# Monidrop® W infusion monitor / General

1. **General**
   - 1.1 General
   - 1.2 Attaching Monidrop to a drip chamber
     - 1.3.1 Purpose of use
     - 1.3.2 Use
     - 1.3.3 Blood transfusion
     - 1.3.4 Other infusion factors
     - 1.3.5 Safety standards

2. **Symbols / Menu structure**

3. **Using the device**
   - 3.1 Activation and start of infusion
     - 3.1.1 Preparations
     - 3.1.2 Starting the device
     - 3.1.3 Starting infusion monitoring
   - 3.2 Starting a new treatment
   - 3.3 Ending an infusion treatment
   - 3.4 Shutting down the device

4. **Notifications**
   - 4.1 Notifications
     - 4.1.1 Infusion rate is less than 6 ml/h
     - 4.1.2 Battery low
     - 4.1.3 Infusion rate is over 1200 ml/h
     - 4.1.4 Self-test program and forced restart
   - 4.2 Managing audio notifications

5. **Battery charge and maintenance**

6. **Cleaning**

7. **Warranty / Inspection and maintenance / Training**

8. **Content of the delivery and inspecting the content / Recycling**

9. **Technical specifications**

10. **Operating instructions for accessories**
1 GENERAL

1 Monidrop® W infusion monitor / General

1.1 General

Green light: Infusion in progress

Yellow light: Notification

Latch, press to open

Brackets for various drip chambers.

Screen

Charging port is located at the bottom of the device.

Power button

Drip size indication

Notification button

Drip detector
1.2 Attaching Monidrop to a drip chamber

1. Open the locking mechanism by pressing the locking lever and open the latch.

2. Put the drip chamber into place as shown in the figures.

Note the correct positioning of different types of drip chamber models! More detailed instructions on page 6.

3. Close the latch by returning it to the locking position.

Note! If the lock feels too tight, check the correct position of the drip chamber.
Correct positions of drip chambers:

Figure 1. Drip chamber with a collar

Figure 2. Drip chamber without a collar

Place the drip chamber in the Monidrop® W device so that the air intake cap points forwards or backwards. If the drip chamber has a collar, guide the edge of the collar to the lower slot (Figure 1). If the drip chamber does not have a collar, guide the drip chamber to the upper slot (Figure 2).
1.3 Patient safety

1.3.1 Purpose of use

The Monidrop® W infusion monitor is intended for the periodic or continuous measurement of parenteral fluids via an intravenous route. The Monidrop® W infusion monitor is used for dosage measurements in intravenous therapy. The device DOES NOT set the IV dosage. Instead, the dosage is set using the roller clamp of the infusion set. The device is intended for use by healthcare professionals at hospitals and health care centres and in home care.

The medical staff must determine the suitability of the device for the intended purpose of use on the basis of device properties and technical specifications.

Monidrop® W DOES NOT ADJUST the infusion rate. ALWAYS use a roller clamp to adjust the infusion rate.

1.3.2 Use

- Users of the Monidrop® W infusion monitor must read this operating manual before using the device. A Monidor salesperson or another authorised person must provide the first guidance of use.
- Make sure that the device is firmly in place during its use.
- Before use, check the device and its fastening mechanisms in case of damage or any missing parts.
- Do not fully open the roller clamp or so that a continuous stream drips into the drip chamber.
- The device is intended for use indoors in a stable operating environment. The device is not suitable for use on vehicles.
- The use of the device is not recommended for children under 10 years of age or patients of less than 20 kg, or in such situations where the measurement accuracy should be better than ±11,8%.
- The device must not be used under direct sunlight, a powerful spotlight or a flashing fluorescent tube. In this case, the measurement accuracy cannot be guaranteed.
1.3 PATIENT SAFETY

- The device must not be used during magnetic resonance imaging (MRI).

- The Monidrop® W infusion monitor is designed as a measuring device for the administration of basic fluids, nutritional solutions, blood products or antibiotics. The device is not applicable for use together with strong medicinal substances requiring high accuracy.

