

Model sam-12 Ultrasonic Diathermy Device

> Directions for Use

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1. Introduction

Thank you for choosing the sam Ultrasonic Diathermy Device! This manual contains general instructions for operation, application, precautions, and maintenance. In order to obtain maximum life and efficiency from the sam Device and to assist in the proper operation of the device, please read and understand this manual thoroughly. This device is only to be used as directed in this manual.

sam was developed as a next generation wearable ultrasound therapy system which combines miniaturization technology into a small and portable ultrasound therapy system. It is designed to work with the human body and maximize the safe and effective delivery of long-duration therapeutic ultrasound. Simple to administer and operate on a broad range of body types, sam allows the delivery of ultrasound treatment for up to four hours. It operates at a preset frequency and allows the use of up to two applicators simultaneously. Applicators are applied and secured to the surface of the body using convenient sam Ultrasound Coupling Bandages.

The specifications put forth in this manual were in effect at the time of publication.

1.1. GENERAL SAFETY

Thoroughly read and understand the precautionary and operating instructions before attempting to operate the sam Ultrasonic Diathermy Device. Know the limitations and hazards associated with using any ultrasound device. Observe the precautionary and operational decals placed on the device. Periodically review the operation procedures and safety precautions outlined in this manual.

1.2. PRESCRIPTION USE ONLY

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

This device complies with 21 C.F.R. § 1050.10

2. Indications for Use

The sam Ultrasonic Diathermy Device is intended to apply ultrasonic energy to generate deep heat within body tissues for the treatment of selected medical conditions such as the relief of pain, the relief of muscle spasms, the treatment of joint contractures and the local increase of circulation.

The sam Ultrasonic Diathermy Device is a prescription use device. sam should only be administered and monitored by a licensed healthcare practitioner.



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3. Safety

3.1. CONTRAINDICATIONS

Contraindications for the use of ultrasound include:

- · Over an area of the body where a malignancy is known to be present
- Over the eyes
- · Over or near growth centers until bone growth is complete
- · Over the reproductive organs
- · Over the pregnant uterus
- · Over a healing bone fracture
- · On the thoracic area if the patient is using a cardiac pacemaker
- Over an active implanted medical device such as an implanted deep brain stimulation device
- · On the brain, spinal cord, or large subcutaneous peripheral nerves
- Ischemic tissues in individuals with vascular disease where the blood supply would be unable to follow the increase in metabolic demand and tissue necrosis might result

3.2. WARNINGS

- If the treatment is reported as painful or too hot at any point during treatment, turn off device and remove the device from the skin.
- Instruct the patient to inform the practitioner if the patient feels any pain or burning during treatment.
- Instruct the patient how to turn off the sam Device and remove the sam Applicator if the patient feels any pain or burning during treatment.
- If the *Lock Switch* is in the locked position it must be placed in the unlocked position to disable the power. Locking the treatment settings is optional and not required for treatment.
- ALWAYS administer treatment using a new sam Ultrasound Coupling Bandage. Use one sam Ultrasound Coupling Bandage per applicator. Use of the sam Ultrasound Applicator without a new sam Ultrasound Coupling Bandage MAY RESULT IN BURN and/or REPEATED SHUTOFF of the sam Applicator.
- The sam Device should be kept out of the reach of children.

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- DO NOT apply the sam Applicator with alternative coupling media as a replacement for the sam Ultrasound Coupling Bandage. Use of alternative coupling media in lieu of the sam Ultrasound Coupling Bandages may reduce the effectiveness of treatment, lead to automatic shutoff of the applicator, or cause a burn.
- DO NOT administer treatment if the applicator is not connected to a sam Ultrasound Coupling Bandage.
- Applicators and sam Ultrasound Coupling Bandages are not sterile. **DO NOT** apply this device over an open wound or inflamed skin.
- DO NOT use the sam Ultrasound Coupling Bandage if the sam Ultrasound Media is dried out. Indications of a dried out bandage include: the cup is not full of gel, there is dry residue or film in the cup, or there is any cut, break, or opening in the bandage or seals.
- DO NOT apply directly over a bone that is near the skin surface.

3.3. PRECAUTIONS

Precaution should be taken when using the device:

- Over an area of the spinal cord following a laminectomy, i.e. when major covering tissues have been removed
- · On patients with hemorrhagic diatheses
- Over areas where metal prosthesis or other metallic implants are embedded in tissue which may form a reflective surface to the ultrasound energy causing unintended irradiation of tissue and excessive heating
- · Over an acute infection or sepsis
- · On patients with peripheral artery disease
- Over a deep vein thrombosis
- Over an anesthetized area or in conjunction with a condition that causes impairment of sensation, such as caused by chemotherapy
- When using the sam Ultrasound Coupling Bandage, ensure the top and bottom seals have been completely removed before attaching the applicator to the skin

3.4. INFLAMMABLE GASES AND ANESTHETICS

 Warning: Explosion hazard if used in the presence of flammable anesthetics, open flame, or oxygen-rich environment



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3.5. ELECTRONICS AND BATTERY

WARNINGS

- This device is rated IPX-0; therefore, it is **Not Waterproof**. **DO NOT** apply a direct stream of any liquid onto the device, submerge the device, or allow any liquid to pool on the surface of the device. **DO NOT** use if device has been submerged in water.
- This device contains a rechargeable lithium-ion battery. **DO NOT** disassemble, **DO NOT** heat above 100°C, **DO NOT** incinerate or expose to water and **DO NOT** ingest.
- The use of accessories, transducers and/or cables other than those specified, with the exception of those sold by the manufacturer as replacement parts for internal components, may result in increased emissions or decreased electrical immunity of the equipment or system.
- DO NOT open or modify any component of the sam Device. Hazards such as shock, burn or inappropriate functionality can result from unauthorized modification of the sam Device.
- Use of the sam Device around electromagnetic interference may negatively affect the output performance and safety of the device. Do not use the device if any abnormal functionality occurs.

