Medical Suction Unit

SAM 35, SAM 36

Operating and Maintenance Manual
Dear Customer,

We take this opportunity to thank you for purchasing a SA Medical Suction Unit. Please read the operating instructions and listed precautions thoroughly before attempting to operate the unit. MGE manufactures its range in accordance with the requirements of BS EN ISO 9001 and BS EN ISO 13485

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Return of Medical Equipment
Should you wish to return any equipment to MGE (Colchester) Limited (MGE), or one of our designated distributors, Health Service Guideline HSG (93) 26 Decontamination of equipment prior to inspection, service or repair must be adhered to. Failure to follow this guideline will invalidate any warranty claims and result in the equipment being destroyed.

Definition of symbols used in these instructions:

- The Instruction for use must be referred to!
- Manufacturers’ details and date of manufacture
- Electricity Warning
- Safety Warning
- Disposal in accordance with directive 2012/19/EU
- Temperature Limits
- Humidity Limits
- Type B applied part
- Ingress Protection
- Equipotential earth point
1. SAFETY INSTRUCTIONS

The safety of the patient and suction unit operator are the first priorities. It is therefore vital that the following precautions are strictly observed:

**WARNING!!!**

Never operate a SAM suction unit without an MGE filter capsule.

The filter capsules are single patient use and cannot be reused. Reuse of such devices can be dangerous for both patient and operator.

A hydrophobic filter must be fitted internally to prevent liquid passing through to the pump. Hydrophobic filters only work once and should be replaced immediately it becomes wetted.

No modification of this equipment is allowed.

Only original and approved spare parts and collection systems must be used with all SAM suction units – failure to use original or approved spares will invalidate the warranty and may cause injury or damage the unit.

No liability can be accepted by MGE for units affected by the occurrence of overflow when Disposable Liner Systems are being used.

Other than for routine daily procedures, any maintenance or repairs to MGE products must be carried out by fully trained and qualified Electro-Biomedical engineer/technician (EBME) or an authorized MGE dealer. Such persons must be familiar with the relevant standards, rules, accident prevention regulations, and operating conditions as a result of their training, experience, and instruction. They are qualified to carry out the required activities and in doing so recognize and avoid potential hazards.

All testing on SAM suction units should be in accordance with ISO 10079-1

Contamination may be present on any components. When cleaning or replacing any part of the SAM unit appropriate protective clothing and gloves MUST be worn to avoid contamination. Disposal of contaminated parts must be according to local protocols.

Store the manual in a safe place, so that it is available to the trained personnel at all times.

All collection containers must be securely mounted when in use.

The overflow valve may not operate fully against frothing. To prevent frothing anti-foam agent maybe used.

When replacing a full collection container, be aware of its weight and ensure handling the container is comfortable to avoid the possibility of spillage.

Transport of the SAM suction unit with a full jar attached is not advised.

Solvent-based cleaning agents or abrasive cleaners must not be used on any SAM suction units. Do not wash any SAM suction units under running water or submerge in water.

Any dismantling and re-assembly of this equipment - for whatever purpose - must be followed by testing in accordance with the manufacturers' recommendations as specified for monthly maintenance.
ELECTRICAL WARNING

Always electrically isolate the SAM suction unit from the mains power supply before carrying out any cleaning, maintenance or repairs. Electrical isolation of the SAM suction unit is by removal of the mains power cable from the mains power supply.

Never operate a SAM suction unit in the presence of flammable gas such as anaesthetic agents. This is an Explosion hazard!

Ensure the power cable does not create a trip hazard or is subject to damage.

The suction pumps may only be opened by qualified technical personnel. Electric shock hazard!

Equipotential bonding may be required in some critical treatment areas. This is intended to minimize any voltage differences between earthed parts of equipment and any other exposed metal in the room. All conductive metal in an equipotential area is connected to a common equipotential earth (reference) point.

The SAM suction unit has on its back panel (adjacent to the mains power cable) an equipotential earth point to allow connection to an earth reference point when required. Installation of this connection must be carried out by a fully qualified engineer. Ref: BS EN 60601-1
2. GENERAL DESCRIPTION

**SAM 35** - General-purpose high vacuum medical suction unit, normally incorporating two collection containers of 2L capacity. Other types of collection container are allowed. Designed specifically for use in medical and healthcare facilities, primarily in operating theatres where high vacuum and high flow are required. These units are not intended for field and transport use.

**SAM 36** - General-purpose high vacuum medical suction unit, incorporating two collection containers of 2L capacity. Other types of collection container are allowed. Designed specifically for use in intra-uterine procedures where high vacuum and high flow are required and is fitted with a wide bore collection container top complete with specimen filter. These units are not intended for field and transport use.

There are no contra-indications.
3. INSTRUCTIONS FOR USE

3.1 Before Operating Unit

BEFORE operating your new SAM suction unit please read the following instructions carefully.

