



moments to care



**Make Infusion
Smarter and Safer**



MEDCAPTAIN MEDICAL TECHNOLOGY CO.,LTD.



Address: 12th Floor, Baiwang Research Building, 5158 West
Shahe Road, Xili, Nanshan, 518055 Shenzhen, Guangdong,
PEOPLE'S REPUBLIC OF CHINA
Tel: +86 755 26953369 E-mail: info@medcaptain.com
Website: www.medcaptain.com

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EN_MP Series -16P-Version 1.3

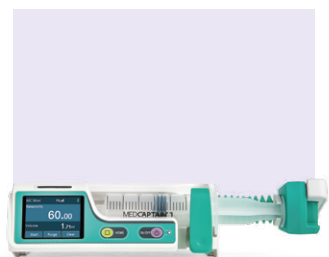
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Touch the future with infusion revolution

MP-30
Syringe Pump



MP-60
Infusion Pump



MP-80
Infusion Workstation



Deeply understanding customers' needs, we persist in providing innovative technologies and products. Such pioneering features as touch screen, multiple infusion modes, and flexible combinations with plug-in model deliver our commitment to the medical community.



Our infusion products and solutions have led to innovation in the infusion field, gaining users' favor and trust in key departments of China's top level university teaching hospitals and medical institutions in more than 30 countries.

Simple and easy to use



Touch operation, easy and convenient.

Innovative touch operation, quick response to various settings, reversed the complicated operation process and brings a different user experience.



Flexible and lightweight

Compact and light body design, flexible and versatile, allowing for free conversion to different clinical circumstances. Pumps can be freely assembled through the guide rail, without any other tools.

Ultra-long standby time

High capacity lithium battery provides reliable support for transport and other emergencies.

Small size, profound capabilities



Automatic pump door

Press lightly and the door opens. Patently designed automatic door guarantees infusion safety under complete closed pump door.

Multiple Infusion Modes

Satisfy needs with several clinical departments and different drug infusion cases. More efficient and intelligent.

Micro Mode

Applications include the use of different syringes for multiple patients that allow different specifications, providing double effectiveness to infusion.

Intelligent Bolus

Three modes in which respond to any type of medical emergency.

Information network

Internal WIFI module, connect to iCIS through hospital LAN, for centralized monitoring of infusion. Or connect to department CIS and HIS system to integrate clinical information data.

Customized calibration

Open consumable, customize and calibrate the consumable brands. Auto-recognize consumable specification, making infusion safe and precise.



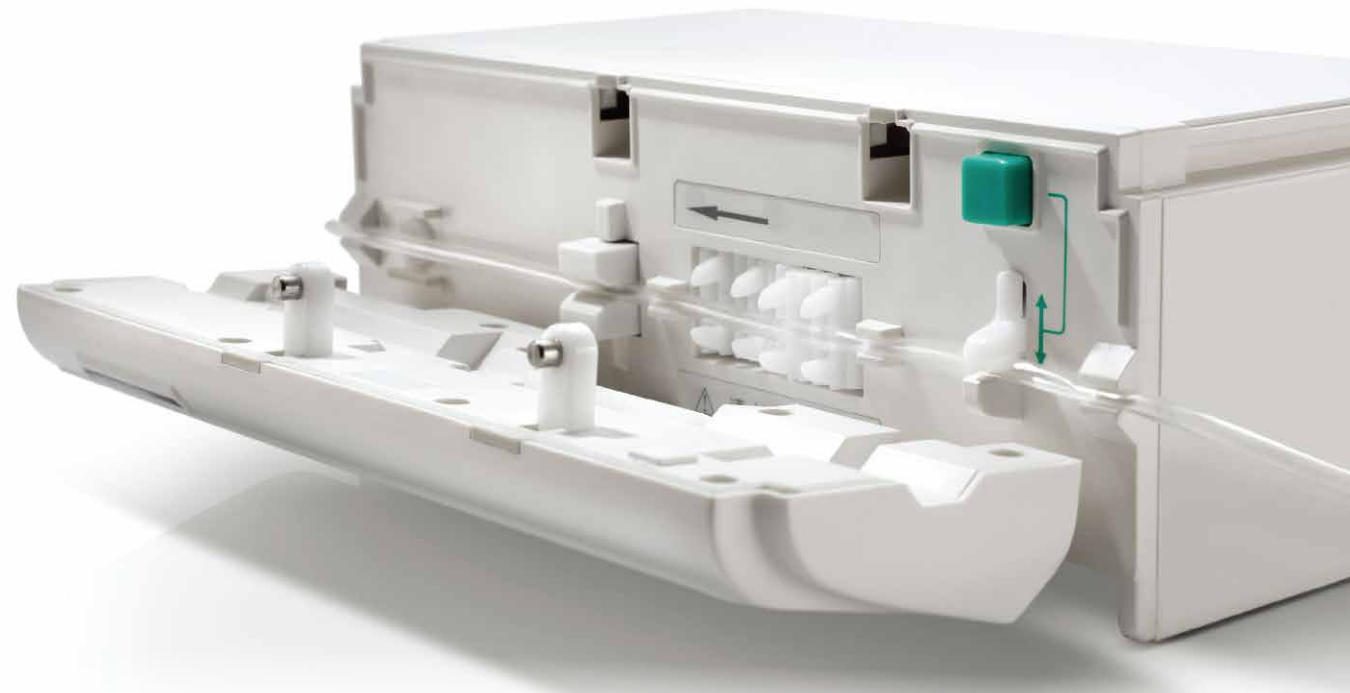
Customized drug library

It may store information of more than 1000 drugs providing a customized drug database for clinics and thus making infusion convenient and safe.

Every innovation targets safer infusion

Automatic antifreeflow system

Press lightly, the anti free-flow clamp will open or close. When pump door opens, the free-flow clamp closes automatically, avoiding risks from liquid spills.



Occlusion alarm and Anti-Bolus

11 occlusion levels setting are available for clinical use of various drugs. When the infusion tube occludes there is automatic decompression, to avoid liquid spills, drug pollution and waste caused by pressure change. Also reducing the risk from over-dose infusion.

Double CPU design, two-way independent sound-light alarm

independent double CPU control guarantees real-control and data reliability, creating strict management of the risk process and a complete alarm system to reassure the clinics.

Emergency alarm system

Internal super-capacitor, providing a 3 minute alarm in the case of a sudden power exhaust, to guarantee safety.

Total squeezing peristaltic system pump

Adopt the most mature total flexible peristaltic technology in the industry, making infusion more safe and precise.

Infusion Consumables

Syringes



Syringe extension tube



IV-sets





Not just simply stackable

MP-80 Infusion Workstation

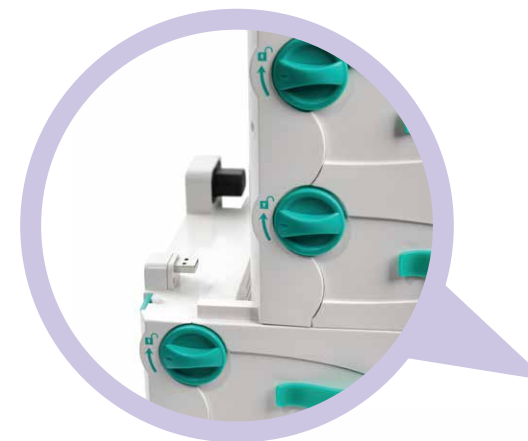


Intelligent relay infusion mode

Auto-continuous infusion, reduced risks from interrupted infusing, satisfying the requirement of non-stop infusion for high-alert medications.

Plug-in and modular design

From 2 to 15 channel infusion workstation for clinical needs can be combined optionally, satisfying requirements of several drugs infusions at the same time for critical patients. Or in some cases, the surplus pump can be removed and used in another bed to improve capacity utilization.





Plug & Play design

Without any tool, remove or plug single pump within 3 seconds, easily initiate continuous infusion while transporting patients.

Information network

Internal WIFI module, connect to iCIS directly, for centralized monitoring of infusion. Or connect to CIS or HIS system to integrate clinical information data.

One click sync for system parameters

Parameters such as volume, date, time and screen brightness, just set them on a pump, and the other pumps in the workstation can be synced automatically.

Space management

Only with one power cord and power is supplied to all the pumps within the workstation.

An infusion tube managing clamp is capable of holding all of infusion tubes and extended tubes, making a tidy working station and saving space, resources, avoiding safety risks.



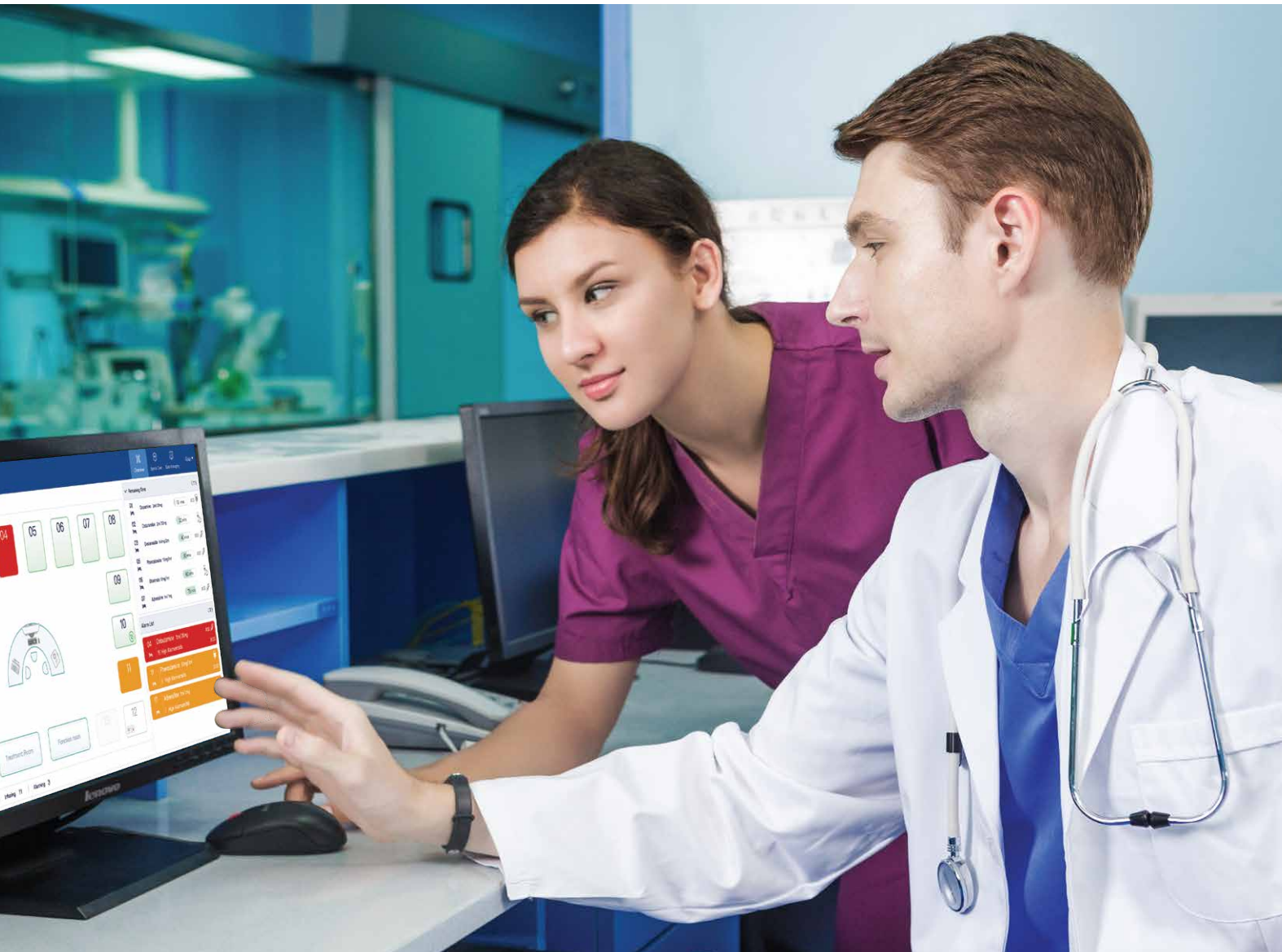
Synchronous alarm

When a single pump alarms, the SCU on the top of workstation alarms at the same time with a loud signal, that reminds doctors or nurses of handling the problem in time.

Intelligent Bi-power supply system

Delicate internal circuit design, allowing for quick responses to sudden power outages.

Visible informatization management



Infusion information central monitoring

The real-infusion information of single pump and workstation can be remotely monitored by iCIS system through WIFI. From your bedside, nurse station, office, corridoranywhere. You can know every patient's infusion status. Maximum 50 infusion beds'with alarms, remaining infusion time, drug name and infusion rate all visible on the interface.

MP-30 Syringe Pump



MP-80 Infusion Workstation

MP-60 Infusion Pump



Data statistics and analysis

Multiple internal statistic modes, real-time information including device use, drug infusion volume, patient quantity and drug flow rate. It is a convenience for the billing and accounting departments to generate the balance of payment and resources. To have a broader, deeper and direct understanding, this provides statistical data and support for clinical surveys and research.

Drug and consumable library

Easily edit and generate more than 1000 drug entries including information, frequently-used consumables in department, and quickly download them to the pump, simplifying clinical personnel job.

Information integration

It connects the CIS and HIS systems to consolidate the data integration of clinical information.

