ProAxis®
Spinal Surgery Table

Owner’s Manual

This manual is supplied in the following versions:

- English (EN)
- Spanish (ES)
- French (FR)
- German (DE)
- Italian (IT)
- Portuguese (PT-BR)
- Japanese (JA)
- Multilingual (ML)

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1 Important Notices

⚠️ CAUTION: To ensure safe operation of the equipment, please READ THESE INSTRUCTIONS COMPLETELY and keep this manual readily available for future reference.

Carefully observe and comply with all warnings, cautions and instructions placed on the equipment or described in this manual.

**NOTE:** *This device is intended for use by trained personnel only. To schedule an in-service, please contact your domestic Mizuho OSI sales representative or call 1-800-777-4674 within the USA or +1-510-429-1500 outside the USA.*

**NOTE:** *The application techniques outlined in these instructions are the manufacturer’s suggested techniques. The final disposition of each patient’s care as related to the use of this equipment rests with the attending surgeon.*

In this manual, the following symbols are used:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="symbol" alt="ECREP" /></td>
<td>This symbol indicates an authorized representative in the European Community.</td>
</tr>
<tr>
<td><img src="symbol" alt="Manufacturer" /></td>
<td>This symbol indicates the Manufacturer of the device.</td>
</tr>
<tr>
<td><img src="symbol" alt="Date" /></td>
<td>This symbol indicates the Date of manufacture.</td>
</tr>
<tr>
<td><strong>NOTE:</strong></td>
<td>This symbol indicates a comment or instruction of importance.</td>
</tr>
<tr>
<td><img src="symbol" alt="CAUTION" /></td>
<td>This symbol is to signify CAUTION. It is intended to alert the user to consult the documentation for safety-related information such as warnings and precautions that cannot, for a variety of reasons, be presented on the device itself.</td>
</tr>
<tr>
<td><img src="symbol" alt="WARNING" /></td>
<td>This WARNING symbol is intended to alert the user of important operation, maintenance, or safety instructions.</td>
</tr>
<tr>
<td><img src="symbol" alt="Disposal" /></td>
<td>This symbol indicates proper disposal instructions.</td>
</tr>
<tr>
<td><img src="symbol" alt="Product Number" /></td>
<td>This symbol indicates a product number.</td>
</tr>
<tr>
<td><img src="symbol" alt="Serial Number" /></td>
<td>This symbol indicates a serial number.</td>
</tr>
<tr>
<td><img src="symbol" alt="Read Owner’s Manual" /></td>
<td>This symbol indicates that you must read the owner’s manual before use.</td>
</tr>
<tr>
<td><img src="symbol" alt="Refer Owner’s Manual" /></td>
<td>This symbol indicates that you are advised to refer to the owner’s manual before use.</td>
</tr>
<tr>
<td><img src="symbol" alt="Battery Safety" /></td>
<td>This symbol directs the user to follow operating instructions (related to battery safety).</td>
</tr>
<tr>
<td><img src="symbol" alt="External Ground Stud" /></td>
<td>This symbol indicates an external ground stud that is required for use when the AC power cable is not connected to a protective earth ground hospital grade AC outlet in your operating room or facility.</td>
</tr>
<tr>
<td>Symbol</td>
<td>Description</td>
</tr>
<tr>
<td>--------</td>
<td>-------------</td>
</tr>
<tr>
<td>![Person]</td>
<td>This symbol indicates this equipment is an applied part TYPE B in accordance with IEC 60601-1 and is generally suitable for applications involving external or internal contact with the patient, excluding the heart. The patient circuit is connected to protective earth and this equipment should be connected only to hospital grade AC outlets with a protective earth ground.</td>
</tr>
<tr>
<td>![Warning]</td>
<td>This symbol represents the visual alarm signal associated with the table being overloaded.</td>
</tr>
<tr>
<td>![Warning]</td>
<td>This symbol represents an Emergency Stop.</td>
</tr>
<tr>
<td>![Warning]</td>
<td>This symbol represents the table’s Weight Limit.</td>
</tr>
<tr>
<td>![Lock]</td>
<td>This symbol indicates that a Warning exists should the Torso Trolley® Chest Tray not be locked in place on the Rail Mounts.</td>
</tr>
<tr>
<td>![Warning]</td>
<td>This symbol indicates that a Warning exists related to actions to be taken, and the owner’s manual must be read before use.</td>
</tr>
<tr>
<td>![Hand]</td>
<td>This symbol represents a Pinch Hazard.</td>
</tr>
<tr>
<td>![No Touch]</td>
<td>This symbol indicates Do not touch, housing energized (related to battery safety).</td>
</tr>
<tr>
<td>![Warning]</td>
<td>This symbol indicates Warning, dangerous voltage (related to battery safety).</td>
</tr>
<tr>
<td>![Moisture]</td>
<td>This symbol indicates a medical device that needs to be protected from moisture.</td>
</tr>
<tr>
<td>![Breakage]</td>
<td>This symbol indicates a medical device that can be broken or damaged if not handled carefully.</td>
</tr>
<tr>
<td>![Orientation]</td>
<td>This symbol indicates “This End Up”, and identifies the correct orientation for safe handling.</td>
</tr>
<tr>
<td>![Temperature]</td>
<td>This symbol indicates the temperature limits to which the medical device can be exposed safely.</td>
</tr>
<tr>
<td>![Humidity]</td>
<td>This symbol indicates the range of humidity to which the medical device can be exposed safely.</td>
</tr>
<tr>
<td>![Pressure]</td>
<td>This symbol indicates the range of atmospheric pressure to which the medical device can be exposed safely.</td>
</tr>
<tr>
<td>Symbol</td>
<td>Description</td>
</tr>
<tr>
<td>--------</td>
<td>------------------------------</td>
</tr>
<tr>
<td><img src="image1.png" alt="Image" /></td>
<td>This symbol indicates Do Not Stack.</td>
</tr>
<tr>
<td><img src="image2.png" alt="Image" /></td>
<td>This symbol identifies POWER OFF.</td>
</tr>
<tr>
<td><img src="image3.png" alt="Image" /></td>
<td>This symbol identifies POWER ON.</td>
</tr>
</tbody>
</table>
WARNING: Proper preoperative and intra-operative procedures must be followed to prevent venous stasis and pooling, pressure sore development, neuropathy, improper electro-surgical tissue grounding, hypertension/hypotension, and hypothermia.

WARNING: To avoid the risk of electric shock, this equipment must only be connected to a supply mains with protective earth.

WARNING: The ProAxis® Table is to be used by personnel that receive training from either Mizuho OSI or from someone qualified by the medical facility to provide this training. Failure to comply with this requirement may result in damage to the table, possible injury to the patient or harm to the healthcare workers.

WARNING: This symbol indicates an external ground stud that is required for use when the AC power cable is not connected to a protective earth ground hospital grade AC outlet in your operating room or facility. To protect the patient, hospital staff, and the device from possible electrical hazards, an external ground wire connection is required between the external ground stud and protective earth ground when the device is in use under battery power or not connected to a protective earth ground.

WARNING: Medical electrical equipment needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided in this manual.

WARNING: Use of the ProAxis® Spinal Surgery Table with patients weighing more than 500 lbs (227 kg) could result in damage to the table, possible injury to the patient, or harm to the healthcare professional.

WARNING: When unlatching the spars from the Head-End of the ProAxis® Spinal Surgery Table for either storage or cleaning, do not use excessive force to unlatch the spars. If after unlocking the spar lock, the spar does not unlatch with hand force, seek assistance from qualified service personnel. Failure to properly unlatch the spars may cause harm to the healthcare professional and/or the device.

WARNING: Before and after each use, inspect the table, components and accessories for possible damage, excessive wear, or non-functioning parts. Carefully inspect all accessible areas, joints, and all moving parts for possible damage or non-function. Damaged or defective parts should not be used or processed. Contact Mizuho OSI Services for repair or replacement (refer to Section 13).

WARNING: The ProAxis® Spinal Surgery Table should not be operated in an oxygen-rich environment nor in the presence of flammable anesthetics, volatile substances, or other explosive gases, liquids, or atmospheres.

WARNING: No modification of the ProAxis® Spinal Surgery Table or its components is
allowed. Any modification to the equipment may result in damage to the table, possible injury to the patient or harm to the healthcare professionals.

⚠️ **CAUTION**: As outlined in the AORN Recommended Practices for Positioning a Patient in the Perioperative Setting, following the positioning of the patient, an assessment of the patient’s alignment, tissue perfusion, and skin integrity should be completed. All contact points of the patient with the table pads should be monitored during the procedure.

**NOTE:** If the integrity of the AC power source is in doubt, the equipment shall be operated from its internal electrical power source (battery).

**NOTE:** If high-frequency surgical equipment, cardiac defibrillators or cardiac defibrillator monitors are to be used with the ProAxis® Spinal Surgery Table, refer to the instructions for use provided by the manufacturer of those devices.

### 1.1 Trademarks and Patents

ProAxis®, IntelliPendant®, Torso Trolley®, GentleTouch®, ShearGuard®, and Orange Aid® are registered trademarks of Mizuho OSI.

Advanced Control Pad System™ is a trademark of Mizuho OSI.

DORO® is a registered trademark of pro med instruments GmbH.

Mayfield® is a registered trademark of Schaerer Mayfield USA, Inc.

O-arm® is a registered trademark of Medtronic, Inc.

ProneView® is a registered trademark of Dupaco, Inc.

Tempur-Pedic® is a registered trademark of Tempur-Pedic North America, Inc.

Velcro® is a registered trademark of Velcro Industries.

**Product protected by:**

- US Patent Number: 8,584,281

(Other patents pending)

### 1.2 Disposal of Electrical Components

⚠️ In accordance with the European Union Waste Electrical and Electronic Equipment (WEEE) Directive, all electrical components and batteries must be disposed of in accordance with local regulations or returned to Mizuho OSI for proper disposal. Please contact Mizuho OSI Services at 1-800-777-4674 within the USA or +1-510-429-1500 outside the USA for further information regarding this requirement.
2 Introduction

2.1 General Description

The ProAxis® Spinal Surgery Table (Figure 1) is designed to support and position a patient undergoing surgical procedures while allowing for the articulation of the patient’s spine intra-operatively through flexion and extension movements. These intra-operative movements allow the natural movement of the spine to be replicated during the surgical procedure. The table provides prone, supine, and lateral positioning capabilities, with enhanced user and patient benefits through key proprietary features. The table’s radiolucent hinge, spars, and two-piece supine top also allow for excellent intraoperative imaging using either a C-arm or O-arm®.

The ProAxis® table has electrically powered positioning and operating functions, which are controlled by means of the IntelliPendant®. The IntelliPendant® provides a visual status display for the Floor Locks, Hinge Mode, Advanced Control Pad System™ (ACP), Battery Status, Height Status, Trendelenburg/Reverse Trendelenburg Angle, Hinge Angle, and Lateral Tilt Angle. The table provides a further feature in providing the user with the option of selecting between two Hinge Mode settings – Fixed Surgical Site (FSS) or Fixed End (FE) mode.

The table includes Mizuho OSI Tempur-Pedic® medical pad technology. The material used in the manufacture of the Mizuho OSI pads has viscoelastic properties and is temperature sensitive, becoming softer where the patient’s body makes the most contact with the surface and remaining firm in the areas where less body contact is being made. Pressure is distributed evenly over the entire surface area. The pads are radiolucent and made without natural rubber latex. The benefits of using Mizuho OSI Tempur-Pedic® Medical Pads are improved pressure management, reduced shear forces, and enhanced patient comfort when used in ambient temperatures. In accordance with AORN recommendations, it is important to limit skin exposures to lower ambient temperatures, protect the patient by initiating passive warming interventions (e.g. applied forced-air warming systems, blankets, drapes and reflective composites), and to maintain an ambient room temperature of 20° to 25°C.

The Advanced Control Pad System™ is utilized when the patient is positioned prone. The pad control system, which is built into the table, is electronically controlled and pneumatically actuated, providing massage action and periodic pressure point stimulation through the Contoured ACP Hip Pads.

Figure 1: ProAxis® Spinal Surgery Table – Prone Patient Setup, Fixed Surgical Site Hinge Mode
2.2 Intended Use

The ProAxis® Spinal Surgery Table is a mobile, dual-column, carbon-fiber hinged frame surgery table designed for temporary support (<24 hours) and positioning of a patient in a prone, supine, or lateral position. The table is intended for use during surgical procedures, including radiographic imaging during such procedures. The table is not intended for use in patient transport.

ProAxis® provides a platform designed to support and position adult and pediatric patients with body weight less than 500 pounds (227 kg) that fall within the height range of 58-81 inches (147-206 cm).

The ProAxis® system, when used with the Torso Trolley® Chest Pad and the Contoured Hip Pads, shall support patients in a prone position with minimal iliac crest to iliac crest distances of 8.0 inches (20.3 cm).

2.3 User Profile

The ProAxis® Spinal Surgery Table is suitable for use by health care professionals, including but not limited to surgeons, radiologists, anesthesiologists, circulating nurses, surgical technicians, biomedical technicians, and radiology technicians.

2.4 Training Requirements

Before using the ProAxis® Spinal Surgery Table, the user must read this ProAxis® Spinal Surgery Table Owner's Manual (REF NW0725).

It is required that personnel using the ProAxis® Spinal Surgery Table receive training by either Mizuho OSI or by someone qualified by the medical facility to provide this training.

⚠️ WARNING: Failure to ensure training prior to use of this device may cause harm to the patient, healthcare professional, or the device.

2.5 Conditions of Use

The ProAxis® Spinal Surgery Table may be used several times throughout the day and night in medical facilities; e.g. hospitals, and outpatient surgery/imaging centers. The ProAxis® Spinal Surgery Table will be used in an operating room or other treatment room, and may be rolled between rooms. It shall not be used for patient transport.

⚠️ WARNING: To maximize patient safety, do not move the table with surgical equipment in vivo that is not free to move with the patient.

2.6 Product Lifetime

The product’s service lifetime is defined as 10 years. At the time of delivery, your product fulfills existing regulations and standards. However, despite proper use, routine inspection, prescribed service, maintenance and repairs, the product is subject to aging and wear. Therefore, Mizuho OSI cannot guarantee the product’s safety after ten (10) years and recommends your product be taken out of service. For product warranty information, refer to Section 13 of this manual.
2.7 Specifications

The ProAxis® Spinal Surgery Table has the following specifications:

- The maximum patient load is 500 pounds (227 kg) in a procedural position at any point within its physical range.
- The patient height range supported by the table is 58-81 inches (147-206 cm).
- Table top width:
  - The open frame used for prone positioning is 19 inches (48 cm).
  - The Supine Tops used for supine and lateral positioning are 21 inches (53 cm).
- Table top length is 81 inches (206 cm).
- When in use, the overall length of the table is 122-138 inches (310 - 351 cm) dependent on where the table is in its range of movements.
- The table can be configured to a length of 80 inches (204 cm) for storage (Figure 55, page 46).
- Defined Home position: Height of 32 inches (81 cm) top of hinged frame to floor with a level table top (i.e. 0° Hinge, Tilt, and Trendelenburg/Reverse Trendelenburg).
- Height Range: 20 inches – 47 inches (51 cm – 119 cm) when table top is Level.
- Lateral Tilt (left/right) is 0 – 15 degrees.
- Trendelenburg is 0 – 15 degrees from the Home position
- Reverse Trendelenburg is 0 – 20 degrees from the Home position
- Two Hinge Modes: Fixed Surgical Site (FSS), Fixed End (FE)
- Hinge Angle Range: -20 to +35 degrees
  - Some height adjustments may be required to achieve the maximum hinge up angle in FSS only.
  - Some height adjustments may be required to achieve the minimum hinge down angle.
- The table trapezoidal rail frame and Supine Tops are made of radiolucent carbon-fiber construction.
- Input power requirement is 100/120V, 50/60 Hz (REF 6988) or 220/240V, 50/60 Hz (REF 6988I) as indicated on the manufacturer’s label.
- As a back-up power source, the table may be operated under battery power. The expected working life of a fully charged battery is approximately eight (8) hours with two procedures averaging six (6) minutes of motion time.
- The table is IPX4 rated per IEC 60529.
- Operating environment: +50 to +86 degrees F (+10 to +30 degrees C), relative humidity 40%-70%.
- Class 1 Equipment, Type B per IEC 60601-1.
- The table is not suitable for use with flammable anesthetic gas mixtures.
2.8 Shipping and Storage

If required to be transported, the ProAxis® Spinal Surgery Table must be transported using the appropriate shipping crate. Unpacking instructions are included with the original shipping crate.

When not in use, the ProAxis® Spinal Surgery Table should be stored in a clean, dry environment.

The following conditions are required of the shipping and or storage environment:

- Ambient temperature -4°F (-20°C) to 122°F (50°C)
- Relative humidity from 10% to 95%, non-condensing
- Atmospheric pressure from 75 to 105 kPa

To prepare the table to be shipped or stored in a more compact configuration, utilize the IntelliPendant® and complete the steps outlined (Section 4.7.5). Transporting or storing the table in the stored configuration reduces the length of the table from 122 inches (310 cm) to 80 inches (204 cm).

When in storage, the table cover provided serves as a dust cover and should be used. The table cover can be adjusted with the Velcro® closures so it will fit the ProAxis® in the standard use table position (fully extended) or stored configuration (reduced length). Also, to ensure that the battery is always fully charged and ready for use, store the table connected to AC power that matches the ratings on the manufacturer’s label, located on the Foot-End column (Figure 10, page 18).
### Glossary of Terms

This glossary of terms assumes that the patient is positioned prone with his or her head oriented towards Head-End Column of the table and his or her feet towards the Foot-End Column. The functions of the IntelliPendant® and Auxiliary Control Panel are oriented for this position.

<table>
<thead>
<tr>
<th>Term</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Head-End Column</strong></td>
<td>The Head-End Column consists of one set of linkage arms and the base. The Power Switch, Emergency Stop Button, Auxiliary Control Panel, and port for the IntelliPendant® are located on the Head-End Column.</td>
</tr>
<tr>
<td><strong>Head-End Frame</strong></td>
<td>The Head-End Frame of the ProAxis® is the patient support structure comprised of the head board, rails and hinge components extending from the Head-End Column. The Head-End Frame is intended to support the patient’s upper torso, and therefore, the Head-End Frame Rails are shorter than the Foot-End Frame Rails.</td>
</tr>
<tr>
<td><strong>Foot-End Column</strong></td>
<td>The Foot-End Column consists of one set of linkage arms and the base. The manufacturer’s label and the port for connecting the Advanced Control Pad System™ Tubing are located on the Foot-End Column.</td>
</tr>
<tr>
<td><strong>Foot-End Frame</strong></td>
<td>The Foot-End Frame of the ProAxis® is the patient support structure comprised of the rails extending from the hinge to the Foot-End Column. When the patient is positioned on the table, his/her lower torso and feet are oriented toward the Foot-End Column.</td>
</tr>
<tr>
<td><strong>Retractable Center Beam</strong></td>
<td>The Retractable Center Beam connects the Head-End and Foot-End Columns. The Beam retracts, allowing the table to be transported or stored in smaller configuration.</td>
</tr>
<tr>
<td><strong>Floor Locks Unlocked</strong></td>
<td>Refers to the Floor Locks being fully retracted. The button on the IntelliPendant® and the Auxiliary Control Panel used to unlock the Floor Locks is labeled [1].</td>
</tr>
<tr>
<td><strong>Floor Locks Locked</strong></td>
<td>Refers to the Floor Locks being fully deployed. The button on the IntelliPendant® and the Auxiliary Control Panel used to lock the Floor Locks is labeled [1].</td>
</tr>
<tr>
<td><strong>Left Side of the Table</strong></td>
<td>Refers to the left side of the table (as you stand at the Head-End Column and look to the Foot-End Column). The button on the IntelliPendant® and the Auxiliary Control Panel used to laterally roll (tilt) the table top to the left is labeled [1].</td>
</tr>
<tr>
<td><strong>Right Side of the Table</strong></td>
<td>Refers to the right side of the table (as you stand at the Head-End Column and look to the Foot-End Column). The button on the IntelliPendant® and the Auxiliary Control Panel used to laterally roll (tilt) the table top to the right is labeled [1].</td>
</tr>
<tr>
<td><strong>Lower the Table</strong></td>
<td>Refers to lowering the Height of the table top. The button on the IntelliPendant® and the Auxiliary Control Panel used to lower the table top is labeled [1].</td>
</tr>
<tr>
<td><strong>Raise the Table</strong></td>
<td>Refers to raising the Height of the table top. The button on the IntelliPendant® and the Auxiliary Control Panel used to raise the table top is labeled [1].</td>
</tr>
<tr>
<td><strong>Trendelenburg</strong></td>
<td>Refers to lowering the height of the Head-End of the table top relative to the Foot-End of the table top. The button on the IntelliPendant® and the Auxiliary Control Panel used to lower the Head-End is labeled [1]. The height of the hinge is maintained during this motion.</td>
</tr>
<tr>
<td><strong>Reverse Trendelenburg</strong></td>
<td>Refers to lowering the height of the Foot-End of the table top relative to the Head-End of the table top. The button on the IntelliPendant® and the Auxiliary Control Panel used to lower the Foot-End is labeled [1]. The height of the hinge is maintained during this motion.</td>
</tr>
<tr>
<td><strong>Hinge Up</strong></td>
<td>Refers to the frame hinging such that the table is flexed upwards. The button on</td>
</tr>
<tr>
<td><strong>Hinge Down</strong></td>
<td>Refers to the frame hinging such that the table is flexed downwards. The button on the IntelliPendant® and the Auxiliary Control Panel used to hinge down the table is labeled [image].</td>
</tr>
<tr>
<td>----------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>Advanced Control Pad System™ (ACP)</strong></td>
<td>The Advanced Control Pad System™ provides user-selectable, pressure point stimulation to the patient’s hips, thighs and chest when positioned prone. The Contoured Hip Pads have built in cells that alternate between inflating and deflating with air.</td>
</tr>
</tbody>
</table>
| **Return to Level** | Refers to the motion of the table required to return the table top to the neutral position. When the Return to Level function is activated, the Lateral Tilt Angle returns to 0 degrees; Trendelenburg / Reverse Trendelenburg Angle returns to 0 degrees and the Hinge Angle returns to 0 degrees. Following a one (1) second pause, if the button continues to be depressed, the table top will return to a height of 32 inches (81 cm) above the floor, which is defined as its Home position.  

**NOTE:** The complete functionality of the Return to Level feature requires either the Rail Mounts or the Supine Tops to be installed on the open frame to be operational. If either of these components is not installed, the Return to Level feature will only return the Lateral Tilt Angle to 0 degrees and the table top height to 32 inches (81 cm) above the floor. |
| **Home Position** | The position of the ProAxis® when the table top is level head-to-foot and side-to-side, the hinge is neutral and the height of the table top is 32 inches (81 cm) above the floor. |
| **Fixed Surgical Site (FSS) Hinge Mode** | The hinge mode of the Frame that raises and lowers the Head-End and Foot-End while maintaining the height of the hinge as it is raised or lowered. This motion approximates “fixing” the surgical site at a constant position.  

When FSS Hinge Mode is selected and the Hinge Angle is positive, the [image] symbol is visible on the display of the IntelliPendant®.  

When FSS Hinge Mode is selected and the Hinge Angle is negative, the [image] symbol is visible on the display of the IntelliPendant®. |
| **Fixed End (FE) Hinge Mode** | The hinge mode of the Frame that maintains the Head-End and Foot-End at a constant height as the hinge is raised and lowered.  

When FE Hinge Mode is selected and the Hinge Angle is positive, the [image] symbol is visible on the display of the IntelliPendant®.  

