Stryker Air I Stryker Air II

Low Air Loss Therapy Mattress (LAL)

Model 2236

Operations/Maintenance Manual







For Parts or Technical Assistance: USA: 1-800-327-0770 (option 2)

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Symbols and Definitions



Warning / Caution - Consult accompanying documentation



Maximum Safe Working Load



Medical Equipment Classified by Underwriters Laboratories Inc. with respect to Electric Shock, Fire, Mechanical and Other Specified Hazards Only in Accordance with UL 60601-1, First Edition (2003) and CAN/CSA C22.2 No. 601.1-M90 with updates 1 and 2.

WARNING / CAUTION / NOTE DEFINITION

The words WARNING, CAUTION, and NOTE carry special meanings and should be carefully reviewed.



WARNING

Alerts the reader about a situation, which if not avoided, could result in death or serious injury. It may also describe potential serous adverse reactions and safety hazards.



CAUTION

Alerts the reader of a potentially hazardous situation, which if not avoided, may result in minor or moderate injury to the user or patient or damage to the equipment or other property. This includes special care necessary for the safe and effective use of the device and the care necessary to avoid damage to a device that may occur as a result of use or misuse.

Note

This provides special information to make maintenance easier or important instructions clearer.

INTENDED USE

This manual is designed to assist you with the safe operation and maintenance of the Stryker Air I and Stryker Air II, Low Air Loss (LAL) mattress system. Carefully read this manual thoroughly before using the equipment or beginning maintenance on it. To ensure safe operation of this equipment, it is recommended that methods and procedures be established for educating and training staff on the safe operation of this Arise® mattress system.

This high quality medical product, when properly installed and cared for will provide many years of trouble free service, and outstanding clinical performance.

PRODUCT DESCRIPTION

The Stryker Air I is an LAL mattress replacement system and the Stryker Air II is an LAL mattress overlay system. Both are a unique therapy system that provides pressure relief by combining low air loss with pulsation. Low air loss therapy has been demonstrated to reduce the risk of pressure ulcers as well as being a valuable aid in the treatment of pressure ulcers.

The Stryker Air I and II mattress system is suitable for both the treatment of existing pressure ulcers stage I through stage IV as well as those who have been assessed at risk from the complications of immobility. The Stryker Air I and II are quiet, comfortable and simple enough for single caregiver installation, featuring rapid inflation in just three to five minutes. The user-friendly controls allow for easy adjustment of patient comfort. The Stryker Air I and II mattress system offers special features to increase peace of mind such as a power interruption alarm and a patient position sensor which optimizes support for seated patients (Fowler Boost).

Additionally, low friction and shear materials, together with average interface pressures below capillary closure levels (32mm of Mercury), means that the Stryker Air I and II mattress system meets the comfort and clinical requirements.

SPECIFICATIONS

Model	Stryker Air I	2236-000-002	LAL Overlay
Wodel	Stryker Air II	2236-000-003	LAL Mattress
D	Width	35"	89 cm
Dimensions (Mattress Replacement)	Length	80", 84"	203 cm, 213 cm
(Mattress Heplacement)	Thickness	8.5"	22 cm
D	Width	35"	89 cm
Dimensions (Mattress Overlay)	Length	80", 84"	203 cm, 213 cm
(Matticss Overlay)	Thickness	5"	13 cm
	Width	13.5"	34 cm
Dimensions (Control Unit)	Height	11"	28 cm
	Depth	7.5"	19 cm
Weight	Mattress	24.5 lbs	10 kg
weight	Control Unit	10 lbs	4.5 kg
Safe Working Load (Max Patient Weight)		500 lbs	227 kg
Electrical Specifications	USA	120VAC, 60Hz, 4A, less than 1A running	
Classification	UL Classified Medical Equipment, UL 60601-1, Can/CSA C22.2 No. 601.1 Note: UL Classification refers to the power unit only.		No. 601.1
		the complete mattress replace	

ENVIRONMENTAL CONDITIONS

Environmental Conditions	Operation	Storage and Transportation
Ambient Temperature	10 °C (104 °F) (50 °F)	-10 °C (104 °F)
Relative Humidity (Non-Condensing)	30%	100%

Stryker reserves the right to change specifications without notice.

Specifications listed are approximate and may vary slightly from unit to unit or by power supply fluctuations.