MP-30A Syringe Pump

Small

- Compact design help space-saving
- Light and easy carrying
- Stackable design, connectable with pumps or workstation



Smart

- User-defined screen brightness and background colors
- Relay function between pumps on any mode and auto shift therapies without pause

Safe

- High accuracy
- Visual, audible alarm information
- Occlusion levels adjustable
- Real-time monitoring infusion states

Sophisticated

- Wi-Fi function available
- Connectivity even in transport
- Compatible with HIS/CIS system
- Connectable with infusion central monitoring system

Simple

- Convenient touchscreen design
- Intuitive operation
- Easy management of infusion
- Clear interface for easy-read information

Specifications

Dimension	252(W) x 74(H) x 133(D) mm
Weight	< 1.6kg (including battery)
Classification	Defibrillation-proof type CF applied part, IPX2
Screen	2.4" LCD touchscreen, 7 colors selectable
Compatible Syringes	All syringes of 5ml,10ml, 20ml, 30ml, and 50/60ml conforming to the standard

Basic Parameter

Infusion mode	Rate, Time, Weight, Sequence, Relay, loading dose, Trapezia, Micro, TIVA	
Infusion rate range	0.1-100ml/h (5ml syringe)	0.1-300.0 ml/h (10ml syringe)
	0.1-600.0ml/h (20ml syringe)	0.1-900.0ml/h (30ml syringe)
	0.1-2000ml/h (50/60ml syringe)	
Increment	0.1 – 99.99 ml/h, minimum increment 0.01 ml/h	
	100 – 999.9 ml/h, minimum increment 0.1 ml/h	
	1000 – 1200 ml/h, minimum increment 1ml/h	
VTBI range	0.1-99.99ml (minimum increment 0.01)	
	100 – 999.9ml(minimum increment 0.1)	
	1000 – 9999ml(minimum increment 1)	
Preset time	00:01 – 99:59 (hh:mm) adjustable, step by 1 min	
Accumulated volume	0 – 9999.99 ml	
Accuracy	Mechanical accuracy: $\pm 1\%$, Accuracy including syringe: $\pm 2\%$	
KVO rate	0.1 – 5.0 ml/h adjustable, step by 0.01 ml/h, default rate 1ml/h	
Bolus rate	0.1-100ml/h (5ml syringe)	0.1-300.0 ml/h (10ml syringe)
	0.1-600.0ml/h (20ml syringe)	0.1-900.0ml/h (30ml syringe)
	0.1-2000ml/h (50/60ml syringe)	
	3 bolus ways selectable: Automatic/ Manual/ Rapid quantitative Bolus	
Purge rate	0.1-100ml/h (5ml syringe)	0.1-300.0 ml/h (10ml syringe)
	0.1-600.0ml/h (20ml syringe)	0.1-900.0ml/h (30ml syringe)
	0.1-2000ml/h (50/60ml syringe)	
Occlusion level	11 levels selectable, (225mmHg-975mmHg)	

Special Function

Recent therapy	20 recent therapies are recorded and can be directly viewed by simple selection
Rate change in infusion	Infusion rate can be changed anytime during infusion without stop
Relay function	Relay function is available with workstation, it allows two pumps that under any infusion mode shift without pause
Auto screen lock	This function can be set to OFF, 15s, 30s, 1min, 10min or 30min The default value is OFF
Drug library	A maximum of 2,000 drug types can be stored.
Repeat alarming	Alarm sounds again in 2min, if there is still any errors after the alarm is muted
Event recording	Maximum 2000 events can be stored for review
Sound Volume	11 levels selectable
Power supply switching	When AC/DC power supply is cut off, the infusion automatically switch to internal battery supply

Power Supply

AC power	100 -240 V, 50/60 Hz, power consumption 45 VA
External DC power	DC 12 V, 1 A
Internal battery	Lithium battery 7.2 V, 2600 mAH
Battery duration	≥ 6 hours @ 5ml/h (with a new battery), 10 hours to fully charge

Alarm

Visual and acoustic alarm	Near Finished, Finished, Syringe Empty, Near empty, OCCL, Low Battery, Battery Empty, No Battery, No Power Supply, Unknown Syringe, Syringe Install Error, Standby Time Expired, Relay Index Duplicate, Syringe Start Fail, Reminder Alarm, OCCL Risk, Tube Off (Only suitable for 30ml, 50/60ml syringes)
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Connectivity

WiFi / Wired	To connect central infusion monitoring system, nurse pager
Barcode scanning	Patient information input by barcode scanner
Data interface	USB*3, LAN, RS485, Wi-Fi (optional)

Mounting

Workstation	Supported
Between pumps	Stackable between any MP series syringe / infusion pumps produced by Medcaptain
Handle	Available
Pole clamp direction	Can be mounted at different directions, 90°/180°/270°/360°

Safe Requirement

Operating conditions	Temperature: 5°C to 40°C Humidity: 15% to 95% RH, non-condensing Pressure altitude: 70.0 kPa – 106.0 kPa
Storage conditions	Temperature: -20°C to +50°C Humidity: 10% to 93% RH, non-condensing Pressure altitude: 22.0 kPa – 107.4 kPa



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Add: 12th Floor, Baiwang Research Building, No.5158 Shahe West Road,
Xili, Nanshan, 518055 Shenzhen, Guangdong, PEOPLE'S REPUBLIC OF CHINA

Tel: +86 755 26953369

E-mail: info@medcaptain.com

Fax: +86 755 26001651

Website: www.medcaptain.com

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EN-MP-30A-2P-Version 1.6

MP-80

Infusion Workstation



Specifications

Name	Infusion workstation
Model	MP-80
Power supply	AC Power supply: 100-240V, 50/60 Hz Power consumption less than 100VA
Cabinet body quantity	1 to 3
Syringe/infusion pump combination channels	1 to 9
QR scanning	Patient information can be input by QR code scanning.
Operating conditions	Temperature: 5°C to 40°C Humidity: 15% to 95% RH, non-condensing Pressure altitude: 22.0 kPa-107.4 kPa
Storage conditions	Temperature: -20°C to +50°C Humidity: 10% to 93% RH, non-condensing Pressure altitude: 22.0 kPa – 107.4 kPa
Classification	Class I, IPX2
Dimensions	Cabinet cover: 237(W) × 34(H) × 160(D) mm Cabinet body: 254(W) × 254(H) × 152(D) mm
Weight	Cabinet cover < 0.62 kg Cabinet body < 1.6 kg



MEDCAPTAIN MEDICAL TECHNOLOGY CO., LTD.



Add: 21-23 F, Unit A, Building 7, Phase 3 of Shenzhen International Innovation Valley,
Liuxin Fourth Street, Xili, Nanshan District, Shenzhen City

Tel: +86 755 26953369

E-mail: info@medcaptain.com

Fax: +86 755 26001651

Website: www.medcaptain.com

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EN-HP-80-1P-Version 3.0

DISTRIBUTOR:





MP-30/MP-30A/MP-30T

Syringe Pump

User Manual

Intellectual Property and Statement

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Statements

MEDCAPTAIN reserves the right for final interpretation of this manual.

MEDCAPTAIN reserves the right to modify the contents of this manual.

The modified contents should be reflected in the newly published manual version.

This manual is applicable to MP-30, MP-30T, and MP-30A syringe pumps.

MEDCAPTAIN is responsible for safety, reliability and performance of this equipment only in the condition that:

- Use in accordance with the user manual.
- All disassembly, replacement, test, modification and repair are conducted by qualified personnel approved by MEDCAPTAIN.
- All replacement parts, supporting accessories and consumables during the maintenance are provided by MEDCAPTAIN.
- Maintenance records for product are reserved.

Please read this “User Manual” carefully and follow “Precautions for Use” before using the MP-30 Series Syringe Pump.



Shanghai International Holding Corp. GmbH (Europe)
Eiffestrasse 80, 20537 Hamburg, Germany



Version Information

- V4.0
- Issued on: June 6, 2023

After-Sale Service

Thank you for using the syringe pump of MEDCAPTAIN MEDICAL TECHNOLOGY CO., LTD.

- During the warranty period, we provide free after-sale services except the following causes:
 - Artificially damaged.
 - Inappropriate use.
 - The voltage of supply network exceeds the range.
 - Natural disasters.
 - Replacement or usage of (non-original) parts, accessories and/or consumables without prior approval of MEDCAPTAIN.
 - Other problems not caused by product itself.

After the warranty period, we continue to provide charged maintenance service. If you have any question when using the syringe pump, please contact your local distributor or directly to MEDCAPTAIN.

Contact our after-sales service department:

After-sales service provider: MEDCAPTAIN MEDICAL TECHNOLOGY CO., LTD.

After-sales address: 12th Floor, Baiwang Research Building, No.5158 Shahe West Road, Xili, Nanshan, 518055 Shenzhen, Guangdong, PEOPLE'S REPUBLIC OF CHINA

Telephone: +86-755-26953369

Postal code: 518055

Website: <http://www.medcaptain.com>

E-mail: mc.service@medcaptain.com

- MEDCAPTAIN MEDICAL TECHNOLOGY CO., LTD. and all local distributors with established after-sales service agencies can effectively resolve your problems in time.



WARNING:

- The device should be operated by clinic medical staff or under the instruction of special clinic medical staff. The operator should be trained on how to use this product.

Preface

Illustrations

All the illustrations provided in this operation manual are for your reference only. The settings or data on the illustrations may differ from the actual settings or data of the product.

Conventions

Italics: Indicates the quoted content.

User password: 1234

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1 Overview

1.1 Purpose

This product is intended for hospitals to infuse at constant rate liquids or liquid medicines through the veins of patients.

1.2 Contraindication

None

1.3 Product Features

MEDCAPTAIN MP-30 Series is a micro-volume continuous-operation syringe pump. It ensures constant infusion rate and accurate dosing volume during longtime infusion.

This syringe pump is used for continuous and micro-volume infusion of liquid or liquid medicine of little volume and high concentration, including, but are not limited to the infusion of chemotherapeutic agents, cardiovascular drugs, antineoplastic, oxytocic, anticoagulant, anesthetic agents.

- Disposable syringes of 5ml, 10mL, 20mL, 30mL, and 50/60mL can be recognized automatically.
- Providing eleven occlusion levels and displaying pressure status of the tube.
- Having large speed range (up to 2000mL/h for 50/60mL syringe).
- Calibration functions are provided for infusion accuracy.
- Safety design by monitoring infusion status of syringe.
- Multiple infusion modes.
- Configure with multi-channel infusion workstation, realizing relay infusion function.
- Wi-Fi function, can be connected to the infusion central monitoring system by intravenous infusion.
- Nurse call function.
- Touchscreen, providing quick and convenient user-friendly interface.
- Night mode display, reducing light interference to patients and environment.
- Supporting connection to a barcode scanner.
- Three types of power supply are supported: AC power supply, DC power supply, and internal lithium battery. The lithium battery can power the syringe pump for not less than 6 hours (at 5ml/h rate at room temperature).
- 2-CPU design and dual-channel real-time monitoring of infusion status for preventing exceptions like insufficient or excessive dose and reporting an alarm in case of an exception in time.
- Modular installation design enables multi-channel pumps among pumps.

Precautions for Use

2 Precautions for Use

In this manual, precautions are classified into warning and caution according to their importance. The meanings are as follows:



WARNING:

The information is about safety and efficiency. Operation against the precautions may cause injuries.

CAUTION:

The information is about guiding suggestions. Operation against the precautions may affect normal use of the product. Read carefully the warnings and cautions in this manual.



WARNING:

- The syringe pump must be operated by clinical professionals.
- Prior to use, please check the status of the pump, power cord and other related accessories to ensure that the device can be used normally and safely.
- The syringe pump does not support air-in-line detection. Always purge air from the system before each use.
- Pay extra attention to twisting of the infusion line when it is used for low-rate infusion. A lower infusion rate indicates a longer time from occurrence to detection of the occlusion, which may suspend the infusion for a long time.
- To avoid the risk of fire or explosion, do not use the syringe pump in a flammable or oxygenated environment.
- The altitude difference between the pump and heart position of the patient should not be larger than 100cm. A smaller altitude difference indicates a higher accuracy of the pressure sensor's result.
- In the event of tube twisting, filter condensation or intubation occlusion during infusion, the internal pressure of the infusion tube increases. Once the causes for occlusion are removed, too much infusion liquid may be infused into the patient. Therefore, proper actions should be taken. For example, clamp the infusion tube before removing the occlusion causes, because the sliding syringe tube is loose.
- To guarantee the infusion safety and proper alarm function, it is recommended that you use only the syringes specified by the manufacturer.
- If a syringe of other brands is used or the syringe parameters are not defined correctly, the infusion accuracy may be affected.
- Only the syringe, tube, syringe needle and other medical parts complying with the local regulations can be used on the syringe pump. Contact your local distributor for more

Precautions for Use

information.

- Operating the syringe pump against the requirements, procedures, warnings or cautions provided in this manual may cause infusion failure, inadequate or excessive dosing, and/or other potential risks.
- There should be a regular monitoring by clinical professionals when using the device.
- The power cord and other affiliated lines should be kept properly to prevent patients from being tripped and avoid electromagnetic interference.
- High-frequency surgical equipment, mobile phone, wireless device and defibrillator may have interference on the syringe pump. Keep away from such devices while operating.
- To avoid the risk of electric shock, this equipment must only be connected to supply mains with protective earthing.
- If the pump and its related accessories are reaching over the life time, they must be scrapped and disposed in accordance with the local laws or hospital ordinances. Please contact your local representative for further details.
- Do not replace any component of this equipment without authorization of the manufacturer.
- When operating the pump or checking the pump's alarm system, the operator shall be in front of the device, no farther than 1 meter.
- There is no patient circuit in this device. The output of the equipment is not allowed to be accessible to patient.
- The operator shall not touch MP-30 Series and the patient simultaneously.



CAUTION:

- The syringe is treated as an applied part of the pump.
- Infusion should be started only when the values on the prescription are the same as the values set on the syringe pump.
- Ensure that the syringe pump is fixed tightly on the stand and the stand is stable.
- Prevent the pump from collision, dropping, mechanical vibration and other external forces to avoid damage on the pump.
- Before pressing the [Start] key, check if the infusion rate is correct, especially the position of the decimal point.
- Do not operate on the screen with sharp objects. Otherwise, the screen may be damaged.
- An occlusion alarm may be generated when high-viscosity liquid is infused at high rate through a thin intravenous needle. In this case, increase the occlusion level or decrease the infusion rate.
- Syringe pump should be placed without the reach of patients and other irrelevant personnel.

