vest size	Size 1 and size V1	Size 2, size 3 and size V2 in in ²
2 electrodes for upper back	16.25 in ²	20.98 in ²
2 electrodes for sides of back	9.75 in ²	11.85 in ²
2 electrodes for lower back	14.4 in ²	19.47 in ²

i-body® straps flex



Electrodes for applying to arms and legs.

Table 2: Electrode sizes for each i-body®straps flex size:

Size of the i-body®straps flex	Electrode size
Size 1 (pair)	16.74 in ² each
Size 2 (pair)	20.77 in ² each
Size 3 (pair)	29.14 in ² each

Fig. 11: i-body® strap flex

i-body® belt



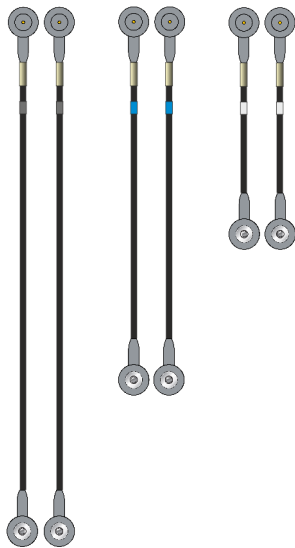
Buttocks electrode, hereinafter referred to as “i-body® belt”, for applying to the buttocks.

Table 3: Electrode sizes for each i-body®belt size:

Size of the i-body®belt	Electrode size
Size 1: 2 buttocks electrodes	each 14.7 in ²
Size 2: 2 buttocks electrodes	each 18.34 in ²

Fig. 12: i-body® belt

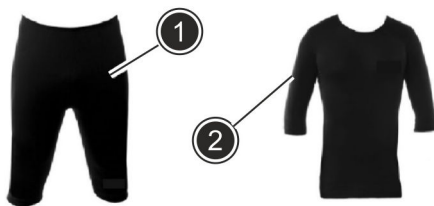
i-body® cable



External cables, hereinafter referred to as “i-body® cables” for connecting the “i-body®” to the external “i-body® straps flex” and “i-body® belt” electrodes.

Fig. 13: i-body® cable

Undergarments



- 1 Pants
- 2 Top

Undergarments for wearing under all electrodes to be applied.

Fig. 14: Undergarments

Pump spray bottle



Pump spray bottle for moistening the electrodes.

Fig. 15: Pump spray bottle

miha bodytec USB flash drive

The miha bodytec USB flash drive is delivered with the device and serves as a data carrier for performing updates as well as saving and transferring settings.

Overview and scope of delivery



In the event of important updates, a miha bodytec USB flash drive will be sent to users of the device who perform the update themselves or have it performed on the device by the trainers.

Customer service

For technical information, please contact us at our headquarters:

Address	miha bodytec Inc. 2171 Executive Drive Suite 200 Addison, Illinois 60101 USA
Telephone	+1 833 367 6442
E-mail	usa@miha-bodytec.com
Website	www.miha-bodytec.us

We are also always interested in information and experience arising from the use of the device which can be valuable for the improvement of our product.

2 Safety

2.1 Intended use

miha bodytec m.ove is a device which performs electronic muscle stimulation based on EMS technology. The device is specifically designed as an addition to other sports and for training muscles.

miha bodytec m.ove is intended to stimulate muscles in order to improve or facilitate muscle performance. In addition it is indicated for the following conditions:

- Re-educating muscles
- Relaxation of muscle spasm
- Retarding or preventing disuse muscle atrophy

The miha bodytec m.ove electrical impulses allow the triggering of action potentials on motoneurons of motor nerves (excitations).

These excitations of motoneurons are transmitted to the muscle fibers via the motor endplate where they generate mechanical muscle fiber responses that correspond to muscle work.

Depending on the parameters of the electrical impulses (pulse frequency, duration of contraction, duration of rest, total session duration), different types of muscle work can be imposed on the stimulated muscles.

miha bodytec m.ove may only be used by persons above the age of 21.

2.2 Misuse

WARNING

Danger in the event of misuse!

- Keep the device out of the reach of children.
- Never apply electrodes to positions other than those described in this manual.
- Never use the electrical stimulation treatment in the following areas of the body:
 - On or through the head
 - Directly on the eyes
 - In areas around the mouth
 - On the front of the neck (especially the carotid artery)
 - With electrode surfaces applied to the chest and the upper back or across the heart
 - Over the menstruating or pregnant uterus
 - Over areas of the skin with a lack of normal sensation.
- Stimulation must not be applied over the carotid sinus nerves, particularly in patients with a known sensitivity to the carotid sinus reflex.
- Stimulation must not be applied over the neck or mouth. Severe spasm of the laryngeal and pharyngeal muscles may occur and the contractions may be strong enough to close the airway or cause difficulty breathing.
- Stimulation shall not be applied transthoracically. The introduction of electrical current into the heart may cause cardiac arrhythmias.
- Stimulation must not be applied transcerebrally.
- Stimulation must not be applied over swollen, infected, or inflamed areas or skin eruptions, e.g., phlebitis, thrombophlebitis, varicose veins, etc.
- Stimulation must not be applied over, or in proximity to, cancerous lesions.
- Whenever using the device, start with an intensity level of “0” and increase slowly.
- Ensure basic tension in the muscle group you are exercising for each pulse to prevent uncontrolled muscle contractions.
- In case of skin irritation and signs of burns beneath the electrodes, stop using the device immediately.
- If you are feeling unwell, dizzy, or have heart pain, stop using the device immediately.
- Some patients may experience skin irritation or hypersensitivity due to the electrical stimulation or electrical conductive medium. The irritation can usually be reduced by alternate electrode placement using an alternate conductive medium.
- Electrode placement and stimulation settings must follow guidance of the prescribing practitioner.
- The long-term effects of repeated electrical stimulation are unknown.
- Only use genuine miha bodytec equipment and spare parts.

Misuse of the device can cause hazardous situations. The device is exclusively intended for use as an electronic muscle stimulator for the usecases described in this manual and must only be used in a dry environment. Skin irritation and burns beneath the electrodes have been reported with the use of powered muscle stimulators.

2.3 Risks due to physical condition: Contraindications

EMS training in the event of existing contraindications

WARNING

Risk of fatal injury in the event of use despite presence of contraindications for specified persons!

All persons training with the device must ensure that none of the following contraindications are present:

- Before each further training session the trainer must check if the health condition of the athletes/patients has changed and/or if they are under influence of alcohol, drugs, narcotics, and/or painkillers.
- In the event of doubts whether any contraindication is present, do not use the device.
- In the event of doubts, consult a doctor before the first or any further trainings.
- If you are taking medication, consult a doctor before the first or any further trainings.

The presence of certain physical conditions can result in a risk of injuries or even death. The existence of such a physical condition is referred to as contraindication.

Never exercise if ...

The device must not be used if the following contraindications are present:

- Acute influence of alcohol, drugs, narcotics, and/or painkillers
- Recent operations
- Cardiac arrhythmias
- Active medical implants
- Epilepsy
- Seizures
- Pregnancy or suspected pregnancy
- Severe circulatory disorders
- Arterial circulatory disorders
- Strong bleeding tendencies (hemophilia)
- Bleeding
- Abdominal wall hernia
- Inguinal hernia
- Tuberculosis
- Tumor diseases
- Arteriosclerosis in advanced stage
- Severe neurological disorders

- Diabetes mellitus
- Febrile diseases
- Acute bacterial or viral infections
- Liver diseases
- Kidney diseases
- Cardiovascular diseases
- Coronary heart diseases
- Infected or wounded areas of the skin
- Skin cancer
- Rhabdomyolysis

2.4 A team for your safety

Acting responsibly

This section describes the requirements for people involved in training with this device. To ensure safe EMS training, the actions described in this manual may only be performed when the relevant person meets the specified requirements.

Athlete/Patient

Athletes/patients use the device actively for EMS training. They make sure that they are instructed by a qualified trainer of the training-relevant control elements and displays and notified of the consequences caused by misuse. They also make sure that they are guided through the training session by the trainer. Athletes/patients must provide the trainer responsible for them with truthful information on current health restrictions (e.g., fever).

Athletes/patients make sure that none of the listed contraindications (🔗 *Chapter 2.3 “Risks due to physical condition: Contraindications” on page 19*) apply to them before they start with the training.

Trainer

The trainer must have taken part in a basic miha bodytec training session and can prove their qualification by presenting a certificate issued in their name. The participation in an additional higher-level EMS training session is recommended. This following information is covered during this training session:

- Effects and special characteristics of EMS training
- Adjustable parameters of the device and their impact
- Identifying and avoiding risks:
 - Function-related residual risks
 - Settings-related residual risks
 - Residual risks due to the condition of the athlete/patient
- Available equipment and its correct usage
- Applying the electrodes
- Function, operation, and programming of the control unit
- Performing updates

The trainer applies the information they learned in the basic miha bodytec training session so that everyone training with the device has a safe EMS workout. In particular, the trainer is responsible for the following:

- Ensuring that updates are integrated promptly following release.
- Ensuring that athletes/patients are healthy enough for EMS training in advance of each training session.
- Ensuring that defective parts are replaced when problems become known.
- Having the equipment properly cleaned after the EMS training session or cleaning it.

The trainer must be qualified in accordance with the local specifications to administer first aid and has a basic knowledge in anatomy, physiology, training theory and training planning.

During a workout, the trainer must focus exclusively on the needs of the athlete(s)/patient(s). They do not actively participate in the workout and are not hooked up to the device but inform and monitor the athletes/patients. A maximum of 2 athletes/patients per 1 trainer is allowed. Before, during, and after the workout, the trainer enquires about the physical condition of the athletes/patients and checks them so risks can be eliminated.

During a workout, the trainer and the athletes/patients must have unimpeded access to the device's control elements. They must be able to use and regulate the device easily, quickly, and with precision.

The full attention of the trainer is required at all times during the workout. If this cannot be ensured, the trainer must reduce the main intensity level of the device/athlete/patient by at least 50 per cent.

The trainer must ensure that all requirements above are met.

Private use of the device

The EMS training with the device must be supervised by a qualified trainer. This also applies to private use of the device. Even if the athlete/patient has qualified as a trainer, they may not use the device without qualified supervision.

Only the mandatory basic miha bodytec training by the manufacturer and an optional additional higher-level EMS training qualify a person as a trainer as described above.

2.5 Warnings

Getting to know the residual risks

The following section specifies risks which can arise from the device even when it is used as intended.

Comply with the safety instructions listed here and with the safety instructions in the other sections of this manual to reduce risks of personal injury and material damage and to avoid dangerous situations.

Interactions

DANGER

Risk of fatal injury from interactions!

Do not connect athletes/patients to a high-frequency surgical appliance simultaneously. This can result in burns under the stimulation current electrodes.

- Keep a minimum distance of 3 ft / 1 m between the device and a short-wave or microwave therapy appliance. Otherwise, this can cause fluctuations in the output values of the stimulation current device.
- Never use the device on people who are fitted with active (e.g., electronic) medical implants.

Interactions with the device can be dangerous and cause injuries including death. Applying the electrode surfaces in the vicinity of the chest can increase the risk of ventricular fibrillation.

Cardiac arrhythmias

So far, there has been no known event of cardiac arrhythmia that can be attributed to training with the device. Since a theoretical residual risk remains, strictly observe the instructions below.



Persons affected will lose consciousness after a brief period of time.

WARNING

Health risk due to cardiac arrhythmia!

- Never exercise by yourself. Always ensure that a qualified trainer is paying attention and can intervene in the event of an emergency.
- Act immediately if the athlete/patient loses consciousness:
 - Disconnect the cables from the electrodes. To do so, unplug the connector plug from the vest.
 - Notify emergency medical services.
 - Administer CPR until emergency medical services arrive.
 - If you have access to a defibrillator, defibrillate the athlete/patient.

The EMS training involves a residual risk which may lead to cardiac arrhythmia. If untreated, cardiac arrhythmia may result in death.

Electrical voltage

**Risk of fatal injury from power supply!**

- Only use the device in a dry environment.
- With the exception of the service cover, never open the control unit.
- Never open the power supply.
- Always inform Customer Service (📞 “Customer service” on page 16) in the event of problems with the device.
- In the event of damage to the insulation, switch off the power supply immediately and unplug the mains plug.
- Observe the safety rules:
 - Pull the mains plug.
 - Make sure that nobody reconnects the mains plug to the power supply.
- Never connect the power supply unit to any voltage other than 100–230 V ~ 47–63 Hz.
- Only use the genuine power supply unit and the genuine power supply cable from the manufacturer. Both are available from Customer Service.
If the power supply unit has to be replaced, only use the genuine power supply unit (GlobTek Medical/ ITE Power Supply; model no.: GTM96600-6518-T2).
- Only use the genuine rechargeable battery from the manufacturer. It is available from Customer Service.
If the rechargeable battery needs to be replaced, use only the genuine rechargeable battery (Battery Pack – miha bodytec m.ove; model no.: RRC2054-2).
For information on how to replace the rechargeable battery, please contact Customer Service (📞 “Customer service” on page 16).

There is an immediate risk of fatal injury due to electrocution in the event of contact with live parts. Damage to the insulation or individual components can have fatal consequences. There is a risk of fatal injury in the event of liquid penetrating the case!

Improper exposure of the electrodes to moisture

**Risk of fatal injury from improper exposure of the electrodes to moisture!**

- When moistening and putting on the electrodes, ensure that there is a sufficient distance to the control unit. No water must penetrate the case.

There is a risk of fatal injury due to electrocution in the event of liquid penetrating the case of the control unit!

Damage to cables and connections

WARNING

Damage to the cables and the connections!


- Do not place any heavy objects on the power supply cable.
- Never kink or pinch the power supply cable.
- Always keep the wall outlet accessible.
- Switch off the device and unplug the mains plug before cleaning and maintenance.

There is the risk of material damage and injuries in the event of damage to the cables or the connections.

Improper routing of connection cables

WARNING

Risk of injury from improper routing of connection cables!


- Always route the connection cables along the “work station m.ove” or the “travel station m.ove” (see  “*travel station m.ove*” on page 11) and secure them with clips.
- If the device is set up to be stationary, secure the connection cable between the device and the wall connection with a cable bridge.

Improper routing of connection cables can lead to trip hazards or strangulation of babies and toddlers/children as well as handicapped persons.

Detaching or connecting electrode cables during operation

WARNING

Risk of injury if detaching or connecting electrode cables during operation!

- Start by making sure that no training program is active ( “*Opening the main menu*” on page 58), then connect or detach the electrodes.
- Never detach electrode cables while training.
- Never connect electrode cables while training.

Detachment or connection of electrode cables can result in injuries that depend on the settings, such as torn muscle fibers or shoulder dislocations.

Lithium-ion battery

WARNING

Danger from the lithium-ion battery!

- Do not open or dismantle batteries.
- Do not subject batteries to heat or fire. Avoid storage in direct sunlight.
- Do not short-circuit the battery.
- Do not store batteries without packaging in containers where short circuits with loose metal objects could occur.
- Do not remove battery from original packaging until immediately before use.
- Avoid mechanical impacts.
- If fluids escape from the battery, avoid contact with the skin or eyes. In case of contact, wash affected areas with plenty of water and consult a doctor.
- Only use suitable chargers.
- To ensure that the device functions correctly, observe the “plus” (+) and “minus” (-) markings on the battery when inserting it.
- Store batteries out of the reach of children.
- Keep batteries clean and dry.
- Charge secondary batteries before use.
- Use batteries only for the use for which they are intended.
- Do not store batteries in a discharged state for more than 1 month.
- Do not store batteries for more than 1 year without charging them.

Misuse or improper storage of the lithium-ion battery can cause dangerous situations.

Hygiene

WARNING

Risk of injury from improper hygienic cleaning of the control elements of the device and equipment!

- Properly clean the control elements of the device and the equipment used after training.
- Before placing the device into intermediate storage, mark the components with the date and name of the last trainer.

In the event of improper hygienic cleaning of the control elements of the device and equipment, bodily fluids can lead to skin reactions and injure athletes/patients with skin problems!

Inhaling or swallowing

WARNING

Risk of injury due to inhaling or swallowing cable clips!

- Keep cable clips in a safe place out of the reach of babies, infants/children, and handicapped persons.

Inhaling or swallowing cable clips can lead to injuries.

Allergic reaction



There are currently no known allergic reactions to materials used by miha bodytec.

What trainers and athletes/patients have to know

WARNING

Danger due to insufficient level of knowledge!

- miha bodytec m.ove may only be used by persons above 21 years.
- Electrodes may only be connected and detached by the trainer or under their supervision.
- Only work out under the supervision of a qualified trainer.
- Only work out under the supervision of a physician or therapist when using the device for the following indications:
 - Re-educating muscles
 - Relaxation of muscle spasm
 - Retarding of preventing disuse muscle atrophy
- Only train with the device if you have been instructed on how to handle it.
- Never allow other persons to operate the device without previous training.
- The manual must have been read completely and fully understood.
- The operating principle of EMS and the effects on the body must be known.
- Children are not allowed to train or play with the device or perform cleaning or maintenance tasks.

Handling the device without training/instruction can result in dangerous situations that can be fatal. Each trainer must take part in a training session held by the manufacturer before using the device and each athlete/patient must have been instructed by the trainer.

Individual adjustment



The individual adjustment of the device is required to support correct training techniques, training positions, and training processes.

Slippery floor

CAUTION

Danger of injury from slipping!

- Clean up accumulations of liquid, such as water and perspiration immediately.
- Wear slip-resistant training shoes.

Floors can be slippery due to perspiration or water. Slipping and falling can result in injuries.

Getting caught

WARNING

Risk of injury due to getting caught in drawers!

- Operate the drawers only with the designated recessed handles.
- Always operate drawers with care.

During operation, fingers can get caught in the drawers of the “work station m.ove” and thus be injured.

Being run over

WARNING

Risk of injury due to running over body parts!

- Move the “work station m.ove” carefully.
- Only use the “work station m.ove” on level ground; secure it to prevent it rolling away if necessary.
- Check the travel routes and keep them clear.
- Wear sturdy shoes when moving the “work station m.ove.”

When the “work station m.ove” is moved, there is a risk that it can run over feet or toes and get them caught under the rollers. Because the “work station m.ove” can be very heavy, this can cause serious injuries.

Unsuitable accessories and incorrect spare parts

WARNING

Risk of injuries from incorrect spare parts and unsuitable accessories!

- Never make changes to the device.
- Never establish connections with other devices not described in this manual.
- Never connect parts other than genuine parts from miha bodytec.
- Only operate the device in combination with the “work station m.ove” or the “travel station m.ove.”
- Never use accessories not described in this manual or approved for use by miha bodytec.

Incorrect spare parts or unsuitable accessories can damage the device and impair its functioning. This can result in injury. In particular, using accessories not obtained from miha bodytec represents an increased safety risk.

Improper use of accessories

WARNING

Risk of injuries due to improper use of accessories!

- Never perform training with the “travel station m.ove” in the transport situation. The control unit can only be applied in order to start training once the device has been converted to the stationary floor stand version.
- Never use the handle of the “travel station m.ove” as a training rod. The handle is designed only for stabilization and for assisting with balance in training situations.
- Never use the handle of the “work station m.ove” as a training rod. The handle is designed only for stabilization and for assisting with balance in training situations.