Become thoroughly familiar with the operation and maintenance of the unit before using and note the information on the control panel and rear of the unit. Only persons trained in its use should operate the unit.

Check that the voltage supply is as printed on the rating plate fitted to the back of the SAM suction unit.

Unscrew and remove the transport screws located on the underside of the unit.

**Warning!** Always electrically isolate the SAM suction unit from the mains power supply before carrying out any cleaning, maintenance or repairs. Electrical isolation of the SAM suction unit is by removal of the mains power plug from the mains power supply.

3.2 Unit Set Up

- Connect the desired collection system securely in place on the bracket fitted to the top of the unit. More detail can be found later in this section.
- Connect the desired equipment to the patient inlet tube on the collection container top. Multi-fit bubble tubing (if supplied) has an I/D ∅7mm. Recommended length of tubing used is 2 meters.
- Connect the unit to the power supply, check label on rear of unit for correct ratings. Do not obstruct access to the mains power supply, and ensure the power cable does not create a trip hazard or is subject to damage. Ensure the exhaust outlet on the side of the unit is kept clear of any restriction.
- Switch the SAM suction unit on by pressing the green rocker switch so that it illuminates. The lamp indicates the SAM suction unit is operating and ready for use.

3.3 Handling

The larger SAM suction unit are not normally intended for lifting or carrying. They will either be designed to installed on a trolley or fitted with lockable wheels. This mobility allows the larger SAM suction unit to be moved during use and easily transported between locations.

3.4 Operating Environment

Operation of this equipment must be within the following ambient condition:

- Temperature: -5°C to +40°C
- Humidity: 30% to 80%

**Warning!** Never operate a SAM suction unit in the presence of flammable gas such as anaesthetic agents. This is an Explosion hazard!

3.5 Gauge (Manometer) - Indicated Vacuum

The unit has a gauge fitted to allow the vacuum to be set at a pre-determined level and monitored during use. To achieve the desired degree of vacuum, occlude the tubing fitted to the patient inlet on the collection container top, and read the indicated vacuum on the gauge. Adjust the vacuum control until the required degree of vacuum is achieved. (Turning the knob clockwise increases the vacuum). The SAM suction unit is now set to operate at the selected vacuum.
3.6 External Filter

**Warning!** Never operate a SAM suction unit without a MGE filter capsule.

The filter capsules are a sealed disposable bacterial or hydrophobic filter capsule. Both the bacterial and hydrophobic filters have bacterial retention of 99.9999% ensuring safety and hygiene for patient and operator. The 0.3 micron particle retention of the filter medium also gives not less than 99.985% Dispersed Oil Particulate (D.O.P.) and so provides an effective barrier against possible aerosol contamination. The hydrophobic filter also has an integrated micro porous membrane on the rear of the filter medium which allows a clear flow of air through one way and an excellent block against any accidental back flow the other. The external filters are fitted between the vacuum connector and the filter capsule mount, and can be either a bacterial or hydrophobic filter. The connector, mount, and ‘O’ rings are autoclavable. However, the filter capsules are ‘Single Patient Use’ and must not be autoclaved. The external filter must be changed with every patient and renewed in accordance with the recommendations detailed in the ‘Daily Procedures’.

**Warning!** The filter capsules are single patient use and cannot be reused. Reuse of such devices can be dangerous for both patient and operator.

3.7 Internal Filter

The SAM suction unit must have fitted internally a hydrophobic filter capsule to prevent ingress of foreign material into the pump. The hydrophobic filter prevents costly pump damage as well as additional protection against contamination from liquids being sucked through due to jar overfill. Liquid sucked through to the pump motor will cause damage and invalidate the warranty. The filter is disposable and must not be autoclaved, but renewed in accordance with the recommended maintenance program.

**Warning!** A hydrophobic filter must be fitted internally to prevent liquid passing through to the pump. Hydrophobic filters only work once and should be replaced immediately it becomes wetted.

3.8 Fluid Collection Containers

The silicon service tube (Ø6mm I/D with 3mm minimum wall thickness) must be connected between the filter and the changeover valve. The changeover valve must then be connected to the vacuum port on the collection container. These tubes have an elbow connector on one end for fitting to the filter and for fitting to the ‘VACUUM’ port of the SAM 2 collection container. The silicon tubing used in conjunction with this equipment is a replaceable item. It should be changed regularly according to the level of usage and where it has become in any way contaminated or damaged.

**Warning!** All collection containers must be securely mounted when in use.

An anti-foam agent may be put into the fluid collection container, without disinfectant solution, before use to reduce the possibility of frothing. It should not however, be placed into the container for extended storage periods.
3.8.1 SAM 2 Collection System

The SAM 2 collection container is fitted as standard to all SAM suction units. They are fully autoclavable Polyester Carbonate (P.E.C.) collection containers, with integral handle and for IU applications, with a wide bore. The collection containers are connected to the SAM suction unit via a changeover valve.