- Only use infusion set suitable for the device, with a drip size ALWAYS of 20 drops (gtt)/ml and accuracy of at least ±10%. Before use, check that the infusion set is attached correctly to the device.

- The Monidrop® W infusion monitor always uses ml/h (millilitres per hour) as the measuring unit for the infusion rate.

- Make sure that fluids can flow freely by straightening all lines.

- The height of the IV bag should be suitable for gravity infusion. The height should be about 60 ... 90 cm above the patient.

- Make sure that the level of the drip chamber does not reach the measurement range. Fill the drip chamber halfway at the most.

- When replacing infusion sets, follow the infusion set manufacturer’s recommendations (note country-specific hygiene regulations).

- The operating conditions of the device must fulfil proper requirements (e.g. VDE 0100, VDE 0107 or IEC regulations). Note country-specific guidelines and differences.

- The device has also been approved for home care in accordance with standard 60601-1-11.

- Do not use the device in a room containing explosive substances.

- Protect the charger against moisture.

- If the device drops or suffers a heavy impact, it must be sent to a service point for inspection.

- Only use the charger delivered with the Monidrop W device.

- Do not tilt the device heavily when using it. If the device is tilted by
more than 20 degrees, its measurement accuracy cannot be guaranteed.

- The user must always evaluate the correctness of the information displayed before making any treatment decisions.
- Regardless of recommended limits, the values selected for treatment must be medically correct.
- The user must ensure that patient information is correct and the selected target concentrations and measured dosage rates comply with the regulations of the country in question.
- Connection is permitted only to devices that Monidor Oy has approved.

**WARNING:** The device does not identify any air or bubbles in the IV line.

**WARNING:** The device does not identify any significantly high flows (continuous drip) if the roller clamp is fully opened. In these cases, the device may not give a direct notification.

**WARNING:** Modifying this device is prohibited.

**WARNING:** The charger cable causes a choking hazard for small children.

**WARNING:** Do not use the device if its charger or cable is damaged.

### 1.3.3 Blood transfusion

- The Monidrop® W infusion monitor can be used to monitor infusion of blood products. Only use disposable products intended and indicated for blood products. The drip size of the blood line must be 20 drops (gtt)/ml.
1.3 PATIENT SAFETY

1.3.4 Other infusion factors

- Check the compatibility of the products and medicine used from the manufacturers’ specifications.

1.3.5 Safety standards

- The Monidrop® W infusion monitor fulfils all safety regulations of standards IEC/EN 60601-1 and EN 60601-1-11:2015 on medical electrical equipment. The Monidrop® W infusion monitor fulfils the sections applicable to infusion monitoring devices of the following chapters of standard EN 60601-2-24: 201.7.9.2, 201.7.9.3 and 201.12.4.4.109.

- The device fulfils electromagnetic compatibility (EMC) requirements in accordance with standards IEC/EN 60601-1-2 and IEC/EN 60601-2-24.

- This product is compliant with Directive 2014/53/EU. The relevant Declaration of Conformity is available at www.monidrop.com.

- If the device is used in the proximity of other devices that may cause significant interference (e.g. high-frequency surgical equipment, computed tomography equipment, infrared transmitters, etc.), follow the safe distances recommended for such equipment.

- If the Monidrop® W infusion monitor is used at temperatures lower than the indicated operating conditions, it must be kept at room temperature for at least one hour before use.