Precautions for Use

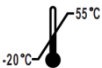
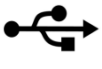
- Avoid direct sunlight, high temperature and high humidity.
- Do not autoclave the syringe pump.
- Before using the internal battery, check the battery to ensure that sufficient power is available. Recharge the battery if required.
- Ensure that a battery is installed in the syringe pump before operation. Otherwise, the system may stop working without reporting an alarm when external power is interruption due to power failure or a short circuit, causing an unsafe condition.
- If the syringe pump cannot work as described in this manual for unknown reasons, stop it and report the details (including syringe, infusion flow, serial number of syringe pump, and type of infusion liquid) to your local distributor or our customer service department.
- Do not disassemble or reconstruct the syringe pump.
- Liquid intrusion into the AC power socket, USB or nurse call socket may cause short-circuit. While connecting the power cable, check if the connecting parts are dry. If liquid spills on the syringe pump, clean the pump with a dry cloth and use the pump after inspection by maintenance personnel.
- The maximum temperature at the applied part of the pump may reach 42.2 °C when the pump runs continuously under the highest environment temperature at the highest infusion rate.
- The highest pressure at the end of the infusion tube may be up to 3000mmHg under the condition of occlusion
- The syringe pump may not generate an infusion pressure that exceeds the maximum occlusion level 1175mmHg.
- The delay time between the onset of the alarm condition and the representation of the alarm is not longer than 150ms.
- After the pump is exposed to a defibrillation voltage, the recovery time of the pump is shorter than 1s (the pump functions properly during exposure to the defibrillation voltage)

Precautions for Use

Symbols:

	Model number
	Medical device
	Authorized Representative in the European Community
	CE Mark: conforms to essential requirements of the Medical Device Directive 93/42/EEC.
	Date of manufacture
	Manufacturer
	Serial number
	Unique device identifier
	Defibrillation-proof type CF applied part
	Alternating current
	Direct current
	DISPOSAL: Do not dispose this product as unsorted municipal waste. Separate collection of such waste for special treatment is necessary.
	CAUTION! Read the accompanying document.
	General warning sign.
	Refer to the operation manual.
IPX2	Protected against vertically falling water drops when enclosure tilted up to 15 °
	Interference may occur near the devices with below sign.
	Nurse call
	ON/OFF

Precautions for Use

	HOME
	Protective earth.
	This way up
	Fragile, handle with care
	Keep dry
	Keep away from sunlight
	Atmospheric pressure limitation
	Temperature limit
	Humidity limitation
	Stacking limit by number (n is 5, 9..., for the specific value, see the product package box.)
	USB2.0 interface

Product Specifications

3 Product Specifications

Product name	Syringe pump
Model	MP-30 Series
Power supply	AC power supply: AC 100-240V, 50/60 Hz, power consumption less than 45 VA External DC power supply: DC 12 V 1A Internal battery: lithium battery 7.2 V 2600 mAh Battery model: 18650-2S1P-02 Continuous use time of battery: not less than 6 hours (for infusion at 5 ml/h with a new battery at room temperature)
Compatible syringes	All syringes of 5ml(Only applicable to MP-30T and MP-30A), 10ml, 20ml, 30ml, and 50/60ml conforming to the standard
Infusion mode	MP-30A : Rate, Time, Weight, Loading Dose, Trapezia, Sequence, Micro, TIVA, Relay (used in combination with the infusion workstation) MP-30 and MP-30T: Rate, Time, Weight, Sequence and Relay (used in combination with the infusion workstation)
Infusion setting range	0.1-100.0ml/h(5ml syringe) 0.1-300.0ml/h(10ml syringe) 0.1-600.0ml/h(20ml syringe) 0.1-900.0ml/h(30ml syringe) 0.1-2000.0ml/h(50/60ml syringe) See the least increment in chart 6-3
VTBI setting range	0.1 - 99.99ml(minimum increment: 0.01) 100 - 999.9ml(minimum increment:0.1) 1000 – 9999ml(minimum increment:1)
Total volume display	0.00-9999.99ml
Accuracy	Mechanical accuracy: $\pm 0.5\%$ Accuracy including syringe: $\pm 2\%$
Infusion time setting range	00:00:01 ~ 99:59:59(h:m:s) (minimum increment: 1s)
Purge operation	100.0ml/h (5ml syringe) (Only applicable to MP-30T and MP-30A)

Product Specifications

	300.0ml/h (10ml syringe) 600.0ml/h (20ml syringe) 900.0ml/h (30ml syringe) 2000ml/h (50/60ml syringe)
Bolus operation	0.1-100.0ml/h(5ml syringe) (Only applicable to MP-30T and MP-30A) 0.1-300.0ml/h(10ml syringe) 0.1-600.0ml/h(20ml syringe) 0.1-900.0ml/h(30ml syringe) 0.1-2000.0ml/h(50/60ml syringe) Automatically calculate the bolus rate by bolus amount, cannot lower than the current rate.
Bolus VTBI	(0.10 ~ 5.00)ml(5ml syringe) (minimum increment:0.01ml) (Only applicable to MP-30T and MP-30A) (0.10 ~ 10.00)ml(10ml syringe) (minimum increment:0.01ml) (0.10 ~ 20.00)ml(20ml syringe) (minimum increment:0.01ml) (0.10 ~ 30.00)ml(30ml syringe) (minimum increment:0.01ml) (0.10 ~ 50.00)ml(50ml/60ml syringe) (minimum increment:0.01ml)
Anti-Bolus	Anti-bolus function, unintended bolus ≤ 1.0 ml
KVO rate	0.1~5ml/h
Occlusion level	MP-30A : 225mmHg ~ 975mmHg, 11 levels are available MP-30 and MP-30T : 300 mmHg ~ 900mmHg , 3 levels are available
Alarm	Near Finished, Finished, Syringe Empty, Near empty, OCCL, Low Battery, Battery Empty, No Battery, No Power Supply, Unknown Syringe, Syringe Install Error, Standby Time Expired, Relay Index Duplicate, Syringe Start Fail, Reminder Alarm, OCCL Risk, Tube Off (Only suitable for 30ml, 50/60ml syringes)
Special function	Repeat alarming: After the sound of an alarm is muted, this alarm is reported again two minutes later if it persists. Event recording: A maximum of 2000 events can be stored for playback.

Product Specifications

	Sound volume: 10 levels are available. Power supply switching: When AC/DC power supply is cut off, the syringe pump automatically switches to internal battery supply. Barcode scanning: Input the patient information by barcode scanning.
Wi-Fi function	Connect Infusion Central Monitoring System to monitor the infusion status.
Drug Library(only for MP-30A)	A maximum of 2,000 drug types can be stored.
Operating conditions	Temperature: 5 ℃ to 40 ℃ Humidity: 15% to 95% RH, non-condensing Atmospheric pressure: 70.0 kPa-106.0 kPa
Operating Altitude	Not higher than 3000m
Storage and shipping conditions	Temperature: -20 ℃ to +55 ℃ Humidity: 10% to 95% RH, non-condensing Atmospheric pressure: 22.0 kPa-107.4 kPa
Classification	1. Class I / Internally powered equipment; 2. Defibrillation-proof type CF applied part; 3. IPX2; 4. No sterilization requirement for pump 5. Not category AP / APG equipment; 6. Mode of operation: continuous
Dimensions	252(W) ×74(H) ×164(D)mm
Weight	About 1.6 kg (including battery)
Service Life	10 years
Main safety standards	IEC60601-1 Medical electrical equipment-Part 1: General requirements for basic safety and essential performance IEC60601-2-24 Medical electrical equipment-Part 2-24: Particular requirements for the safety of infusion pumps and controllers

Product Specifications

	IEC60601-1-8 Medical electrical equipment-Part 1-8: General requirements for basic safety and essential performance -- Collateral standard: General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems IEC60601-1-2 Medical electrical equipment-Part 1-2: General requirements for basic safety - Collateral standard: Electromagnetic compatibility requirements and tests
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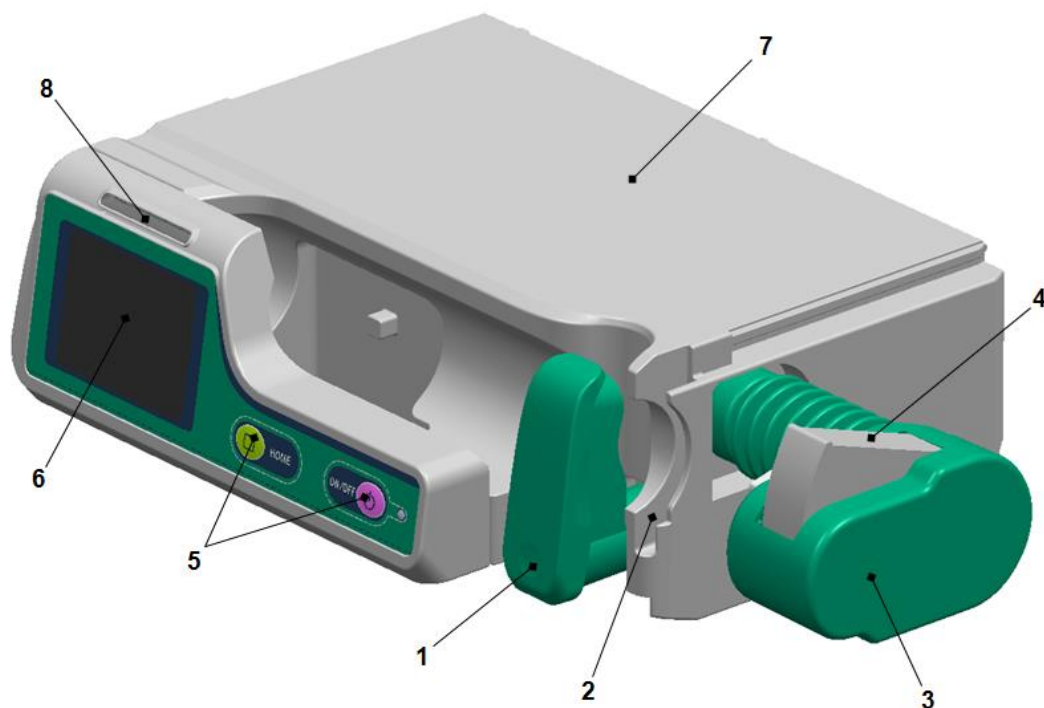
4 Product Description

4.1 Operating Principle

The MP-30 Series syringe pump mainly consists of the pump shell, display and operating system, monitoring system, alarm system, motor drive system, drive module, power supply system, Wi-Fi communication module (optional), handle (optional) and pole clamp (optional).

The syringe pump adopts the dual processor structure, controls the motor precisely, drives the peristaltic sheet to infuse through the mechanical drive device, monitors the sensors and infusion process, and provides sound and light alarms.

4.2 Composition of Syringe Pump



1 - Clamp

2 - Slit

3 - Slider

4 - Clutch

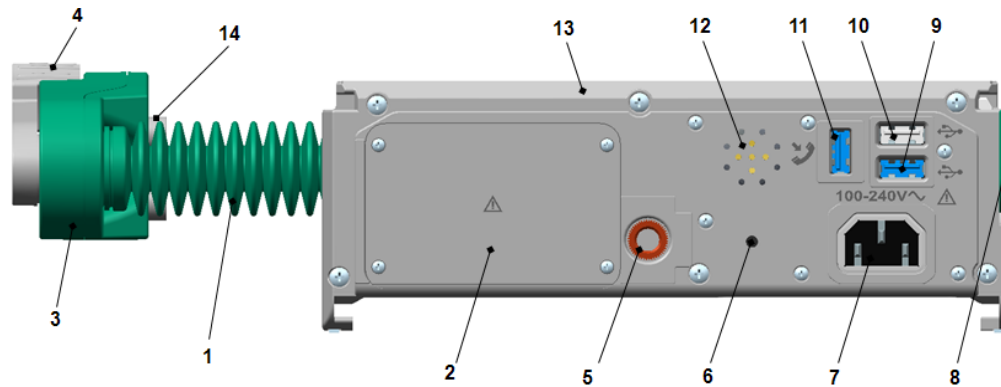
5 - Operation button

6 - Touch screen

7 - Shell

8 - Alarm indicator

Product Description



1 –Protective cover	2 –Battery chamber	3 – Slider
4 -Clutch	5 –Threaded hole	6 –Auxiliary alarm
7 –AC power inlet	8 -Multi-function connector	9 –External inlet 1
10 –External inlet 2	11 –External inlet 3	12 –Buzzer
13 –Shell	14 –Syringe clamp	

- Battery chamber. Replaceable battery inside the chamber.
- Threaded hole. To fix the pole clamp, then fix the pump to the IV pole via the pole clamp.
- Auxiliary alarm. Audible alarm sounds when product functions abnormally.
- Buzzer. To alarm in high, medium or low level during infusion.
- AC power inlet. To connect the external AC power source.
- External inlets 1: Used to connect barcode scanner, wired networking module (optional), DC power supply line and infusion workstation. Data can be transmitted externally through the wired networking module.
- External inlets 2: Used to connect wired networking modules.
- External inlets 3: Used to connect the nurse caller.

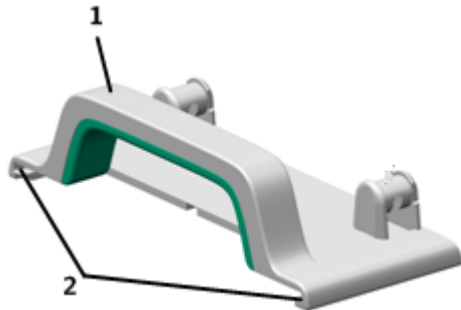
CAUTION:

- Only the accessories or devices specified by the manufacturer are allowed to be connected to the Pump. Otherwise, an electric shock may occur. See Table 4-1.
- Additional equipment connected to medical electrical equipment through the network/data coupling (USB or LAN port) must comply with the respective IEC or ISO standards (e.g. IEC 60950 and IEC 62368-1 for data processing equipment). Furthermore all configurations shall comply with the requirements for medical electrical systems (see clause 16 of the 3Ed. of IEC 60601-1).
- Anybody connecting additional equipment to medical electrical equipment configurations a medical system and is therefore responsible that the system complies with the requirements for medical electrical systems. Attention is drawn to the fact that local laws take priority over the above mentioned requirements. If in doubt, consult your local representative or the technical service department.