THERAPY MATTRESS SYSTEM ILLUSTRATION



THERAPY CONTROL UNIT ILLUSTRATION



PRODUCT FEATURES

The Stryker Air I and II are comprised of two components:

- 1. Therapy Control Unit
- 2. Therapy Mattress System

Therapy Control Unit Features

- True low air loss with up to 100 liters of airflow per minute.
- Three modes of operation Autofirm, Therapy and Pulsate.
- Pulsating feature that oscillates the air throughout the mattress every 30 seconds.
- Autofirm mode provides maximum air inflation designed to assist both patients and caregivers during patient transfer and treatment.
- Patient position sensor optimizes support for seated occupants (Fowler Boost).
- Compact lightweight control unit is quiet, robust and powerful, with a reusable air filter and integrated carrying handle for portability.
- · Crisp, easy to read graphics for intuitive set up and therapy control.
- · Automatic panel lock out to avoid unwanted or accidental adjustments.
- · Eight therapeutic comfort control settings to maximize patient compliance and promote healing.
- Closed loop pressure sensor control system eliminates concerns of changes in mattress interface pressure due to ambient temperature and pressure changes.
- · Integrated swing out hanging brackets for fixing to most bed frames.
- · Quick disconnect hose feature allows for rapid attach and CPR deflate at the control unit.
- · Audible and visual alarms for power interruptions.

Therapy Mattress System (LAL Mattress and Overalay)

- Twenty (80") or twenty one (84") individual therapy cells help to evenly distribute the patient's weight and maximize pressure relief. Modular cell design mattress for ease of cleaning, re-assembly and cost effective service.
- · Lower two-inch support cell provides support in the event of a power failure.
- 6.5" deep mattress therapy cells and 5" deep overlay therapy cells are constructed of highly durable, polyurethane coated nylon to provide adequate support and prevent bottoming out for most patients within weight limit.
- · Integrated low-pressure heel section provides lower interface pressures.
- Durable base tub is constructed from 100% heavy weight 1680 denier nylon with a 1.5 oz. urethane coating and incorporates bed attachment loops for stability.
- Maximum weight capacity: Up to 500 lbs. based on model.
- CPR deflation in 30 seconds or less.
- · Anti-kink, easy clean air supply hose set.
- Integrated CairRails risk management side bolsters—two inch side bolsters that inflate on both sides of the patient
 along the mattress edge to provide additional support and to provide a gentle reminder to the patient that they are
 near the edge of the mattress. Constructed with lower ingress/egress area.

INDICATIONS FOR USE

Note: The selection of a pressure-relieving surface needs to be based on each individual patient's clinical condition, diagnosis and/or co-morbidities. The choice and use of a support surface is one factor in a holistic program of wound care and treatment.

Pressure Relief

Pressure Relief	Rehabilitation	Dermatology
 Neueology 	Burns	Amputations

Pain Management

• AIDS	Arthritis	Oncology

The Stryker Alr I and II mattress system provides uniform distribution of weight over a wide surface area, which relieves pressure against bony prominences and provides a soft, gentle therapy surface to lie on. For patients experiencing severe pain and discomfort due to pressure and/or positioning limitations, consider the Arise as an adjunct to pain management interventions.

Note: Pressure relief and pain management are conditions and diagnoses for which the Stryker Alr I and II mattress system may be indicated. Occasionally, there are orthopedic and neurological patients that require body positioning to be maintained in specific alignment. The Stryker Alr I and II mattress system has safety features to prevent deflation of the therapy cells and to keep patients from bottoming out at all times. The use of the Stryker Alr I and II mattress system for these patients should be considered on an individual basis and discussed with the attending physician.

THERAPY PADS

Many healthcare facilities are facing the challenge of infection control. This products quilted therapy pads are treated with an antimicrobial to protect the therapy pad from the growth of mold, mildew and bacteria.

Key features and benefits:

- · Treated with a highly effective bacteriostat agent to inhibit the growth of bacterial and fungus
- Constructed from a very smooth nylon fabric with low friction and low shear properties to protect the patient's skin from damaging friction/shearing forces.
- Breathable, moisture vapor permeable fabric allows air to circulate beneath the pad and wicks away excess moisture. This keeps your patient dry and helps to prevent skin maceration.
- Two-way stretch therapy pad is designed for optimal comfort, moisture vapor transfer, stain resistance and ease of laundering.

KEY AREAS IN THE TREATMENT OF COMPROMISED SKIN

Pressure Relief

The Arise® is divided into three distinct anatomical zones: head, seat, and foot. Each zone is adjusted to ensure optimal pressure relief and provide a comfortable sleeping surface.

Maximum pressure relief is achieved through delivering a specific amount of air to each therapy cell and allowing controlled amounts of air to escape, thus equalizing the pressure between the patient and the therapy cells. This distributes the patient's weight evenly over a wide surface area resulting in average pressure readings below capillary closure levels.

Moisture Control

Patients are at risk for skin maceration if excess moisture is permitted to accumulate beneath the patient. This may be due to perspiration, incontinence or wound drainage.