- Place the collection containers in the brackets. Connect the tubing from the changeover valve to the angle connector in the centre of the lid.
- Connect the patient tube to the patient connection.
- Turn on the SAM suction unit.
- Check the desired vacuum is established.
- After the suction procedure – disconnect the patient tube.

The SAM 2 is fitted with an overflow valve designed to shut off the vacuum to the SAM 2 collection container when the fluid level reaches 1750ml. When the valve operates, the SAM suction unit must be switched off and the full container replaced by an empty one.

It should be noted that even after the valve has shut off, fluid might continue to be drawn into the container to an extent dependent upon the level of vacuum in the container at the time when the valve closes.

⚠️ Warning! The overflow valve may not operate fully against frothing. To reduce frothing an anti-foam agent may be used.

⚠️ Warning! When replacing a full collection container, be aware of its weight and ensure handling the collection container is comfortable to avoid the possibility of spillage. Transport of the SAM suction unit with a full jar attached is not advised.

3.8.2 SAM 4 Collection System

The SAM 4 collection container is fitted as standard to all SAM suction units. They are fully autoclavable Polyester Carbonate (P.E.C.) collection containers. The collection containers are connected to the SAM suction unit via a changeover valve.

- Place the collection containers in the cradle. Connect the tubing from the changeover valve to the connector on top of the lid (marked ‘VACUUM’).
- Connect the patient tube to the ‘PATIENT’ connection.
- Turn on the SAM suction unit.
- Check the desired vacuum is established.
- After the suction procedure – disconnect the patient tube.

The SAM 4 is fitted with an integral shut off for overflow protection.

⚠️ Warning! The overflow valve may not operate fully against frothing. To reduce frothing an anti-foam agent may be used.
Warning! When replacing a full collection container, be aware of its weight and ensure handling the collection container is comfortable to avoid the possibility of spillage. Transport of the SAM suction unit with a full jar attached is not advised.

3.9 Disposable Liner Systems

The SAM suction units have been designed to accept various disposable liner systems. When fitted, protection of the unit against overflow is entirely dependent upon the correct use of the appropriate liner in accordance with the manufacturers’ instructions.

Warning! No liability can be accepted by MGE for units affected by the occurrence of overflow when Disposable Liner Systems are being used.

3.9.1 VacSax suction system

- Place liner into collection container and push firmly.
- Push the taper connector on the vacuum tubing from the suction controller into the vacuum port with a twisting motion.
- Connect the patient tubing firmly to the patient port to ensure a good fit.
- Turn on the SAM suction unit.
- Confirm suction is present at the patient tube by occluding the patient tube end.
- After the suction procedure – disconnect the patient tube and fit stopper located on rim into the port.
- Turn OFF the SAM suction unit and remove the liner for disposal.

The collection container may be sterilised by autoclaving at 121°C or by washing with water based disinfectants. Before autoclaving, rinse the collection container well to remove any detergent.

3.9.2 Abbott Receptal Disposable Suction System

- Place the collection container in the bracket making sure that the tee connection on the collection container is tight. Fully extend the liner, and place it into the collection container. Make sure that the liner is tightly secured in the collection container.
- The liner incorporates an internal shut off valve, which protects the vacuum source from contamination. Connect the vacuum source tubing (from the filter) to one side of the tee connection and the lid to the other side using the ‘yellow to yellow’ coding.
- Connect the patient tube to the patient connection, either directly or by using the optional elbow piece, which prevents kinking of the patient tubing.
- Turn on the SAM suction unit to inflate the liner and the collection container is now ready for use.
- For maximum safety and to minimise the risk of contamination when dismantling, the fluid level must not rise above the ‘DO NOT FILL ABOVE THIS LINE’ mark on the collection container.
- At the end of the procedure, the SAM suction unit should remain switched on while the patient suction tubing is removed from the patient port and discarded.
- Disconnect the liner lid tubing from the tee connection and immediately reconnect the yellow connector to the patient port with a push and twist motion.
• Turn the vacuum off and use the thumb tab to remove the liner for disposal.

**Warning!** The liner lid tubing must not be used as a carry handle.

### 3.9.3 Serres Suction Bag System

- The system comprises the suction bag (A), the collection container (B), and the angle connector (C).
- Place the collection container (B) in the bracket. Connect the tubing from the **SAM** filter to the angle connector (C).
- Place the suction bag into the collection container ensuring bag tail is not trapped between the lid and collection container.
- Turn on the **SAM** suction unit.
- Install the suction bag by using the vacuum – Occlude the patient connection and simultaneously press lightly from the middle of the lid.
- Establish the desired vacuum and ensure the bag is fully inflated.
- Connect the patient tube to the patient connection.
- After the suction procedure – disconnect the patient tube and close the connection with the plug provided on the lid.
- Turn OFF the **SAM** suction unit and remove the liner for disposal.

If necessary, the collection container only, may be washed (85°C) and autoclaved (121°C). Before washing, disconnect the angle connector. If autoclaving, rinse the collection container well to remove any detergent.