PRECAUTIONS

- Only recharge the sam Device using the sam Electrical Charger. Use with any other recharging device may result in damage to the system and void all warranties.
- When not in use, power 'OFF' the device to protect the functionality of the components.
- Avoid dropping the applicator or power controller and avoid scratching the lens of the applicator. Rough handling may reduce the device's acoustic output power, thereby reducing the effectiveness of therapy.
- The power controller and applicators should be routinely checked for cracks and other damage before each use to determine that the device functions normally.
- DO NOT place the device in a location where the power charging cord could be a trip hazard.
- DO NOT use sharp objects such as a pencil point or ball point pen to operate the buttons on the control panel as damage may result.
- Prevent potential electromagnetic or other interference. **DO NOT** open the sam Device or connect the device or components of the device to any non-sam part. Keep the device clean and make sure no exposed non-insulated wires are visible. If damage is present, **do not** administer treatment.

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4.1. PRESET TREATMENT

The sam Device is preconfigured to provide continuous ultrasonic output at a preset frequency and intensity which cannot be modified by the user. The user can set the treatment duration to be 1, 2, 3, or 4 hours.

4.2. sam ULTRASOUND APPLICATORS

The sam Ultrasound Applicators serve as the ultrasound transducers of the sam Device. The applicators offer low-profile design with light emitting diode (LED) on/ off notification. The ergonomic plastic housing and smooth contours provide enhanced comfort.

4.3. sam SENSING

sam is designed to work with the human body and maximize the safe and effective delivery of long-duration therapeutic ultrasound. Each sam Applicator is equipped with closed-loop continuous temperature monitoring which maintains treatment site temperatures below 44°C during normal operation. See section 13.4 on page 24 for more information.

4.4. sam ULTRASOUND COUPLING BANDAGES

The sam Device utilizes ultrasound coupling bandages which are manufactured with ultrasound coupling media sealed inside. The ultrasound coupling bandages ARE REQUIRED to secure the sam Applicators to the body.

4.5. BATTERY OPERATION

Powered by a rechargeable lithium-ion battery, the sam Device can provide 4 hours of therapy on a single battery charge.

4.6. LOCK SWITCH

The sam Power Controller Module includes a slide lock switch that allows the locking of treatment settings so that they cannot be inadvertently modified during a 1 – 4 hour treatment session. Locking the sam Power Controller Module will disable both the treatment time toggle button and the power ON/OFF button. To disable the device or modify treatment time once in locked mode, the lock switch must be moved to the unlocked position. Similarly, the device cannot be turned on while in locked mode and must be unlocked first. The use of the locking feature is OPTIONAL.



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5. sam Components

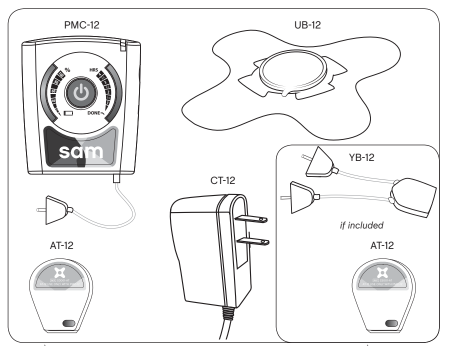


figure 1: sam Model-12 Components

sam MODEL 12 COMPONENTS

| Ultrasound Applicators |
|------------------------------|
| Ultrasound Coupling Bandages |
| User Manual |
| Power Controller Module |
| Electrical Charger |
| Y-branch Adapter |
| |

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6. Operator Interface

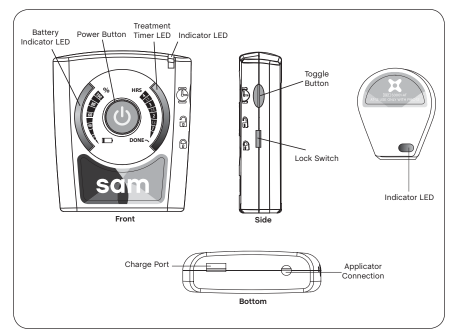


figure 2: sam Interface Components

7. Accessories

The sam Device may be used with any of the following accessories:

sam Belt Clip

sam Armband

sam Carrying Case

* Additional sam Ultrasound Coupling Bandages may be obtained by contacting the manufacturer



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8. LED Display

The sam Device contains Light Emitting Diode (LED) displays that indicate the functions of the device (see Figure 2).

8.1. POWER CONTROLLER INDICATOR LED

The Indicator LED provides power, charging and error checking information.

| Color of Indicator LED | Meaning |
|---------------------------|---|
| Opaque (absence of light) | Power Controller is OFF. |
| | No power is being generated |
| Blue | Power Controller is ON. Note – Ultrasonic |
| | energy is generated only when the Power |
| | Controller is connected to the sam |
| | Applicator(s) |
| Green | Power Controller battery is fully charged |
| Amber | Power Controller battery is charging |
| Red | Too many applicators are attached. A |
| | maximum of two applicators may be |
| | attached to a single power controller at |
| | any time |

Table 1. Power Controller LED Color Definitions

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8.2. CONTROL PANEL

The front face of the sam Power Controller Module contains a control panel. The LEDs on the control panel are divided into a field for *Battery Indicator* and a field for *Treatment Timer*. The operator is able to make selections by pressing the *Power Button* and the *Toggle Button*.