- If the Monidrop® W infusion monitor is used at temperatures higher than the indicated operating conditions, it must be kept at room temperature for at least 30 minutes before use.
2 Symbols / Menu structure

Device symbols:

Power switch On/Off

Notification button

Näytön symbolit:

Battery status

Mute

Sound on

Drip indicator

Remotely set target limit

or limit (if used with IV Screen)

Liquid volume

Treatment duration

Infusion rate

Notification

Device or drip chamber tilted

Infusion rate more than 1,200 ml/h

Infusion rate less than 6 ml/h

Unstable drip detection
2 SYMBOLS / MENU STRUCTURE

Label:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description of symbols</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Information" /></td>
<td>Note: Read the instructions delivered with the product.</td>
</tr>
<tr>
<td><img src="image" alt="Drip size" /></td>
<td>Drip size: 20 drops (gtt)/ml</td>
</tr>
<tr>
<td><img src="image" alt="Markings of electrical and electronic equipment" /></td>
<td>Markings of electrical and electronic equipment in accordance with directive 2002/96/EC (waste electrical and electronic equipment)</td>
</tr>
<tr>
<td>CE0598</td>
<td>CE marking in accordance with directive 93/42/EEC</td>
</tr>
<tr>
<td><img src="image" alt="Temperature limit" /></td>
<td>Temperature limit</td>
</tr>
<tr>
<td><img src="image" alt="Humidity limit" /></td>
<td>Humidity limit</td>
</tr>
<tr>
<td><img src="image" alt="Air pressure limitation" /></td>
<td>Air pressure limitation</td>
</tr>
<tr>
<td><img src="image" alt="Non-ionising electromagnetic radiation" /></td>
<td>Non-ionising electromagnetic radiation (IEC 60417-5140)</td>
</tr>
</tbody>
</table>
Main screen (measurement view)

The top row indicates the device status; battery status, infusion rate and notification sound.

Treatment duration

Current infusion rate

Liquid volume administered during treatment

If the line in the adjacent figure is shown for the current infusion rate, the measurement result is uncertain or the drip cannot be detected. Check that the drip chamber and device are straight and correctly attached. Remove any foam from the drip chamber.
3 Using the device

3.1 Activation and start of infusion

3.1.1 Preparations

1. Insert the pointed tip of the IV line into the bottle in an upright position. Fill at most 1/2 of the drip chamber to keep the drip detection function operational.

2. Fill the drip chamber and close the roller clamp.

3. Check that the Monidrop® W device is undamaged and contains all the necessary parts.

3.1.2 Starting the device

1. Start the device by pressing the power button. If the device is connected to the charger, a battery charge symbol will be displayed on the screen and an orange light will indicate that the device is being charged.

Note the self-test program: The device performs a self-test program during start-up.

3.1.3 Starting infusion monitoring

1. Attach the Monidrop® W device to the drip chamber as instructed.

2. Make sure that the device is firmly in place and in an upright position.

3. Use the roller clamp to set the infusion rate. The measurement result is shown by the Monidrop® W device.

4. The Monidrop® W device gives notification if the infusion rate is over 1200 ml/h or the battery status is low.

5. Press the notification button on the front of the unit if you want to activate or deactivate notifications when infusion rate is less than 6 ml/h or when battery status is low.
3 USING THE DEVICE

Note: Attaching different types of infusion sets to the Monidrop® W device must be done in accordance with the instructions.

WARNING: Do not apply any force when attaching the drip chamber to the Monidrop® W device. If the fastening mechanism seems stiff, check that the position and installation of the infusion set and clamp is correct.

- Only use infusion sets suitable for the device, with a drip size ALWAYS of 20 drops (gtt)/ml and accuracy of at least ±10%. Before use, check that the line is attached correctly to the device.

3.2 Starting a new treatment
To start a new treatment, reset the previous treatment by turning the power off for three (3) consecutive seconds . Then, restart your device to be ready to use again.

3.3 Ending an infusion treatment
You can stop monitoring the treatment in progress by switching off the device by pressing the power button for three (3) seconds.

3.4 Shutting down the device
Shut down the device by pressing the power button for three (3) seconds. When the device is switched off, any previously set data and settings will be reset.
4 Notifications

The Monidrop® W infusion monitor is equipped with its own notification system (notification sound and optic notification).