Product Description

- The plug is used to disconnect from the mains supply. Install the pump at a position where an operator can insert and remove the plug conveniently.

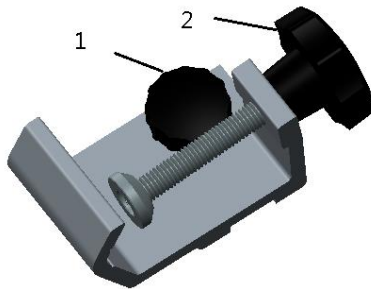
4.3 Handle



1 –Handle

2 –Slide rail

4.4 Pole Clamp



1 - Mounting screw

2 - Mounting knob of infusion stand

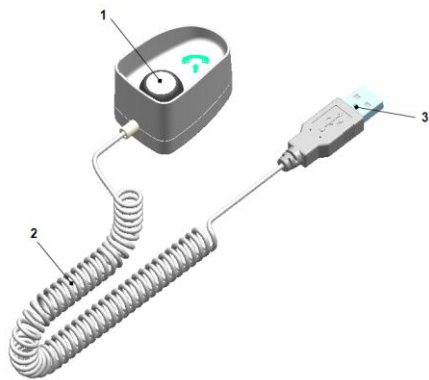


CAUTION:

- The diameter of the drip stand should be 16~36mm. Any size beyond this range may cause insecure installation.
- Please ensure that the floor is solid and level. Don't fix other equipment on the IV pole to prevent the pole from falling over.
- During installation, ensure that the knob and mounting screw are tightened up.

Product Description

4.5 Nurse Call



1 –Button

2 –Cable

3 –Socket

4.6 Accessories Accompanied

1 –AC power cord

1

2 - Pole clamp

1

3 – Handle

1

4 - Operation manual

1

5 - Quick operation instruction

1

6 -Packing list

1

4.7 Optional Accessories

Table 4-1 List of optional accessories

Options	Description
Power cable	Standard configuration by factory
Lithium battery pack	7.2V @ 2600mAh
Handle	MP-1
Nurse pager	MP-2
Barcode scanner	MP-4
Pole clamp	——
DC power cable	——

5 Preparations for Use

- Before using the syringe pump, read carefully the operation procedures and precautions in this manual.
- Before using the syringe pump for the first time, set the date and time to ensure that history can be recorded correctly.
- Before using the syringe pump for the first time, set the brand of syringe.
- Before using the syringe pump for the first time, recharge the internal battery fully. If the syringe pump is off, the battery can be charged fully at least 10 hours after being connected to an external power supply.
- Place the syringe pump on a stable platform.
- Alternatively, use the provided pole clamp to mount the syringe pump on an infusion stand.
 - Put the syringe pump on the pole clamp while aligning the retaining knob with the threaded hole, and rotate the handle to fix the syringe pump on the pole clamp.
 - Clamp the pole clamp on the infusion stand, adjust the syringe pump to an appropriate position, and tighten the retaining knob for infusion stand on the pole clamp.
- Connect external power supply.
 - Insert the supplied AC power cord into the AC inlet on the right side of the syringe pump. Plug the cord into an AC power outlet with grounding terminal.
 - To power the syringe pump with external DC power supply, contact your local distributor for help.



CAUTION:

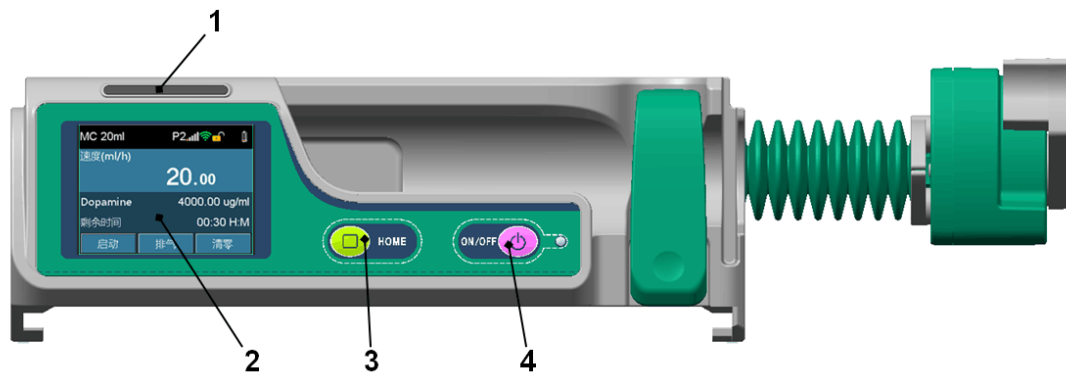
- Do not install the syringe pump at a place not convenient for an operator to connect the AC power.

Operating Instructions

6 Operating Instructions

6.1 Display and Keys

● Display



1 –Alarm indicator

2 –TFT touch screen

3 –Home key

4 –ON/OFF key

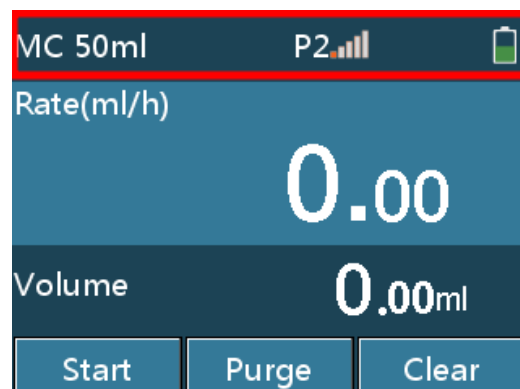
The alarm indicator indicates the alarm severity (high, medium, or low) with three colors: red, yellow and green. The alarm severity is determined based on the criticality of infusion information.

TFT touchscreen, resolution: 320x240 pixels

The display is divided into three areas: information area, work data area and function key area.

See below for further description.

Information area: Displays the syringe brand and specifications, occlusion pressure level, real-time pressure, external power supply, battery volume, and Wi-Fi signal. Touch the brand and specifications zone to enter the page for syringe brand adjustment. Touch the occlusion pressure level to enter the page for occlusion level selection.



Operating Instructions

See below for further description.

P2 Occlusion pressure level: 2

 Real-time occlusion pressure: Five bars in total. A larger number of illuminated bars indicates a higher pressure.

 External power source symbol. This symbol is displayed when the pump is connected to an external AC/DC power source.

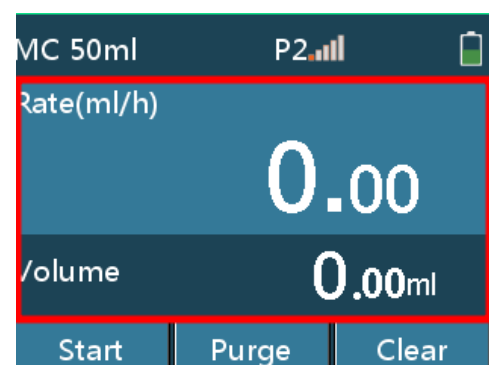
 Screen lock symbol. Two states exist: lock and unlock.

 Battery volume and charging status. Four bars in total. A larger number of illuminated bars indicates a higher remaining battery capacity.

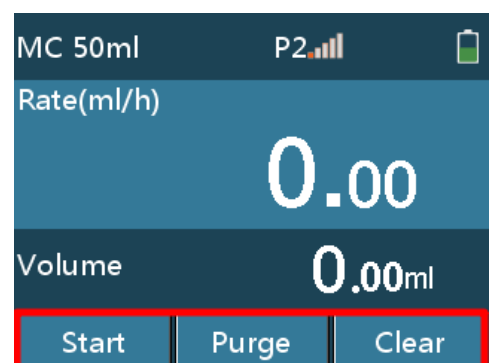
 Wi-Fi signal

 This symbol is displayed when the pump is connected to the workstation.

Work data area: Displays the current infusion rate and infusion volume or displays different infusion work data based on different infusion modes. The work data can be adjusted by touching the specific zone in different working modes.



Function key area: Displays such keys as [Start], [Purge], [Clear], [Stop], and [Bolus]. Setting keys such as numbers and letters appear on corresponding interfaces.



Operating Instructions

Keys

In addition to touchscreen keys, two keys are available on the key panel: [HOME] and [ON/OFF].

- [HOME]: Main menu key. Before infusion, press [HOME] once to enter a setting menu, such as patient information, syringe setting, event recall, device setting, etc. To return to the infusion preparation screen, press [HOME] once again on any setting interface.
- [ON/OFF]: Key for switching on/off the pump. When the pump is off, press and hold [ON/OFF] for 1 second to switch it on. When the pump is on, press [ON/OFF] and select [Power Off] or press and hold [ON/OFF] for 3 seconds to shut down the pump.

6.2 Powering On the Syringe Pump

CAUTION:

- Power on the pump and then install the syringe.
 - Press [ON/OFF] to start the pump.
 - The self-test starts and startup interface appears.
 - After self-test finishes, the infusion preparation screen is displayed.
 - The screen displays patient information, infusion brand and occlusion level stored last time the device is powered off.
 - If the self-test is abnormal, corresponding information is displayed in the information area.

Infusion preparation interface:



WARNING

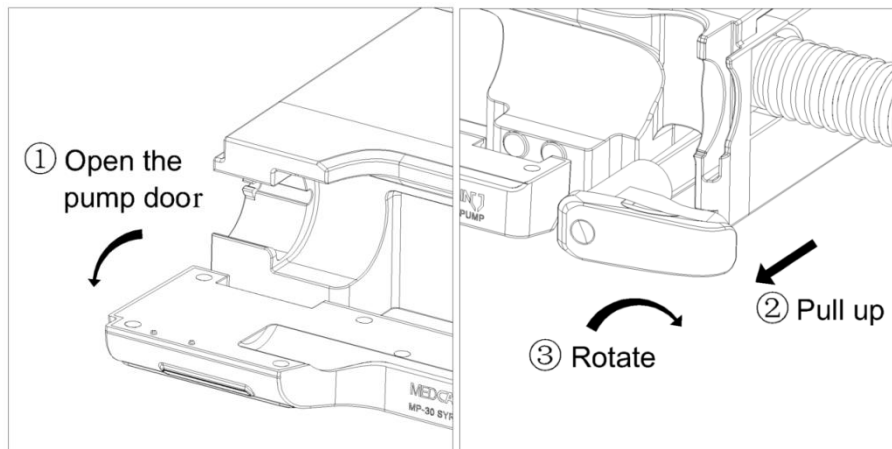
- After the pump is powered on, confirm that the loudspeaker and alarm indicator work properly. In addition, check if the self-test is finished and no error messages appear. (Refer to Chapter 8 Troubleshooting.)
- Ensure that the displayed syringe brand and specification are consistent with those of the installed syringe. Otherwise, infusion accuracy is not guaranteed and the alarm function may fail.
- If the syringe brand set differs from the brand of the syringe actually used, the infusion

Operating Instructions

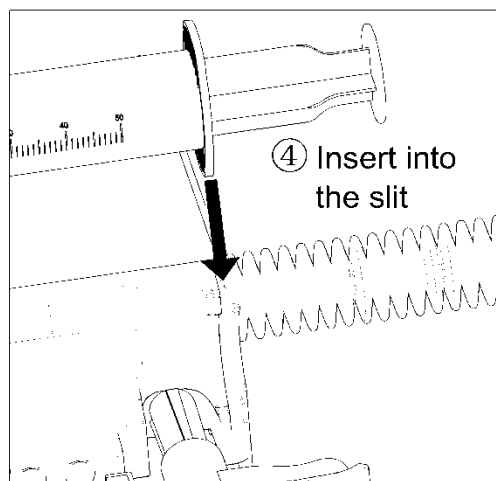
accuracy and alarm function cannot be guaranteed.

6.3 Syringe Installation

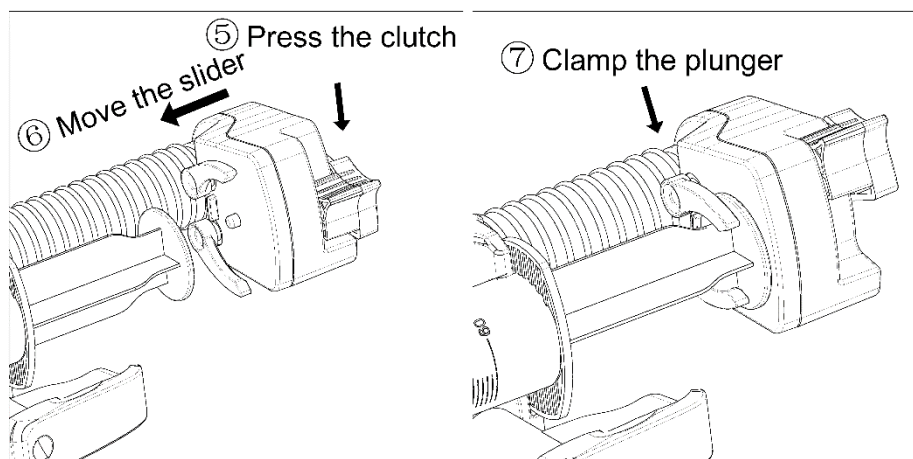
- Open the pump door, pull the clamp and turn rightwards (①/②/③)



- Press the clutch and move the slider fully to the right.
- Attach the syringe. Insert the flange of the syringe into the slit (④)

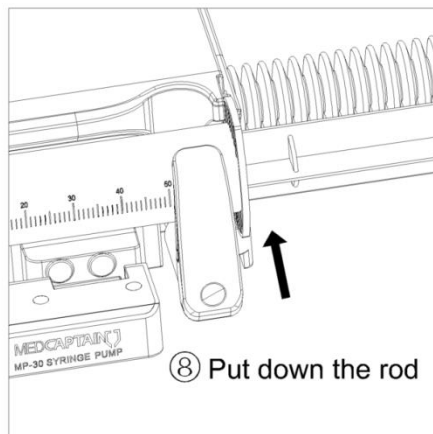


- Press the clutch, and move the slider until the contact pin of the slider hits the syringe plunger. (⑤/⑥ /⑦)

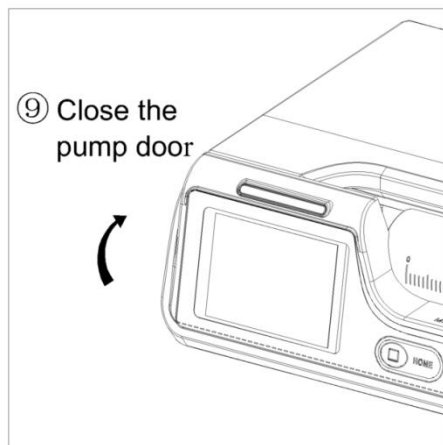


Operating Instructions

- Turn back the clamp and lower it slowly to hold the syringe securely (⑧).