| Color of Battery Indicator LED* | Meaning | |
|---|---------------------------------------|--|
| Red Only | Low Battery, Re-Charge required | |
| Blinking Red | Re-Charge required | |
| Amber | Power Controller is partially charged | |
| Green Power Controller fully charged | | |
| *These lights remain on for 4 seconds and subsequently turn off | | |

Table 2. Battery Indicator LED Color Definitions

| Treatment Time Indicator LED | Meaning |
|------------------------------|---|
| Blue | Treatment time remaining. Treatment |
| | timer is set in one hour increments and |
| | will count down in 30 minute increments |
| Amber | Treatment complete. This single "done" |
| | light will activate only when a treatment |
| | cycle has completed and will remain |
| | illuminated for 8 hours, then turn off |

Table 3. Treatment Time Indicator LED Color Definitions

8.3. APPLICATOR INDICATOR LED

| LED Color | Meaning |
|--------------------------------|--|
| Opaque (absence of light) | Applicator is not receiving power |
| Blue | Applicator is receiving power from the |
| | Power Controller. Ultrasonic energy is |
| | being generated at all times this light |
| | is illuminated |
| Red (accompanied by vibration) | sam Sensing Mode. No Ultrasonic energy |
| | is being generated from this applicator while this light is illuminated |

Table 4. Applicator LED Color Definitions



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9. Initial Setup Instructions

Remove the sam Device components (Figure 1) from the packaging and inspect for any damage that may have occurred during shipment. Charge the device for up to 6 hours or until the Power Controller Indicator LED is Green. The sam Device arrives only partially charged due to generally accepted shipping practices.

9.1. CHARGING THE POWER CONTROLLER MODULE

- A. Plug the micro USB end of the sam Electrical Charger into the charging port on the bottom of the sam Power Controller.
- B. Plug the electrical charger into a 120/230 VAC wall outlet. The sam Power Controller Module indicator LED will be amber. When the device is fully charged the top right indicator LED will change from amber to green.

10. Treatment Options

10.1. TREATMENT DURATION

The sam Ultrasonic Diathermy Device provides ultrasound therapy at a preset frequency and intensity. The user can program the treatment duration to be 1, 2, 3, or 4 hours. The maximum treatment duration setting on the sam Device is 4 hours. Treatment duration should be set to the minimum increment required for effective therapy. Due to individual differences in skin type and tolerance, it is recommended to begin with a 1 hour treatment. Increase treatment duration in subsequent applications only as tolerated. Maximum usage time is 4 hours per day per treatment area.

The sam Device may be used while charging. Charging the device during treatment has no impact on device frequency or intensity output settings. If using sam while charging, position the device in such a way that the charger cable is easily removed from the power controller module if needed.

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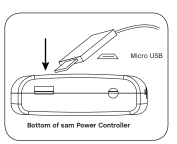


figure 3A

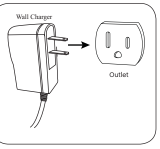


figure 3B

10.2. TREATMENT LOCATIONS

Figures 4 and 5 depict examples of device placement on two treatment locations. Figure 4A and 4B illustrates using the sam Device with one applicator on the shoulder and knee. Figure 5A and 5B illustrates using the sam Device with two applicators on the shoulder and knee. When determining treatment location, consider that the maximum diathermic effect will occur directly under the applicator face; therefore, placing the applicator near or directly over the target area while following all warning and cautionary instructions is advised.

- Note: These figures are examples and are not intended to be the suggested or only allowable applicator configurations for those body locations.
- Warning: Do Not apply directly over a bone that is near the skin surface.

10.3. ONE OR TWO APPLICATORS:

The sam Device may be used with one or two applicators simultaneously. The decision to use one or two applicators is dependent on the size and anatomical area of treatment. Using two applicators allows a larger area of tissue to receive ultrasound treatment. For example, large anatomical regions, such as the shoulder could benefit from two applicators, whereas smaller treatment areas such as the forearm may only require one.

10.3.1. SINGLE-APPLICATOR MODE

Connect the power controller wire to a single applicator by inserting the power controller wire jack into the matching cavity at the base of the applicator. The port for the power controller wire is described as 'Applicator connection' on the Power Controller drawing shown in Figure 2, page 10.

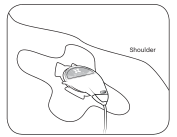


figure 4A: Single Shoulder Application

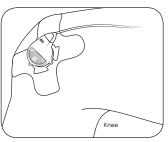


figure 4B: Single knee Application



figure 5A: Dual Shoulder Application

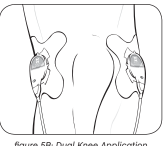


figure 5B: Dual Knee Application

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10.3.2. DUAL-APPLICATOR MODE

The Y-adapter (optional) may be used to power two Applicators simultaneously. As shown in Figure 8, connect the Y-adapter directly to the Power Controller wire and then connect each Applicator to a wire jack at the other end of the Y-adapter.

When using two sam Applicators and two sam Ultrasound Coupling Bandages simultaneously, the sam Ultrasound Coupling Bandages should be positioned so that their footprints do not overlap.

Caution: DO NOT overlap sam Ultrasound Coupling Bandages (Figure 6).

Caution: DO NOT use the Y-adapter when using only one applicator for treatment.

Caution: Do NOT attach more than one Y-adapter to the sam Device.

Note: If two applicators will be used, two sam Ultrasound Coupling Bandages (one for each applicator) and the Y-adapter must be used for the treatment.