Improper use of accessories can cause damage to the device and accessories. This can result in injury. In particular, incomplete assembly for the training situation and improper stress on the accessories present an increased safety risk.

Pre-existing conditions

CAUTION


Caution must be exercised in the presence of the following:

- When there is a tendency to hemorrhage following acute trauma or fracture.
- Following recent surgical procedures where muscle contraction may disrupt the healing process.
- Over areas of the skin which lack normal sensation.

Improper maintenance or repair

WARNING

Risk of injury from improper maintenance or repair!

- Never perform unauthorized repairs of the device.
- In case of defects on the device, contact Customer Service ( “Customer service” on page 16).
- Only perform the maintenance tasks described in this manual.

If maintenance or repair is performed improperly, there is a risk of injury during training and operation. Additionally, mistakes during maintenance or repair can cause permanent damage to the device.

Interruption of the update process

NOTICE

Interrupting the update process can cause material damage!

- Do not disconnect the power supply during an update.
- Do not switch off the device during an update.
- Wait for the update to finish and do not input anything.

The device can be permanently damaged if it is turned off during the update process or if the power supply is interrupted.



If the update is interrupted due to an interruption of power, contact Customer Service (📞 “Customer service” on page 16).

Condensation

NOTICE

Risk of damage to the device from condensation!

- Only use the device in a dry environment.
- Never store the device in damp places.
- Let the device acclimatize in the event of location changes.
- In the event of faults, immediately disconnect the device from the power supply and notify Customer Service (📞 “Customer service” on page 16).
- Never switch on the device if condensation is visible.

Condensation can form in the event of changes in humidity and temperature. Moisture in the control unit can result in damage or even total destruction.

2.6 Reporting of adverse events

MedWatch (FDA)

MedWatch is the Food and Drug Administration's (FDA) program for reporting serious reactions, product quality problems, therapeutic inequivalence/failure, and product use errors with human medical products, including drugs, biologic products, medical devices, dietary supplements, infant formula, and cosmetics.

If you think you or someone in your family has experienced a serious reaction to a medical product, you are encouraged to take the reporting form to your doctor. Your health care provider can provide clinical information based on your medical record that can help FDA evaluate your report.

However, we understand that for a variety of reasons, you may not wish to have the form filled out by your health care provider, or your health care provider may choose not to complete the form. Your health care provider is not required to report to the FDA. In these situations, you may complete the Online Reporting Form yourself.

You will receive an acknowledgement from FDA when your report is received. Reports are reviewed by FDA staff. You will be personally contacted only if we need additional information.




Submitting adverse event reports to the FDA

Use one of the methods below to submit voluntary adverse event reports to the FDA:

- MedWatch Online Voluntary Reporting Form:
<http://www.accessdata.fda.gov/scripts/medwatch/index.cfm?action=reporting.home>
- Consumer Reporting Form FDA 3500B:
<http://www.fda.gov/downloads/AboutFDA/ReportsManualsForms/Forms/UCM349464.pdf>
Follow the instructions on the form to either fax or mail it in for submission. For help filling out the form, see MedWatchLearn:
<http://www.accessdata.fda.gov/scripts/MedWatchLearn/>
- Call FDA at 1-800-FDA-1088 to report by telephone.

2.7 Symbols on the miha bodytec m.ove

2.7.1 Symbols on device

Symbol	Standard Reference	Standard Title	Symbol Title	Explanatory Text
	ISO 7010 Second Edition 2011-06-01, P007	Graphical Symbols – Safety Colours And Safety Signs – Registered Safety Signs [Including AMENDMENT 1 (2012) Through AMENDMENT 7 (2016)]	Not for persons with pacemakers or implanted defibrillators	The device must never be used by persons with cardiac pacemakers or other active medical implants. In this connection, also observe the contraindications.
	ES60601-1:2005/ (R)2012 And A1:2012, Table D.2, Symbol 10	Medical electrical equipment – Part 1: General requirements for basic safety and essential performance.	Follow instructions for use	Read and observe this manual carefully before start-up and use.
	ISO 7010 Second Edition 2011-06-01, P019	Graphical Symbols – Safety Colours And Safety Signs – Registered Safety Signs [Including AMENDMENT 1 (2012) Through AMENDMENT 7 (2016)]	No stepping on surface	Do not step on the surfaces of the device or its accessories at any time.

2.7.2 Rating plates

Rating plate miha bodytec II



Fig. 16: Rating plate miha bodytec m.ove, example

NOTICE

The rating plate is located on the underside of the control unit and contains the following information:

- Manufacturer
- Type
- Model year
- Barcode
- Serial number
- Supply connection (control unit)
- Current consumption (control unit)
- Protection class
- Protection type
- Device type
- UDI Number
- FCC IDs
- IC IDs
- Hardware version identification number (HVIN)
- Mandatory sign: Read manual
- Mandatory sign: Prohibited for persons fitted with cardiac pace-makers
- Mandatory sign: Waste of electrical and electronic equipment (WEEE)
- Mandatory sign: CE marking: Sign of European technical conformity
- Mandatory sign: IEC protection Class II
- Mandatory sign: RX only
- Mandatory sign: CAN ICES-3 (B)/NMB-3(B): Compliance with Canadian ICES-3 Rules.
- Mandatory sign: FCC marking: Compliance with part 15 of the FCC Rules

Federal law restricts this device to sale by or on the order of a practitioner licensed by the law of the State in which he/she practices to use or order the use of the device.

Rating plate equipment






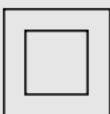



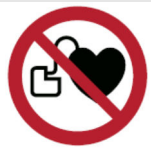





Fig. 17: Rating Plate equipment (example: i-body strap flex)

The equipment's rating plate contains the following information:



- Manufacturer
- Type
- Model year
- Serial number
- Device type
- Cleaning symbols

Symbols on rating plates






Symbol	Standard Reference	Standard Title	Symbol Title	Explanatory Text
	ISO 15223-1 Third Edition 2016-11-01 Clause 5.1.1	Medical Devices – Symbols To Be Used With Medical Device Labels, Labelling, And Information To Be Supplied – Part 1: General Requirements	Manufacturer	Indicates the device manufacturer.
	ISO 15223-1 Third Edition 2016-11-01 Clause 5.1.3	Medical Devices - Symbols To Be Used With Medical Device Labels, Labelling, And Information To Be Supplied – Part 1: General Requirements	Date of manufacture	Indicates the date when the device was manufactured.
	ES60601-1:2005/ (R)2012 And A1:2012, Table D.1, Symbol 11	Medical electrical equipment – Part 1: General requirements for basic safety and essential performance.	Consult instructions for use	Indicates the need for the user to read this manual.
	ISO 15223-1 Third Edition 2016-11-01 Clause 5.1.7	Medical Devices – Symbols To Be Used With Medical Device Labels, Labelling, And Information To Be Supplied – Part 1: General Requirements	Serial number	Indicates the manufacturer's serial number so that a specific device can be identified.
	ES60601-1:2005/ (R)2012 And A1:2012, Table D.1, Symbol 4	Medical electrical equipment – Part 1: General requirements for basic safety and essential performance.	Direct current	Indicates direct current.
	ES60601-1:2005/ (R)2012 And A1:2012, Table D.1, Symbol 9	Medical electrical equipment – Part 1: General requirements for basic safety and essential performance.	Insulation protection Class II	Identifies an IEC Class II Insulation Protection.
	ES60601-1:2005/ (R)2012 And A1:2012, Table D.1, Symbol 20	Medical electrical equipment – Part 1: General requirements for basic safety and essential performance.	Type BF applied part	Identifies a type BF applied part complying with IEC 60601-1.


Symbol	Standard Reference	Standard Title	Symbol Title	Explanatory Text
	ISO 7010 Second Edition 2011-06-01, P007	Graphical Symbols – Safety Colours And Safety Signs – Registered Safety Signs [Including AMENDMENT 1 (2012) Through AMENDMENT 7 (2016)]	Not for persons with pacemakers or implanted defibrillators	The device must never be used by persons with cardiac pacemakers or other active medical implants. In this connection, also observe the contraindications.
	765/2008/EC 768/2008/EC MDD 93/42/EEC Articles 4,11,12,17, Annex II)	The requirements for accreditation and market surveillance relating to the marketing of products; Medical Device Directive.	CE marking	Signifies European technical conformity.
	21 CFR 801.109	Labeling-Prescription devices.	Prescription only	Requires prescription in the United States.
	EN 50419	Marking of Electrical and Electronic Equipment in accordance with Article 11(2) of Directive 2002/96/EC (WEEE).	Recycle: Electronic Equipment	Waste of electrical and electronic equipment (WEEE). Electrical and electronic parts may contain toxic materials. These parts must be collected separately and turned in at communal collection points, or disposed of by a specialist company.
	ISO 15223-1 Third Edition 2016-11-01 Clause 5.1.5	Medical Devices - Symbols To Be Used With Medical Device Labels, Labelling, And Information To Be Supplied – Part 1: General Requirements	Batch code	Indicates the manufacturer's batch code so that the batch or lot can be identified.
	ES60601-1:2005/ (R)2012 And A1:2012, Table D.3, Symbol 2	Medical electrical equipment – Part 1: General requirements for basic safety and essential performance.	Degree of Ingress Protection Provided by Enclosure	Manufacturer-determined degree of particle and water ingress protection, where... N1= degree of protection from particulates (scale of 0 – 6); and N2 = degree of protection from water (scale of 0 – 8)

Safety

Symbol	Standard Reference	Standard Title	Symbol Title	Explanatory Text
	N/A	N/A	Compliance with US and Canadian electrical safety standards	Indicates the compliance with the minimum requirements of US and Canadian electrical safety standards tested by accredited certification body "TÜV Rheinland"
	47 CFR Part 15	Code of Federal Regulations, Title 47: Telecommunication, Part 15 - Radio Frequency Devices	Compliance with US electromagnetic interference standards	Indicates the compliance with part 15 of the FCC Rules for Class B digital devices.
CAN ICES-3 (B)/ NMB-3(B)	CAN ICES-3 (B)/ NMB-3(B)	Interference-causing Equipment Standard: "Digital Apparatus," ICES-003 of the Canadian Department of Communications	Compliance with Canadian electromagnetic interference standards	Indicates the compliance with the exemption from the routine evaluation limits in section 2.5 of RSS 102 and with RSS-102 RF exposure for Class B digital devices.

2.7.3 Cleaning symbols

Symbol	Standard Reference	Standard Title	Symbol Title	Explanatory Text
	7000 Fifth Edition 2014-01-15, 3125	Graphical Symbols For Use On Equipment - Registered Symbols	Wash by hand	Hand wash only.
	7000 Fifth Edition 2014-01-15, 3114	Graphical Symbols For Use On Equipment - Registered Symbols	Do not dry clean	Do not dry clean.
	7000 Fifth Edition 2014-01-15, 3113	Graphical Symbols For Use On Equipment - Registered Symbols	Do not iron	Do not iron.
	7000 Fifth Edition 2014-01-15, 3109	Graphical Symbols For Use On Equipment - Registered Symbols	Do not tumble dry	Do not tumble dry.
	7000 Fifth Edition 2014-01-15, 3124	Graphical Symbols For Use On Equipment - Registered Symbols	Do not bleach	Do not bleach.

Symbol	Standard Reference	Standard Title	Symbol Title	Explanatory Text
	7000 Fifth Edition 2014-01-15, 3089	Graphical Symbols For Use On Equip- ment - Registered Symbols	Washing, normal process, max- imum 40 °C	Wash using normal cycle at 105 °F.

2.8 Environmental protection

Disposal

NOTICE

Environmental hazard from incorrect disposal!

- Arrange for disposal of electrical scrap and electronic components by locally authorized specialist companies.
- In case of doubt, obtain information about disposal in accordance with environmental regulations from local municipal authorities or specialized disposal companies.

Environmental hazards can be caused by incorrect disposal.

The device must not be disposed of via domestic waste. Recycle dismantled components:

- Have metals turned into scrap.
- Recycle plastic elements.
- Sort and dispose of all other components according to material condition.


Electronic components

Electronic components such as liquid crystal display, printed circuit boards and cabling can contain toxic substances. These must not get into the environment. Disposal must be carried out by a specialized disposal company.



3 Technical data

Table 4: Device information

Name	miha bodytec m.ove
Manufacturer	miha bodytec GmbH Siemensstr. 1 86368 Gersthofen Germany
Initial importer	miha bodytec Inc. 2171 Executive Drive Suite 200 Addison, Illinois 60101 USA
Intended use	<p>miha bodytec m.ove is a device for electronic muscle stimulation based on EMS technology. Regarding its use, the device is specifically designed as an addition to other sports and for training muscles.</p> <p>miha bodytec m.ove is intended to stimulate muscles in order to improve or facilitate muscle performance. In addition, it is indicated for the following conditions:</p> <ul style="list-style-type: none"> ■ Re-educating muscles ■ Relaxation of muscle spasm ■ Retarding or preventing disuse muscle atrophy <p>The miha bodytec m.ove's electrical impulses allow the triggering of action potentials on motoneurons of motor nerves (excitations). These excitations of motoneurons are transmitted to the muscle fibers via the motor endplate where they generate mechanical muscle fiber responses that correspond to muscle work. Depending on the parameters of the electrical impulses (pulse frequency, duration of contraction, duration of rest, total session duration), different types of muscle work can be imposed on the stimulated muscles.</p>
Dimensions (control unit)	16.73 × 11.41 × 2.76 (W × D × H in inches) 42.5 × 29 × 7 (W × D × H in cm)
Dimensions (work station m.ove)	22 × 37.4 × 44.88 (W × D × H in inches) 56 × 95 × 114 (W × D × H in cm)
Dimensions (travel station m.ove)	19.29 × 11.81 × 37.8 (W × D × H in inches) 49 × 30 × 96 (W × D × H in cm)
Weight (control unit)	7.5 lb 3.4 kg
Weight (i-body® with cable set)	3.3 lb 1.5 kg

Weight (i-body® belt)	0.9 lb 0.4 kg
Weight (i-body® strap flex)	0.44 lb 0.2 kg
Weight (work station m.ove with maximum load)	max 110 lb max. 50 kg
Weight (travel station m.ove with maximum load)	max 55 lb max. 25 kg
Electrode size	9.75 – 64.36 in² 62.93 – 415.25 cm²
Connection to power supply (power supply unit)	100 – 240 V ~ 50 – 60 Hz
Connection to the network (control unit)	17 V – 19 V 
Current consumption (control unit during operation with the rechargeable battery)	0.6 – 1 A
Current consumption (control unit while battery is charging)	3.2 A ±5%
Essential performance characteristics:	(at 500 ohm)
Output power	< 300 mJ
Effective current	< 50 mA
Output voltage	< 500 mJ
Permissible ambient temperature (operation)	+40 °F – +105 °F +5 °C – +40 °C
Permitted battery charging temperature	+32 °F – +113 °F 0 °C – +45 °C
Time for warming up the device from the lowest storage temperature (-4 °F / -20 °C) up to operating temperature for intended use of +68 °F (+20 °C) ambient temperature	10 hours
Time for cooling down the device from the maximum permissible storage temperature (+122 °F / +50 °C) down to operating temperature for intended use of +68 °F (+20 °C) ambient temperature	10 hours
Permissible humidity (operation)	15% – 90%, non-condensing Note: Do not expose the device to the maximum humidity of 90% for more than 96 hours.
Maximum operating altitude above sea level	10,000 ft (20.67 Hg) 3000 m (700 hPa)

Technical data

Number of fixed programs	11
Number of program memories	Up to 200
Number of channels	8
Pulse width setting	50 µs – 400 µs in steps of 25 µs
Pulse rise setting	0.0 s – 1.0 s in steps of 0.1 s
Frequency setting	2.0 Hz – 10.0 Hz in steps of 0.5 Hz; 10.0 Hz – 150.0 Hz in steps of 1.0 Hz
Pulse pause setting	0.0 s – 1.0 s in steps of 1.0; 1.0 s – 10.0 s in steps of 0.1
Pulse time setting	1.0 s – 10.0 s in steps of 0.1
Time specification setting	0.5 min – 5.0 min in steps of 0.5; 5.0 min – 30.0 min in steps of 1.0; 30.0 min – 90.0 min in steps of 2.0
Protection type	<p>IP22</p> <ul style="list-style-type: none"> ■ The device is protected against solid foreign objects with a diameter of ≥ 0.5 in (12.5 mm). ■ The device is protected against dripping water.
Device type	<p>Type BF</p> 
Protection class	<p>Protective insulation II</p>  <p>Protection class II equipment has an increased or dual insulation at the level of the rated insulation voltage between active and touchable parts. In most cases, they are not connected to the protective conductor.</p>
Transponder card	RFID (13.56 MHz), according to ISO/IEC 14443 Mifare Classic 4k; can be formatted on the device
Synchronized start	2.4 GHz M2M wireless mesh network
Number of persons training with synchronized start, maximum	2
Cloud service	<p>5 GHz Wi-Fi</p> <p>(can only be used with an access point that is available upon request)</p>
Bluetooth®	2.4 GHz
Power supply	GlobTek Medical/ITE Power Supply; model no.: GTM96600-6518-T2
Expected service life of the device	The device is designed for a 10-year service life. With continuous product monitoring and regular maintenance, the service life can be significantly longer.

Expected service life of accessories (contact points, Velcro fasteners, etc.)	The service life of the accessories depends on how they are used during training. As soon as the function is no longer ensured or the hygienic requirements are no longer met, the accessories must be replaced.
Expected service life of the rechargeable battery	Approx. 600 discharge cycles

Compliance

The miha bodytec m.ove complies with the applicable requirements of the following international and national standards:

- IEC 60601-2-10:2012 – Medical electrical equipment – Part 2 – 10: Particular requirements for the basic safety and essential performance of nerve and muscle stimulators
- IEC 60601-1-2:2014 – Medical electrical equipment - Part 1 – 2: General requirements for basic safety and essential performance – Collateral standard: Electromagnetic disturbances – Requirements and tests
- AAMI ANSI ES 60601-1_2005/(R)2012 and A1:2012 – Medical electrical equipment – Part 1: General requirements for basic safety and essential performance
- IEC 60601-1-11:2010 – Medical electrical equipment – Part 1 – 11: General requirements for basic safety and essential performance – Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment
- IEC 62304:2015 – Medical device software – Software life-cycle processes
- ISO 14971:2007 – Medical devices – Application of risk management to medical devices
- IEC 62366-1:2015 – Medical devices – Part 1: Application of usability engineering to medical devices
- ISO 10993-1:2009 – Biological evaluation of medical devices – Part 1: Evaluation and testing within a risk management process
- ISO 10993-5:2009 – Biological evaluation of medical devices – Part 5: Tests for in vitro cytotoxicity
- ISO 10993-10:2010 – Biological evaluation of medical devices – Part 10: Tests for irritation and skin sensitization

FCC Statement

This device complies with part 15 of the FCC Rules. Operation is subject to the following two conditions:

- (1) This device may not cause harmful interference, and
- (2) This device must accept any interference received, including interference that may cause undesired operation.

Any changes or modifications not expressly approved by the party responsible for compliance could void the user's authority to operate the equipment.

Note: This equipment has been tested and found to comply with the limits for a Class B digital device, pursuant to part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference in a residential installation.