### 3.10 Effect of altitude

Altitude effects all vacuum pumps and it should be noted that there will be a reduction in the maximum achievable vacuum / negative pressure level equivalent to approximately 3.5% per 300m (approximately 1000ft) rise in altitude.

### 3.11 Cleaning Procedure

**Warning!** Contamination may be present on any components. When cleaning or replacing any part of the **SAM** suction unit appropriate protective clothing and gloves MUST be worn to avoid contamination. Disposal of contaminated parts must be according to local protocols.

**Warning!** Always electrically isolate the **SAM** suction unit from the mains power supply before carrying out any cleaning, maintenance or repairs. Electrical isolation of the **SAM** suction unit is by removal of the mains power cable from the mains power supply.

**Warning!** Solvent-based cleaning agents or abrasive cleaners must not be used on any **SAM** suction units. Do not wash any **SAM** suction units under running water or submerge in water.

To clean the outside case of the **SAM** suction unit, disconnect the unit from the power supply, wipe over with a clean damp cloth or use an appropriate mild disinfectant solution. Following the manufacturers’ instructions on cleaning product, and avoiding excessive moisture.

### 3.12 Transport

The **SAM** suction unit will be adequately boxed and protected to ensure no damage occurs during normal transportation of goods, providing the ambient conditions are within the following parameters:

Temperature max +60°C min -20°C  
Humidity max +80% min +30%
There are no restrictions for land, air, or sea transport.

3.13 Storage

SAM suction units must be stored in a dry, dust-free, well ventilated environment. The storage environment should not exceed the temperature and humidity conditions stated below. Avoid direct sun or UV exposure and shield nearby sources of heat. The equipment should be stored in its original packaging providing no damage is evident. Protect against ground moisture by storing on a shelf or wooden pallet.

Temperature max +60°C min -20°C
Humidity max +80% min +30%

3.14 Long Term Storage

When the units are held in storage, or used very infrequently, the three-monthly maintenance period may be extended to twelve months. Particular care must be taken when inspecting flexible components, such as valves, diaphragms, to ensure embrittlement has not occurred. SAM suction units should be stored in a cool, dry environment, as stated above.

3.15 Instructions by Medical Staff to Patients

When the equipment is required by a patient for home use, Medical Staff must fully instruct the patient on the safe operation of the equipment. In the event of equipment contamination or failure, the patient must be advised to switch off the unit and contact the authority from which the unit is loaned.

3.16 Troubleshooting

<table>
<thead>
<tr>
<th>Problem:</th>
<th>Cause:</th>
<th>Solution:</th>
</tr>
</thead>
<tbody>
<tr>
<td>No power to unit</td>
<td>Unit not turned on</td>
<td>Turn Main Switch „ON/OFF“ ON</td>
</tr>
<tr>
<td></td>
<td>Wrong operating voltage</td>
<td>Check mains voltage output</td>
</tr>
<tr>
<td></td>
<td>Mains not connected</td>
<td>Connect mains cable</td>
</tr>
<tr>
<td></td>
<td>Defective Fuse</td>
<td>Check fuse and replace if indicated</td>
</tr>
<tr>
<td>Pumps fail to run</td>
<td>Leak in vacuum system</td>
<td>Check all connection and tubing.</td>
</tr>
<tr>
<td></td>
<td>Collection Jar is full</td>
<td>Replace Jar</td>
</tr>
<tr>
<td>Pump suction power too weak</td>
<td>Vacuum leaks</td>
<td>Check all seals and hoses. Make sure Lid is securely on Jar.</td>
</tr>
<tr>
<td></td>
<td>Vacuum not to required setting</td>
<td>Turn vacuum control clockwise until the desired suction power is reached</td>
</tr>
<tr>
<td></td>
<td>Tubing is plugged, bent and/or disconnected</td>
<td>Replace tubing if plugged and eliminate any bends</td>
</tr>
<tr>
<td></td>
<td>Internal hydrophobic filter is blocked or wet</td>
<td>Replace internal hydrophobic filter. – Must be done by EBME</td>
</tr>
<tr>
<td></td>
<td>External filter is blocked</td>
<td>Replace External filter.</td>
</tr>
</tbody>
</table>

If a problem cannot be solved, contact a fully trained and qualified engineer (EBME), authorized MGE dealer or MGE (sales@mgelectric.co.uk).
4. MAINTENANCE

Warning! Other than for routine daily procedures, any maintenance or repairs to MGE products must be carried out by fully trained and qualified engineers (EBME) or an authorized MGE dealer. Such persons must be familiar with the relevant standards, rules, accident prevention regulations, and operating conditions as a result of their training, experience, and instruction. They are qualified to carry out the required activities and in so doing recognize and avoid potential hazards. All testing on SAM suction units should be in accordance with ISO 10079-1

Warning! No modification of this equipment is allowed.