<table>
<thead>
<tr>
<th>Notification type</th>
<th>Notification type</th>
<th>Optic notification</th>
<th>Acknowledgment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Yellow LED</td>
<td>Red LED</td>
</tr>
<tr>
<td>Visual notification</td>
<td>No</td>
<td>Continuous</td>
<td>Blinking triangle</td>
</tr>
<tr>
<td>Sound notification</td>
<td>Yes</td>
<td>Blinking</td>
<td>Blinking triangle</td>
</tr>
<tr>
<td>Self test notification</td>
<td>Yes</td>
<td>Continuous</td>
<td>Error code (Check chapter 4.1.3.)</td>
</tr>
</tbody>
</table>

4.1 Notifications

4.1.1 Infusion rate is less than 6 ml/h
You can activate the notification by pressing the notification button on the front of the device. The message is displayed with a flashing yellow light and sound and the screen displays a flashing notification triangle at the infusion rate symbol.

- Notification when infusion rate is less than 6ml / h
4.1.2 Battery low
The device gives notification with a blinking yellow light, a notification tone and a flashing triangle at the infusion rate symbol. The message is automatically cleared when the device is connected to the charger.

• The device gives notification when the battery status is down to 10%.

4.1.3 Infusion rate is over 1200 ml/h
The message is displayed with a flashing yellow light and sound and the screen displays a flashing notification triangle at the infusion rate symbol.

• Notification when infusion rate is more than 1200 ml/h

4.1.4 Self-test program and forced restart
The device performs a self-test program during start-up. If there are any problems during the test, the device will give an notification by emitting a continuous red light and tone, and all device functions will be locked. The device can be switched off and restarted, in which case the device will perform self-test program again.

If the device freezes, it may be forced to restart by pressing the power button for an continuous period of eight 8 seconds. Contact maintenance if the device does not start regardless of the restart.

4.2 Managing audio notifications
Audio notifications can be muted and activated using the notification button. At the top of the screen, the mute symbol is displayed when the message is less than 6ml / h. Notifications with an infusion rate less than 6 ml/h are always muted by default when the device is turned on.

By pressing the audio notification button a second time, the battery charge status notification sound is also muted. Pressing the button again will enable audible notifications for battery status and infusion rate less than 6 ml / h.
5 Battery charge and maintenance

The Monidrop® W infusion monitor is equipped with a rechargeable lithium ion polymer battery. A new battery can be used for 72 hours on one charge. To ensure optimal use, the device is equipped with a protective function against overcharging and excess battery discharge (prevention of deep discharge).

At the top of the screen, the battery status is indicated in percentage, and the battery symbol indicates the battery status in steps. Symbols indicate the battery status as shown below. On the left, the battery is fully discharged and on the right, the battery is fully charged. The percentage displayed below indicates the battery status on a rough level.

| 0% | 25% | 50% | 75% | 100% |

The Monidrop® W infusion monitor can also be used when its battery is being charged, and it will operate normally. When the device is connected to the charger, the middle LED at the top of the screen will be orange when the battery is being charged and the symbol shown below will be displayed in place of the battery status symbol.

If the battery is fully discharged, the device will go off automatically. If the battery is fully discharged, keep the charger plugged up to one hour until the charging light comes on. The battery can then be fully charged.

Note: Only use the charger delivered with the device to charge the Monidrop® W infusion monitor. Note that the device has a limited movement range when the charger is connected.
Instructions for the optimal use of the battery:

• Charge the battery at least once every three months.

• At the regular operating temperature, the battery can be charged and discharged approximately 500 times before its capacity decreases below 80% of the maximum value.

• The battery will slowly discharge if the infusion monitor is not connected to the power grid and even if the device is not in use.

• The full operating period of the battery can only be achieved when the device is operated uninterrupted at room temperature and the battery is fully charged. The battery capacity displayed is indicative. If the battery is older, the actual operating period may differ from that shown on the screen.

Note: The battery may explode or leak if it is opened or burned/heated. Note waste handling guidelines.

Note: While the device is charged, disconnection from the mains supply occurs by removing the charger from the AC power socket.

**WARNING:** While charging Monidrop during use, ensure that access to the mains plug is not blocked so that the charger can be disconnected from the mains power receptacle in the event of an emergency.

