

USER MANUAL

OAKWORKS®

Arm Hammock



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PRODUCT USE DESCRIPTION / IMPORTANT SAFETY INSTRUCTIONS

PRODUCT USE DESCRIPTION

The Arm Hammock is used to comfortably and easily support the patients arms in the prone position during procedures. It is for use on Imaging/Pain Management tables with the Integrated Headrest tabletop style. It is intended to be used by a healthcare professional in a medical environment. 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, warns against an electrical shock hazard. Be sure to observe and comply with all 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 and on product labels, indicates that the table and components are a Type B Applied Part pursuant to IEC 60601-1.



This symbol, when used in this manual or on product labels, indicates a Protective Earth (Ground) Terminal.



This symbol, when used in this manual and on product labels, indicates the name and address of the manufacturer.



This symbol, when used in this manual or on product labels, indicates the country of manufacture along with date of manufacture of the device next to it.



This symbol, when used in this manual or on product labels, indicates alternating current (AC).



This symbol, when used in this manual or on product labels, indicates direct current (DC).



This symbol is used to indicate that the operator should consult the user manual.



Sitting is prohibited in this area.

CONTRAINDICATIONS

There are no known contraindications to the use of this equipment.

IMPORTANT SAFETY INSTRUCTIONS



CAUTION READ AND SAVE THESE INSTRUCTIONS

The Arm Hammock is not designed to support the entire weight of the patient. Do not allow the patient to use in a manner to adjust themselves using their weight on the table or aid in the dismounting of the table.

The Arm Hammock is designed to be a standalone device. It must not be modified or incorporated into any other equipment.

It is the responsibility of the operator of this equipment to ensure the proper operation of this accessory and it is very important for the safety of the operator and patient. Directions for use of this equipment are described in this manual. The operator should read these sections carefully.

Weight Limit: (patient's arms) 30 lbs. / 13 kg. Do not exceed.

Disposal of waste products, residues, etc., and accessories at the end of the expected service life are listed in the Med-RA-PM-71 Imaging and Pain Management Tables User Manual listed on www.oakworksmed.com.

To reduce the risk of injury to persons:

1. Never operate this device if it is not working properly, or if it has been damaged. Contact OAKWORKS® Customer Service before use if damage is found.
2. Do not use it outdoors.



MISE EN GARDE

LIRE ET CONSERVER CES INSTRUCTIONS

Le hamac à bras n'est pas conçu pour supporter tout le poids du patient. Ne laissez pas le patient s'utiliser de manière à s'ajuster en utilisant son poids sur la table ou à l'aider à démonter la table.

Le Arm Hammock est conçu pour être un appareil autonome. Il ne doit pas être modifié ou incorporé à tout autre équipement.

Il est de la responsabilité de l'opérateur de cet équipement de s'assurer du bon fonctionnement de cet accessoire et c'est très important pour la sécurité de l'opérateur et du patient. Les instructions d'utilisation de cet équipement sont décrites dans ce manuel. L'opérateur doit lire attentivement ces sections.

Limite de poids : (les bras du patient) 30 livres. / 13 kg. Ne dépasse pas.

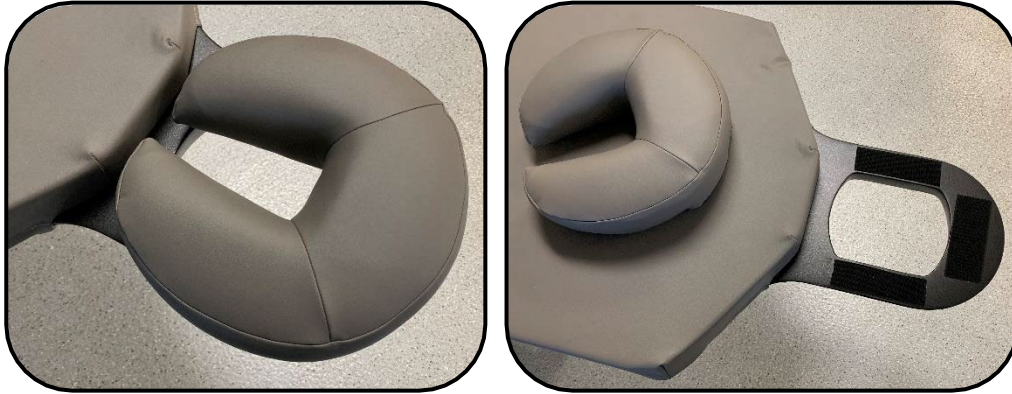
L'élimination des déchets, résidus, etc. et des accessoires à la fin de la durée de vie prévue est répertoriée dans le manuel d'utilisation des tables d'imagerie et de gestion de la douleur Med-RA-PM-71 répertorié sur www.oakworksmed.com.

