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INTRODUCTION

This manual is designed to assist you with the operation of the 1231 Emergency Care Stretcher, and the 1731 PACU Bed. Read it thoroughly before using the equipment or beginning any maintenance on it.

SPECIFICATIONS

<table>
<thead>
<tr>
<th></th>
<th>1231</th>
<th>1731</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall Bed Length \ Width</td>
<td>83”/30”</td>
<td>83”/30.5”</td>
</tr>
<tr>
<td>Minimum \ Maximum Bed Height</td>
<td>22”/35.5”</td>
<td></td>
</tr>
<tr>
<td>Maximum Weight Capacity</td>
<td>500 pounds</td>
<td></td>
</tr>
<tr>
<td>Fowler Angle</td>
<td>0 to 90°</td>
<td></td>
</tr>
<tr>
<td>Knee Gatch Angle</td>
<td>0 to 30°</td>
<td></td>
</tr>
<tr>
<td>Trendelenburg \ Reverse Trendelenburg</td>
<td>+18° to –18°</td>
<td></td>
</tr>
</tbody>
</table>

Stryker reserves the right to change specifications without notice.

WARNING / CAUTION / NOTE DEFINITION

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

WARNING

The personal safety of the patient or user may be involved. Disregarding this information could result in injury to the patient or user.

CAUTION

These instructions point out special procedures or precautions that must be followed to avoid damaging the equipment.

NOTE

This provides special information to make maintenance easier or important instructions clearer.
A) Pump to raise litter.

B) Depress in the center of pedal (B) to lower both ends of the stretcher together.

C) Depress the side of pedal (B) closest to the foot end of the stretcher to lower the foot end.

D) Depress the side of pedal (B) closest to the head end of the stretcher to lower the head end.

E) Brake and Steer functions (foot end).

F) Brake and Steer functions (head end).
OPERATING BASE CONTROLS – DUAL LOWERING PEDALS

A Pump to raise litter.

B Depress to lower head end.

C Depress to lower foot end.

Note: Depress B & C together to lower both ends of the litter simultaneously.

D Brake and Steer functions (foot end).

E Brake and Steer functions (head end).
Stretcher Operation

RAISING AND LOWERING LITTER HEIGHT – UNI–LOWER PEDAL

CAUTION
Be sure to move any equipment that may be in the way before raising or lowering the litter height.

To raise the litter height, pump pedal (A) repeatedly until the desired height is achieved (see illustration on page 3).

To lower both ends of the litter together, depress the center of pedal (B). To lower only the head end of the litter, depress the side of pedal (B) closest to the head end. To lower only the foot end of the litter, depress the side of pedal (B) closest to the foot end (see illustration on page 3). The base may be equipped with optional variable descent controls. With variable descent controls, the farther you press down on the pedal, the faster the litter will lower.

RAISING AND LOWERING LITTER HEIGHT – DUAL LOWERING PEDALS

CAUTION
Be sure to move any equipment that may be in the way before raising or lowering the litter height.

To raise the litter height, pump pedal (A) repeatedly until the desired height is achieved (see illustration on page 4).

To lower both ends of the litter simultaneously, depress pedal (B) and (C) together using the same foot. To lower only the head end of the litter, depress pedal (B). To lower only the foot end of the stretcher, depress pedal (C) (see illustration on page 4). The base may be equipped with optional variable descent controls. With variable descent controls, the farther you press down on the pedal, the faster the litter will lower.

TRENDELENBURG/REVERSE TRENDELENBURG – UNI–LOWER PEDAL

NOTE
Litter height must be raised first in order to achieve a trend. or reverse trend. position.

CAUTION
Be sure to remove any equipment that may be in the way before lowering stretcher.

For Trendelenburg positioning (head down), depress the side of pedal (B) closest to the head end of the stretcher (see illustration, page 3).

For Reverse Trendelenburg positioning (foot down), depress the side of pedal (B) closest to the foot end.

NOTE
The higher the litter is before pedal (B) is activated, the greater the trend. or reverse trend. angle will be. (Maximum trend. angle is +18°. Maximum reverse trend. angle is −18°).