This equipment generates, uses and can radiate radio frequency energy and, if not installed and used in accordance with the instructions, may cause harmful interference to radio communications. However, there is no guarantee that no interference will occur in a particular installation.

If this equipment causes harmful interference to radio or television reception, which can be determined by turning the equipment off and on, the user is encouraged to try to correct the interference by one or more of the following measures:

- Reorient or relocate the receiving antenna.
- Increase the distance between the equipment and receiver.
- Connect the equipment to an outlet on a circuit different from that to which the receiver is connected.
- Consult the dealer or an experienced radio/TV technician for help.

This equipment complies with FCC radiation exposure limits set forth for an uncontrolled environment.

IC Caution

The device meets the exemption from the routine evaluation limits in section 2.5 of RSS 102 and compliance with RSS-102 RF exposure, users can obtain Canadian information on RF exposure and compliance. This device contains licence-exempt transmitter(s)/ receiver(s) that comply with Innovation, Science and Economic Development Canada's licence-exempt RSS(s).

Operation is subject to the following two conditions:

- (1) This device may not cause interference.
- (2) This device must accept any interference, including interference that may cause undesired operation of the device.

Le dispositif rencontre l'exemption des limites courantes d'évaluation dans la section 2.5 de RSS 102 et la conformité à l'exposition de RSS-102 rf, utilisateurs peut obtenir l'information canadienne sur l'exposition et la conformité de rf. L'émetteur/récepteur exempt de licence contenu dans le présent appareil est conforme aux CNR d'Innovation, Sciences et Développement économique Canada applicables aux appareils radio exempts de licence.

L'exploitation est autorisée aux deux conditions suivantes:

- (1) L'appareil ne doit pas produire de brouillage;
- (2) L'appareil doit accepter tout brouillage radioélectrique subi, même si le brouillage est susceptible d'en compromettre le fonctionnement.

Environment

The miha bodytec m.ove is designed for use in domestic environments and professional health facilities, but not near active systems of high-frequency surgical devices or in high-frequency shielding rooms used for magnetic resonance imaging where high-intensity electromagnetic interferences occur.

Open source software

This product also contains open source software, among other things, which was developed by third parties. Specifically, they include AFL, AGPL, Artistic, Beerware, BSD, Combined_OpenSSL+SSLeay, copyleft-next, eCos, FSLA, FTL, Google-BSD, GPL, IBM-pibs, IETF, IJG, ISC, LGPL, Libpng, MIT, MPL, OpenSSL, RHeCos, Sun, WebM, X11, Zlib.

The open source software is protected by copyright. You are entitled to use the open source software in accordance with the relevant open source software license terms, which we will gladly send you upon request.

The open source software is provided free of charge. You can request the source code for up to three years from the purchase date of the device, if provided in the relevant open source software license terms. Any liability for the use of the open source software beyond the program cycle provided by us for the product as well as any liability for defects caused by changes to the open source software is excluded. We do not provide technical support for this product if it was changed.

4 Electromagnetic compatibility

4.1 Compliance details

As noted in section 3, the miha bodytec m.ove EMS training device complies with IEC 60601-1-2:2014 and IEC 60601-2-10:2012. According to these standards, the requirements apply as listed in the tables below.

Guidance and manufacturer's declaration – electromagnetic emissions

The miha bodytec m.ove EMS training device has been designed for use in electromagnetic environments as outlined below.

The buyer or user of the miha bodytec m.ove device must ensure that the device is operated under such conditions.

Emission tests	Compatibility	Electromagnetic environment – guidance
RF emissions CISPR 11	Group 1	The miha bodytec m.ove EMS training device uses low RF power for internal operation. Generating very low RF interference emissions, it is not likely to interfere with electronic equipment in its vicinity.
RF emissions CISPR 11	Class B	The miha bodytec m.ove device is suitable for usage in any building, including residential buildings and those connected up direct to low-voltage power supply networks installed for residential buildings.
Harmonic distortion. IEC 61000-3-2	Class A	
Voltage fluctuations and flicker. IEC 61000-3-3	Complies	

Guidance and manufacturer's declaration - electromagnetic immunity

The miha bodytec m.ove EMS training device has been designed for use in electromagnetic environments as outlined below.

The buyer or user of the miha bodytec m.ove device shall make sure that the device is used under such conditions.

Immunity test	IEC-60601 test level	Compliance level	Electromagnetic environment – instructions
Electrostatic discharge (ESD) IEC 61000-4-2	±8 kV on contact discharge ±2 kV, ±4 kV, ±8 kV, ±15 kV air discharge	±8 kV contact discharge ±2 kV, ±4 kV, ±8 kV, ±15 kV air discharge	The miha bodytec m.ove EMS training device can be used in rooms without humidity control. Floors can be covered with synthetic material. If so, the relative humidity should be at least 5 %.

Guidance and manufacturer's declaration - electromagnetic immunity

The miha bodytec m.ove EMS training device has been designed for use in electromagnetic environments as outlined below.

The buyer or user of the miha bodytec m.ove device shall make sure that the device is used under such conditions.

Immunity test	IEC-60601 test level	Compliance level	Electromagnetic environment - guidance
Radiated RF IEC 61000-4-3	10 V/m 80 MHz to 2.7 GHz modulation 80% AM at 1 kHz	10 V/m	The minimum separation distance d in m to a RF emitting device with a maximum power P in W not operating in a frequency band covered by the proximity fields test can be calculated using the equation in clause 8.10 of IEC 60601-1-2 with E as the field strength in V/m taken from the compliance level: $E = 6 / d \cdot \sqrt{P}$
Proximity fields from RF wireless communications equipment IEC 61000-4-3	385 MHz: 27 V/m 450 MHz: 28 V/m 710 MHz: 9 V/m 745 MHz: 9 V/m 780 MHz: 9 V/m 810 MHz: 28 V/m 870 MHz: 28 V/m 930 MHz: 28 V/m 1720 MHz: 28 V/m 1845 MHz: 28 V/m 1970 MHz: 28 V/m 2450 MHz: 28 V/m 5240 MHz: 9 V/m 5500 MHz: 9 V/m 5785 MHz: 9 V/m	385 MHz: 27 V/m 450 MHz: 28 V/m 710 MHz: 9 V/m 745 MHz: 9 V/m 780 MHz: 9 V/m 810 MHz: 28 V/m 870 MHz: 28 V/m 930 MHz: 28 V/m 1720 MHz: 28 V/m 1845 MHz: 28 V/m 1970 MHz: 28 V/m 2450 MHz: 28 V/m 5240 MHz: 9 V/m 5500 MHz: 9 V/m 5785 MHz: 9 V/m	Portable RF communications equipment (including peripherals such as antenna cables and external antennas) shall be used no closer than 30 cm (12 inches) to any part of the miha-bodytec m.ove EMS training device, including cables specified by the manufacturer. The field strength test levels are calculated using the equation in clause 8.10 of IEC 60601-1-2: $E = 6 / d \cdot \sqrt{P}$ with E as the field strength in V/m, the maximum power P in W of and the minimum separation distance d in m to the RF.

Electromagnetic compatibility

Guidance and manufacturer's declaration - electromagnetic immunity

Mains-frequency (50/60 Hz) magnetic field IEC 61000-4-8	30 A/m, 50 Hz or 60 Hz	30 A/m 50 Hz or 60 Hz	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.
Electrical fast transients/bursts IEC 61000-4-4	Input a.c. power port: ± 2 kV Input d.c. power port: ± 2 kV Signal input/output parts port: ± 1 kV (with 100 kHz repetition frequency)	Input a.c. power port: ± 2 kV Input d.c. power port: ± 2 kV Signal input/output parts port: ± 1 kV (with 100 kHz repetition frequency)	Mains power quality should equal that of usual commercial or hospital environments. Note: The miha bodytec m.ove EMS training device has no input d.c. port and no signal input/output parts ports with a maximum cable length of 3 meters or more.

Guidance and manufacturer's declaration - electromagnetic immunity

The miha bodytec m.ove EMS training device has been designed for use in electromagnetic environments as outlined below.

The buyer or user of the miha bodytec m.ove device shall make sure that the device is used under such conditions.

Immunity test	IEC-60601 test level	Compliance level	Electromagnetic environment - guidance
Surges IEC 61000-4-5	Input a.c. power port: ± 0.5 kV, ± 1 kV, ± 2 kV line-to-ground ± 0.5 kV, ± 1 kV, line to line Input d.c. power port: ± 0.5 kV, ± 1 kV, ± 2 kV line-to-ground ± 0.5 kV, ± 1 kV, line to line Signal output parts port: ± 2 kV	Input a.c. power port: ± 0.5 kV, ± 1 kV, ± 2 kV line-to-ground ± 0.5 kV, ± 1 kV, line to line Input d.c. power port: ± 0.5 kV, ± 1 kV, ± 2 kV line-to-ground ± 0.5 kV, ± 1 kV, line to line Signal output parts port: ± 2 kV	Mains power quality should equal that of usual commercial or hospital environments

Guidance and manufacturer's declaration - electromagnetic immunity

Conducted disturbances induced by RF fields IEC 61000-4-6	3 V 0.15 MHz to 80 MHz 6 V in ISM and amateur radio bands between 0.15 MHz and 80 MHz Modulation 80 % AM at 1 kHz	3 V 0.15 MHz to 80 MHz 6 V in ISM and amateur radio bands between 0.15 MHz and 80 MHz Modulation 80 % AM at 1 kHz	As described in annex A.3 of IEC 60601-1-2, the voltage induced on a cable in the frequency range 150 kHz to 80 MHz from a field strength of 10 V/m is unlikely to exceed 6 V r.m.s. Therefore, assuming a field strength E of 10 V/m in the ISM and amateur radio bands between 0.15 MHz and 80 MHz (see note 2), the minimum separation distance d in m to a RF communications equipment with a maximum power P in W can be calculated using the equation in clause 8.10 of IEC 60601-1-2: $d = 6 / E \cdot \sqrt{P}$
Voltage dips IEC 61000-4-11	0% U _T ; 0.5 cycle at 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0% U _T ; 1 cycle; 70% U _T ; 25/30 cycles; Single phase: at 0°	0% U _T ; 0.5 cycle at 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0% U _T ; 1 cycle; 70% U _T ; 25/30 cycles; individual phase: at 0°	Mains power quality should be that of a typical commercial or hospital environment. If the use of the miha-bodytec m.ove EMS training device requires continued operation during power mains interruptions, an uninterruptible power supply (UPS) sufficiently fast and powerful to provide the energy needed shall be used.
Voltage interruptions IEC 61000-4-11	0% U _T ; 250/300 cycles	0% U _T ; 250/300 cycles	

Electromagnetic compatibility


Guidance and manufacturer's declaration - electromagnetic immunity

Electrical transient conduction along supply lines (input d.c. power port)	Not applicable	Not applicable	The miha bodytec m.ove EMS training device is not intended to be installed in passenger cars and light commercial vehicles including ambulances fitted with 12 V electrical systems or commercial vehicles including ambulances fitted with 24 V electrical systems.
--	----------------	----------------	--

Note 1: U_T is the a.c. mains voltage prior to application of the test level.

Note 2: The ISM (industrial, scientific and medical) bands between 150 kHz and 80 MHz are 6.765 MHz to 6.795 MHz, 13.553 MHz to 13.567 MHz, 26.957 MHz to 27.283 MHz and 40.66 MHz to 40.70 MHz. The amateur radio bands between 150 kHz and 80 MHz are 1.8 MHz to 2.0 MHz, 3.5 MHz to 4.0 MHz, 5.3 MHz to 5.4 MHz, 7 MHz to 7.3 MHz, 10.1 MHz to 10.15 MHz, 14 MHz to 14.2 MHz, 18.07 MHz to 18.17 MHz, 21.0 MHz to 21.4 MHz, 24.89 MHz to 24.99 MHz, 28.0 MHz to 29.7 MHz and 50.0 MHz to 54.0 MHz.

Replaceable parts:

All cables and equipment shall be obtained from miha bodytec or authorized dealers. For replaceable parts available see  *Chapter 1.2 "Equipment" on page 10*. Using other cables or equipment can void the authority to operate the equipment.

Essential performance:

In the presence of an electromagnetic disturbance, the electronic muscle stimulation (EMS) operates as intended.

In case of voltage interruptions, continuous operation is not guaranteed. If continuous operation is required, an uninterruptible power supply (UPS) sufficiently fast and powerful to provide the energy needed shall be used.

4.2 EMC Warnings

WARNING

Risk of injury if other equipment is used!

Use of this equipment adjacent to or stacked with other equipment must be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment shall be observed to verify that they are operating normally.

NOTICE

Risk of damage due to the use of third-party equipment!

Use of equipment, transducers and cables other than those specified or provided by the manufacturer of this equipment could result in increased electromagnetic emissions or decreased electromagnetic immunity of this equipment and result in improper operation.

NOTICE

Risk of damage due to portable RF communications equipment!

Portable RF communications equipment (including peripherals such as antenna cables and external antennas) must be used no closer than 30 cm (12 inches) to any part of the miha bodytec m.ove EMS training device, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment can result.

5 Basic information on EMS training

5.1 Safety first

Increasing loads slowly



Fig. 18: Control unit 0 position

Safety and controlled training take priority over experimentation and hurried and uncontrolled increase of the stress parameters. For every training session, start at “0” with the stress for the main level and then increase slowly using the main controller.

5.2 EMS training

What is EMS training?

EMS stands for **E**lectro**M**yo**S**timulation. This means the muscles are stimulated by electrical pulses. Complete body training which addresses all muscle groups is possible with up to 8 pairs of electrodes.

During conventional training, the muscles are controlled via electrical signals from the brain to initiate a contraction and thus movement of the muscles. With EMS, electrical pulses from outside activate the muscles. It makes no difference to the muscle whether electrical stimulations are sent from the brain or from electrodes: It reacts by contracting!

Special features and benefits:

- All muscle groups can be activated using an electrode system with up to 10 pairs of electrodes.
- Static and dynamic training with interaction of deliberate muscle contraction and EMS.
- Exercise postures increase the contraction of the deliberately stimulated muscles.
- Positive and negative electrodes are not on the same muscle.
- Agonist and antagonist are stimulated simultaneously.
- EMS training can cause more intense muscle contractions than classic strength training. At the same time, there is comparatively very little stress on the joints.

- Deeper muscle groups can be easily reached.
- The training is very intense and thus accordingly short (10 – 20 minutes).

To be observed during EMS training

On the one hand, EMS training can be used on a large part of the body. On the other hand, EMS training permits athletes/patients to attain a training intensity level that is substantially higher than that of conventional workout methods. This is why a few special considerations must be observed during the training process. Regardless of the athletic background and physical condition, the instructions on the training frequency and regeneration – in particular for EMS beginners – must be observed *⚡ Chapter 5.3 “Training frequency and regeneration” on page 49.*

5.3 Training frequency and regeneration

Overtraining

⚠ WARNING

Risk from overtraining!

- Always give your body sufficient regeneration time.
- Do not train if suffering from continued severe muscle ache.
- In the event of overload, make cyclical alternating intensive and regenerative workouts on different days.

Overtraining can result in physical weakness, sleep disorders, headaches, increased resting and active pulse rate, muscle and tendon pains, and also depression among other things.

The recommended training frequency is 1 – 2 times per week. Do not train in the event of severe muscle ache, feeling unwell, or other contraindications (*⚡ “Never exercise if ...” on page 19).*

Training if fatigued

⚠ WARNING

Danger to health from EMS training if performed while ill or otherwise fatigued!

- If ill, only train after consulting with your doctor.
- Always eat and drink sufficiently before training *⚡ “Always eat and drink sufficiently” on page 50.* Never train on an empty stomach.
- Follow the instructions on training frequency and regeneration *⚡ Chapter 5.3 “Training frequency and regeneration” on page 49.*

EMS training is an intense workout that challenges the body. If the athlete/patient is ill or otherwise fatigued due to health reasons, their body is unable to cope with the stress and their illness may become worse as a result

Illness and training

Just as with intense conventional workouts, EMS training may only be performed if the body is in adequate physical condition.

Basic information on EMS training

Persons suffering from fever in particular are not permitted to train with the device.

The first workout

Do not let the first workout exceed 15 minutes in length, and use the following programs:

- Program 1: Pulse familiarization for 3 approx. minutes
- Program 6: Test workout for 12 minutes

Training in the first 10 weeks

In the first 10 weeks, do not exceed a training frequency of one training session per week.

Do not exceed a training time of 20 minutes.

Training after the first 10 weeks

After an acclimatization period of 10 weeks, workouts must still be separated by a regeneration phase of at least 4 days. Depending on the athlete's/patient's physical condition and fitness level, the regeneration phase may have to be longer.

Do not exceed a training time of 20 minutes.



If clarification is necessary, sports physicians can provide information about the optimum regeneration phase for optimum training results.

Always eat and drink sufficiently

EMS training is an intense workout and stimulates a large number of muscles simultaneously. This is why the athlete's/patient's body has elevated energy requirements during training. Before an intense workout, ensure that you replenish your body's reserve of carbohydrates. This can be done by eating a meal high in carbohydrates (but not heavy) 2 – 3 hours before a workout.

Your body also requires adequate fluids. Drink 500 ml before and after a workout to replenish your hydration levels.

Effect of training on blood count

Training close to your physical limit can initially lead to extremely elevated levels of creatine kinase and myoglobin, especially during the first training sessions. Even in healthy people, this is connected with stress on the kidneys.

EMS beginners in particular must therefore proceed carefully. On the one hand, this particular stress factor must be increased slowly. On the other hand, athletes/patients must ensure sufficient regeneration between individual workouts.

After an acclimatization phase of 8 – 10 weeks with one workout per week, the body has adjusted to the situation. This means that the effects of EMS training on the body's creatine kinase and myoglobin levels are comparable to the effect of a very intense conventional muscular workout.

Regeneration



Muscle growth takes place in the resting phase between workouts. Therefore, always allow your body the necessary rest.

You should drink a high-protein energy drink such as a protein shake for better regeneration and for stimulation of muscle building. Take a break after the workout and do not complete any further workouts to guarantee best possible regeneration and to avoid overtraining.

6 Understanding the device

6.1 Explanation of terms

Programs

Programs are defined as settings that are based on various parameters such as the pulse time, pulse width, pause time, etc., which generate a pulse within the desired intensity range that is specific to the selected workout. While the program is running, the desired pulse will be output for the previously selected duration. This makes it possible to engage in “freestyle” training.

Training plan

A training plan is the combination of a program and a sequence of different exercises which are demonstrated on the display by an avatar (male or female). You can select from different training plans, each composed of different exercises and sequences.

Loading the level



“Load level” is only available in combination with the transponder card. The function must be enabled in the device settings and under card management. The function is also only available for training plans and training programs that are defined on the customer card.

“Load level” is a function that automatically sets the intensity for the level controller.

When you complete a workout, the intensity levels currently set for the level controller are stored on the transponder card. These intensity levels can be automatically accessed on the transponder card during future workouts.

For this to work, the “Load level” function must be active in the device settings. In addition, this option must also be set on the transponder card of the athlete/patient.

When a workout is started with a transponder card in place, level controllers 1 – 10 are automatically set to the saved levels.

To increase safety for the athlete/patient, there are threshold values for the main controller if the “Load level” function is active. If the intensity level is increased using the main controller, the device automatically stops this increase at the value levels of 30, 40, and 50. To increase the intensity level above these thresholds, the main controller must be pressed once (each time).