On the Stryker Alr I and II mattress system, moisture is controlled via the specially treated breathable, fluid-proof, urethane coated nylon therapy pad. The moisture vapor permeable fabric of the therapy pad allows a sufficient amount of air to circulate beneath the pad and wicks away excess moisture.

Shear and Friction Reduction

Shearing occurs when the skin is stationary in relation to the support surface, while the underlying tissues and vessels are stretched and damaged. When a patient's skin rubs against another surface, the result is friction. The top surface of the Stryker Alr I and II mattress system therapy pad is constructed from a very smooth nylon fabric with low friction and shear properties to protect the patient's skin from these damaging forces.

Summary of Safety Precautions

Important Notice: The equipment has to be installed and operated in the manner for which it was intended. Facility staff/user is responsible for reading and understanding the product user manual prior to performing any operations or maintenance on this product.

A

WARNING

- The safe use of the equipment is maximized when used in conjunction with bed rails; there may be an increased risk of falls when such bed rails are not present. Serious injury or death can result from the use (potential entrapment) or non-use (potential patient falls) of side rails or other restraints. Local policies regarding the use of side rails should be taken into account. Whether and how to use side rails is a decision that should be based on each patient's individual needs and should be made by the physician, caregivers and responsible parties.
- When using the Stryker Alr I and II mattress system, always ensure that the patient is positioned properly within the
 confines of the bed. Do not let any extremities protrude over the side or between the bed rails when the mattress
 is being used.
- Do not use in the presence of flammable anesthetics. Do not use in the presence of smoking materials or open flame. Air flowing through the air mattress will support combustion.
- To reduce the risk of shock, adhere to the following instructions. Failure to do so could result in personal injury or equipment damage.
 - Immediately after using the Arise®, unplug it from its power source.
 - Do not place or store the product where it can fall or be pulled into a tub or sink.
 - Do not place or drop the product into water or other liquid.
 - · Do not remove the back of the control unit.
- To reduce the risk of burns, shock, fire, or personal injury, adhere to the following instructions. Failure to do so could result in personal injury or equipment damage.
 - Use this product only for its intended purpose as described in this manual. Only use attachments and/or accessories that are recommended by the manufacturer.
 - If this product has a damaged power cord or plug, is not working properly, has been dropped or damaged, or has been dropped into water, do not operate it. For examination and repair, return the product to the manufacturer.
 - · Keep the control unit and power cord away from heated surfaces, e.g. space heaters.
 - Never block the air openings of the product. Do not place the control unit on a surface, such as a bed or couch, where the air opening and/or filter compartment, located on the back of the control unit, may be blocked. Keep the air openings free of lint and hair.
 - Never drop or insert any object into any opening or hose.
 - Do not spill food or liquids onto the control unit. If a spillage does occur, turn off the unit, disconnect it from its
 power supply and allow at least 24 hours for drying.
 - · Do not use the product outdoors, or where aerosol-spray products are used.
 - Plug this product only into a properly grounded outlet. Refer to Grounding Instructions on page 13.
 - Be sure nothing is placed on the power cord and ensure it is not located where it can be stepped on or tripped over.
 - Do not attempt to service the control unit. Please call Stryker at 1-800-327-0770 for any service requests.
 - The therapy pad (top cover) of this product is not air permeable and may present a suffocation risk. It is the responsibility of the caregiver to ensure that the patient can use this product safely.
- The risk of entrapment can develop when equipment is placed on bed frames that leave gaps of even a few inches between the mattress and the headboard, footboard, and bed or side rails. The equipment is NOT to be used when such gaps are present.



CAUTION

• CairRails (side air bolsters), when inflated, offer additional support and provide a gentle reminder to the patient that they are nearing the side edge of the mattress.

Summary of Safety Precautions

BED SYSTEM ENTRAPMENT INFORMATION

Although common in the practice of long-term care, bedside rails, in recent years, have also been a subject of regulatory review and evolution in design and use.

That focus includes not only the challenge of achieving an appropriate balance between resident security and unnecessary restraint, but also the additional safety issue of entrapment.

The U.S. Food and Drug Administration (FDA), has addressed the potential danger of entrapment with new safety guidelines for medical beds. These guidelines recommend dimensional limits for critical gaps and spaces between bed system components.

Entrapment zones involve the relationship of bed components often directly assembled by the healthcare facility rather than the manufacturer. Therefore, compliance is the responsibility of the facility.

GROUNDING INSTRUCTIONS



WARNING

Use a properly grounded, three-prong, 120V AC outlet for this product. Failure to use a grounded outlet could result in personal injury or damage to equipment or house wiring, including risk of fire. A qualified electrician should be contacted to correct the wiring and ensure a properly grounded outlet.