**Battery maintenance and warranty:**

The device manufacturer provides the battery with a warranty of two (2) years. This is the typical service life of the battery (without causing any significant reduction in the operating period). After two years, the battery will be replaced during scheduled maintenance.

The battery can only be replaced by a service provider authorised by Monidor.
6 Cleaning

Wipe the surface clean using a piece of cloth and a mild soap-water solution. If required, disinfect the device using ApoWipe or a similar product.

The device can also be disinfected as follows:

- 80% isopropanol
OR
- Mild alkaline disinfectant
OR
- Phenoxyethanol + quaternary disinfectant, dilution 1:10
OR
- Chlorine-containing disinfectant, with an active chlorine content of up to 2%, dilution 1:10

Note: Do not immerse the device in any fluid or spray any water or cleaning agent on the device.

Note: Do not use a cleaning stick to touch or press the protective film behind the speaker holes and bars.
7 Warranty / Inspection and maintenance / Training

The manufacturer’s liability

The manufacturer, assembly or installation company or importer is only responsible for device safety when:

- An authorised person has installed the device or made expansions, additions, modifications or repairs.
- Electrical installations in the operating facilities are in accordance with regulations (e.g. VDE 0100, 0107 and/or IEC standards), taking any country-specific requirements into account.
- The device is used in compliance with the operating manual.
- Regular maintenance has been performed regularly.

Warranty

- Monidor provides all Monidrop® W infusion monitors and accessories it has manufactured with a warranty of 12 months. The warranty covers installation and the replacement of defective parts if the defect is caused by a structural, manufacturing or material error. The warranty becomes void if the owner/user or any third party makes modifications to the device.
- The warranty does not cover any defects caused by an incorrect or improper use or regular wear and tear.
- The device must be disposed of as electrical and electronic waste (currently in EU area only).

Technical inspection and maintenance

- It is recommended that a technical safety inspection and scheduled maintenance be performed every two years. Any markings in accordance with the checklist must be entered in the device maintenance register. Maintenance can only be performed by maintenance employees trained/authorised by Monidor or its distributor.
- Monidor will provide authorised maintenance organisations with maintenance instructions at request.
Periodic inspections:

Check that the device is clean, undamaged and in an operating condition. Only use the device in accordance with this operating manual.

The areas at the back of the device inside the white frames in the figure below are intended for the labels of hospitals or health care centres.

Training

Training for the use of the Monidrop® W infusion monitor is available. Contact the device seller for more information.
8 Content of the delivery and inspecting the content / Recycling

Content of the delivery

Monidrop® W infusion monitor, charger, quick user guide

Inspecting the content of the delivery

Regardless of protective packaging, there may be transportation damage. Check that nothing is missing from the delivery. Do not use the device if it is damaged. Contact device maintenance.

Test that the device works correctly before its first use. This is a statutory procedure in many countries.

Recycling

The device and battery must be disposed of as electrical and electronic waste.

Note any hygiene and waste handling guidelines related to disposable products and IV fluids.
## 9 Technical specifications