- Close the pump door (⑨).



WARNING

- Ensure there are no air bubbles in the syringe.
- If the syringe flange is not properly installed in the slit, flow rate accuracy and alarm function cannot be guaranteed.

6.4 Purging

WARNING:

- After loading a syringe on the syringe pump, remove the air bubbles from the syringe and the IV line.
- Before purging the IV line, ensure that the IV line is not connected to patients.
- Purging can be done only in non-infusion process.
- Stop purging after ensuring that liquid is drained out from the needle.

Operating Instructions

- Press [PURGE], the purge interface pops up. Click [Stop], the purge stops.



- The green indicator flashes during purging.

CAUTION:

- When high viscosity IV fluids are infused through thin vein needle by bolus operation, occlusion alarm may be generated. In this case, reduce the infusion rate to purge.
- The purge function can be used to remove any mechanical gap. Otherwise, a considerable delay may be caused in the start of the infusion.
- Total volume cannot be cleared after infusion starts.
- The volume under the purge function will not be calculated into the total volume.

6.5 Setting the Infusion Rate

- Press the rate area on the touchscreen to enter the setting interface.

Rate (ml/h)			
20.0_			
1	2	3	←
4	5	6	C
7	8	9	Cancel
.	0	Shift	Confirm

- Press the [CLEAR] key to reset total volume.



CAUTION:

- If you enter an infusion rate that is out of the valid range, the infusion rate is invalid and you need to enter again.
- When a syringe of different size is installed, if the flow rate is out of range, please reset the rate to maximum valid rate.
- The flow rate can be set while the pump is infusing.

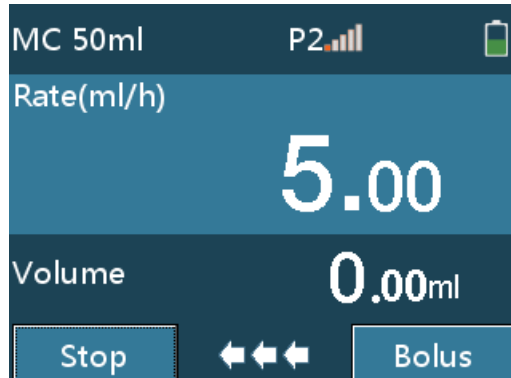
Operating Instructions

6.6 Puncture

Insert an IV cannula into the patient's vein. Connect the cannula to the syringe in the pump.

6.7 Starting Infusion

Click the [START] key to start infusion at the setting rate. The green indicator flashes.

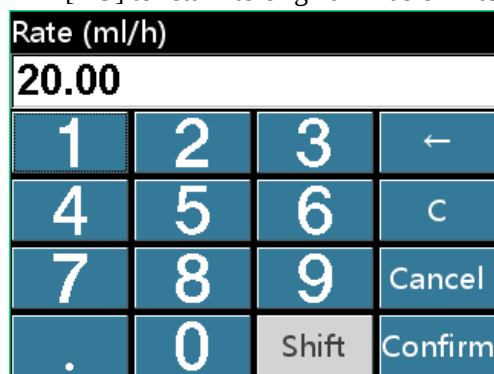


CAUTION:

- Infusion should be started only when the values on the prescription are the same as the values set on the syringe pump.
- If no operation is performed after syringe installation for more than 2 minutes, the START-REMINDER alarm is reported.

6.8 Changing Rate During Infusion

- Click the rate display area on the screen, click [OK] on the pop-up interface, and then click [NO] to return to original infusion interface.



- After entering the rate, if you click [Cancel], the system returns to the original infusion interface without change. If you click [Confirm], the system returns to original infusion interface and operates at the new rate.

CAUTION:

- If no operation is performed on the reference or rate setting interface for more than 10 seconds, the system returns to the infusion interface automatically.

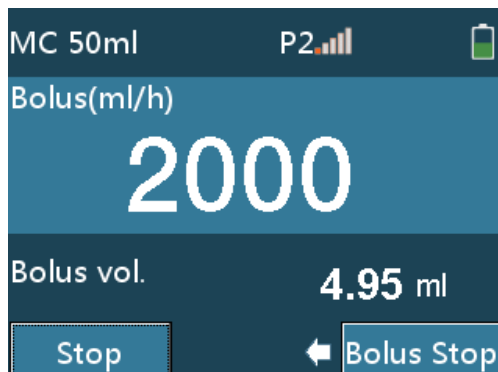
Operating Instructions

6.9 Bolus

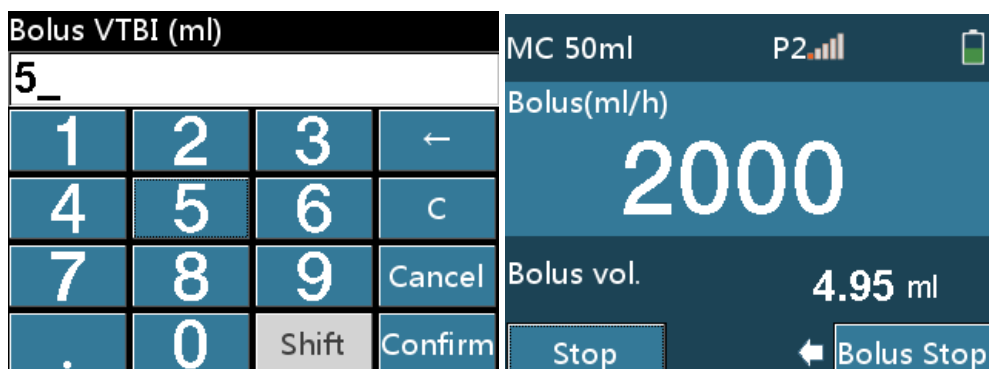
MP-30T and MP-30A: Support three bolus modes, that is, manual bolus, rapid quantitative bolus, and automatic bolus.

MP-30: Only supports automatic bolus.

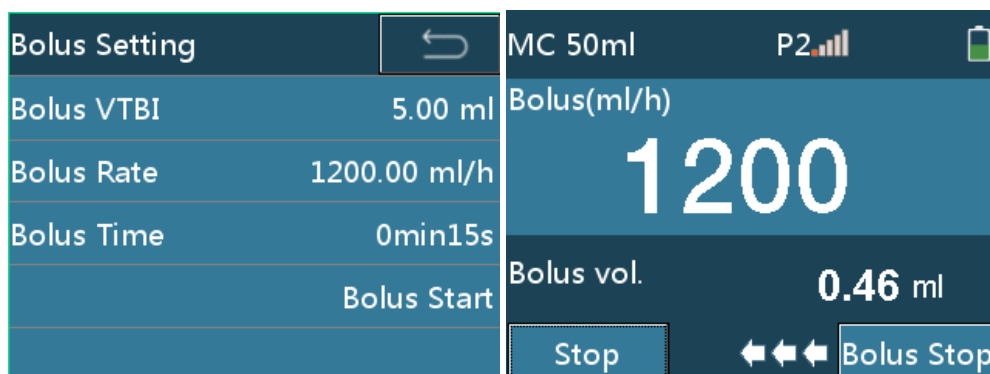
- Choose “Manual bolus”, during the infusion, press and hold [Bolus] for 1s to enter the bolus interface. Bolus continues while the button is pressed and held and stops immediately when the button is released.



- Choose “Rapid quantitative Bolus”, during the infusion, click [Bolus] to enter the bolus VTBI interface, set the bolus volume, click [Confirm] to start and click [Bolus Stop] to stop the bolus and return to infusion interface.



- Choose “Automatic Bolus”, during infusion; click [Bolus] to enter the bolus setting interface. Set any two of Bolus VTBI, Bolus rate and Bolus Time, click [Bolus Start] to enter the bolus interface, click [Bolus Stop] to stop the bolus



Operating Instructions



CAUTION:

- Current bolus volume is displayed when bolus is running.
- Bolus volume will be accumulated into the total volume.

6.10 Stopping Infusion

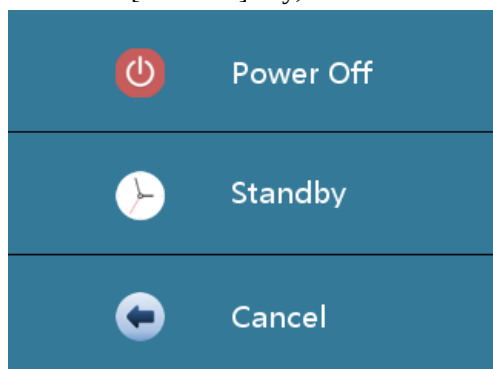
In the infusion process or after infusion, click [STOP] to stop the operation.

6.11 Replacing Syringe

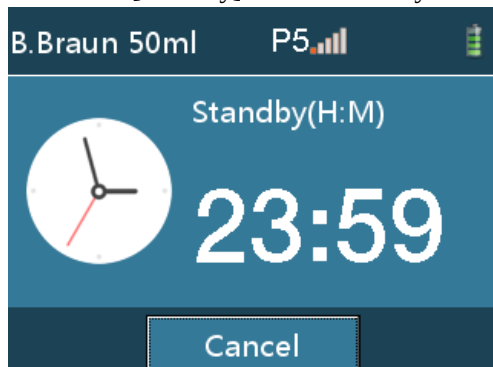
If the amount of solution in the syringe is getting low, repeat the steps in sections 6.3 and 6.6 to replace the syringe.

6.12 Powering Off the Syringe Pump

- Press [ON/OFF] key, and choose Power Off, Standby or Cancel.



- Click [Power Off] to shut down.
- Click [Standby] to enter standby interface, the standby time can be modified.



- Click [Cancel] to return to the previous interface.

7 Setting the Syringe Pump



- After the pump is powered off, all parameter settings will be automatically saved.
- Parts of parameters will not be saved if the device is forced to shut down.

7.1 Infusion Set

Press the [HOME] key to enter the setting interface, click [Infusion Set] to enter the detailed infusion setting interface. Infusion Mode, occlusion level, bolus mode, KVO rate, brand, relay set, micro mode, near finished and recent therapy can be set and adjusted here.

7.1.1 Infusion Mode

■ Rate mode

In Rate Mode, set the drug name, rate and VTBI, and click [Confirm] to operate.

Rate Mode	
Drug Name	None
Rate	---- ml/h
VTBI	---- ml
Volume	38.16 ml
Confirm	

■ Time mode

In Time Mode, set the drug name, VTBI and Time, and click [Confirm] to operate.

Time Mode	
Drug Name	None
VTBI	---- ml
Time	--h--min
Rate	---- ml/h
Confirm	

■ Weight mode

In Weight Mode, set the drug Info, Dose Rate, Weight and VTBI, wait for the device to calculate the rate automatically, and then click [Confirm] to operate.

Setting the Syringe Pump

Weight Mode	
Drug Info	---- mg/ml
DoseRate	---- ug/kg/min
Weight	---- kg
VTBI	---- ml
Confirm	---- ml/h

■ Trapezia mode

In Trapezia Mode, set the drug name, VTBI, rate, rise time and fall time, wait for automatic calculation of the rate, and then click [Confirm] to operate

Trapezia Mode	
Drug Name	None 
VTBI	---- ml
Rate	---- ml/h
RiseTime	--h--min
FallTime	--h--min 

■ Loading Dose mode

In Loading Dose mode, set the drug name, VTBI, maintain rate, loading rate and loading time, wait for automatic calculation of the rate, and then click [Confirm] to operate

LoadingDose Mode	
Drug Name	None 
VTBI	---- ml
MaintainRate	---- ml/h
LoadingRate	---- ml/h
LoadingTime	-- h-- min 

■ Sequence mode

In Sequence Mode, set the drug name, many groups of sequence rate, and time, and then click [Confirm] to operate in sequence.

MP-30: Supports 5 groups of sequence.

MP-30T and MP-30A: Support 10 groups of sequence.

Setting the Syringe Pump

Sequence Mode		↩
Drug Name	None	▲
Rate1	---- ml/h	
Time1	--h--min	
Rate2	---- ml/h	
Time2	--h--min	▼

■ TIVA mode

In TIVA Mode, set the drug info, loading dose, loading time dose rate and weight, wait for automatic calculation of the rate, and then click [Confirm] to operate.

TIVA Mode		↩
Drug Info	---- mg/ml	▲
LoadingDose	---- mg/kg	
LoadingTime	--h--min	
DoseRate	---- ug/kg/min	
Weight	---- kg	▼



CAUTION:

- The drug library function (only for MP-30A) can be applied to all working modes. The therapy data of the drug library is not edited by the manufacturer.

Setting the Syringe Pump

7.1.2 Occlusion Level

- MP-30A: Eleven occlusion levels are available (Factory setting is level 6).
- MP-30 and MP-30T: Three occlusion levels are available (Factory setting is level 2).