TRENDELENBURG/REVERSE TRENDELENBURG – DUAL LOWERING PEDALS

NOTE
Litter height must be raised first in order to achieve a trend. or reverse trend. position.

CAUTION
Be sure to remove any equipment that may be in the way before lowering stretcher.

For Trendelenburg positioning (head down), depress pedal (B) (see illustration, page 4).

For Reverse Trendelenburg positioning (foot down), depress pedal (C) (see illustration on page 4).
APPLYING THE BRAKE SYSTEM

NOTE
For user convenience, the brake/steer pedal is located on both ends of the stretcher.

WARNING
Always apply the caster brakes when a patient is getting on or off the stretcher. Push on the stretcher to ensure the brakes are securely locked. Always engage the brakes unless the stretcher is being moved. Injury could result if the stretcher moves while a patient is getting on or off the stretcher. If brakes do not hold properly, refer to your stretcher maintenance manual for a brake adjustment procedure.

To engage the brakes on the head (non–control) end, push fully down on the left side of pedal (E) (see illustration on page 4).

To engage the brakes on the foot (control) end, push fully down on the right side of pedal (D) (see illustration on page 4).

NOTE
Your stretcher may be equipped with optional side control brake and steer functions in addition to the standard head and foot end controls. The side control brakes operate the same as the head and foot end versions. They are indicated by a label on each side of the stretcher base.

OPERATING DIRECTIONAL STEERING CASTER/5TH WHEEL OPTIONS
The purpose of the steer wheel and 5th wheel options are to help guide the stretcher when transporting a patient along a straight line and also for pivoting at corners.

To engage the steer wheel/5th wheel, push the proper side of the brake/steer pedal to the full down position.

NOTE
Stretchers cannot be equipped with both steering caster and fifth wheel options. The choice was made at the time the stretcher was purchased.

The steer wheel is located at the foot end of the stretcher on the patient’s left. The 5th wheel is located underneath the center of the base assembly.
OPERATING THE BIG WHEEL OPTION

When the brake/steer pedal is in the neutral or brake position, the Big Wheel is elevated approximately 1” and the stretcher rests on the four casters.

NOTE
The two Big Wheels do not pivot. The stretcher cannot be moved directly sideways with the Big Wheel activated. With the pedal in the neutral position, the stretcher can be moved in any direction including sideways.

When the brake/steer pedal is in the steer position, the foot end casters are elevated approximately 1/4” and the stretcher rests on the two head end casters and the two Big Wheels. This provides increased mobility and ease of steering the stretcher.
USING PUSH HANDLES

CAUTION
The push handles should always be used when transporting patients. Avoid using other parts of the stretcher as pushing devices or damage may occur.

To use the push handles:
Pivot the handles up and push down until they are locked into position.

To store the push handles:
Lift the handles up and pivot them down to store in the handle rests.
USING GLIDEAWAY™ SIDERAIRS

NOTE
Raising and lowering siderails is a two–hand-ed operation. Use one hand to hold and position the siderail and the other hand to operate the siderail latch.

WARNING
When lowering the siderail to the collapsed position, keep extremities of patients and staff away from the siderail spindles or injury could occur.

To raise siderails: Pull up on the siderail (A) and raise it to full up position until the latch (B) engages.

To lower siderails: Pull up on the latch (B) and guide the siderail to the full down position.

NOTE
The latches (B) are colored red for easy identification.

WARNING
To avoid injury or damage to the equipment, Do not allow siderail to lower on its own.

NOTE
There is a dual siderail latch option available where there are latches on both ends of the stretcher.
OPERATING FOWLER AND KNEE GATCH

Operating Pneumatic Fowler

Squeeze handle (A) for pneumatic assist in lifting the Fowler to the desired height. Remove hands from handle when desired height is achieved.
To lower, squeeze handle (A) and push down until the Fowler has reached the desired height. Remove hands from handle when desired height is achieved.

NOTE
Handle (A) is colored red for easy identification.

CAUTION
If pneumatic system appears difficult to operate, refer to the stretcher maintenance manual for "Pneumatic Fowler Adjustment".

WARNING
Keep fingers/hands clear of the area around the Fowler release handle and the Fowler frame when lowering. Injury could result if care is not taken when lowering the Fowler.