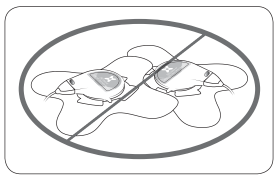


figure 6: NO Bandage Overlap

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11.1. CHECK DEVICE CHARGE

- A. To prevent accidental activation during shipping, the device will arrive to the user in the 'locked' position. To unlock the device, slide the *Lock Switch* on the sam Power Controller 'up' into the 'unlocked' position (Figure 7A).
- B. Check to make sure the Power Controller holds enough charge to provide the desired duration of treatment. Press the *Toggle Button* on the sam Power Controller to view the battery indicator lights (Figure 7B). The lights will illuminate to show how much battery life is remaining in the device. If the device is not fully charged it may not be able to deliver treatment for 4 hours. The device will only permit the user to set a treatment duration for which the battery charge is capable of fulfilling.
 - i. To check the treatment duration for which the battery charge is capable of fulfilling, simply continue to press the *Toggle Button* sequentially to view all timer settings allowed by the current battery charge. See section 9 for charging instructions.

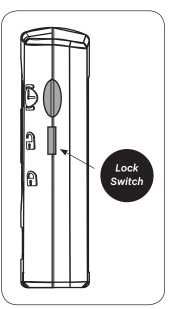
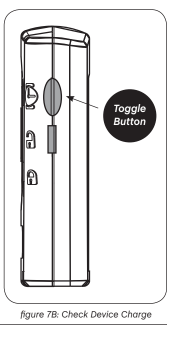


figure 7A: Check Device Charge





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11.2. CONNECT APPLICATOR TO POWER CONTROLLER

* If using 2 applicators: first connect the Y-adapter directly to the power controller wire jack and then connect each applicator to a wire jack at the other end of the Y-adapter (Figure 8*).

- A. Insert the *power controller wire* into the matching cavity at the base of the applicator at a 45 degree angle (Figure 8A).
- B. **Gently** twist the wire jack clockwise until the applicator edge is flush with the wire jack (Figure 8B).

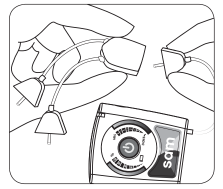


figure 8*: Connecting Y-Adapter

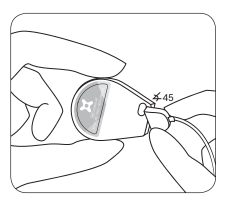


figure 8A: Insert power controller wire at 45° angle

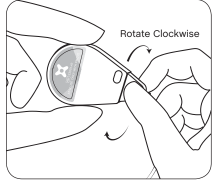


figure 8B: Gently twist to Lock

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11.3. ATTACH APPLICATOR TO BANDAGE

- A. Remove the circular seal from the top of the bandage to reveal the coupling media within the gel cup (Figure 9A).
- B. Attach the applicator, face down, into the coupling media (Figure 9B).
- C. Firmly press the applicator down onto the center of the gel cup until a clicking noise is heard or clicking sensation is felt (Figure 9C).

If using dual applicators, repeat these steps with the second applicator and the second bandage.

- Note: When the applicator is pressed down onto the gel cup, do not be concerned if coupling media flows out the edges of the gel cup. Wipe any excess coupling media away with a tissue or towel.
- Note: Ensure the applicator is fully attached to the bandage.

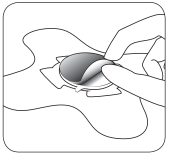


figure 9A: Removing Foil Seal

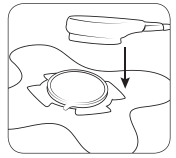


figure 9B: Attach Applicator

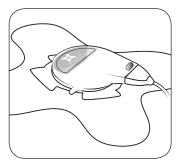


figure 9C: Press Applicator Down



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11.4. APPLY ULTRASOUND COUPLING BANDAGE TO TREATMENT LOCATION

- A. Hold the applicator so the bottom of the bandage faces up. Remove the paper liner from the back of the bandage, revealing the bandage adhesive (Figure 10A).
 - i. The circular seal should peel off the bottom of the gel cup along with the paper bandage liner. If it does not peel with the paper, be sure to peel away the circular seal so the coupling media is completely uncovered.
 - ii. Add more sam Coupling Media as needed.
- B. Turn the bandage over and adhere the bandage to the desired treatment location (Figure 10B).
 - i. The sam Coupling Media must be in direct contact with the skin.

If using dual applicators, repeat these steps to adhere the second bandage and applicator to the second treatment site.

Note: When applying bandages, both sides of the ultrasound coupling media should be uncovered. One side should be in contact with the skin. The other side should be in contact with the face of the applicator.

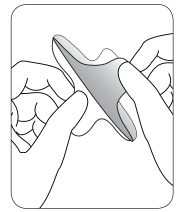


figure 10A: Removing Paper Liner

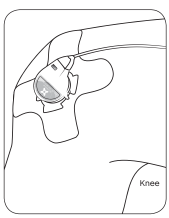


figure 10B: Adhere Bandage to Site

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11.5. TURN THE DEVICE 'ON'

- A. Check to make sure the *Lock Switch* is in the 'unlocked' position (Figure 11A).
- B. Press and hold the *Power Button* on the sam Power Controller for at least 1 second to turn the device 'ON' (Figure 11B).
 - i. As confirmation that the device is 'ON', the LEDs on the sam Power Controller will illuminate blue (treatment timer and indicator LEDs) and the indicator LED on the applicator will illuminate blue.
 - ii. The battery indicator LEDs will remain 'ON' for about 4 seconds after turning 'ON' the device.
 - iii. The treatment timer LEDs and blue indicator LEDs (power controller and applicator) will remain 'ON' for the duration of treatment.
- C. Press the *Toggle Button* to select 1, 2, 3 or 4 hour treatment duration (as allowed by current battery charge) (Figure 11C).
 - i. If the *Toggle Button* is pressed up to the 4 hour setting and a decreased treatment time is desired, pressing it again will return the treatment timer to the 1 hour setting.