<table>
<thead>
<tr>
<th>Device type</th>
<th>Infusion monitor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Classification (according to EN 60601-1)</td>
<td>Internally powered medical device. Charger and device complies with class II requirements while connected to mains supply.</td>
</tr>
<tr>
<td>Class (Directive 93/42/EEC)</td>
<td>Im</td>
</tr>
<tr>
<td>Moisture protection</td>
<td>Monidrop® W IP 52 (dustproof, protection against water dropping vertically or at a maximum angle of 15 degrees) (Charger IP 22)</td>
</tr>
<tr>
<td>External power source:</td>
<td>5 V</td>
</tr>
<tr>
<td></td>
<td>100–240V ~47–63Hz</td>
</tr>
<tr>
<td></td>
<td>0.30–0.15 A</td>
</tr>
<tr>
<td></td>
<td>Using the charger delivered with the Monidrop® W device</td>
</tr>
<tr>
<td>Operating principle</td>
<td>Device is attached mechanically directly to the drip chamber of the infusion administration set. It calculates drops that are dropping inside the drip chamber. Drop detection is based on transmitting IR light through a drip chamber. When drop breaks the light, the event can be measured by receiving light sensor. Result are calculated and showed to user in convenient units (ml/h and ml).</td>
</tr>
<tr>
<td>Interference protection EMC</td>
<td>IEC/EN 60601-1-2 / 60601-2-24</td>
</tr>
<tr>
<td>Operating rate</td>
<td>100% (continuous operation)</td>
</tr>
<tr>
<td>Operating conditions</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Relative humidity, %</td>
</tr>
<tr>
<td></td>
<td>20% ... 85% (of air humidity condensation)</td>
</tr>
<tr>
<td></td>
<td>Temperature</td>
</tr>
<tr>
<td></td>
<td>+5 ... +40 °C</td>
</tr>
<tr>
<td></td>
<td>Air pressure</td>
</tr>
<tr>
<td></td>
<td>80 kPa ... 106 kPa</td>
</tr>
</tbody>
</table>
### Storage conditions:
- Relative humidity, %: 20 % ... 85 % (of air humidity condensation)
- Temperature: -20 ... +55 °C
- Air pressure: 80 kPa ... 106 kPa

<table>
<thead>
<tr>
<th>Specification</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Battery type</td>
<td>Li-ion-polymer</td>
</tr>
<tr>
<td>Operation time</td>
<td>Approx. 72 hours</td>
</tr>
<tr>
<td>Charge time</td>
<td>Approx. 2 hours</td>
</tr>
<tr>
<td>Weight</td>
<td>Approx. 110 g</td>
</tr>
<tr>
<td>Dimensions (W × H × D)</td>
<td>54 x 166.1 x 43.4 mm</td>
</tr>
<tr>
<td>Measurement range</td>
<td>6 ml/h ... 1200 ml/h</td>
</tr>
<tr>
<td>Volume range</td>
<td>0 ml ... 9999 ml</td>
</tr>
<tr>
<td>Treatment duration limit</td>
<td>99 hours 59 minutes</td>
</tr>
<tr>
<td>Measurement accuracy</td>
<td>Typical measurement accuracy ± 1.8%</td>
</tr>
<tr>
<td></td>
<td>Furthermore, the measuring accuracy is affected by the accuracy of the IV infusion set, which is 0% ± 10% while connected to the Monidrop</td>
</tr>
<tr>
<td>Regular maintenance (operating safety)</td>
<td>Recommended once in two years</td>
</tr>
<tr>
<td>Expected service life</td>
<td>5 years</td>
</tr>
<tr>
<td>Radio module (optional)</td>
<td>Compatible with WiFi protocols 802.11 b/g/n</td>
</tr>
<tr>
<td></td>
<td>Frequency range 2.4 GHz ~ 2.5 GHz (2400M ~ 2483.5M)</td>
</tr>
<tr>
<td></td>
<td>Standards FCC/CE</td>
</tr>
</tbody>
</table>
### Manufacturer's instructions and declaration of electromagnetic compatibility

<table>
<thead>
<tr>
<th>Disturbance test</th>
<th>Compatibility</th>
<th>Electromagnetic environment – instructions</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions</td>
<td>Group 1</td>
<td>Monidrop® W only uses RF energy in internal functions. Therefore, it only has very low emissions, and probably does not disturb any nearby electrical devices.</td>
</tr>
<tr>
<td>CISPR 11</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| RF discharge     | Class B       | Monidrop® W and its parts are suitable for use in all facilities, including households and residential buildings connected to the public low-voltage grid. |
| CISPR 11         |               |                                            |

| Harmonic emissions | Class A       |                                      |
| IEC 6100-3-2       |               |                                      |

| Voltage changes/flicker emissions | Compatible     |
| IEC 61000-3-3             |               |

**Note:** Maximum emissions were measured with a complete system. (Monidrop® W and its parts)
## Manufacturer’s instructions and declaration – electromagnetic interference tolerance

The Monidrop® W infusion monitor is intended for use in the electromagnetic environments defined below. The owner or user of the Monidrop® W infusion monitor and its parts must ensure that the device is used in such an environment.