Operating Optional Crank Knee Gatch

Remove the crank handle (A) from its storage magnet under the litter, pivot it up and push in to engage it.
Turn crank handle (A) clockwise to raise the Knee Gatch and counterclockwise to lower it.
USING THE SAFR™ PATIENT TRANSFER BOARD

WARNING
When using the SAFR™ Board to transfer a patient, always lock the brakes on all stretchers, beds, etc. being used and always be certain the transfer board is placed securely on the surface of the mating stretcher or bed. The patient stretcher and the mating surface must be at the same height before the patient is transferred. Failure to confirm the brakes are locked and the transfer board is securely placed could result in patient injury.

The SAFR™ patient transfer board stores flat under the litter. To use it, pull it straight out and pivot it up into place. Using a sheet, draw the patient onto the mating bed or stretcher.

The siderail can be raised and lowered with the SAFR™ Board in the up position but the board must be pivoted up so it is perpendicular to the litter.

![STRETCHER SIDE VIEW](image1)

![STRETCHER END VIEW](image2)
OPERATING OPTIONAL 2–STAGE PERMANENTLY ATTACHED I.V. POLE

NOTE
The 2–stage permanently attached I.V. pole is an option and may have been installed at either the head, foot or both ends of the stretcher. The choice was made at the time the stretcher was purchased.

To use the 2–stage permanently attached I.V. pole:
1. Lift and pivot the pole from the storage position and push down until it is locked into the receptacle.
2. To raise the height of the pole, pull up on the telescoping portion (A) until it locks into place at its fully raised position.
3. Rotate the I.V. hangers (B) to desired position and hang the I.V. bags.
4. To lower the I.V. pole, turn the latch (C) clockwise until section (A) lowers.

CAUTION
The weight of the I.V. bags should not exceed 40 pounds.
NOTE
The 3–stage permanently attached I.V. pole is an option and may have been installed at either the head, foot or both ends of the stretcher. The choice was made at the time the stretcher was purchased.

To use the 3–stage permanently attached I.V. pole:
1. Lift and pivot the pole from the storage position and push down until it is locked into the receptacle.
2. To raise the height of the pole, pull up on the telescoping portion (A) until it locks into place at its fully raised position.
3. For a higher I.V. pole, pull up on section (B). Release section (B) at any desired height and it will lock into place.
4. Rotate the I.V. hangers (C) to the desired position and hang the I.V. bags.
5. To lower the I.V. pole, push up on the red portion of grip (D) while holding onto section (B) until it lowers. Turn latch (E) clockwise until section (A) lowers.

CAUTION
The weight of the I.V. bags should not exceed 40 pounds.
OPERATING OPTIONAL TETHERED I.V. POLE

To use the tethered I.V. pole:

1. Remove the I.V. pole from the storage trough under the litter and insert into the receptacle on the corner of the litter frame.

2. To raise the height of the pole, turn knob (A) counter-clockwise and pull up on the telescoping portion (B) of the pole to raise it to the desired height.

3. Turn knob (A) clockwise to lock the telescoping portion in place.
OPERATING OPTIONAL HEEL STIRRUPS

1. To use the optional heel stirrups, turn the handle (A) on the lock screw located under the litter frame and swing the stirrup assembly into position. Tighten the handle (A) to hold the assembly in place.

2. Loosen knob (B) and pull out the extension tube (C) to the desired length. Tighten knob (B).

3. Loosen knob (D) and raise or lower the stirrup (E) to the desired height. Tighten knob (D).

OPERATING OPTIONAL FOOT EXTENSION/DEFIBRILLATOR TRAY

To use as a defibrillator tray, pull out top knob (A) and pivot tray (B) over the foot extension until tray extends flat over foot end of stretcher.

To use as a foot extension, pull out knob (A) and pivot the defibrillator tray back until it locks against the foot extension (C). While holding onto the assembly, pull out bottom knob (D) and lower the foot extension down until it is flat.

WARNING

If the stretcher is equipped with the optional foot end I.V. pole, the I.V. pole must be in the raised position when the foot extension/defibrillator tray is installed. If the I.V. pole is not raised, the foot extension will not function properly and injury could occur.