When FE Hinge Mode is selected and the Hinge Angle is negative, the [image] symbol is visible on the display of the IntelliPendant®. |
| **Hospital Grade AC Outlet** | Refers to specially designated outlets (receptacles) that include additional grounding reliability, assembly integrity, strength, and durability. A hospital grade outlet in the United States may be indicated by a green colored dot on the face of the outlet. |
3 Component Identification

3.1 Table Orientation

The ProAxis® Spinal Surgery Table is described as having a Head-End and a Foot-End (Figure 2). When the components are set up and the patient is positioned, the patient’s head is oriented towards the Head-End of the device and his/her feet are oriented towards the Foot-End of the device.

![Figure 2: ProAxis® Spinal Surgery Table](image)

The linkage arms of the columns raise, lower, and rotate when the table is in use, and may come in contact with items located on or near the table, potentially causing damage to the table or item. The base surfaces at the Head-End and Foot-End columns are not intended to be used for storage.

⚠️ CAUTION: Do not place any items on the base of the Head-End or Foot-End columns or below the table. Storing or placing anything in these areas may result in damage to the device or the item being stored.

An important safety label regarding Patient Weight Capacity is also located at the Head-End of the table top (Figure 3).

![Figure 3: Patient Weight Capacity Label at Head-End](image)
**WARNING**: Use of the ProAxis® Spinal Surgery Table with patients weighing more than 500 pounds (227 kg) could result in damage to the table, possible injury to the patient, or harm to the healthcare professionals.

The Power Switch, Battery, and AC Power Indicator LEDs, Power Receptacle, and Circuit Breakers are located on the Head-End table base.

![ProAxis® Table Base, Head-End](image)

**Figure 4: ProAxis® Table Base, Head-End**

The Hand Pendant, called the IntelliPendant®, the Auxiliary Control Panel, the Emergency Stop, and Cord Wrap are located on the Head-End column.
3.1.1 IntelliPendant®

The ProAxis® Spinal Surgery Table functions are controlled by the single hand pendant referred to as the IntelliPendant® (Figure 5).

The graphical display of the IntelliPendant® provides real time information about the table’s position: Height, Trendelenburg / Reverse Trendelenburg angle, Lateral Tilt angle and Hinge angle. The IntelliPendant® also informs the user of the table’s status for Power, Hinge Motion, ACP Setting, and Floor Locks. The table’s functions and settings, which include Hinge Mode, Memory Position, Advanced Control Pad System™ Cycles, Floor Lock Override, and Storage Configuration, are managed through the use of the IntelliPendant®.

When the table is in use, the current Height is identified on the display screen in inches/centimeters. The angle of Trendelenburg/Reverse Trendelenburg, Lateral Tilt, and the Hinge is displayed in degrees. The Hinge angle displayed when the hinge is raised or lowered will range from -20 to +35 degrees as the hinge angle changes.

Detailed instructions for utilizing the IntelliPendant® are provided in Section 4.

3.1.2 Cord Wrap

A Cord Wrap is provided on the inside of the Head-End column for convenient storage of the Power Cord when not in use (Figure 5).

3.1.3 Emergency Stop

An Emergency Stop is provided on the Head-End column above the connection port for the IntelliPendant® (Figure 5). Pressing this button will immediately stop any motion of the ProAxis® Spinal Surgery Table.

Detailed instructions for utilizing the Emergency Stop are provided in Section 4.
3.1.4 Auxiliary Control Panel

The Auxiliary Control Panel (Figure 5) is intended to provide a second option for controlling primary table motions and Floor Lock function. The Auxiliary Control Panel may be utilized to control table Height, Trendelenburg/Reverse Trendelenburg, Lateral Tilt, Hinge motion, and Floor Lock operation.

**NOTE:** *Should the IntelliPendant® stop functioning, the Auxiliary Control Panel will allow the user to execute the basic motions of the table and lock/unlock the Floor Locks.*

Detailed instructions for utilizing the Auxiliary Control Panel are provided in Section 4, Basic Operation.

3.1.5 Advanced Control Pad System™

![Figure 6: ProAxis® Foot-End Column](image)

The Advanced Control Pad System™ Connection Port is located on the Foot-End column (Figure 6). Detailed instructions for utilizing the Advanced Control Pad System™ are provided in Section 4.
3.2 Storage Cart

A Storage Cart is provided for ease of storing the table’s standard components and optional accessories when not in use. The silhouettes on the Cart provide guidance on where each component or accessory should be stored. The front side of the Cart is primarily dedicated to the storage of standard components (Figure 7), while the back side of the Cart is designed for the storage of most optional accessories (Figure 8).

![Figure 7: Front Side of Storage Cart](image)

**NOTE:** Ensure the Chest Tray is stored as shown within the ledge of the Cart to prevent damage to the pieces of the component when removing or handling.
3.3 Component Part Number

Each standard component of the ProAxis® Spinal Surgery Table is identified and labeled with its respective part number.

![ProAxis® Component Label](image)

Figure 8: Back Side of Storage Cart

Figure 9: ProAxis® Component Label
3.4 Model Number and Serial Number

In addition to the product number and serial number, the following information is provided on the manufacturer’s label:

![Manufacturer's Label, Examples](image)

**NOTE:** Additional product information may also be found on the label.
4 Basic Operation

4.1 Control Operation

For use with AC power:

1. Plug the Power Cord into a properly grounded receptacle. Refer to the manufacturer’s label at the Foot-End column for input voltage requirements (Figure 10, page 18).

   **NOTE:** *Only use the Power Cord supplied by Mizuho OSI with the table or if necessary, one of equivalent rating.*

2. Depress the Power Switch and observe that the switch illuminates with a green ring indicating that power is applied to the table.

3. If the AC power cable is not connected to an outlet with a protective earth ground, then the external ground stud should be connected to a protective earth ground (Figure 4, page 13).

   **WARNING:** This symbol indicates an external ground stud that is required for use when the AC power cable is not connected to a protective earth ground hospital grade AC outlet in your operating room or facility or when the table is in use under battery power. Failure to ensure ground may cause harm to the patient, healthcare professionals, or the device.

   **NOTE:** *If the integrity of the AC power source is in doubt, the equipment shall be operated from its internal electrical power source (battery).*

Should the table be operated on battery power:

1. Ensure that the battery has been properly charged as outlined in Section 12.3. Depress the Power Switch and observe that the switch illuminates with a green ring indicating that power is applied to the table. The battery LED at the base of the Head-End column will not be illuminated, indicating that the table is being used on battery power (Figure 4, page 13).

2. The battery status symbol on the screen of the IntelliPendant® indicates the level of charge in the batteries. If the battery status symbol with one green bar appears, the battery is not charged and the table should only be used with AC power.

   **WARNING:** Failure to ensure the table is properly charged when used on battery power may cause the table to malfunction and may harm the patient, healthcare professional and the table.

3. To protect the patient, hospital staff, and the table from possible electrical hazards, an external ground cable connection is required between the external ground stud and protective earth ground when the table is in use under battery power (Figure 4, page 13).

The expected working life of a fully charged battery is approximately eight (8) hours with two procedures averaging a total of six (6) minutes of motion time. While in storage, it is recommended that the table be plugged in to charge the battery. A fully depleted battery will take 24 hours to fully charge. This can be accomplished when using the table under AC power.

If during use, the battery symbol on the IntelliPendant® appears, the table should be connected to AC power, and it can continue to be utilized while the battery is charging.
4.2 Emergency Stop

To immediately interrupt the motion of the ProAxis® Spinal Surgery Table, press the Emergency Stop button located on the side of the Head-End column (Figure 12). The IntelliPendant® will display if the Emergency Stop is engaged (Figure 14), and the IntelliPendant® will vibrate. No further table functions will be allowed until the Emergency Stop is reset. To reset the table, turn the Emergency Stop button in the direction of the arrows until it returns to the original position (Figure 13).

4.3 Auxiliary Control Panel

The Auxiliary Control Panel provides function and motion buttons labeled with symbols identifying what function or motion can be achieved (Figure 15, page 21). The motion buttons control the Height, Trendelenburg/Reverse Trendelenburg, Lateral Tilt Right/Left, Hinge Up/Down, and Return to Level. The two (2) function buttons control the table’s Floor Lock/Unlock capabilities.

A green Light Emitting Diode (LED) illuminates on the face of the Auxiliary Control Panel when the Floor Locks are fully deployed and locked, when the table frame has returned to a level position, and when the table is prepared for storage.

In order for the motion buttons on the Auxiliary Control Panel to function, the Floor Locks must be fully deployed. To lock the Floor Locks, press the Floor Lock button on the Auxiliary Control Panel or the IntelliPendant®. Confirm the Floor Locks are engaged by observing that the Floor Lock LED illuminates green and the Floor Lock symbol is visible on the IntelliPendant®.
To utilize the Auxiliary Control Panel, press and hold a button until the desired motion is achieved. A display on the IntelliPendant® will show the symbol of the motion button when it is activated (Figure 23, page 26).

![Diagram of Auxiliary Control Panel]

**Figure 15: Auxiliary Control Panel**

### 4.4 Floor Locks

When the ProAxis® Spinal Surgery Table is powered on, the Floor Locks will automatically return to their previous state of being either fully deployed (locked) or fully retracted (unlocked).

To lock the Floor Locks via the IntelliPendant® or the Auxiliary Control Panel, press the Floor Lock button once and release to deploy the Locks (Figure 16). The IntelliPendant® will display when the Floor Locks are fully deployed and locked (Figure 22, page 25). On the Auxiliary Control Panel, the Floor Lock LED will illuminate (Figure 15). To unlock the Floor Locks via the IntelliPendant® or the Auxiliary Control Panel, press the Floor Unlock button once to completely retract the Floor Locks (Figure 17). The IntelliPendant® will display when the Floor Locks are fully retracted and unlocked (Figure 21, page 25). It will take approximately 10 to 15 seconds for the Floor Locks to change position.

**NOTE:** The ProAxis® table will not function until the Floor Locks are completely deployed and the symbol is visible on the IntelliPendant® display.

![Floor Lock Deployed (Locked)](image1)

**Figure 16: Floor Lock Deployed (Locked)**

![Floor Lock Retracted (Unlocked)](image2)

**Figure 17: Floor Lock Retracted (Unlocked)**
4.4.1 IntelliPendant® Activated Floor Lock Override

If the Floor Lock systems fail during table use such that powered table motion is prevented, the IntelliPendant® can be used to activate the Floor Lock Override function (see Section 4.7.4.4, page 35). Use of the Floor Lock Override function will allow for 60 seconds of activity during which multiple actions can be taken.

**NOTE:** Should a particular motion take longer than 60 seconds, the action will be completed as long as the function button continues to be depressed.

**NOTE:** The Floor Lock Override function allows for changes to Height, Trendelenburg/Reverse Trendelenburg, Lateral Roll, Hinge, and Return to Level functions via the Hand Pendant and is intended for use in emergency situations only.

4.4.2 Manual Floor Lock Override

In the event the table’s floor locks do not retract properly to allow for table transport, the Floor Locks can be manually raised or lowered (see Section 0).

**NOTE:** The manual floor lock override is intended only as a means of transporting the table should the Floor Lock systems fail. Use of this override will not allow the user to activate any of the table functions.

4.5 Moving the Table

Before moving the ProAxis® Spinal Surgery Table or preparing it for its storage sequence of movements, ensure the table is returned to its Home position such that it is neutral and at a maximum height of 32 inches (81 cm).

⚠️ **CAUTION:** When moving the ProAxis® Spinal Surgery Table in its normal configuration, ensure that the table is at a height no greater than its home position of 32 inches (81 cm) and in a neutral state with the IntelliPendant® reading 0º for Lateral Roll, Trendelenburg/Reverse Trendelenburg and Hinge to prevent the table from being unstable during relocation.

After all four (4) Floor Locks are unlocked, the table can be rolled for relocation. The ProAxis® Spinal Surgery Table is heavy, and a minimum of two people is required to move it. Position one person at the Head-End of the table and one person at the Foot-End. Care should be taken to control the ProAxis® when rolling it.

⚠️ **CAUTION:** If the ProAxis® Spinal Surgery Table is allowed to roll too fast, it may be difficult to stop or turn. Impact of the table with a stationary object may cause serious damage to the table or the other object. If an impact does occur the table must be visually inspected for damage, and a Function Check must be performed (refer to Section 6). If damage is discovered or the table does not successfully complete the Function Check, call Mizuho OSI Services (refer to Section 13).

**NOTE:** For easier moving and storage, the ProAxis® Spinal Surgery Table may also be configured to its retracted position of 80 inches (203 cm) (see Section 4.7.5).
4.6 Torso Trolley®

The Torso Trolley® is designed to support the patient and simultaneously move the prone patient’s upper torso in conjunction with the hinge angle being changed, preventing any spinal compression or distraction.

The Torso Trolley® is formed by utilizing the Chest Tray to which the chest pad is mounted. The 5-Way Adjustable Arm Boards are also mounted to the Chest Tray utilizing the Arm Board Brackets provided. Refer to Section 8 for detailed instructions on the setup and use of the Torso Trolley®.

4.7 IntelliPendant®

The ProAxis® Spinal Surgery Table functions are controlled by the IntelliPendant®. When the table is in use, the display screen provides the user with detailed real time information as to the status of each motion or function. This includes specific data on the table’s position including Height, Trendelenburg/Reverse Trendelenburg Angles, Lateral Tilt Angle, and Hinge Angle. To activate a function or motion, a button must be pressed and held until the function is complete or the desired position is achieved. The IntelliPendant® also provides important information about the system’s operational state, including Power status, Hinge Motion, Advanced Control Pad setting, and Floor Lock status (Figure 18).

Utilizing the Soft Keys allows the user to access and select the following functions and settings: Hinge Mode, Memory Position, Advanced Control Pad System™ Cycles, and Floor Lock Override (Figure 18).

Figure 18: IntelliPendant®
The IntelliPendant® also provides haptic feedback to the user when a requested action is not possible. This may occur under the following conditions:

- Floor Locks are Unlocked and must be engaged before continuing.
- The maximum range of a particular motion has been reached.
- A system error has occurred.

A visual indication of the constrained movement will also be displayed on the IntelliPendant® screen informing the user of the table’s state. For instances where a particular table function has reached its limit, the image depicted on the screen will show the location of the error or limit through the use of an orange circle (see Section 4.7.2).

To connect:

1. Align the pins with the port located at the Head-End column of the table (Figure 19). Insert until the locking collar clicks, securing the pins in the port.

![Figure 19: Connecting the IntelliPendant®](image)

**NOTE:** Only connect the IntelliPendant® supplied with the ProAxis® Spinal Surgery Table to the IntelliPendant® port.

2. The IntelliPendant® should remain connected and be stored on the hand pendant clip on the Head-End column when not in use (Figure 5, page 14).

Press the Power Switch on the Head-End base once the IntelliPendant® has been connected. A green ring will illuminate around the switch confirming power is applied to the table (Figure 4, page 13). The Graphical Display that appears when the ProAxis® table is turned on is identified as the IntelliPendant® screen (Figure 20).
The Floor Locks must be fully deployed for the IntelliPendant® or Auxiliary Control Panel motion buttons to function. To lock the Floor Locks, press the Floor Lock button once on the IntelliPendant® or the Auxiliary Control Panel and confirm the Floor Lock symbol is visible on the IntelliPendant®, indicating the Floor Locks have been engaged (Figure 22).
4.7.1 Table Functions

When in use, the table’s position at any given time is identified on the IntelliPendant® (Figure 18, page 23) as follows:

- Height reported in inches/centimeters
- Angle of Trendelenburg/Reverse Trendelenburg reported in degrees
- Angle of Lateral Tilt reported in degrees
- Angle of the Hinge reported in degrees

The function being executed is also reflected on the IntelliPendant® screen, informing the user of the action taking place (Figures 23 and 24).

Figure 23: Hinge Up Action Displayed

Figure 24: Hinge Down Action Displayed
4.7.2 Defined Function and Setting Symbols

The following symbols are utilized on the function buttons of the IntelliPendant®, the Auxiliary Control Panel, or on the display screen itself when in use.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Height Up" /></td>
<td>Height Up</td>
</tr>
<tr>
<td><img src="image" alt="Height Down" /></td>
<td>Height Down</td>
</tr>
<tr>
<td><img src="image" alt="Trendelenburg" /></td>
<td>Trendelenburg</td>
</tr>
<tr>
<td><img src="image" alt="Reverse Trendelenburg" /></td>
<td>Reverse Trendelenburg</td>
</tr>
<tr>
<td><img src="image" alt="Lateral Tilt, Right" /></td>
<td>Lateral Tilt, Right</td>
</tr>
<tr>
<td><img src="image" alt="Lateral Tilt, Left" /></td>
<td>Lateral Tilt, Left</td>
</tr>
<tr>
<td><img src="image" alt="Hinge Up" /></td>
<td>Hinge Up</td>
</tr>
<tr>
<td><img src="image" alt="Hinge Down" /></td>
<td>Hinge Down</td>
</tr>
<tr>
<td><img src="image" alt="Lock (Floor Locks) and Locked Position" /></td>
<td>Lock (Floor Locks) and Locked Position</td>
</tr>
<tr>
<td><img src="image" alt="Unlock (Floor Locks) and Unlocked Position" /></td>
<td>Unlock (Floor Locks) and Unlocked Position</td>
</tr>
<tr>
<td><img src="image" alt="Half-lock Position; indicates the transitional stage of the Floor Locks while they are in the process of being either locked or unlocked" /></td>
<td>Half-lock Position; indicates the transitional stage of the Floor Locks while they are in the process of being either locked or unlocked</td>
</tr>
<tr>
<td><img src="image" alt="Lock Error; indicates the Floor Locks must be locked before executing any table function" /></td>
<td>Lock Error; indicates the Floor Locks must be locked before executing any table function</td>
</tr>
<tr>
<td><img src="image" alt="Represents Return to Level on function button, also appears on the screen display to indicate height position" /></td>
<td>Represents Return to Level on function button, also appears on the screen display to indicate height position</td>
</tr>
<tr>
<td><img src="image" alt="Menu" /></td>
<td>Menu</td>
</tr>
<tr>
<td><img src="image" alt="ACP ON/OFF Function" /></td>
<td>ACP ON/OFF Function</td>
</tr>
<tr>
<td><img src="image" alt="Save to Memory Position" /></td>
<td>Save to Memory Position</td>
</tr>
<tr>
<td><img src="image" alt="Go to Memory Position" /></td>
<td>Go to Memory Position</td>
</tr>
<tr>
<td><img src="image" alt="Cancel" /></td>
<td>Cancel</td>
</tr>
<tr>
<td><img src="image" alt="Accept / Select" /></td>
<td>Accept / Select</td>
</tr>
<tr>
<td>Symbol</td>
<td>Description</td>
</tr>
<tr>
<td>--------</td>
<td>-------------</td>
</tr>
<tr>
<td>![image]</td>
<td>Advanced Control Pad Cycle, Fast</td>
</tr>
<tr>
<td>![image]</td>
<td>Advanced Control Pad Cycle, Slow</td>
</tr>
<tr>
<td>![image]</td>
<td>Fixed End Hinge Mode, Hinge Up</td>
</tr>
<tr>
<td>![image]</td>
<td>Fixed End Hinge Mode, Hinge Down</td>
</tr>
<tr>
<td>![image]</td>
<td>Fixed Surgical Site Hinge Mode, Hinge Up</td>
</tr>
<tr>
<td>![image]</td>
<td>Fixed Surgical Site Hinge Mode, Hinge Down</td>
</tr>
<tr>
<td>![image]</td>
<td>Home (Return to Level) Position Achieved</td>
</tr>
<tr>
<td>![image]</td>
<td>Floor Lock Override</td>
</tr>
<tr>
<td>![image]</td>
<td>Indicates the system is running on battery power or AC power, and the battery voltage meets or exceeds the 100% charged threshold</td>
</tr>
<tr>
<td>![image]</td>
<td>Indicates the system is running on battery power, and the battery voltage exceeds the 75% threshold but not the 100% threshold</td>
</tr>
<tr>
<td>![image]</td>
<td>Indicates the system is running on battery power, and the battery voltage exceeds the 50% threshold but not the 75% threshold</td>
</tr>
<tr>
<td>![image]</td>
<td>Indicates the system is running on battery power, and the battery voltage is below the 50% threshold. Requires Charging and AC Power Connection</td>
</tr>
<tr>
<td>![image]</td>
<td>Battery Charging</td>
</tr>
<tr>
<td>![image]</td>
<td>AC Power Connected</td>
</tr>
<tr>
<td>M1</td>
<td>Displayed when user has saved the current position as M1, or when the table has arrived at the stored position M1</td>
</tr>
<tr>
<td>M2</td>
<td>Displayed when user has saved the current position as M2, or when the table has arrived at the stored position M2</td>
</tr>
<tr>
<td>![image]</td>
<td>Indicates that the requested motion would exceed a boundary for height up, hinge angle increase, or Trendelenburg angle increase</td>
</tr>
<tr>
<td>![image]</td>
<td>Indicates that the requested motion would exceed a boundary for height down, hinge angle decrease, or Trendelenburg angle decrease</td>
</tr>
<tr>
<td>![image]</td>
<td>Indicates that the requested left tilt motion would exceed a boundary</td>
</tr>
<tr>
<td>![image]</td>
<td>Indicates that the requested right tilt motion would exceed a boundary</td>
</tr>
<tr>
<td>![image]</td>
<td>Appears when a &quot;hinge up&quot; button is held, and the maximum hinge angle has been reached</td>
</tr>
<tr>
<td>or</td>
<td>Appears when a &quot;hinge up&quot; button is held, and a boundary has been reached</td>
</tr>
<tr>
<td>Condition</td>
<td>Description</td>
</tr>
<tr>
<td>--------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Appears when a “hinge down” button is held, and a boundary has been reached</td>
<td></td>
</tr>
<tr>
<td>Appears when a “height up” button is held, and a boundary has been reached</td>
<td></td>
</tr>
<tr>
<td>Appears when a “height down” or “home” button is held, and a boundary has been reached</td>
<td></td>
</tr>
<tr>
<td>Appears when a “Trendelenburg” button is held, and a boundary has been reached</td>
<td></td>
</tr>
<tr>
<td>Appears when a “Reverse Trendelenburg” button is held, and a boundary has been reached</td>
<td></td>
</tr>
<tr>
<td>Appears when a “Lateral Roll Right” button is held, and a boundary has been reached</td>
<td></td>
</tr>
<tr>
<td>Appears when a “Lateral Roll Left” button is held, and a boundary has been reached</td>
<td></td>
</tr>
<tr>
<td>Appears when an upward hinge motion is requested, and hinge lockout is in effect (due to the hinge enable cables on Torso Trolley® or supine top not being plugged in) Note that hinge and memory buttons can do this.</td>
<td></td>
</tr>
<tr>
<td>Appears when a downward hinge motion is requested, and hinge lockout is in effect (due to the hinge enable cables on Torso Trolley® or supine top not being plugged in) Note that hinge and memory buttons can do this.</td>
<td></td>
</tr>
<tr>
<td>Appears when the Emergency Stop is enabled</td>
<td></td>
</tr>
</tbody>
</table>
4.7.3 Return to Level

The IntelliPendant® is equipped with a Return to Level button that allows the table to be returned to a neutral position from any current state. When the Return to Level function is activated, the Lateral Tilt Angle returns to 0 degrees, Trendelenburg/Reverse Trendelenburg Angle returns to 0 degrees, and the Hinge Angle returns to 0 degrees. Following a one (1) second pause, if the button continues to be depressed, the table top will return to a height of 32 inches (81 cm) above the floor, which is defined as its Home position.

**NOTE:** The complete functionality of the Return to Level feature requires either the Rail Mounts or the Supine Tops to be installed on the open frame to be operational. If neither of these components is installed, the Return to Level feature will only return the Lateral Tilt Angle to 0 degrees and the table top height to 32 inches (81 cm) above the floor. This is intended to prevent unintentional traction via the automatic return of Trendelenburg/Reverse Trendelenburg to neutral when using Cervical Management. The Hinge capability is not enabled when neither Rail Mounts nor the Supine Tops are installed.

