OAKWORKS®
Spine Positioning System II

CRESCENT FACE PAD

CONToured torso support pad

RADiOLUCENT FRAME WITH ADJUSTABLE FACE REST

CONToured torso wedge

8" x 22" x 2"
(20 x 56 x 5 cm)
LARGE RECTANGULAR ADJUSTER PAD

7" x 12" x 1½"
(18 x 30 x 4 cm)
SMALL RECTANGULAR ADJUSTER PAD

CARRY CASE

8" (20cm) SEMI-ROUND BOLSTER

www.oakworksmed.com · 717.235.6807

made in the USA with US & imported parts
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INTRODUCTION / PRODUCT USE DESCRIPTION / SYMBOL IDENTIFICATION

INTRODUCTION

The Spine Positioning System II is an integral component of the pain management fluoroscopy suite. With this system procedural set up time is reduced, patient comfort is enhanced and unwanted movement is minimized. Most importantly, the target anatomy is more readily visualized which allows the physician to perform spine procedures in a more efficient and secure manner. In collaboration with leading pain management physicians, Oakworks designed the Spine Positioning System II in an effort to achieve the critical balance between optimal imaging and patient comfort. The radiolucent adjustable frame and versatile padding system provide a metal free imaging support platform capable of quickly positioning a wide variety of patient physiques for extended periods of time. The adjustable face rest position provides individualized positioning for all types of cervical procedures and anatomy. The contoured torso support pad is complimented by a host of uniquely shaped and sized adjuster pads and wedges that enable a multitude of positioning combinations for ideal patient comfort and imaging needs for all spinal column procedures.

PRODUCT USE DESCRIPTION

The Oakworks® Spine Positioning System II is a patient cradle device for use in diagnostic and therapeutic procedures of the spine. It is intended to be used by a healthcare professional in a medical environment solely for the purpose of aiding in patient positioning and comfort during non-surgical imaging or spinal injection procedures. It may also be used during minimally invasive surgical procedures such as vertebroplasty or kyphoplasty. The Spine Positioning System II, its secondary components, and optional components are suitable for use in fluoroscopy suites. No special training is required but a review of the following Safety Instructions is important for the safety of the operator and patient. The healthcare professional should read and understand this entire manual before use with a patient.

SYMBOL IDENTIFICATION

This symbol, when used in this manual and on product labels, represents a caution warning. Be sure to read and comply with all precautions and warnings.

This symbol, when used in this manual and on product labels, indicates the potential of exposure to harmful x-rays. Be sure to read and comply with all warnings.

This symbol when used in this manual or on product labels, warns that when stacking containers during transport and storage, there should be do not stack more than 5 containers high.

This symbol, when used in this manual or on product labels, indicates that the product should be protected from moisture. The humidity specifications for Transport & Storage are listed on page 21.

This symbol, when used in this manual or on product labels, indicates that information is given regarding the recommended temperature limits during transport and storing.

This symbol, when used in this manual or on product labels, indicates the date of manufacture of the device.

This symbol is used to indicate that the operator should consult the user manual.
A patient safety strap is required during all procedures. Follow normal and required safety protocol for all procedures where the patient is in an elevated position for the procedure (straps, attendants, etc.). Always be certain that attending staff is aware of the patient’s position while the device is in use. Reposition the patient if necessary to promote stability. Due to the increased distance between the patient and the table surface, additional safety measures are recommended when the table top is not used in a level position due to the risk of the patient falling off the table.

The Oakworks® Spine Positioning System II is not designed for use with diagnostic x-ray systems where the x-ray generator is located above the radiographic table and the film cassette or image intensifier is located below the radiographic table. The X-Ray generator must be located below the radiographic table. The Spine Positioning System II is not designed for use with magnetic resonance imaging systems. The Spine Positioning System II is not intended for use in cranial procedures.

Do not overhang the radiolucent frame beyond the warning line on the frame.

Operate the C-arm of the fluoroscopy system with the Spine Positioning System II in place before using the device with a patient for the first time. Make sure there is adequate clearance to permit free C-arm rotation for both the patient and the positioning device.

Do not permit the patient to push down on the Crescent Face Pad in an effort to lift themselves up while dismounting the platform and/or the table.

The Spine Positioning System II should generally not be used when a patient is under general anesthesia, especially when prolonged cases are performed. This will reduce the risk of ocular or facial nerve injury.

The cushioning foam contained within the Torso Support will lose its ability to spring back to the original position over time and the amount of foam compression will increase. Therefore, the Torso Support should be replaced periodically to ensure the device functions as intended.

To prevent the potential of cross-contamination, it is strongly advised to use barrier techniques when the device is in use. A disposable or laundered patient gown, or disposable pad are satisfactory for use as a barrier for the Torso Support and other components and accessories, except when the patient presents with pathology that would indicate otherwise. A disposable face rest cover should be used to cover the Face Rest Pad. Contact Oakworks for ordering information. Barrier techniques should be used in addition to disinfection procedures, not in lieu of them.