If the stretcher is equipped with the optional foot end push handles, use caution while the foot extension/defibrillator tray is installed to avoid pinching your fingers.
Preventative Maintenance

CHECKLIST

____ All fasteners secure
____ Siderails move and latch properly
____ All casters lock with brake pedal engaged
____ Steer function working properly
____ All casters secure and swivel properly
____ Body restraints working properly
____ I.V. pole intact and operating properly
____ Oxygen bottle holder intact and operating properly
____ Fowler operates and latches properly
____ Knee Gatch operates properly
____ Trendelenburg/Reverse Trendelenburg operating properly
____ No rips or cracks in mattress cover
____ Transfer boards intact and operating properly
____ Arm board support levers intact and operating properly
____ Ground chain intact
____ No leaks at hydraulic connections
____ Hydraulic jacks holding properly
____ Hydraulic drop rate set properly
____ Hydraulic oil level sufficient
____ Lubricate where required, including the brake adjuster assembly and brake cam

Serial No.____________________

____________________

____________________

____________________

____________________

Completed By:__________________________  Date:_____________

NOTE
Preventative maintenance should be performed at a minimum of annually. A preventative maintenance pro-
gram should be established for all Stryker Medical equipment. Preventative maintenance may need to be
performed more frequently based on the usage level of the product.
Model 1231/1731 stretchers are designed to be power–washable. The unit may show some signs of oxidation or discoloration from continuous washing. However, no degradation of the stretcher’s performance characteristics or functionality will occur due to power washing as long as the proper procedures are followed.

- Follow the cleaning solution manufacturer’s dilution recommendations exactly.
- Remove the mattress prior to washing the unit; do not wash the mattress with the stretcher.
- Position the Fowler at 45°, place the unit in full reverse Trendelenburg (foot end down), raise the siderails, and place the I.V. poles and push handles in the up position.
- Stryker Medical recommends the standard hospital surgical cart washer for power washing Model 1231/1731 stretchers.
- Do not replace the mattress on the stretcher until the unit is completely dry.

DO NOT STEAM CLEAN THE UNIT. Use a maximum water temperature of 180°F/68°C. Maximum air dry temperature (cart washers) is 240°F/115°C. Water pressure – 1500 psi/130.5 bar. If a hand held wand is being used to wash the unit, the pressure nozzle must be kept a minimum of 24 inches/.61m from the unit. Stretchers must have maintenance performed after a minimum of every fifth washing. Refer to the maintenance manual for specific lubrication instructions.

Failure to comply with these instructions may invalidate any/all warranties.
In general, when used in those concentrations recommended by the manufacturer, either phenolic type or quaternary type disinfectants can be used. Iodophor type disinfectants are not recommended for use because staining may result. The following products have been tested and have been found not to have a harmful effect WHEN USED IN ACCORDANCE WITH MANUFACTURERS RECOMMENDED DILUTION.*