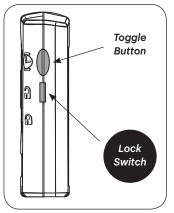


figure 11A: Slide to Unlocked Position



figure 11B: Press Power Button

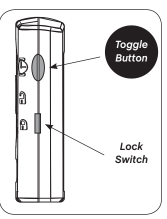


figure 11C: Select Duration



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11.6. PREVENT UNINTENTIONAL SHUTOFF

Once satisfied with the treatment settings, the settings may be locked to prevent unintentional shutoff during treatment by sliding the *Lock Switch* down into the 'locked' position as indicated by the symbols on the sam Power Controller (see Figure 11A). In this state the settings cannot be changed until the *Lock Switch* is moved back to the 'unlocked' position.

Note: The system is not required to be locked in order to operate. Always be aware of how to disable a locked device. Always notify the patient if their device is locked.

11.7. TREATMENT DELIVERY

Treatment will be delivered for the duration set by the user. The treatment timer LEDs will count down in 30 minute increments as treatment progresses. Upon completion, the device will automatically shut off and the 'treatment complete' LED will illuminate for approximately 8 hours before turning off. See section 12 for how to manually power 'OFF' the device.

- Warning: If the sam Lock Switch is in the locked position, the sam Power Button will not turn off the device until the sam Lock Switch is placed into the unlocked position.
- Warning: ALWAYS instruct the patient to alert the practitioner in the event of discomfort during treatment.
- Warning: As an additional safety measure, ALWAYS instruct the patient how to turn off the sam Device and remove the sam Applicator in the event of discomfort or emergency.
- Warning: ALWAYS keep the sam Power Controller within easy reach so that power may be ceased at any time.

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12.1. POWER OFF THE POWER CONTROLLER

Move the *Lock Switch* up to the 'unlocked position' (if locked) then press and hold the *Power Button* on the sam Power Controller *for at least 1 second* to turn the device 'OFF' (See Figure 11B for device diagram).

12.2. REMOVE DEVICE FROM SKIN

- A. Remove the bandage(s) from the skin.
- B. Remove the applicator from the sam Ultrasound Coupling Bandage by pulling the tab on the side of the gel cup. This unlocks the cup from the applicator (See Figure 12).
- C. Throw away the sam Bandages.
- Warning: NEVER use a sam Ultrasound Coupling Bandage for more than one use. Misuse of sam Ultrasound Coupling Bandages or use for more than the intended treatment duration MAY RESULT IN BURN OR REPEATED SHUTOFF of the applicator.

12.3. CLEAN THE APPLICATORS

Clean any residual ultrasound coupling media off of the applicator and off of the skin. See section 14 for 'Cleaning and Maintenance' details.

12.4. RECHARGE THE DEVICE

Connect the sam Power Controller to the sam Electrical Charger for recharging. Prior to the next use, allow up to 6 hours to recharge the battery to 100% charge. See section 9 for charging instructions.

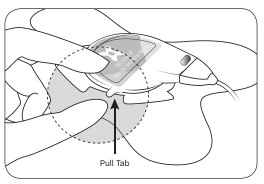


figure 12: Removing Applicator



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13.1. ON (UNLOCKED) MODE

To turn the device 'ON', first ensure the Lock Switch is in the unlocked position as indicated by the unlocked symbol on the sam Power Controller Module. Press and hold the *Power Button* for at least 1 second. Press the *Toggle Button* on the right side of the sam Power Controller Module to set the treatment duration. The treatment duration may be set for 1, 2, 3 or 4 hours. The battery indicator LED and treatment timer LED will be displayed on the control panel (see section 8: LED Display). The applicator indicator LED will be displayed on each attached applicator (See Figure 2, page 10). Each connected applicator will emit ultrasound energy immediately upon the sam Power Controller Module entering the 'ON' mode of operation. The sam Applicator will emit ultrasonic energy whether the device is locked or unlocked (locking is optional). A powered sam Applicator will always emit ultrasonic energy unless it enters the sam Sensing Mode or the treatment duration concludes.

13.2. ON (LOCKED) MODE

Once the device is powered 'ON', the user may lock the control panel settings by sliding the *Lock Switch* on the sam Power Controller Module to the 'locked position' as indicated by the lock symbol on the sam Power Controller Module. This will prevent the device from unintentional shutoff during treatment. While locked, the *Power Button* and the *Toggle Button* cannot be modified. The sam Applicator will emit ultrasound whether the device is locked or unlocked. To change the treatment duration or turn the power off, simply slide the *Lock Switch* to the 'unlocked position' as indicated by the symbols on the sam Power Controller Module then press the desired user button.

Warning: If the Lock Switch is in the locked position, the Power Button will not turn off the device until the Lock Switch is placed into the unlocked position. Locking the treatment settings is not required.

13.3. OFF MODE

When the device is 'OFF', the LEDs on the applicator will not be illuminated, the LEDs on the sam Power Controller Module will not be illuminated, and no ultrasound energy will emanate from the device. Always turn the device 'OFF' when not in use or if pain or uncomfortable heating is indicated by the patient.