Note

- Before installing this product, have the electrical system checked to make sure the electrical circuits and the electrical service are properly grounded.
- Having a three-prong outlet does not necessarily mean it is grounded. Sometimes two-prong outlets are replaced with a three-prong type even though there is no ground wire.
- There is always a chance of a loose connection or poor installation of a ground wire that causes the loss of proper ground at the outlet. Inadequate grounding at electrical outlets can occur even if there is a ground wire. Wires can become loose over time at the connection to the outlet.



CAUTION

A qualified electrician is required for the installation of new wires on a circuit.

How to Determine if Your Outlet has the Proper Grounding

Most hardware stores sell circuit testers (see Figure 2) that can be used to test an outlet for proper grounding. The tester plugs into an outlet and by observing the indicator lights you can determine if the outlet is properly grounded. For a higher level of assurance, an electrician should be requested to thoroughly test the electrical system with more reliable equipment.

If repair or replacement of the cord or plug is necessary, please contact Stryker at 1-800-327-0770.



Figure 2

Setup Procedures



WARNING

Ensure you have read and understand all Safety Precautions listed on page 12 -13 prior to performing the setup procedures below.



CAUTION:

Do not place the control unit on the floor. Position the power cord to keep personnel from tripping over it. When the product is not in use, properly store the power cord. Failure to do so could result in personal injury.

Stryker Air I (LAL Mattress)

- 1. Remove the existing mattress from the bed.
- 2. Place the Arise LAL mattress with the hose connection at the foot end of the bed and the therapy cells facing up. Secure the six (6) straps on the mattress securely to the movable part of the bed frame.
- 3. If the therapy pad is not already on the mattress, attach it securely to the mattress.
- 4. Hang the control unit on the foot of the bed facing away from the bed (see Figure 3). Attach the hose connector marked CPR to the underside of the control unit (see Figure 4).



Figure 3







Figure 4

- 5. Plug the Fowler Boost sensor into the side of the control unit.
- 6. Plug in the control unit and the yellow Standby light will illuminate. Press the Power key. The control unit will start and the green light will illuminate.
- 7. Allow three to five minutes for full inflation. Place the patient on the mattress. Mattress can be inflated with patient on it, but will take longer, depending on patient weight/size. Note: Keep the control unit on while the patient is on the mattress.
- Position the patient's head in the position in which the patient will be for the largest portion of the day. If the patient is lying flat, please use three (3) fingers for the hand check. If the patient will be sitting up for the majority of the day, please use two (2) fingers.
- 9. Perform a hand check by placing fingers locally under the patient's buttocks between two cushions. The patient should not bottom out. If they do, increase the therapy control by one comfort level, until they no longer bottom
- 10. The CairRails air side bolsters can be inflated or deflated as required. Locate the turn valve on the hose assembly between the mattress and the control unit. Next, inflate/deflate the CairRails by moving the turn valve to the up (inflate) or down (deflate) position. Note: When inflating CairRails, it is recommended that the control unit be in Autofirm mode to achieve optimal results.

Setup Procedures



CAUTION:

Do not place the control unit on the floor. Position the power cord to keep personnel from tripping over it. When the product is not in use, properly store the power cord. Failure to do so could result in personal injury.

Stryker Air I (LAL Overlay)

- Place the Stryker Air II LAL overlay on top of the mattress with the hose connection at the foot end of the bed and the therapy cells facing up. Secure the straps around the head and foot ends of the mattress. Fasten the two straps by the seat section securely to the sleep deck of the bed frame.
- 2. If the therapy pad is not already on the mattress, attach it securely using the zippers.
- 3. Hang the control unit on the foot of the bed facing away from the bed. Attach the hose connector marked CPR to the underside of the control unit (see Figure 5).
- 4. Plug the Fowler boost sensor into the side of the control unit.



Figure 5

- 5. Plug in the control unit and the yellow Standby light will illuminate. Press the Power key. The control unit will start and the green light will illuminate.
- 6. Allow three to five minutes for full inflation. Place the patient on the overlay. Overlay can be inflated with patient on it, but will take longer, depending on patient weight/size. Note: Keep the control unit on while the patient is on the overlay.
- 7. Perform a hand check by placing fingers locally under the patient's buttocks between two cushions. The patient should not bottom out. If they do, increase the therapy control by one comfort level, until they no longer bottom out.
- 8. The CairRails air side bolsters are currently not available with the Stryker Air II LAL overlay.

Operation Guide



CAUTION

The patient's head should be positioned in the center of the top section of the mattress. When using the mattress system always ensure that the patient is positioned properly within the confines of the bed. Do not let any extremities protrude over the side or between the bed rails when the mattress is being used.

PATIENT COMFORT CONTROLS AND MONITORING

(Refer to Figure 5)

Power Switch

The power switch is used to turn the power on and off.

