# MD System 1.0





# $\mu$ dialysis

# Instructions for use MD System

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#### MD System

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The manufacturer will be considered responsible for the device with respect to safety, reliability and function only, if:

- the device is used in conformity with these instructions for use.
- the installation, the extensions, the re-adjustments, the modifications and the repairs have been carried out by the manufacturer or authorized representatives.
- M Dialysis AB is only responsible for replacement of defective parts, not wear of parts.
- M Dialysis AB is not responsible for any personal injury or any damage resulting from incorrect use of the MD System.

Note: At the time of printing this manual correctly describes the device and its functions. However, as modifications may have been carried out since the production of this manual, the system package may contain one or more amendments to this manual. This manual including any amendments must be thoroughly read before using the device.

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### 1. Safety-related information

Prior to using the MD System, attentively read the entire instructions for use and become familiar with the operation of the device.

Due to the lack of a limit value monitoring function, the MD System may not be used in standalone mode for intensive medical patient monitoring.

A therapy decision may not be made based solely on the information provided by MD System but must be accompanied by another method, e.g. the ICP measurement, pbtiO2 and other.

In order to ensure safety, reliability and performance of the system, the following notes shall have to be observed carefully:

- The MD System must be operated by qualified staff only. (See intended user 2.1.1.)
- Prior to using the MD System, you must be completely familiar with the operation of the system.
- If two or more devices with separate main power connections are used on the same patient, the device leakage current adds up which may cause a potential hazard. In this case, use shall be permitted only, if it has been secured that patient and operator are safe, and that the requirements of standard IEC 60601-1-1 are fulfilled.
- Prior to using the MD System, the correct condition of the device and any accessories shall have to be checked. Check the plausibility of the readings before using the device for monitoring. The device and the accessories must not be damaged or soiled; otherwise the device must not be used.
- Do not use the device if it reveals obvious signs of a mal function; in this case forward the unit to the after-sales service of the manufacturer.
- For service-related questions, contact the manufacturer directly. If required, the manufacturer will direct you to a local authorized service partner.
- The MD System should be positioned and/or fastened in a manner by which the device is secure and cannot be damaged or cause hazards. Care must be taken to prevent cables attached to the MD System from tipping or moving the system.
- After use, clean, disinfect and/or sterilize the accessories according to the instructions for use. If the accessory has been provided with separate instructions for use, the instructions rendered there shall be applicable.
- No liquids or fluids shall enter the device. If this should happen nonetheless, first remove the main adapter from the port and switch off the system using the power switch. Provide the device to the after-sales service department for inspection. Subsequently a safety-related check is necessary.

- The MD System has been designed in conformity with IEC 60601-1. It is a class II product with an internal power source, an external main power adapter and has been allocated to class IIa (MPG).
- The MD System must not be used in conjunction with the MRT.
- To avoid leakage currents the following instructions should be observed for the installation of the system:
  - Moveable multiple sockets must not lay on the floor.
  - Additional moveable multiple sockets or extension cables must not be connected directly to the device.
  - Devices, which are not part of the system, must not be connected.
  - The moveable multiple sockets have to be appropriate for the load of the system.

Warning! MD System is an adjunct analytical device which must always be used together with other means of assessing the patient's clinical condition. Diagnosis and change in therapy must not be based solely on data from MD System.

**Warning**! The MD System is intended to be used in hospital environments except:

- during magnetic resonance imaging (MRI).
- during hyperbaric oxygen therapy.
- in rooms warmer than +38°C

# 2. Description of the device

#### 2.1. Intended use

MD System is a microdialysis analysing system, which is used in conjunction only with M Dialysis catheters, performing continuous monitoring of Glucose, Lactate and Pyruvate levels in microdialysates.

#### 2.1.1. Intended user

The MD System is intended to be used by experienced health care professionals

#### 2.1.2. Intended purpose

The MD System is a monitoring device for measuring and displaying the levels of concentration for Glucose, Lactate and Pyruvate in tissue or blood. This information support clinical decisions or can be of use for clinical research. The device provides the information only when connected to the MD Catheter.

#### 2.1.3. Intended use environment

The MD System is intended to be used in a clinical environment by experienced healthcare professionals and also in clinical research environment. The product is not intended for outside hospital use such as helicopters or ambulances. The MD System is not intended for home use. Qualified physicians experienced in the field of application must always assess whether the use of MD System is appropriate for a specific patient or not.

#### 2.2. Indications for use

#### 2.2.1. Condition

The MD System is indicated when the clinician decides there is a need to measure and display the metabolic changes via analyses of Glucose, Lactate and Pyruvate.

#### 2.2.2. Part of Body or Type of Tissue with Which the Device Interacts

The MD Catheter is in vivo. MD Amplifier can come in contact with the body of the patient.

#### 2.2.3. Frequency of Use

The MD System is indicated for use when prescribed by a clinician.