![Figure 25: Display when Return to Level Function is Completed](image)

4.7.4 Soft Keys Functions and Settings

To control system functions and settings, the Soft Keys corresponding to the icons at the bottom of the screen are utilized (Figure 26).
Pressing the Menu Key provides the user with a list of setting choices for the table, which may be selected and customized as needed for each procedure (Figure 27).

![Set Hinge Mode]

Set ACP Mode
Override Floor Locks
Store Table
Set Language
Set Units
Version Info

Figure 27: Menu Key Selections

4.7.4.1 Memory Positioning

The ProAxis® Spinal Surgery Table has the ability to store two specified table positions in memory for ease of returning to the desired position at a later time. The table positions can be stored to either Memory Position 1 or Memory Position 2.

Assignment of the table position to memory is completed through the use of the IntelliPendant®.

To assign the table’s current position to a memory position:

1. From the Home Screen, select the Save to Memory Position using the corresponding Soft Key (Figure 26).
2. Assign the table position to either the Memory Position 1 or Memory Position 2 location (Figure 28).

![Save Current Position to Memory 1 or Memory 2 by Selecting Below]

Figure 28: Memory Position Selection

3. The table position is now saved.
To move the table from its current position to a saved memory position:

1. From the Home Screen, select the Go To Memory Position using the corresponding Soft Key (Figure 26).
2. Press and hold the Soft Key corresponding to either Memory Position 1 or Memory Position 2 to move to the desired location.

**NOTE:** If the hinge capability is not enabled (see Section 8 for setup instructions) and the desired Memory Position requires a change in hinge angle, the selection of Memory Positions will not provide the desired motion to the stored position requested.

### 4.7.4.2 Hinge Mode

The ProAxis® Spinal Surgery Table has two operating modes for the hinged frame, Fixed Surgical Site (FSS) or Fixed End (FE), which can be selected depending on the surgeon's preference for positioning the patient.

FSS mode is designed to ensure the height of the top at the hinge remains constant when the hinge is flexed or extended. This is accomplished through the simultaneous motions of the Head-End and Foot-End of the table. This mode is utilized primarily for prone patient positioning.

FE mode is designed to ensure the height of the Head-End and Foot-End of the table top remains at a constant height when the hinge is flexed or extended. This mode is utilized primarily for lateral patient positioning.

**NOTE:** For the hinge of the frame to be operational, the Rail Mounts or the Supine Top, Head-End needs to be installed and the cables connected. Refer to Section 8 for setup instructions.

Selection of the hinge mode is completed through the use of the IntelliPendant®.

To activate the desired Hinge Mode:

1. Press the Menu Soft Key on the IntelliPendant® and observe that the Set Hinge Mode setting is selected (Figure 27).
2. Upon confirming that Set Hinge Mode is highlighted, press the Soft Key that corresponds to the check mark.
3. Utilize the arrow Soft Keys to highlight the FSS or FE mode, and press the check mark Soft Key to select the desired hinge mode (Figure 29).
4. Selection of the desired hinge mode can be visually confirmed through the corresponding symbol depicted on the IntelliPendant® display (Figures 30 and 31).

![Figure 30: FSS Mode, Positive Hinge Angle](image1)

![Figure 31: FE Mode, Negative Hinge Angle](image2)

### 4.7.4.3 Advanced Control Pad System™ (ACP)

The ProAxis® provides the Advanced Control Pad System™ (ACP) for use when the patient is positioned prone. The ProAxis® Contoured ACP Hip Pads, the Bilateral Winged ACP Chest Pad, and the Flat ACP Chest Pad are designed specifically with Tempur-Pedic® medical material and air cells within the pads. The cells inflate and deflate at selectable intervals providing massage action and periodic pressure point stimulation to the patient’s chest, hips, and thighs when positioned prone. At the discretion of the surgeon, the pads can also be used for patient support on the ProAxis® without the massage action being activated.

The ProAxis® Spinal Surgery Table has two speed settings, Fast and Slow, for the ACP System.

- The Fast Cycle, alternates inflation and deflation of the air cells every four (4) seconds. The Fast Cycle is represented with the symbol.
- The Slow Cycle, alternates inflation and deflation of the air cells every six (6) seconds. The Slow Cycle is represented with the symbol.

**NOTE:** Selection of the ACP System cycle is at the discretion of the surgeon.

Selection of the ACP System cycle is completed through the use of the IntelliPendant®.

To activate the ACP System:

1. Press the Menu Soft Key on the IntelliPendant®.
2. Utilize the Down arrow Soft Key to select **Set ACP Mode** (Figure 32). Press the Soft Key that corresponds to the check mark.
3. Utilize the Up or Down arrow Soft Keys to highlight either the Fast Cycle or Slow Cycle, and press the check mark Soft Key to select the desired speed (Figure 33).

4. Return to the Main IntelliPendant® Screen (Figure 26, page 30) and press the ACP Soft Key to activate.

**NOTE:** The ACP system will not function if the ACP function is not selected on the main screen of the IntelliPendant®.

5. Selection of the desired ACP setting can be visually confirmed through the corresponding symbol depicted on the IntelliPendant® display (Figures 34 and 35).
   a. The Fast Cycle is represented with the symbol (1 cycle every four (4) seconds).
   b. The Slow Cycle is represented with the symbol (1 cycle every six (6) seconds).

**NOTE:** If the ACP system has not been selected, no symbol will appear on the screen, indicating the ACP function is not in use.
4.7.4.4 Floor Lock Override

To allow for the use of the Floor Lock Override functionality:

1. Press the Menu Soft Key on the IntelliPendant®.
2. Utilize the Down arrow Soft Key to select the Override Floor Locks option (Figure 36).
3. Upon confirming that Override Floor Lock is highlighted, press the Soft Key that corresponds to the check mark.

4. Utilize the Down arrow Soft Key to select the Override mode (Figure 37).
5. Upon confirming that the Override Floor Lock mode Override is highlighted, press the check mark Soft Key (Figure 37). The Override functionality will now be enabled.
6. Press the appropriate motion button on the IntelliPendant® to achieve the desired table motion. The Floor Lock Override capability will allow for 60 seconds of activity during which multiple actions can be taken. Should a desired motion take longer than 60 seconds, the action will be completed as long as the motion button continues to be depressed.

7. Should additional time be required beyond the 60 seconds to complete an additional function or if the motion button has been released once the 60 seconds have elapsed, the Floor Lock Override capability will have to be activated starting with Step 1 above.

**NOTE:** The Floor Lock Override function allows for changes to Height, Trendelenburg/Reverse Trendelenburg, Lateral Roll, Hinge, and Return to Level motions via the Hand Pendant and is intended for use in emergency situations only.

### 4.7.4.5 Language Selection

To select the preferred language display for the IntelliPendant®:

1. Press the Menu Soft Key on the IntelliPendant®.
2. Utilize the Down arrow Soft Key to select Set Language (Figure 38). Press the Soft Key that corresponds to the check mark.
3. Utilize the arrow Soft Keys to highlight the desired language, and press the check mark Soft Key to set the language mode (Figure 39).

![Figure 39: Language Selection Menu](image)

**NOTE:** The language setting is preserved when the table is powered off.

### 4.7.4.6 Display Units Selection

To select the preferred units display for the IntelliPendant®:

1. Press the Menu Soft Key on the IntelliPendant®.
2. Utilize the Down arrow Soft Key to select **Set Units** (Figure 40). Press the Soft Key that corresponds to the check mark.

![Figure 40: Set Units Selection](image)

3. Utilize the arrow Soft Keys to highlight the desired units, and press the check mark Soft Key to set the units mode (Figure 41).
4.7.5 Configuration of Table for Transport or Storage

The ProAxis® table can be configured to a smaller footprint for transport or storage. To prepare the table to be shipped or stored in a more compact configuration, utilize the IntelliPendant® and complete the steps outlined. Transporting or storing the table in the stored configuration reduces the length of the table from 122 inches (310 cm) to 80 inches (204 cm).

**NOTE:** Do not attempt to disconnect the spars from their connection point without performing the exact procedure described below.

**WARNING:** Failure to properly disconnect the spars using the process below may cause injury to the healthcare professional and/or damage to the table.

To configure the table to its stored position:

1. Press the Menu Soft Key on the IntelliPendant®.
2. Utilize the Down arrow Soft Key to select **Store Table** (Figure 42). Press the Soft Key that corresponds to the check mark.
NOTE: If the ProAxis® Table is under too much load such that it is unsafe to disconnect the spars, a blue warning screen identifying an “ERROR” will appear on the IntelliPendant® and the user will not be able to proceed with the storage sequence (Figure 43).

![ERROR: Main Controller]

Figure 43: Store Table Screen

WARNING: Attempting to disconnect the spars when this “ERROR” message appears may cause injury to the healthcare professional and/or damage to the table.

3. Following the instructions depicted on the IntelliPendant®, complete the following steps to configure the table to its stored position.
   a. STEP 1: Remove any components and accessories attached to the table (Figure 44). If the Supine Top or Chest Tray is detected, the user will be notified via the IntelliPendant® screen.

![Figure 44: ProAxis® with All Components Removed in Preparation for Storage]

b. STEP 2: Confirm the next step to allow the table to move to its storage position. Continue to hold down the Soft Key until the configuration is completed (Figure 45).
c. **STEP 3:** Confirm selection of storage position using the Soft Key that corresponds to the check mark. After continuing with this selection, confirm the Storage LED light on the Control Panel is illuminated green.

**NOTE:** Upon confirmation, the remaining steps must be completed to store the table.

d. **STEP 4:** Disconnect the Spars by:
   
i. Loosening the Spar Lock on either side of the spars by turning the knob counterclockwise (Figure 46).

**WARNING:** Do not use excessive force when disconnecting the Spars. If after loosening the Spar Lock, the Spar does not disconnect with hand force, seek assistance from qualified service personnel. Failure to do so may cause harm to the healthcare professional or device.
ii. Simultaneously pressing down on the black tabs on each side of the table to disconnect the spars from their locked position (Figure 47) and raise them out of the latches.

![Figure 47: Pressing down on the black tab to disconnect the Spar](image)

iii. Remove the spars by lifting up and shifting the pair of spars to either side such that they can drop down for storage (Figure 48).

![Figure 48: Disconnecting the Spars](image)
iv. Move spars to the side such that one is positioned on the outside of the Head-End column while the other is positioned between the posts of the cervical receptacle (Figure 49).

Figure 49: Spars in process of being lowered to their storage position

**WARNING:** Take note of the pinch points at the spar hinges. Ensure no items are in the path of the spar hinges and healthcare professionals position themselves away from the linkage arms when moving the table to a storage position. Failure to do so may cause injury to the healthcare professional or possible damage to the table.
Figure 50: Spars Positioned Down for Storage

e. STEP 5: Install the protective Spar Cover on each Spar (Figure 51).

Figure 51: Spar Covers Being Installed
STEP 6: Loosen the Beam Lock (Figure 53).

Using the IntelliPendant®, continue to the next step, which allows the table to configure the Floor Locks such that those at the Head-End are unlocked while those at the Foot-End remain locked.
h. **STEP 7:** Retract the table (Figure 54).

![Figure 54: Collapsing the ProAxis® Table for Storage](image)

i. **STEP 8:** Secure the Beam Lock.

j. Using the IntelliPendant® or the Auxiliary Panel, unlock the Floor Locks. With the assistance of two people, the ProAxis® table may now be moved to its storage location.
To expand the table from its retracted position:

1. With the assistance of two people, move the table to the desired location.
2. Press the Power Switch on the Head-End base, and observe that a green ring illuminates around the switch, confirming power is applied to the table.
3. The IntelliPendant® will now provide step-by-step instructions on how to configure the table to its full position.
4. Complete the following steps in order as directed on the Hand Pendant:
   a. Loosen the Beam Lock.
   b. Continue to the next step, which allows the table to configure the Floor Locks such that those at the Head-End are unlocked while those at the Foot-End remain locked.
   c. Expand the table.
   d. Tighten the Beam Lock.
   e. Remove the protective Spar Cover from each Spar.
   f. Attach the Spars by:
      i. Raising the spars such that one is positioned on the outside of the Head-End column while the other is positioned between the posts of the cervical receptacle (Figure 49, page 42).
      ii. Lifting the spars above the latch position.
      iii. Lowering the spars into the respective latches.
      iv. Tightening the Spar Locks to secure the spars in place.
5. Using the IntelliPendant®, lock the Floor Locks. Complete the Function Check (see Section 6).

6. Confirm Storage LED is no longer illuminated on the Auxiliary Control Panel.

NOTE: Upon expanding the table, the table will still be in the hinged state required for storage. To level the table top, install the Head-End Supine Top or the two rail mounts to activate the table’s hinge functionality. Then, press either the Hinge Down button or the Return to Level button until the Hinge Angle reads 0°.

4.7.6 Version Info

To access information on the table’s software version:

1. Press the Menu Soft Key on the IntelliPendant®.

2. Utilize the Down arrow Soft Key to select Version Info (Figure 56). Press the Soft Key that corresponds to the check mark.

3. Information on the table’s current software version will be displayed.

NOTE: Software version information will be primarily used by a service technician should it need to be accessed.

![Figure 56: Version Info Menu Selection](image)

4.8 Independent Monitoring System (IMS)

The ProAxis® Spinal Surgery Table includes an Independent Monitoring System that monitors the load on the table and notifies the user in the event the table is overloaded. The system works by monitoring the load on the Head-End plate to which the spars are attached. If the load exceeds the allowable amount for the maximum-weight patient, the system will notify the user through the use of an error message and table motion will be interrupted.

The table can be overloaded for three (3) reasons, any of which will be detected by the IMS and prevent motion of the table. No other conditions are monitored that will trigger the alarm system.

1. A patient whose weight is over the maximum weight limit is placed on the table.

2. Extreme external forces.

3. A loss of coordination between the Head-End and Foot-End, which can place additional load or tension on the table.
Should the table be overloaded due to any of these three conditions, the overload notification of the IMS will be triggered and the user will be notified via the Overload screen (Figure 57). In this state, the user has the opportunity to reverse the table’s overloaded state by removing the load.

![Image](image_url)

**Figure 57: IntelliPendant® Display when the Overload Notification of the IMS is Triggered**

If the load on the table increases beyond the overload notification level, the IMS will trigger an overload alarm to notify the user and prevent any further motion of the table (Figure 58).

![Image](image_url)

**Figure 58: IntelliPendant® Display when the IMS Overload Alarm is Triggered**

For the user, there is no detectable delay from the time the alarm system detects the table is overloaded to when a warning is displayed on the IntelliPendant®.

The specific load on the table that triggers the alarm system is preset at the factory and is not adjustable. The functionality of the alarm system should be verified as part of the periodic maintenance of the table.
5 Inspection

5.1 Acceptance and Transfer

1. Upon receipt of your ProAxis® Spinal Surgery Table, remove it from the shipping crate. Remove any protective wrapping or packaging. Visually inspect all surfaces for freight damage.

NOTE: Any freight damage must be reported to the freight carrier immediately upon delivery. It is the responsibility of the recipient to make freight damage claims.

2. Read the model number, serial number and confirm the power requirements on the manufacturer’s label located on the Foot-End column of the table.

3. Perform Function Check (refer to Section 6).

5.2 Pre-Procedure/Post-Procedure

Before and after each use of the ProAxis® Spinal Surgery Table, visually inspect all accessible areas, electrical cords, the table and all movable parts and components for possible damage that may adversely affect the proper operation of the table. Verify Beam Lock and Silver Spar Knobs are tightened. Examine the patient safety straps for any sign of wear or damage. Examine the covers of all pads for tears or other damage that might cause the pads to trap fluids or other contaminants. Damaged or defective products should not be used or processed. Contact Mizuho OSI Services for repair or replacement (refer to Section 13).

5.3 Semi-Annual Preventative Maintenance

A Semi-Annual Preventative Maintenance (PM) check (refer to Section 10.3) on your ProAxis® Spinal Surgery Table is required at least once every six (6) months.

To obtain the PM checklist call Mizuho OSI Services at 1-800-777-4674 within the USA or +1-510-429-1500 outside the USA.

5.4 Product Lifetime

At the time of delivery, your product fulfills existing regulations and standards; however, in spite of proper use, routine inspection, prescribed service, maintenance and repairs, the product is subject to aging and wear. Therefore, Mizuho OSI cannot guarantee the product’s safety after ten (10) years and recommends your product be taken out of service. For product warranty information, refer to Section 13.6 of this manual.
6 Function Check

**NOTE:** Perform all steps of the Function Check before every use of the ProAxis® Surgery Table.

For a complete definition of terms used in this procedure, please refer to the Glossary of Terms in Section 2.9.

1. Confirm the Beam Lock is tightened.
2. Confirm the silver Spar Locks at the Head-End of the table are tightened, securing the spars in place.
3. Connect the IntelliPendant® cable to the IntelliPendant® port on the Head-End column (Figure 5, page 14).
4. Plug the Power Cord into a properly grounded receptacle to use with AC power. Refer to the manufacturer’s label on the Foot-End column for input voltage requirements (Figure 10, page 18).
5. Press the Power Switch on the Head-End base, and observe that a green ring illuminates around the switch, confirming power is applied to the table.
6. If the AC power cable is not connected to an outlet with a protective earth ground, then the external ground stud should be connected to a protective earth ground (Figure 4, page 13).
7. If the table needs to be used under battery power due to AC outage, ensure that the battery has been properly charged as outlined in Section 12.3. Press the Power Switch on the Head-End base, and observe that a green ring illuminates around the switch confirming power is applied to the table. The battery status symbol on the IntelliPendant® should be visible indicating a charged battery. If the battery status symbol on the IntelliPendant® is the battery is not adequately charged and the table should only be used with AC power. To protect the patient, hospital staff, and the table from possible electrical hazards, an external ground cable connection is required between the external ground stud and protective earth ground when the table is in use under battery power (Figure 4, page 13).
8. Lockout mode: The ProAxis® Surgery Table motions are only operational if the Floor Locks are locked. The green indicator located on the Auxiliary Control Panel is illuminated and the Floor Lock symbol is visible on the IntelliPendant® display screen when the Floor Locks are completely engaged. Unlock the Floor Locks and verify that neither the IntelliPendant® nor the Auxiliary Control Panel functions operate except to activate the Floor Locks.
9. Complete a Floor Lock check using the Auxiliary Control Panel (Figure 15, page 21).
   a. Press the Floor Lock button and observe the LED indicator is illuminated on the Auxiliary Panel and the Floor Lock symbol is visible on the IntelliPendant® display. The table is locked if it cannot be moved on its wheels.
   b. Press the Floor Unlock button and observe the LED is no longer illuminated on the Auxiliary Panel and the Floor Unlock symbol is visible on the IntelliPendant® display. The table is unlocked if it can move on its wheels.
c. Floor Lock Override Check:
   i. With the Floor Locks unlocked, utilize the IntelliPendant® and select the **Override Floor Locks** setting.
   ii. With the Override function active, utilize the Height Up, Height Down, Lateral Tilt Right and Left, or Trendelenburg/Reverse Trendelenburg buttons. These buttons should function without the Floor Locks being locked while the Override function is utilized.

10. **IntelliPendant® Check:**
   a. Utilizing the IntelliPendant®, press the **Floor Lock** button. Observe on the Auxiliary Control Panel that the LED illuminates and that the Floor Lock symbol displays on the IntelliPendant® display after the floor locks are completely deployed and locked. Look under the Head-End and Foot-End base, and observe that all four feet have moved down and are in direct contact with the floor (Figure 16, page 21).
   b. Press the **Floor Unlock** button. Observe that all four locking feet move up. Confirm the LED is not illuminated on the Auxiliary Control Panel and the Floor Unlock symbol displays on the IntelliPendant® display after the floor locks are completely retracted and unlocked. Look under the Head-End and Foot-End base and observe that all four feet have moved up and are not contacting the floor (Figure 16, page 21).
   c. Repeat Step (a) to ensure the Floor Locks are fully engaged such that the rest of the table’s functionality may be tested.
   d. Press and hold the **Height Up** button. Observe that the table top raises through its entire range. Observe the IntelliPendant® display, and confirm that the Height Up symbol is visible in the action display area of the screen. The number beside the Height Up symbol in the motion status area of the screen should be changing, corresponding to where the table top is within the height range.
   e. Press and hold the **Height Down** button. Observe that the table top lowers through its entire range. Observe the IntelliPendant® display, and confirm that the Height Down symbol is visible in the action display area of the screen. The number beside the Height Down symbol in the motion status area of the screen should be changing, corresponding to where the table top is within the height range.
   f. Press and hold the **Reverse Trendelenburg** button. Observe that the Foot-End of the table top moves down through its entire range. Observe the IntelliPendant® display, and confirm the Reverse Trendelenburg symbol is visible in the action display area of the screen. The number beside the Reverse Trendelenburg symbol in the motion status area of the screen should be changing, corresponding to where the table top is within the Reverse Trendelenburg range.
   g. Press and hold the **Trendelenburg** button. Observe that the Head-End of the table top moves down through its entire range. Observe the IntelliPendant® display, and confirm that the Trendelenburg symbol is visible in the action display area of the screen. The number beside the Trendelenburg symbol in the motion status area of the screen should be changing, corresponding to where the table top is within the Trendelenburg range.
   h. Press and hold the **Lateral Tilt, Left** button. Observe that the table top tilts to the left through its entire range. Observe the IntelliPendant® display, and confirm that the Lateral Tilt, Left symbol is visible in the action display area of the screen. The number beside the Lateral Tilt, Left symbol in the motion status area of the screen should be changing, corresponding to where the table top is within the Lateral Tilt range.
i. Press and hold the Lateral Tilt, Right button. Observe that the table top tilts to the right through its entire range. Observe the IntelliPendant® display, and confirm that the Lateral Tilt, Right symbol is visible in the action display area of the screen. The number beside the Lateral Tilt, Right symbol in the motion status area of the screen should be changing, corresponding to where the table top is within the Lateral Tilt range.

j. Install the Two-Piece Supine Top on the frame (refer to Section 8) and test the table’s hinge functionality:

1. Select the FSS Hinge Mode Setting using the IntelliPendant®.

2. Press and hold the Hinge Up button. Observe that the table top hinge angle increases through its entire range. Observe the IntelliPendant® display, and confirm that the FSS Hinge Up symbol is visible on the screen. The number beside the Hinge Up symbol in the motion status area of the screen should be changing, corresponding to where the hinge is within the range of travel.

3. Press and hold the Hinge Down button. Observe that the table top hinge angle decreases through its entire range. Observe the IntelliPendant® display, and confirm that the FSS Hinge Down symbol is visible on the screen. The number beside the Hinge Down symbol in the motion status area of the screen should be changing, corresponding to where the hinge is within the range of travel.

4. Select the FE Hinge Mode Setting using the IntelliPendant®.

5. Press and hold the Hinge Up button. Observe that the table top hinge raises through its entire range. Observe the IntelliPendant® display, and confirm the FE Hinge Up symbol is visible on the screen. The number beside the Hinge Up symbol in the motion status area of the screen should be changing, corresponding to where the hinge is within the range of travel.

6. Press and hold the Hinge Down button. Observe that the table top hinge lowers through its entire range. Observe the IntelliPendant® display, and confirm that the FE Hinge Down symbol is visible on the screen. The number beside the Hinge Down symbol in the motion status area of the screen should be changing, corresponding to where the hinge is within the range of travel.

