PMI Single Use REM Grounding Pad w/ Attached Cord

ITEM NUMBER P4500

Section and Preparation of Placement Site
1. Select a well-vascularized, muscular, and convex site in proximity to the surgical site. Do not place over scar, inflamed skin, fatty tissue, bony prominences, metal prostheses, ECG electrodes and cables, pacemakers, or fluids may pool.
2. Shave all hair from selected site. Dry skin thoroughly.

Placement of Electrode
1. Remove the backing from the pad.
2. Apply to skin starting from one end of the pad and smooth out to the opposite end.
3. Apply finger pressure on the adhesive border and massage entire pad area to ensure adequate contact with the skin.
4. At the completion of the surgical procedure, remove the pad slowly and with care to avoid skin trauma.

WARNING
If the patient is repositioned for the surgical procedure, verify that the pad remains in proper contact with the skin and the cable remains connected to the pad. Failure to do so may result in patient burns.

WARNING
During the surgical procedure, assure that no object obstructs the pad site.

WARNING
If higher than normal power settings are required, a problem may exist. Before increasing the power setting, verify that the pad has full contact with the patient's skin. Check the cable and connectors and inspect all active accessories.

WARNING
Do not relocate the pad after initial application.

WARNING
Do not reuse the pad.

WARNING
Do not cut or modify the pad in any way.

WARNING
Review and follow any additional instructions, warnings, and precautions in the Operating Instructions for the generator in use.

Warranty
The manufacturer warrants that reasonable care has been used in the manufacture of this device. This warranty is in lieu of and excludes all other warranties not expressly set forth herein, whether expressed or implied by operation of law or otherwise, including but not limited to any implied warranties of MERCHANTABILITY OR FITNESS, since handling, storage, cleaning and sterilization of this device as well as factors relating to the patient, his diagnosis, treatment, surgical procedures and other matters beyond the manufacturer's control directly affect this device and the results obtained from its use. The manufacturer shall not be liable for any incidental or consequential loss, damage or expense, directly or indirectly, arising from the use of this device other than replacement of all or part of it. The manufacturer neither assumes liability for it, any other or additional liability or responsibility in connection with this device. Because of continuing product improvements, prices, specifications and model availability are subject to change without notice.

Distributed by:
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Single Use
Non-Sterile

NOT MADE WITH NATURAL RUBBER LATEX
Professional Use Only
Read Instructions Before Use

CAUTION: U.S. Federal law restricts this device to sale by or on the order of a physician.