#### 2.2.4. Physiological Purpose

The MD System is indicated when the purpose is to gain information for treatment, to assess adequacy of treatment, or to rule out causes of symptoms.

#### 2.2.5. Patient Population

Adult, pediatric, and neonatal non-ambulatory patients. The user should refer to each Microdialysis catheter's instructions for use.

#### 2.3. Terminology and definitions

**Applied parts** – MD Amplifier and MD Sensor are applied parts, placed close to the patient. MD Unit is regarded as applied part with a contact duration of 10s < t < 1min.

**Biosensor** – Coated electrodes which generate the raw current from the different analytes in the microdialysate. The biosensor is included in the MD Sensor.

**MD Unit** – Central unit containing process control and analyzing unit

**MD Amplifier** – Non-consumable part (multiple use) placed close to the patient. Contains amplifier and A/D converter. Amplifies the raw signal from the biosensor and sends digital values to the MD Unit

**MD Cartridge** – Part containing filled syringes connected to the MD Sensor. The MD Cartridge is, prepared with syringes, inserted in the MD Unit and the syringes are actuated by the syringe motors. MD Cartridge is for multiple use.

**MD Sensor** – Part to which the patient Microdialysis catheter outflow is connected to. Contains a biosensor and connective tubing to the MD Cartridge. MD Sensor is a single use part.

**Warning!** Be sure to check that MD Sensor package is undamaged prior use.

**106 Syringe** – Special syringe for Microdialysis supplied by M Dialysis AB. 106 Syringe is for single use.

**Syringe Orange** – Special syringe for calibration of MD System, supplied by M Dialysis. Syringe orange is for single use.

**Perfusion Fluid CNS** – Perfusion Fluid is a sterile, isotonic fluid especially developed for brain Microdialysis.

**Perfusion Fluid T1** – Perfusion Fluid is a sterile, isotonic fluid especially developed for Microdialysis in peripheral tissues.

Sensor docking – Where the MD Sensor is connected to the MD Amplifier.

**MD Catheter** - Microdialysis catheter supplied by M Dialysis. For example, the 70 Brain Microdialysis Catheter, 63 Microdialysis Catheter and others. For handling of the MD Catheter read the MD Catheter IFU.

**Catheter Extension** - Sterile disposable extension tubing with luer lock connections connected between the 106 Syringe and the Microdialysis Catheter. The Catheter Extension is for single use.

#### 2.4. System overview

MD System consists of:

- 1.MD Unit, consisting of a power supply unit, syringe pumps and a computer.
- 2.MD Sensor, consisting of a Biosensor and tubing
- 3.MD Cartridge, consisting of an interface for the MD Unit and a syringe retainer.
- 4.MD Amplifier, consisting of an amplifier with cable.
- 5.MD Catheter, consisting of a variety of Microdialysis catheters supplied by M Dialysis, for example the 70 Microdialysis Brain Catheter, 63 Microdialysis Catheter and others.
- 6.MD Monitor, consisting of a pre-configured Windows PC with integrated monitor and a power supply unit.



#### 2.5. System block diagram



MD System has five main functions:

- A controlled flow of perfusion liquid to the MD Catheter.
- A controlled flow for Calibration fluid to perform recurring calibrations.
- A Biosensor that converts the content in the microdialysate to an electric current.
- An amplifier that amplifies the signal from the biosensor.
- Analysis software that displays the signal from the biosensor.

To facilitate the perfusion flow of 0.3  $\mu$ l/min – 2.0  $\mu$ l/min, one standard 106 Syringe (7c) is actuated by a Piezo motor with sufficient resolution. The 106 syringe supplies the MD catheter (5) with perfusion fluid that allows molecules to diffuse over the catheter dialysis membrane from the surrounding tissue, creating the so called microdialysate or dialysate.

The analytes pass through the Biosensor in the MD Sensor (2) where enzyme coated electrodes each generate a current that is proportional to the number of endogenous molecules of the specific analyte. After the Biosensor, the dialysate continues to the housing tube as waste (7a). The MD Amplifier (4) amplifies and converts the analogue signal to a digital signal which is processed in the MD Unit (1) and displayed on the MD Monitor (6).

Recurring calibrations where Calibration Fluid is pumped through the MD Sensor, instead of the regular microdialysis perfusate, will ensure that any drift in the analyte signals is compensated for.

#### 2.6. MD Unit parts





#### 2.7. **MD Unit power supply**

The MD Unit comes with a power supply, 18V DC. Manufacturer: Powerbox Model code: EXM 80 5120



Warning! Use only specified power supply identified on the power inlet. If any suspicion of damage or other fault of power supply replace with new. If in doubt consult M Dialysis AB.

#### 2.8. **MD Sensor**

MD Sensors are for single use only and should be disposed, according to the hospital routines, as biohazardous waste after used.



Warning! Don't use if package is broken or damaged.