Table 7-1 MP-30A Relationship between occlusion level and pressure

Occlusion level	Display	Pressure (mmHg)	Pressure (KPa)	Pressure (bar)	Pressure (psi)
1	P 1	225	30	0.3	4.35
2	P 2	300	40	0.4	5.80
3	P 3	375	50	0.5	7.25
4	P 4	450	60	0.6	8.70
5	P 5	525	70	0.7	10.15
6	P 6	600	80	0.8	11.60
7	P 7	675	90	0.9	13.05
8	P 8	750	100	1.0	14.50
9	P 9	825	110	1.1	15.95
10	P 10	900	120	1.2	17.40
11	P 11	975	130	1.3	18.85

Table 7-2 MP-30 and MP-30T Relationship between occlusion level and pressure

Occlusion level	Display	Pressure (mmHg)	Pressure (kPa)	Pressure (bar)	Pressure (psi)
1	P 1	300	40	0.4	5.80
2	P 2	550	73	0.7	10.64
3	P 3	900	120	1.2	17.40



CAUTION:

- When an occlusion alarm is reported, the motor automatically rotates reversely to release the tube pressure (Anti-Bolus) for preventing infusion of extra amount of pills into a patient after the occlusion alarm is cleared. When the tube pressure drops to 20% of the occlusion pressure, Anti-Bolus automatically stops.

Setting the Syringe Pump

- When you infuse viscous solution with the Occlusion Level setting under 4 and the tubing is clear, occlusion alarm tends to be generated. Carefully observe the  symbol in the upper information area, and change the occlusion level if more than 2 bars are illuminated.
- When you operate the pump with the Occlusion Level setting over 8, the in-line pressure builds up substantially until Occlusion alarm is generated. Always make sure that the IV line is securely connected to the syringe.
- An occlusion alarm may be generated when high-viscosity liquid is infused at high rate through a thin intravenous needle. In this case, increase the occlusion level or decrease the infusion rate.
- The maximum infusion pressure produced by the syringe pump does not exceed 1125mmHg.

Setting the Syringe Pump

7.1.3 Bolus Mode

- Three bolus modes are available: Manual Bolus, Rapid quantitative Bolus and Automatic Bolus. Please refer to the chapter 6.9 for further instructions.

7.1.4 KVO Rate

- KVO-rate can be adjusted from 0.1ml/h to 5ml/h (Step by 0.01ml/h). The default rate is 1ml/h.

7.1.5 Brand

- You can choose the consumable brand in the following sequence: [Home] -> [Infusion Set] -> [Brand]
- Several brands syringe of 5ml (Only applicable to MP-30T and MP-30A), 10mL, 20mL, 30mL, and 50/60mL have been preset and customized. Select the syringe accordingly for clinical use.



CAUTION:

- Users must use the consumable brand which is specified by the manufacturer.
- To add a syringe of other brand, users are strongly recommended to contact the manufacturer or manufacturer's representative to set and test, so as to ensure the infusion accuracy.

7.1.6 Relay Setting

- In relay mode, switch on relay and set the relay number.

Relay Set	
Relay On-Off	☒
Relay Index	0
Enable Relay Cycle	☐



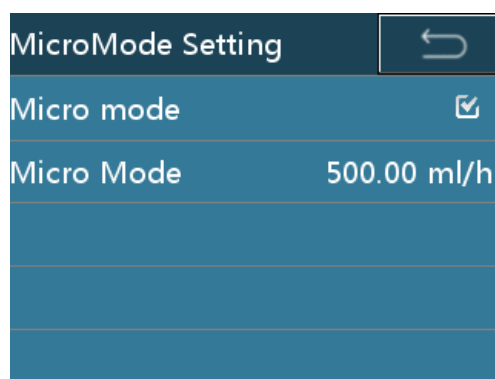
CAUTION:

- The relay number must be set in sequence from number 1 when there are several syringe pumps or infusion pumps.
- When “Enable Relay Cycle” is activated, infusion will cycle from the first pump after the last pump finishes the infusion.

7.1.7 Micro Mode Setting

- After Micro Mode is selected, the maximum rate can be set in the mode.

Setting the Syringe Pump



7.1.8 Near Finished

- The Near Finished alarm is generated when infusion is almost completed. The duration from generation of this alarm to infusion completion can be adjusted from 1 min to 30 min (adjustment step: 1 min). By default, this alarm is generated 3 minutes before infusion completion

7.1.9 Recent Therapy

- The therapies of the recent 20 times are recorded. The recorded therapy can be started directly by simple selection.



7.2 System Set

7.2.1 Volume Setting

- Ten volume levels are available (the factory setting is level 5).

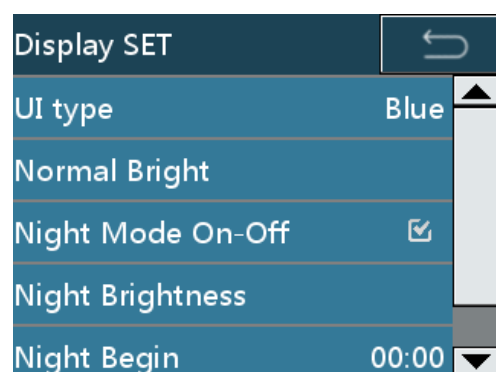


CAUTION:

- Do not set the alarm volume to a level lower than the ambient noise to ensure that the alarm can be recognized correctly.
- The alarm system may fail when the alarm volume is set to an extreme value. Check alarm limit values based on clinical conditions.
- After the pump is inserted into a workstation, when the volume setting on the pump changes, the volume setting on the workstation changes simultaneously.

Setting the Syringe Pump

7.2.2 Display SET



- Seven different color options are available for UI type.
- The brightness can be adjusted in [Normal Bright].
- All the parameters of the night mode can be adjusted here.

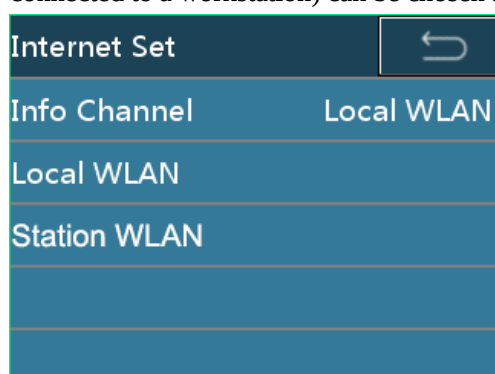


CAUTION:

- In night mode, the setting range of start time is 17:00-09:00, and the setting range of finish time is the same as that of start time. By default, the start time is 00:00 and finish time is 00:00.
- After the pump is inserted into a workstation, when the display setting on the pump changes, the display setting on the workstation changes simultaneously.

7.2.3 Internet Set

- [Info Channel], [Local WLAN], and [Workstation WLAN] (not available if the pump is not connected to a workstation) can be chosen and set.



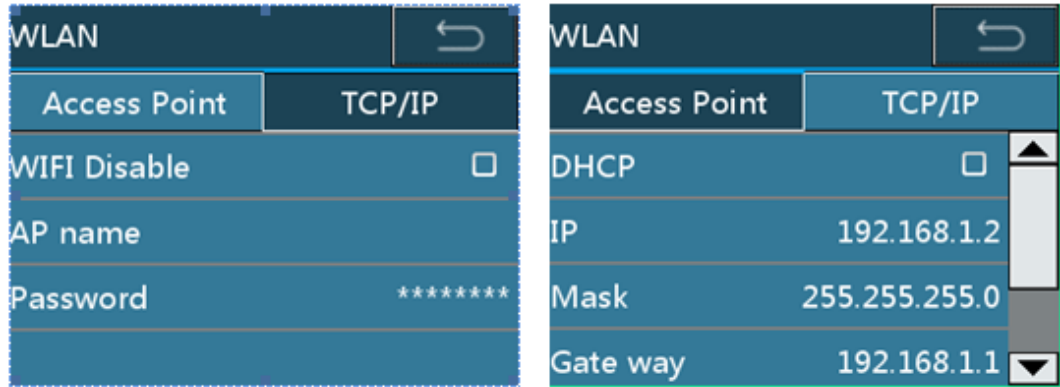
- Click [Info Channel] to choose the channel type
 - Choose [Local WLAN] to use local WLAN channel to connect to the network, and the local WLAN parameters can be set
 - Choose [Station WLAN] to use station WLAN channel to connect to the network, and the station WLAN parameters can be set
 - Choose [Local RS485] to use local RS485 cable to connect to the network
 - Choose [Station RS485] to use station RS485 cable to connect to the network

Setting the Syringe Pump



 CAUTION:

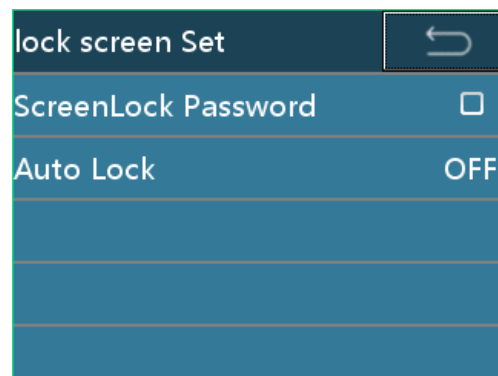
- The pump can communicate with working station, and the working station can manage the alarm and implement relay function between pumps.
- The setting of [Local RS485] and [Station RS485] must be done by the manufacturer's representatives. Please contact the manufacturer or your local distributor for further information.
- Only the accessory and devices supplied or specified by the manufacturer are allowed to be connected to the pump. Otherwise, pump exception and other unpredictable hazards may be incurred.
- Click [Local WLAN]/[Station WLAN] to set WLAN parameters.
- [Wi-Fi Disable] must be deselected, the AP name and Password of the network must be input, and the TCP/IP's information must be set.



7.2.4 Lock Screen Set

- Click [Screen Lock Password] to enable/disable the screen lock password function. When the function is enabled, a password is required to unlock the screen. When the function is disabled, no password is required to unlock the screen
- Click [Auto Lock] to set the screen auto lock function. This function can be set to OFF, 15s, 30s, 1min, 2min, 5min, 10min, or 30min. The default value is OFF, indicating that the screen auto lock function is disabled.

Setting the Syringe Pump



7.2.5 Collection Setting



- [Mode Collection]: Choose the frequently used infusion mode from the [Infusion mode] option. Once the frequently used infusion modes are chosen, the unnecessary modes will not appear in the list of 7.1.1 [Infusion mode] option. The default setting is “all the four infusion modes are chosen”.
- [Brand Collection]: Choose the frequently used syringe’s brand from the [Brand] option. Once the frequently used brands are chosen, the unnecessary brands will not appear in the list of 7.1.5 [Brand] option. The default setting is “all the preloaded brands are chosen”
- [Drug Collection]: Choose the frequently used drug from the drug library. Once the frequently used drugs are chosen, the chosen drugs will be shown on the top of the drug library list. The default setting is “none of the drug is chosen”. The data of the drug library is not edited by the manufacturer.

7.2.6 Pressure Unit

- Choose the measurement unit for the pressure. The optional units are: mmHg, kPa, bar and PSI. The default setting is mmHg

Setting the Syringe Pump

PressureUnit	
mmHg	☒
kPa	☐
bar	☐
PSI	☐

7.2.7 Date & Time Setting

- Set the date, time, and their format.

Date&Time Set	
Date Set	2016-01-28
Time Set	10:37 AM
Use 24-hour Format	☐
Select Date Format	

7.2.8 Maintenance

- Click [Maintenance] option to do the [Language Select], [Touch Adjust], [Factory Data Reset], and check the version information.
- To check the version information, follow the route of: [Home] -> [System Set] -> [Maintenance] -> [Version Info].

Setting the Syringe Pump

7.3 History

History		History:1	
01-28 10:18AM	Alarm	Time	2016-01-28 10:18:41AM
01-21 07:17PM	Alarm	Event: Alarm(Reminder Alarm)	
01-21 05:32PM	Alarm	Rate: 0.00 ml/h	
01-20 03:58PM	Alarm	Volume: 0.00 ml	
01-20 03:55PM	Alarm		

The history records are listed in Table 7-2.

Table 7-2 History records

Event	Record Parameters
Start up	Occurrence time
Shutdown	Occurrence time
Standby	Occurrence time, standby set time
Start	Occurrence time, rate, VTBI
Bolus	Occurrence time, Bolus rate, Bolus way
Bolus stop	Occurrence time, Bolus rate, Bolus accumulated volume
Stop	Occurrence time, rate, accumulated volume
KVO	Occurrence time, accumulated volume, KVO rate
KVO stop	Occurrence time, KVO rate, KVO accumulated volume
Flow rate change	Occurrence time, Flow rate before and after change
Alarm	Occurrence time, alarm event, system trouble with trouble code
Purge	Occurrence time, purge rate, accumulated volume
Purge stop	Occurrence time, purge rate, purge accumulated volume



CAUTION:

- The history records can be saved when power is cut.
- A maximum of 2000 history records can be stored. If the amount reaches storage limit, the earliest record would be removed.
- Alarm system can't be powered off separately by an operator unless the pump is powered off. The time of powering off is captured in the history records.

Setting the Syringe Pump

7.4 Patient File

- Click [Patient File] to enter the patient file page. The [Department], [Room No.], [Bed No.] and [Patient Data] can be set.

Patient File	↩
Department	
Room No.	
BedNo.	
Patient Data	

- Click [Patient Data] option to enter patient data setting page. Choose [New] to build a new patient data and the previous patient data will be cleared automatically. Choose [Modify] to modify the current patient data.

Patient Data	↩
New	Hospital No.
Modify	Name
	Sex Male
	Age 0
	Weight 0.0 kg



CAUTION:

- If the pump is inserted to a working station, once the patient file on the pump is changed, the data on the station will be synchronized at the same time.

7.5 Using the Internal Battery

- If no AC/DC power supply is available, the internal battery operates.
- When external power stops working, the internal battery starts and the yellow indicator lights with a short alarm sound.
- Before using the pump for the first time or using the pump after the pump is not used for a long time, please charge the battery for at least 10 hours.
- The approximate remaining power in the built-in battery is displayed by [battery] indicators. During battery operation, battery discharged is shown by a decreasing number of active indicators.

Setting the Syringe Pump

- When the syringe pump is connected to any external AC or DC power supply, the charge of the built-in battery starts. When battery is charging, a lightning symbol will be displayed at the left side of the battery symbol on the screen.