7. Return to the FSS Hinge Mode Setting using the IntelliPendant® if desired.

k. With the Two-Piece Supine Top installed on the frame, re-position the table top by raising the Height to 36 inches (91 cm); laterally Tilt the table top left or right six (6) degrees; raise or lower the Hinge 10 degrees; and place the table top in five (5) degrees of Trendelenburg. Confirm this position by reading the display on the IntelliPendant® screen. Press and hold the Return to Level button. Observe that the table top levels side to side and head to foot. Continue to depress the button, and after a one-second delay, the table top will move down to its Home height. Observe that the table top is level and at the Home height position of 32 inches (81 cm) off the floor when movement stops. Observe the IntelliPendant® display, and confirm the Height display reads 32 inches (81 cm), the Hinge angle reads 0 degrees, the Tilt angle reads 0 degrees, and the Trendelenburg/Reverse Trendelenburg angle reads 0 degrees.

l. Using the Menu key, navigate to the ACP setting and select the Fast setting. Using the ACP Soft Key on the Main Screen, activate the ACP function. Observe the Fast ACP symbol is present on the IntelliPendant® display screen. Repeat the process, this time selecting the Slow setting. Observe that the Slow ACP symbol is now present on the IntelliPendant® display screen.
11. Battery operation check:
   a. Observe the Battery Status on the base of the Head-End column. If the LED is illuminated green - OK, then the battery is properly charged. Confirm that the Battery Status icon on the IntelliPendant® is \[\text{\textbullet}\] indicating the battery is fully charged and the table is ready to operate on battery power.
   
   b. If the LED is illuminated red or the Battery Status icon on the IntelliPendant® is \[\text{\textbullet}\], the battery must be charged prior to using the table. To charge the battery, ensure the Power Cord is plugged into a functioning receptacle. The AC LED at the base of the Head-End column will illuminate green indicating that appropriate AC power is applied to the table. The table must remain plugged in for a minimum of three (3) hours to ensure sufficient charging of the battery to operate the table.
   
   c. If the LED remains red and the Battery Status icon is still \[\text{\textbullet}\] after three (3) hours, continue to charge the battery for up to 24 hours.
   
   d. If the LED does not illuminate green-OK and the Battery Status icon does not reflect \[\text{\textbullet}\] after 24 hours, refer to Section 12.3 for servicing instructions.

**NOTE:** The table may be used with AC power even when the Battery Status icon is \[\text{\textbullet}\] indicating the battery needs charging. Confirm the green AC power LED at the base of the Head-End column is illuminated.
7 ProAxis® Spinal Surgery Table
Standard Components

7.1 ProAxis® Standard Components

In addition to the IntelliPendant® and the Power Cord, the following standard patient positioning components are shipped with the ProAxis® Spinal Surgery Table. Pictures of the individual components and instructions for use are provided in Section 8.

NOTE: Use proper lifting and carrying techniques when moving components and accessories due to their weight and size.

<p>| REF 6988-899 | Rail Mount, Left | The Rail Mount, Left slides on the left side of the Head-End hinged, open frame and works in conjunction with the Chest Tray to translate the patient’s upper torso when the hinge of the frame is raised or lowered. |
| REF 6988-887 | Rail Mount, Right | The Rail Mount, Right slides on the right side of the Head-End hinged, open frame and works in conjunction with the Chest Tray to translate the patient’s upper torso when the hinge of the frame is raised or lowered. |
| REF 6988-950 | Chest Tray | The Chest Tray sits on the Rail Mounts and provides the platform to which to attach one of the three available Chest Pads used to support the patient. The Chest Tray has a numeric scale to allow for ease in patient positioning. |
| REF 6988-530 | ProAxis® 5-Way Articulating Arm Board (2) - includes REF 6988-526 Arm Board Bracket (2) | The 5-Way Articulating Arm Boards provide support for the patient’s arms when positioned prone on the ProAxis®. When in place, the Arm Board can be translated, articulated, raised, lowered, and laterally adjusted depending on the positioning needs of the patient. The Ball Joint controls the articulating motion of the Arm Board, which may be required for positioning a patient with decreased shoulder or arm range of motion. The 1-inch (2.5 cm) pad provided is of Mizuho OSI Tempur-Pedic® medical construction. The bracket required to attach the Arm Board on the Chest Tray is provided with the Arm Board assembly. |
| REF 6988-758 | ProAxis® Chest Pad | The ProAxis® Chest Pad is of Mizuho OSI Tempur-Pedic® medical construction. When properly positioned, the top edge of the pad should rest at the bottom of the patient’s supra-ternal notch. |
| REF D28502CE | Re-usable ProneView® Adjustable Protective Helmet | The re-usable ProneView® Adjustable Protective Helmet is part of the ProneView® system and supports the one-time use disposable cushion insert provided in the Patient Care Kit. The ProneView® Adjustable Protective Helmet has four posts with 2.2 inches (5.6 cm) of height adjustment. |
| REF D28580CE | Re-usable ProneView® Mirror Platform with Post Holes | The re-usable Mirror Platform with Post Holes provides support for the ProneView® Adjustable Protective Helmet and allows the patient’s eyes to be viewed from three sides when positioned prone. The ProneView® Adjustable Protective Helmet posts seat in the post holes of the mirror. |
| REF 6988-969 | Cervical Chest Tray | The Cervical Chest Tray sits directly on the rails of the open frame and the ProAxis® Chest Pad attaches to the tray to support the patient. The Chest Tray has Velcro® straps for securing it to the frame. |
| REF 6988-725 | Cervical Traction Vector Adjustor | The Cervical Traction Vector Adjustor mounts on the Head-End of the table and maintains alignment of the traction rope when cervical traction is utilized. |</p>
<table>
<thead>
<tr>
<th>REF</th>
<th>Description</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>6988-209</td>
<td>Cervical Management Mounting Bracket</td>
<td>The Cervical Management Mounting Bracket installs in the Head Board and allows the Mizuho OSI optional accessory, the Cervical Management Base Unit, to be utilized with an aluminum or radiolucent DORO® or Mayfield® skull clamp.</td>
</tr>
<tr>
<td>6988-387</td>
<td>Contoured ACP Hip Pad, Left</td>
<td>The Contoured ACP Hip Pad, Left is of Mizuho OSI Tempur-Pedic® medical construction and provides support for the patient when positioned prone on the open frame. When positioned correctly, the patient’s left iliac crest should be positioned in the top third of the pad labeled Left.</td>
</tr>
<tr>
<td>6988-388</td>
<td>Contoured ACP Hip Pad, Right</td>
<td>The Contoured ACP Hip Pad, Right is of Mizuho OSI Tempur-Pedic® medical construction and provides support for the patient when positioned prone on the open frame. When positioned correctly, the patient’s right iliac crest should be positioned in the top third of the pad labeled Right.</td>
</tr>
<tr>
<td>6977-739</td>
<td>Buttocks Strap</td>
<td>The Buttocks Strap rests across the patient’s lower gluteal area when positioned prone to secure the patient in place on the Contoured ACP Hip Pads and open frame.</td>
</tr>
<tr>
<td>6844-148</td>
<td>Leg Board with Pad (2)</td>
<td>The two (2) Leg Boards, when used together, provide support for the patient’s legs when positioned prone on the open frame. The 1-inch (2.5 cm) pad is made of Mizuho OSI Tempur-Pedic® medical construction.</td>
</tr>
<tr>
<td>5840-450</td>
<td>Leg Sling</td>
<td>The Leg Sling is placed over the Foot-End section of the hinged, open frame and is secured in place with Velcro® straps. Positioning pillows should be utilized in the sling to achieve the desired positioning and support of the patient’s legs when positioned prone.</td>
</tr>
<tr>
<td>6988-880</td>
<td>Retractor Adaptor</td>
<td>The Retractor Adaptor provides six (6) inches (15 cm) of side rail and is required for mounting ancillary equipment onto the carbon fiber open frame.</td>
</tr>
<tr>
<td>6946</td>
<td>Tempur-Pedic® Medical Pillow, Standard Size</td>
<td>A standard size pillow of Tempur-Pedic® medical construction is provided for patient positioning. A cleanable, zippered cover encases the pillow.</td>
</tr>
<tr>
<td>6947</td>
<td>Tempur-Pedic® Medical Pillow, Queen Size (2)</td>
<td>A queen size pillow of Tempur-Pedic® medical construction is provided for patient positioning. A cleanable, zippered cover encases the pillow.</td>
</tr>
<tr>
<td>6988-623</td>
<td>Supine Top, Head-End Assembly</td>
<td>The Supine Top, Head-End is of carbon fiber construction and mounts on the Head-End section of the hinged, open frame. It is secured to the frame with locking brackets and accommodates supine and lateral patient positioning.</td>
</tr>
<tr>
<td>6988-621</td>
<td>Supine Top, Foot-End Assembly</td>
<td>The Supine Top, Foot-End is of carbon fiber construction and mounts on the Foot-End section of the hinged, open frame. It is secured to the frame with locking brackets and accommodates supine and lateral patient positioning.</td>
</tr>
<tr>
<td>6988-442</td>
<td>Tempur-Pedic® Medical 2-inch (5 cm) Supine Top Pad (one piece)</td>
<td>The one piece Tempur-Pedic® Medical 2-inch (5 cm) Pad mounts on the Supine Tops. The pad is of viscoelastic foam construction and has a built-in translation slide on the bottom of the pad that interfaces with the surface of the Supine Top, Foot-End, allowing for support and movement when the hinge is raised or lowered.</td>
</tr>
<tr>
<td>6977-959</td>
<td>Universal Side Rail Adaptor (4)</td>
<td>Utilized on the Supine Top, the Universal Side Rail Adaptor is designed to protect the carbon fiber and provide 6-in. (15 cm) of side rail for mounting ancillary equipment.</td>
</tr>
<tr>
<td>5840-43</td>
<td>Safety Strap, 60 inches (152 cm) (2)</td>
<td>The Safety Straps are provided to secure the patient when positioned prone on the ProAxis®.</td>
</tr>
</tbody>
</table>
Safety Strap for Supine Top (2)

The Safety Strap slides on the carbon fiber rails of the Supine Top and is provided to secure the patient when positioned supine or lateral on the ProAxis®.

ProAxis® Spinal Surgery Table ProneView® Patient Care Kit (3)

The Patient Care Kits are single-use pad covers intended to prevent cross-contamination and protect the equipment. Three (3) ProneView® Patient Care Kits are included with the ProAxis® table.

ProAxis® Component Cart

The ProAxis® Component Cart is provided to safely store all the standard components of the table. It is also designed to store optional accessories which may be acquired for use with the ProAxis®. A cover is provided with the cart and serves as a dust cover when the cart is not in use.

ProAxis® Table Cover

The ProAxis® Table Cover is provided to serve as a dust cover when the table is not in use. The cover fits the table when set up (fully extended) or it can be adjusted with the Velcro® closures to cover the table when it is configured for transport or storage.

Spar Cover (2)

The Spar Covers are provided to serve as a protective cover for the spars when the table is not in use and configured for transport or storage.

7.2 ProAxis® Patient Care Kits

The ProAxis® Surgery Table ships with three (3) individual ProAxis® ProneView® Patient Care Kits. A case of six (6) individual kits is provided when ordering REF 6988A-PV-ACP.

Each individual ProAxis® ProneView® Patient Care Kit includes the following:

- ProneView® Cushion Insert
- Chest Pad Cover for ProAxis® Chest Pad
- Pair of Hip Pad Covers with Aprons
- Pair of Foam Arm Cradles
- 1 Set of Single-Use, Disposable Advanced Control Pad System™ Tubing

**WARNING:** Patient Care Kits are single use-only. Discard after use. Re-use of any component in the Patient Care Kit may result in cross contamination.

The components should be used as described in the Patient Care Kit instructions included in each kit. The ProneView® Patient Care Kit works with the provided ProneView® Adjustable Protective Helmet System. The system is designed by the manufacturer to support the patient’s head while in a prone position. Additionally, the position of the patient’s facial alignment in the ProneView® Adjustable Protective Helmet can be readily viewed while in the prone position.

**NOTE:** Remove the Patient Care Kit components from the packaging and visually inspect the components. The ProneView® Cushion Insert should be opened sixty (60) minutes prior to use and the Arm Cradles should be opened twenty (20) minutes prior to use for complete expansion of the foam.

**NOTE:** The Mirror Platform with Post Holes is a rectangular shaped mirror with four holes patterned to accept the adjustable posts on the helmet. The mirror can be seated on the Face Plate with the longer side oriented either parallel or perpendicular to the overall length of the table.
8 Setup of ProAxis® Spinal Surgery Table for Surgical Procedures

NOTE: This device is intended for use by trained personnel. Prior to setup and use of the ProAxis® Spinal Surgery Table, ensure personnel have been trained. To schedule an in-service, please contact your domestic Mizuho OSI sales representative or call 1-800-777-4674 within the USA or +1-510-429-1500 outside the USA.

To prepare the ProAxis® Spinal Surgery Table for patient positioning, complete the following steps:

1. Connect the IntelliPendant® to the port on the Head-End column (Figure 19, page 24).
2. Press the Power Switch and observe that the switch illuminates with a green ring indicating that power is applied to the table.
3. Utilizing the IntelliPendant®, press the Floor Unlock button. Observe that the Floor Locks are completely retracted and the Floor Unlock symbol is visible on the display of the hand pendant (Figure 21, page 25).
4. With the assistance of a second person, move the table to where it will be used. One person should be located at the Head-End and the other person at the Foot-End of the table. Care should be taken to control the table when rolling.

WARNING: If the ProAxis® Spinal Surgery Table is allowed to roll too fast, it may be difficult to stop or turn. Impact of the table top with a stationary object may cause serious damage to the table top. If an impact does occur, the table must be visually inspected for damage and a Function Check must be performed (refer to Section 6). If damage is discovered or the table does not successfully complete the Function Check, call Mizuho OSI Services (refer to Section 13).

5. Orient the Head-End of the table in the room where the anesthesiologist work station is located.
6. Plug the Power Cord into a properly grounded receptacle. Refer to the manufacturer’s label on the Foot-End column for input voltage requirements (Figure 10, page 18). If the AC power cable is not connected to an outlet with a protective earth ground, then the external ground stud should be connected to a protective earth ground (Figure 10, page 18).
7. Using the IntelliPendant®, press the Floor Lock button. Observe that the Floor Lock symbol is visible on the hand pendant display (Figure 22, page 25).
8. Complete a Function Check with the IntelliPendant® (refer to Section 6).

NOTE: Perform all steps of the Function Check before every use of the ProAxis® Spinal Surgery Table.

NOTE: The ProAxis® Spinal Surgery Table is now ready for completing the setup of the patient components prior to patient transfer. Components have been designed to specifically mount on the carbon-fiber, trapezoidal frame or the Supine Top, while safely supporting the patient and not damaging the frame.
Figure 59: ProAxis® Spinal Surgery Table Open Frame Ready for Setup
8.1 ProAxis® Prone Patient Positioning

**NOTE:** Only Mizuho OSI components and accessories have been tested and approved for use with the ProAxis® Spinal Surgery Table. Other manufacturers’ products have not been tested for proper performance when used with the ProAxis® Spinal Surgery Table, and therefore are not endorsed for use by Mizuho OSI. Use of other manufacturers’ products may void the warranty.

**WARNING:** Use of components not manufactured by Mizuho OSI may result in harm to the patient, the table top, the device, or the healthcare professional.

![Figure 60: ProAxis® Prone Setup – Fixed Surgical Site Hinge Mode](image)

The following instructions provide the recommended steps for setup. Final determination of patient positioning and setup is at the discretion of the surgeon and surgical team.

1. Confirm the ProAxis® Spinal Surgery Table is prepared for setup (refer to Section 8). When the patient is positioned prone, their head is oriented to the Head-End of the table (Figure 1, page 6).

2. Confirm that the Floor Locks are locked. Ensure the Floor Lock symbol is visible on the IntelliPendant® display.

3. Open the following: the Patient Care Kit, the ProneView® Cushion Insert or the GentleTouch® Face Pillow, and the package of Arm Cradles and set aside.

**NOTE:** The surgeon may decide to use the ProneView® Cushion or the GentleTouch® Face Pillow depending on which option allows for the achievement of proper cervical spine position.
4. Slide the Rail Mount, Right and Rail Mount, Left onto the corresponding Head-End sections of the Hinged Frame. The nipple will engage the bracket, locking the Rail Mount in place. To confirm, pull back on the Rail Mount; it should remain secure in the Hinged Frame bracket (Figure 61).

![Figure 61: Installing the Rail Mount](image)

5. Connect the cable connector extending from the Rail Mount, Right/Left to the port on the corresponding side of the Head-End Assembly. Align the pins of the cable connector with the port, insert and turn the locking collar to the right securing the pins in the port (Figure 62).

**NOTE:** To remove the Rail Mount, turn the locking collar to the left and pull the cable connector out of the port, disconnecting it from the Head-End Assembly. With the cable connector disconnected, depress the Black Tab on the frame while sliding the Rail Mount towards the hinge.

**NOTE:** When both rail mounts are connected, the Torso Trolley® drive pins may reposition to align properly.
6. The Torso Trolley® consists of a Chest Tray to which the Flat ACP or non-ACP Chest Pad or optional Bilateral Winged ACP Chest Pad and 5-way Adjustable Arm Boards are mounted (Figure 69, page 66).
   a. Prior to installing the Chest Tray, ensure the Rail Mounts are correctly installed and the Hinged Frame is in the neutral position.
b. Confirm the display on the IntelliPendant® shows the Hinge, Trendelenburg, and Lateral Tilt angles are all 0° (Figure 22, page 25). If necessary, utilize the IntelliPendant® and press the Return to Level button to return the Hinged Frame to the neutral position.

![Image of ProAxis Spinal Surgery Table with yellow strip highlighted]

**Figure 64: Left and Right Rail Mounts Attached**

7. To install the Chest Tray:
   a. Orient the Chest Tray over the posts of the Rail Mounts and lower into place. The Chest Slide can be placed anywhere on the Rail Mounts that allows the posts to sit between the teeth of the Chest Slide (Figure 65).
   b. Confirm the Chest Tray is not placed on the rails at a point lower than Position 1 on the Rail Mounts in order to ensure the Chest Tray is properly engaged.

**NOTE:** When mounting the Chest Tray onto the rails, ensure the yellow strip on each of the Rail Mounts is not visible. Appearance of the yellow strips indicates that the Chest Tray is not properly engaged with Rail Mounts.

**WARNING:** Visibility of the yellow strips on the Rail Mounts indicate that Chest Tray is not properly engaged with the Rail Mounts, which will prevent it from sliding correctly during hinge motion. Failure to ensure the Chest Tray is correctly mounted may result in harm to the patient or the device.

   c. The Chest Tray may be moved up or down the length of the Rail Mounts by simultaneously squeezing the grips of the Chest Tray and moving it to the desired location. Release both grips to lock the Chest Tray into place.
8. To attach the ProAxis® Chest Pad, orient the opening of the pad bracket so it surrounds the lower surface of the Chest Tray. Slide onto the Chest Tray until the tab of the bracket is secure on the tray. The indentation in the pad for the supra-sternal notch should be oriented towards the Head-End of the table (Figure 66). Confirm the tab of the bracket is secure on the tray. To remove, press the tab of the bracket and slide the pad towards the Foot-End of the table.
NOTE: Also at the discretion of the physician, additional foam or padding can be placed between the patient and the pad as needed to accommodate varying sizes of patients.

NOTE: With ideal positioning, the Chest Pad should support the patient such that the load is borne predominantly by the sternum. When the patient is positioned correctly, the top edge of the pad should rest below the patient’s anatomical landmark, the supra-ternal notch.

WARNING: Positioning the Chest Pad superior to the supra-ternal notch may apply pressure to the throat and the patient’s airway. Failure to ensure the patient is positioned correctly on the Chest Pad may result in harm to the patient or the device.

9. Install the 5-Way Adjustable Arm Boards.
   a. Slide the Arm Board Mounting Brackets into each opening of the bracket on the underside of the Chest Slide (Figure 67). Advance the brackets fully and tighten the T-Handles.
   b. Attach the Articulating Arm Board to the non-patient transfer side of the table. Insert the arm board assembly into the Arm Board Bracket and tighten the T-Handle on the Arm Board Mounting Bracket.
   c. Attach the Articulating Arm Board to the patient transfer side of the table. Insert the arm board assembly into the Arm Board Bracket and tighten the T-Handle on the Arm Board Mounting Bracket.
   d. Adjust the arm boards as needed in preparation for patient transfer (Figure 68). The Height, Translation, and Angle of the arm board surface can be adjusted. To adjust the Height, loosen the Black Knob and raise or lower the assembly, or raise or lower the assembly on the mounting bracket. Tighten the Black Knob and or the T-Handle when the desired Height is achieved. To translate the arm board towards the Head-End or Foot-End of the table, pull the pin on the underside of the arm board, translate and then re-position and release the pin to lock in place. To articulate the arm board, loosen the drop handle, pivot in the ball joint to the desired position, and tighten T-Handle.

CAUTION: Use care when positioning the arm board to ensure that it does not come in contact with the floor or surrounding equipment positioned near the table. Failure to allow for proper clearance of the arm board with its surroundings may result in damage to the arm board or other equipment.

   e. Pivot the Articulating Arm Board on the patient transfer side out of the way by turning the Black Knob and moving the arm board. Secure in place by re-tightening the Black Knob (Figure 68).
T-Handle Allows Lateral Adjustment of Arm Board in Place

T-Handle Allows Height Adjustment of Arm Board

Bracket on the bottom of the Chest Tray includes T-Handles to secure the Arm Board Mounting Bracket.

Figure 67: Arm Board Mounting Bracket Installed

Pull knob to Release and Translate Arm Board North/South.

Pivot & Lateral Adjustment

Articulating Ball Joint

Height Adjustments

Figure 68: 5-Way Adjustable Arm Board Installed
10. Confirm the Torso Trolley® slides properly. Simultaneously squeeze the grips of the Chest Tray (Figure 70). This releases the post in the Rail Mount from the teeth of the Chest Tray and allows for the Chest Tray with Arm Boards to be moved easily toward the Head-End or Foot-End of the table. The 1-8 scale on the Chest Tray assists in identifying where the Chest Tray is in its range of movement.
11. Attach the Contoured ACP Hip Pads. The pads are labeled right and left specific to be oriented to the patient’s iliac crest when lying prone. Open the Locking Bracket, orient the Pad with Locking Bracket over the rail of the open frame on the Foot-End section of the frame, lower into place and slide towards the Hinge until it rests against the Pad Stop (Figure 71).

![Figure 71: Mounting the Contoured ACP Hip Pad on the Frame](image)

12. Press up on the white tab until the black component of the clamp fits into the open notch of the white tab. Press down on the lip of the black component of the clamp to lock the pad in place (Figure 72).

![Figure 72: Locking the Contoured ACP Hip Pad Bracket](image)

13. When installed correctly, the pad should rest against the Pad Stop and the arrow on the side of the Bracket should be pointed toward the Foot-End of the table (Figure 73). To remove the Contoured Hip Pad, lift up on the lip of the Bracket so that it disconnects, and lift the pad off the rail of the frame.
NOTE: Depending on desired area of flexion and patient’s anatomy, the hip pads may be moved towards the Foot-End of the table at the surgeon’s discretion to achieve proper positioning.

Figure 73: Contoured ACP Hip Pad Locked in Place

14. To install the Adjustable Leg Sling, place the sling over the Foot-End section of the open frame (Figure 74). Feed the four straps through their respective buckles on the opposite side of the sling and secure. The sling should be adjusted for each patient. The straps on the sling consist of a double-sided hook and loop closure that allows for infinite adjustment along the length of the strap. The hook end of the strap may be folded over onto the loop portion of the strap to allow for greater adjustability. The strap may be folded over itself again to shorten, thereby raising the sling and decreasing the degree to which the patient’s hips are flexed. If the double-sided hook and loop closure is worn from repetitive use or misuse, replacement of the sling is required.

Figure 74: Sling Attached to ProAxis® Spinal Frame

15. Place two (2) or three (3) 6946 or 6947 Tempur-Pedic® Medical Pillows in the sling approximately where the patient’s lower legs will rest to achieve the desired amount of knee flexion and support (Figure 75).
16. The Leg Boards may be used in place of the Leg Sling to support the legs. Attach the Leg Boards with the 1-inch (2.5 cm) Tempur-Pedic® Medical Pad. Release the Velcro® Straps on the sides of the Leg Board. Place one Leg Board over the rails of the open frame and slide into place. Secure with the Velcro® Straps. Repeat this process with the second Leg Board. Ensure that the two Leg Boards abut each other (Figure 76).