Warning! The MD Sensor is single use only. MD Sensor Catheter Tubing to MD Connection cartridge (Microvial holder)

#### 2.9. MD Amplifier

The MD Amplifier can be reused for a period of up to 1 year.



Warning! Only clean housing with disinfectant (70% ethanol or equivalent). Do not clean or touch MD Sensor connection socket. Do not clean using sharp objects.

#### 2.10. MD Cartridge



#### 2.11. Battery back-up

The MD Unit contains a battery back-up if MD System should be disconnected from the mains power supply. Also, a safety feature in case of mains power failure.

#### 2.12. MD Unit holder

The MD Holder is an accessory for mounting the MD Unit on an ICU-rail or pole.



## 3. Operation of MD System

#### 3.1. General

MD Unit is controlled by MD Software. The user operates the MD Unit using the touch screen of the connected MD Monitor. The user shall <u>read this manual</u> before operating the instrument.

#### 3.2. Manual and Help (On-line manual)

This manual describes the most important MD System functions and how to operate MD System.

#### 3.3. Unpacking and installation

Trained clinical staff or personnel designated by M Dialysis must perform all installation of the MD Unit. The MD Unit should be placed on a table or rack or attached to any of the ICU-poles used for instruments at the intensive care unit using the MD Unit Holder.

Note! All communication ports and on/off switch are located on the bottom side of the MD Unit, make sure that these are easily accessible after installation. For ranges of acceptable temperature and humidity, see the technical specifications.

Warning! Make sure to use a trolley or table with sufficient stability for placement of the MD Unit. When mounting the MD unit shall only be done to a part with sufficient load capacity.

#### 3.4. EN 60601 requirements

To fulfill the EN 60601 requirements for leakage current and electrical separation the installation must satisfy the EN 60601-1-1 standard.

**Warning!** Multiple socket-outlet or extension cord shall not be connected to the MD System.

#### 3.5. Handling of MD Sensor

The MD Sensors are single use only and should be stored in a fridge, +2 - +8 °C. Before usage they should be taken out in room temperature about 20 minutes prior connecting to MD System.

#### 3.6. Edit settings in the MD Software

Editing MD System settings should only be performed by trained personnel or designated by M Dialysis.

#### 3.7. Exchange of MD Sensor

**Warning!** Exchange of MD Sensor must be done under clean conditions, use gloves.

**Warning!** MD Sensor and its parts, syringes and extension tubing, are for single use and should not be reused on other patients.

To exchange the MD Sensor first follow the procedure for temporary disconnecting of patient section 3.18 Prepare a new MD Sensor according to 3.16 (the MD Cartridge itself may be reused). The orange calibration fluid syringe, is usually not depleted but cannot be reused. Neither can remaining perfusion fluid be reused. Reconnect the MD Sensor and start MD System as described in section 4.7.2.

#### 3.8. Exchange of perfusion fluid and Calibration fluid

Exchange of fluids may be done in the same manner as described in section 3.7 but shifting fluids instead of MD Sensor.

#### 3.9. Disconnecting and storage of the system

After ending the microdialysis process according to section 3.19, reconnect the empty MD Cartridge to the MD Unit, disconnect the power adaptor and the MD Amplifier. The unit is now ready for storage.

#### 3.10. Calibration

The MD Unit will automatically perform calibration every 6 hours. This can be postponed 30 minutes.

Manual calibration can be performed from the settings menu.

#### 3.11. Indicator lights MD Unit

#### 3.11.1. MD System status

MD Unit status is indicated by LEDs on the MD Unit in the following way:

Steady green	MD Unit ready for patient, standby
Flashing green	MD Unit running
Steady blue	System startup
Steady orange	Measurement or
	technical error

Warning! MD System does not have an alarm system compliant with IEC 60601-1-8.

#### 3.11.2. Battery status

Battery status is indicated by LEDs on the MD Unit in the following way:

Steady green	>1h left, charging	
Flashing green	>1h left, not charging	
Steady blue	<1h left, charging	
Flashing blue	<1h left, not charging	
Steady orange	<15min left, charging	
Flashing orange	<15min left, not charging	

#### 3.12. Installing the MD Unit holder

The MD Unit holder is attached to the bottom side of the MD Unit.

#### 3.12.1. Installation on rail

The MD Unit holder is suitable for EU design rail (25x10mm) and shall only be attached to a rail with sufficient load capacity.

Note! If the knob is mounted for pole mounting, see below, unscrew the knob and reattach it as shown in the picture. Make sure that the clamp is attached in the right position on the rail.

**Warning!** If attached to other types of rails be sure to check that the MD Unit is firmly attached. If in doubt, do not attach to the rail.



Attach the MD Unit to the rail. Lock knob firmly but tighten only by hand.



#### 3.12.2. Installation on ICU-pole, diameter 24-30mm.

Note! If the knob is mounted for ICU-rail mounting, see above, unscrew the knob and reattach it as shown in the picture below.