<table>
<thead>
<tr>
<th>Interference tolerance test</th>
<th>Test level IEC 60601-1-2 IEC 60601-2-24</th>
<th>Compatibility level</th>
<th>Electromagnetic environment – instructions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic discharge (ESD) in accordance with standard IEC 60601-4-2</td>
<td>Contact: ± 6 ± 8 ± 15</td>
<td>± 6 kV no interference ± 8 kV stop with an alarm possible</td>
<td>The floor must be made of wood, concrete or ceramic tiles. If the floor is covered with a synthetic material, the relative humidity must be at least 30%.</td>
</tr>
<tr>
<td>Electrostatic transient/burst in accordance with standard IEC 61000-4-4</td>
<td>± 2 kV power supply lines ± 1 kV input/output lines</td>
<td>± 2 kV</td>
<td>The quality of supply current must correspond with the quality in a typical commercial environment or hospital environment.</td>
</tr>
<tr>
<td>Surge in accordance with standard IEC 61000-4-5</td>
<td>± 1 kV differential mode ± 2 kV common mode</td>
<td>± 2 kV</td>
<td>The quality of supply current must correspond with the quality in a typical commercial environment or hospital environment.</td>
</tr>
<tr>
<td>Voltage decreases, brief interruptions and voltage changes in power supply lines in accordance with IEC 61000-4-11</td>
<td>&lt;5 % Ut (&gt;95% decrease in Ut) For 0.5 periods 40 % Ut (60% decrease in Ut) For 5 periods 70 % Ut (30% decrease in Ut) For 25 periods &lt;5 % Ut (&gt;95% decrease in Ut) For five seconds &lt;5% Ut for five seconds (&gt;95% decrease)</td>
<td>Compatible when using an internal battery</td>
<td>The quality of supply current must correspond with the quality in a typical commercial environment or hospital environment.</td>
</tr>
</tbody>
</table>
**9 TECHNICAL SPECIFICATIONS**

**Note 1:** Different testing values of standard IEC 60601-2-24 are indicated in the table. These testing values do not permit any hazardous interference, while interference is permitted using the lowest testing values of standard IEC 60601-1-2.

<table>
<thead>
<tr>
<th>Magnetic field at a network frequency (50/60 Hz) in accordance with standard IEC 61000-4-8</th>
<th>IEC 60601-2-24: 400 A/m</th>
<th>400 A/m</th>
<th>Magnetic fields at a network frequency must be at a level characteristic to a typical commercial environment or hospital environment.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IEC 60601-1-2: 3V_{eff} normal and 10V_{eff} Within ISM frequency range</td>
<td>10V_{eff} 150 KHz – 80 MHz</td>
<td>Mobile and movable RF communications devices cannot be used at a distance closer to Monidrop and its parts than the distance calculated using the formula applied to the transmitter frequency.</td>
</tr>
</tbody>
</table>
| | IEC 60601-2-24: 10V_{eff} 150 kHz – 80 MHz | 10 V/m 80 MHz – 3 GHz | **Recommended distance**

\[ d = 1.2 \sqrt{P} \]

Field intensity must be less than 10V/m

\[ d = 1.2 \sqrt{P} \]

800 MHz – 2.5 GHz

where P is the maximum output of the transmitter in watts (W) in accordance with the transmitter manufacturer and d is the recommended distance in meters (m).

The field intensity caused by fixed RF transmitters which can be determined by means of an electromagnetic survey must be lower than the compatibility level at each frequency range.

There may be interference close to devices furnished with the following symbol:

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**Note 2:** A higher frequency range is applied to 80 MHz and 800 MHz.

**Note 3:** These instructions may not apply to all situations. Electromagnetic spreading is also affected by absorption to structures, objects and people, and reflection from them.