17. If using the ProneView® Mirror and Helmet System, place the Mirror Platform with the ProneView® Adjustable Helmet on the Chest Tray (Figure 77). Ensure that the adjustable posts are in their lowest position in preparation for patient transfer. Fit the ProneView® Cushion Insert into the ProneView® Helmet and within the tabs provided at the chin end of the ProneView® Helmet. Adjust the Cushion Insert openings to be centered relative to the ProneView® Helmet openings.
18. If using the GentleTouch® Face Pillow, place the Pillow on the Chest Tray (Figure 78).

Figure 78: GentleTouch® Face Pillow Properly Positioned

NOTE: In some cases if use of tongs or a head halter is desired in conjunction with the use of the Torso Trolley®, an optional short Cervical Traction Vector (REF 6988-720) is available.
19. After the desired prone components are in place on the frame, the tubing can be connected to the Contoured ACP Hip Pads.
   a. Secure the ACP tubing with the Velcro® wraps along the frame and by routing it through the underside of the Leg Sling. Ensure the tubing is secure and out of the way such that the tubing will be able to move when the Hinged Frame is moved.
   b. Ensure the third port that does not connect to the Hip Pads is cinched to prevent air from escaping the system.
   c. Connect the tubing to the port at the Foot-End of the table.
   d. Utilize the IntelliPendant® and raise and lower the Hinge to the extent of its travel. Re-adjust the ACP tubing as may be required.

**NOTE:** Verify the ACP tubing will not be stretched or pinched during hinge movement, which may interrupt or obstruct air flow to the pads. Failure to do so may result in ACP malfunction.

20. Place the Patient Kit Covers on the Chest and Hip Pads (Figure 79).

![Figure 79: ProAxis® Spinal Surgery Table Prone Setup with Patient Care Kits Applied](image)

21. Trial position the imaging equipment if it will be used. If necessary, re-locate the ProAxis® Surgery Table in the room. Unlock the Floor Locks by using the IntelliPendant® and with the assistance of a second person move the ProAxis® Surgery Table. When the table is located where it will be used, lock the Floor Locks.

⚠️ **CAUTION:** The ProAxis® Surgery Table is not for patient transport. The location of the table relative to any imaging equipment that may be used needs to be confirmed prior to transferring the patient to the table.
22. Position the patient bed or stretcher so the patient’s anatomical landmarks are aligned with the support pads on the hinged frame of the ProAxis® Surgery Table. It may be necessary to re-position the Chest Pad by manually moving the Torso Trolley®. The top edge of the Chest Pad should align with the patient’s anatomical landmark, the bottom of supra-sternal notch, and the top third of the Contoured ACP Hip Pad should align with the patient’s anatomical landmark, the iliac crest.

23. The height of the patient bed or stretcher should be level with the top of the Contoured ACP Hip Pad. To assist the patient transfer process, you may choose to use the IntelliPendant® and laterally tilt the table toward the patient bed or stretcher by pressing the appropriate tilt button.

**NOTE:** *When the patient is correctly positioned prone on the ProAxis®, the hinge will flex or extend the patient’s spine at L3.*

24. Lock the patient stretcher or bed. Confirm by observing on the IntelliPendant® display that the table Floor Locks are locked. The Floor Locks Locked symbol should be visible (Figure 22, page 25).

25. Ensure that the ProneView® Adjustable Protective Helmet legs are set to the lowest position.

26. Place the ProneView® Adjustable Protective Helmet with Cushion Insert on the patient’s face. Ensure the patient’s eyes are visible. With the assistance of others, log-roll the patient onto the ProAxis® Spinal Surgery Table supporting the patient’s head, torso and legs.

27. After log-rolling the patient to a prone position and onto the Mirror Platform on the Chest Tray, verify there is adequate clearance between the patient’s nose and the mirror, and verify the endotracheal tube is not kinked.

⚠️ **WARNING:** Failure to confirm the positioning of the patient and patient tubes may cause harm to the patient.

28. Continue to support the patient on the table until the safety straps are in place. DO NOT leave the patient unattended on the table.

⚠️ **WARNING:** Patient must be restrained at all time by a safety strap(s) when positioned on the table.

29. Remove the patient bed or stretcher while supporting the patient on the table.

30. Install and secure the Articulating Arm Board on the patient transfer side of the table (Figure 68, page 65). Position the patient’s arms in the Arm Cradles and on the Arm Boards. Final positioning of the patient’s arms should be completed after confirming the patient’s orientation on the Chest Pad and the Contoured ACP Hip Pads.

31. Confirm that the patient’s face is seated in the Cushion Insert; the eyes should be visible in the mirror; the neck should be in a neutral position (Figure 80). Raise the height of the ProneView® Adjustable Protective Helmet until a neutral neck position is achieved by turning the posts of the helmet. There is a 2.3-inch (5.6 cm) height adjustment range available.
WARNING: Frequently monitor the patient’s neck, head, eyes, nose, and mouth to ensure that a safe position is maintained and the ET tube has not been kinked or displaced. Failure to do so can lead to serious adverse consequences including blindness and failure to ventilate.

WARNING: Failure to maintain neutral neck position can result in neck damage and/or chin abrasions. Some erythema may result from skin contact with the materials in this product or from compression during prolonged cases.

Figure 80: Patient Positioned Properly in ProneView® Helmet

32. Visually confirm the placement of the Contoured ACP Hip Pads. The pad should be mounted abutting the Pad Stop (Figure 71, page 68). When positioned correctly, the patient’s anatomical landmark, the iliac crest, should rest in the top one-third of the pad. The Patient Kit Hip Pad Cover should be smooth against the patient’s skin. Ensure that the patient’s skin is smooth and flat against the Patient Kit Hip Pad Cover. To re-position the patient on the Pad, the patient should be lifted and moved such that the iliac crest contacts the top third of the pad. Ensure the Hip Pad Cover is smoothed and lower the patient back onto the Pad. Confirm correct placement of the Pads such that the each of the patient’s iliac crests rest in the top one-third of each respective Hip Pad (Figure 81).

CAUTION: Failure to ensure that the patient is positioned correctly on the Contoured ACP Hip Pad, may result in harm to the patient or the device.
33. Visually confirm the placement of the Chest Pad. When positioned correctly, the edge of the Pad should rest below the patient’s anatomical landmark, the supra-sternal notch (Figure 82). Breasts should be down and lying flat with the nipples oriented towards the Foot-End of the table. The Patient Kit Chest Pad Cover should be smooth against the patient’s skin. Ensure that the patient’s skin is smooth and positioned flat against the cover. If necessary, re-position the Chest Pad. While lifting and supporting the patient, move the Torso Trolley® with the attached Chest Pad to the desired location and smooth the Chest Pad Cover. The patient should then be lowered back onto the Chest Pad (Figure 83).

**NOTE:** With ideal positioning, the Chest Pad should contact the patient such that the load is borne predominantly by the sternum.

**WARNING:** Positioning the Chest Pad superior to the supra-sternal notch may apply pressure to the throat and patient’s airway. Failure to ensure the patient is positioned correctly on the Chest Pad may result in harm to the patient or the device.
34. Visually confirm that the Leg Sling or Leg Boards provide support to the patient’s knees, with legs supported on pillows. If using the Leg Sling, the patient’s knees should be flexed and legs supported on the pillows. Re-position the legs and add pillows as required under the patient’s tibias to ensure that the ankles are not hyper-extended and there is no pressure on the patient’s feet (Figure 81, page 74).

⚠️ **CAUTION:** Failure to ensure that the patient is positioned correctly with the Leg Sling or Leg Boards may result in harm to the patient or the device.

⚠️ **CAUTION:** When the table is positioned at a low height, take care in positioning the patient in the Leg Sling such that the Sling does not come in contact with the center beam. Failure to properly monitor the position of the Leg Sling with the patient’s legs positioned in it may result in harm to the patient or the device.

35. Place the Buttocks Strap across the lower gluteal area of the patient. A blanket may be used under the Buttocks Strap to protect the patient from direct contact with the strap. The strap can be fed through the buckle on the sides of the Contoured ACP Hip Pads and then back on itself, securing the Velcro®. Confirm that the Buttocks Strap is positioned low on the gluteal area, and that it surrounds both Contoured ACP Hip Pads, and is secured snugly around the buttocks. Ensure that the patient does not directly contact the frame of the table. Use foam or gel padding as needed to pad the area between the patient and the rails of the frame.


**NOTE:** The placement and location of the Straps are at the discretion of the surgeon and vary by procedure.
NOTE: Due to the open nature of the ProAxis® frame and the long duration of many spinal procedures, the use of forced air warmers, fluid warmers, and blankets should be considered to help prevent hypothermia. Blankets may be placed over the feet up to the gluteal area and across the shoulders and arms out of the surgical field to aid in maintaining the patient’s body temperature. These devices should be used according to the manufacturer’s directions and at the discretion of the surgeon.

37. Position the patient’s arms and adjust the Articulating Arm Boards as needed. Support the arms in the foam Arm Cradles. The V of the cradle should rest against the inside of the patient’s elbow. Translate, raise, lower, or articulate the Arm Board as needed and then secure in place (Figure 84). Confirm that all the T-Handles and the Drop Handle are tightened. Confirm that the patient’s arms are positioned so there is no more than 90 degrees flexion at the shoulder and no more than 90 degrees flexion at the elbow. Use the Velcro® Arm Board Straps provided to secure the arms.

![Figure 84: Articulating Arm Board Functionality](image)

**WARNING:** Hyperextension of the shoulder may cause compression of the brachial plexus resulting in a potential nerve or vascular injury. Failure to ensure proper arm positioning may result in harm to the patient.

![Figure 85: Final Neutral Prone Positioning](image)
WARNING: Proper pre-operative and intra-operative procedures must be followed to prevent venous stasis and pooling, pressure sore development, neuropathy, improper electrosurgical tissue grounding, hypertension/hypotension, and hypothermia.

CAUTION: Use care and monitor the patient when changing the position of the ProAxis® Surgery Table (Height Up/Down, Hinge Up/Down, Trendelenburg/Reverse Trendelenburg, and Lateral Roll) to ensure that the patient or table does not interfere with other equipment and the components and accessories have proper clearance in relation to the surrounding areas. Failure to do so may cause harm to the patient or device.

38. Confirm that the patient is properly positioned. If a Retractor Adaptor is required to mount auxiliary equipment, mount it on the rail frame. Open the Adaptor by turning the black knob and releasing the bottom of the Adaptor. Place the Adaptor around the frame, align the black knob, turn to tighten and secure the Adaptor in place (Figure 86). Ensure that the patient does not directly contact the Retractor Adaptor. Use foam or gel padding as needed to pad the area between the patient and the Adaptor.

![Retractor Adaptor](image)

Figure 86: Retractor Adaptor Mounted on Carbon Fiber Rail

WARNING: Only the metal Retractor Adaptor may be positioned in the area above the hinge and below the rail mounts. Use of the optional Radiolucent Retractor Adaptor at this location results in a pinch point when using the hinge capabilities of the table, which may cause harm to the patient or the device.

NOTE: Use of the Radiolucent Retractor Adaptor at any location below the Hip Pads is allowed when the hinge functionality of the table is engaged. The Radiolucent Retractor Adaptor can also be used above the hinge when the table is configured for cervical and the hinge functionality is deactivated.

39. Refer to Section 4 for instructions on how to utilize the IntelliPendant® and operate the table.
CAUTION: The linkage arms of the columns raise, lower, and rotate when the table is in use. Ensure no items are in the path of the moving arms and healthcare professionals position themselves away from the linkage arms when the table is in motion. Failure to do so may cause injury to the healthcare professional or possible damage to the table.

Figure 87: Prone Positioning with Fixed Surgical Site Mode, Hinge Up
### 8.1.1 Components Used for Prone Patient Positioning

<table>
<thead>
<tr>
<th>Rail Mount</th>
<th>Chest Tray</th>
<th>5 Way Articulating Arm Board</th>
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<td><img src="image_url" alt="Chest Tray" /></td>
<td><img src="image_url" alt="5 Way Articulating Arm Board" /></td>
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<td><strong>6988-899, Left</strong></td>
<td><strong>6988-530, Arm Board Assembly with Bracket &amp; Pad (2)</strong></td>
</tr>
<tr>
<td><strong>Right</strong></td>
<td><strong>6988-887, Right</strong></td>
<td><strong>5379-7722 1-inch (2.5)</strong></td>
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<td><strong>6988-950</strong></td>
<td><strong>Tempur-Pedic® Medical Arm Board Pad</strong></td>
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<th>Contoured ACP Hip Pad, Left</th>
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<td><strong>6988-387</strong></td>
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<thead>
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<th>Leg Sling</th>
<th>Buttocks Strap</th>
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<td><strong>5840-450</strong></td>
<td><strong>6977-739</strong></td>
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<td>Safety Strap, 60 in (152 cm)</td>
<td>Retractor Adaptor</td>
<td>ProAxis® ProneView® Patient Care Kit</td>
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<th>ProAxis® Component Cart</th>
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<tr>
<td>ProAxis® GentleTouch® Patient Care Kit</td>
<td>Radiolucent Retractor Adaptor</td>
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Mizuho OSI 2016
### Optional Accessory

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<td>Cervical Traction Vector, Short</td>
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![Cervical Traction Vector, Short](image)
8.2 ProAxis® Prone Patient Positioning with Cervical Management

The following instructions provide the recommended steps for setup. Final determination of patient positioning and setup is at the discretion of the surgeon and surgical team.

**NOTE:** When setting up the ProAxis® Table for prone patient positioning with cervical management, the hinge capability of the table should not be enabled.

### 8.2.1 Prone Positioning with Cervical Management Base Unit

1. Confirm that the ProAxis® Spinal Surgery Table is prepared for setup (refer to Section 8).
2. Confirm that the Floor Lock symbol is visible on the display of the IntelliPendant® (Figure 22, page 25). Observe the display on the IntelliPendant® and confirm the frame is level and the hinge neutral. The Hinge, Trendelenburg, and Lateral Tilt angles should read 0° (Figure 22, page 25). If necessary, install the Right and Left Rail Mounts (refer to Section 8.1) to temporarily enable the hinge functionality. Utilizing the IntelliPendant®, press the Return to Level button to return the Hinged Frame to a neutral position. Remove the Rail Mounts and return them to their storage location on the ProAxis® Component Cart.

**NOTE:** DO NOT install the Left and Right Rail Mounts and the Torso Trolley® Chest Tray. The absence of these components will allow the hinge capabilities of the table to be disabled and the cervical components to be properly mounted.

3. Install the ProAxis® Cervical Management Mounting Bracket on the Head-End Assembly (Figure 90).
4. Orient the posts of the Bracket over the Head-End Assembly receptacle. Ensure the T-Handles of the receptacle are unscrewed. Slide the posts into the receptacle. The Bracket will rest on the top of the receptacle. Tighten both T-Handles (Figure 90).

5. To use the optional Mizuho OSI Cervical Management Base Unit on the ProAxis® Spinal Surgery Table:
   a. Remove the Cervical Management Base Unit Sub-Assembly from the Table Adaptor Assembly by:
      i. First, loosening the Horizontal Slide Locking Lever on the Table Adaptor Assembly.
      ii. Aligning the Stop Cap of the Horizontal Slide so it can pass through and out of the Table Adapter Assembly. Pull the Sub-Assembly away from the Table Adaptor Assembly to remove (Figure 91).
b. Attach the Cervical Management Base Unit Sub-Assembly into the ProAxis® Cervical Management Mounting Bracket. To attach:
   i. Ensure the Horizontal Slide Locking Lever on the ProAxis® Cervical Management Mounting Bracket is loosened and ready to receive the Cervical Management Base Unit Sub-Assembly.
   ii. Align the Stop Cap and insert the posts of the Base Unit into the ProAxis® Bracket (Figure 92).
   iii. Tighten the Horizontal Slide Locking Lever when the desired position of the Horizontal Slide is achieved.
6. In preparation for attaching a Skull Clamp, attach the appropriate adaptor to the end of the Cervical Management Base Unit. The Cervical Management Base Unit includes the aluminum Swivel Adaptor used to mount an aluminum DORO® or Mayfield® Skull Clamp. A specific Radiolucent Adaptor is required to mount a radiolucent DORO® or a radiolucent Mayfield® device. Refer to the Cervical Management Base Unit Owner’s Manual for ordering information and detailed instructions on the use of the Radiolucent Adaptors.

7. With the Cervical Management Base Unit mounted on the ProAxis®, adjustments can be made to the position of the Adaptor to which the Skull Clamp is connected (Figure 93).
   a. Refer to the Mizuho OSI Owner's Manual [REF NW0658 provided with the Cervical Management Base Unit for detailed information on the functionality of the Cervical Management Base Unit Sub-Assembly and its various adjustment options.
   b. Each of the Black Knobs (#1, 2, and 3 in Figure 93) on the Sub-Assembly allow for making adjustments to the position of the assembly until the desired position is achieved.

   ![Figure 93: Cervical Management Components](image)

   c. To raise the height of the Yoke and Transitional Arms, turn the Black Knob #1 until the desired height of the assembly is achieved. Loosen the Black Knobs, #2 and #3, controlling the position of the Transitional Arms holding the Adaptor. Adjustments can be made to the position and angle of the Assembly utilizing the Black Knobs. Tighten the Black Knobs when the desired position is achieved.

8. The components of the Cervical Management Base Unit Sub-Assembly have a starburst locking mechanism. Ensure that all teeth of the starburst are aligned and the corresponding knob is properly tightened when assembling or repositioning the device. All knobs should be tightened by hand; DO NOT use a wrench of any kind.

**WARNING:** Failure to ensure that all teeth of the starbursts are aligned and the knobs are properly tightened may cause harm to the patient, healthcare professional or the device.
9. Attach the Contoured ACP Hip Pads. The pads are labeled right and left specific to be oriented to the patient’s iliac crest when lying prone. Open the Locking Bracket, orient the Pad with Locking Bracket over the rail of the open frame on the Foot-End section of the frame, lower into place and slide towards the Hinge until it rests against the Pad Stop (Figure 71, page 67). Press up on the white tab until the black component of the clamp fits into the open notch of the white tab. Press down on the lip of the black component of the clamp to lock the pad in place. (Figure 72, page 67). When installed correctly, the pad should rest against the Pad Stop and the arrow on the side of the Bracket should be pointed toward the Foot-End of the table. To remove the Contoured Hip Pad, lift up on the lip of the Bracket so that it disconnects and lift the pad off the rail of the frame.

NOTE: Depending on the patient’s anatomy, the hip pads may be moved towards the Foot-End of the table at the surgeon’s discretion to achieve proper positioning.

10. Attach the Cervical Chest Tray by releasing the Velcro® Straps on both sides and orienting the Cervical Chest Tray above the rails of the frame in the desired location. Slide the Chest Tray over the Frame until it sits on the Frame, and secure with the Velcro® Straps (Figure 94).

![Cervical Chest Tray Installed on ProAxis® Frame](image)

11. To attach the Flat ACP or Non-ACP Chest Pad, orient the opening of the pad bracket so it surrounds the edge of the Cervical Chest Tray. Slide onto the Cervical Chest Tray until the tab of the bracket is secure on the tray (Figure 95). To remove, press the tab of the bracket and slide the pad towards the Foot-End of the table.

NOTE: With ideal positioning, the Chest Pad should support the patient such that the load is borne predominantly by the sternum. When the patient is positioned correctly, the top edge of the pad should rest below the patient’s anatomical landmark, the supra-sternal notch.

WARNING: Positioning the Chest Pad superior to the supra-sternal notch may apply pressure to the throat and the patient’s airway. Failure to ensure the patient is positioned correctly on the Chest Pad may result in harm to the patient or the device.
12. To install the Adjustable Leg Sling, place the sling over the Foot-End section of the open frame (Figure 74, page 68). Feed the four straps through their respective buckles on the opposite side of the sling and secure. The sling should be adjusted for each patient. The straps on the sling consist of a double-sided hook and loop closure that allows for infinite adjustment along the length of the strap. The hook end of the strap may be folded over onto the loop portion of the strap to allow for greater adjustability. The strap may be folded over itself again to shorten, thereby raising the sling and decreasing the degree to which the patient’s hips are flexed. Place two (2) or three (3) Tempur-Pedic® Medical Pillows in the sling approximately where the patient’s lower legs will rest to achieve the desired amount of knee flexion and support (Figure 75, page 69).

13. The Leg Boards may be used in place of the Leg Sling to support the legs. Attach the Leg Boards with the 1-inch (2.5 cm) Tempur-Pedic® Medical Pad. Release the Velcro® Straps on the sides of the Leg Board. Place one Leg Board over the rails of the open frame and slide into place. Secure with the Velcro® Straps. Repeat this process with the second Leg Board. Ensure that the two Leg Boards abut each other (Figure 76).

14. After the desired Flat Chest Pad and Contoured ACP Hip Pads are in place on the frame, the tubing can be connected to the pads.
   a. Align the tubing with the long end directed to the pad furthest away from the port at the Foot-End column (Figure 6, page 15).
   b. Secure the ACP tubing with the Velcro® wraps along the frame and by routing it through the underside of the Leg Sling. Ensure the tubing is secure and out of the way.
   c. Connect to the tubing to the port at the Foot-End of the table.
   d. Re-adjust the ACP tubing as may be required.

15. Place the Patient Kit Covers on the Chest and Hip Pads.
16. Trial position the imaging equipment if it will be used. If necessary, re-locate the ProAxis® Spinal Surgery Table in the room. Unlock the Floor Locks by using the IntelliPendant® and with the assistance of a second person move the ProAxis® Spinal Surgery Table. When the table is located where it will be used, lock the Floor Locks.

⚠️ CAUTION: The ProAxis® Spinal Surgery Table is not for patient transport. The location of the table relative to any imaging equipment that may be used needs to be confirmed prior to the patient being transferred to the table.

17. Refer to the specific manufacturer’s instructions for use regarding the Skull Clamp being attached to the Cervical Management Base Unit. The surgeon determines the best approach to secure the patient to the Skull Clamp.

18. Confirm the Black Knobs of the Cervical Management Base Unit are loosened so that the Transitional Arms and Adaptor can be readily positioned when connecting the Skull Clamp.

19. Position the patient bed or stretcher so the patient’s anatomical landmarks are aligned with the support pads on the hinged frame of the ProAxis® Spinal Surgery Table. It may be necessary to re-position the Chest Pad. The top edge of the Chest Pad should align with the patient’s anatomical landmark, the bottom of supra-sternal notch, and the top third of the Contoured ACP Hip Pad should align with the patient’s anatomical landmark, the iliac crest.

20. The height of the patient’s bed or stretcher should be level with the top of the Contoured ACP Hip Pad. To assist the patient transfer process, you may choose to use the IntelliPendant® and laterally tilt the table toward the patient bed or stretcher.

21. Lock the patient stretcher or bed. Confirm by observing the IntelliPendant® display that the table Floor Locks are locked. The Floor Locks Locked symbol should be visible (Figure 22, page 25).

22. The surgeon supports the patient’s head in the Skull Clamp while others assist to log-roll the patient onto the ProAxis® Spinal Surgery Table. The surgeon continues to support the patient’s head and Skull Clamp until final positioning is achieved with the Cervical Management Base Unit.

⚠️ WARNING: After log-rolling the patient to a prone position, confirm that the endotracheal tube is not kinked. Failure to confirm may cause harm to the patient.

23. Continue to support the patient on the table until the safety straps are in place. DO NOT leave the patient unattended on the table.

⚠️ WARNING: Patient must be restrained at all times by a safety strap(s) when positioned on the table.