Be sure to support the weight of the patient’s head while making adjustments to the cervical positioning feature of the Platform Frame. Make sure all cam locks are secure before relinquishing support of the positioning assembly.

The Cervical Support System has metal parts that can cause back scatter of x-rays, see Product Description for photo.

When x-rays are present, wear a suitable radiation barrier.

The Spine Positioning System II is constructed using metal pins in the Quick Cam Locks and aluminum tubing in the support structure. These are out of the field of view in most A-P and oblique tilted views. Place the positioning assembly according to the recommendations in the directions for use to eliminate, or reduce any artifacts. If artifacts still remain to the extent that they would compromise the efficacy of needle placement, discontinue use of the device during the affected procedure.

The Spine Positioning System II is designed to be a standalone product used with radiographic equipment. It must not be modified or incorporated into any other equipment.

All materials used in the construction of the device and accessories are safe for temporary and moderately frequent human contact. The device is not intended for prolonged contact.

Do not use the Face Rest Support Arms as a handle to carry the Spine Positioning System II.

Follow maintenance instructions found near the end of this manual. Mechanical components should be checked periodically to insure that they are functioning properly to insure the safety of the patient.

SPS II weight limit: 350 lbs. (159 kg.) Crescent Face Pad Support weight limit: 25 lbs. (11 kg.)
PRODUCT DESCRIPTION

Spine Positioning System II

**STANDARD SPECIFICATIONS**

<table>
<thead>
<tr>
<th>Specification</th>
<th>Measurement</th>
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<tbody>
<tr>
<td>Weight</td>
<td>16 lbs. (7 kg.)</td>
</tr>
<tr>
<td>Frame with Face Rest</td>
<td>12” (30 cm.) Wide x 32.5” (84 cm.) Long</td>
</tr>
<tr>
<td>Crescent Face Pad</td>
<td>12” (30 cm.) diameter</td>
</tr>
<tr>
<td>Contoured Torso Support Pad</td>
<td>6.5” x 23” x 30” (17 x 58 x 76 cm.)</td>
</tr>
<tr>
<td>Contoured Torso Wedge</td>
<td>22” x 29” x 2” (56 x 74 x 5 cm.)</td>
</tr>
<tr>
<td>Large Rectangular Adjuster Pad</td>
<td>8” x 22” x 2” (20 x 56 x 5 cm.)</td>
</tr>
<tr>
<td>Small Rectangular Adjuster Pad</td>
<td>7” x 12” x 1.5” (18 x 30 x 4 cm.)</td>
</tr>
<tr>
<td>8” (20 cm.) Semi-Round Bolster</td>
<td>6” x 8” x 26” (15 x 20 x 66 cm.)</td>
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<tr>
<td>Carry Case</td>
<td>Transports the SPS II System</td>
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<td>Warranty</td>
<td>2 years - Frame, Fabric and padding</td>
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<tr>
<td>Safety Listings</td>
<td>FDA and CE marked</td>
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</table>
PRODUCT DESCRIPTION

RADIOLUCENT FRAME
Used to support the Torso Support and Crescent Face Pad. One cam lock facilitates cervical flexion and extension.

Crescent Face Pad
The Crescent Face Pad supports the patient’s face in a prone position without compromising air space for breathing. The face pad can be moved in situations to prevent imaging of the locking mechanism when performing upper cervical procedures that require substantial imaging angulation.

Contoured Torso Support Pad
The Contoured Torso Support is constructed of dense foam in the center, flanked by softer foam. The softer foam accommodates to the patient’s shoulders and/or breasts to maximize comfort. This helps provide enhanced patient stability while allowing for the shoulders to descend for optimal cervical and thoracic imaging.

The Distal end of the torso support pad is hollowed out under the abdomen to enhance patient comfort and stability. Additionally, the Distal end of the torso support pad is wider to enhance patient stability by reducing sway while in the device.
PRODUCT DESCRIPTION

CONToured TORSO WEDGE

The Contoured Torso Wedge is constructed of dense foam. This provides enhanced patient stability and conveniently reduces shoulder interference during cervical procedures.

SMALL RECTANGULAR ADJUSTER PAD

The 7” x 12” (18 x 30 cm.) Small Rectangular Adjuster pad is used to reduce lumbar lordosis and/or increase chest height to allow for shoulders to naturally descend out of the plane of the cervical and thoracic spine. This pad offers a wide range of flexibility for general patient positioning and stabilization.

LARGE RECTANGULAR ADJUSTER PAD

The wider 8” x 22” (20 x 56 cm.) Large Rectangular Adjuster pad can be used to allow those with a shorter humerus to allow the forearm and elbow to rest and stabilize on a flat surface. Additionally, this can be used as the Small adjuster pad is utilized, with a wider support.

