

Instruction for Use

Gutta Percha Points(Obturator)

Used to fill a prepared root canal prior to tooth restoration.

Composition

Gutta Percha, Zinc Oxide, Zirconium Oxide, Barium Sulfate, Coloring Agent

Application

1. Check the product if it is damaged, presence of impurities, and shape is suitable for use.
2. Check if the product is within the expiration date.
3. After treating the root canal of the damaged tooth, fill the root canal with the product by using obturation gun.
4. This product is disposable and discarded after use.

Intended Purpose

1. Intended Use: Used to fill a prepared root canal prior to tooth restoration.
2. Indication: Used to fill a prepared root canal prior to tooth restoration.
3. Contraindication:
Do not use for patient who is allergic to this product.
4. Patient Population: All age group
5. Intended user: Dentist
6. Side Effects/Adverse Effects: There are no side effects or adverse effects reported.
7. Potential risk if re-used: Re-use of used Gutta percha obturator may cause inflammation.
8. Can be used with sealer or cement and it shall be used under the control of the dentist according to the clinical indication.
9. Single Use Device

Precaution

1. Do not use other than dental specialist.
2. Do not use if the product is expired.
3. Before use, check for infection inside the root canal.
4. Do not use for patient who is allergic to this product.

Storage

Store at room temperature away from direct sunlight

Disposal : Dispose of contents/container according to the related regulations of the country.

Shelf Life : 4 years from the date of manufacture

Radiopaque : Equivalent to 6mm aluminium stepwedge

Symbols



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■ In case that any serious incident that has occurred in relation to the device, please report to the manufacturer and the competent authority of the Member State in which the user and/or patient is established.

■ Size Designation and Length: Please contact the manufacturer for size designation and length.