24. Remove the patient bed or stretcher while supporting the patient on the table.
25. Visually confirm the placement of the Contoured ACP Hip Pads. The pad should be mounted abutting the Pad Stop (Figure 73, page 68). When positioned correctly, the patient’s anatomical landmark, the iliac crest, should rest in the top one-third of the pad. The Patient Kit Hip Pad Cover should be smooth against the patient’s skin. Ensure that the patient’s skin is smooth and flat against the Patient Kit Hip Pad Cover. To re-position the patient on the Pad, the patient should be lifted and moved such that the iliac crest contacts the top third of the pad. Ensure the Hip Pad Cover is smoothed and lower the patient back onto the Pad. Confirm correct placement of the Pads such that each of the patient’s iliac crests rest in the top one-third of each respective Hip Pad (Figure 81, page 74).

**CAUTION:** Failure to ensure that the patient is positioned correctly on the Contoured ACP Hip Pad may result in harm to the patient or the device.

26. Visually confirm the placement of the Chest Pad. When positioned correctly, the edge of the Pad should rest at the patient’s anatomical landmark, the supra-sternal notch (Figure 82, page 74). Breasts should be down and lying flat, with the nipples oriented towards the Foot-End of the table. The Patient Kit Chest Pad Cover should be smooth against the patient’s skin. Ensure that the patient’s skin is smooth and positioned flat against the cover. If necessary, re-position the Chest Tray. While lifting and supporting the patient, move the Chest Tray to the desired location and smooth the Chest Pad Cover. The patient should then be lowered back onto the Chest Pad.

**NOTE:** With ideal positioning, the Chest Pad should contact the patient such that the load is borne predominantly by the sternum.

**WARNING:** Positioning the Chest Pad superior to the supra-sternal notch may apply pressure to the throat and patient’s airway. Failure to ensure the patient is positioned correctly on the Chest Pad may result in harm to the patient or the device.

27. Visually confirm that the Leg Sling or Leg Boards provide support to the patient’s knees, and the legs are supported on the pillows. If the Leg Sling is being used, the patient’s knees should be flexed and legs supported on the pillows. Re-position the legs and add pillows as required under the patient’s tibias to ensure that the ankles are not hyper-extended and there is no pressure on the patient’s feet (Figure 81, page 74).

**CAUTION:** Failure to ensure that the patient is positioned correctly with the Leg Sling or Leg Boards may result in harm to the patient or the device.

28. Position the patient's arms back to his/her sides and secure. The disposable Positioning Straps may be used to assist in final positioning. The Arm Cradles may be used to provide padding.

29. Place the Buttocks Strap across the lower gluteal area of the patient. A blanket may be used under the Buttocks Strap to protect the patient from direct contact with the strap. The strap can be fed through the buckle on the sides of the Contoured ACP Hip Pads and then back on itself, securing the Velcro. Confirm that the Buttocks Strap is positioned low on the gluteal area, and that it surrounds both Contoured ACP Hip Pads, and is secured snugly around the buttocks. Ensure that the patient does not directly contact the frame of the table. Use foam or gel padding as needed to pad the area between the patient and the rails of the frame.

Figure 96: Final Prone Positioning with Cervical Management Base Unit

**NOTE:** The placement and location of the Straps are at the discretion of the surgeon and vary by procedure.

31. Due to the open nature of the ProAxis® frame and the long duration of many spinal procedures, the use of forced air warmers, fluid warmers, and blankets should be considered to help prevent hypothermia. Blankets may be placed over the feet up to the gluteal area and across the shoulders and arms out of the surgical field to aid in maintaining the patient's body temperature. These devices should be used according to the manufacturer’s directions and at the discretion of the surgeon.

**WARNING:** Proper pre-operative and intra-operative procedures must be followed to prevent venous stasis and pooling, pressure sore development, neuropathy, improper electrosurgical tissue grounding, hypertension/hypotension, and hypothermia.

**WARNING:** Use care and monitor the patient when changing the position of the ProAxis® Surgery Table (Height Up/Down, Trendelenburg/Reverse Trendelenburg, and Lateral Roll) to ensure that the patient or table does not interfere with other equipment, and the components and accessories have proper clearance in relation to the surrounding areas. Failure to do so may cause harm to the patient or device.

32. Confirm that the patient is in their final position. If a Retractor Adaptor is required to mount auxiliary equipment, mount it on the rail frame. Open the Adaptor by turning the black knob and releasing the bottom of the Adaptor. Place the Adaptor around the frame, align the black knob, turn to tighten and secure the Adaptor in place. Ensure that the patient does not directly contact the Retractor Adaptor. Use foam or gel padding as needed to pad the area between the patient and the Adaptor.

33. Refer to Section 4, Basic Operation, for instructions on how to utilize the IntelliPendant® and operate the table.

**CAUTION:** The linkage arms of the columns raise, lower, and rotate when the table is in use. Ensure no items are in the path of the moving arms and healthcare professionals position themselves away from the linkage arms when the table is in motion. Failure to do so may cause injury to the healthcare professional or possible damage to the table.
8.2.2 Prone Patient Positioning with Cervical Traction Vector Adjustor

The Cervical Traction Vector Adjustor is designed to provide adjustability of the angle of pull of cervical traction as well as to maintain alignment of the traction rope when skull tongs or a head halter is used.

1. Confirm that the ProAxis® Surgery Table is prepared for setup (refer to Section 8).
2. Confirm that the Floor Lock symbol is visible on the display of the IntelliPendant® (Figure 22, page 25). Observe the display on the IntelliPendant® and confirm the frame is level and the hinge neutral. The Hinge, Trendelenburg, and Lateral Tilt angles should read 0° (Figure 22, page 25). If necessary, temporarily install the Right and Left Rail Mounts (refer to Section 8.1) in order to enable the hinge functionality. Utilizing the IntelliPendant®, press the Return to Level button to return the Hinged Frame to the neutral position. Remove the Rail Mounts and return them to their storage location on the ProAxis® Component Cart.
3. Install the ProAxis® Cervical Traction Vector Adjustor. Extend the Drop Lock on the post of the Cervical Traction VectorAdjustor, align the posts with the Head-End Assembly receptacle, and lower the Vector into place (Figure 98). Position the Vector at the desired height and tighten both T-Handles on the Head-End Assembly (Figure 99).
Figure 98: Installation of Cervical Traction Vector Adjustor

**NOTE:** DO NOT install the Left and Right Rail Mounts and the Torso Trolley® Chest Tray. The absence of these components will allow the hinge capabilities of the table to be disabled and the cervical components to be properly mounted.

Figure 99: Cervical Traction Vector Adjustor Properly Installed on ProAxis®

4. A traction rope can be directed over the Traction Vector Adjustor, with the length of rope adjusted so the attached weight(s) can hang freely. While monitoring the position of the weights, utilize the IntelliPendant® to raise and lower the table through its full range of motion. Ensure the weight(s) continues to hang freely in all table positions. Adjust the length of traction rope if necessary.

5. Install the remaining prone positioning components as instructed in Section 8.2.2.
### 8.2.3 Components for Prone Patient Positioning with Cervical Accessories

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<tr>
<td>Cervical Chest Tray</td>
<td>6988-969</td>
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<tr>
<td>Flat Chest Pad</td>
<td>6988-758 ProAxis® Chest Pad</td>
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<td>Contoured ACP Hip Pad, Left</td>
<td>6988-387</td>
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<td>Contoured ACP Hip Pad, Right</td>
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<td>Leg Sling</td>
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<td>Leg Board</td>
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<td>Buttocks Strap</td>
<td>6977-739</td>
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<tr>
<td>Safety Strap, 60 in. (152 cm)</td>
<td>5840-43 (2)</td>
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<tr>
<td>Tempur-Pedic® Medical Pillow</td>
<td>Retractor Adaptor</td>
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<td>REF 6910-3034 DORO® Radiolucent Skull Clamp (pictured)</td>
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<td></td>
<td>REF 6910-1001 DORO® QR3 Aluminum Skull Clamp</td>
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<td>REF 6988-449 Radiolucent Retractor Adaptor</td>
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<tr>
<td>REF 5979-200 Radiolucent Mayfield® Adaptor for Cervical Management Base Unit</td>
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8.3 ProAxis® Supine Patient Positioning

The following instructions provide the recommended steps for setup. Final determination of patient positioning and setup is at the discretion of the surgeon and surgical team.

1. Confirm the ProAxis® Surgery Table is prepared for setup (refer to Section 8).

2. Confirm that the Floor Lock symbol is visible on the display of the IntelliPendant® (Figure 22, page 25). Observe the display on the IntelliPendant® and confirm the frame is level and the hinge neutral. The Hinge, Trendelenburg, and Lateral Tilt angles should read 0° (Figure 22, page 25). If necessary, temporarily install the Right and Left Rail Mounts (refer to Section 8.1) in order to enable the hinge functionality. Utilizing the IntelliPendant®, press the Return to Level button to return the Hinged Frame to a neutral position. Remove the Rail Mounts and return them to their storage location on the ProAxis® Component Cart.

3. Confirm that there are no accessories mounted to the rails before attempting to mount the Two-Piece Supine Top.

4. The ProAxis® Two-Piece Supine Top is made of carbon fiber construction and mounts on the open rail frame. Each section (Head-End and Foot-End) has a set of Locking Brackets that secure the top to the frame. The Head-End section also has a Cable Connector that plugs into the Head-End port, allowing the hinge to function (Figure 104, page 97).
   a. To install the Head-End section on the frame, open the Locking Brackets, align the Locking Brackets over the rails, and lower into place (Figure 101). The top should rest flat on the frame. Slide the Top toward the Head-End Column until it rests against the Brackets of the rail frame on both sides (Figure 107, page 99).
b. To engage the Locking Brackets on each side, first press up on the white tab until the black portion of the clamp fits into the open notch of the white tab. Press down on the lip of the black component of the clamp to lock it into place (Figure 102).
c. Connect the Cable Connector to the port on the Head-End Assembly and turn the locking collar (Figure 104).

d. To install the Foot-End Supine Top on the open frame, open the Locking Brackets, align the Locking Brackets over the rails, and lower into place. The top should rest flat on the frame.

e. Slide the top toward the Foot-End Column until it rests against the Brackets of the rail frame on both sides (Figure 105). When the Two-Piece Supine Top is correctly installed, there will be a gap between the two tops such that the hinge is exposed (Figure 106).
f. To engage the Locking Brackets on each side, first press up on the white tab until the black portion of the clamp fits into the open notch of the white tab. Press down on the lip of the black component of the clamp to lock it into place (Figure 102, page 96).

g. When necessary to remove the Supine Top, first disconnect the Cable Connector from the port. Then, open the Locking Brackets and lift the each top off the frame.

5. Install the Tempur-Pedic® Medical Pad on the table tops. Align the “A” label on the pad over the corresponding label on the Head-End Supine Top (Figure 107). The Velcro® on the pad and the top should also be aligned. Lower the pad onto the top securing the Velcro®.
6. Trial position the imaging equipment if it will be used. If necessary, re-locate the ProAxis® Surgery Table in the room. Unlock the Floor Locks by using the IntelliPendant® and with the assistance of a second person move the ProAxis® Surgery Table. When the table is located where it will be used, lock the Floor Locks.

⚠️ **CAUTION:** The ProAxis® Surgery Table is not for patient transport. The location of the table relative to any imaging equipment that may be used needs to be confirmed prior to the patient being transferred to the table.

7. Position the patient bed or stretcher so that the patient is aligned with the table top. The height of the patient bed or stretcher should be level with the top of the table pad.

**NOTE:** When the patient is correctly positioned prone on the ProAxis®, the hinge will flex or extend the patient’s spine at L3. To achieve the same when the patient is supine or lateral, align the patient’s umbilicus with the hinge. To access L4-L5 during a lateral approach, positioning the patient such that the hinge is centrally located between the patient’s iliac crest and greater trochanter may be required.

8. Lock the patient stretcher or bed. Confirm that the floor locks are locked. The Floor Lock symbol should be visible on the display of the IntelliPendant®.

9. Transfer the patient in a supine position using a standard drawsheet method.

10. Secure the patient with the Safety Straps. DO NOT leave the patient unattended on the table.
WARNING: Patient must be restrained at all times by a safety strap(s) when positioned on the table.

11. To mount the Safety Strap or any optional accessories such as Pivoting Arm Boards, Clark Sockets or a Cross Arm Support, a Universal Side Rail Adaptor should always be used. Universal Side Rail Adaptors are used to protect the carbon-fiber rails of the top from unnecessary damage.

12. To install the Universal Side Rail Adaptor:
   a. Open the clamp using the black knob.
   b. Place the Adaptor in the desired position, with the black knob facing down.
   c. Turn the black knob to tighten the clamp (Figure 108).
   d. Ancillary equipment can now be mounted on the 6-inch (15 cm) of side rail provided by the Universal Side Rail Adaptor.

   ![Figure 108: Universal Side Rail Adaptor Mounted on ProAxis® Supine Top](image)


WARNING: Proper pre-operative and intra-operative procedures must be followed to prevent venous stasis and pooling, pressure sore development, neuropathy, improper electrosurgical tissue grounding, hypertension/hypotension, and hypothermia.

WARNING: Use care and monitor the patient when changing the position of the ProAxis® Surgery Table; Height Up/Down, Hinge Up/Down, Trendelenburg/Reverse Trendelenburg, and Lateral Roll to ensure that the patient or table does not interfere with other equipment. Failure to do so may cause harm to the patient or device.
14. Refer to Section 4 for instructions on how to utilize the IntelliPendant® and operate the table.

⚠️ CAUTION: The linkage arms of the columns raise, lower, and rotate when the table is in use. Ensure no items are in the path of the moving arms and healthcare professionals position themselves away from the linkage arms when the table is in motion. Failure to do so may cause injury to the healthcare professional or possible damage to the table.
### 8.3.1 Components Used for Supine Patient Positioning

<table>
<thead>
<tr>
<th>Item</th>
<th>Reference(s)</th>
</tr>
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<tbody>
<tr>
<td>Two-Piece Supine Top with 2-inch (5 cm) Tempur-Pedic® Medical Pad</td>
<td></td>
</tr>
<tr>
<td></td>
<td><a href="#">REF 6988-623 Head-End Section</a></td>
</tr>
<tr>
<td></td>
<td><a href="#">REF 6988-621 Foot-End Section</a></td>
</tr>
<tr>
<td></td>
<td><a href="#">REF 6988-442 2-inch (5 cm) Tempur-Pedic® Medical Supine Top Pad</a> (only)</td>
</tr>
<tr>
<td>Universal Side Rail Adaptor (4)</td>
<td><a href="#">REF 6977-959</a></td>
</tr>
<tr>
<td>Safety Straps (2)</td>
<td><a href="#">REF 5855-550</a></td>
</tr>
<tr>
<td>Mizuho OSI Tempur-Pedic® Medical Positioning Pillows (3)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>[REF 6946 Standard (1)]</td>
</tr>
<tr>
<td></td>
<td>[REF 6947 Queen (2)]</td>
</tr>
<tr>
<td>Optional Accessory</td>
<td><a href="#">REF 6950 Mizuho OSI Tempur-Pedic® Medical 6 pc. Positioning Pad Set</a></td>
</tr>
<tr>
<td>Optional Accessory</td>
<td><a href="#">REF 5356 Pivoting Arm Board with 2-inch (5 cm) Pad</a></td>
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<tr>
<td></td>
<td><a href="#">REF 69441 Arm Board 1-inch (2.5 cm) Pad Only</a></td>
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<tr>
<td>Optional Accessory</td>
<td></td>
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<tr>
<td></td>
<td><a href="#">REF 5393 Clark Socket</a></td>
</tr>
<tr>
<td>Optional Accessory</td>
<td><a href="#">REF 5857 Cross Arm Support with 1-inch (2.5 cm) Pad</a></td>
</tr>
</tbody>
</table>
8.3.2 Supine Patient Positioning with Cervical Traction Vector Adjustor

The Cervical Traction Vector Adjustor is designed to provide adjustability of the angle of pull of cervical traction as well as to maintain alignment of the traction rope when skull tongs or a head halter is used.

1. Confirm that the ProAxis® Surgery Table is prepared for setup (refer to Section 8).
2. Install the Two-Piece Supine Top and Tempur-Pedic® Medical Pad for supine positioning as instructed in Section 8.3.
3. Install the ProAxis® Cervical Traction Vector Adjustor. Extend the Drop Lock on the post of the Cervical Traction Vector Adjustor, align the posts with the Head-End Assembly receptacle, and lower the Vector into place (Figure 111). Position the Vector at the desired height and tighten both T-Handles on the Head-End Assembly (Figure 99, page 92).

4. A traction rope can be directed over the Traction Vector Adjustor, with the length of rope adjusted so the attached weight(s) can hang freely.
5. While monitoring the position of the weights, utilize the IntelliPendant® to raise and lower the table through its full range of motion. Ensure the weight(s) continues to hang freely in all table positions. Adjust the length of traction rope if necessary.
### 8.3.3 Components for Supine Patient Positioning with Cervical Accessories

<table>
<thead>
<tr>
<th>Two-Piece Supine Top with 2-inch (5 cm) Tempur-Pedic® Medical Pad</th>
<th>Safety Straps (2)</th>
<th>Cervical Traction Vector Adjustor</th>
</tr>
</thead>
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<tr>
<td><img src="image1" alt="Two-Piece Supine Top with 2-inch (5 cm) Tempur-Pedic® Medical Pad" /></td>
<td><img src="image2" alt="Safety Straps (2)" /></td>
<td><img src="image3" alt="Cervical Traction Vector Adjustor" /></td>
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<td>REF 6988-623 Head-End Section</td>
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<td>REF 6988-725</td>
</tr>
<tr>
<td>REF 6988-621 Foot-End Section</td>
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<td></td>
</tr>
<tr>
<td>REF 6988-442 2-inch (5 cm) Tempur-Pedic® Medical Supine Top Pad</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Mizuho OSI Tempur-Pedic® Medical Positioning Pillows (3)</th>
<th>Universal Side Rail Adaptor (4)</th>
<th>Optional Accessory</th>
</tr>
</thead>
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<td><img src="image5" alt="Universal Side Rail Adaptor (4)" /></td>
<td><img src="image6" alt="Optional Accessory" /></td>
</tr>
<tr>
<td>REF 6946 Standard (1)</td>
<td>REF 6977-959</td>
<td>REF 6950 Mizuho OSI Tempur-Pedic® Medical 6 pc. Positioning Pad Set</td>
</tr>
<tr>
<td>REF 6947 Queen (2)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
8.4 ProAxis® Lateral Patient Positioning

The following instructions provide the recommended steps for setup. Final determination of patient positioning and setup is at the discretion of the surgeon and surgical team.

1. Confirm the ProAxis® Spinal Surgery Table is prepared for setup (refer to Section 8).

2. Confirm that the Floor Lock symbol is visible on the display of the IntelliPendant® (Figure 22, page 25). Observe the display on the IntelliPendant® and confirm the frame is level and the hinge neutral. The Hinge, Trendelenburg, Reverse Trendelenburg, and Lateral Tilt angles should read 0° (Figure 22, page 25). If necessary, temporarily install the Right and Left Rail Mounts (refer to Section 8.1) in order to enable the hinge functionality. Utilizing the IntelliPendant®, press the Return to Level button to return the Hinged Frame to a neutral position. Remove the Rail Mounts and return them to their storage location on the ProAxis® Component Cart.

3. Confirm that there are no accessories mounted to the rails before attempting to mount the Two-Piece Supine Top.

4. The ProAxis® Two-Piece Supine Top is made of carbon fiber construction and mounts on the open rail frame. Each section (Head-End and Foot-End) has a set of Locking Brackets that secure the top to the frame. The Head-End section also has a Cable Connector that plugs into the Head-End port, allowing the hinge to function (Figure 104, page 97).
   a. To install the Head-End section on the frame, open the Locking Brackets, align the Locking Brackets over the rails, and lower into place (Figure 101, page 96). The top should rest flat on the frame. Slide the Top toward the Head-End Column until it rests against the Brackets of the rail frame on both sides (Figure 107, page 99).
   b. To engage the Locking Brackets on each side, first press up on the white tab until the black portion of the clamp fits into the open notch of the white tab. Press down on the lip of the black component of the clamp to lock it into place (Figure 102, page 96).
   c. Connect the Cable Connector to the port on the Head-End Assembly and turn the locking collar (Figure 104, page 97).
d. To install the Foot-End Supine Top on the open frame, open the Locking Brackets, align the Locking Brackets over the rails, and lower into place. The top should rest flat on the frame.

e. Slide the top toward the Foot-End Column until it rests against the Brackets of the rail frame on both sides. (Figure 105, page 98). When the Two-Piece Supine Top is correctly installed, there will be a gap between the two tops such that the hinge is exposed (Figure 106, page 98).

f. To engage the Locking Brackets on each side, first press up on the white tab until the black portion of the clamp fits into the open notch of the white tab. Press down on the lip of the black component of the clamp to lock it into place (Figure 102, page 96).

g. When necessary to remove the Supine Top, first disconnect the Cable Connector from the port. Then, open the Locking Brackets and lift the each top off the frame.

5. Install the Tempur-Pedic® Medical Pad on the table tops. Align the “A” label on the pad over the corresponding label on the Head-End Supine Top (Figure 107, page 99). The Velcro® on the pad and the top should also be aligned. Lower the pad onto the top securing the Velcro®.

6. Trial position the imaging equipment if it will be used. If necessary, re-locate the ProAxis® Surgery Table in the room. Unlock the Floor Locks by using the IntelliPendant® and with the assistance of a second person move the ProAxis® Surgery Table. When the table is located where it will be used, lock the Floor Locks.

⚠️ CAUTION: The ProAxis® Surgery Table is not for patient transport. The location of the table relative to any imaging equipment that may be used needs to be confirmed prior to the patient being transferred to the table.

7. Position the patient bed or stretcher so that the patient is aligned with the table top. The height of the patient bed or stretcher should be level with the top of the table pad.

**NOTE:** *When the patient is correctly positioned prone on the ProAxis®, the hinge will flex or extend the patient’s spine at L3. To achieve the same when the patient is supine or lateral, align the patient’s umbilicus with the hinge. To access L4-L5 during a lateral approach, positioning the patient such that the hinge is centrally located between the patient’s iliac crest and greater trochanter may be required.*

8. Lock the patient stretcher or bed. Confirm that the floor locks are locked. The Floor Lock symbol should be visible on the display of the IntelliPendant®.

9. Transfer the patient in a supine position using a standard drawsheet method.

10. Stabilize and maintain the patient in the lateral decubitus position by means of at least two (2) patient safety straps or a lateral positioning device (REF 5300 or REF 5301).

⚠️ WARNING: Patient must be restrained at all time by a safety strap(s) when positioned on the table.

11. To mount safety straps or optional accessories such as Lateral Arm Boards, Clark Sockets or Lateral Positioners, a Universal Side Rail Adaptor should always be used. Universal Side Rail Adaptors are used to protect the carbon-fiber rails of the top from unnecessary damage.
12. To install the Universal Side Rail Adaptor:
   a. Open the clamp using the black knob.
   b. Place the Adaptor in the desired position, with the black knob facing down.
   c. Turn the black knob to tighten the clamp (Figure 108, page 100).
   d. Ancillary equipment can now be mounted on the 6 inches (15 cm) of side rail provided by the Universal Side Rail Adaptor.

13. To attach the Lateral Positioner Set (REF 5300):
   a. Ensure that three Universal Side Rail Adaptors have been attached ready to receive the attachments.
   b. To mount the rectangular Positioners, first place the Aluminum Side Rail Sockets on the Universal Side Rail Adaptors.
   c. Mount the rectangular Positioners such that one pad is between the scapulae, the second at the sacrum, and the third over the pubis. Tighten the T-Handles of the Aluminum Side Rail Sockets.
   d. Adjust each pad position until it rests against the patient.
   e. Tighten the pads in place by turning the Drop Handles.

14. To use the pivoting pubic pad of the Deluxe Lateral Positioner Set (REF 5301):
   a. Mount a Clark Socket on the Universal Side Rail Adaptor. Place the post of the Pubic Pad in the Clark Socket and tighten.
   b. To adjust the position of the Pubic Pad, loosen the Drop Handles, pivot to the desired position, and tighten in place by turning the Drop Handles.