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13.4. SAM SENSING MODE

In the event that the treatment site underneath the applicator reaches the temperature threshold, the applicator will pause ultrasound output and vibrate once with a red LED notification to signify that the device has begun a cooling or rest cycle. sam Applicator will automatically resume treatment with a blue LED notification after the site has cooled.

Warning: Should the patient ever report discomfort or a painful sensation from the site under the sam Applicator, the applicator should be removed immediately.

Caution: At no time during treatment should the applicator be covered by thick insulating material such as a coat, blanket or sports wrap. This may cause sam to disable and remain disabled throughout the therapy session.

13.5. END OF TREATMENT MODE

At the end of treatment, an amber-colored "Treatment Complete" LED will illuminate on the sam Power Controller Module and all other displays (battery indicator, treatment timer, indicator LED, and applicator LEDs) will be inactive. All ultrasound emissions from the applicator will cease. The "Treatment Complete" LED will remain illuminated for approximately 8 hours after completion of therapy, at which point it will time out and turn off. To set the sam Device for a new treatment, remember to move the lock switch to the unlocked position and assess whether the sam Device requires recharging.

14. Cleaning and Maintenance

The exterior of the sam Power Controller and the applicator surfaces may be cleaned with a soft cloth, tissue, or towel and one of the following cleaning agents: mild detergent and water or disinfecting medical wipes.

Caution: Properly clean the applicator between treatments.

- Caution: The device is not waterproof. Do not apply a direct stream of liquid onto the device, submerge the device, or allow any liquid to pool on the surface of the device.
- Caution: DO NOT USE: Phenolic-based disinfectants, quaternary ammonium, chlorinebased disinfectants, solvent-based cleaners, or abrasive materials. Doing so may damage the plastic housing and void the warranty.

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15. Storage & Operating Conditions

15.1. STORAGE

Store the sam Device in the following conditions:

| Temperature: 5-57 °C; | | |
|--|--|--|
| Humidity: 10-80%; | | |
| Atmospheric pressure range: 700-1060 hPa | | |

Store the sam Ultrasound Coupling Bandages in the following conditions:

| Temperature: 5-30 °C; | |
|--|--|
| Humidity: 10-80%; | |
| Atmospheric pressure range: 700-1060 hPa | |

15.2. OPERATION

Only operate the sam Device in the following conditions:

| Temperature: 1-44 °C; | |
|--|--|
| Humidity: 10-80%; | |
| Atmospheric pressure range: 700-1060 hPa | |

Caution: Do not keep the device in extreme hot or cold temperatures (above 50°C or below 0°C). Do not leave the sam Device in a hot or freezing car. Do not leave the device in direct sunlight for extended periods. UV light may damage or discolor the device.

16. Disposal of Waste Products

sam Ultrasound Coupling Bandages are one time use and may be disposed of in regular sanitation trash. No special disposal procedures are necessary.

Old, damaged or expired sam Power Controller Modules and applicators should be recycled or returned to the manufacturer for proper disposal.

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17. Appendix

A. SYMBOLS

| | Consult User Manual/Instructions for Use |
|------------------|---|
| | Manufacturer/Date of Manufacture |
| Li-ion | Lithium-ion battery inside |
| * | Class BF Applied Part |
| | Do not use if package is damaged |
| \otimes | Do not reuse |
| $((\mathbf{c}))$ | Non-ionizing radiation |
| X | Separate collection for electrical and electronic equipment. Must not be disposed of in unsorted municipal waste. |
| \triangle | Caution, consult accompanying documents |
| ((`• | Diverging beam |
| ·vvvv | Continuous wave (CW) |
| W | Watts, ultrasonic power |
| MHz | Frequency in megahertz |
| ERA | Effective radiating area |
| BNR | Beam non uniformity ratio |
| | Use by date |
| X | Temperature limits |
| <u>%</u> | Humidity limitation |
| SN | Serial number |
| REF | Catalogue/Reorder Number |
| LOT | Lot number |
| | |



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B. SPECIFICATIONS

| Maximum Acoustic | 0.65 W ±20% per transducer | |
|---------------------------------|--|--|
| Power Output | 1.3 W ±20% for 2 transducers | |
| Maximum Intensity | 0.132 W/cm ² ±20% | |
| Frequency | 3 MHz ±20% | |
| Duty Cycle | 100% - continuous wave | |
| Beam Form | Wide Beam - 5 degree diverging lens | |
| Individual Transducer Dimension | 5 cm ² emitting surface area (circular) | |
| BNR | <5:1 | |
| ERA | 6 cm ² ±20 % | |
| Maximum Treatment Duration | 4 hours | |

The sam Device does not contain microprocessor or software

Other Electrical Ratings for the sam Device Components Electrical Charger (Model GS2U-006-050-A)

| _ | |
|------------------|---------------------|
| Input Voltage | 100-240 V, 50-60 Hz |
| Input Current | 0.2 A |
| Output Voltage | 5 V DC |
| Output Current | 1.2 A |
| Max Output Power | 6 W |

Power Controller

| Output Voltage | 3.7 V DC ±10% |
|------------------------|---------------------|
| Max Output Amperes | 700 mA |
| Battery Protection | Max Current: 2 Amps |
| | Max Voltage: 4.2 V |
| | Min Voltage: 3.1 V |
| Cable Voltage Rating | 300 V |
| Cable 20°C Resistance: | 94/km |

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Applicator

| Input Voltage: | 3.7 V DC ±10% | |
|--------------------------|------------------------------------|--|
| Ultrasound Frequency: | 3 MHz ±20% | |
| Ultrasonic Output Power: | 1 Transducer: 0.65 W ±20% | |
| | 2 Transducers Together: 1.3 W ±20% | |
| Max Input Current: | 400 mA | |