Warning! Contamination may be present on any components. When cleaning or replacing any part of the SAM suction unit appropriate protective clothing and gloves MUST be worn to avoid contamination. Disposal of contaminated parts must be according to local protocols.

Warning! Any dismantling and re-assembly of this equipment - for whatever purpose - must be followed by testing in accordance with the manufacturers’ recommendations as specified for Monthly maintenance.

Warning! Isolate the SAM suction unit from the mains power supply before carrying out any maintenance or repairs. Electrical isolation of the SAM suction unit is by removal of the mains power cable from the mains power supply.

4.1 Daily Procedures

- The external filter capsule should be changed after any of the following:
  - Each day’s use.
  - If it becomes wetted by froth.
  - After aspiration of any infective material.
  - Before being used on a new patient.

- Examine the collection container for damage. Replace if necessary.
- Examine the external tubing for ageing, damage, or contamination. Replace if necessary using equivalent tubing (Ø6mm I/D with 3mm minimum wall thickness).
4.2 Monthly Maintenance

- Carry out the daily maintenance procedure.
- Check the vacuum and the flow at the collection container top.
- With the collection container empty and the vacuum control valve set to maximum, switch on the unit and read the indicated vacuum on the gauge. Occlude the suction inlet (Patient connection) on the collection container top and note the time taken for the gauge to indicate an increase to 450mm Hg (60 kPa) vacuum. This time should not exceed 10 seconds.
- The gauge should continue to rise and stabilise to indicate the maximum vacuum available, and this should be not less than 590mm Hg (78.7 kPa).
- Replace the exhaust filters in the exhaust outlet housing located on the side of the casing and marked 'Exhaust Outlet'.

4.3 Three Monthly Maintenance

- Carry out the monthly procedure. Poor performance of the unit would indicate the internal pump requires maintenance

**Warning!** Always electrically isolate the SAM suction unit from the mains power supply before carrying out any cleaning, maintenance or repairs. Electrical isolation of the SAM suction unit is by removal of the mains power cable from the mains power supply.

- Remove the back cover.
- Check the internal and external tubing for ageing or wear. Replace with equivalent tubing where necessary. Tubing to gauge Ø5mm I/D, wall thickness min 2mm, all other tubing on the SAM suction unit Ø6mm I/D, wall thickness min 3mm.
- Replace the internal hydrophobic filter. Details in section 4.6 Replacing the Internal Hydrophobic Filter
- Check the internal wiring for ageing or wear. Replace with equivalent material and terminations where necessary.
- Examine the mains cable for wear or damage and replace if necessary using equivalent cable.
- Replace the back cover. Do not overtighten the rear cover fixings. (Recommended torque 20 cNm)
- Examine the overflow valve float mechanism (SAM 2) to ensure that this has free movement.
- Check the vacuum and flow performance at the collection container as previously described.
4.4 General Layout
4.5 Pump (If Required)

- Remove the pump from the SAM suction unit.
- Remove the pump head from the pump casing and examine the diaphragm for signs of wear.
- Check the balance within the connecting rod for excessive movement. Replace the connecting rod and balance if necessary.
- Remove both pump head nozzles from the pump head.
- Remove the transfer valves.
- Clean the valve seats and check the valves for signs of wear.
- Replace any worn or damaged components as necessary and re-assemble.
- Replace the pump back into the SAM suction unit and re-connect the internal tubing.

**Note:** Replacement kits are available to change an old LQ45 Pump with a new MQ24 Pump. Details found in Options and Recommended Spares section.
4.6 Replacing the Internal Hydrophobic Filter

**Warning!** A hydrophobic filter must be fitted internally to prevent liquid passing through to the pump. Hydrophobic filters only work once and should be replaced immediately it becomes wetted.

When replacing filters, particularly the hydrophobic filter, it is important to ensure the filter is connected correctly to suit the direction of flow. The filters will only operate efficiently in one direction. The flow diagram indicates the correct orientation installed in the pipework.

![Hydrophobic Filter Installation](image)

4.7 Replacing the Main Cable

**Warning!** Always electrically isolate the SAM suction unit from the mains power supply before carrying out any cleaning, maintenance or repairs. Electrical isolation of the SAM suction unit is by removal of the mains power cable from the mains power supply.

- Isolate the SAM suction unit from the mains power supply!
- Detach the back panel to access the internal terminal block.