Standby Light

The *Standby* indicator will illuminate when the unit is plugged into a power outlet. Press the **Power** button to turn the unit on and inflate the mattress. Therapy is the default mode at startup. When the *Standby* light is on, it may also indicate that there has been a power interruption and the therapy control unit is ready to be turned back on. Press the **Power** button and reset the preferred mode of therapy and comfort level.

Modes

The Arise® has two low air loss therapy modes, Therapy and Pulsate and an Autofirm mode

- Therapy: The unit starts in the Therapy mode, which is the standard low air loss therapy.
- Pulsate: Pulsate will slightly decrease the pressure in all cushions every 30 seconds then return to the programmed comfort adjust level.
- 3. Autofirm: Autofirm mode provides maximum air inflation designed to assist both patients and caregivers during patient transfer and treatment. The unit will automatically return to the mode it was in prior to Autofirm (either therapy or pulsate) in approximately 12 minutes.

Power Autofirm Therapy Pulsate Position Sensor Alarm Reset Low Air Loss Pressure Relief System

Figure 5

Comfort Adjust

The Comfort Adjust function is located on the right side of the control panel. The Stryker Air I and II can be customized to meet individual patient needs within a therapeutic window. Use the up and down keys to simultaneously increase or decrease pressure in all three zones (head, seat and foot). This function will not work in Autofirm mode.

Lockout

This feature is to prevent any unauthorized changes to the patient settings. To unlock and make adjustments to the settings press both **Up** and **Down** comfort arrows at the same time to disengage the Lockout function. The Lockout function will return in approximately five minutes.

Note: The unit is designed to lock out all the adjustment controls after the patient has been positioned correctly. In approximately five minutes after the last button push, the power on light begins to flash indicating Lockout is enabled.

Operation Guide

CPR

The hose connection at the control unit is marked CPR. Disconnect the hose from the control unit. CPR connection style may vary by model. To re-inflate the air cells, reattach the hose to the control unit.

TRANSPORT

To transport the patient in bed, turn the control unit off. Unplug the power cord from the outlet. Do not disconnect the hose connection at the control unit. The lower support cell will prevent the patient from bottoming out for up to 12 hours. Some models incorporate a foam safety cell.

POWER FAILURE

After a power failure the unit will start up in *Standby* mode. Reset the preferred mode and comfort level as described on page 16. In the event of a power failure, the lower support cells will hold air for up to 12 hours. Some models incorporate a foam safety cell. In case of an extended power failure, transfer the patient to a hospital mattress or other surface.

CAIRRAILS

Integrated CairRails risk management air bolsters offer a bilateral side bolster solution designed to address healthcare's growing concerns of liability in relation to patient falls and entrapment. CairRails are being recognized by some of the nations leading healthcare systems for improving their patient safety and risk management programs.

CairRails are recommended for patients requiring additional support during patient care and transfer. CairRails can help reduce costs while ensuring optimal clinical outcomes and increasing patient safety.

Note: When inflating CairRails, it is recommended that the control unit be in Autofirm mode to achieve optimal results.

Features and Benefits

- · A bilateral side air bolster solution which can enhance your facilities entrapment/risk management program.
- · Easy to engage Ready Valve for instant inflation and deflation.
- · Transfer friendly-deflate for ease of assisted transfer or when bolsters are not required.
- Unique contoured design allows ease of ingress/egress, while providing additional protection, comfort and supports patient compliance.
- Designed to fit on most therapeutic support surfaces.
- · Promotes maximum independence by allowing caregiver to decide when added protection is required.

Note: Side bolsters are meant to be used with side rails and to provide a documentable and functional intervention for the risk management issues of falls and entrapment but in no way guarantees the prevention of falls or entrapment occurrences.

Nursing Procedures

RECOMMENDED LINEN

Special linens are not necessary for the Stryker Air I and II. While there is no need for a bottom sheet the therapy pad should be covering the therapy cells at all times. The patient should never be lying directly on the therapy cells. Depending upon the patient's specific needs, the following linens may be utilized:

- Draw or slide sheet to aid in positioning and to further minimize friction and shearing.
- · Incontinence barrier pad for patients incontinent of urine and/or stool, and patients with heavily draining wounds.
- · Add top sheet, blanket and/or bedspread as needed for patient comfort.
- · Keep the amount of padding between the patient and bed to a minimum for optimum performance.

CHANGING THE THERAPY PAD

- 1. Place the therapy pad over the therapy cells, fitting the corner of the cushions into the corner of the therapy pad. (Similar to a fitted sheet)
- 2. Zip the therapy pad along each side of the mattress tub.