Favorites menu

The favorites menu is the first menu tab. It enables the trainer to create a customized choice of workout options.

The favorites menu can be used for storing frequently used programs and/or training plans. This lets trainers restrict the range of choices available to the athlete/patient and therefore select the programs and training plans to be used. Creating a summary in the favorites menu also makes it possible to select choices faster, which facilitates the use of transponder cards.



If favorites have been selected, only these favorite workouts will be displayed in the card management menu.

Data backup/restoration

You use the data backup function to create a backup copy of the device settings and programs as well as of the training plans on an external device (USB flash drive). You can use the “Restore” function to transfer device settings and/or programs and training plans (depending on your selection) to other devices.



Use only the genuine USB flash drives supplied by the manufacturer on the USB port of the device.

Master device/managed device

With synchronized start, one device is assigned a managing function. This device controls the training procedure for all other devices involved and is referred to as the master device. Devices participating in the synchronized start that are not master devices are referred to as managed devices.

The athlete/patient must strictly adhere to the instructions of the trainer. If physical or device-specific problems arise during training, notify the trainer immediately and follow the trainer's instructions.



To ensure the trainer and persons training can communicate as required, the managed devices must be oriented towards the master device to allow the trainer to monitor each person training. To ensure that training with synchronized start is effective and safe, a maximum of 2 persons training per trainer is recommended.

Brand code



Risk of injury due to incorrect input of brand codes!

- **Never enter brand codes yourself.**
- **If you want to use brand codes, contact Customer Service** (📞 “Customer service” on page 16).

If brand codes are entered without adequate background knowledge, athletes can be injured due to malfunctions or incorrect parameterization and the device can be damaged.

The brand code (parameterization) is used to select or import default factory settings, e.g., color settings, graphic displays, and network functions. This selection option may only be used by service employees of miha bodytec or under their supervision.

i-body® connect wireless

The “i-body connect wireless” menu item can be used to switch the Bluetooth® connection to the i-body® connect wireless device on and off.

In the event of faults in a connected i-body® connect wireless, device information such as serial number, firmware, and error codes can be output. This is important information for the service employee at miha bodytec. Moreover, the “Perform self-test” and “Export log” functions can be performed under the guidance of our service employee.

Understanding the device

System status

The system status can be used for identifying different device parameters. It is an important source of information for our service employees in the event of a defect. Provide service employees with system status information if asked.

Export log





“Export log” is a function that allows relevant device information to be exported to a USB flash drive in the event of an error.



The “Export log” function must be used only under the guidance of miha bodytec service personnel.

Symbolic level controller assignment

The individual connections of the electrodes are assigned symbolically to the level controllers. The strength for the area is adjusted using the respective level controller.

Symbol	Channels/application area
	Channel 1 – legs
	Channel 2 – buttocks
	Channel 3 – lower back
	Channel 4 – upper back
	Channel 5 – side back
	Channel 6 – abdomen
	Channel 7 – chest
	Channel 8 – arms

Symbol	Channels/application area
9 	Channel 9
10 	Channel 10

6.2 Opening menu tabs

Main menu

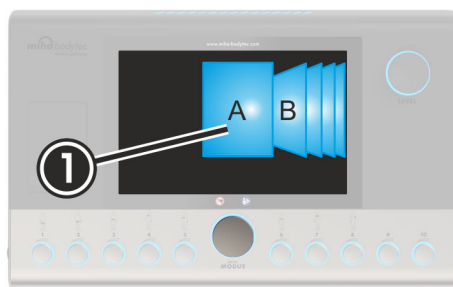


Fig. 19: Main menu

The menu level where the menu tabs are displayed (Fig. 19/1) is the main menu. When the main menu is displayed, the device is in a safe condition. In this context, “safe condition” means that the electrode connections are voltage-free. Electrodes may be attached and detached from the device only when the device is in this safe condition.



Menu tabs (Fig. 19/1) offer convenient access to different configuration options and special types of training sessions. You can start navigating the available options with the help of menu tabs. After selecting a menu tab, you will be shown the associated menus or secondary menu tabs.

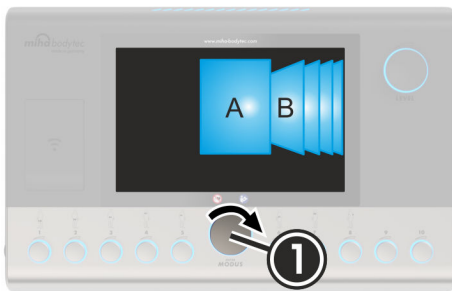


Fig. 20: Selecting menu tabs

1. ➡ Turn the multi-function button (Fig. 20/1) clockwise.



Fig. 21: Selected menu tab

⇒ The following menu tab (Fig. 21/B) is selected.

2. ➡ Press the multi-function button (Fig. 20/1).

⇒ The selected menu tab is opened.

***i** When you turn the multi-function button counterclockwise, you can select the menu tab displayed to the left next to the selected menu tab. In the example shown, tab "A" (Fig. 21/A) is selected.*

6.3 Opening menus

Structure of menus

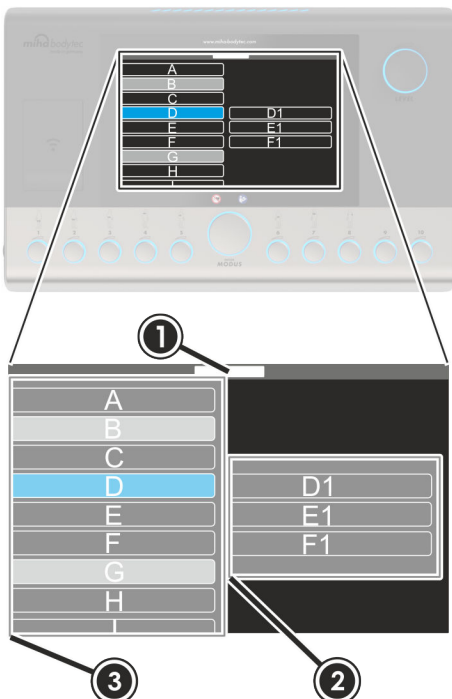


Fig. 22: Structure of menus

The device menus always have the same structure. The status line displays the name of the menu (Fig. 22/1).

The items of the respective menu (Fig. 22/3) are listed one below the other on the left side.



If there are more items in a menu than can be shown simultaneously on the display, only some of the menu items are shown. In the example shown, this is the lowest menu item (Fig. 22/1).

The current settings (Fig. 22/2) of the respective menu items are shown on the right half of the display. The current settings are always displayed next to the associated menu items.

Menu colors

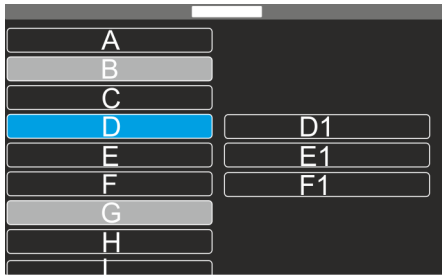


Fig. 23: Example menu

Menus are labelled with different colors. The color of a menu item indicates whether the menus are locked or can be selected using the multi-function button. The device also uses the color of the menu items to provide feedback on which menu is currently selected:

Color	Example	Meaning
Black background	A	Menu can be selected.
Gray background	B	Menu is locked. Locked menus cannot be opened.
Colored background*	D	Menu is currently selected.
Gray background and colored frame*	G	Locked menu is currently selected**.

* The indicated color corresponds to the color selected in the "Device color" menu item.

** By turning the multi-function button, only the position is shown.

Selecting and opening menus

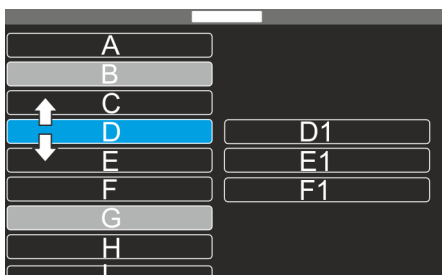


Fig. 24: Selecting a menu

1. Turn the multi-function button until the desired menu item is displayed against a colored background.
⇒ Menu item with colored background has been selected.
2. Press the multi-function button.
⇒ The menu opens and the submenu associated with the menu item is selected.

Settings in submenus

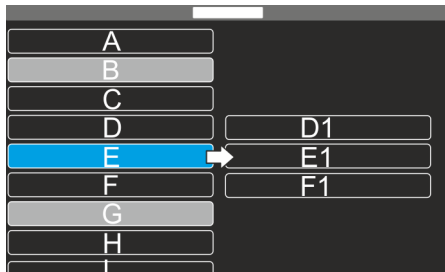


Fig. 25: Setting has been selected

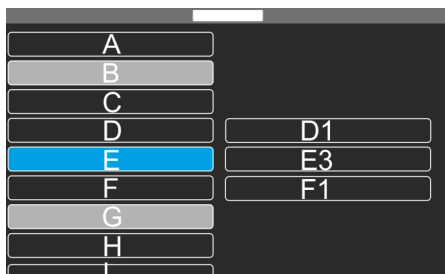


Fig. 26: Setting has been changed

1. ➤ Open the desired menu item containing a submenu
↳ “Selecting and opening menus” on page 57.
2. ➤ Press the multi-function button.
⇒ Submenu has been selected (Fig. 25/white arrow).
3. ➤ Turn the multi-function button.
⇒ The settings options available for the submenu are displayed one at a time (example here “E1 → E2 → E3”).
4. ➤ Press the multi-function button.

⇒ The displayed submenu setting (here Fig. 26/E3) is applied.

Opening the main menu



How the main menu opens depends on the current device status. The following cases are possible:

- Open the main menu while a menu is displayed ↳ “From a menu” on page 58.
- Open the main menu while a training session is active ↳ “During training” on page 59.

From a menu

1. ➤ Turn the multi-function button to select the top or bottom menu item.
2. ➤ Press the multi-function button to open the top or bottom menu item.
3. ➤ Repeat steps 1 and 2 until the “Main menu” menu item is displayed.

4. ➤ Select the “Main menu” menu item.
⇒ The main menu is displayed.



Fig. 27: Main menu

During training

1. ➤ Press the multi-function button.
⇒ The training session is cancelled or ended. The training screen remains open.
2. ➤ Press the multi-function button again.
⇒ Without transponder card: The main menu is displayed.
With transponder card: Data is written on the card and the programs or training plans saved on the card are displayed. The main menu is only displayed after the card has been removed from the device.



Fig. 28: Main menu

6.4 Entering text

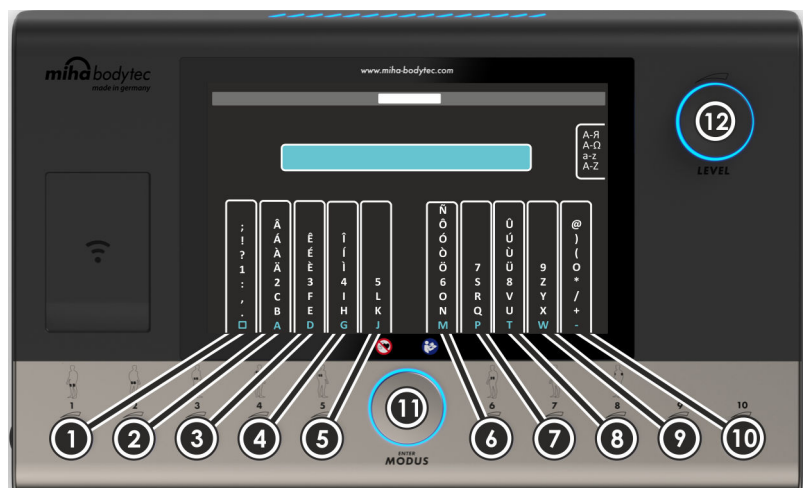


Fig. 29: Entering text

When creating customized settings or personal data, you have the option to store information in text form (e.g., “Create customer card” → “Name”).

When you select these text boxes with the multi-function button (Fig. 29/11), you have the following input options:

- When you turn the main controller (Fig. 29/12), you can switch between individual typefaces and upper- and lowercase letters.
- When you press the main controller briefly (Fig. 29/12), the last letter in the text box is deleted. When you keep it depressed, the entire text box is deleted.
- When you turn the individual level controller (Fig. 29/1 – 10) (channels 1 – 10), the letters are selected. When you press the respective controllers, they are adopted into the text box.
- When you press the multi-function button (Fig. 29/11), the entered text is adopted.

7 Customization and setup

7.1 Adjusting the device setup

Displaying device settings

Personnel: ■ Trainer

1. ➤ Open the “Setup” menu tab.
 2. ➤ Open the “Device settings” menu item.
- ⇒ The “Device settings” menu opens.

Menu items device settings

You can set the following parameters in the “Device settings” menu:

- **Load level:** Enables or disables the option “Load level” in general, but requires additional activation for each customer on the customer card.
- **Device color:** Lets you adjust the color of all LED lights in the device.
- **Active color:** Sets the color of the decreasing LED strip in the display and on the rear while a pulse is being output.
- **Pause color:** Sets the color of the increasing LED strip in the display and on the rear while no pulse is being output.
- **Pause transition color:** Specifies the color of the LED strip in the display and on the rear if a pause transition of more than 0 LEDs was selected.
- **Pause transition:** Specifies how many LEDs are supposed to change color at the beginning of the pulse.
- **Brightness:** Specifies the brightness of the display in the device.
- **Language:** Specifies the language to be used for display texts.
- **Gender of avatar:** Specifies the gender of the avatar for workouts without a transponder card.

Changing controller colors



Any change of the controller color will not take effect until you exit the “Setup” menu. Pause transitions can be used for instance to demonstrate to the athlete/patient with the device when they have to contract their muscles.

7.2 Selecting the language



The device lets you choose from the following menu language options:

- German
- English
- French
- Spanish
- Italian
- Russian
- Greek
- Polish
- Portuguese
- Dutch
- Hungarian
- Romanian
- Turkish
- Chinese
- Czech

Selecting the default language for display texts



The selected default language is the default option applying to all athletes/patients.

Personnel: ☒ Trainer

1. ➤ Open the “Setup” menu tab.
2. ➤ Open the “Device settings” menu item.
 - ⇒ The “Device settings” menu opens.
3. ➤ Open the “Language” menu item.
4. ➤ Select the desired language and confirm your selection by pressing the multi-function button.
 - ⇒ Display texts are shown in the language you have selected.



Every athlete/patient can customize the language settings and store these personalized settings on the transponder card
🔗 “Selecting individual language settings for the athletes/patients” on page 101.

7.3 Program memory settings

Displaying the program memory

Personnel: ■ Trainer

1. ➞ Open the “Setup” menu tab.
2. ➞ Open the “Program memory settings” main menu item
 ↳ *Further information on page 59.*
 ⇒ The “Program memory settings” opens.

Menu items

You can set the following parameters in the “Program memory settings” menu:

- **Name:** Defines a custom name for the training program. Instructions on how to write names ↳ *Further information on page 59.*
- **Training time:** Specifies the total training time.
- **Pulse time:** Specifies the duration for which pulses will be emitted. The pulse time is indicated in seconds.
- **Pause time:** Specifies the time in between pulses. Pause time is indicated in seconds.
- **Frequency:** The pulse frequency in Hz indicates how many individual pulses will be emitted per contraction phase.
- **Pulse width:** Specifies the length of an individual pulse. The longer the pulse width, the deeper the pulse will penetrate the muscle tissue. The pulse width is indicated in microseconds.
- **Rise time:** Specifies the time that will elapse between the beginning of the pulse to the pulse reaching its peak.
- **Save program:** Saves the program on the device. Once saved, a program can no longer be modified or deleted. It can only be deactivated. It will then only be displayed in the training plan memory settings and program memory settings. The program can be activated again in these settings.

The “Program memory settings” menu displays in numerical order all training programs that have been created.



While you can view the programs’ memory settings by selecting them, you will not be able to modify or delete them.

7.4 Setting up the training plan memory

Viewing/creating training plan memory

Personnel: ■ Trainer

1. ➞ Open the “Setup” menu tab.
 ⇒ The “Setup” menu tab opens.
2. ➞ Turn the multi-function button to the right to select the “Set up training plan memory” menu item.
 ⇒ The “Set up training plan memory” menu item is selected.

3. ➤ Confirm your selection by pressing the multi-function button.
⇒ The “*Set up training plan memory*” menu opens.
4. ➤ Turn the multi-function button to the right to select the “*Create training plan memory*” menu item.
⇒ The “*Create training plan*” menu item is selected.
5. ➤ Confirm your selection by pressing the multi-function button.
⇒ The “*training plan*” menu opens.

Menu items of the training plan memory

You can set the following parameters in the “*Set up training plan memory*” menu:

- **Name:** Defines a custom name for the training program. Instructions on how to write names ↪ *Further information on page 59*.
- **Program:** Specifies the training program including pulse width, rise time, etc., which will be stored for the training plan.
- **Training time:** Specifies the duration of the entire training plan.
- **Repetitions:** This screen shows the number of repetitions/exercises after selecting the exercise.
- **Exercises:** Allows you to select the individual exercises. “Pause” is an exercise.
- **Remaining repetitions/time:** Depending on the training time, this shows the number of repetitions that can still be completed during the remaining time.
- **Save training plan:** The training plan cannot be saved as long as the value of the remaining repetitions is not “0.”



- *Only training programs with 4-second cycles are displayed.*
- *During training, there will be a remaining time that elapses with no exercising if the training time cannot be divided by 8 seconds (4-second exercise/4-second pause).*
- *If a training plan is deactivated, it is no longer visible in the “Training plans” selection or in the favorites menu.*
- *After storing the training plan, you can change the exercises and the name, but you cannot alter the corresponding program for the training plan.*
- *If you select “Pause” as the exercise, the main controller value is automatically set to 30. You can only increase this value during or after the pause if you press the main controller. The LED strip turns orange while the intensity level is being reduced.*

7.5 Setting up synchronized start

What is training with synchronized start?

During training with synchronized start, several devices are networked to form a training group. One device in the training group provides the start command and enables simultaneous training with several devices.



To ensure that the trainer and athletes can communicate as specified, the managed devices must be oriented towards the master device so that the trainer can monitor each athlete. A maximum of 2 athletes per trainer are recommended to ensure that training with synchronized start is effective and safe.

Steps to set up synchronized start

2 steps are required to set up synchronized start:

- Setting up the master device ➤ “Setting up the master device” on page 65
- Setting up managed devices ➤ “Setting up a managed device” on page 66

Overview of synchronized start settings

You can set the following parameters in the “Synchronized start” menu:

- **Synchronized start:** Enables or disables the synchronized start option in general (if synchronized start is enabled, the synchronized start symbol lights up blue).
- **Master device:** Specifies whether the device is supposed to control the training as the master device or participate in the training as a managed device.
- **Group number:** Defines a group number for the training session.
- **Countdown:** Specifies the delay with which the training session will begin following the start command.
- **Flexible training:** When the flexible training option is activated, all participating devices only have to be set to the same cycle (e.g., 4 s of pulse and 4 s of pause). In addition, the users of all devices must train using a training plan or a program.
When flexible training option is deactivated, a device can only participate in the training with synchronized start session if it is set to the same program or same training plan as the master device.