Pour réduire le risque de blessures aux personnes :

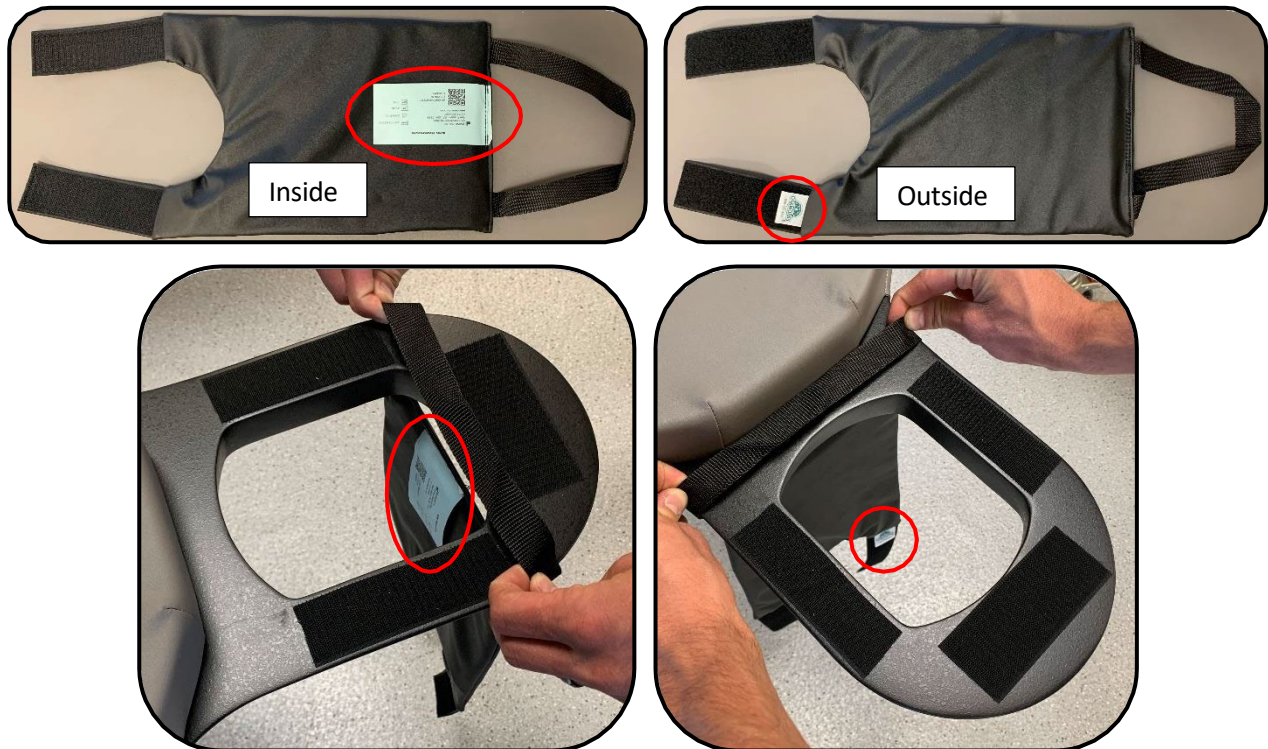
1. N'utilisez jamais cet appareil s'il ne fonctionne pas correctement ou s'il a été endommagé. Contactez le service client OAKWORKS® avant utilisation si des dommages sont constatés.
2. Ne l'utilisez pas à l'extérieur.

ARM HAMMOCK ATTACHMENT INSTRUCTIONS

- 1) Remove the Face Rest Pad from the head end of the Imaging and Pain Management table.



- 2) Slip the looped end of the Arm Hammock around the exposed end of the top as shown. The blue label should be facing inside while the small Oakworks logo tag should be facing outside.



ARM HAMMOCK ATTACHMENT INSTRUCTIONS

- 3) Flip up Arm Hammock and attach the Velcro to the Velcro on the exposed tabletop. If the Velcro does not latch, then the Arm Hammock is backwards.



- 4) Re-attach the Face Rest Pad onto the head end of the table using the Velcro from the Arm Hammock.



- 5) The Arm Hammock is ready for use.



CLEANING & DISINFECTION

RECOMMENDED CLEANERS/DISINFECTANTS

Reference the Recommended Cleaners and Disinfectant list (MMINML0008-EN) that came with the table. This information can also be found at www.oakworksmed.com under product information.

All cleaners and disinfectants have the ability to degrade the upholstery to some extent. However, following the recommended cleaner and disinfectant list and cleaning process will provide the best care for your table and support a long product life.