Y-Adapter

| Voltage Rating | 300 V |
|-----------------|-------|
| 20°C Resistance | 94/km |

C. NOTICE OF COMPLIANCE, CALIBRATION AND OPERATIONAL PERIOD

The sam Device meets performance standards under 21 C.F.R. § 1050.10 -PERFORMANCE STANDARDS FOR SONIC, INFRASONIC, AND ULTRASONIC RADIATION-EMITTING PRODUCTS. The sam Device meets IEC 60601-1, 3 ed. (2012); IEC 60601-1-2, 3 ed. (2007); Medical Electrical Equipment, General Requirements for Safety; Electromagnetic Compatibility. The sam Device is calibrated to provide 3 MHz ±20% ultrasound diathermy at 0.65 W ±20% per ultrasound applicator.

Operational Period:

The sam Power Controller Module is intended to maintain calibration for up to 300 charging cycles and each ultrasound applicator is intended to withstand 1500 hours of run time, after which the system should be recycled or returned to ZetrOZ, Inc. for proper disposal.



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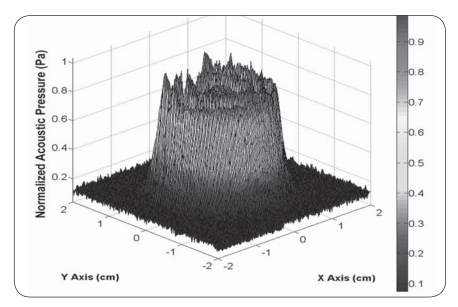


figure A. Ultrasound Field Scan Across Applicator Face

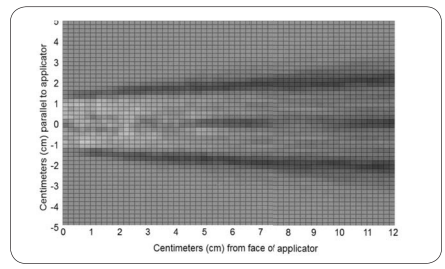


figure B. Ultrasound Field Scan Away From the Applicator Face

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E. ELECTRICAL IMMUNITY AND EMISSIONS

The following components of the sam Ultrasonic Diathermy Device are compliant with the requirements of IEC 60601-1-2 ed 3.0 (2007-03):

| sam Model 12 | |
|-------------------------------------|------------------|
| | Cable Length |
| AT-12: Ultrasound Applicators | |
| UB-12: Ultrasound Coupling Bandages | |
| PMC-12: Power Controller Module | 48 inches ±1.2 |
| CT-12: Electrical Charger | 60 inches ±2.0 |
| YB-12: Y-branch Adapter | 8 inches ±.6 x 2 |

Guidance and Manufacturer's Declaration - Emissions

| Emissions Test | Compliance | Electromagnetic Environment – Guidance | |
|----------------|------------|--|--|
| RF Emissions | Group 1 | The sam uses RF energy only for its internal | |
| CISPR 11 | | function. Therefore, its RF emissions are very low | |
| | | and are not likely to cause any interference in | |
| | | nearby electronic equipment. | |
| RF Emissions | Class B | The sam is suitable for use in all establishments, | |
| CISPR 11 | | including domestic, and those directly connected | |
| | | to the public low-voltage power supply network | |
| | | that supplies buildings used for domestic | |
| | | purposes. | |
| Harmonics | Class A | | |
| IEC 61000-3-2 | | | |
| Flicker | | | |
| IEC 61000-3-3 | | | |



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Guidance and Manufacturer's Declaration - Immunity All

ME Equipment and ME Systems

The sam is intended for use in the electromagnetic environment

specified below. The customer or user of the sam should ensure

that it is used in such an environment.

| Immunity Test | IEC 60601 | Compliance | Electromagnetic |
|----------------|-------------------|------------------|-----------------------------------|
| | Test Level | Level | Environment - Guidance |
| ESD | ±6kV Contact | ±6kV Contact | Floors should be wood, concrete |
| IEC 61000-4-2 | ±8kV Air | ±8kV Air | or ceramic tile. If floors are |
| | | | synthetic, the r/h should be at |
| | | | least 30%. |
| EFT | 2kV Mains | ±2kV Mains | Mains power quality should be |
| IEC 610000-4-4 | ±1kV I/Os | ±1kV I/Os | that of a typical commercial |
| | | | or hospital environment. |
| Surge | ±1kV Differential | ±1kV Differentia | Mains power quality should be |
| IEC 61000-4-5 | ±2kV Common | Common N/A | that of a typical commercial or |
| | | (no ground) | hospital environment. |
| Voltage | >95% Dip for | >95% Dip for | Mains power quality should be |
| Dips/Dropout | 0.5 Cycle | 0.5 Cycle | that of a typical commercial or |
| IEC 61000-4-11 | 60% Dip for | 60% Dip | hospital environment. If the user |
| | 5 Cycles | 5 Cycles | of the sam requires continued |
| | 30% Dip for | 30% Dip for | operation during power mains |
| | 25 Cycles | 25 Cycles | interruptions, it is recommended |
| | >95% Dip for | >95% Dip for | that the sam be powered from |
| | 5 Seconds | 5 Seconds | an uninterruptible power supply |
| | | | or battery. |
| Power | 3A/m | 3A/m | Power frequency magnetic fields |
| Frequency | | | should be that of a typical |
| 50/60Hz | | | commercial or hospital |
| Magnetic Field | | | environment. |
| IEC 61000-4-8 | | | |