PATIENT POSITIONING AND COMFORT

<u>General Repositioning:</u> Patients should be turned and repositioned per individual turning schedule or per facility policy. It may be helpful to activate the Autofirm mode to achieve a firm surface for repositioning purposes. The unit will automatically return to the mode it was in prior to Autofirm in approximately 12 minutes or you can manually return to therapy mode once patient has been repositioned. Unless counter indicated, it is desirable to keep the head of the bed in the low position to provide optimal pressure relief and minimize the risk of shearing injuries.

Elevating Patient into Sitting Position: The special properties of the Stryker Air I and II therapy pad reduce the opportunity for shear and friction that may occur when raising the head of other beds. As with any surface, sliding can be expected, therefore patients should be repositioned after elevation. The knee gatch or foot of the bed may be elevated first, to help prevent the patient from sliding when the head of the bed is elevated.

INCONTINENCE

Moisture against the skin surface leads to maceration, or softening of the tissues. To prevent maceration, we recommend you use an incontinence barrier pad to absorb the excess moisture. In the event of incontinence or excess drainage on the therapy pad, you should wipe off the excess fluid from the bed surface.

SAFETY INFORMATION

Patient Migration

Specialty bed products are designed to reduce/relieve pressure and the shearing/friction forces on the patient's skin. The risk of gradual movement and/or sinking into hazardous positions of entrapment and/or inadvertent bed exit may be increased due to the nature of these products.

Traction

With any traction or unstable fractures, maintain physician-directed angle of articulation and guard against risks of patient migration or inadvertent deflation of patient surface.

Skin Care

Monitor skin conditions regularly, particularly in areas where incontinence and drainage occur or collect, and consider adjunct or alternative therapies for high acuity patients. Early intervention may be essential to preventing serious skin breakdown.

Bed Height

To minimize the risks of falls or injury the patient surface should always be in the lowest practical position when the patient is unattended. Make sure areas under and around the frame are clear of objects, persons and parts of body before adjusting height.

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Cleaning, Storage and Care

\wedge

WARNING

- Unplug the control unit from its power source. Failure to do so could result in personal injury or equipment damage.
- Do not expose the unit to excessive moisture that would allow for liquid pooling. Personal injury or equipment damage could occur.



CAUTION

- Do not use harsh cleansers/detergents, such as scouring pads and heavy-duty grease removers, or solvents, such as acetone. Equipment damage could occur.
- · Do not use any steam cleaning device on the unit.

CONTROL UNIT

Wipe off dust. If necessary, clean the housing exterior with a disinfectant solution or a mild detergent and a damp cloth. Then wipe dry.

GENERAL CLEANING

If there is no visible soilage with possible body fluids, we recommend that you clean the mattress system with a mild detergent and warm water. If disinfection is desired, you may use a combination cleanser/disinfectant as explained in "Disinfecting" section.

- Patient care equipment that does not come in contact with mucous membranes or non-contact skin requires low-level disinfection. Wiping surfaces with a properly prepared detergent or disinfectant carries out low level disinfecting.
- Processing of dirty patient care equipment should take place in a designated area away from clean or sterile supplies and food preparation areas.
- Detergent/disinfectants should not be mixed with other germicides or detergents. Using the proper dilution insures the most effective killing power of the disinfectant.
- · Wash hands often and well, including after removal of gloves.
- Patient care equipment that is used in isolation areas should be disinfected in accordance with all internal policies and procedures regarding such equipment.

DISINFECTING

When there is visible soilage, and between patients, it is recommend that the unit and mattress be disinfected with a tuberculocidal disinfectant. Disinfectant should be registered with the Environmental Protection Agency (EPA).

- Use rubber gloves and eye protection.
- Prepare detergent/disinfectant (registered by EPA as hospital disinfectant) solution according to instructions on label for correct use-dilution.
- With support surface deflated, thoroughly wipe down entire mattress, as air cells will lie flat. Be sure to reach all areas underneath and in-between air cells. Allow to air dry.
- If dust or other soiling has accumulated along air hoses, remove using swabs moistened with detergent/disinfectant as necessary. Allow all components to air dry. Wrap mattress in plastic and return to storage area.
- · Thoroughly wipe down outside of control unit and allow to air dry. Cover with plastic and return to storage area
- Remove gloves and dispose; wash hands.

Cleaning, Storage and Care

THERAPY PAD

The therapy pad can be wiped down with a disinfectant solution or a mild detergent with a damp cloth. If heavily soiled, the therapy pad can be laundered in a washer and dryer with warm water (no more than 120° Fahrenheit). A non-bleach detergent should be used sparingly. Wipe dry or allow to air dry.

STEAM CLEANING

Do not use any steam cleaning device on the unit. Excessive moisture can damage mechanisms in this unit.