- Disconnect the brown and blue conductors of the cable from the terminal block, and withdraw the cable.
- Fit the replacement cable through the access hole.
- Connect the conductors to their respective terminals (Blue to Blue and Brown to Brown) in the terminal block. Provide sufficient length of cable between the gland and terminal block.
- Fit the cable into the labyrinth.
- Ensure all conductors are fitted as the original.
- Check continuity of each circuit.
- If reconnection is correct - replace the back panel.
4.8 Flow Diagram

![Flow Diagram](image-url)
## 5. OPTIONS AND RECOMMENDED SPARES

### 5.1 Options
User Instruction Manuals are offered in English. Other languages are available on request.

<table>
<thead>
<tr>
<th>Model</th>
<th>Electrical Specification</th>
<th>Collection Jars</th>
<th>Power Cable</th>
<th>Version</th>
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<tbody>
<tr>
<td>35</td>
<td>01 230Vac 50/60Hz</td>
<td>00 No Jar</td>
<td>01 UK</td>
<td>01 Static</td>
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<td>36</td>
<td>02 110Vac 50/60Hz</td>
<td>01 SAM 2 (2Ltr)</td>
<td>05 European</td>
<td>02 Mobile</td>
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<td>04 Pennine</td>
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<td>05 SAM 4 (1Ltr)</td>
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<td>08 Cardinal Medline</td>
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<td>10 SAM 2 Holder</td>
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<td>11 MTP</td>
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<td>13 Flowmeter Flowvac</td>
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</tbody>
</table>

**Example:** SAM 35 / 01 01 01 02

### 5.2 Recommended spares
Only original and approved spare parts must be used with all SAM suction units – failure to use original spares will invalidate the warranty and may cause injury and/or damage to unit.

All spares can be purchased by the user from MGE directly. Not all available spares are listed below. Please contact the MGE sales team for a full list. (sales@mgelectric.co.uk)

It is recommended that only competent persons should undertake the replacement of spare parts.

#### 5.2.1 Collection Container Components
- Disposable bacterial filters – 24pk ................................................................. MSP1002
- SAM 2 bottle .......................................................... SAM 2
- SAM 2 IU bottle .......................................................... SAM 2 IU
- SAM 4 bottle .......................................................... MSP1290
- SAM 4 lt Jar Conversion ................................................................. MSP1317
- SAM 2 bottle top assembly .......................................................... MSP1047
- SAM 2 IU plastic bottle top assembly .................................................. MSP1071
- SAM 2 overflow valve – 10pk .......................................................... MSP1048
- Silicone tube (O/D ∅12mm) – 25M .................................................. MSP1156
- SAM patient multi-fit bubble tubing (I/D ∅7mm) – 30M .................................. MSP1155
### Unit Components

- Disposable hydrophobic filters – 24pk  
  MSP1003
- Elbow connector and ‘o’ rings – 10pk  
  MSP1004
- Exhaust filter – 10pk  
  MSP1015
- Filter capsule mounts with ‘o’ rings and fixings – 10pk  
  MSP1017
- 2amp 20mm fuses – 10pk  
  MSP1225
- Power cable – 2M  
  MSP1567

### Pump Components

- MQ45 pump (Pump & motor 230V 50/60Hz)  
  MSP1226
- Motor 230V 50/60Hz  
  MSP1227
- MQ45 pump head connector set  
  MSP1230
- MQ45 pump head service kit  
  MSP1229
- MQ45 consumable spares service kit  
  MSP1231
- Con rod/diaphragm assembly  
  MSP1210
- **SAM 35 MQ45 replacement for LQ45**  
  MSP1236
- **SAM 36 MQ45 replacement for LQ45**  
  MSP1240

### Accessories

- Foot-switch conversion kit  
  MSP1151
6. TECHNICAL SPECIFICATION

6.1 General Dimensions

- **Free air-flow (litres/minute):** *50 lt/min** nominal *25 lt/min* nominal
- **Vacuum:** *675mmHg** *730mmHg*
- **Performance classification:** High vacuum High flow (ISO 10079-1)
- **Nominal collection container capacity:** 2 Litres or 4 Litre**
- **Power input (watts):** 100W
- **Standard supply voltages available:** 220V-240V 50/60 Hz, 1Ph 110V-120V 50/60 Hz, 1Ph
- **Capacitor:** 220V-250V 4uF 110V-120V 12uF
- **Fuse type: 5 x 20mm** 220V-250V: F2.0A, 250V 110V-120V F 3.0A, 250V
- **Electrical protection:** Type B, Applied part to EN60601-1
  Class I insulated (Earth Required) to EN60601-1
- **Electric motor classification:** BS2757 Class B (insulated). Auto reset thermal cut out
- **Basic Weight:** 21Kg
- **Maximum allowable weight:** 30kg all units.
- **Maximum Noise level:** 55 dB (a)

*Vacuum measurements are quoted at sea level, and flow rates are taken at the filter point.
** SAM 35 Options
* SAM 36 Options
6.2 End of life

IMPORTANT INFORMATION
Correct disposal of the product in accordance with EC directive 2012/19/EU

At the end of its life, the product must not be disposed of as urban waste.
It must be taken to a special local authority differentiated waste collection centre or to a dealer providing this service.

6.3 Electromagnetic Compatibility (EMC)

Special Instructions / Notes regarding the SAM suction unit and Electromagnetic compatibility (EMC) testing to EN60601-1-2: 2014

The SAM suction unit has been tested regarding its ability to operate in an environment containing other electrical/electronic equipment (including other medical devices).