8” (20 cm.) SEMI-ROUND BOLSTER

This bolster may be placed under the patient’s ankles to enhance positioning stability.
PREPARATION FOR USE

**CAUTION**

Do not overhang the platform frame beyond the WARNING line on the frame.

Unpack and inspect all components. Identify the components and their use with the pictures located in the Product Description Section of this manual.

All components are shipped in a clean but not sterile condition. If the Spine Positioning System II will be used for an indicated surgical procedure, be sure to disinfect the components prior to use. Disinfectants that can be used are described in the Cleaning & Disinfecting Section of this manual.

FACE REST PLATFORM ADJUSTMENT

Step 1 - Open cam

Step 2 - Grasp platform and raise to desired position

Step 3 - Begin to close the cam making sure that the small locking pins enter corresponding positioning holes (Minor platform “rocking” may be necessary for the pins to enter the holes). Do not force the cam to close.

Step 4 - Continue to close cam.

Step 5 - Close cam to the final position.

Step 6 - Double check platform by applying downward force to ensure your face rest platform is securely locked.
**DIRECTIONS FOR USE**

**TORSO PAD STRAP**

**WARNING**  
A patient safety strap must be used during all procedures.

To secure the Torso Pad to the table, wrap the Velcro® strap under the table top and attach securely.

**TRANSPORTING THE SPINE POSITIONING SYSTEM II**

Open the cam lock on the adjustable face rest and rotate the face rest flat against the base frame. This will protect the face rest support platform during transport.

When placing the Spine Positioning System II in the Carry Case, put some pads, wedges or bolsters on both sides of the base frame.
The following imaging scenarios of patients will demonstrate:
1. Various body types using the Spine Positioning System II (SPS II)
2. Their positioning and specific configurations of the SPS II used in particular clinical situations
3. Various fluoroscopic images of these factitious patients that exemplify the value of the SPS II

PATIENT - ALICIA

Components used: Radiolucent Frame, Crescent Face Pad, Contoured Torso Support Pad, Contoured Torso Wedge, Small Adjuster Pad, 8” Semi-Round Bolster (not pictured)
**Patient - Don**

Components used: Radiolucent Frame, Crescent Face Pad, Contoured Torso Support Pad, Large Adjuster Pad, 8” Semi-Round Bolster (not pictured)

SPS II set up for Don shown here

Don in the SPS II while obtaining an oblique image of the lumbar spine

Lateral image of the lumbar spine

Right oblique image of the lumbar spine

AP image of the lumbar spine

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**Imaging Scenarios**

**Warning**

A patient safety strap must be used during all procedures.
**IMAGING SCENARIOS**

**WARNING**

A patient safety strap must be used during all procedures.

**PATIENT - LIZ**

Components used: Radiolucent Frame, Crescent Face Pad, Contoured Torso Support Pad, Contoured Torso Wedge, 8” Semi-Round Bolster (not pictured)

Liz in the SPS II while obtaining an AP image of the upper thoracic spine.

Liz in the SPS II while obtaining a lateral image of the upper thoracic spine.

AP image visualizing the T1-2 interlaminar space.

Lateral image primarily through the C7—T2 segments.

Contralateral oblique showing the upper thoracic facet joints.
!! WARNING!!

A patient safety strap must be used during all procedures.

PATIENT - MARY

Components used: Radiolucent Frame, Crescent Face Pad, Contoured Torso Support Pad, Contoured Torso Wedge, 7” x 12” Rectangular Adjuster Pad, 8” Semi-Round Bolster (not pictured)

SPS II set up for Mary shown here

Mary in the SPS II while obtaining an AP image through the C1-2 segment

AP image through the C1-2 joints

Lateral image through the C1-3 segments
**WARNING**

A patient safety strap must be used during all procedures.

**PATIENT - CARL**

Components used: Radiolucent Frame, Crescent Face Pad, Contoured Torso Support Pad, Contoured Torso Wedge, 8” Semi-Round Bolster (not pictured)

![SPS II set up for Carl shown here](image)

Carl in the SPS II while obtaining an AP image of the mid-thoracic spine

Right thoracic oblique image to visualize the trajectory for a transforaminal injection

AP image of the mid-thoracic spine for planning the trajectory for a left thoracic facet injection

Contralateral oblique showing the trajectory for targeting the mid-thoracic facet joint
A patient safety strap must be used during all procedures.

**PATIENT - DEBBIE**

Components used: Radiolucent Frame, Crescent Face Pad, Contoured Torso Support Pad, Contoured Torso Wedge, Small Adjuster Pad, 8” Semi-Round Bolster (not pictured)

Debbie in the SPS II while obtaining a lateral cervical image.