Setting up the master device

Personnel: ■ Trainer

1. ➤ Open the “Setup” menu tab.
⇒ The “Setup” menu tab opens.
2. ➤ Open the “Synchronized start” menu item.
3. ➤ Set the current “Synchronized start” setting to “YES”
➤ “Settings in submenus” on page 58.
4. ➤ Set the current “Master device” setting to “YES”.

5. ➤ Set the current “Group number” setting to the desired group number.

i The group number must be between 1 and 16. The group number you selected also needs to be set for all other managed devices participating in the training session.

⇒ The device is ready to control the training session as the master device.

Setting up a managed device



All managed devices that are supposed to participate in the training session have to be set up as described below.

Personnel: ■ Trainer

1. ➤ Open the “Setup” menu tab.
⇒ The “Setup” menu tab opens.
2. ➤ Open the “Synchronized start” menu item.
3. ➤ Set the current “Synchronized start” setting to “YES”
 ⚙ “Settings in submenus” on page 58.
4. ➤ Set the current “Master device” setting to “NO”.
5. ➤ Set the current “Group number” setting to the desired group number.

i The selected group number must match the group number of the master device.

⇒ The device is ready to participate in the training session as a managed device.



All athletes also have to specify that they want to participate in the training with synchronized start session ⚙ “Setting up the transponder card for training with synchronized start” on page 100.

7.6 Adjusting the favorites menu



The favorites menu allows trainers to store their preferred training plan selections. These selection options are called favorites and represent a preselection of training plans. Preselections can be stored on transponder cards. When a transponder card is inserted, an athlete/patient can only choose from the set preselections.

1. ➤ Open the “Setup” menu tab.
⇒ The “Setup” menu tab opens.
2. ➤ Open the “Favorites menu settings” menu item.
⇒ The “Favorites menu settings” menu item opens.

3. ➤ Use the multi-function button to select a workout of your choice and press the button to confirm your selection.
i You can select all types of workouts (including all training programs and training plans you have created).
 ⇒ The corresponding workout is selected.
4. ➤ Turn the multi-function button to set the “YES/NO” selection box to the “YES” option.
 ⇒ The corresponding workout has been assigned to the favorites.
5. ➤ Confirm the assignment by pressing the multi-function button.
6. ➤ Use the multi-function button to select the “Setup menu” menu item and press the button to confirm your selection.
 ⇒ The “Favorites menu settings” menu is closed.

7.7 Backing up data

Personnel: ■ Trainer
 Materials: ■ miha bodytec USB flash drive

i “Backup” will not be enabled unless you insert the miha bodytec USB flash drive into the device before switching to the “Setup” menu tab.

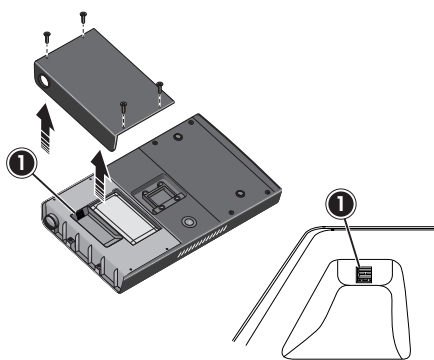


Fig. 30: USB port

1. ➤ Remove the service cover with the help of a screwdriver.
2. ➤ Insert the miha-bodytec USB flash drive into the USB port (Fig. 30/1).
3. ➤ Open the “Setup” menu tab.
 ⇒ The “Setup” menu tab is opened.
4. ➤ Use the multi-function button to select the “Data backup” menu item and press the button to confirm your selection.
5. ➤ Turn the multi-function button to the left to set the selection box to “YES” and press the multi-function button to back up the data.
i The backup files written to the USB flash drive may only be copied or moved. Subsequently edited files will no longer be recognized by the device and can therefore no longer be used.
 ⇒ The device settings, the created training sessions, and the device statistics are saved to the miha bodytec USB flash drive as backup files.
6. ➤ Remove the miha bodytec USB flash drive.
7. ➤ Close the service cover with the help of a screwdriver.

7.8 Restoring/importing data

Personnel: ■ Trainer

Materials: ■ miha bodytec USB flash drive

i “Restore data” will not be enabled until you insert the miha bodytec USB flash drive containing the backup files into the device before switching to the “Setup” menu tab.

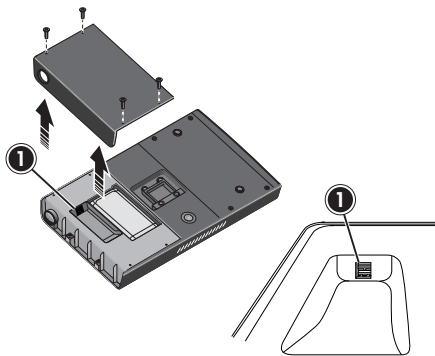


Fig. 31: USB port

Importing settings

1. ➤ Remove the service cover with the help of a screwdriver.
2. ➤ Insert the miha bodytec USB flash drive in the device.
3. ➤ Open the “Setup” menu tab.
⇒ The “Setup” menu tab is opened.
4. ➤ Use the multi-function button to select the “Restore data” menu item and press the button to confirm your selection.

5. ➤ Use the multi-function button to select the “Import settings” menu item and press the button to confirm your selection.
6. ➤ **i** You can use the selection box to define whether or not the device settings stored in the backup files are supposed to be imported and applied.

Use the multi-function button to select the “YES” or “NO” selection box and press the button to confirm your selection.

Importing training programs

7. ➤ Use the multi-function button to select the “Import training programs” menu item and press the button to confirm your selection.
8. ➤ **i** You can use the selection box to define whether or not the training plans and programs stored in the backup files are supposed to be imported and applied.

Use the multi-function button to select the “YES” or “NO” selection box and press the button to confirm your selection.

Starting data restoration


9. ➤ **i** After selecting the data packages you wish to import, you can start restoring the data stored on the miha bodytec USB flash drive.

Use the multi-function button to select the “Start process” menu item and press the button to confirm your selection.

10. ➤ **i** All settings and training programs as well as all training plans stored on the device will be erased and replaced with the contents of the backup files.

Use the multi-function button to select “YES” selection box and press the button to confirm your selection.

⇒ Data restoration starts.

11.  *Once the data has been restored/imported successfully, it is necessary to restart the device.*

Press the on/off switch on the rear of the device.

⇒ The device is switched off.

12. ➤ Remove the miha bodytec USB flash drive.

13. ➤ Close the service cover with the help of a screwdriver.

14. ➤ Press the on/off switch on the rear of the device.

⇒ The device starts up. The data has been restored.

7.9 Installing updates

Data errors due to computer viruses

WARNING

Risk of injury due to data errors caused by computer viruses!

- Only use the specified download paths for update files provided in the miha bodytec newsletter.
- Only use computers with updated anti-virus software when handling update files
- Before copying the update file to the miha bodytec USB flash drive, scan the computer for viruses.

Computer viruses can change the update files. This can result in errors in the control unit that affect the electrical outputs.

Interruption of the update process

NOTICE

Interrupting the update process can cause material damage!

- Do not disconnect the power supply during an update.
- Do not switch off the device during an update.
- Wait for the update to finish and do not input anything.

The device can be permanently damaged if it is turned off during the update process or if the power supply is interrupted.



If the update is interrupted due to an interruption of power, contact Customer Service (📞 “Customer service” on page 16).

How do I receive an update file?

miha bodytec can send important updates by mail on a miha bodytec USB flash drive. Important updates must be installed immediately.

Installing updates

- Personnel: ■ Trainer
- Materials: ■ Latest update file
■ miha bodytec USB flash drive



Keep the device updated at all times.

1. ➤ Save the relevant update file on the miha bodytec USB flash drive.
2. ➤ Remove the service cover with the help of a screwdriver.
3. ➤ Insert the miha bodytec USB flash drive with the update file into the USB port on the underside of the device (Fig. 32/1).
4. ➤ Open the “Setup” menu tab.
⇒ The “Setup” menu tab is opened.
5. ➤ Use the multi-function button to select the “Update” menu item and press the button to confirm your selection.

i If the “Update” menu item has a gray background, the update file on the miha bodytec USB stick is defective, or the miha bodytec USB flash drive has not been fully inserted. Ensure that the update file is valid and that the miha bodytec USB flash drive is fully inserted.

⇒ A prompt appears asking if you want to perform an update.

6. ➤ Answer the prompt with “Yes”.
⇒ The update process starts. This can take several minutes.
7. ➤ Wait for the update, do not disconnect the power supply, and do not switch off the device.
⇒ After the update process is complete, you will see a prompt to manually turn the device off and on again.
8. ➤ Switch off the device.
9. ➤ Remove the miha bodytec USB flash drive.
10. ➤ Close the service cover with the help of a screwdriver.
11. ➤ Switch on the device.

i It takes approx. 17 seconds after the device has been switched on before it is ready for use.

⇒ The update is now complete and the device is ready for use.

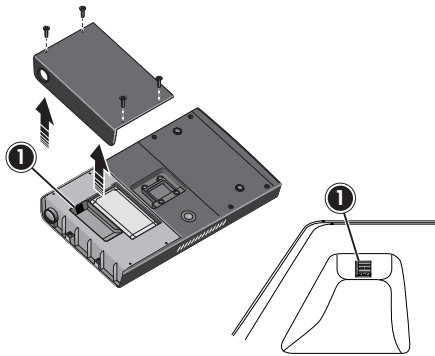


Fig. 32: USB port

7.10 Using networks

Wi-Fi

The device is fitted with a Wi-Fi module so it can exchange data with the miha bodytec LOGX portal.

It can only be used in networks with the designated and preconfigured Wi-Fi access point, which is available upon request from miha bodytec.

miha bodytec LOGX portal



The device's network capability is used for communicating with the miha bodytec LOGX portal. The miha bodytec LOGX portal is a control center for commercial use. It is provided by miha bodytec.

7.11 Resetting to factory settings



A security code can be used for resetting the device settings to the state they were in when it was delivered. This process deletes customized settings. If you wish to reset the device, contact Customer Service (📞 “Customer service” on page 16).

7.12 Displaying statistics

1. ➡ Open the “Setup” menu tab.
⇒ The “Setup” menu tab opens.
2. ➡ Use the multi-function button to select the “Statistics summary” menu item and press the button to confirm your selection.
⇒ The “Statistics summary” menu opens.

You can view the following statistics in the “Statistics summary” menu:

- Device runtime (operating hours counter)
- Programs (total runtime divided across the programs)
- Customers (total runtime divided across customer card numbers)



Example:

- 25:30 equals 25 hours and 30 minutes.

Statistics are displayed in hours and minutes.

8 Before the training

8.1 Eating and drinking

Nutrition

Eat a high carbohydrate meal 2 – 3 hours before the training to achieve the best possible training effect.

Drink at least 17 fl oz of liquid 30 – 45 minutes beforehand. Luke-warm water is ideal. Avoid beverages containing caffeine and alcoholic drinks due to the increased fluid excretion.

Mineral drinks and energy drinks

Drink mineral drinks or high-carbohydrate energy drinks immediately before the training. Sufficient liquid supply with carbohydrate drinks must be guaranteed during as well as after the training to prevent hypoglycemia (low blood sugar).

8.2 Selecting undergarments and shoes

Wearing miha bodytec undergarments

The miha bodytec undergarments available as equipment have been specially developed for use with the device with respect to material composition and strength. It is mandatory to wear the genuine miha bodytec undergarments (🔗 *“Undergarments” on page 15*).

Ensuring that undergarments fit correctly

Always pay attention to the correct size. The undergarments must fit closely everywhere, but must not cut into the skin or be too tight. Pay attention that pants and shirt do not overlap under the electrode areas on the abdomen and back. Doubled fabric layers change pulse transfer, making the actual workout different from what was planned.

Choosing the correct undergarment size

Basically, the sizes of the undergarments correspond to the usual clothing sizes. In addition, the hip width can be used as a more detailed indication:

	XS	S	M	L	XL	XXL
Hip width men	2'11"	3', 3'1"	3'2", 3'3", 3'4"	3'5", 3'6"	3'7", 3'8"	3'9", 3'10", 3'11"
Hip width women	2'9", 2'10"	2'11", 3', 3'1"	3'2", 3'3", 3'4"	3'5", 3'6"	3'7", 3'8", 3'9", 3'10"	3'11", 4', 4'1", 4'2", 4'3"

Wear slip-resistant training shoes

Always wear sport shoes with slip-resistant soles during training.

8.3 Moistening and applying electrodes

Moistening and applying the electrodes



Electrodes may only be moistened and applied by the trainer or under their supervision.

Incorrectly attached cables

WARNING

Risk of injury in the event of incorrectly attached cables!

- Electrodes may only be moistened and applied by the trainer or under their supervision.

Cables not attached to the correct electrodes present a risk.

Changing attachment during operation

WARNING

Risk of injury if detaching or connecting electrode cables during operation!

- Start by making sure that no training program is active (🔗 “Opening the main menu” on page 58), then connect or detach the electrodes.
- Never detach electrode cables while training.
- Never connect electrode cables while training.

Detachment or connection of electrode cables can result in injuries that depend on the settings, such as torn muscle fibers or shoulder dislocations.

Damages

WARNING

Risk of injury due to damaged cables!

- Check the cables and electrodes for damage before every workout.
- Never use the device if cables or electrodes are damaged.

Training with damaged cables or damaged electrodes can lead to injuries.

Unsuitable accessories

WARNING

Risk of injury from wearing accessories that are too small or too large!

- Select the i-body® in accordance with the instructions 🔗 “Choosing the correct i-body® size” on page 74.
- Select i-body® straps flex and i-body® belt in accordance with the instructions 🔗 “Selecting the size of the i-body® straps flex” on page 74.

Wearing accessories that are too small or too large (i-body®, i-body® straps flex, i-body® belt) can lead to contact injuries.

Before the training

Choosing the correct i-body® size



You can find the tag at the lower end of the zipper of the i-body®.

The i-body® is available in the following unisex sizes:

i-body® version	Size
1	Small
2	Medium
3	Large
V1	Small + additional adjustment options
V2	Large + additional adjustment options

Select the i-body® vest so that it fits snugly on your body and does not dig in anywhere. If the vest is loose, select a smaller size. If the vest is too tight, select a larger size.

Selecting the size of the i-body® straps flex

i-body® straps flex are available in 3 different versions:

i-body® straps flex version	Size
1	Small
2	Medium
3	Large

Select i-body® straps flex so that they fit your arms and legs closely and do not dig in anywhere when closed.

The version information provides a guideline. If i-body® straps flex never fit snugly when closed, select a smaller size. If the hook and eye cannot be closed, select a larger size.

Selecting the size of the i-body® belt

i-body® belts are available in 2 different versions:

i-body® belt version	Size
1	Small
2	Large

Select the i-body® belt so that the buttocks belt fits snugly on your body and does not cut in anywhere. If the buttocks belt is loose, select a smaller size. If the buttocks belt is too tight, select a larger size or use an extension piece to enlarge it.

Moistening instructions



Fig. 33: Pump spray bottle

Personnel: ■ Trainer

1. ➤ Spread out the opened i-body® and the i-body® straps flex on a level surface for moistening.
2. ➤ Moisten the electrode surfaces evenly using the pump spray bottle filled with water for a minimum of 3 seconds per electrode.

⚠ DANGER

Risk of fatal injury from improper exposure of the electrodes to moisture!

- When moistening and putting on the electrodes, ensure that there is a sufficient distance to the control unit. No water must penetrate the case.

There is a risk of fatal injury due to electrocution in the event of liquid penetrating the case of the control unit!

i

Do not use distilled water. Do not exceed a water temperature of +86 °F. Only fill the pump spray bottle up to the max. marking visible on the side. Otherwise, no pressure can be built up. Pump air into the bottle by moving the pumping piston up and down (Fig. 33/1).

Before the training

Moistening and applying electrodes



If the electrodes are not moistened correctly, the pulses cannot be transmitted correctly. If the electrodes are moved, always moisten them again.

Personnel: ■ Athlete/Patient
 ■ Trainer

Preparing the i-body®

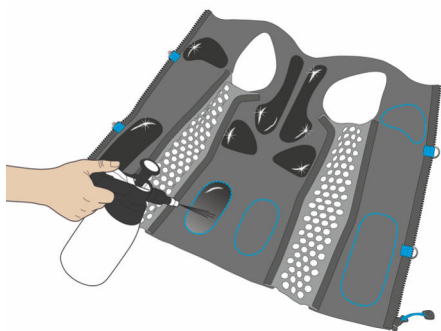


Fig. 34: Moistening the i-body®

1. ➤ Have an i-body® of the correct size ready for use
 ↳ “Choosing the correct i-body® size” on page 74.
2. ➤ Spray the i-body® with water until a visible film of water has formed on the electrodes (Fig. 34).

Putting on the i-body®

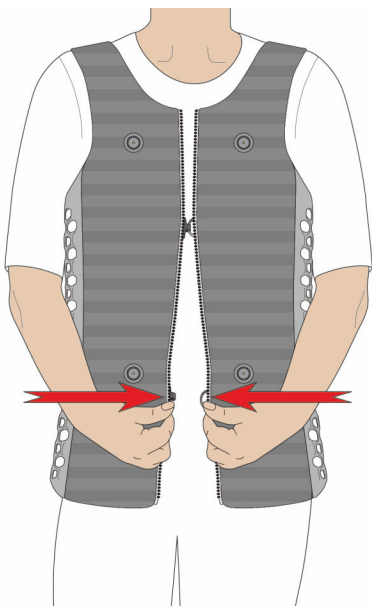


Fig. 35: Wearing the i-body®

3. ➤ Put on the i-body®.

i Close the “fitting aid” (hook and loop in the area of the chest and abdomen) in accordance with Fig. 35.

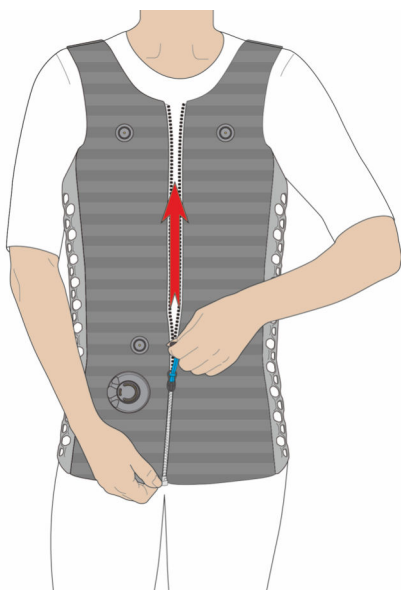


Fig. 36: Closing the i-body®

4. ➔ Close the i-body® with the zipper (Fig. 36).



Fig. 37: Smoothing out the i-body®

5. ➔ Smooth out the i-body® (Fig. 37).

i For optimum fit of the i-body® and the electrodes, it can be pulled into place on the fabric in the front and rear. However, never pull on the silicone on the side to smooth out the i-body®.

Putting on and closing the i-body® straps flex (arms)

Before the training

6. ➔ Moisten 2 i-body® straps flex (Fig. 38).



Fig. 38: i-body® straps flex

7. ➔ Pre-stretch the i-body® strap flex and strap it on the arm.

i The pushbutton and the loops (Fig. 39/1) on the i-body® strap flex must face outwards.