The purpose of this testing is to ensure the SAM suction unit is not likely to adversely affect the normal operation of other such equipment and that other such equipment is not likely to adversely affect the normal operation of the SAM suction unit.

Despite the testing of the SAM suction unit that has been undertaken, normal operation of the SAM suction unit can be affected by other electrical/electronic equipment and portable and mobile RF communications equipment.

As the SAM suction unit is medical equipment, special precautions are needed regarding EMC (electromagnetic compatibility).

It is important that the SAM suction unit is configured and installed/put into service, in accordance with the instructions/guidance provided herein and is used only in the configuration as supplied.

If the SAM suction unit is used with cables / accessories other than those supplied, this may result in increased emissions or decreased immunity of the SAM suction unit in relation to EMC performance.

It should be noted that cables / accessories provided with the SAM suction unit should not be used on other equipment. To do so may result in increased emissions or decreased immunity of the other equipment in relation to EMC performance.

The SAM suction unit should not be used adjacent to or stacked with other equipment. If adjacent or stacked use with other equipment is necessary, the SAM suction unit and the other equipment should be observed/monitored, to verify normal operation in the configuration in which it will be used.

For the purposes of EN60601-1-2, the SAM suction unit has no essential performance.

The user should be aware that if the SAM suction unit is subject to electromagnetic interference in the form of electrical fast transients/bursts (IEC61000-4-4), electrostatic discharges (IEC61000-4-2) or conducted / radiated RF interference (IEC 61000-4-6 / IEC61000-4-3), incorrect operation of the SAM suction unit may occur. This could cause the vacuum to stop.
Guidance and manufacturer’s declaration – electromagnetic emissions

It is important that the SAM 35/36 is configured and installed/put into service, in accordance with the instructions/guidance provided herein and is used only in the configuration as supplied. For the purposes of EN60601-1-2:2014, the SAM 35/36 has no essential performance.

It should be noted that cables / accessories provided with the SAM 35/36 should not be used on other equipment. To do so may result in increased emissions or decreased immunity of the other equipment in relation to EMC performance.

<table>
<thead>
<tr>
<th>Emission test</th>
<th>Compliance</th>
<th>Electromagnetic environment - guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CISPR 11</td>
<td>Group 1</td>
<td>The SAM 35/36 uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment. If the SAM 35/36 is used with cables / accessories other than those supplied, this may result in increased emissions or decreased immunity of the SAM 35/36 in relation to EMC performance. The SAM 35/36 should not be used adjacent to or stacked with other equipment. If adjacent or stacked use with other equipment is necessary, the SAM 35/36 and the other equipment should be observed/monitored, to verify normal operation in the configuration in which it will be used.</td>
</tr>
<tr>
<td>Harmonic emissions</td>
<td>Class B</td>
<td></td>
</tr>
<tr>
<td>IEC61000-3-2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Voltage fluctuations / flicker</td>
<td>Complies</td>
<td></td>
</tr>
<tr>
<td>emissions IEC61000-3-3</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Guidance and manufacturer’s declaration – electromagnetic immunity

The purpose of this testing is to ensure the SAM 35/36 is not likely to adversely affect the normal operation of other such equipment and that other such equipment is not likely to adversely affect the normal operation of the SAM 35/36.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601 test level</th>
<th>Compliance level</th>
<th>Electromagnetic environment - guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic discharge (ESD)</td>
<td>±2kV, ±4kV, ±6kV and ±15kV air discharge</td>
<td>±2kV, ±4kV, ±6kV and ±15kV air discharge</td>
<td>The SAM 35/36 should not be used adjacent to or stacked with other equipment. If adjacent or stacked use with other equipment is necessary, the SAM 35/36 and the other equipment should be observed/monitored, to verify normal operation in the configuration in which it will be used.</td>
</tr>
<tr>
<td>IEC60601-1-2:2014 IEC61000-4-2:2008</td>
<td>±6kV contact discharge</td>
<td>±6kV contact discharge</td>
<td></td>
</tr>
<tr>
<td>Electrical fast transient / burst</td>
<td>± 2 kV for power supply lines</td>
<td>± 2 kV for power supply lines</td>
<td>Mains power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>IEC60601-1-2:2014 IEC61000-4-4:2012</td>
<td>± 1 kV for input / output lines</td>
<td>Input / output line tests not applicable.</td>
<td></td>
</tr>
<tr>
<td>Surge</td>
<td>Test level 1 ±0.5kV, ±1.0kV</td>
<td>Test level 1 ±0.5kV, ±1.0kV</td>
<td>Mains power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>IEC60601-1-2:2014 IEC61000-4-5:2005</td>
<td>Test level 2 ±2 kV</td>
<td>Test level 2 ±2 kV</td>
<td></td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines</td>
<td>Test level 1 Reduction of the supply voltage of 100% for a half and 1 periods</td>
<td>Test level 1 Reduction of the supply voltage of 100% for a half and 1 periods</td>
<td>Mains power quality should be that of a typical commercial or hospital environment. If the user of the SAM 35/36 requires continued operation during power mains interruptions, it is</td>
</tr>
</tbody>
</table>
### Guidance and manufacturer's declaration – electromagnetic immunity

