

CERAMENT® G

Product Fact Sheet



COMPOSITION:

| CERAMENT Powder | Liquid | Gentamicin Powder |
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| <p>60 wt% α-calcium sulfate hemihydrate (CaS)</p> <p>40 wt% hydroxyapatite (HA)</p> <ul style="list-style-type: none"> A calcium phosphate, with a chemically and structural similarity to the mineral phase of bone Osteoconductive, which means it forms a direct bond with osteoblasts that form new bone Engineered in R&D to have a specific size and crystallinity that confers high injectability and slow resorption rate | <ul style="list-style-type: none"> Saline: sodium chloride 9 mg/mL | <ul style="list-style-type: none"> Gentamicin sulfate; providing 17.5 mg of gentamicin/mL of CERAMENT paste (both 5 and 10 mL product) |

TIP EXTENDERS:

11G, 50mm length
 11G, 100mm length
 Tapered tip

SIZE AND ORDER CODES:

| Volume | Order code |
|--------|------------|
| 5mL | A0450-03 |
| 10mL | A0450-01 |

| | |
|------------|-------|
| GMDN code | 47255 |
| UMDNS code | 37286 |

Manufacturer:

BONESUPPORT AB
 Scheelevägen 19
 IDEON Science Park
 SE-223 70 Lund
 Sweden

Email: info@bonesupport.com
www.bonesupport.com

REGULATORY INFORMATION:

Regulatory Status

Medical Device Classification: Class III by rule 8 and 13 of the Council Directive 93/42/EEC amended by Directive 2007/47/EC

Intended Use

CERAMENT G is a resorbable ceramic bone graft substitute intended to fill gaps and voids in the skeletal system to promote bone healing.

CERAMENT G provides a void/gap filler that during the surgical procedure can augment hardware and bone alignments.

Gentamicin is included in CERAMENT G to prevent colonization of Gentamicin sensitive microorganisms in order to protect bone healing.

Indications

CERAMENT G is indicated to be placed into bone voids or gaps in the skeletal system, i.e. extremities, spine, and pelvis that is not intrinsic to the stability of the bony structure. These osseous defects may be e.g. spontaneous occurring, surgically created, resulting from traumatic injury to the bone during primary surgery and revision surgery, or osseous defects identified around hardware devices.

Description

CERAMENT G is bone graft substitute which is injectable and moldable into beads, consisting of Calcium sulfate, Hydroxyapatite and Gentamicin sulfate. CERAMENT G delivers 17.5 mg Gentamicin/ml paste. By combining Calcium sulfate and Hydroxyapatite an optimal balance is achieved between implant resorption rate and bone in growth rate. Calcium sulfate acts as a resorbable carrier for hydroxyapatite. Hydroxyapatite has a slow resorption rate, high osteoconductivity promoting bone in growth, and gives long term structural support to the newly formed bone. By adding Gentamicin, colonization with Gentamicin sensitive microorganisms can be prevented in order to protect bone healing. The ceramic bone substitute material is placed into the bone void under visual inspection.

Contraindications

- Hypersensitivity to any amino-glycoside antibiotics
- Myasthenia gravis
- Severe renal impairment
- Pre-existing calcium metabolism disorder
- Pregnancy
- Breastfeeding

Warnings in IFU

Addition of Gentamicin does not negate the need for systemic antibiotics

PACKAGING MATERIAL SPECIFICATIONS:

| | |
|-------------------------------|--|
| Latex | Not made with natural rubber latex |
| Animal tissue | Commission regulation No 722/2012 does not apply |
| Phthalates | Not made with phthalates |
| Storage conditions | 15–25°C / 59–77°F |
| Shelf-life | 24 months |
| Sterilization | CERAMENT G is supplied sterile, except for CERAMENT MIXING LIQUID that has a non-sterile outer surface |
| Sterile | Yes |
| Single Use/disposable: | Yes |
| Sterilization methods: | EO, Steam and Gamma Irradiation |
| EO residuals: | Fulfills ISO 10993-7:2008 |

Packaging dimensions

46cm (l) x 18.9cm (w) x 6cm (d)

CERAMENT MATERIAL PROPERTIES:

Setting temperature <43°C
pH Physiological pH (6 ≤ pH ≤ 10)

Biocompatibility

The product has been evaluated to be biological safe to use (no unacceptable biological risks have been found). The components of the implantable paste have been used in humans for decades. The plastic components of the product, that come in direct or in indirect contact with the patient, are all of USP Class VI or equivalent. The glass components that come in direct or in indirect contact with the patient, are of Type I. The final product has been extensively tested with biocompatibility in-vitro and in-vivo tests, following the requirements of ISO 10993-1.

Handling

- ✓ Injectable
- ✓ For use with a bead mold (not included in pack)*
- ✓ Drillable

*If beads are prepared, wait until final setting at 20 mins

Compatibility

- ✓ Autograft, allograft, hardware