15. Standard positioning aids (Mizuho OSI REF 6946 or REF 6947 Tempur-Pedic® Medical Pillows, REF 6950 Positioning Pad Set, foam or gel padding) may be used to provide additional patient stability and comfort.

WARNING: Proper pre-operative and intra-operative procedures must be followed to prevent venous stasis and pooling, pressure sore development, neuropathy, improper electrosurgical tissue grounding, hypertension/hypotension and hypothermia.

WARNING: Use care and monitor the patient when changing the position of the ProAxis® Surgery Table (Height Up/Down, Hinge Up/Down, Trendelenburg/Reverse Trendelenburg, and Lateral Roll) to ensure that the patient or table do not interfere with other equipment, and the components and accessories have proper clearance in relation to the surrounding areas. Failure to do so may cause harm to the patient or device.
16. Refer to Section 4 for instructions on how to utilize the IntelliPendant® and operate the table.

⚠️ CAUTION: The linkage arms of the columns raise, lower, and rotate when the table is in use. Ensure no items are in the path of the moving arms and healthcare professionals position themselves away from the linkage arms when the table is in motion. Failure to do so may cause injury to the healthcare professional or possible damage to the table.
### 8.4.1 Components Used for Lateral Patient Positioning

<table>
<thead>
<tr>
<th>Two-Piece Supine Top with 2-inch (5cm) Tempur-Pedic® Medical Pad</th>
<th>Universal Side Rail Adaptor (4)</th>
<th>Safety Straps (2)</th>
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</thead>
<tbody>
<tr>
<td><img src="image1.png" alt="Image" /></td>
<td><img src="image2.png" alt="Image" /></td>
<td><img src="image3.png" alt="Image" /></td>
</tr>
<tr>
<td>REF 6988-623 Head-End Section</td>
<td>REF 6977-959</td>
<td>REF 5855-550</td>
</tr>
<tr>
<td>REF 6988-621 Foot-End Section</td>
<td>REF 6988-442 2-inch (5 cm) Tempur-Pedic® Medical Supine Top Pad</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Mizuho OSI Tempur-Pedic® Medical Positioning Pillows</th>
<th>Optional Accessory</th>
<th>Optional Accessory</th>
</tr>
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<tr>
<td><img src="image4.png" alt="Image" /></td>
<td><img src="image5.png" alt="Image" /></td>
<td><img src="image6.png" alt="Image" /></td>
</tr>
<tr>
<td>REF 6946 Standard</td>
<td>REF 6950 Mizuho OSI Tempur-Pedic® Medical 6 pc. Positioning Pad Set</td>
<td>REF 5364 Lateral Arm Board Unit with 2-inch (5 cm) Pads</td>
</tr>
<tr>
<td>REF 6947 Queen</td>
<td></td>
<td></td>
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</tbody>
</table>

<table>
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<th>Optional Accessory</th>
<th>Optional Accessory</th>
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<td><img src="image8.png" alt="Image" /></td>
<td><img src="image9.png" alt="Image" /></td>
</tr>
<tr>
<td>REF 5394 Aluminum Side Rail Socket</td>
<td>REF 5393 Clark Socket</td>
<td>REF 5301 Deluxe Lateral Positioner with Ball Joint (pictured)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>REF 5300 Lateral Positioner Set</td>
</tr>
</tbody>
</table>
# ProAxis® Optional Accessories

Many optional accessories are available for use on the ProAxis® Spinal Surgery Table. For more information, please contact your local Mizuho OSI Sales Representative or Mizuho OSI Sales Operations at 1-800-777-4574 within the USA or +1-510-429-1500 outside the USA. At the prompt, select Option 1.

<table>
<thead>
<tr>
<th>REF</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>6988A-GT-ACP</td>
<td>ProAxis® GentleTouch® Patient Care Kit Each ProAxis® GentleTouch® Patient Care Kit includes the following: 6-inch (15 cm) GentleTouch® Face Pillow; Chest Pad Cover for ProAxis® Chest Pad; Pair of Hip Pad Covers with Aprons; Pair of Positioning Straps; Pair of Foam Arm Cradles; one (1) Set of Single-Use, Disposable Advanced Control Pad System™ Tubing. A case of six (6) individual kits is provided when ordering 6988A-GT-ACP. <strong>NOTE:</strong> Remove the Patient Care Kit components from the packaging and visually inspect the components. The GentleTouch® Face Pillow and the Arm Cradles should be opened twenty (20) minutes prior to use for complete expansion of the foam.</td>
</tr>
<tr>
<td>6988-449</td>
<td>Radiolucent Retractor Adaptor The Radiolucent Retractor Adaptor provides six (6) inches (15 cm) of side rail and can be used as an alternative to the standard Retractor Adaptor for mounting ancillary equipment onto the carbon fiber open frame.</td>
</tr>
<tr>
<td>6988-720</td>
<td>Cervical Traction Vector, Short The Cervical Traction Vector, Short may be used when a head halter or tongs is needed to support the head in conjunction with the Torso Trolley®. This shorter version allows for use with the Prone Torso Trolley® configuration.</td>
</tr>
<tr>
<td>5979-1</td>
<td>Cervical Management Base Unit The Cervical Management Base Unit accommodates patients being positioned for cervical procedures with the use of a DORO® and/or Mayfield® Skull Clamp. The unit provides a swivel adaptor for use with an aluminum Mayfield® or DORO® Skull Clamp.</td>
</tr>
<tr>
<td>6910-1001</td>
<td>Aluminum DORO® QR3 Skull Clamp The aluminum DORO® QR3 Skull Clamp provides 3-point fixation and can be mounted on the Cervical Management Base Unit.</td>
</tr>
<tr>
<td>6910-3034</td>
<td>Radiolucent DORO® Skull Clamp The Radiolucent DORO® Skull Clamp provides 3-point fixation and can be mounted on the Cervical Management Base Unit utilizing the Radiolucent DORO® Adaptor, which is sold separately.</td>
</tr>
<tr>
<td>5979-300</td>
<td>Radiolucent DORO® Adaptor for Cervical Management Base Unit The Radiolucent DORO® Adaptor is required to attach the Radiolucent DORO® Skull Clamp to the Cervical Management Base Unit.</td>
</tr>
<tr>
<td>5979-200</td>
<td>Radiolucent Mayfield® Adaptor for Cervical Management Base Unit The Radiolucent Mayfield® Adaptor is required to attach the Radiolucent Mayfield® Skull Clamp to the Cervical Management Base Unit.</td>
</tr>
<tr>
<td>6910-3006-5</td>
<td>Stainless DORO® Skull Pins (Adult) Skull pins provided in sterile packages of three (3). There are 12 packages in 1 box. Stainless pins are for use with the aluminum skull clamp.</td>
</tr>
<tr>
<td>6910-3006-2</td>
<td>Titanium DORO® Skull Pins (Adult) Skull pins provided in sterile packages of three (3). There are 12 packages in 1 box. Titanium pins are for use with the radiolucent skull clamp.</td>
</tr>
<tr>
<td><strong>REF</strong></td>
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<td><strong>6977-973</strong></td>
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</tr>
</tbody>
</table>
10 Cleaning, Storage, and Maintenance

10.1 Cleaning and Disinfecting

**NOTE:** Clean and disinfect the ProAxis® Spinal Surgery Table after each use.

10.1.1 Table Exterior

The exterior surface should be regularly wiped clean with a mild detergent solution and wiped dry with a soft lint-free cloth. This includes the table pad, table top, and all components and accessories. Care should be taken to avoid exposing the table to excessive moisture. Flooding, fogging, or steam cleaning should not be used to clean the table.

⚠️ **CAUTION:** Never pour any liquid directly onto the ProAxis® Spinal Surgery Table. Never subject the ProAxis® Spinal Surgery Table to an equipment washing machine or any pressurized stream of water.

Blood or other fluids, etc., if allowed to remain on the table for a long period of time, will require special cleaning to remove. A 5% acetic acid solution (or white vinegar) is especially good for this purpose.

**NOTE:** Use of iodophors will cause staining.

To remove staining or discoloration of plated or stainless steel surfaces, clean with a commercial cleaning compound labeled for stainless steel and then buff the surface by hand.

To disinfect exterior surfaces, use a quaternary ammonium disinfectant compound according to manufacturer's directions for use. Wipe dry with a soft lint-free cloth.

**NOTE:** Failure to thoroughly dry the surface after cleaning and disinfecting may result in rust.

⚠️ **WARNING:** Should it be necessary to disconnect the spars from the Head-End of the table for cleaning and disinfecting purposes, do not use excessive force to disconnect the spars. If after unlocking the spar lock, the spar does not disconnect with hand force, seek assistance from qualified service personnel. Failure to properly disconnect the spars may cause harm to the healthcare professional or the device.

10.1.2 ProneView® Helmet and Mirror System

Clean the ProneView® Helmet and Mirror after each use by spraying and wiping with a germicidal cleaner then wipe dry, or wash with anti-bacterial soap and water, rinse, then wipe dry.

**NOTE:** Do not autoclave the ProneView® Helmet or Mirror Platform.
10.1.3 Mizuho OSI Tempur-Pedic® Medical Pads

**NOTE:** The Mizuho OSI Tempur-Pedic® Medical Pads should always be stored in a flat position. The pads can become stiff in cold temperatures and can crack and break if in a rolled position. It is important that you allow the pads to warm to room temperature before attempting to use or handle.

When handling a Tempur-Pedic® Medical Pad, always grasp by the entire thickness of the pad.

**CAUTION: DO NOT** lift, slide, or carry Mizuho OSI Tempur-Pedic® Medical Pads by grabbing the fabric cover. The cover may tear or rip. Do not leave the pads folded, as the pad construction may take a permanent crease in the bent area. The permanent crease will cause performance issues of the pad and may cause patient harm or injury.

The pads are intended to be cleaned in place. They do not need to be rotated or removed.

Clean with standard hospital disinfectants labeled for use on table pads. Always dilute and rinse per manufacturer’s label instructions. Wipe dry with a lint free cloth. DO NOT soak or autoclave the pads.

**NOTE:** The use of bleach or highly concentrated chemicals may discolor the cover and will void the warranty on the pad.

When cleaning the bottom of the pad or the table top, simply lift one end of the pad, and fold it over onto the other end. Clean the pad or the table top, wipe dry with a lint free cloth then reposition the pad to flat on the table top.

10.2 Storage

When not in use, the ProAxis® Spinal Surgery Table should be stored in a clean, dry environment.

The following conditions are required of the shipping and/or storage environment:

- Ambient temperature -4°F (-20°C) to + 122°F (50°C).
- Relative humidity from 10% to 95%, non-condensing.
- Atmospheric pressure from 75 to 105 kPa.

When in storage, the table cover provided serves as a dust cover and should be used. Also, to ensure that the battery is always fully charged and ready for use, store the table connected to AC power that matches the ratings on the manufacturer’s label on the Foot-End column of the table (Figure 10, page 18).

10.3 Maintenance

Cleaning the table surfaces and table pad after each use will assure many years of trouble-free service.

Mizuho OSI recommends customers inspect and maintain Tempur-Pedic® pads per Mizuho OSI’s pad inspection program guidelines. Contact your Mizuho OSI sales representative or Mizuho OSI Services for a pad inspection checklist.

All components are lubricated for life at the factory and no other lubrication is required.

Preventative Maintenance is required every six (6) months. Contact Mizuho OSI Services for a complete Preventative Maintenance Checklist.
For detailed repair information or to order replacement parts, call or contact via the web Mizuho OSI Services:
1-800-777-4674, Option 2 (for calls within the USA)
+1-510-429-1500, Option 2 (for calls outside the USA)
+1-510-429-8324 (Fax)
service@mizuhosi.com
www.mizuhosi.com

Mizuho OSI Services is available from 5 AM to 5 PM Pacific Time, Monday through Friday. Please leave a message after normal business hours.
11 Software Troubleshooting

In the event that the IntelliPendant® presents an error message, first identify the code and error message that appears on the screen (Figure 114). The error codes contain four hexadecimal digits, SDCC.

- The first two characters displayed (SD) are numerical codes representing the source of the error, and correspond to text information provided on the top of the screen. In the example in Figure 114, the SD code is 21, which indicates the source of the error is the Head Shoulder Motor Controller.

- The last two characters displayed (CC) indicate the specific error, and correspond to the text information provided in the center of the screen. In the example in Figure 114, the CC code is 02, which indicates the error is a Motor Stall.

The error messages received can be categorized into three different types of errors:

1. Recoverable Error: An error that can be cleared by pressing the check mark Soft Key and continuing with the desired table functions
2. Reboot Error: An error that can be cleared by turning the table power off and back on.
3. Call for Service Error: An error that prevents the table from properly operating. In such instances, the user must contact Mizuho OSI Services immediately.

The following tables list the error codes and message that may appear on the IntelliPendant® along with the corresponding potential cause(s) and actions to take in order to troubleshoot the problem.

NOTE: The User Action listed as “Acknowledge Error” corresponds to the act of pressing the check mark Soft Key on the IntelliPendant®.

NOTE: Should the problem persist after the user presses the check mark Soft Key on the IntelliPendant® to “Acknowledge Error,” contact Mizuho OSI Services (Section 13).
### Code (SD) and Error Source and Detail

<table>
<thead>
<tr>
<th>Code (SD)</th>
<th>Error Source and Detail</th>
</tr>
</thead>
<tbody>
<tr>
<td>00</td>
<td>None – general</td>
</tr>
<tr>
<td>10</td>
<td>Main Controller</td>
</tr>
<tr>
<td>21</td>
<td>Motor Controller, Head Shoulder</td>
</tr>
<tr>
<td>22</td>
<td>Motor Controller, Head Elbow</td>
</tr>
<tr>
<td>23</td>
<td>Motor Controller, Head Wrist</td>
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<tr>
<td>24</td>
<td>Motor Controller, Foot Shoulder</td>
</tr>
<tr>
<td>25</td>
<td>Motor Controller, Foot Elbow</td>
</tr>
<tr>
<td>26</td>
<td>Motor Controller, Foot Wrist</td>
</tr>
<tr>
<td>27</td>
<td>Motor Controller, Tilt</td>
</tr>
<tr>
<td>30</td>
<td>Chest Tray</td>
</tr>
<tr>
<td>41</td>
<td>Floor Lock, Head-End</td>
</tr>
<tr>
<td>42</td>
<td>Floor Lock, Foot-End</td>
</tr>
<tr>
<td>50</td>
<td>Hand Pendant</td>
</tr>
<tr>
<td>60</td>
<td>Auxiliary Control Panel</td>
</tr>
<tr>
<td>70</td>
<td>Pump Controller</td>
</tr>
</tbody>
</table>

### Code (SD) and Error Message

<table>
<thead>
<tr>
<th>Code (SD)</th>
<th>Code (CC)</th>
<th>Error Message</th>
<th>Potential Cause(s)</th>
<th>User Action(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>00</td>
<td></td>
<td>General Errors</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>C0</td>
<td></td>
<td>Board reset (Power on)</td>
<td>Board was powered on.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>C1</td>
<td></td>
<td>Brown out reset</td>
<td>Board temporarily lost power.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>C2</td>
<td></td>
<td>Watchdog reset</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>C3</td>
<td></td>
<td>Software reset</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>C4</td>
<td></td>
<td>External reset</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>C5</td>
<td></td>
<td>Illegal opcode reset</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>C6</td>
<td></td>
<td>Trap conflict reset</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>C7</td>
<td></td>
<td>Unknown reset</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>C8</td>
<td></td>
<td>Stack overflow reset</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>C9</td>
<td></td>
<td>Stack underflow reset</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>D0</td>
<td></td>
<td>Low logic supply voltage</td>
<td>The logic voltage into the board is low.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>D2</td>
<td></td>
<td>Low drive supply voltage</td>
<td>The drive voltage into the board is low.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>E0</td>
<td></td>
<td>Invalid state transition</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>CODE (SD)</td>
<td>CODE (CC)</td>
<td>ERROR MESSAGE</td>
<td>POTENTIAL CAUSE(S)</td>
<td>USER ACTION(S)</td>
</tr>
<tr>
<td>----------</td>
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</tr>
<tr>
<td>10</td>
<td>Main Controller Errors</td>
<td>23</td>
<td>Move not stopped</td>
<td>User requested motion but the system timed out waiting for one or more boards to be ready.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>25</td>
<td>Head wrist maximum height</td>
<td>The table moved past the allowed maximum height of the Head-End wrist.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>26</td>
<td>Foot wrist maximum height</td>
<td>The table moved past the allowed maximum height of the Foot-End wrist.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>27</td>
<td>Head wrist minimum height</td>
<td>The table moved past the allowed minimum height of the Head-End wrist.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>28</td>
<td>Foot wrist minimum height</td>
<td>The table moved past the allowed minimum height of the Foot-End wrist.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>29</td>
<td>Minimum hinge height</td>
<td>The table moved past the allowed minimum height of the hinge.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2A</td>
<td>Maximum hinge angle</td>
<td>The table moved past the allowed maximum hinge angle.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2B</td>
<td>Minimum hinge angle</td>
<td>The table moved past the allowed minimum hinge angle.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2C</td>
<td>Maximum tilt angle</td>
<td>The table moved past the allowed maximum tilt angle.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2D</td>
<td>Minimum tilt angle</td>
<td>The table moved past the allowed minimum tilt angle.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2E</td>
<td>Table not ready to start new motion</td>
<td>The user requested motion but one of the boards was not ready for motion for a period that exceeded the timeout.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>41</td>
<td>Communication lost with Motor board</td>
<td>One or more motor boards are not communicating with the main controller.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>42</td>
<td>Communication lost with Chest Tray board</td>
<td>The chest tray board is not communicating with the main controller.</td>
</tr>
<tr>
<td>Code (SD)</td>
<td>Code (CC)</td>
<td>Error Message</td>
<td>Potential Cause(s)</td>
<td>User Action(s)</td>
</tr>
<tr>
<td>-----------</td>
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</tr>
<tr>
<td>43</td>
<td></td>
<td>Communication lost with Floor Lock board</td>
<td>One or both floor lock boards are not communicating with the main controller.</td>
<td>Turn table power off and back on. Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>44</td>
<td></td>
<td>Communication lost with Pump board</td>
<td>The ACP board is not communicating with the main controller.</td>
<td>Turn table power off and back on. Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>45</td>
<td></td>
<td>Table failed Initialize Stage One. If problem persists, call for service</td>
<td>The main controller failed to properly initialize.</td>
<td>Turn table power off and back on.</td>
</tr>
<tr>
<td>46</td>
<td></td>
<td>Table failed Initialize Stage Two. If problem persists, call for service</td>
<td>The main controller failed to properly initialize.</td>
<td>Turn table power off and back on.</td>
</tr>
<tr>
<td>47</td>
<td></td>
<td>Table failed to initialize. If problem persists, call for service</td>
<td>The main controller failed to properly initialize.</td>
<td>Turn table power off and back on.</td>
</tr>
<tr>
<td>48</td>
<td></td>
<td>Self-Test timed out &lt;stage&gt; failed</td>
<td>The main controller failed to properly initialize.</td>
<td>Turn table power off and back on.</td>
</tr>
<tr>
<td>60</td>
<td></td>
<td>Motor board version mismatch</td>
<td>The software programmed into one or more motor boards is incompatible with the main controller software.</td>
<td>Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>61</td>
<td></td>
<td>Chest tray version mismatch</td>
<td>The software programmed into the chest tray board is incompatible with the main controller software.</td>
<td>Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>62</td>
<td></td>
<td>Floor lock version mismatch</td>
<td>The software programmed into one or both floor lock boards is incompatible with the main controller software.</td>
<td>Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>63</td>
<td></td>
<td>ACP version mismatch</td>
<td>The software programmed into the ACP board is incompatible with the main controller software.</td>
<td>Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>64</td>
<td></td>
<td>Hand pendant version mismatch</td>
<td>The software programmed into the hand pendant is incompatible with the main controller software.</td>
<td>Contact Mizuho OSI Services to replace the hand pendant.</td>
</tr>
<tr>
<td>65</td>
<td></td>
<td>Control panel version mismatch</td>
<td>The software programmed into the auxiliary control panel board is incompatible with the main controller software.</td>
<td>Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>70</td>
<td></td>
<td>ADC failed self-test</td>
<td>The analog to digital converter timed out during self-test.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>71</td>
<td></td>
<td>Multiple keys depressed at the same time</td>
<td>Multiple keys were depressed for longer than a specific period.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>CODE (SD)</td>
<td>CODE (CC)</td>
<td>ERROR MESSAGE</td>
<td>POTENTIAL CAUSE(S)</td>
<td>USER ACTION(S)</td>
</tr>
<tr>
<td>----------</td>
<td>----------</td>
<td>-------------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------------</td>
<td>-----------------------------------------------------</td>
</tr>
<tr>
<td>72</td>
<td></td>
<td>Possible stuck key on Pendant or Control Panel. If problem persists, call for service.</td>
<td>A single key was still depressed for longer than a specific period after a multiple key press.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>A0</td>
<td></td>
<td>Clock battery failure</td>
<td>The small coin-cell battery on the main controller board is not working.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>A1</td>
<td></td>
<td>Clock oscillator not running</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>A2</td>
<td></td>
<td>Clock not incrementing</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>A3</td>
<td></td>
<td>Clock failed self-test</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>A4</td>
<td></td>
<td>Clock communication error – Invalid request</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>A5</td>
<td></td>
<td>Clock communication error – Queue full</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>A6</td>
<td></td>
<td>Clock failed to initialize</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>A7</td>
<td></td>
<td>RTC SRAM checksum failure (internal error)</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>A8</td>
<td></td>
<td>RTC EEPROM checksum failure (internal error)</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>A9</td>
<td></td>
<td>RTC SRAM invalid hinge mode (internal error)</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>AA</td>
<td></td>
<td>RTC SRAM invalid IMS state (internal error)</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>AB</td>
<td></td>
<td>RTC SRAM invalid storage mode (internal error)</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>AC</td>
<td></td>
<td>RTC SRAM invalid version (internal error)</td>
<td>Internal software error.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>B0</td>
<td></td>
<td>Reduce load, cycle Table power Call service if error persists OVERLOAD!</td>
<td>The IMS has detected a loading condition that exceeds the red threshold.</td>
<td>Turn table power off and back on. Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>B1</td>
<td></td>
<td>Reduce load Turn table off and back on OVERLOAD!</td>
<td>The IMS has detected a loading condition that exceeds the yellow threshold.</td>
<td>Turn table power off and back on.</td>
</tr>
<tr>
<td>B2</td>
<td></td>
<td>Table is still overloaded after cycling power. Continue operation?</td>
<td>The IMS has detected a loading condition that still exceeds the yellow threshold after a reboot.</td>
<td>Acknowledge the condition.</td>
</tr>
</tbody>
</table>
## 2X Motor Controller Errors