CAUTION:

- If AC or DC power is connected, the battery will be recharged.
- Use AC power to charge the battery. If recharged by an external 12 VDC power supply, the battery cannot be fully charged (50% at most).
- During infusion and the pump powered by battery, if a low-battery alarm occurs, press [SILENT] to silence the alarm the alarm will repeat in two minutes, connect the pump to AC/DC power supply immediately. If battery empty alarm occurs, the silence does not function and syringe pump will stop.
- 3 minutes before the battery empty, the pump will auto power off.
- The actual battery duration may be different and affected by the ambient temperature, flow rate, external communication, etc.
- If the battery is aging, the actual battery duration may be shorter. Periodically check the battery.

7.6 Connecting to the <Infusion Central Monitoring System> (Optional)

Syringe pumps can be connected to the < Infusion Central Monitoring System >, which can obtain working states of pumps remotely.



CAUTION:

- Syringe pump must not be operated through the < Infusion Central Monitoring System>.

7.7 Nurse Call (Optional)

After syringe pump is connected to the infusion central monitoring system, patient can press nurse pager in bed, and then the infusion central monitoring system in nurse station will give out a sound tip and display patient's information in screen, so that the nurse can take care of the patient in time.

7.8 Connecting a Barcode Scanner (Optional)

After a barcode scanner is connected to the pump, the patient information, such as record No. and hospital No., can be scanned, and the patient information in the pump will update automatically by pump prompts.

The barcode scanner can scan maximum 18 figures.

8 Troubleshooting

8.1 Alarm

The syringe pump provides users with a variety of status information about itself and its injection process. If any abnormality is detected, the syringe pump generates an alarm and informs users in the form of sound, light, and character.

All the alarms on this pump are of technical type.

Considering the importance of abnormal information, alarm information is classified into three levels from the viewpoint of security: low-level, mid-level, and high-level alarms. For audio and visual expressions of alarms at three levels, see Table 8-1. The alarm volume ranges from 45 dB to 85 dB.

Table 8-1 Alarm severity and the audio and visual expressions of each level

Alarm	Sound	Light
Low-level alarm	Give out three beeps at intervals of 25 seconds.	The yellow indicator is steady on.
Mid-level alarm	Give out three beeps at intervals of 25 seconds.	The yellow indicator is flashing.
High-level alarm	Give out a series of beeps at intervals of 15 seconds.	The red indicator is flashing.

When an alarm (except **Battery Empty**) occurs, press [SILENT] to pause the alarm sound.

But the buzzer beeps again if you do not eliminate the alarm within 2 minutes.



CAUTION:

- The setting of the alarm will be saved when the power is cut. When the pump restarts from a power failure situation, the alarm setting will be reloaded to the system and remains the same as it was before the power failure.



WARNING:

- There will be a potential risk if the same or similar devices are using different alarm setting in any specialized region.

Troubleshooting

8.2 Faults and Troubleshooting

Table 8-2 Alarm symptom, alarm level, fault cause, and troubleshooting

Alarm Symptoms	Alarm level	Causes	Troubleshooting
No Power Supply	Low-level	No external AC/DC power supply is connected.	Immediately connect the AC power supply or the external DC power supply.
No Battery	Mid-level	The syringe pump has no internal battery or the internal battery operates abnormally.	Replace the internal battery.
Low Battery	Low-level	The internal battery is running critically low.	Immediately connect an AC power supply or an external DC power supply.
Battery Empty	High-level	The battery is out.	Immediately connect an AC power supply or an external DC power supply.
Near End	Low-level	The infusion is end within the Near Finished Alarm setting period.	Wait until the infusion finishes.
Occlusion Alarm	High-level	1. The syringe IV line is occluded. 2. The OCCL level is too low for high viscosity drug's infusion.	Press [STOP] to stop the injection. Check and remove the cause, continuous to inject.
Near Empty	Low-level	It takes less than three minutes to complete the infusion.	Wait till syringe is purged to be empty.
Syringe Empty	High-level	The syringe is empty.	Press [STOP] to clear the alarm.
Unknown Syringe	High-level	Syringe disengagement	Press [STOP] to clear the

Troubleshooting

Alarm Symptoms	Alarm level	Causes	Troubleshooting
		from slit during infusion.	alarm. Check if the syringe pump clamp or syringe is installed correctly.
Syringe Install Error	High-level	The slider is loosen during the infusion or the slider is not installed in a right way.	Press [STOP] to clear the alarm. Check if the holder is installed correctly.
Finished	High-level	The limit amount or the infusion time is complete	Press [STOP] to clear the alarm.
Reminder Alarm	Low-level	Forget to operate the alarm (no key operation is made two minutes after the syringe is installed).	Press any key to clear the alarm.
Standby Time Expired	Mid-level	Standby mode has ended	Press [Cancel] to exit Standby mode.
Relay Index Duplicate	High-level	Relay Index Duplicate	Reset Relay Index
Syringe Start Fail	High-level	Syringe pump cannot be started in relay mode.	Check the syringe pump, and solve the problem that causes the failure of start.
OCCL Risk	Mid-level	Occlusion pressure reaches about 70% of the set occlusion level.	Release the pressure to eliminate the alarm.
The tubing falls off (only for 30, 50/60ml syringes)	Mid-level	The syringe extension tube has fallen off.	Check if the pipeline falls off, if it is, reinstall the pipeline; if not, press the [Stop] key to continue the infusion.

8.3 Troubles and Troubleshooting

When the device is faulty, a corresponding trouble code appears on the interface and a high-level alarm is generated.

Table 8-3 Troubles and troubleshooting

Trouble code	Alarm level	Troubleshooting
Sensor Error	High-level	Record the trouble code, power off the pump, and contact manufacturer or manufacturer's representatives.
Motor Error	High-level	
Circuitry Error	High-level	
Driver COM Error	High-level	
System Error	High-level	

9 Maintenance

9.1 Cleaning and Disinfection

- Before cleaning the pump, be sure to turn off the power and disconnect the AC or DC power cables, disconnect the device from the patient.
- If there is dirt on the pump, wipe it with wet soft cloth dampened with cold or lukewarm water.
- Use a piece of dry soft cloth to clean the AC power supply socket, USB socket or the nurse call socket, ensure that the socket is dry before using it.
- If the clamp or clutch needs to be removed for cleaning contact your local distributor.
- Do not use organic solvent such as alcohol or thinner.
- If disinfection is necessary, using the common disinfectors such as Chlorhexidine gluconate and Benzalkonium chloride. After using the agent with a soft cloth, wipe off it with a soft cloth dampened with water or warm water. When using the disinfecting agent, follow the caution of each agent.
- The syringe pump must not be autoclaved.
- Never use a dryer or similar device to dry the syringe pump.
- If liquid spills onto the pump, check whether the pump still functions normally. Test the insulation and leakage current when necessary.
- Do not soak the syringe pump into water.



WARNING:

- Do not clean or disinfect the pump when it is running.

9.2 Periodic Maintenance

Perform a periodic maintenance inspection to ensure safe operation and the longest possible life of the syringe pump, and check the syringe pump once every 2 years. Contact manufacturer or manufacturer's representatives for any doubt.

9.2.1 Checking the Appearance

- Appearance checking: There are no cracks or damages.
- Key operations: If you can press the keys smoothly, they are available.

9.2.2 Checking the Power Cable

- Check the appearance of the power cable. If the appearance is damaged and the plug and the socket are in poor contact, contact manufacturer or manufacturer's representatives for replacement in time.
- If you connect the syringe pump to the AC/DC power and there is no indication of powering on, contact manufacturer or manufacturer's representatives for maintenance in time.

Maintenance

9.2.3 Checking the Infusion Rate

- Check the infusion flow once every 2 years by using the graduate and stopwatch.

Checking condition:

Syringe	Infusion rate	Infusion time	Volume in graduate
MC /B.Braun50/60ml	60mL/h	10min	9.8-10.2mL

9.2.4 Alarms

- Syringe unknown

Pull of the syringe clamp during infusion, alarm information will be visible on the display and audible.

- Syringe Install Error

Press the clutch during infusion, alarm information will be visible on the display and audible.

- Occlusion

Checking condition:

Syringe	Infusion rate	Occlusion level	Alarm time
MC /B.Braun50/60ml	25mL/h	The default occlusion level	Within 2 min

9.2.5 Electric and Mechanical Safety

To ensure safety, test the insulation voltage, leakage current and earthen resistance according to the IEC 60601-1.

9.2.6 Checking the Internal Battery

- Perform the following inspections on the battery every 2 years:
- Connect to the AC power supply to recharge the battery for over 10 hours.
- Turn on the power and attach the syringe (50/60 mL syringe).
- Set the infusion rate to 5 mL/h and start the infusion. Record the start time.
- Operate the system until it stops infusing due to low battery alarm. Record the finish time.
 - If the continuous operation duration reaches 80% or more of the asserted battery operation time, the battery is in good condition.
 - If the continuous operation duration is shorter than half of the asserted battery operation time, the battery reaches the end of its service life. In this case, you must replace the battery. You are suggested to contact the local distributor for battery replacement.
 - Record how long the operation can last after a Low Battery alarm and Battery Empty alarm are triggered respectively. If the operation lasts for less than 30min after a Low Battery alarm is triggered or lasts for less than 3min after a Battery Empty alarm is triggered, contact the local distributor for battery replacement.
- After the battery level check is completed, recharge the battery for next use.

Maintenance

9.2.7 Replacing the Battery

- **Remove an internal battery.**
 - Turn the power off and disconnect the power cord.
 - Use a screwdriver to loosen the battery cover fixing screws at the bottom of the pump.
 - Remove the battery cover.
 - Disconnect the battery cable connector.
 - Remove the battery.
- **Install the internal battery.**
 - Insert the connector of the battery cable into the battery.
 - Insert the new battery into the battery compartment.
 - Attach the battery cover.
 - Use a screwdriver to tighten the screws securing the battery cover.



CAUTION:

- Remove the battery if the syringe pump is not likely to be used for some time.



WARNING:

- The battery's replacement must be done by specialist who has been trained to finish such operation. Otherwise there will be a risk of danger.
- Please strictly follow the instruction to replace the battery, and the battery should be provided by the manufacturer. Otherwise there will be a risk of danger.
- Do not disassemble or short circuit the battery, do not through the battery into the fire. Otherwise there will be a risk of danger caused by the battery linkage or explosion.
- Please follow the local laws to dispose the old battery.

9.3 Maintenance

- If any trouble, explain the situation to the manufacturer or manufacturer's representative and request for repair.
- Never disassemble or try to repair the syringe pump. Doing so may cause a serious failure. The manufacturer and the distributor shall not be responsible for any syringe pump that has been disassembled, modified or used for any purpose other than that for which it is intended.
- If the syringe pump is dropped or subjected to impact, remove it from service even if it doesn't appear damaged externally. Request the manufacturer or manufacturer's representative to inspect it for a possible internal problem.



CAUTION:

- Maintenance staff can request for the related service manual from the manufacturer if

Maintenance

needed.



WARNING:

- The accessories' replacement must be done by specialist who has been trained to finish such operation. Otherwise there will be a risk of danger.
- Parts of the Pump are not serviced or maintained while in use with the patient

9.4 Storage

- Avoid water spills.
- Never store in a hot and humid place.
- Store the pump out of excessive vibration, dust, and corrosive gas.
- Store the pump out of direct sunlight and ultraviolet ray as discoloration may result.

9.5 Transportation

The syringe pump can be transported by common vehicles and shall be protected from being clashed, shook, or wetted by rain and snow. Transportation method shall be in accordance with the order contract specifications.

9.6 Environmental Protection and Recycling

At the end of the product's service life, please contact the manufacturer or manufacturer's representative for disposal advice. Alternatively, dispose of the product and its battery according to the local laws and regulations.

10 Infusion Accuracy Characteristics

The following test is performed in accordance with the IEC60601-2-24:2012 standard. It is used to observe the infusion accuracy and the occlusion response. (For detailed test conditions, see the IEC60601-2-24:2012 standard.)



- The infusion accuracy and the occlusion response may be affected by the use conditions including the pressure, temperature, humidity, syringe, and infusion tube.
- The infusion accuracy does not reflect the clinical standards, for example, patients' age and weight and medicine taken.
- The experiment data only represents the measurement data in the lab.
- Under single fault conditions, the maximum volume of liquid that might be infused by the syringe pump is 5ml and the maximum infusion volume inaccuracy is up to $\pm 40\%$.