Attach the MD Unit to pole. Lock knob firmly but tighten only by hand. Note! the MD Unit shall only be attached to a pole with sufficient load capacity.



#### 3.13. Installing the MD Monitor

The MD Monitor has a standard VESA mount interface for installation in the vicinity of the patient. For more information see MD Monitor Manual. This product is certified as a Class I product per the MDD and IEC 60601-1. MD Monitor must only be mounted to a part with sufficient load capacity.



There is also an MD Monitor Arm available as an accessory. For MD Monitor arm installation see MD Monitor Arm manual.

**Warning!** The Monitor arm should never be mounted on a mobile pole





The power adaptor is connected to (9) DC-in, MD Monitor cable is connected to (2) and USB memory is connected to (6).

#### 3.14. Turn on the MD Unit

1. Connect the power adaptor to mains and the MD Unit.



2. Connect the MD Monitor to the MD Unit using the Ethernet cable.



3. Turn on the MD Unit by pushing the Power-ON/OFF switch underneath the unit.



#### 3.15. Turn on the MD Monitor

Turn on the MD Monitor by pushing the Power-ON/OFF switch on the MD Monitor. MD Cartridge will be ejected.



Hotkey and LED definition at front panel

#### 3.16. Start-up process and connecting the MD System

Check flow setting, default 0.5  $\mu$ L/min.

Tap the START button on the MD Monitor's touch screen. Instructions of how to connect the complete MD System and the full starting procedure will then follow on the MD Monitor, follow these instructions. The instructions are also described below:



After each step is completed the circle to the top right will turn green. There is also an option to skip a step when applicable.

Note! The on-screen pictures are for guidance; always follow the instructions in this manual.

**Warning!** Be sure to check for damages on MD System prior to each patient session.

#### 3.16.1. Pre-use check

**Pre-use check** is described in step 1–5 on the Monitor Screen. Press "SKIP" to go directly to step 6, "Enter the patient ID".

Note! Pre-use check should be done when starting a new patient



#### 3.16.2. Preparation of MD Sensor and MD Cartridge

Note! The preparation of the MD Cartridge and sensor should be conducted under clean conditions. The preparation can be done prior to starting MD System.

The preparation requires the following items,

- 1 Bottle of Calibration Fluid
- 1 Syringe Orange (Orange colored piston)
- 1 single use MD Sensor
- 1 MD Cartridge
- 1 106 Syringe (non-colored piston)
- 1 pcs of Catheter Extension
- 1 Perfusion Fluid (see Microdialysis catheter IFU what perfusion fluid to use)
- 1 Microdialysis Catheter

**Warning!** It is important that the color coding of syringes and Luer-connectors is followed throughout the whole process from filling syringes to connecting and placing the syringes in the MD Cartridge.

**I** Warning! Be sure to check that MD Sensor package is undamaged prior use.

1. Penetrate the membrane of the Calibration fluid vial with the cannula of the orange piston syringe and fill the syringe with 2.5 ml of Calibration fluid. Remove air bubbles.



2. Connect orange syringe to orange Luer connector on the MD Sensor.



3. Fill the 106 Syringe (non-colored piston) with sterile 2.5 ml Perfusion Fluid. Make sure it is room tempered and that all air bubbles are removed.



Warning! Using wrong fluids could impair or give wrong analysis result

4. Connect the Catheter Extension to the 106 Syringe (non-colored piston)

**Warning!** Do not remove the sterile cap from the Catheter Extension.

Warning! Ensure that all luer-lock connections are firmly tightened.



- 5. Place the Syringe Orange in the first (1) slot in the cartridge. The slots are color coded.
- 6. Place the 106 Syringe (non-colored piston) in the third (3) slot in the cartridge.



Warning! Ensure that all luer-lock connections are firmly tightened.

7. Connect the MD Sensor to the MD Amplifier.



8. Connect the MD Amplifier cable to the MD Unit.



9. Connect the MD Catheter's inlet to the MD Cartridge via the Catheter Extension.

Warning! Confirm that the sterile caps on the extension tubing and MD Catheter have <u>not</u> been removed until this step.

**Warning!** Ensure that there is enough slack on the Catheter Extension between the MD Cartridge and the MD Catheter.



Warning! Pay close attention to the placement and handling of the MD Cartridge when not inserted to the MD Unit.

Warning! if the catheter is accidently extracted from the patient, reinsertion must be done with a new catheter.

10. Insert the MD Cartridge in the MD Unit. Align the MD Cartridge in the center and insert it straight into the cavity.



- 11. Press start and follow the instructions on the MD Monitor to proceed.
- 12. The system will now automatically start to calibrate and warm up. This procedure could last up to 60 minutes. After the calibration process, the first values will be obtained. The data will be analyzed and presented on the MD Monitor.