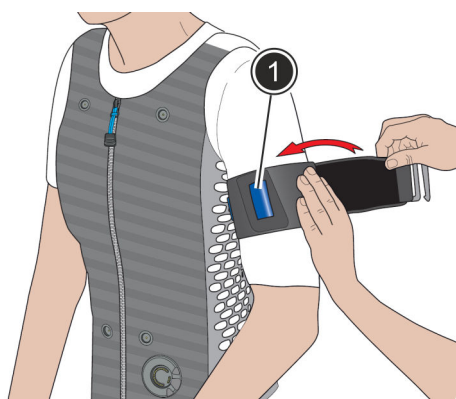


Fig. 39: Putting on the i-body® strap flex

8. ➔ Insert the hook into one of the loops (Fig. 40). Adjust the width so that it is comfortable for the athlete and at the same time provides enough contact pressure.

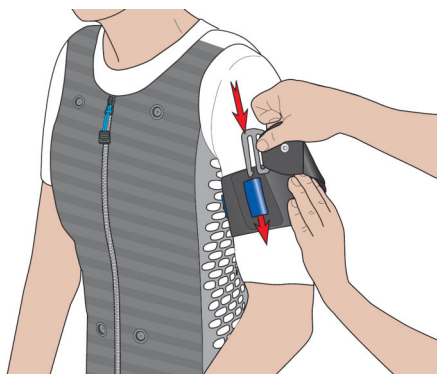


Fig. 40: Closing the i-body® strap flex

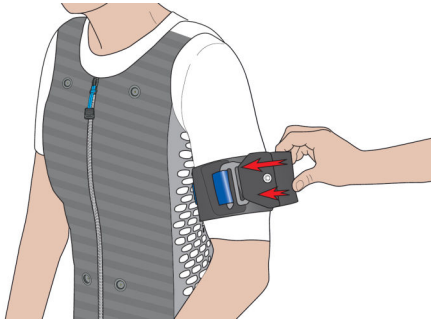


Fig. 41: Positioning the i-body® strap flex

9. ➤ Turn the i-body® strap flex so that the pushbutton faces outwards (Fig. 41).
10. ➤ If the i-body® strap flex does not fit properly, select a different loop to make it tighter or wider. If necessary, select a different i-body® straps flex size.
11. ➤ Put the i-body® strap flex on the right arm the same way as on the left arm.

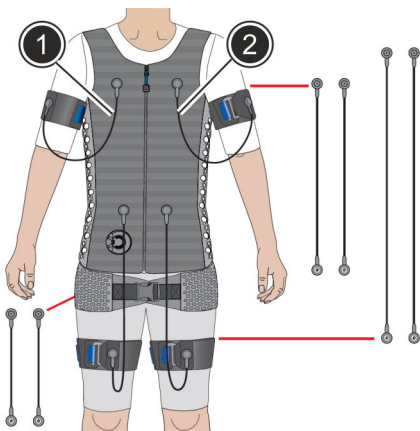


Fig. 42: Connecting the i-body® straps flex

12. ➤ Connect both i-body® straps flex to the i-body® (Fig. 42/1 and 2) to the i-body® cable.

! When connecting the i-body® straps flex, always ensure that the corresponding cables (medium length) are used, in accordance with Fig. 42.

Putting on and closing the i-body® straps flex (legs)

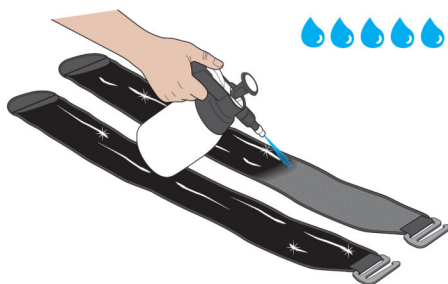


Fig. 43: i-body® straps flex

13. ➤ Moisten 2 more i-body® straps flex (Fig. 43).

Before the training

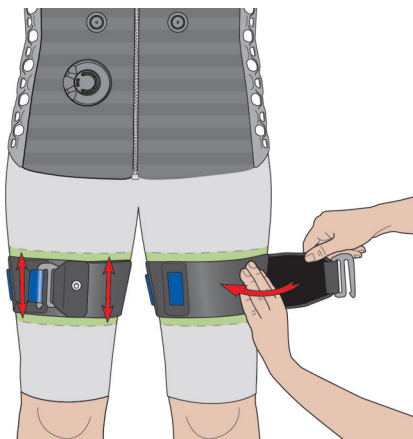


Fig. 44: Putting on the i-body® strap flex

14. ➤ Pre-stretch one i-body® strap flex and strap it on the leg (Fig. 44).

i The pushbutton and the loops on the i-body® strap flex must face outwards.

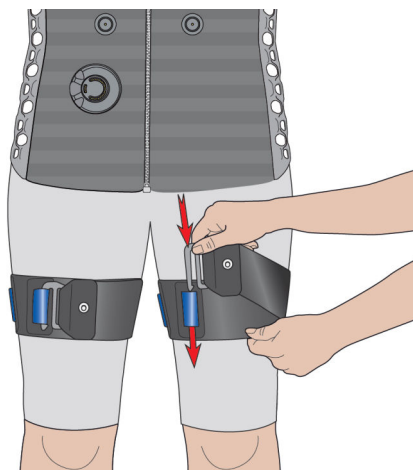


Fig. 45: Closing the i-body® strap flex

15. ➤ Insert the hook into one of the loops (Fig. 45). Adjust the width so that it is comfortable for the athlete and at the same time provides enough contact pressure.

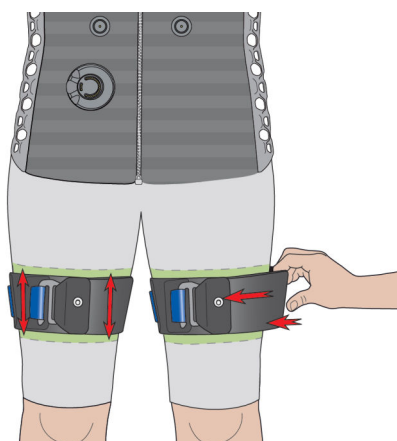


Fig. 46: Positioning the i-body® strap flex

16. ➤ Turn the i-body® strap flex so that the pushbutton faces forwards (Fig. 46).
17. ➤ If the i-body® strap flex does not fit properly, select a different loop to make it tighter or wider. If necessary, select a different i-body® straps flex size.
18. ➤ Put the i-body® strap flex on the right leg in the same way as described for the left leg.

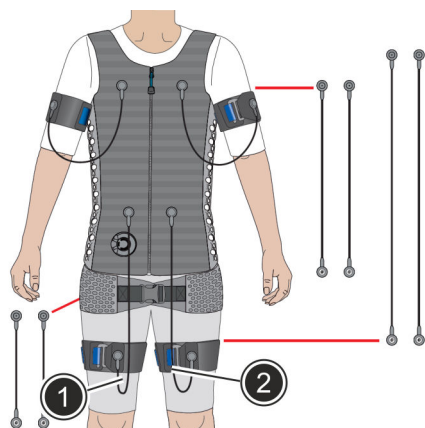


Fig. 47: Connecting the i-body® straps flex

- 19.** Connect both i-body® straps flex to the i-body® (Fig. 47/1 and 2) with the i-body® cable.

***i** When connecting the i-body® straps flex, always ensure that the corresponding cables (longest length) are used, in accordance with Fig. 47.*

Putting on and connecting the i-body® belt

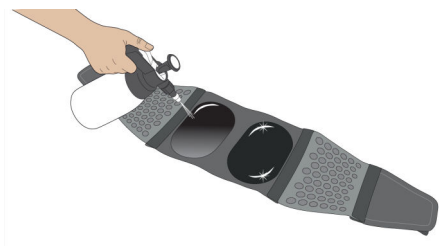


Fig. 48: i-body® belt with buttocks electrodes

- 20.** Moisten the electrodes of the i-body® belt (Fig. 48).

***i** Connect the electrodes to the cables; then put on the i-body® belt.*

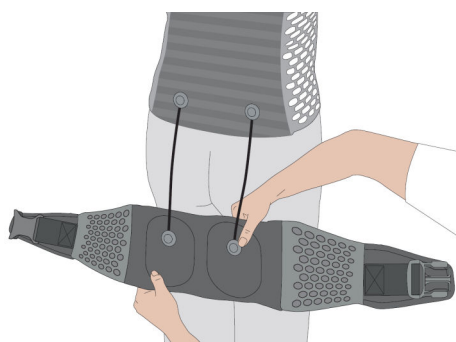


Fig. 49: Connecting the i-body® belt

- 21.** Connect both electrodes of the i-body® belt to the i-body® (Fig. 49) to the i-body® cable.

***i** When connecting the i-body® belt, always ensure that the corresponding cables (shortest length) are used, in accordance with Fig. 47.*

Before the training

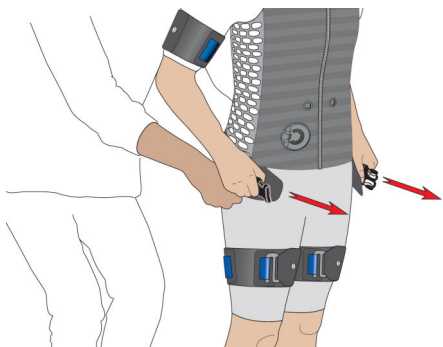


Fig. 50: Fastening the i-body® belt

22. ► Securely fasten the i-body® belt (Fig. 50).

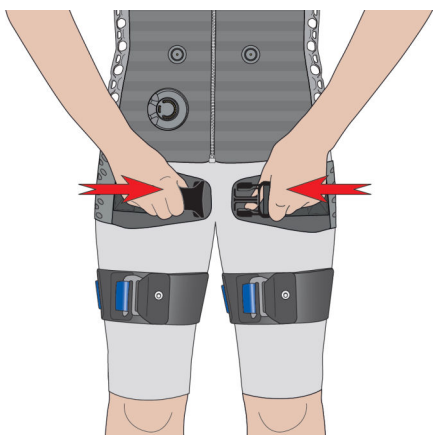


Fig. 51: Securing the i-body® belt

23. ► Secure the i-body® belt with the release buckle (Fig. 51).

24. ► **⚠ WARNING! Risk of injury if connecting while the training program is running!**

Make sure that no training program is active.

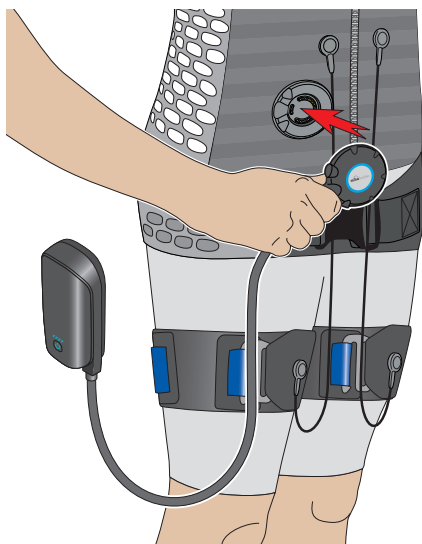


Fig. 52: Connecting

25. ► Connect the cable of the i-body® connect wireless to the magnetic connector of the i-body® (Fig. 52). See the “i-body® connect wireless” manual for further information.

⇒ All electrodes are moistened, attached, and connected as instructed. The i-body® connect wireless has been safely applied to the athlete according to the “i-body® connect wireless” manual and the cable of the i-body® connect wireless is firmly connected to the vest. Training can now start.

9 Training with the device

9.1 Safety during training

Electrical voltage



Risk of fatal injury from power supply!

- Only use the device in a dry environment.
 - Never open the control unit and power supply.
 - Always inform Customer Service (☎ *“Customer service” on page 16*) in the event of problems with the device.
 - In the event of damage to the insulation, switch off the power supply immediately and unplug the mains plug.
 - Observe the safety rules:
 - Unplug the mains plug.
 - Ensure that nobody reconnects the mains plug to the power supply.
 - Never connect the power supply to any voltage other than 100 – 230 V ~ 47 – 63 Hz.
 - Only use the genuine power supply unit provided by the manufacturer
- If the power supply must be replaced, only use the genuine power supply (GlobTek Medical/ITE Power Supply; Model No.: GTM91099-6024-6.0-T2).

There is an immediate risk of fatal injury due to electrocution in the event of contact with live parts. Damage to the insulation or individual components can be fatal. There is a risk of fatal injury in the event of liquid penetrating the case!

Operation without proper attachment



Risk of injury from operating the device without proper attachment!

- The device must not be placed on a table or the ground during training.
- Only use the device if it is connected to the attachment of the “work station m.ove” or the “travel station m.ove” (see ☎ *“travel station m.ove” on page 11*) as described in the assembly instructions of the “travel station m.ove” or the “work station m.ove”.

If the device is not secured by an attachment, it can fall to the ground or be damaged by impacts. This can cause malfunctions injuring the athlete/patient.

Condensation

NOTICE

Risk of damage to the device from condensation!

- Only use the device in a dry environment.
- Never store the device in damp places.
- Let the device acclimatize in the event of location changes.
- In the event of faults, immediately disconnect the device from the power supply and notify Customer Service (🔗 “Customer service” on page 16).
- Never switch on the device if condensation is visible.

Condensation can form in the event of changes in humidity and temperature. Moisture in the control unit can result in damage or even total destruction.

Observing the operating temperatures

NOTICE

Risk of damage to the device from not observing the operating temperatures!

- Before the workout, ensure that the device is at the prescribed operating temperature 🔗 Chapter 3 “Technical data” on page 36.
- Observe the time needed to cool down or warm up the device 🔗 Chapter 3 “Technical data” on page 36.

9.2 Connecting the device

Connecting the control unit

Personnel: ■ Trainer

The control unit can be operated with or without a power supply unit. The connection between the control unit and the person training is always wireless. For more information, see the “i-body® connect wireless” manual.

1. ➔ Remove the connector cover of the electrical connection on the control unit (Fig. 53/1). Connect the magnetic plug of the power supply unit (Fig. 53/2) into the magnetic socket of the control unit.

2. ➔ **i** When connecting the power cable (Fig. 53/2), make sure the magnetic plug connection is safely locked in place.

Connect the power cable to the power supply unit (Fig. 53/2).

- ⇒ Attach the control unit to either the “travel station m.ove” or the “work station m.ove” accessory. For more information, see the assembly instructions for the “travel station m.ove” or the “work station m.ove”.

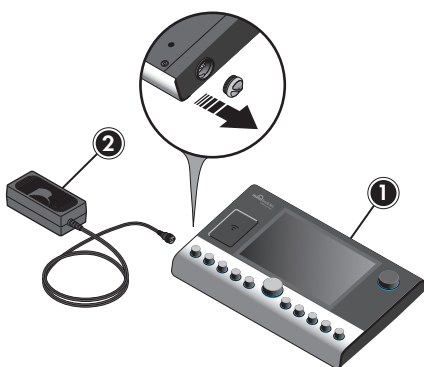



Fig. 53: Connecting the control unit

⚠ CAUTION

Danger due to improper attachment of control unit!

- The miha bodytec m.ove may only be used for training in combination with the “work station m.ove” or the “travel station m.ove” (see  “travel station m.ove” on page 11).

The device must not be placed on a table or the ground during training.

9.3 Operation with the rechargeable battery

Charging the rechargeable battery

Personnel: ■ Trainer

The “miha bodytec m.ove” is equipped with a rechargeable battery for mobile use. It can be charged via the supplied power supply unit both when the device is switched on and when it is switched off. During the charging process, the control unit can be operated as usual.

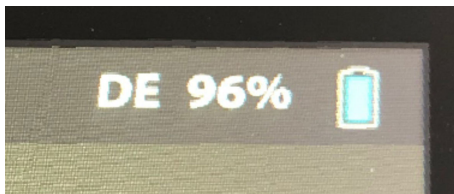


Fig. 54: State of charge of the rechargeable battery

When the device is switched on, the current state of charge of the rechargeable battery is indicated as a percentage via the battery symbol in the upper right corner of the display (Fig. 54).

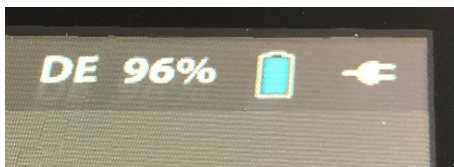


Fig. 55: Rechargeable battery is charging

When the power supply is connected, the rechargeable battery is charged automatically. This is indicated at the same place by an additional plug symbol (Fig. 55).



If the rechargeable battery is used as intended and after approximately 300 discharge cycles per year, it is recommended to replace the rechargeable battery after approximately 24 months. For information on how to replace the rechargeable battery, please refer to the “miha bodytec m.ove” service manual.

9.4 Individual training settings

Specifying settings



The individual adjustment of the device is required to support correct training techniques, training positions, and training processes. This is why these settings must always be determined by a trainer.



Every person reacts differently to current intensity. Therefore, the current intensity must be regulated individually depending on the respective current sensitivity.

10 Overview of programs/training plans

10.1 Selecting individual training settings

Specifying settings



The individual adjustment of the device is required to support correct training techniques, training positions, and training processes.



The individual adjustment of the device is required to support correct therapy techniques, therapy positions, and therapy processes.



Every person reacts differently to the current intensity. Therefore, the current intensity must be regulated individually depending on the respective current sensitivity.

10.2 Stress standards

Pulse intensity

- Currents in the range of thousandths of an ampere (mA) are used in the EMS.
- The pulse intensity determines the degree to which the nerves and muscle fibers are stimulated.
- The stronger the pulse, the stronger the contraction.
- Every person reacts differently to the current intensity. Therefore, the current intensity must be regulated individually depending on the respective current sensitivity.

Pulse duration

The pulse duration specifies the time in seconds (s) or milliseconds (ms) during which a current is output and the muscle is held in the shortened (contracted) condition.

Pulse frequency

The pulse frequency is displayed in Hertz (Hz). The pulse frequency specifies how many individual pulses per second act on the muscles during the contraction phase.

- Each pulse results in a muscle contraction.
- The number of muscle contractions is increased by increasing the pulse frequency.
 - 7 Hz: Improved circulation and metabolic stimulation
 - 85 Hz: Optimum stimulation of striated muscles
 - 100 Hz: Relaxation and pain relief

Pulse width

- The pulse width describes the time duration of a single pulse.
- The longer a single pulse lasts, the deeper it penetrates the tissue and increases the stimulation of the moving parts.

Pulse rise

- The pulse rise specifies in which time a current pulse increases to its maximum value.
- Rectangular pulses are preferred due to the rising slope of the pulse and the resulting effective stimulation.

Overview of programs/training plans

Pause time

- The pause time is the time in seconds (s) or milliseconds (ms) during which no current is flowing.
- The recommendations for an optimum pulse-pause ratio vary between 1:1 and 1:5.
- Guide values for general strength training:
 - A good recovery between the individual contractions.
 - High performance for each individual contraction.
- Similar principles for stress and recovery apply as for conventional strength training (🔗 [Chapter 5.3 “Training frequency and regeneration” on page 49](#)).

10.3 Training programs

Overview of training programs

No.	Program	Duration	Frequency	Pulse duration	Pulse pause	Pulse rise	Pulse width
1	Impulse familiarization	5 min	85 Hz	Continuous	0 s	0 s	350 µs
2	Invigoration - Basic	20 min	85 Hz	4 s	4 s	0.4 s	350 µs
3	Invigoration - Advanced	20 min	85 Hz	4 s	4 s	0 s	350 µs
4	Muscular endurance	20 min	7 Hz	Continuous	0 s	0 s	350 µs
5	Body relax	10 min	100 Hz	1 s	1 s	0 s	150 µs
6	Test training	12 min	85 Hz	4 s	4 s	0.4 s	350 µs



The device uses bipolar pulses and supplies all channels equally during all programs.