OAKWORKS® recommends a prepackaged wipe for cleaners/disinfectants to ensure best distribution of disinfectant for the required kill time, without leaving excess residue and/or overexposing components therefore minimizing the potential for damage to materials. Please read and follow disinfectants manufacturers' directions for cleaning and disinfection.

OAKWORKS® does NOT recommend the use of cleaners/disinfectants containing Hydrogen Peroxide, Acetic Acid, or Phenolics. These chemicals can cause damage to the appearance and/or material integrity of various components. Also, while the recommended cleaners/disinfectants list includes products containing Quaternary Ammonium compounds (“quats”), not all products containing quats are approved for use. Some contain additional detergents and/or surfactants which can damage some materials.

Use of non-approved cleaners or disinfectants may lead to damage to upholstery and other materials found on the table and will void the warranty.

CLEANING PROCESS

Follow the cleaners/disinfectant manufacturers' directions for use. Please note that cleaning and disinfecting an OAKWORKS® table is a two-part process. First it must be cleaned of any visible soil, then it can be disinfected. Please follow this procedure for best results:

1. Using an approved cleaner or mild liquid soap and water, clean any visible soil off the Arm Hammock. It is recommended that the upholstery be cleaned at least once a week to prevent disinfectant build-up.
2. Rinse with clean water and dry with a clean cloth or towel.
3. Using an approved disinfectant, thoroughly disinfect all surfaces of the Arm Hammock, making sure they remain wet for the disinfectant manufacturer's recommended contact time. Do not allow disinfectant to pool on the upholstery after the recommended contact time.
4. Wipe off any excess liquid with a cloth or towel and clean water.
5. Dry all surfaces with a clean cloth or towel.

Avoid using writing instruments or other similar instruments around the upholstery as it can cause permanent staining. If this does occur, do not wipe with an alcohol based cleaner. Instead, blot the stain with a clean cloth/ paper towel. Use a recommended cleaner or disinfectant to remove the stain. Follow this with a rinse of clean water.

INSPECTIONS & WARRANTY

INSPECTIONS

RECOMMENDED REGULAR INSPECTIONS (monthly or local standard)

- Check for damage to the fabric and Velcro straps.
- Visually inspect for obvious damage that could cause problems during operation.

RECOMMENDED PERIODIC INSPECTIONS (yearly or local standard)

- Visually inspect the fabric and Velcro straps components for obvious damage that could cause problems during operation.
- Check that all fasteners are present and can fasten securely.
- Clean unusual buildup of dirt on the Arm Hammock not normally cleaned on a regular basis.
- Check for tears in the seams.

WARRANTY

View complete warranty details at www.oakworksmid.com under the “INFO” tab, “Warranty and Return Policy”.

UNIQUE DEVICE IDENTIFICATION (UDI) INFORMATION



Arm Hammock ← Model Name

Manufacturer Info →

Manufacturer: Oakworks, Inc.
923 East Wellspring Road
New Freedom, PA, USA 17349
+1 717-235-6807
www.oakworks.com

GTIN 00817463021325
Manuf. date: 2016-10-26
SN FL4636744
MD QTY: 1 EA

(01)00817463021325(11)20161026(21)FL4636744

GTIN Manufacture Date Serial Number

Unique Device Information:

- GTIN - 14 digit number unique for each variation of a model
- Manufacture Date - Country of manufacture and date the device goes into production in YYYY-MM-DD format
- SN - Serial Number
- MD - Medical Device Symbol
- QTY - Quantity of the product

SPECIFICATIONS

PRODUCT SPECIFICATIONS

Weight	0.5 lbs. (0.2 kg.)
Shipping Weight	1 lb. (0.4 kg.)
Support Capacity	30 lbs. (13 kg.)

ENVIRONMENTAL CONDITIONS

CONDITIONS	TEMPERATURE	HUMIDITY	ATMOSPHERIC PRESSURE
Normal Use	50° (10°C) to 104° (40°C)	20% to 60% RH	98 to 105 kPa
Storage & Transport	-20° (-29°C) to 135° (57°C)	20% to 95% RH	98 to 105 kPa

USER MANUAL

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CoYoMe B.V.

Edisonstraat 4

3261 LD Oud-Beijerland, The Netherlands

www.coyome.eu

Enrico Cohen

Enrico@coyome.nl

info@coyome.nl

Phone: +31 613.886.424

www.oakworksmed.com



EMERGO EUROPE
Westervoortsedijk 60
6827 AT Arnhem
The Netherlands



Notice: Any serious incident that has occurred in relation to the device should be reported to OAKWORKS®, Inc. and to the competent authority of the Member State in which the user and/or patient is established.

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