**Note 4:** Different testing values of standard IEC 60601-2-24 are indicated in the table. These testing values do not permit any hazardous interference, while interference is permitted using the lowest testing values of standard IEC 60601-1-2.
The Monidrop® W infusion monitor is intended for use in electromagnetic environments where radiated RF emissions are controlled. The owner or user of Monidrop® W and its parts can help to prevent electromagnetic interference by keeping the minimum distance between mobile and movable RF communications devices (transmitters) and Monidrop® W as defined in the following recommendations in accordance with the maximum output of the communications device.

<table>
<thead>
<tr>
<th>Rated power of radio transmitter (W)</th>
<th>Distance in accordance with transmitter frequency (m)</th>
<th>150 KHz – 80 MHz</th>
<th>80 MHz – 800 MHz</th>
<th>800 MHz – 2.5 GHz</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.01</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.1</td>
<td></td>
<td>0.12</td>
<td>0.12</td>
<td>0.23</td>
</tr>
<tr>
<td>1</td>
<td></td>
<td>0.38</td>
<td>0.38</td>
<td>0.73</td>
</tr>
<tr>
<td>10</td>
<td></td>
<td>1.2</td>
<td>1.2</td>
<td>2.3</td>
</tr>
<tr>
<td>100</td>
<td></td>
<td>3.8</td>
<td>3.8</td>
<td>7.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>12</td>
<td>12</td>
<td>23</td>
</tr>
</tbody>
</table>

**Note 1:** If the maximum output of the transmitter is not presented above, the recommended distance (d) in meters (m) can be defined using the formula applied to the transmitter frequency, where P is the maximum transmitter output in watts (W) as indicated by the transmitter manufacturer.

**Note 2:** An additional factor of 10/3 is used to calculate the recommended distance for transmitters that operate within the frequency range of 0.15 MHz – 2.5 GHz. This reduces the possibility that mobile/movable communications devices cause interference if they are accidentally carried to a patient area.

**Note 3:** These instructions may not apply to all situations. Electromagnetic spreading is also affected by absorption to structures, objects and people, and reflection from them.
10 Operating instructions for accessories

Charger

The Monidrop® W infusion monitor is equipped with a charger, with which the battery can be charged during the use of the device and when the device is not in use.

The Monidrop® W infusion monitor can only be charged using the charger delivered with the package. Always check that the correct charger is used.

Technical specifications: 100–240 V ~47–63 Hz 0.30–0.15 A

Manufacturer: Sinpro, made in Taiwan
Model number: HPU10C-102 or HPU10B-102
IP 22

Important charger safety instructions:
This product is intended for indoor use only. When using an electrical appliance, basic safety precautions should always be followed, including the following: Read all instructions before using power supplies.

DANGER - To reduce the risk on electric shock: A power supply should never be left unattended when plugged in. Always unplug this power supply from the electric outlet immediately after using.

WARNING - To reduce the risk of burns, fire, electric shock, or injury to persons:
1. Do not allow to be used as a toy. Pay close attention when this power supply is used by or near children.
2. Use this power supply only for its intended use as described in this manual. Use only attachments recommended by the manufacturer as contained in this manual.
3. Never operate this power supply if it has a damaged cord or plug, if it is not working properly, if it has been dropped or damaged, or fallen into water. In such cases, return the appliance to the nearest authorizes dealer or service center for examination, repair, electrical or mechanical adjustment.
4. Never drop or insert any object into any openings.
5. Do not use outdoors.
6. Do not operate where aerosol (spray) products are being used or where oxygen is being administered.
7. The socket-outlet should be installed near the power supply and should be easily accessible.
8. Do not unplug by pulling on cord. To unplug, pull on the plug, not on the cord.
9. Units without level V Efficiency may no longer be used for electronic household and office equipment.
10. Manufacturer can be contacted: Sinpro Electronics Co., Ltd. No.5, Yanxi St. P.E.P.Z. Pingtung City, Pingtung County 90093, Taiwan