Impulse familiarization (1)

Continuously emitted pulse to start off the workout.

Invigoration (2 and 3)

The invigoration programs are used for muscle building and tightening and strengthening the connective tissue and the skin. At the same time, the strengthening programs increase the basal metabolic rate (BMR), thus ensuring that a lot more calories are burnt.

The basic option (2) allows beginners to experience an adequate workout feeling by providing for a more gently rising pulse wave. When selecting the advanced option (3), users will experience an instantaneous pulse rise, resulting in an even stronger contraction of the muscles.

Muscular Endurance (4)

The muscular endurance program is intended to stimulate weight loss among other things. The requirement for this is a combination with the strengthening programs (2 and 3) to increase muscle mass and reduce body fat levels by building muscle and increasing muscular endurance. Activation and tightening of the connective tissue also takes place due to the pulse type used here.

Body Relax (5)

The "Body Relax" relaxation program provides body relaxation, stress reduction, and slightly improved blood circulation of the tissue for removal of metabolic waste products. The muscles function imperceptibly. The feeling the athlete/patient should experience is entirely pleasant.

Test training (6)

The test training program has the identical parameters like the invigoration-basic program. The only difference is the length of the program and is therefore not to be regarded as another output mode. It is ideal for starting EMS training thanks to its shorter length and gentle familiarization process. For EMS beginners in particular, it is important to follow the instructions on training frequency and regeneration ↗ *Chapter 5.3 "Training frequency and regeneration" on page 49.*

10.4 Training plans

Optimized sequences

The training plans provided by the device combine the training programs (↗ *Chapter 10.3 "Training programs" on page 88*) with specially designed exercises. The sequences perfectly complement the intended training targets.

Overview of programs/training plans

Training plan 1

Color	Red
Name	Fitness
Program	Invigoration – basic (2)
Time	20 min

Exercise sequence	Exercise	Number of repetitions
0	Basic position	18
1	Dynamic squats	12
2	Dynamic forward lunges, left	12
3	Dynamic forward lunges, right	12
4	Dynamic diagonal crunches, left	12
5	Dynamic diagonal crunches, right	12
6	Dynamic knee compression	12
7	Dynamic body rotation, left	12
8	Dynamic body rotation, right	12
9	Dynamic crunches	12
10	Dynamic forward fold	12
11	Dynamic biceps	12

Overview of programs/training plans

Training plan 2

Color	Green
Name	Invigoration back
Program	Invigoration – basic (2)
Time	20 min

Exercise sequence	Exercise	Number of repetitions
0	Basic position	18
1	Dynamic squats	15
2	Dynamic butterfly reverse diagonal, left	12
3	Dynamic butterfly reverse diagonal, right	12
4	Dynamic body rotation, left	12
5	Dynamic body rotation, right	12
6	Dynamic diagonal crunches, left	12
7	Dynamic diagonal crunches, right	12
8	Dynamic crunches	15
9	Dynamic overstretching	15
10	Dynamic forward fold	15

Overview of programs/training plans

Training plan 3

Color	Blue
Name	Sports performance
Program	Invigoration – advanced (3)
Time	20 min

Exercise sequence	Exercise	Number of repetitions
0	Basic position	18
1	Dynamic squats	15
2	Dynamic forward lunges, left	10
3	Dynamic forward lunges, right	10
4	Dynamic side lunges, left	10
5	Dynamic side lunges, right	10
6	Dynamic knee compression	13
7	Dynamic diagonal crunches, left	10
8	Dynamic diagonal crunches, right	10
9	Dynamic body rotation, left	10
10	Dynamic body rotation, right	10
11	Dynamic crunches	13
12	Dynamic forward fold	13

Overview of programs/training plans

Training plan 4

Color	Orange
Name	Body forming
Program	Invigoration – basic (2)
Time	20 min

Exercise sequence	Exercise	Number of repetitions
0	Basic position	18
1	Dynamic squats	13
2	Dynamic knee compression	13
3	Dynamic forward lunges, left	10
4	Dynamic forward lunges, right	10
5	Dynamic side lunges, left	10
6	Dynamic side lunges, right	10
7	Dynamic diagonal crunches, left	10
8	Dynamic diagonal crunches, right	10
9	Dynamic crunches	13
10	Dynamic body rotation, left	10
11	Dynamic body rotation, right	10
12	Dynamic forward fold	13

Overview of programs/training plans

Training plan 5

Color	Yellow
Name	Test training
Program	Test training (6)
Time	12 min

Exercise sequence	Exercise	Number of repetitions
0	Basic position	10
1	Dynamic butterfly reverse	8
2	Dynamic shoulder rotation	8
3	Static forward lunges, left	8
4	Static forward lunges, right	8
5	Static knee compression	8
6	Dynamic crunches	8
7	Static forward fold	8
8	Static single-leg stand, left	8
9	Static single-leg stand, right	8
10	Static broad squats	8

10.5 Training

Incorrect programming

WARNING

Risk of injury from incorrect program selection!

- Only perform pre-programming after extensive experience with EMS training.
- Always be careful with pre-programming.
- Always check the program parameters & Chapter 10 “Overview of programs/training plans” on page 87.

Selecting an incorrect program according to its programming can result in injuries.

Rechargeable battery sufficiently charged



Before starting the application, make sure that the rechargeable battery is sufficiently charged.

When to interrupt the training?

Safety first!

Stop a workout immediately if you feel unwell, have muscle cramps, or are dizzy. Drink fluids and, if problems persist, see a doctor if necessary.

What to do in emergency situations?

A basic requirement of EMS training is that the athlete is supervised by their trainer. The trainer can therefore intervene appropriately in the event of an emergency. Act immediately if an athlete loses consciousness:

Personnel: ■ Trainer

1. ➔ Press the multi-function button.
⇒ The current training stops.
2. ➔ Turn off the device via the on/off switch on its underside (Fig. 56/1).
3. ➔ Detach the i-body® connect wireless incl. the cable from the athlete. See the “i-body® connect wireless” manual for further information.
4. ➔ Initiate first aid measures.

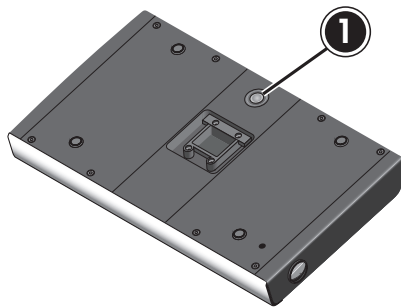


Fig. 56: On/off switch

Stop button



Stop button

The multi-function button is not only used for choosing and confirming the programs, but also as a stop button. The training can be stopped at any time with the multi-function button.

Turning up the intensity



Turning up the intensity during the pulse pause is limited to 5 units both on the level controllers and the main controller.



Main controller

If the “Load level” function is enabled, the device is programmed such that the main controller cuts out at 30, 40, and 50 for safety reasons. Regardless of whether or not the “Load level” function is enabled, an additional cut-out is set for all programs and training plans when the device reaches a value of 85. Before the athlete can increase the intensity beyond this value, the main controller must be acknowledged by pressing it once.

Pausing the training session

Personnel: ■ Athlete/Patient
 ■ Trainer

The training session can be interrupted between individual exercises to pause or discuss the next exercise with the trainer.

1. ➤ Press the main controller (Fig. 57/14).
 - ⇒ The main level jumps to “0” and all LEDs are flashing.
The training time continues running during the pause.
2. ➤ Turn the main controller to end the pause.
 - ⇒ The training session continues as usual.

Electrode detection/channel shutoff



Before starting with the workout (countdown) and during the pulse phases, the electrodes connected via the channel outputs are detected.



If a pair of electrodes is not detected before starting with the workout (countdown), the corresponding channel is shown in gray on the display and cannot be reactivated during the training session. If this happens even though something was connected to this channel, remove the transponder card again right away at the start of the workout. This ends the training session. Next, check to make sure that the contacts are properly connected and the electrodes are sufficiently moistened. Correct the problem, if necessary.



If the contact of a channel is interrupted unexpectedly during the workout, the intensity of the channel drops to “0” and cannot be increased again afterwards. The channel is shown in gray on the display. The channel intensity can only be increased again once the electrode is properly reconnected.

Working out

Personnel: ■ Athlete/Patient
 ■ Trainer



Fig. 57: Control unit

1. ➤ Select the program by turning the multi-function button (Fig. 57/13).
2. ➤ Read and check the parameters of the selected program by turning the main controller (Fig. 57/14).
3. ➤ Confirm the desired program by pressing the multi-function button (Fig. 57/13).
4. ➤ Adjust the application time if necessary by turning the multi-function button.
5. ➤ Press the multi-function button to start the program.
 - ⇒ The time at the top right in the display starts running.



Fig. 58: Basic position

6. ➤ **⚠ WARNING! Risk of injury from uncontrolled muscle contractions!**
- Always build up basic tension right before the pulses.
 - Always pay attention to the light bar on the control unit to see the coming pulse type.
 - Relax for a short time during the interval pauses.
 - Do not exercise unless a second person can intervene in case of an emergency. Never exercise by yourself.

If insufficient basic body tension is established before the pulses, muscles can contract uncontrollably. This can cause muscle injuries.

Adopt the basic position (Fig. 58).

7. ➤ Turn the main controller to increase the pulse intensity to 65%.
8. ➤ Increase the intensity by turning the individual level controllers (Fig. 57/1 – 8) until the pulses have reached the required intensity.
9. ➤ Always pay attention to the light bar on the control unit to see the coming pulse type.
10. ➤ If necessary, adjust the pulse strength during the course of the training.
- ⇒ The pulses stop automatically after the completion of the program. The program is ended prematurely by pressing the main controller.

10.6 Transponder system

Transponder card

⚠ WARNING

Risk of injury if transponder cards are mixed up!

- Always ensure that the correct transponder card is used.
- Always check the program parameters.
- Always start with a low level and then increase it slowly.

Incorrect program parameters can result in the respective settings being too high. This can cause injuries.

The device comes standard with a transponder card reader to maximize operation convenience. The following personalized data can be saved on the transponder card:

- Program and/or training plan
- Time specification
- Last value of the main controller
- Last value of the individual level controllers



The “Load level” function will not be enabled unless the following criteria have been met:

- The “Load level” function on the corresponding customer card must be selected and confirmed with “YES” ↗ Chapter 10.6 “Transponder system” on page 98.
- The “Load level” function must be selected in the device settings and confirmed with “YES”.

The “Load level” function will remain disabled until both of these criteria are have been met. The function will also remain disabled if only one of the confirmations has been selected.

First use of the transponder card

Personnel: ■ Trainer

Materials: ■ Transponder card

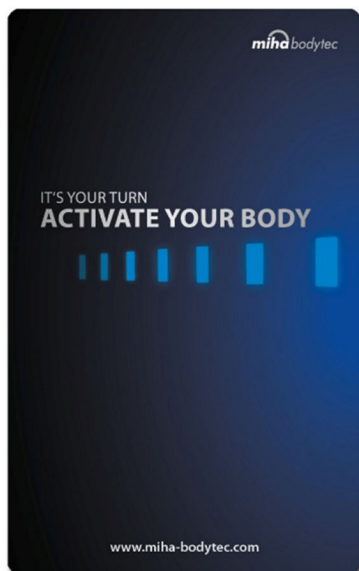


Fig. 59: Transponder card

1. ➞ Place the new transponder card (Fig. 59) in the main menu on the designated contact surface.
2. ➞ If the transponder card is still unformatted, you will be prompted to format the transponder card. Confirm the prompt by selecting “YES” using the multi-function button.
3. ➞ If the transponder card is empty, the following question will be displayed: “Empty card. Create customer card?”. Confirm the prompt by selecting “YES” using the multi-function button.
 - ⇒ The system will automatically switch to the “Card management” menu.
4. ➞ Use the multi-function button to set the different default parameters in the “Card management” menu and save them to the transponder card using the “Write card” function.
 - ⇒ The transponder card will now be formatted with the values you have defined and is ready for use.

Overview of programs/training plans

Using the transponder card

Personnel: ■ Athlete/Patient

Materials: ■ Transponder card

1. ➤ Place the transponder card in the main menu on the designated contact surface.
 - ⇒ The system automatically switches to the selection of the workouts stored on the transponder card. The “*Card management*” menu shows the programs/training plans stored on the transponder card and allows you to select them using the multi-function button.
2. ➤ Press the multi-function button again to start the program/training plan (↪ *Chapter 10.5 “Training” on page 94*).
3. ➤ Complete your training or press the multi-function button again to terminate the workout prematurely.
4. ➤ Remove the transponder card. To do so, press the lower end of the transponder card and slide it upward.

Changing data on the transponder card

Personnel: ■ Trainer

Materials: ■ Transponder card

1. ➤ Remove the transponder card if a card is currently inserted in the device.
2. ➤ Use the multi-function button to select the “*Card management*” menu item.
3. ➤ Place the transponder card on the designated contact surface and change its settings as needed.
4. ➤ Use the multi-function button to select “*Write card*”.
 - ⇒ The changes will be saved on the transponder card.
5. ➤ Remove the card from the designated contact surface.
 - ⇒ The data on the transponder card has been changed.

Setting up the transponder card for training with synchronized start



After the device has been set up as a master or managed device, the user settings for every athlete have to be defined as well. These user settings are linked to the transponder card associated with the athlete. This ensures that athletes will only participate in the training with synchronized start if they wish to do so.

Personnel: ■ Trainer

Materials: ■ Transponder card

1. ➤ Remove the transponder card if a card is currently inserted in the device.
2. ➤ Open the “*Card management*” menu tab.
⇒ The “*Card management*” menu opens.
3. ➤ Insert the transponder card.
4. ➤ Open the “*Synchronized start*” menu item.
5. ➤ Set the current “*Synchronized start*” setting to “YES”.
⇒ The transponder card stores the information that the athlete will participate in the training with synchronized start.
6. ➤ Use the multi-function button to select “*Write card*”.

***i** If the synchronized start settings are correct and a transponder card is placed in position, the synchronized start symbol on the master device lights up blue on the outside and red on the inside, and the managed device lights up green on the outside and blue on the inside.*

Selecting individual language settings for the athletes/patients



Each athlete/patient has the option of selecting an individual language. Once an individual language has been selected, a customized language will automatically be used for all display texts as soon as an athlete/patient logs into the device.

Personnel: ■ Trainer

Materials: ■ Transponder card

1. ➤ Open the “*Card management*” menu tab.
⇒ The “*Card management*” menu opens.
2. ➤ Insert the transponder card.
3. ➤ Open the “*Language*” main menu item.
4. ➤ Select the language by turning the multi-function button.
5. ➤ Select the desired language by pressing the multi-function button.
⇒ The transponder card of the athlete/patient is linked to an individual language setting. Display texts will be shown in the individually selected language when the transponder card has been inserted.
6. ➤ Use the multi-function button to select “*Write card*”.

Selecting training programs from favorites



If favorites have been specified, the athlete/patient can only access the training plans that have been set up as favorites (↪ Chapter 7.6 “Adjusting the favorites menu” on page 66).

10.7 Training with synchronized start

Personnel: ■ Trainer

Materials: ■ Transponder card

Prerequisites

- A master device as well as at least one managed device must be set up correctly.
- Synchronized start must be activated on all participating transponder cards.

1. ➤ Place the transponder cards on the devices.
2. ➤ Select a workout on the master device by pressing the multi-function button once.

***i** If “Flexible training” is activated, the corresponding programs or training plans must be selected on the managed devices. If “Flexible training” is not activated, the managed devices immediately switch to the program selected on the master device.*

3. ➤ Start the workout by pressing the multi-function button on the master device once.
 - ⇒ The countdown begins on all the participating devices. The synchronized start was successful.

11 After the training

11.1 Taking off electrodes

Taking of the electrodes



Electrodes may only be detached by the trainer or under their supervision.

Pulling on the cable

NOTICE

Damage due to pulling on the cable!

- Do not pull on the cable to detach the i-body® straps flex and the i-body® belt.
- Always detach the pushbutton connections directly at the closure.

Removing the i-body® straps flex and electrodes by pulling on the cable can cause damage to the cables and the material.

Taking off electrodes

- Personnel:
- Athlete/Patient
 - Trainer

Detach the electrodes from the cables one after the other.

1. ➔ **▲ WARNING! Risk of injury if detaching the electrodes while the training program is running!**

Make sure that no training program is active. To do so, switch to the main menu (☰ “Opening the main menu” on page 58).

2. ➔ Disconnect the connection between cable of the i-body® connect wireless and the i-body® (Fig. 60). See the “i-body® connect wireless” manual for further information.
3. ➔ Disconnect all i-body® straps flex on arms and legs from the i-body® connect wireless.
4. ➔ Disconnect and remove the i-body® belt from the i-body® connect wireless.

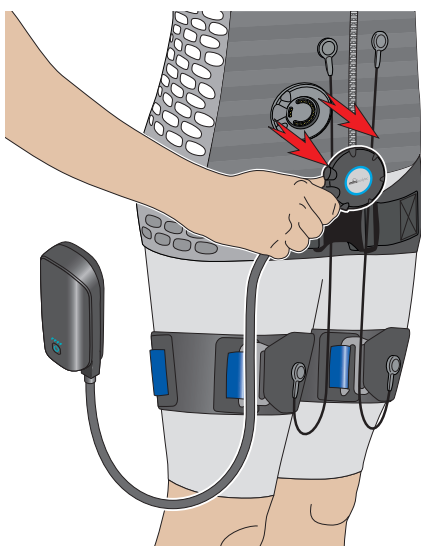
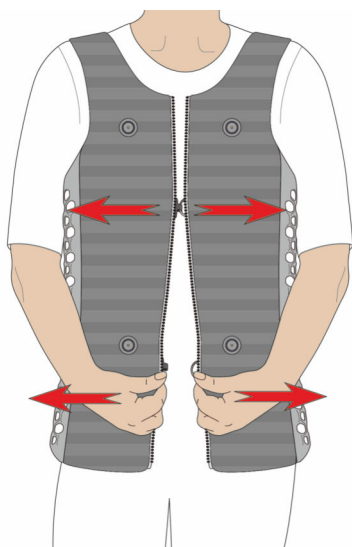


Fig. 60: Detaching the main connection



5. ➡ Open the zipper and the fitting aid and take off the i-body® (Fig. 61).

Fig. 61: Taking off the i-body®

11.2 Cleaning and storage

11.2.1 Cleaning the i-body® and the individual electrodes

Improper cleaning of electrodes

NOTICE

Danger of material damage!

- Remove cables.
- Never use the following cleaning agents:
 - Synthetic cleaning agents
 - Solvents
 - Chlorides
 - Polishing agents
 - Washing/polishing agents
 - Aerosol sprays
- Observe the cleaning symbols ☞ Chapter 2.7.3 “Cleaning symbols” on page 34.
- Do not dry clean the electrodes.
- Do not exceed a water temperature of 105 °F / 40 °C.
- Do not use any fabric softeners and bleach.
- Do not tumble dry.
- Do not iron or treat with steam.
- Do not spin-dry.
- Do not wring out.

