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**SYNTHETIC CALCIUM PHOSPHATE SURGICAL MATERIAL  
FOR RESTORATION OF HARD TISSUES DEFECTS**

# **GDT SYNTHETIC BONE GRAFT**

## **INSTRUCTIONS FOR USE**

**GDT Synthetic bone material** supplied in the form of granules based on  $\beta$ -tricalcium phosphate (60%) with hydroxyapatite (40%).

### **PURPOSE**

GDT Synthetic Bone Graft is osteoplastic material, for optimisation of bone regeneration in general dental and maxillofacial surgery, as well as in traumatology and orthopedics. Prepared by sintering of synthetic materials, it does not contain substances of animal origin, biologically compatible with body tissues. Granules have a high micro-, macro- and intergranular porosity, creating ideal conditions for the fast recovery of the bone. The material is radiopaque.

### **FIELD OF USE**

**Parodontics:** filling a two- or multi-bone pockets and bi- and tri-furcations of the teeth, augmentation of the atrophied sinus.

**Implantology:** sinus lifting or sub-antral augmentation (mixed with patient bone or allograft), filling of alveolar defects for supporting of sinus base after the tooth extraction, filling extraction defects prior to the implant placement.

**Cyst defects:** Defects after extirpation of the bone cyst, defects after resection of the root apex and defects after removal of impacted teeth surgically, as well as other multi-grid bone defects of the alveolar processes and facial bones.

### **FEATURES**

Sterile and gradually resorbable bone substitute material.

Releases calcium and phosphate ions to help promote new bone formation.

GDT Synthetic Bone Graft arrives in the different particle size as following:

- 100-500  $\mu\text{m}$  – small periodontal bone defects;

- 500-1000 µm – universal particle size for most cyst and alveolar defects;
- 1000-2000 µm – recommended for large defects and sinus lifting.

The choice of granules size depends on the size and the location of the defect.

GDT Synthetic Bone Graft granules have an osteoconductive micro-and-macroporous structure that fosters dense new bone growth. Depending on the size of the granules and tissue regeneration potential of the materials, GDT Synthetic Bone Graft is completely absorbed in 6 to 9 months. Granules dissolving process takes place in parallel with the regeneration.

#### **DIRECTION FOR USE**

After the necessary preparatory surgical procedures, the granules can be mixed with the patient's blood or with saline solution in a sterile Dappen dish before being placed onto the operative site using a dental spatula. For large bone defects, mix with bone particles from the same patient (autologous bone) for the best results. The operative site will have to be closed by joining together the wound edges (coaptation) with suture stitches.

#### **PACKAGING:**

Type	Size	Products	REF
Granules	0.25 cc	Bits 500-1000 µm	GD-SG025
Granules	0.5 cc	Bits 500-1000 µm	GD-SG050
Granules	1.0 cc	Bits 500-1000 µm	GD-SG100
Granules	2.0 cc	Bits 500-1000 µm	GD-SG200
Granules	5.0 cc	Bits 500-1000 µm	GD-SG500
Blocks	□ 5 x 5 x 5 mm	4 pcs in vial	GD-SG5x5
Blocks	□ 5 x 5 x 10 mm	2 pcs in vial	GD-SG5x10
Cones	∅ 5 x L 15 mm	2 pcs in vial	GD-SG515

Sterile vial in blister package (sterilised by gamma radiation – minimal dose: 25 kGy).

#### **STORAGE**

Store in a dry and dark place in tightly closed containers at temperature (+5°C...+25°C). Avoid contact with moisture! Do not use after the expiry date mentioned on the outer package.



**STERILE R**

**ATTENTION!** The sterility of the product cannot be guaranteed if the package bears evidence of damage, has been opened previously or wet.

*Failure to comply with the conditions of storage leads to a change of the working characteristics of the material and decrease the shelf life of the material.*

*The manufacturer is not responsible for any loss of quality caused by the failure to comply with terms of transportation, storage and use established by the manufacturer for this product. Responsibility for the use of the material for purposes other than those specified by the manufacturer falls on the user.*