The **SAM 35/36** is intended for use in the electromagnetic environment specified below. The customer or the user of the **SAM 35/36** should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity to RF test</th>
<th>Electromagnetic environment - guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Basic standard</strong></td>
<td>If the <strong>SAM 35/36</strong> is used with cables / accessories other than those supplied, this may result in increased emissions or decreased immunity of the <strong>SAM 35/36</strong> in relation to EMC performance.</td>
</tr>
<tr>
<td>IEC60601-1-2:2014 (table 9)</td>
<td><strong>The <strong>SAM 35/36</strong> should not be used adjacent to or stacked with other equipment. If adjacent or stacked use with other equipment is necessary, the <strong>SAM 35/36</strong> and the other equipment should be observed/monitored, to verify normal operation in the configuration in which it will be used.</strong></td>
</tr>
<tr>
<td><strong>Test setup</strong></td>
<td>Despite the testing of the <strong>SAM 35/36</strong> that has been undertaken, normal operation of the <strong>SAM 35/36</strong> can be affected by other electrical/electronic equipment and portable and mobile RF communications equipment.</td>
</tr>
<tr>
<td>IEC61000-4-3:2006; Amd, 1:2007 and Amd, 2:2010</td>
<td><strong>Interference may occur in the vicinity of equipment marked with the following symbol:</strong> <img src="image" alt="Symbol" /></td>
</tr>
</tbody>
</table>

**NOTE U<sub>T</sub>** is the a.c. mains voltage prior to application of the test level.
These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

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.

a Field 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 SA 35/36 is used exceeds the applicable RF compliance level above, the SA 35/36 should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orientating or relocating the SA 35/36.

b Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.

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**Recommended separation distances between Portable and mobile RF communications equipment and the SA 35/36**

The SA 35/36 is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the SA 35/36 can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the SA 35/36 as recommended below, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>Rated maximum output power of transmitter W</th>
<th>Separation distance according to frequency of transmitter m</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>150 KHz to 80 MHz</td>
</tr>
<tr>
<td>0.01</td>
<td>$d = 1.2 \sqrt{P}$</td>
</tr>
<tr>
<td>0.1</td>
<td>0.12</td>
</tr>
<tr>
<td>1</td>
<td>0.38</td>
</tr>
<tr>
<td>10</td>
<td>1.2</td>
</tr>
<tr>
<td>100</td>
<td>3.8</td>
</tr>
<tr>
<td>100</td>
<td>12</td>
</tr>
</tbody>
</table>

For transmitters rated at a maximum output power not listed above, the recommended separation distance $d$ in metres (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.
7. NOTES
8. OTHER PRODUCTS IN THE SAM RANGE

Mains Powered Suction

SAM 12 - General ward high vacuum suction unit
SAM 14 – Twin jar minor operating theatre high vacuum suction unit
SAM 15 - Intra-uterine aspirator suction unit
SAM 16 – Twin jar intra-uterine aspirator suction unit
SAM 17T - Thoracic Theatre suction unit
SAM 17W - Thoracic Ward suction unit
SAM 18 - Intensive care low vacuum suction unit
SAM 19 - Twin jar intensive care low vacuum suction unit
SAM MS – Micro suction unit

Portable Suction

SAM HOSPY - General high vacuum suction unit
SAM EPS - Battery powered portable suction unit (Neonatal Option available)
SAM MANUVAC - Portable foot operated suction unit
SAM TVAC - Disposable, hand operated suction unit

Oxygen Flowmeters

SAM OXYFLOW - Oxygen flowmeter
SAM OXYHUM - Oxygen humidifier

Pipeline Regulators

SAM 50 - High vacuum pipeline regulator with remote probe
SAM 51 - High vacuum pipeline regulator with direct probe
SAM 52 - Low vacuum pipeline regulator with remote probe
SAM 53 - Low vacuum pipeline regulator with direct probe
SAM 54 - High vacuum pipeline regulator - remote probe & mobile trolley

Research and Development

Since 1954, when MGE produced their first surgical suction units, the SAM range has become accepted as the industry standard, both in the U.K. and throughout the world. In recent years, the SAM range has been greatly extended, with models now available for portable, electrical suction and central suction requirements. They have been completely re-designed, using lightweight, robust materials, achieving greater efficiency, and making them easier to clean and operate. All MGE equipment is manufactured and assembled to very high standards of quality at the modern factory in Colchester in accordance with BS EN ISO 9001 Quality Management System, BS EN ISO 13485 and the Medical Device Directive 93/42/EEC.

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