<table>
<thead>
<tr>
<th>CODE (SD)</th>
<th>CODE (CC)</th>
<th>ERROR MESSAGE</th>
<th>POTENTIAL CAUSE(S)</th>
<th>USER ACTION(S)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B3</td>
<td></td>
<td>Spars under stress, not ready for storage. If problem persists, call for service.</td>
<td>The IMS has detected a loading condition that still exceeds the white threshold while attempting to go through the table storage process, after moving into the storage position.</td>
<td>Acknowledge error. Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>B4</td>
<td></td>
<td>IMS Tension gauge failure</td>
<td>The IMS tension reading is erroneous.</td>
<td>Turn table power off and back on.</td>
</tr>
<tr>
<td>B5</td>
<td></td>
<td>IMS Moment gauge failure</td>
<td>The IMS moment reading is erroneous.</td>
<td>Turn table power off and back on.</td>
</tr>
<tr>
<td>01</td>
<td></td>
<td>Motor fault</td>
<td>The motor board detected a problem with the motor driver. The system may have attempted to change motor speed too abruptly or the motor hall effect sensors may have failed, among other potential causes.</td>
<td>Acknowledge error.</td>
</tr>
<tr>
<td>02</td>
<td></td>
<td>Motor stall</td>
<td>The motor board detected no motor movement while driving the motor very hard. The relative encoder may have failed, the gear train may be sluggish or locked, or the table may be overloaded, among other potential causes.</td>
<td>Acknowledge error. Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>03</td>
<td></td>
<td>Brake engaged current too high</td>
<td>The motor board detected more current flowing than expected in the brake while it should be engaged.</td>
<td>Acknowledge error. Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>04</td>
<td></td>
<td>Brake disengaged current too low</td>
<td>The motor board detected less current flowing than expected in the brake while it should be disengaged.</td>
<td>Acknowledge error. Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>05</td>
<td></td>
<td>Absolute encoder fault</td>
<td>The motor board failed to read the absolute encoder.</td>
<td>Acknowledge error. Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>06</td>
<td></td>
<td>Relative encoder fault</td>
<td>During self-test, the motor board determined the relative encoder was disconnected.</td>
<td>Acknowledge error. Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>07</td>
<td></td>
<td>Encoder synchronization</td>
<td>The relative encoder and the absolute encoder disagree about the position of the joint more than what is acceptable.</td>
<td>Acknowledge error. Contact Mizuho OSI Services.</td>
</tr>
<tr>
<td>CODE (SD)</td>
<td>CODE (CC)</td>
<td>ERROR MESSAGE</td>
<td>POTENTIAL CAUSE(S)</td>
<td>USER ACTION(S)</td>
</tr>
<tr>
<td>----------</td>
<td>-----------</td>
<td>-----------------------------------</td>
<td>----------------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>30</td>
<td>Chest Tray Errors</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>05</td>
<td>Motor 1 stalled</td>
<td>The motor is not moving when commanded to move.</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>06</td>
<td>Motor 2 stalled</td>
<td>The motor is not moving when commanded to move.</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>09</td>
<td>Motor 1 disconnect</td>
<td>The motor was disconnected while the user was requesting the table to move.</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>0A</td>
<td>Motor 2 disconnect</td>
<td>The motor was disconnected while the user was requesting the table to move.</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>0B</td>
<td>Motor 1 encoder failure</td>
<td>The encoder could not be read while the chest tray was connected.</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>0C</td>
<td>Motor 2 encoder failure</td>
<td>The encoder could not be read while the chest tray was connected.</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>4X</td>
<td>Floor Lock Errors</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>03</td>
<td>Motor 1 overcurrent</td>
<td>The motor was drawing more current than expected.</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>04</td>
<td>Motor 2 overcurrent</td>
<td>The motor was drawing more current than expected.</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>05</td>
<td>Motor 1 unlock timeout</td>
<td>The motor took longer than expected to unlock the floor lock and has not fully unlocked.</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>06</td>
<td>Motor 2 unlock timeout</td>
<td>The motor took longer than expected to unlock the floor lock and has not fully unlocked.</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>07</td>
<td>Motor 1 lock timeout</td>
<td>The motor took longer than expected to lock the floor lock and has not fully locked.</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>08</td>
<td>Motor 2 lock timeout</td>
<td>The motor took longer than expected to lock the floor lock and has not fully locked.</td>
<td>Acknowledge error.</td>
<td></td>
</tr>
<tr>
<td>50</td>
<td>Hand Pendant Errors</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>01</td>
<td>Stuck pendant keypad button</td>
<td>The hand pendant detected a button press during self-test.</td>
<td>Turn table power off and back on. Contact Mizuho OSI Services to replace pendant.</td>
<td></td>
</tr>
</tbody>
</table>
11.1 Software Release Notes

11.1.1 Software Version 1.105

11.1.1.1 Known Software Issues

Item 501 – E-stop generates low current errors

When pressing the Emergency Off button abruptly stopping all table motion, the table will report
warnings on the Hand Pendant. The warnings are of two varieties: Brake disengaged current too
low (Code 04) from one of the motors or Low Drive supply voltage from one of the boards. These
warnings are informational and completely normal. The user can continue operation of the table
after disengaging the Emergency Off button and acknowledging the warning.

Item 607 – In storage mode, table incorrectly boots to the Main screen

The assistant wizard for bringing the table out of the storage configuration is not easily accessed.
The user may access the wizard by going through the Store Table wizard all the way through
Step 8. Note that, since the table is presumably already stored, the steps in the instructions for the
Store Table wizard have already been followed, so the user can quickly acknowledge each of the
Store Table steps without taking mechanical action with the table to get to the instructions for
deploying the table.

Item 593 – Hinging incorrectly locked out upon reboot with trolleys attached

Occasionally directly after the table has been turned on with the Torso Trolley® installed, pressing
the Hinge Up or Hinge Down button will incorrectly indicate that the hinge motion is locked out.
Pressing the Home button briefly to initiate motion will correctly allow hinge motion and thereafter
the Hinge Up and Hinge Down button will operate correctly.

Item 595 – Storage sequence fails to detect single chest trolley attached

The first step in the Store Table wizard reminds the user to remove all accessories from the table.
Once the step is acknowledged, the software detects whether the Supine Top or the Torso Trolley®
is installed and, if so, informs the user that the respective accessory was detected, returning the
user to the first step. If the user only removes one of the two Torso Trolley® support spars from the
table, this condition is not detected.
11.1.2 Software Version 1.110

11.1.2.1 Software Enhancements

Language Support
The Hand Pendant now supports the English, French, Italian, German, and Spanish languages. Through a menu selection, the user may change the display language at any time. The language setting is preserved when the table is powered off.

Units Support
The Hand Pendant now supports displaying the table height in inches or centimeters. Through a menu selection, the user may change the units in which the table height is displayed at any time. The units setting is preserved when the table is powered off.

Independent Monitoring System Notifications
The table notifies the user when the Independent Monitoring System detects that the table is overloaded. Previously, even momentary loading conditions that occur, for example, when a user jumps onto the table configured for prone surgery, could lead to these notifications. The table now reduces the occurrence of these nuisance notifications.

11.1.2.2 Resolved Software Issues

Item 607 – In storage mode, table incorrectly boots to the Main screen
The assistant wizard for bringing the table out of the storage configuration was not easily accessed. This is resolved in version 1.110 so that when the table is powered on in the storage configuration, the hand pendant displays the Ready to Store/Undo screen.

Item 593 – Hinging incorrectly locked out upon reboot with trolleys attached
Previously, occasionally directly after the table has been turned on with the Torso Trolley® installed, pressing the Hinge Up or Hinge Down button would incorrectly indicate that the hinge motion is locked out. This is resolved in version 1.110.

Item 595 – Storage sequence fails to detect single chest trolley attached
The first step in the Store Table wizard reminds the user to remove all accessories from the table. Once the step is acknowledged, the software detects whether the Supine Top or the Torso Trolley® is installed and, if so, informs the user that the respective accessory was detected, returning the user to the first step. Previously, if the user only removes one of the two Torso Trolley® support spars from the table, this condition is not detected. This condition is now detected in version 1.110 and the software informs the user that the Torso Trolley® was detected.

11.1.2.3 Known Software Issues

Item 501 – E-stop generates low current errors
When pressing the Emergency Off button abruptly stopping all table motion, the table will report warnings on the Hand Pendant. The warnings are of two varieties: Brake disengaged current too low (Code 04) from one of the motors or Low Drive supply voltage from one of the boards. These warnings are informational and completely normal. The user can continue operation of the table after disengaging the Emergency Off button and acknowledging the warning.
11.1.3 Software Version 1.111

11.1.3.1 Software Enhancements

New Torso Trolley® Support

The ProAxis® has a newly designed Torso Trolley® for ProAxis® version 1.2. The software supports this new Torso Trolley® as well as the previous generation Torso Trolley®.

11.1.3.2 Resolved Software Issues

There were no issues previously reported as known software issues that are now resolved in this revision.

11.1.3.3 Known Software Issues

Item 501 – E-stop generates low current errors

When pressing the Emergency Off button abruptly stopping all table motion, the table will report warnings on the Hand Pendant. The warnings are of two varieties: Brake disengaged current too low (Code 04) from one of the motors or Low Drive supply voltage from one of the boards. These warnings are informational and completely normal. The user can continue operation of the table after disengaging the Emergency Off button and acknowledging the warning.
11.2 Manual Floor Lock Override

In the event of an error where the floor locks cannot be unlocked through the use of the IntelliPendant® or Auxiliary Panel, the floor locks can be manually unlocked.

To manually raise or lower the floor locks:

1. Remove the grey plastic plugs on the upper surface of each base cover. There are a total of four (4) gray plastic plugs, with two (2) located on the Head-End base cover and two (2) located on the Foot-End base cover (Figure 115).

2. Insert a 13 mm socket with 150 mm minimum extender to capture a hex feature on the top surface of the floor lock motor drive pulley (Figure 116).
   a. Rotate the pulley in a clockwise direction to RAISE the floor locks.
   b. Rotate the pulley in the counter-clockwise directions to LOWER the floor locks.
Figure 116: Use of Socket to Raise/Lower Floor Locks

NOTE: Observe the motion of the floor lock as the pulley is rotated. Once the lock has retracted into the base of the caster, you should stop rotating. Do not over-torque.

3. Remove the socket wrench and reinstall the grey plastic plugs, taking care to ensure they form a water tight seal.

NOTE: Take care to make sure the socket does not detach from the extender during removal.
12 Electrical System

12.1 Description

The electrical system provides control of all the table functions and is comprised of the following:

- Detachable Power Cord
- Power Switch
- Power/Charging Indicator
- Circuit Breakers
- The Emergency Stop button
- IntelliPendant®
- Batteries

12.2 Component Identification and Function

12.2.1 Detachable Power Cord

The table is equipped with a standard IEC Power Cord with the appropriate hospital grade connector. The Power Cord is connected to the table at the IEC power receptacle located on the inside of the Head-End base near the Power Switch. Do not store material near the power inlet nor otherwise impede access to the detachable power cord.

**WARNING:** Ensure the IEC Power Cord provided is utilized and connected to a properly grounded receptacle that meets the input voltage requirements identified on the manufacturer’s label on the Foot-End column of the table. Failure to do so may result in harm to the patient, healthcare professional, or the device.

12.2.2 Power Switch

A Power Switch is located on the Head-End base of the table. When the switch is illuminated green, it indicates that the table is powered ON (Figure 117).

12.2.3 Power/Charging Indicator

Two indicators (located in the Power Inlet Assembly) provide status of the AC and Battery charger. When connected to AC Power, the AC Indicator will illuminate green. The batteries are automatically charged if AC is present. The Battery Indicator will illuminate yellow if the batteries are charging and then turn green when they are fully charged (Figure 117).
12.2.4 Circuit Breakers

Two circuit breakers provide overload protection to the table (Figure 117). They are located in the Power Inlet assembly. The Circuit Breakers pop out if the table experiences power overload. Before proceeding to reset the Circuit Breakers, investigate the cause of the occurrence. To reset the Circuit Breakers, push them in.

12.2.5 The Emergency Stop Button

The Emergency Stop button is a switch that turns off power to all the motors in the table if pressed. For more information on the Emergency Stop button, refer to Section 4.2.

12.2.6 IntelliPendant®

The IntelliPendant® provides primary user interface to the table. The IntelliPendant® has a single processor with software that sends commands to the Main Controller PCA and displays status from the Main Controller PCA.

12.3 Battery System

The ProAxis® Surgery Table uses three (3) sealed, lead-acid, rechargeable batteries in series to provide 30 volts (There are two batteries are 12-volts and one battery at 6-volts).

Normal battery operation is approximately eight (8) hours. This will allow for two procedures averaging six (6) minutes of motion time.

Check the battery for signs of corrosion. The battery terminals should be clean and free from corrosion, oil, grease, dirt, or other contaminants.

Battery Pack ordering number: REF 6988-750

⚠️ CAUTION: Failure to use an approved Mizuho OSI battery voids the warranty and can cause harm to the table.
12.3.1 Charging the Batteries

When the Battery Status symbol on the IntelliPendant® indicates less charge, the batteries require charging. To charge, plug the Power Cord into an appropriate hospital grade AC outlet.

It is imperative that they be charged within 6 months of storage, otherwise permanent loss of capacity might occur as a result of sulfation. Using the table while on AC power will also allow the batteries to charge.

The battery will be fully charged from a completely depleted state in 24 hours. If the battery fails to hold a charge it should be replaced (refer to Section 13).

⚠️ CAUTION: If the table is running on battery and is not charged when this Battery Status Symbol ⚡ displays on the IntelliPendant®, the table will run out of charge and stop functioning.

12.3.2 Replacing the Batteries

When replacing the batteries for the ProAxis® Spinal Surgery Table:

- Replace only with an identical type and size battery pack.
- Replace all three batteries at the same time.
- At a minimum, replace all batteries once every five (5) years.

**NOTE:** Failure to use an approved Mizuho OSI battery voids the warranty and can cause harm to the table.

⚠️ WARNING: Electrical shock hazard exists with access covers removed; use caution when working in exposed areas. Servicing and replacement of batteries should only be performed by trained personnel.

To remove and replace the batteries:

1. Raise the height of the table to its highest position.
2. Ensure that the Power Switch is pressed to OFF and unplug the Power Cord from the wall outlet.

**NOTE:** Disconnect batteries only with Power Cord unplugged and Power Switch pressed to OFF.

3. Remove Cover, Head-End Column, Inner. Next, remove the Cover Assembly, Head-End Base, Left.
4. Below is a picture of the battery pack, as installed on the table.

Figure 119: Battery Pack, Installed on the Table

5. Remove the Base Cover Mount Brackets on both sides of the battery pack.
Figure 120: Removing Base Cover Mount Brackets

6. Before removing the battery pack, remove the Black and Red terminals from the Main Controller PCA Assembly.

Figure 121: Red and Black Battery Terminals
7. Remove the battery pack from the battery compartment by pulling on the black battery strap. The batteries are attached to the base with Velcro®.

8. Install the new battery pack into the battery compartment. The batteries will be attached to the base with Velcro®.

9. Replace the battery pack and reassemble in reverse order.
10. Test to confirm proper battery connection:
   a. Verify that the Power Cord is unplugged from the AC wall outlet.
   b. Press the Power Switch to the ON position (utilizing battery power).
   c. Verify that the Floor Locks are deployed by viewing the Floor Lock symbol ⛽ on the
      IntelliPendant®.
   d. With properly connected batteries, the Battery Status LED on the Power Inlet
      Assembly illuminates, and the IntelliPendant® displays this symbol - ☑️.
      
      • **A green LED** indicates batteries are charged and installed correctly.
      • **A blinking yellow LED** indicates that the table is connected to AC wall outlet and
        the batteries are charging.

Upon receipt, the batteries will have a charge; however, it is required prior to use to charge the batteries. To charge the batteries, plug the Power Cord into wall receptacle. The battery will be fully charged from a completely depleted state in 24 hours.

12.3.3 Lithium Battery

The ProAxis® Spinal Surgery Table also uses a non-rechargeable, lithium CR2032 battery to provide backup power to the real-time clock when the table is not powered. This battery is located on the Main Controller PCA.

**NOTE:** *Should the lithium battery ever need to be replaced, contact Mizuho OSI Services (Section 13).*

This battery contains perchlorate and special handling may apply. Your local regulations may impose special handling of this battery related to disposal. Please follow your local regulations if you dispose of this battery.
13 Technical Support

13.1 Contact for Parts and Service

For detailed repair information or to order replacement parts, call Mizuho OSI Services at 1-800-777-4674 for calls within the USA or +1-510-429-1500 for calls outside the USA. At the prompt, select Option 2.

A Services line is available from 5 AM to 5 PM Pacific Time, Monday through Friday. After business hours, please leave a message.

Please state slowly your name, phone number, your facility name and city, affected equipment model number, and serial number.

An email message may be left anytime at service@mizuhosi.com or through the Web Support Form on the web site: www.mizuhosi.com.

13.2 Further Technical Information

Mizuho OSI will supply circuit diagrams, parts lists, technical descriptions, calibration instructions, or other information necessary to qualified service personnel to service those parts of the ProAxis® Spinal Surgery Table that can be serviced without returning the unit to the manufacturer or distributor. Qualified service personnel are defined as those individuals who have been properly trained and approved by Mizuho OSI to complete service work on the table.

13.3 Order Replacement Parts

If unable to identify a part, please telephone, fax, or email Mizuho OSI Services before placing an order. Once the part number is obtained, follow the instructions below to order the replacement part.

If part number is known, please telephone, fax, or email Mizuho OSI Services with the part number and description for price and availability.

To place a Replacement Parts (RP) order, please telephone, fax, or email Mizuho OSI Services with the part number, description, price, customer number, and method of shipment with the purchase order. Indicate that the order is for Replacement Parts (RP).

13.4 Return Damaged Parts

Identify part or part number to be returned.

Telephone, fax, or e-mail Mizuho OSI Services with the part number and description of the part for return to obtain a Return Goods Authorization (RGA) number.

Complete the Certificate of Decontamination provided by Mizuho OSI Services.

Return the part with RGA number clearly marked on the outside of the package, and include the Certification of Decontamination with the shipment.
13.5 Send a Part for Repair

If unable to identify part, please telephone, fax, or e-mail Mizuho OSI Services for assistance.

If the part number is known, please telephone, fax, or e-mail Mizuho OSI Services with the part number and description of problem to obtain a Repair Authorization (RA) number.

Clean/disinfect parts prior to shipping and complete the Certificate of Decontamination provided by Mizuho OSI Services.

Ship the part with the RA number clearly marked on the outside of the package, and include the Certificate of Decontamination with the shipment.

The part will be evaluated and the customer will be contacted with the cost of repair, if not covered under table warranty.

After customer approval of repair cost, part will be repaired and returned to the customer.

13.6 Warranty

Mizuho OSI guarantees all of its products for a period not to exceed one (1) year from date of invoice for defects in material and workmanship with the exception of products that are misused, altered, or damaged. All Tempur-Pedic® medical pads come with a two (2) year warranty against manufacturer defect. Defective merchandise will be credited or replaced.

13.7 European Union EC Representative

Emergo Europe
Molenstraat 15
2513 BH The Hague
The Netherlands
Tel: (31) (0) 70 345-8570
Fax: (31) (0) 70 346-7299
# 14 Appendix

## 14.1 Electromagnetic Emissions

The ProAxis® Spinal Surgery Table is intended for use in the electromagnetic environment specified below. The customer or the user of the ProAxis® Spinal Surgery Table should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Emissions test</th>
<th>Compliance</th>
<th>Electromagnetic environment - guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions</td>
<td>Group 1</td>
<td>The ProAxis® Spinal Surgery Table uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>CISPR 11</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RF emissions</td>
<td>Class A</td>
<td>The ProAxis® Spinal Surgery Table is suitable for use in all establishments other than domestic and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>CISPR 11</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Harmonic emissions</td>
<td>Class A</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Voltage fluctuations /</td>
<td>Complies</td>
<td></td>
</tr>
<tr>
<td>flicker emissions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-3</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
# 14.2 Electromagnetic Immunity

## Guidance and manufacturer's declaration – electromagnetic immunity

The ProAxis® Spinal Surgery Table is intended for use in the electromagnetic environment specified below. The customer or the user of the ProAxis® Spinal Surgery Table should ensure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601 test level</th>
<th>Compliance level</th>
<th>Electromagnetic environment - guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic discharge (ESD)</td>
<td>± 6 kV contact</td>
<td>± 6 kV contact</td>
<td>Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.</td>
</tr>
<tr>
<td>IEC 61000-4-2</td>
<td>± 8kV air</td>
<td>± 8kV air</td>
<td></td>
</tr>
<tr>
<td>Electrical fast transient / burst</td>
<td>± 2kV for power supply lines</td>
<td>± 2kV for power supply lines</td>
<td>Mains power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>IEC 61000-4-4</td>
<td>± 1kV for input / output lines</td>
<td>± 1kV for input/output lines</td>
<td></td>
</tr>
<tr>
<td>Surge</td>
<td>± 1 kV line(s) to line(s)</td>
<td>± 1 kV line(s) to line(s)</td>
<td>Mains power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>IEC 61000-4-5</td>
<td>± 2 kV line(s) to earth</td>
<td>± 2 kV line(s) to earth</td>
<td></td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines</td>
<td>&lt;5% $U_T$ (&gt;95% dip in $U_T$) for 0.5 cycle</td>
<td>&lt;5% $U_T$ (&gt;95% dip in $U_T$) for 0.5 cycle</td>
<td>Mains power quality should be that of a typical commercial or hospital environment. If the user of the ProAxis® Spinal Surgery Table requires continued operation during power mains interruptions, it is recommended that the ProAxis® Spinal Surgery Table be powered from an uninterruptible power supply or a battery.</td>
</tr>
<tr>
<td>IEC 61000-4-11</td>
<td>40% $U_T$ (60% dip in $U_T$) for 5 cycles</td>
<td>40% $U_T$ (60% dip in $U_T$) for 5 cycles</td>
<td></td>
</tr>
<tr>
<td></td>
<td>70% $U_T$ (30% dip in $U_T$) for 25 cycles</td>
<td>70% $U_T$ (30% dip in $U_T$) for 25 cycles</td>
<td></td>
</tr>
<tr>
<td></td>
<td>&lt;5% $U_T$ (&gt;95% dip in $U_T$) for 5 s</td>
<td>&lt;5% $U_T$ (&gt;95% dip in $U_T$) for 5 s</td>
<td></td>
</tr>
<tr>
<td>Power frequency (50/60 Hz) magnetic field</td>
<td>3 A/m</td>
<td>3 A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>IEC 61000-4-8</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: $U_T$ is the a.c. mains voltage prior to application of the test level.
Guidance and manufacturer’s declaration – electromagnetic immunity

The ProAxis® Spinal Surgery Table is intended for use in the electromagnetic environment specified below. The customer or the user of the ProAxis® Spinal Surgery Table should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601 test level</th>
<th>Compliance level</th>
<th>Electromagnetic environment - guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF</td>
<td>IEC 61000-4-6</td>
<td>3 Vrms 150 kHz to 80 MHz</td>
<td>3 V</td>
</tr>
<tr>
<td>Radiated RF</td>
<td>IEC 61000-4-3</td>
<td>3 V/m 80 MHz to 2.5 GHz</td>
<td>3 V/m</td>
</tr>
</tbody>
</table>

\[
d = 1.2 \sqrt{P} \\
\]

80 MHz to 800 MHz

\[
d = 2.3 \sqrt{P} \\
\]

800 MHz to 2.5 GHz

Where \( P \) is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and \( d \) is the recommended separation distance in meters (m).

Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, a should be less than the compliance level in each frequency range. b

Interference may occur in the vicinity of equipment marked with the following symbol:

Note 1: At 80 MHz and 800 MHz, the higher frequency range applies.
Note 2: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

a Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the ProAxis® Spinal Surgery Table is used exceeds the applicable RF compliance level above, the ProAxis® Spinal Surgery Table should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the ProAxis® Spinal Surgery Table.

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

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

<table>
<thead>
<tr>
<th>Rated maximum output power of transmitter (W)</th>
<th>Separation distance according to frequency of transmitter (m)</th>
<th>150 kHz to 80 MHz</th>
<th>80 MHz to 800 MHz</th>
<th>800 MHz to 2.5 GHz</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.01</td>
<td>d = 1.2 \sqrt{P}</td>
<td>0.12</td>
<td>0.12</td>
<td>0.23</td>
</tr>
<tr>
<td>0.1</td>
<td></td>
<td>0.37</td>
<td>0.37</td>
<td>0.74</td>
</tr>
<tr>
<td>1</td>
<td></td>
<td>1.2</td>
<td>1.2</td>
<td>2.3</td>
</tr>
<tr>
<td>10</td>
<td></td>
<td>3.7</td>
<td>3.7</td>
<td>7.4</td>
</tr>
<tr>
<td>100</td>
<td></td>
<td>12</td>
<td>12</td>
<td>23</td>
</tr>
</tbody>
</table>

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

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

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