FILTER CLEANING

Check the air filter on the rear of the unit regularly for buildup of dust/dirt. If buildup is visible turn off the control unit and disconnect the power cord from the wall outlet. Remove the filter by grasping the filter pulling outward. Replace with the second supplied filter. Ensure the replaced filter covers the entire filter region.

Hand-wash the removed filter in warm soapy water and allow to air dry. When dry, store the filter in a safe place for the next filter maintenance.

STORAGE

<u>Control Unit:</u> The power cord may be wrapped around the unit for convenience. Wrap the unit in a plastic bag for dust resistance then store the unit in an area appropriate for an electronic medical device.

<u>Support Surface:</u> Gently roll up the support surface, expelling any residual air, for temporary storage. The mattress should be wrapped in plastic and/or a clean bag for storage.

Preventative Maintenance

MAINTENANCE CHECKLIST All fasteners secure (reference all assembly drawings). Hose assembly is not damaged or leaking. Power switch is working properly. Power cord is not frayed and is attached to blower assembly. All electrical connections function properly. Control Unit buttons function properly. Plastic on the control unit assembly is not damaged. Mattress holds air and all straps are intact. Inspect all product labeling for signs of degradation. Unit Serial Number:

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Date: _____

Completed by:

Troubleshooting

Problem / Failure	Recommended Action	
Therapy surface is not inflating.	1.	Ensure the hose connection from the therapy mattress system (mattress) to
		the control unit is securely connected.
	2.	Ensure that the control unit is plugged into an A/C outlet.
		Ensure that the power is not on <i>Standby</i> . If on <i>Standby</i> , press the Power
		button.
Unable to change therapy mode	1.	Make sure the <i>Lockout</i> function is disabled. To disable, press
or adjust comfort control.		the Up and Down comfort adjust arrows simultaneously.

Note

The parts and accessories listed on this page are all currently available for purchase. Please call Stryker Customer Service (800)-327-0770.

Stryker Air I Power Unit: 2236-000-005			
Part Name	Part View	SYK Part Number	
Blower assembly		2236-100-020	
Bumper Feet (Pkg 10 Sets)		2236-100-019	
Fuse, External, 5A (Pkg 10)		2236-100-035	
Fuse, Internal, 0.5A (Pkg 10)	internal Fuse o	2236-100-036	
Filter Foam, pkg (10)		2236-100-017	
Kit, E-Ring (pkg 100) for 3G Bed Hooks	E-ring	2236-100-022	
Kit, Bed Hook Assembly		2236-100-014	

Stryker Air I Power Unit: 2236-000-005			
Part Name	Part View	SYK Part Number	
Kit, Power Cord Kit		2236-100-018	
Mattress Hose Connector		2236-100-021	

Stryker Air I Mattress: 2236-000-003			
Part Name	Part View	SYK Part Number	
Air Cell, Stryker Air I Foot		2236-100-010	
Air Cell, Main Support (5x36")		2236-100-008	
Back Raise Sensor Assembly		2236-100-032	
External Hose Set Assy, Stryker Air I		2236-100-037	
External Hose Cover		2236-100-039	

Stryker Air I Mattress: 2236-000-003			
Part Name	Part View	SYK Part Number	
Hinge Set	TOTAL STATE OF THE PARTY OF THE	2236-100-038	
Hose Clamp (Pkg 25)	6	2236-100-034	
Loop Strap, Stryker Air I		2236-100-043	
Mattress Tub, Stryker Air I		2236-100-040	
Therapy Pad, Stryker Air I (Overlay)		2236-100-006	

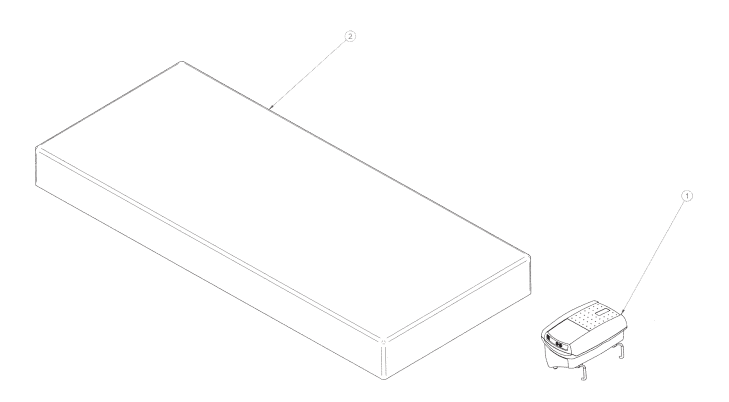
Stryker Air II Power Unit: 2236-000-005			
Part Name	Part View	SYK Part Number	
Blower assembly		2236-100-046	
Bumper Feet (Pkg 10 Sets)		2236-100-019	

