

ComfortGel SE

Stretcher Support Surface

ComfortGel SE is designed to improve patient outcomes through: **pressure redistribution & enhanced comfort.**¹

When we developed the SE series of stretcher surfaces, we focused our innovative design on aiding in proactive prevention.

ComfortGel SE was designed with the overall patient experience in mind and has specific features to help redistribute pressure and enhance patient comfort.

Standard features

- CoreGel Technology
- Polycarbonate polyurethane cover
- Welded seams
- Sloped heel section
- Three-sided zipper
- Slip-resistant bottom

Warranty

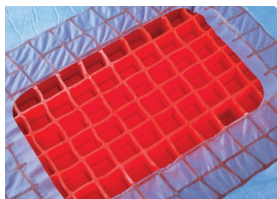
- Internal components: three (3) years
- Fire barrier: one (1) year
- Support surface cover: one (1) year

Specifications

Model number	1805
Overall width	66.0 cm (26") 76.2 cm (30")
Overall length	193.0 cm (76")
Overall thickness	14.0 cm (5.5")
Weight capacity	317.5 kg (700 lb.)
Fire retardant inner cover	Optional

CoreGel Technology

Located in the sacral region, CoreGel creates a positioning pocket to help prevent patient migration.² Its design helps absorb and disperse patient weight to aid in pressure redistribution.



Welded seams

Our covers are sealed with a specific type of RF welding utilising electromagnetic bonding to limit fluid ingress and infiltration.



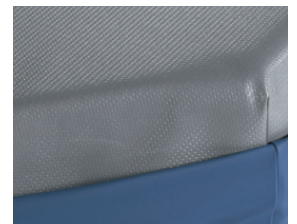
Cover design

ComfortGel SE's surface cover provides increased chemical resistance; high-frequency sealed seams, and a three-sided zipper for better visibility during internal component inspections.



Slip-resistant bottom

ComfortGel SE's bottom features a slip-resistant material to eliminate Velcro while maintaining affixation to the stretcher during patient movement and transfer.



*Flammability standards: CALTB117; 16CFR1632; CGSB CAN 2-4.2 Method 27.7-M77; BS EN 597-1; BS EN 597-2; CAL TB129; 16CFR1633; BS 6807, Clause 9; BFD IX-11; UNI9175

1. Recommended to be implemented in combination with clinical evaluation of risk factors and skin assessments made by a health care professional.

2. Best when used in combination with a stretcher equipped with Lift Assist Backrest.

A healthcare professional must always rely on his or her own professional clinical judgment when deciding whether to use a particular product when treating a particular patient. Stryker does not dispense medical advice and recommends that healthcare professionals be trained in the use of any particular product before using it. The information presented is intended to demonstrate the breadth of Stryker product offerings. A healthcare professional must always refer to the package insert, product label and/or instructions for use before using any Stryker product. Products may not be available in all markets because product availability is subject to the regulatory and/or medical practices in individual markets. Please contact your Stryker representative if you have questions about the availability of Stryker products in your area. Stryker Corporation or its divisions or other corporate affiliated entities own, use or have applied for the following trademarks or service marks: Stryker. All other trademarks are trademarks of their respective owners or holders.

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