<table>
<thead>
<tr>
<th>TRADE NAME</th>
<th>DISINFECTANT TYPE</th>
<th>MANUFACTURER</th>
<th>*MANUFACTURER'S RECOMMENDED DILUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>A33</td>
<td>Quaternary</td>
<td>Airwick (Professional Products Division)</td>
<td>2 ounces/gallon</td>
</tr>
<tr>
<td>A33 (dry)</td>
<td>Quaternary</td>
<td>Airwick (Professional Products Division)</td>
<td>1/2 ounce/gallon</td>
</tr>
<tr>
<td>Beaucoup</td>
<td>Phenolic</td>
<td>Huntington Laboratories</td>
<td>1 ounce/gallon</td>
</tr>
<tr>
<td>Blue Chip</td>
<td>Quaternary</td>
<td>S.C. Johnson</td>
<td>2 ounces/gallon</td>
</tr>
<tr>
<td>Elimstaph</td>
<td>Quaternary</td>
<td>Walter G. Legge</td>
<td>1 ounce/gallon</td>
</tr>
<tr>
<td>Franklin Phenomysan F2500</td>
<td>Phenolic</td>
<td>Purex Corporation</td>
<td>1 1/4 ounce/gallon</td>
</tr>
<tr>
<td>Franklin Sentinel</td>
<td>Quaternary</td>
<td>Purex Corporation</td>
<td>2 ounces/gallon</td>
</tr>
<tr>
<td>Galahad</td>
<td>Phenolic</td>
<td>Puritan Churchill Chemical Company</td>
<td>1 ounce/gallon</td>
</tr>
<tr>
<td>Hi–Tor</td>
<td>Quaternary</td>
<td>Huntington Laboratories</td>
<td>1/2 ounce/gallon</td>
</tr>
<tr>
<td>LPH</td>
<td>Phenolic</td>
<td>Vestal Laboratories</td>
<td>1/2 ounce/gallon</td>
</tr>
<tr>
<td>Matar</td>
<td>Phenolic</td>
<td>Huntington Laboratories</td>
<td>1/2 ounce/gallon</td>
</tr>
<tr>
<td>Omega</td>
<td>Quaternary</td>
<td>Airwick (Professional Products Division)</td>
<td>1/2 ounce/gallon</td>
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<tr>
<td>Quanto</td>
<td>Quaternary</td>
<td>Huntington Laboratories</td>
<td>1 ounce/gallon</td>
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<tr>
<td>Sanikleen</td>
<td>Quaternary</td>
<td>West Chemical Products</td>
<td>2 ounces/ gallon</td>
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<tr>
<td>Sanimaster II</td>
<td>Quaternary</td>
<td>Service Master</td>
<td>1 ounce/gallon</td>
</tr>
<tr>
<td>Vesphene</td>
<td>Phenolic</td>
<td>Vestal Laboratories</td>
<td>1 1/4 ounce/ gallon</td>
</tr>
</tbody>
</table>

Quaternary Germicidal Disinfectants, used as directed, and/or Chlorine Bleach products, typically 5.25% Sodium Hypochlorite in dilutions ranging between 1 part bleach to 100 parts water, and 2 parts bleach to 100 parts water are not considered mild detergents. These products are corrosive in nature and may cause damage to your stretcher if used improperly. If these types of products are used to clean Stryker patient handling equipment, measures must be taken to insure the stretchers are rinsed with clean water and thoroughly dried following cleaning. Failure to properly rinse and dry the stretchers will leave a corrosive residue on the surface of the stretcher, possibly causing premature corrosion of critical components.

NOTE
Failure to follow the above directions when using these types of cleaners may void this product’s warranty.

Clean Velcro AFTER EACH USE. Saturate Velcro with disinfectant and allow disinfectant to evaporate. (Appropriate disinfectant for nylon Velcro should be determined by the hospital.)

REMOVAL OF IODINE COMPOUNDS
This solution may be used to remove iodine stains from mattress cover and foam footrest pad surfaces.

1. Use a solution of 1–2 tablespoons Sodium Thiosulfate in a pint of warm water to clean the stained area. Clean as soon as possible after staining occurs. If stains are not immediately removed, allow solution to soak or stand on the surface.

2. Rinse surfaces which have been exposed to the solution in clear water before returning bed to service.
Warranty

Limited Warranty:
Stryker Medical Division, a division of Stryker Corporation, warrants to the original purchaser that its products should be free from defects in material and workmanship for a period of one (1) year after date of delivery. Stryker’s obligation under this warranty is expressly limited to supplying replacement parts and labor for, or replacing, at its option, any product which is, in the sole discretion of Stryker, found to be defective. Stryker warrants to the original purchaser that the frame and welds on its beds will be free from structural defects for as long as the original purchaser owns the bed. If requested by Stryker, products or parts for which a warranty claim is made shall be returned prepaid to Stryker’s factory. Any improper use or any alteration or repair by others in such manner as in Stryker’s judgement affects the product materially and adversely shall void this warranty. Any repair of Stryker products using parts not provided or authorized by Stryker shall void this warranty. No employee or representative of Stryker is authorized to change this warranty in any way.

Stryker Medical stretchers are designed for a 10 year expected life under normal use conditions and appropriate periodic maintenance as described in the maintenance manual for each device.

