



# Surgical Drapes

## EENT/ Neurology

CODE	DESCRIPTION	CASE QTY	CASE DIMENSIONS
 80-06001G	Craniotomy Drape w/ Pouch, 122x134" (310x340cm)	20	21x16x17" (54x40x42cm)
 80-06001G-S	Craniotomy Drape w/ Pouch, 122x134" (310x340cm)	12	18x11x19" (45x29x48cm)

### Materials

- Multilayered SMS Material with Antistatic Properties and Alcohol Repellency
- HydraGuard® Absorbent Reinforcement
- Incise Film
- Clear Polyethylene (PE) Film
- Fluid Collection Pouch, Clear Polyethylene (PE) Film

### Product Attributes

- Fluid collection pouch with filtering screen and suction port
- Clear gusseted anesthesia panels
- Absorbent reinforcement material for fluid management
- Oval fenestration

### Sterilization

**Sterile:** Sterilized by EO (ethylene oxide). Product is sterile unless the package has been damaged or opened.

**Non-Sterile:** If a product is delivered non-sterile and is ultimately for use in sterile environments, it is intended to be sterilized by the re-packager. This includes further packaging and sterilization according to the validated processes of the re-packager. These products are to be sterilized by ethylene oxide (EO). Other methods of sterilization have not been validated by Alleset.

### Shelf Life

**Sterile:** 5 years

**Non-Sterile:** Established at 5 years based upon sterile packaged product aging studies and normal warehouse storage conditions. After 5 years, the product should be performance tested to ensure compliance to specifications.

### Product Certifications/Standards

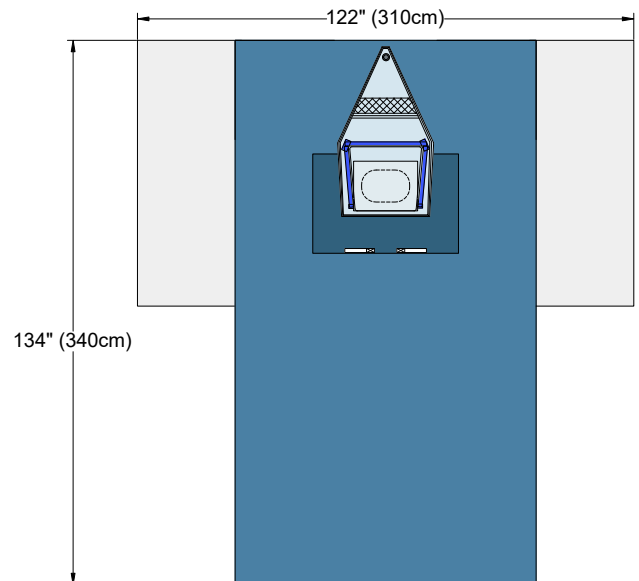
- CE certified as Class I, according to MDR 2017/745 Annex VIII, Rule 1

### Manufacturing Certifications/Registrations

- EN ISO 13485:2016
- Medical Device Single Audit Process Certified

### Storage

- General warehouse conditions



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Country of Origin Abbreviations per ISO 3166-1, Alpha-2 Codes.



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