**Warning!** Pay close attention to tubing and cables between the patient and MD Unit. Do not to squeeze the tubing.

Place the MD Amplifier near the catheter insertion point. Fix with bandage or other means. The distance is limited by the length of the catheter tubing.

**Note!** Place MD Unit at a fixed level during measurements and MD Sensor shall always be placed lower than the catheter membrane **Note!** Make sure that the MD Amplifier is not in direct contact with the patient (by padding) to avoid decubitus.

**Note!** Make sure that the MD Unit is at similar height level to the MD Catheter.

Note! Make sure that the MD Amplifier is properly secured to the patient.



**Warning**! Inspect skin condition beneath MD Amplifier mounting regularly with respect to infection and avoid decubitus. MD Amplifier shall not be continuously attached for a period longer than 5 days.

#### 13. Flush check

Make sure that there is perfusion fluid in the Microvial holder on the outlet of the MD Catheter prior to connecting it to MD Sensor.

**Note!** The MD Sensor should be connected to the MD Amplifier PRIOR to connecting the MD Catheter to the MD Sensor.

Connect the MD Catheter's outlet to the MD Sensor, using the Microvial holder.



#### 3.17. Using the GUI: Graphic User Interface

#### 3.17.1. Home screen



MD Monitor graphic user interface:

Home – Will display the main panel
View – Display of current measured microdialysis in graphs and figures
Settings – General settings such as date, time and language
Service – A password protected panel for MD System service
Patient data – Management of patient data
Start/Pause/Stop – For starting a new patient/ temporary pause in microdialysis/ termination of patient session
Battery icon – Current battery status
MD System icon – Status of MD System
Logotype – Displays system software version

#### 3.17.2. System status display on the home screen

Tap the icons, current system status will be displayed.



MD Unit connected to mains. Battery charging.

MD Unit disconnected from mains. Back-up battery in use.

Remaining perfusion fluid in the cartridge and the set flow rate for the Microdialysis.



Current software version.

#### 3.17.3. Display of analyte values

To display graphs of the microdialysis values tap the view symbol on the home screen.






## 3.17.4. Change system settings

The Settings menu is not password protected.



The Settings menu handles System settings.

## 3.17.4.1. System settings menu



Settings menu

- Date and time
- Language
- Flow rate

The analyte is transported from the catheter to the Biosensor which will create a time delay. A higher flow rate will reduce the lag time but will shorten the total monitoring time. A flow of 0.3  $\mu$ l/min will give a monitoring time of approx. 5 days whilst a flow of 2.0  $\mu$ l/min gives a monitoring time of approx. 18 hours before the 106 Syringe in the MD Cartridge needs to be replaced.

- Graph range setting
- Reset Default Settings

#### 3.17.5. Service menu

The Service menu is password protected and should only be accessed by M Dialysis authorized personnel. For further information, contact M Dialysis or your local representative.

### 3.17.6. Patient data

The patient data menu is for managing the analysis data that has been saved in the system. MD System only saves the current patient session, after each patient the data should be exported. Data saved in MD System is erased when patient session is terminated.



3.17.6.1.

Edit patient data:

	PATIENT DATA	
	I	Lac <b>-0,0</b>
John		0,0
Last Name		
Doe		Glu
Catheter Location		0,0
Brain		0,0
Comments		
124 characters remaining		
Example comment.		Pyr <b>-0,0</b>
Energie control .	h.	
	Save	-0,0
	Save	
		L/P
		500,0
	I	
VIEW SETTINGS SERVICE PA	TIENT DATA	🥢 🗲 📈 dialysis

#### 3.17.6.2. Export patient data

Data can be exported by plugging in a portable USB memory to the MD Monitor using the ports in the picture below.



To export data, tap the [download or export the patient data as csv], found under Patient Data.



The save dialog will open in a separate window. Save the file to the USB memory.

## 3.17.7. Stop or pause microdialysis session

- 1. To stop the session, tap STOP (1).
- To temporary pause the analysis and reconnect same patient later, use the Pause option (3) and confirm in next window by pressing "Yes" Note! Current patient's data remains stored in the system during pause.
- 3. To stop microdialysis and change patient, tap Stop (2) and confirm in next window by pressing "Yes".
- 4. To abort stop and continue microdialysis, tap Cancel (4).



Notice		Х
Are you sure?		
	Yes	No

## 3.18. Temporary disconnection of the patient (Pause session)

Note! Current patient's data remains stored in the system during pause.

## 3.18.1. Disconnecting the patient

- 1. After tapping STOP (1) tap the pause button (3) on the MD Monitor and follow the instructions on screen.
- 2. Wait until MD Cartridge is ejected.

Warning! Do not to squeeze the tubing when MD Cartridge is ejected from MD Unit.