Stryker Air II Power Unit: 2236-000-005			
Part Name	Part View	SYK Part Number	
Fuse, External, 5A (Pkg 10)		2236-100-035	
Fuse, Internal, 0.5A (Pkg 10)	Internal Fuser of	2236-100-018	
Filter Foam, pkg (10)		2236-100-044	
Kit, E-Ring (pkg 100) for 3G Bed Hooks	E-ring	2236-100-047	
Kit, Bed Hook Assembly		2236-100-014	
Kit, Power Cord Kit		2236-100-045	
Mattress Hose Connector		2236-100-021	

Stryker Air II Mattress: 2236-000-004			
Part Name	Part View	SYK Part Number	
Air Cell, Cair Rail		2236-100-053	
Air Cell, LAL (6"x36")		2236-100-052	
Back Raise Sensor Assembly		2236-100-032	
External Hose Set Assy, Stryker Air I		2236-100-048	
External Hose Cover, LAL w/Cair Rail		2236-100-049	
Hinge Set		2236-100-038	
Hose Clamp (Pkg 25)		2236-100-034	
Loop Strap, Stryker Air II		2236-100-011	

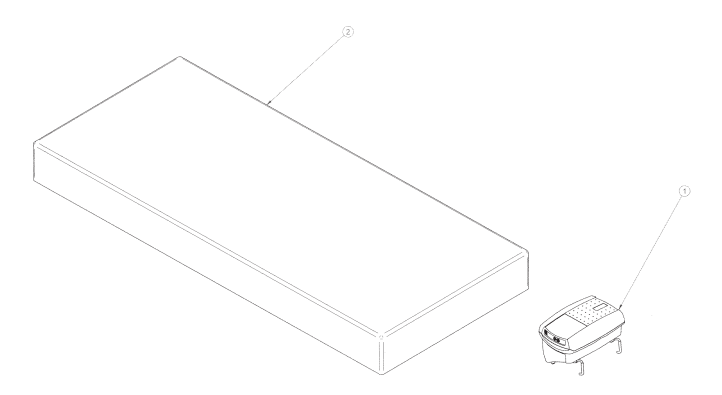
Stryker Air II Mattress: 2236-000-004			
Part Name	Part View	SYK Part Number	
Mattress Tub, Stryker Air II		2236-100-050	
Therapy Pad, Stryker Air II		2236-100-051	

For Reference Only: 55-0060 Rev-00



Item	Part No.	Part Name	Qty.	
1	2236-100-005	Stryker Air I Power Unit	1	
2	2236-100-003	Stryker Air I Mattress	1	

For Reference Only: 55-0061 Rev-00



Item	Part No.	Part Name	Qty.
1	2236-100-005	Stryker Air II Power Unit	1
2	2236-100-004	Stryker Air II Mattress	1

Warranty

LIMITED WARRANTY

The Stryker Air I & II mattress system is guaranteed for a period of one year from the date of delivery, against defects in materials and workmanship, under normal use and service. This one-year warranty includes all mechanical and electrical components. Steel structural components on beds are covered under warranty for a period of 10 years from the date of delivery. Welds are covered under warranty for the lifetime of the product.

Damage caused by use in unsuitable environmental conditions, abuse or failure to maintain the product in accordance with user and service instructions is not covered. Any alteration, modification, or repair unless performed by or authorized in writing by Stryker Medical, will void this warranty. Damages arising from improper use will not be covered under this warranty.

Improper use is defined as, but not limited to, those caused by

- Burns
- Use of improper chemical agents
- · Needle punctures, cuts or abrasions
- Excessing loads
- Staining
- Negligent or excessive usage
- · Improper maintenance, handling and/or cleaning
- Failure to use in the manner indicated in this user manual

PARTS

The Stryker Air I & II mattress system contain various parts that wear from normal use. Stryker Medical's obligation under this warranty is limited to supplying replacement parts, servicing or replacing, at its option, any product which is found by Stryker Medical to be defective.

Warranty replacement parts are covered by the terms of this warranty until the product's original one-year warranty period expires. Part replacements under warranty may be requested to be returned back to the manufacturer. When requested by Stryker Medical, parts must be returned for inspection at the customer's expense. Credit will be issued only after inspection.

SERVICE

Most service requests can be handled by the facility Maintenance Department with assistance from the Stryker Medical Technical Support Service Department. Most parts requested can be shipped next day air at the customer's expense. Should a technician be required, one will be provided by Stryker Medical, at our discretion. Only the Stryker Medical Technical Support Service Department can dispatch authorized technicians. This warranty is extended to the original purchaser of the equipment.

For further service needs please call Stryker at 1-800-327-0770.

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