This statement constitutes Stryker’s entire warranty with respect to the aforesaid equipment. STRYKER MAKES NO OTHER WARRANTY OR REPRESENTATION, EITHER EXPRESSED OR IMPLIED, EXCEPT AS SET FORTH HEREIN. THERE IS NO WARRANTY OF MERCHANTABILITY AND THERE ARE NO WARRANTIES OF FITNESS FOR ANY PARTICULAR PURPOSE. IN NO EVENT SHALL STRYKER BE LIABLE HEREUNDER FOR INCIDENTAL OR CONSEQUENTIAL DAMAGES ARISING FROM OR IN ANY MANNER RELATED TO SALES OR USE OF ANY SUCH EQUIPMENT.

To Obtain Parts and Service:
Stryker products are supported by a nationwide network of dedicated Stryker Field Service Representatives. These representatives are factory trained, available locally, and carry a substantial spare parts inventory to minimize repair time. Simply call your local representative, or call Stryker Customer Service at (800) 327–0770.

Service Contract Coverage:
Stryker has developed a comprehensive program of service contract options designed to keep your equipment operating at peak performance at the same time it eliminates unexpected costs. We recommend that these programs be activated before the expiration of the new product warranty to eliminate the potential of additional equipment upgrade charges.

A SERVICE CONTRACT HELPS TO:
- Ensure equipment reliability
- Stabilize maintenance budgets
- Diminish downtime
- Establish documentation for JCAHO
- Increase product life
- Enhance trade–in value
- Address risk management and safety
Stryker offers the following service contract programs:

<table>
<thead>
<tr>
<th>SPECIFICATIONS</th>
<th>GOLD</th>
<th>SILVER</th>
<th>PM* ONLY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annually scheduled preventative maintenance</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>All parts,** labor, and travel</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Unlimited emergency service calls</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Priority one contact; two hour phone response</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Most repairs will be completed within 3 business days</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>JCAHO documentation</td>
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<tr>
<td>On–site log book w/ preventative maintenance &amp; emergency service records</td>
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<td></td>
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</tr>
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<td>Factory–trained Stryker Service Technicians</td>
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<tr>
<td>Stryker authorized parts</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>End of year summary</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stryker will perform all service during regular business hours (9–5)</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>

* Replacement parts and labor for products under PM contract will be discounted.
** Does not include any disposable items, I.V. poles (except for Stryker HD permanent poles), mattresses, or damage resulting from abuse.

Stryker Medical also offers personalized service contracts.

Pricing is determined by age, location, model and condition of product.

For more information on our service contracts, please call your local representative or call (800) 327–0770 (option #2).

Return Authorization:

Merchandise cannot be returned without approval from the Stryker Customer Service Department. An authorization number will be provided which must be printed on the returned merchandise. Stryker reserves the right to charge shipping and restocking fees on returned items.

SPECIAL, MODIFIED, OR DISCONTINUED ITEMS NOT SUBJECT TO RETURN.

Damaged Merchandise:

ICC Regulations require that claims for damaged merchandise must be made with the carrier within fifteen (15) days of receipt of merchandise. DO NOT ACCEPT DAMAGED SHIPMENTS UNLESS SUCH DAMAGE IS NOTED ON THE DELIVERY RECEIPT AT THE TIME OF RECEIPT. Upon prompt notification, Stryker will file a freight claim with the appropriate carrier for damages incurred. Claim will be limited in amount to the actual replacement cost. In the event that this information is not received by Stryker within the fifteen (15) day period following the delivery of the merchandise, or the damage was not noted on the delivery receipt at the time of receipt, the customer will be responsible for payment of the original invoice in full.

Claims for any short shipment must be made within thirty (30) days of invoice.

International Warranty Clause:

This warranty reflects U.S. domestic policy. Warranty outside the U.S. may vary by country. Please contact your local Stryker Medical representative for additional information.
European Representative
Stryker EMEA RA/QA Director
Stryker France
ZAC Satolas Green Pusignan
Av. De Satolas Green
69881 MEYZIEU Cedex
France

Stryker Medical
6300 S. Sprinkle Road, Kalamazoo, MI 49001-9799
(800) 327-0770
www.strykermedical.com