3. Disconnect the Microdialysis catheter vial holder from the MD Sensor and connect a microvial to the vial holder.

Sensor Historic Strike via la via connector Remove the 106 syringe from the MD Cartridge Remove the WD Amplifier with the MD Sensor from the patient Remove the vial and connect the MD Sensor for the MD Sensor Reinser 106 syringe in the MD Cartridge Reduct the MD Cartridge in the MD Cartridge Reduct the MD Cartridge in the MD Cartridge Reduct the MD Sensor with the MD Amplifier to the patient NEXT		Retracting motors and ejecting cartridge Disconnect the catheter vial connector from th	Lac N/A
Remove MD Cartridge Remove the 106 syringe from the MD Cartridge Remove the WD Amplifier with the MD Sensor from the patient Remove the vial and connect the MD Catheter's outlet to the MD Sensor Reinsert 106 syringe in the MD Cartridge Related the MD Cartridge in the MD Amplifier to the patient			
Remove the 106 Syringe from the MD Catridge Remove the WD Amplifier with the MD Sensor from the MD Sensor Reinsert 106 Syringe in the MD Catridge Reload the MD Catridge in the MD Unit Reattach the MD Sensor with the MD Amplifier to the patient			
Remove the MD Amplifier with the MD Sensor from the patient Remove the vial and connect the MD Catheter's outlet to the MD Sensor Reinsert 106 syringe in the MD Catridge Reload the MD Catridge in the MD Unit Reattach the MD Sensor with the MD Amplifier to the patient		-	
Reload the MD Cartridge in the MD Unit  Reattach the MD Sensor with the MD Amplifier to the patient  VP  VA			ntient N/A
Reload the MD Cartridge in the MD Unit  Reattach the MD Sensor with the MD Amplifier to the patient  VP  VA		Remove the vial and connect the MD Catheter's outlet to	
Reload the MD Cartridge in the MD Unit  Reattach the MD Sensor with the MD Amplifier to the patient  VP  VA		Reinsert 106 syringe in the MD Cartridge	
		Reload the MD Cartridge in the MD Unit	Pvr
		Reattach the MD Sensor with the MD Amplifier to the patie	
		_	L/P N/A

- **Warning!** A microvial must be connected to the MD Catheter's outlet.
- 4. Remove the MD Cartridge from the MD Unit. Make sure that the MD Cartridge has been ejected by the MD Unit prior to removal.



5. Remove the 106 Syringe with perfusion fluid from the MD Cartridge.

Warning! During e.g MR investigation the 106 Syringe must be removed from the MD Cartridge but it must <u>not</u> be disconnected from the Catheter Extension.



6. Remove the MD Amplifier with the MD Sensor from the patient. Alternatively, the MD Sensor is disconnected from the MD Amplifier and the MD Amplifier remains attached to the patient.



7. The patient is now disconnected from MD System.



## 3.18.2. Reconnecting the patient

1. Remove the vial and connect the MD catheter connector to the MD Sensor.



2. Reinsert the 106 Syringe with perfusion fluid into the MD Cartridge.



3. Insert the MD Cartridge in the MD Unit. Align the MD Cartridge in the center and insert it straight into the cavity.



4. Reconnect the MD Sensor to the Amplifier, or if both were removed from the patient, reattach the MD Sensor with the MD Amplifier to the patient.



5. Press "RUN" on the MD Monitor and MD System will continue with the Microdialysis monitoring.

### 3.19. Ending the Microdialysis process

### 3.19.1. Shutting down the system

- 1. Press Stop on the MD Monitor and confirm.
- 2. Follow the instructions on the MD monitor on how to save patient data and shut down the system.

### 3.19.2. Disconnecting the patient

1. Disconnect the MD catheter from the MD Sensor and the extension tube.



2. Remove MD Sensor and MD Amplifier from patient.



### 3.19.3. Disposal of the MD Sensor

- 1. The MD Unit will eject the cartridge at shut down, after which it is possible to remove the MD Cartridge.
- 2. Remove the MD Cartridge from the MD Unit. Make sure that the MD Cartridge has been ejected by the MD Unit prior to removal.



3. Remove all syringes from the MD Cartridge



4. The MD Sensor should be taken care of as biological hazard waste.



5. The MD Cartridge can be reused. *Warning! Do not use a damaged MD Cartridge, replace a damaged cartridge immediately.* 

# 4. Routine maintenance

## 4.1. Cleaning the instrument

Wipe the outside of MD System components, except MD Sensor, with disinfectant (70% ethanol or equivalent).

Note! Sodium hypochlorite shall not be used on MD Amplifier.

Sodium hypochlorite may cause miss colorations on aluminum surfaces and should only be used to a limited extent.