AP image of the cervical spine while imaging through the C7-T1 interlaminar space.

Complete lateral image of the cervical spine including the C7-T1 segment.

Contralateral oblique of the cervical spine.
A patient safety strap must be used during all procedures.

PATIENT - JANE

Components used: Radiolucent Frame, Crescent Face Pad, Contoured Torso Support Pad, Contoured Torso Wedge, 8” Semi-Round Bolster (not pictured)

Jane in the SPS II while obtaining a lateral image of the cervical spine.

The lower cervical interlaminar spaces are seen without visualization of the mandible over the target interspaces.

Lateral collimated image of the cervical spine. The C6-7-T1 interspaces are appreciated for all posterior approach cervical procedures such as interlaminar epidural steroid injections, facet injections, medial branch blocks and medial branch radio frequency neurotomy.
**CLEANING & DISINFECTION / INSPECTIONS & MAINTENANCE**

### CLEANING & DISINFECTION

**WARNING** Before cleaning with any liquid cleaner be sure to unplug the power cord from the outlet.

Use a 10% sodium hypochlorite (bleach) solution or *Recommended Disinfectants* on all surfaces. Clean all sides of each upholstered section. Follow the directions on the disinfectant and wipe off excess.

*Recommended Disinfectants*

Protex, MadaCide, Accell TB, Virox®

**Note:** Damage caused by unapproved substances will not be covered under the warranty.

*DO NOT* use citrus based cleaners or other strong cleaners, such as alcohol, acetone, higher concentrations of bleach or other products that contain high concentrations of these substances.

*DO NOT* expose the fabric to temperatures below 50°F/10°C or above 104°F/40°C.

*DO NOT* expose the fabric to direct sunlight, adhesives, liquids, or abrasive materials.

### INSPECTIONS & MAINTENANCE

Inspect Torso Support Pad monthly to be sure that the foam has not lost shape or firmness to the extent that patient support would be compromised.

Inspect the base and components monthly to ensure that they have not been damaged. Replace any damaged or worn components.

Inspect face rest platform locking mechanism weekly. Use the following procedure:

1. **Step 1 - Lock the platform cam**
2. **Step 2 - Rock platform up & down**

   Gently rock platform up and down and note any “looseness” (some flexing is normal). Look for gaps between the aluminum parts. If you feel “looseness” or see gaps, see Face Rest Platform Cam Tightening.

Inspect joints:

- **Bad (gap)**
- **Good (no gap)**
1. Use 1/2” socket wrench to grasp the locknut.
2. Hold the cam with other hand.
3. Tighten the cam until there is no gap between the 2 metal parts.

WARRANTY

View complete warranty details at www.oakworks.com

PRODUCT SPECIFICATIONS

<table>
<thead>
<tr>
<th>Component</th>
<th>Aluminum Equivalence</th>
<th>Dimensions</th>
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<tbody>
<tr>
<td>Radiolucent Frame</td>
<td>1.20 mm @ 100 kVp, HVL of 3.6 mm</td>
<td>1/4” x 12” x 32.5” (.6 x 30 x 84 cm.)</td>
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<tr>
<td>Crescent Face Pad</td>
<td>.72 mm @ 100 kVp, HVL of 3.6 mm</td>
<td>12” (30 cm.) diameter</td>
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<tr>
<td>Contoured Torso Support Pad</td>
<td>1.10 mm @ 100 kVp, HVL of 3.6 mm</td>
<td>23” x 30” x 6.5” (58 x 76 x 17 cm.)</td>
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<td>Contoured Torso Wedge</td>
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<td>Small Adjuster Pad</td>
<td>.35 mm @ 100 kVp, HVL of 3.6 mm</td>
<td>7” x 12” x 1.5” (18 x 30 x 4 cm.)</td>
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<tr>
<td>Large Adjuster Pad</td>
<td>.35 mm @ 100 kVp, HVL of 3.6 mm</td>
<td>8” x 22” x 2” (20 x 56 x 5 cm.)</td>
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<tr>
<td>8” (20 cm.) Semi-Round Bolster</td>
<td>N/A</td>
<td>6” x 8” x 26” (15 x 20 x 66 cm.)</td>
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ENVIRONMENTAL CONDITIONS

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<tr>
<th>Conditions</th>
<th>Temperature</th>
<th>Humidity</th>
<th>Atmospheric Pressure</th>
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<tr>
<td>Normal Use</td>
<td>50° (10°C) to 104° (40°C)</td>
<td>20% to 60% RH</td>
<td>98 to 105 kPa</td>
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<td>Storage &amp; Transport</td>
<td>-20° (-29°C) to 135° (57°C)</td>
<td>20% to 95% RH</td>
<td>98 to 105 kPa</td>
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</table>
USER MANUAL

OAKWORKS®
Spine Positioning System II

CONTACT INFORMATION:

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FDA Listed