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Guidance and Manufacturer's Declaration - Immunity

The sam is intended for use in the electromagnetic environment

specified below. The customer or user of the sam should ensure

that it is used in such an environment.

| Immunity Test | IEC 60601 | Compliance | Electromagnetic |
|---------------|------------|---------------|---|
| | Test Level | Level | Environment - Guidance |
| | | | Portable and mobile communications |
| | | | equipment should be separated from |
| | | | the sam by no less than the distances |
| | | | calculated/listed below: |
| Conducted RF | 3 Vrms | (V1) = 3 Vrms | D = (3.5/V1)(Sqrt P) |
| IEC 61000-4-6 | 150 kHz to | | 150 kHz to 80 MHz |
| | 80 MHz | | |
| Radiated RF | 3V/m | | D = (3.5/Eq)(Sqrt P) |
| IEC 61000-4-3 | 80 MHz to | (E1) = 3 V/m | 80 to 800 MHz |
| | 2.5 GHz | | |
| | | | D=(7/E1)(Sqet P) |
| | | | 800 MHz to 2.5 GHz |
| | | | Where P is the max power in watts and D |
| | | | is the recommended separation distance |
| | | | in meters. |
| | | | Field strength from fixed transmitters, |
| | | | as determined by an electromagnetic |
| | | | site survey, should be less than the |
| | | | compliance levels (V1 and E1). |
| | | | Interference may occur in the vicinity of |
| | | | equipment containing a transmitter. |



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Recommended Separation Distances between portable and

mobile RF Communications equipment and the sam ME Equipment and

ME Systems that are NOT Life-supporting

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

| Max Output | Separation (m) | Separation (m) | Separation (m) |
|---------------|--------------------|--------------------|--------------------|
| Power (Watts) | 150kHz to 80MHz | 80 to 800MHz | 800MHz to 2.5GHz |
| | D=(3.5/V1)(Sqrt P) | D=(3.5/E1)(Sqrt P) | D=(3.5/E1)(Sqrt P) |
| 0.01 | 0.11667 | 0.11667 | 0.23333 |
| 0.1 | 0.36894 | 0.36894 | 0.73785 |
| 1.0 | 1.1667 | 1.1667 | 2.3333 |
| 10.0 | 3.6894 | 3.6894 | 7.3785 |
| 100.0 | 11.667 | 11.667 | 23.333 |

F. WARRANTY

ZetrOZ, Inc. offers a 1-year manufacturer's warranty for the sam Device. If the sam Device fails due to defects in material or workmanship, ZetrOZ, Inc., at its discretion, will:

- 1. REPAIR the sam Device OR
- 2. REPLACE the sam Device with another sam Device

THIS LIMITED WARRANTY AND ANY IMPLIED WARRANTIES THAT MAY EXIST UNDER STATE LAW APPLY ONLY TO THE ORIGINAL PURCHASER OF THE sam DEVICE AND ARE NON-TRANSFERABLE.

Extent of Limited Warranty

This limited warranty does not cover damages due to external causes, including, without limitation, accident, usage not in accordance with product instructions, misuse, neglect, alteration or repair.

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G. TROUBLESHOOTING

| Question or Problem | Solution |
|--|---|
| 1. How to determine if the device is delivering treatment | As confirmation that the device is delivering ultrasound, the LEDs on the Power Controller will illuminate blue (treatment timer and indicator LEDs) and the indicator LED on the applicator will illuminate blue. See section 8 on LED Display. |
| 2. The device isn't turning ON and the buttons aren't working | Ensure the power controller lock switch is in the UNLOCKED position. If the device still fails to power ON/OFF or toggle treatment duration, contact the manufacturer with serial and model number details. |
| 3. The indicator LED on the Applicator does not illuminate when the device is turned ON | Ensure that the applicator is fully connected to the power controller wire jack and/or the Y-adapter wire jack. If recently in sam Sensing Mode, ensure that the applicator has had time to reach normal operating temperature and the indicator LED on the applicator has turned from red back to blue. If the applicator LED still fails to illuminate blue, contact the manufacturer with serial and model number details. |
| 4. How to observe the Battery level during treatment | The Battery Indicator display can only be viewed when the device is powered ON or OFF by pressing the <i>Power</i> <i>Button</i> , or when the <i>Toggle Button</i> is pressed. (See section 11.1) The lights remain illuminated for approximately 4 seconds after the device is turned ON or OFF. The Battery Indicator LEDs do not remain illuminated during treatment. The treatment duration is indicated by the blue treatment timer LEDs (See section 8 on LED display). |

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G. TROUBLESHOOTING

| Question or Problem | Solution |
|---|---|
| 5. The Applicator is not staying secured during treatment | Turn off the device. Remove the sam Ultrasound Coupling Bandage from the applicator. Replace with new sam Ultrasound Coupling Bandage. See section 11.3 for gel cup application instructions. |
| 6. The Applicators are vibrating and red | The device has entered 'sam Sensing Mode' (See section 13.4) Each sam Applicator is equipped with closed-loop continuous temperature monitoring which maintains treatment site temperatures below 44°C during normal operation. In the event that the treatment site underneath the applicator reaches the temperature threshold, the applicator will pause ultrasound output and vibrate once with a red LED notification that the device has begun a cooling or rest cycle. sam will automatically resume treatment with a blue LED notification after the site has cooled. Caution: At no time during treatment should the applicator be covered by thick insulating material such as a coat, blanket or sports wrap. This may cause sam to disable and remain disabled throughout the therapy session. |

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x sam

sam Manufactured by ZetrOZ, Inc. 56 Quarry Road Trumbull, CT 06611 USA

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