**Caution!** Do not to use sharp objects for cleaning device or accessories

**Caution!** Do not to use excessive amounts of water or other liquids during cleaning

# 5. Troubleshooting

Error:	Message on MD Monitor; Waiting for a connection
Action:	<ul> <li>Check that MD Unit is on and is running</li> <li>Check that ethernet cable between MD Unit and MD Monitor is properly connected and without visual damage</li> <li>Restart MD Monitor</li> <li>Restart MD Unit</li> <li>If possible, try using another ethernet cable</li> <li>If problem remains, please contact M Dialysis Service Center or your local distributor</li> </ul>
Error:	Message on MD Monitor; MD Sensor is not connected
Action:	<ul> <li>Check that MD Sensor is properly connected to the MD Amplifier</li> <li>Check that the cable between MD Unit and MD Amplifier is properly connected and without visual damage.</li> <li>Check that MD Sensor is without visual damage</li> <li>Check that MD Amplifier is without visual damage</li> <li>Check that MD Amplifier is without visual damage</li> <li>If possible, try connecting another MD Sensor</li> <li>If possible, try using another MD Amplifier</li> <li>If problem remains, please contact M Dialysis Service Center or your local distributor</li> </ul>
Error:	No message, system freeze
Action:	<ul> <li>Restart MD Monitor</li> <li>Restart MD Unit</li> <li>If problem remains, please contact M Dialysis Service Center or your local distributor</li> </ul>

# 6. Classification and Regulations

## 6.1. In accordance with the MDD (Medical Device Directory)

In accordance with the MDD, Article 9, MD System is classified as a Class IIa medical device.

### 6.2. In accordance with EN 60601-1:2006

#### 6.2.1. Type of protection against electric shock

Complete system except MD Monitor is Class II equipment. MD Monitor is a class I equipment.

#### 6.2.2. Degree of protection against electric shock

Type CF equipment

### 6.2.3. Degree of protection against harmful ingress of water

IPX1 – Only with MD Cartridge mounted and MD Unit in upright position.

### 6.2.4. Mode of operation

Continuous operation

## 6.2.5. Mechanical strength

Portable equipment

### 6.2.6. Suitability for use in an oxygen rich environment

Not suitable for use in an oxygen rich environment

### 6.3. Regulations

MD unit meets international standard EN 60601 class IIa.

The MD unit bears the CE label in accordance with the provisions of the European Directives for EU Directive 93/42/EC (MDD), EU Directive (2014/30/EU) (EMC), EU Directive 2012/19/EC (WEEE II) and EU Directive 2011/65/EC (RoHS II)

# 7. Technical specifications

### 7.1. Technical data

Voltage: Power consumption: Compliance: Insulation: Dimensions W x D x H: Weight: Principle: Samples:	100 - 240, VAC 50 / 60 Hz max 40 VA Power adaptor complies with 60601-1 MD Unit power adaptor is double isolated 220 mm x 220 mm x 332 mm 5 kg Electrochemical biosensor Continuous on-line measurement of Microdialysate
Monitor cable: Recovery time after exposure to	Standard ethernet, shielded (Cat 5e or later)
defibrillation voltage:	5 minutes

**Warning!** MD Unit is heavier than it may appear. Exercise caution upon moving or lifting MD Unit.

MD System measures Glucose, Lactate and Pyruvate within the following intervals and with the following accuracies: Glucose 0.2–15 mM,  $\pm$  30 % or 0.1 mM whichever is greatest Lactate 1–10 mM,  $\pm$  30 % or 2 mM whichever is greatest Pyruvate 10–150 µM,  $\pm$  30 % or 10 µM whichever is greatest Calculated parameters: The Lactate-Pyruvate ratio

M Dialysis AB reserves the right to make changes in the specifications without prior notice.

### 7.2. Duration of contact for applied parts and accessible parts

MD Amplifier and MD Sensor are applied parts and have continuous contact with the patient for up to 5 days. MD Unit is regarded as applied part with a contact duration of 10 s < t < 1 min.

Other components are regarded as accessible parts with a duration of contact 10 s < t < 1 min.

### 7.3. Expected service life

Expected service life; MD Unit 7 years MD Amplifier 1 year MD Cartridge 1 year

### 7.4. Operating environment

- The MD Unit is manufactured for indoor use and should be placed in a draught free place and not in direct sunlight.
- No radio transmitters, cellular phones or other wireless communication devices should be used in the vicinity of the MD Unit.

- The MD Unit should not be submitted to higher levels of disturbance as specified in EN 60601-1-2 and EN 61010.
- MD System should not be used with flammable anesthetics category AP or APG.