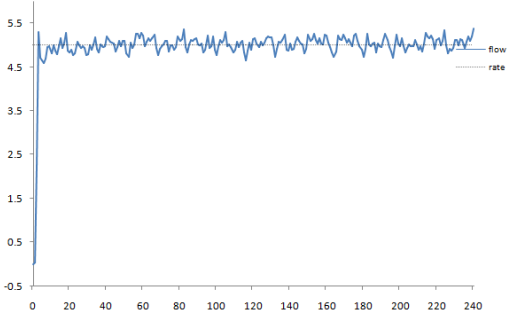
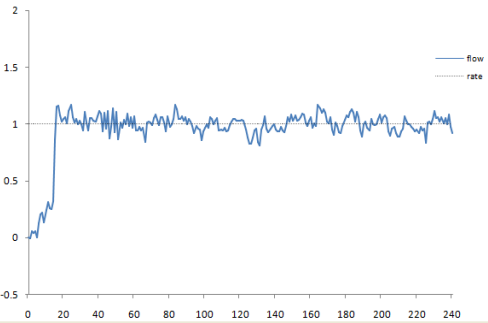
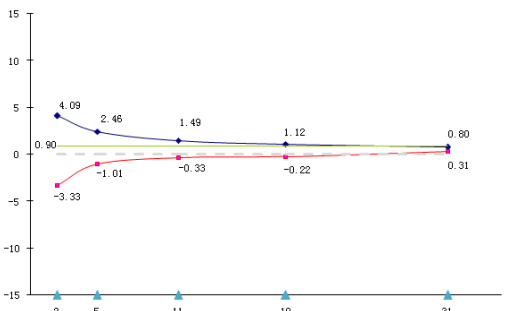
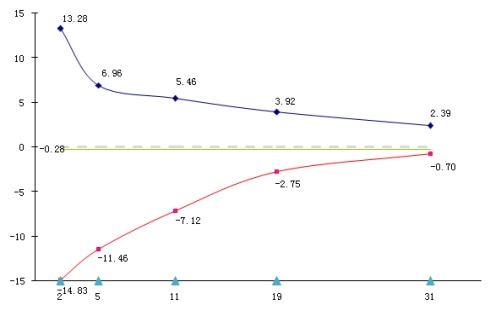
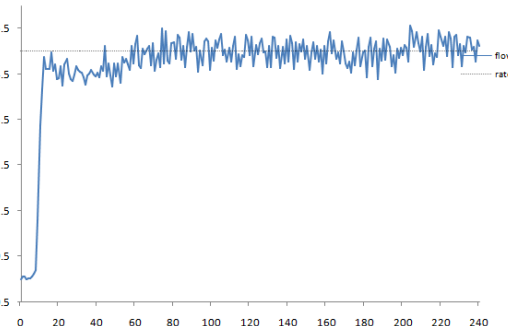
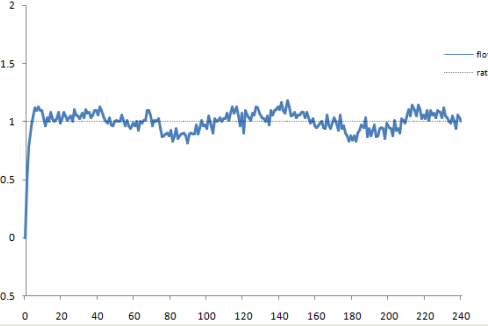
10.1 Flow Rate Characteristics

Start-up and Trumpet curves show the characteristics of the syringe pump after the injection begins and the injection changing status after the syringe pump reaches a normal flow rate.

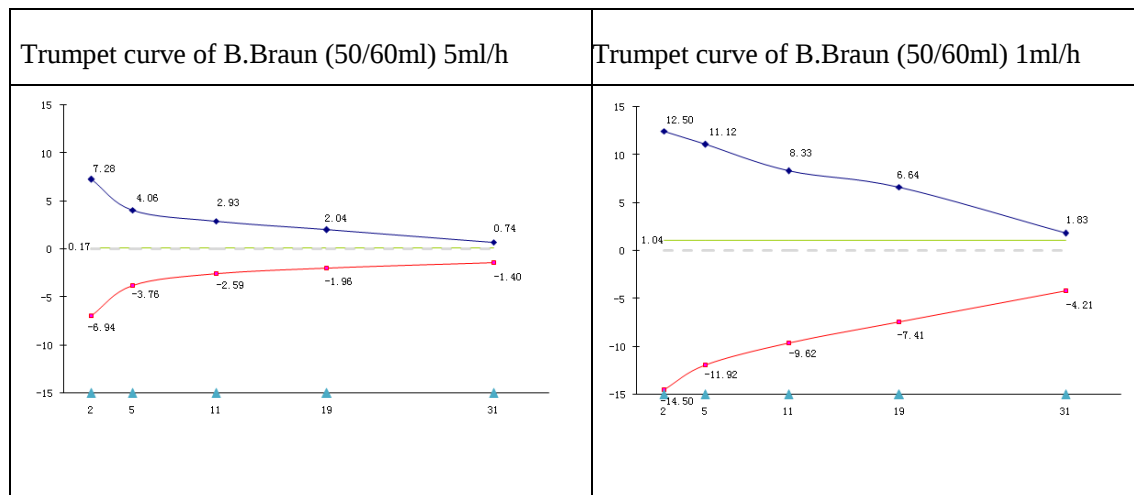
The following test method is performed in accordance with the method mentioned in chapter 201.12.1.102 of the standard IEC 60601-2-24:2012 (Please check above chapter for further details.).

- Accuracy test conditions:
 - Temperature: 21 °C;
 - Relative humidity: 60%;
 - Syringe type: MC (5ml, 10ml, 20ml, 30ml, and 50/60ml), B.Braun (20ml and 50/60ml); 4 sets each.
 - Syringe pump: 1 set
 - Sampling interval: 0.5min
 - Test Period: 120min
 - Test Liquid: ISO 3696:1987 Class III water

Infusion Accuracy Characteristics

Start-up curves of MC (50/60ml) 5ml/h	Start-up curves of MC (50/60ml) 1ml/h
	
Trumpet curve of MC (50/60ml) 5ml/h	Trumpet curve of MC (50/60ml) 1ml/h
	
Start-up curves of B.Braun (50/60ml) 5ml/h	Start-up curves of B.Braun (50/60ml) 1ml/h
	

Infusion Accuracy Characteristics



10.2 Occlusion Characteristics

The occlusion characteristics are reflected by the longest delay time to start an alarm.

The following test method is accordance with the method mentioned in chapter 201.12.4.4.104 of the standard IEC 60601-2-24:2012 (Please check above chapter for further details.).

Occlusion test conditions:

Temperature: 21 ℃;

Relative humidity: 65%;

Syringe type: MC (50/60ml); 3 set

Syringe Pump: 1 set

Length of the extension tube: 1m

Table10-1 Occlusion level, alarm delay time and pill amount under the rate of 5mL/h (MP-30A)

Infusion rate	Occlusion pressure level	Occlusion pressure (mmHg)	Occlusion alarm time (hh:mm:ss)	Bolus (ml)
5ml/h	P1	225 ±150	00:07:46	0.18
	P11	975 ±150	00:25:50	0.68

Infusion Accuracy Characteristics

Table 10-2 Occlusion level and alarm delay time under the rate of 1mL/h (MP-30A)

Infusion rate	Occlusion pressure level	Occlusion pressure (mmHg)	Occlusion alarm time (hh:mm:ss)
1ml/h	P1	225 ±150	00:32:00
	P11	975 ±150	01:45:31

Table10-3 Occlusion level and alarm delay time under the rate of 0.1mL/h (MP-30A)

Infusion rate	Occlusion pressure level	Occlusion pressure (mmHg)	Occlusion alarm time(hh:mm:ss)
0.1ml/h	P1	225 ±150	09:01:59
	P11	975 ±150	21:16:35

Table 10-4 Occlusion level, alarm delay time and pill amount under the rate of 5ml/h (MP-30/MP-30T)

Infusion rate	Occlusion pressure level	Occlusion pressure (mmHg)	Occlusion alarm time (hh:mm:ss)	Bolus (ml)
5ml/h	P1	300 ±100	00:12:24	0.18
	P3	900 ±200	00:22:39	0.55

Table 10-5 Occlusion level and alarm delay time under the rate of 1ml/h (MP-30/MP-30T)

Infusion rate	Occlusion pressure level	Occlusion pressure (mmHg)	Occlusion alarm time (hh:mm:ss)
1ml/h	P1	300 ±100	00:35:00
	P3	900 ±200	1:19:34

Infusion Accuracy Characteristics

Table 10-6 Occlusion level and alarm delay time under the rate of 0.1ml/h (MP-30/MP-30T)

Infusion rate	Occlusion pressure level	Occlusion pressure (mmHg)	Occlusion alarm time (hh:mm:ss)
0.1ml/h	P1	300±100	04:13:16
	P11	900±200	24:37:05



CAUTION:

Unit conversion list

Description	Unit	Unit conversion
Pressure	kPa	1kPa=7.5mmHg
	psi	1psi=51.715mmHg
	bar	1bar=750mmHg

Appendix A

Appendix A Electromagnetic Compatibility (EMC)

The MP-30 Series syringe pump conforms to EMC standard IEC 60601-1-2.

Guidance and manufacturer's declaration – electromagnetic emissions		
The MP-30 Series syringe pump should be used under the regulation electromagnet environment. The user should operate the MP-30 Series syringe pump under following electromagnet environment.		
Emission test	Emission test	Electromagnetic environment-instructions
RF emission CISPR 11	Group 1	MP-30 Series syringe pump only use radio-frequency while operating its internal functions, therefore, the radio-frequency is much low and has little interference to the electronic devices nearby.
Radio-frequency emission CISPR 11	Class A	The MP-30 Series syringe pump can be used in any building including civil residence.
Harmonic emission IEC61000-3-2	Class A	
Voltage fluctuation and flashing IEC 61000-3-3	Complies	

Guidance and manufacturer's declaration – electromagnetic immunity			
The [MP-30 Series] is intended for use in the electromagnetic environment specified below. The customer or the user of the [MP-30 Series] should assure that it is used in such an environment.			
IMMUNITY test	IEC60601test level	Compliance level	Electromagnetic environment –guidance
Electrostatic discharge (ESD) IEC 61000-4-2	±8 kV contact discharge ±15 kV air discharge	±8 kV contact discharge ±15 kV air discharge	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.
Electrical fast	±2 kV power cable	±2 kV power cable	Mains power quality should

Appendix A

transient (EFT) IEC61000-4-4	±1 kV I/O cable		be that of a typical commercial or hospital environment.
Surge IEC 61000-4-5	±1 kV difference mode ±2 kV common mode	±1 kV difference mode ±2 kV common mode	
The voltage dropping, short interruption and voltage change IEC 61000-4-11	<5% U_T (dropping>95% U_T)0.5 period 40% U_T (dropping 60% U_T)5 period 70% U_T (dropping 30% U_T)25 period <5% U_T (dropping>95% U_T)5seconds	<5% U_T (dropping>95% U_T)0.5 period 40% U_T (dropping 60% U_T)5 period 70% U_T (dropping 30% U_T)25 period <5% U_T (dropping>95% U_T)5seconds	Mains power quality should be that of a typical commercial or hospital environment. If the user of the [MP-30 Series] requires continued operation during power mains interruptions, it is recommended that the [MP-30 Series] be powered from an uninterruptible power supply or a battery.
Power frequency magnetic fields (50/60Hz) IEC 61000-4-8	3 A/m	3A/m	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment
NOTE U_T is the AC mains voltage prior to application of the test level.			

Appendix A

Guidance and manufacturer's declaration – electromagnetic immunity			
The [MP-30 Series] is intended for use in the electromagnetic environment specified below. The customer or the user of the [MP-30 Series] should assure that it is used in such an environment.			
Immunity Test	IEC 60601 test level	Compliance level	Electromagnetic environment –guidance
Conducted RF IEC61000-4-6	3Vrms 150k~80MHz	3Vrms	Portable and mobile RF communications equipment should be used no closer to any part of the [MP-30 Series], including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.
Radiated RF IEC61000-4-3	3V/m 80M~2.5GHz	3V/m	Recommended separation distance: $d = 1.2\sqrt{P}$ $d = 1.2\sqrt{P} \quad 80M\sim 800MHz$ $d = 2.3\sqrt{P} \quad 800M\sim 2.5GHz$ where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance d in meters (m). Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey^a, should be less than the compliance level in each frequency range^b. Interference may occur in the vicinity of equipment marked with the following symbol: 
NOTE 1 At 80 MHz and 800 MHz, the higher frequency range applies. NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.			
^aField strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the [MP-30 Series] is used exceeds the applicable RF compliance level above, the [MP-30 Series] should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the [MP-30 Series]. ^b Over the frequency range 150 kHz to 80 MHz, field strengths should be less than [3] V/m.			

Appendix A

Recommended separation distances between portable and mobile RF communications equipment and the [MP-30 Series]

The [MP-30 Series] is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the [MP-30 Series] can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the [MP-30 Series] as recommended below, according to the maximum output power of the communications equipment.

Rated maximum output power of transmitter (W)	Separation distance according to frequency of transmitter (m).		
	150k~80MHz $d = 1.2\sqrt{P}$	80M~800MHz $d = 1.2\sqrt{P}$	800M~2.5GHz $d = 2.3\sqrt{P}$
0.01	0.12	0.12	0.23
0.1	0.38	0.38	0.73
1	1.2	1.2	2.3
10	3.8	3.8	7.3
100	12	12	23

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in meters (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

NOTE 1 At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

Appendix B

Appendix B Default Factory Settings

This chapter lists some default factory settings of syringe pump. Users can not modify the default factory settings, but if necessary, they can recover the syringe pump to the default factory settings state.

Parameters

Parameters setting	Default factory setting
KVO flow rate	1ml/h
Pressure unit	mmHg
Occlusion pressure	MP-30A: 600mmHg MP-30 and MP-30T: 550mmHg
Near end	3min
Built-in consumable brand	MC (5, 10, 20, 30, 50/60ml), B.Braun (20, 50/60ml)

System time

System time and date	The default factory setting
Time	00:00
Date	2014-1-1
Time form	24 hours
Date form	Year-month-day

Parameter Units

Parameter Name	Unit
Acti Agent	ng, ug, mg, g, U, kU, IU, EU, mmol, mol, kcal, mIU, kIU, mEq
Conc.	ng/ml, ug/ml, mg/ml, g/ml, U/ml, kU/ml, IU/ml, EU/ml, mmol/ml, mol/ml, kcal/ml, mIU/ml, kIU/ml, mEq/ml
DoseRate	x/min, x/kg/min, x/h, x/kg/h. (x is n ng, ug, mg, g, U, kU, IU, EU, mmol, mol, kcal, mIU, kIU, mEq)

MEDCAPTAIN MEDICAL TECHNOLOGY CO., LTD.

Manufacturer: MEDCAPTAIN MEDICAL TECHNOLOGY CO., LTD.
Manufacturer Address: 12th Floor, Baiwang Research Building,
No.5158 Shahe West Road, Xili, Nanshan, 518055
Shenzhen, Guangdong, PEOPLE'S REPUBLIC OF CHINA
Production Address: Building C, Jiale Science and Technology
Industrial Park, Matian Street, Guangming, 518106
Shenzhen, Guangdong, PEOPLE'S REPUBLIC OF CHINA
Tel: +86-755-26953369
Website: <http://www.medcaptain.com>
E-mail: mc.service@medcaptain.com

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