

## NeoDrys® Flex Saliva Absorbents Instructions for Use

**CAUTION:**

Rx Only. These instructions, in whole or in part, are not a substitute for formal training. Appropriate professional education is **STRONGLY RECOMMENDED** prior to using this device clinically.

**DESCRIPTION:**

NeoDrys Flex is a sublingual and parotid saliva absorbent. It is designed to fit on the floor of the mouth to contact the sublingual mucosa, next to the lingual molars, or over the parotid gland on the cheek. The product is manufactured with permeable materials that entrap saliva within particles of the absorbent.

**INDICATIONS**

NeoDrys Flex absorbent is indicated to absorb moisture for up to 14 times its weight. Based on its placement, Flex will absorb from the sublingual, submandibular, and parotid glands for up to 15 minutes depending on the volume of saliva production.

**CONTRAINDICATIONS TO USE**

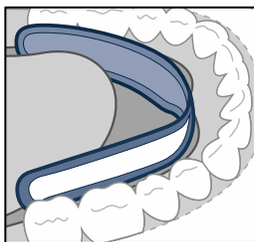
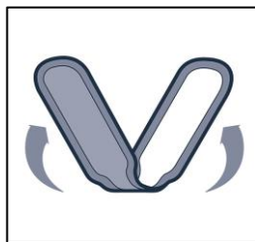
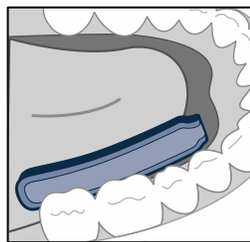
Use of Microcopy NeoDrys Flex is contraindicated on any patient who is allergic to any of the components of the product. Do not reuse. The NeoDrys Flex is single-use. 

**CLINICAL PRECAUTIONS AND WARNINGS:**

- a) Discard any damaged NeoDrys Flex product immediately.
- b) Microcopy NeoDrys Flex are for **SINGLE-PATIENT-USE ONLY** and should be used exclusively in a dental setting.
- c) To remove without tissue irritation, release adhesion with ample water spray to the tissue contacting side of the product.
- d) Non-toxic by ingestion. If product bursts, remove as much as possible from mouth. Rinse mouth thoroughly with plenty of water. If adverse symptoms appear, seek medical attention.
- e) Always keep track of Lot Numbers of NeoDrys Flex to ensure traceability.

**CLINICAL USE:**
**Sublingual Steps**

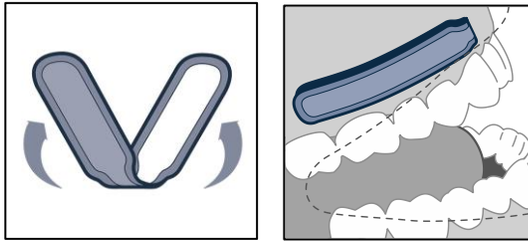
- Rinse and dry underside of tongue
- Blue side faces tongue.


**1.**

**2.**


- **1. Anterior:** Position below tongue on the sublingual floor.
- **2. Posterior:** Fold white sides together and position next to lingual molars.
- **Important:** To remove, gently spray with water.

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### Parotid Steps



1.

2.

- 1. Parotid: Fold white sides together.
- 2. Parotid: Position above facial molars.
- **Important:** To remove, gently spray with water.

### STORAGE

Microcopy NeoDrys Flex should be stored in a dry, closed container. Improper storage conditions will shorten the shelf life and may cause malfunction of the product.

### TRACEABILITY

Each package includes **Lot number** LOT on its label.

This number must be quoted in any correspondence regarding the product.

### SYMBOLS:

	Manufacturer	Indicates the medical device manufacturer.		Do not reuse	Indicates a medical device that is intended for one use, or for use on a single patient during a single procedure.
<span style="border: 1px solid black; padding: 2px;">LOT</span>	Lot Code	Indicates the manufacturer's batch code so that the batch or lot can be identified.		Consult instructions for use	Indicates the need for the user to consult the instructions for use.
	CE marking	Signifies European technical conformity.		Do not use if package is open or damaged	Indicates a medical device that should not be used if the package has been damaged or opened.
<span style="border: 1px solid black; padding: 2px;">REF</span>	Catalog Number	Indicates the manufacturer's catalog number so that the medical device can be identified.		Date of Manufacture	Symbol for date of manufacture.

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	Medical Device	Indicates device is designed and intended for medical use.		Keep Dry	Indicates a medical device that needs to be protected from moisture.
	DEVICE for professional use only	(ref US FDA CDRH) Indicates device shall only be used by a trained professional.		Authorized European representative	Indicates the Authorized representative in the European Community.
	Non-Sterile	Indicates a medical device that has not been subjected to a sterilization process.			

**CONTACT INFORMATION:**



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