MD System excluding the MD Sensor shall be kept under the following environmental condition:

#### Operating

Temperature; +15 °C to +30 °C Humidity; 10% to 90% (non-condensing) Pressure; 700 hPa to 1060 hPa **Storage** Temperature; -10 °C to +60 °C Humidity; 10% to 90% (non-condensing) Pressure; 500 hPa to 1060 hPa **Transport** Temperature; -10 °C to +60 °C Humidity; 10% to 90% (non-condensing)

Pressure: 500 hPa to 1060 hPa

MD Sensor shall be kept under the following environmental condition:

#### Operating

Temperature; +15 °C to +30 °C Humidity; 10% to 90% (non-condensing) Pressure; 700 hPa to 1060 hPa **Storage and transport** Temperature; +2 °C to +8 °C Humidity; 10% to 90% (non-condensing) Pressure; 500 hPa to 1060 hPa

**Warning!** Do not operate/store/transport MD System and MD Sensor outside the above noted environmental conditions.

#### 7.5. MD System materials and ingredients patient or operator is exposed to

MD Amplifier:	Medical grade PC, Poly Carbonate
MD Sensor cable:	PVC, Polyvinyl Chloride
MD Sensor:	Medical grade PC, Poly Carbonate
MD Sensor tubing:	Medical grade TPE, Thermo Plastic Elastomer
Perfusion fluid:	Perfusion Fluid is a non-toxic, sterile, isotonic fluid
Calibration fluid:	Fluid containing defined concentrations of Glucose, Lactate
	and Pyruvate to support sensor calibrations.

#### 7.6. Disturbances

Devices transmitting on radio frequencies can affect the performance of MD System if used in close proximity of MD System.

# 8. Consumables and Options

**Warning!** Only consumables from M Dialysis shall be used.

For up to date information, please visit our website: http://www.mdialysis.com

# 9. Spare Parts and accessories

**Warning!** Only Spare Parts and accessories from M Dialysis shall be used.

For up to date information on approved spare parts and accessories, please visit our website: <u>http://www.mdialysis.com</u>

8050187 MD System Sensor LPG Kit

- 8070203 MD Sensor LPG
- 8072001 Catheter Extension

8050181 MD System Syringe Kit, Brain Tissue

- One 106 Syringe from **8010191** 106 Syringe 20/pkg
- One Syringe Orange from 8070081 Syringe Orange 20/pkg
- One bottle from 8072009 MD Calibrators
- One ampoule from P000151 Perfusion Fluid CNS 10x5 mL

**8050182** MD System Syringe Kit, Peripheral Tissue

- One 106 Syringe from **8010191** 106 Syringe 20/pkg
- One Syringe Orange from 8070081 Syringe Orange 20/pkg
- One bottle from 8072009 MD Calibrators
- One ampoule from P000034 Perfusion Fluid T1 10x5 mL

# 10. Liability, warranty and service

#### 10.1. Liability

M Dialysis AB is liable for the safety and reliability of its equipment only if:

- a) repair, maintenance and modifications are carried out by authorized personnel;
- b) components are replaced with M Dialysis approved spare parts;
- c) devices are used only with M Dialysis approved accessories and consumables;
- d) devices are used in accordance with M Dialysis's operating instructions.
- e) MD Unit is not repaired on site, unit should be replaced and sent to M Dialysis AB

#### 10.2. Warranty

M Dialysis AB offers a one-year warranty from the day of purchase on defective material and assembly.

The warranty does not cover damage resulting from incorrect use or maintenance, or from unauthorized software modification.

M Dialysis AB is only responsible for replacement of defect parts, not of consumables or wear parts. M Dialysis AB is not responsible for any personal injury or any damage resulting from incorrect use of the MD Unit.

### 10.3. Service

Routine service by authorized personnel shall be done with a twelve-month interval. Routine service allows an expected service life of seven (7) years.

Routine service includes:

- General inspection of the MD Unit.
- Function test.
- Software upgrade, if applicable.

A service agreement may be purchased after the warranty period has ended.

For more information, please contact your local supplier or the M Dialysis service department:

M Dialysis AB Hammarby Fabriksväg 43 SE-120 30 Stockholm Sweden E-mail: service@mdialysis.se Tel. +46 8 470 10 20 Web: http://www.mdialysis.com

### 10.4. Arrival inspection

MD System must be inspected for damage upon arrival and unpacking according to the following points:

- Check the outer packaging for damage, any visible damage must be noted to the courier upon delivery.
- Any visible damage upon unpacking must be reported to M Dialysis immediately.

### 10.5. Functional test

There is an option to perform a pre-use check in the start guide (see section 3.16.1).

# 11. Text and symbol explanation



Leven	ETL- listed product. Conforms to UL 60601-1:2003 Rev. 2006 and Certified to CSA C22.2#601.1 (R2001)
<b>CE</b> 0413	CE marked according to the Medical Device Directive (MDD 93/42/EEC), EU Directive (2014/30/EU) (EMC), EU Directive 2012/19/EC (WEEE II) and EU Directive 2011/65/EC (RoHS II)
STERILE A	Sterile Using Aseptic Processing Techniques
STERILE R	Sterilized Using Irradiation
STERILE EO	Sterilized Using Ethylene Oxide

# 12. Disposal of product and accessories

Discontinued MD Sensors shall be handled according to the hospital routines for biohazard material.



Disposal of the MD Unit, or parts thereof, shall be in accordance with local regulations.