Incorrect cleaning can cause material damage of the i-body® electrode vest and individual electrodes.

Observing the cleaning intervals

Interval	Component	Cleaning type
As necessary or after use.	Accessories: ■ i-body® ■ i-body® belt ■ i-body® straps flex	Disinfection <i>☞ Chapter 11.2.1 “Cleaning the i-body® and the individual electrodes” on page 104</i>
If visibly dirty, but at least once a month.	Device	Wipe the control unit, surfaces, and grab handles with a moist cloth. <i>☞ Chapter 11.2.3 “Cleaning the control unit” on page 107</i>
As necessary, but at least once a month.	Accessories: ■ i-body® ■ i-body® belt ■ i-body® straps flex	Cleaning <i>☞ Chapter 11.2.1 “Cleaning the i-body® and the individual electrodes” on page 104</i>

Cleaning the i-body®, i-body® straps flex, and the i-body® belt



Use ISOPROPANOL 70% for disinfection.

The electrode surfaces have an anti-bacterial layer. If necessary, i-body® straps flex, i-body® belt, i-body®, and additional electrodes can be cleaned. To do so, proceed as follows:

Personnel: ■ Trainer

1. Wash the i-body® by hand at a maximum water temperature of 105 °F / 40 °C.

📌 If a washer with a hand wash setting is available, you can also wash the i-body® in the washer using the supplied laundry net.

2. Hang the i-body® on a clothes hanger and let it dry in a well-ventilated place.

3. Wash the i-body® belt by hand at a maximum water temperature of 105 °F / 40 °C.

📌 If a washer with a hand wash setting is available, you can also wash the i-body® belt in the washer using the supplied laundry net.

4. Let the i-body® belt dry in a well-ventilated place.

5. Wash the i-body® straps flex by hand at a maximum water temperature of 105 °F / 40 °C.

📌 If a washer with a hand wash setting is available, you can also wash the i-body® straps flex in the washer using the supplied laundry net.

6. ➡ Let the i-body® straps flex dry in a well-ventilated place.

⇒ The washed electrodes are now clean and can dry. They can be reused once they are dry.

Storing the i-body®, i-body® straps flex and the i-body® belt



All parts except the control unit are subject to wear and must be inspected regularly and replaced if necessary. Only intact equipment guarantees optimal use of the device.

Ambient conditions for storing and safekeeping the accessories
🔗 “Storage of the packages” on page 109.

- Hang the i-body® on a clothes hanger to dry.
- Always attach the wash protection to the i-body® straps flex for storing and drying.
- Sort the i-body® straps flex and the i-body® belts by size and lay them out open to dry in a well-ventilated place.
- Check the hygienic condition of the i-body®, the i-body® straps flex, and the i-body® belts regularly.
- Clean or replace accessories that are in inadequate hygienic condition.

11.2.2 Cleaning the external cables

Improper cleaning of external cables.

NOTICE

Danger of material damage!

- Disconnect cables from control unit and external electrodes
- Never use a washing mashine or dryer for the external cables
- Never use the following cleaning agents:
 - Synthetic cleaning agent
 - Solvents
 - Chlorides
 - Polishing agents
 - Washing/polishing agents
 - Aerosol sprays
- Do not use any fabric softeners and bleach.

Incorrect cleaning can cause material damage of the cables.

Only use a soft, lightly moistened cloth for cleaning the control unit connection cable and the cables for the external electrodes. Never use abrasives or solvents.

Personnel: ■ Trainer

1. ➤ Clean all cables with a soft, slightly moist cloth.

2. ➤ Dry all cables surfaces with a soft, dry cloth.

⇒ All external cables are now dry and clean. This completes the cleaning of the external cables.

11.2.3 Cleaning the control unit

Incorrect cleaning of the control unit

NOTICE

Material damage from incorrect cleaning!

- Unplug the mains plug.
- Only clean with a moist cloth.
- Do not use any abrasives.
- Do not use any solvents.

Incorrect cleaning may cause material damage. The surface can become unsightly and the control unit can even be destroyed.

Cleaning the control unit

Only use a soft, slightly moistened cloth to clean the case parts. Never use abrasives or solvents.

Personnel: ■ Trainer

1. ➤ Clean all control unit surfaces with a soft, slightly moist cloth.

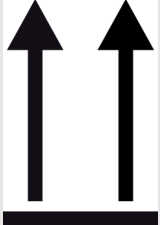


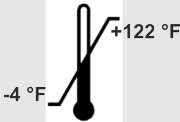
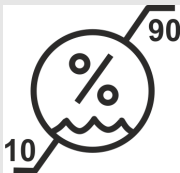
2. ➤ Dry all control unit surfaces with a soft, dry cloth.

3. ➤ Wipe the LC display with a soft polishing cloth.

⇒ All control unit surfaces are now dry and clean. This completes the cleaning of the control unit.

12 Packaging and storage

12.1 Symbols on the packaging

Symbol	Standard Reference	Standard Title	Symbol Title	Explanatory Text
	ISO 7000 Fifth Edition 2014-01-15, 0623	Graphical Symbols For Use On Equipment – Registered Symbols	This way up	The arrows of this symbol show which side of the package is up. Packages must be transported and stored with the arrows pointing upward.
	ISO 15223-1 Third Edition 2016-11-01 Clause 5.3.4	Medical Devices – Symbols To Be Used With Medical Device Labels, Labelling, And Information To Be Supplied – Part 1: General Requirements	Keep dry	Packages with this symbol are sensitive to moisture. These packages must be protected from moisture (e.g., rain) when being transported and stored.
	ISO 15223-1 Third Edition 2016-11-01 Clause 5.3.1	Medical Devices – Symbols To Be Used With Medical Device Labels, Labelling, And Information To Be Supplied – Part 1: General Requirements	Fragile, handle with care	This symbol indicates that the package contains fragile objects. These packages must be transported carefully and protected against shocks. Do not throw packages.
	ISO 15223-1 Third Edition 2016-11-01 Clause 5.3.7	Medical Devices – Symbols To Be Used With Medical Device Labels, Labelling, And Information To Be Supplied – Part 1: General Requirements	Storage temperature range	This symbol on the packages shows the safe temperature limits a device may be exposed to for storage purposes.
	ISO 15223-1 Third Edition 2016-11-01 Clause 5.3.8	Medical Devices – Symbols To Be Used With Medical Device Labels, Labelling, And Information To Be Supplied – Part 1: General Requirements	Storage humidity range	This symbol on the packages shows the safe limits for non-condensing relative humidity within the temperature limits a device may be exposed to for storage purposes. Chapter 12.2 “Transport and storage” on page 109.

Symbol	Standard Reference	Standard Title	Symbol Title	Explanatory Text
	Class 9 – UN3481	Lithium ion batteries contained in or packed with the equipment, installed / integrated at the source.	Caution, lithium ion batteries	This symbol indicates that the device or it's package contains a lithium ion battery.

12.2 Transport and storage

Transporting packages

The symbols listed in [Chapter 12.1 “Symbols on the packaging” on page 108](#) are attached to the packaging. Observe these symbols when handling packages.

Storage of the packages

Store the packages under the following conditions:

- Do not store outdoors.
- Store in a dry and dust-free location.
- Do not expose to aggressive media.
- Protect against direct sunlight.
- Avoid mechanical shocks.
- Storage temperature:
 - -4 °F – +122 °F (-20 °C – +50 °C)
 - +41 °F – +95 °F (+5 °C – +35 °C) at a relative humidity of up to 90%, non-condensing
 - +41 °F – +122 °F (+5 °C – +50 °C) at a water vapor pressure of up to 50 hPa
- Relative air humidity: max. 90%, non-condensing.
- Permissible air pressure: 10 psi – 15.5 psi (680 – 1,070 hPa).
- If storing for longer than 3 months, check the general condition of all parts and the packaging regularly. If necessary, renew or replace preserving agents.



Under certain circumstances, storage instructions may be affixed to packages that extend beyond the requirements specified here. Comply with these instructions accordingly.



The period of storage of the electrodes should not be longer than 24 months.

12.3 Environmental protection

Incorrect disposal of packaging.

ENVIRONMENT!

Incorrect disposal can be hazardous for the environment.

- Dispose of packaging materials in accordance with environmental regulations.
- Comply with local disposal regulations. If necessary, outsource the disposal to a specialist company.

Packaging materials are valuable raw materials. In many cases, they can be re-used or recycled for other uses. Environmental hazards can result from the incorrect disposal of packaging materials.

Disposal

NOTICE

Environmental hazard from incorrect disposal!

- Return electronic scrap and electronic components to the manufacturer.

Contact customer service in case of questions (☎ “Customer service” on page 16).

Environmental hazards can result from incorrect disposal.

The device must not be disposed of via domestic waste.



Battery disposal

The circuit board inside the control unit houses a CMOS battery which is used for storing the device settings when the device is unplugged. In addition, a lithium-ion battery is located behind the service cover on the underside of the control unit, which allows the control unit to be operated in mobile use without a mains power connection. The following battery types are installed:

- Lithium battery CR2032 (3 V, 220 mAh)
- Lithium-ion battery – miha bodytec m.ove; model no.: RRC2054-2 (14.4 V, 6.80 Ah)

The installed batteries must not be disposed of via domestic waste. They must be disposed of in the discharged state in accordance with the regulations applicable at the place of use.

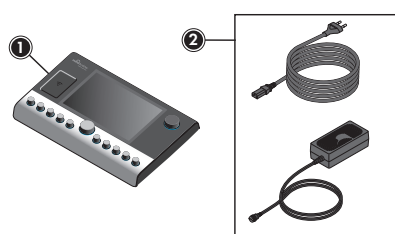
Our products contain rechargeable lithium-ion batteries.

By recycling batteries, it is possible to recover valuable resources such as zinc, iron, nickel etc.

Our registration number with the Umweltbundesamt (German Environment Agency): 21010366

13 Ensuring optimum function

13.1 Maintaining the device



- 1 Control unit
- 2 Power supply and power cable

Fig. 62: Conducting a visual inspection



Fig. 63: Examining the i-body® electrode vest


The device and the i-body® electrode vest must be serviced at regular intervals to ensure that they can assist you successfully during therapy.

⚠ WARNING

Danger from maintenance or service work during use!

- Perform maintenance only when the device is not being used.

No maintenance or service activities may be performed while the device is being used by a patient/athlete. Maintenance of the device during training can cause dangerous situations.

The maintenance tasks described in the  Table on page 112 must be performed at the specified interval to ensure basic safety and compliance with the performance characteristics described in the technical data.

Ensuring optimum function

When must the device be serviced?	What must be serviced?	Who must service the device?
Before every workout	<p>Check the device for signs of damage:</p> <ul style="list-style-type: none"> ■ Control unit (Fig. 62/1) ■ Electrical lines and plug connectors (Fig. 62/2) ■ i-body® (Fig. 63) <p>Replace damaged parts immediately with genuine spare parts of the manufacturer ↗ “Customer service” on page 16.</p> <p>Do not use the device if there are signs of damage.</p>	Trainer
Every 24 months	Check the electrical system.	Licensed electrician

13.2 Maintaining the electrodes

To ensure optimum function, it is recommended to replace the electrodes after 12 months of use.

13.3 Handling error messages

If a malfunction was identified that affects the operational readiness, the active training session (voltage generation) is blocked and an error code is displayed in the status line as well as in the LED line. Configurative operation functions remain enabled.



Notify the manufacturer and the relevant authority of the member state in which the operator is located of all major incidents that occur in connection with the device.

Proceed as follows to fix the error:

1. ➞ Remember or write down the error code.
2. ➞ Switch off the device with the on/off switch.
3. ➞ Switch on the device with the on/off switch.

***i** It takes approx. 17 seconds after the device has been switched on before it is read for use.*

4. ➞ Check whether the error is displayed again.
 - If the error is no longer displayed, continue with the workout.
 - If the error is displayed again or another error code is displayed, notify the manufacturer or Customer Service and report the error codes (↗ “Customer service” on page 16).

14 Index

A

Abdominal wall hernia	19
Accessories	
i-body® cable	15
i-body® connect charger	13
i-body® connect wireless	12
Improper use	28
Rechargeable battery	11
travel station m.ove	11
unsuitable accessories	27
work station m.ove	12
Active medical implants	19
Adjusting the grab handle height	85, 87
Adverse events	29
Arterial circulatory disorders	19
Arteriosclerosis	19
Athlete	20

B

Backup	67, 68
Bacterial infections	19
Basic position	97
Battery	
Disposal	110
Bleeding tendencies	19
Body Relax	89
Brand code	53
Buttocks electrode	14

C

Cardiac arrhythmias	19, 22
Cardiovascular diseases	19
Changing controller color	61
Channel shutoff	96
Channels	36
Circulatory disorders	19
Cleaning	105
Cleaning intervals	105
Cleaning the control unit	107

Cleaning the external cables	106
Compliances	36
Computer viruses	69
Contact data	16
Contraindications	19
Control unit	
Connecting	84
Connections	10
On/off switch	10
Rechargeable battery	10
Top side	8
USB port	10
Copyright	4
Coronary heart diseases	19
Cramps	19
Creatine kinase	50
Current consumption	36
Customer service	16
Customization and setup	
Device settings	61
Displaying device settings	61
Displaying the program memory	63
Displaying training plan memory	63
Favorites menu settings	66
Selecting the default language	62
Statistics	71

D

Data backup	53
Data restoration	53, 68
Device setting	61
Device settings	61
Device update	70
Diabetes mellitus	19
Dimensions	36
Displaying device setting	61
Displaying the program memory	63
Displaying training plan memory	63
Disposal	35, 110

Dizziness	19	Data restoration	53
E		Favorites menu	52
Electrical voltage	23	Loading the level	52
Electrode vest	13	Programs	52
Electrodes		Training plan	52
Connecting	76	F	
Detection	96	Factory settings	
Moistening	76	Resetting	71
Taking off	103	Favorites menu	52
Electronic components	35, 110	Favorites menu settings	66
Emergency	95	Feeling unwell	19
EMS training	7, 48	Fever	19, 49
After the first 10 weeks	50	Flu	49
Blood count	50	G	
Creatine kinase level	50	General information	
During private use	21	Explanation of terms	52, 53
Fluid balance	50	Text boxes	59
Food intake	50	H	
In the first 10 weeks	50	Hazards	7, 48
Myoglobin level	50	Heart pain	19
Special considerations	49	Hemophilia	19
The first workout	50	Hip strap	14
Environmental protection	35, 110	I	
Epilepsy	19	i-body®	13
Equipment		Choosing the correct size	74
Electrode vest	13	Cleaning	105
i-body®	13	Connecting	76
i-body® belt	14	Moistening instructions	75
i-body® straps flex	14	Putting on	76
Pump spray bottle	15	Storing	106
Transponder card	13	i-body® belt	14
Undergarments	15	Choosing the correct size	74
Error code in the status line	112	Cleaning	105
Error display	112	Connecting	76
Error messages	112	Putting on	76
Essential performance	46	Storing	106
Explanation of symbols	32	i-body® cable	15
Explanation of terms			
Data backup	53		

i-body® connect wireless	12, 53	Menus	
i-body® straps flex	14	Colors	57
Choosing the correct size	74	Opening	57
Cleaning	105	Settings in submenus	58
Connecting	76	Structure	56
Moistening instructions	75	miha bodytec	
Putting on	76	Contact data	16
Storing	106	Customer service	16
Implants	19	LOGX portal	71
Impulse familiarization	88	USB flash drive	70
Inflammations	19	Misuse	17, 18
Inguinal hernia	19	Moistening	75
Intended use	17, 18, 36	Multi-function button	8
Interactions	22	Muscular Endurance	89
Invigoration	88	Myoglobin	50
K		N	
Kidney diseases	19	Network	70
Kidneys	19	Neurological disorders	19
L		O	
Level controller	8, 54	On/off switch	10
Lithium-ion battery	25	Opening menu tabs	55
Liver	19	Operating the device	
Liver diseases	19	Opening menu tabs	55
Loading the level	52	Opening menus	57
LOGX portal	71	Selecting menus	57
M		Settings in submenus	58
Main menu	55	Other applicable documents	4
Opening	58	Overview	
Maintenance		Synchronized start settings	65
Checking the mounting screws	111	Training programs	88
Having the device tested	111	P	
Sending in the device	111	Parametrization	53
Manufacturer	36	Pause	96
Medical implants	19	Personalization and setup	
Memory card	36	Backup	67
Menu language		Choice of languages	62
Selecting	62	Data restoration	68
		Menu items of the program memory	63

Index

Menu items of the training plan memory	64	Settings	85, 87
Synchronized start settings	65	Resetting to factory settings	71
Power supply	36	Severe circulatory disorders	19
Pregnancy	19	Skin cancer	19
Preparation	72	Software update	70
Program memory	36, 63	Spare parts	11
Programs	52	unsuitable spare parts	27
Protection class	36	Statistics	71
Protection type	36	Storage	106, 109
Pulses		Stress standards	48
Pulse acclimatization	87	Strong bleeding tendencies (hemophilia)	19
Pulse duration	87	Symbols	
Pulse frequency	87	In this manual	3
Pulse intensity	87	on the device	54
Pulse pause	87	On the packaging	108
Pulse rise	36, 87	Synchronized start	
Pulse time	36	Overview of settings	65
Pulse width	36, 87	Setting up a managed device	66
Pump spray bottle	15	Setting up the master device	65
Moistening electrodes	75	Setting up the transponder card	100
Putting on	76	Steps to set up	65
Q		Training	102
Qualification	26	Training with synchronized start	64
R		System status	54
Rating plate	31	T	
Recent operations	19	Technical data	36
Rechargeable battery	11	Test training	89
Charging	85	Text boxes	59
Regeneration	51	Trainer	20
Replaceable Parts	46	Training	85, 87, 97
Residual risks	7, 48	Choosing the correct shoes	72
Rhabdomyolysis	19	Regeneration	51
Rotary switch	8	Training frequency	48
Symbolic assignment	54	What is EMS training?	49
S		What to do in emergency situations?	95
Service	16	When to interrupt the training?	94
Setting up a managed device	66	Training plan	52
Setting up the master device	65	Body forming	93
		Fitness	90

Invigoration back	91	USB	
Sports performance	92	miha bodytec USB flash drive	15, 70
Test training	94	Use	17, 18
Training plan memory	63	Using USB flash drives	53
Training program		V	
Body Relax	89	Viral infections	19
Muscular Endurance	89	W	
Pulse acclimatization	88	Weight	36
Strengthening	88	Who is allowed to do what?	20
Test training	89	Wi-Fi	36, 70
Training session pause	96	Wireless Mesh Network	36
Transponder card	13	work station m.ove	12
Changing data	100	Wounds	19
Deleting	98		
First use	99		
Formatting	98		
Programming	98		
Selecting language settings for the athletes	101		
Setting up synchronized start	100		
Training plans	102		
Using	100		
Transport	109		
Transporting the device	109		
travel station m.ove	11		
Tuberculosis	19		
Tumor diseases	19		
U			
Under the influence of alcohol	19		
Under the influence of drugs	19		
Under the influence of narcotics	19		
Under the influence of painkillers	19		
Undergarments	72		
Correct fit	72		
Pants	15		
Top	15		
Update			
How do I receive an update file?	69		
Importing	70		