# IDODENT FLOURIDE MINT- fluoride paste United Exchange Corp.

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

-----

## Active Ingredient Purpose

Sodium monofluorophosphate 0.76%------ Anticavity (Total Fluoride Content - 1000 ppm Approx.)

#### Uses

Regular brushing with fluoride toothpaste helps protect teeth and roots against cavities

## Warnings

## **Warnings**

### Keep out of reach of children under 6 years of age.

In case of accidental overdose, seek professional assistance or contact a Poison Control Center immediately.

#### **Directions**

Adults and children 2	Brush Teeth thoroughly after meals, twice a day, or use as
years and older	directed by a dentist or physician.
Children under 6	To minimize swallowing use a pea-sized amount and supervise
years	brushing until good habits are established.
Children under 2	Ask a dentist or physician.
years	Ask a deficise of prhysician.

#### Other Information

Store at room temperature 20-25°C (68-77°F)

## Inactive ingredients

calcium carbonate, water, sorbitol, hydrated silica, sodium lauryl sulfate, flavor, cellulose gum, sodium silicate, tetrasodium, propphosphate, sodium saccharin, titanium dioxide, methylparaben, PEG-8, propylparaben

## Distributed by:

United Exchange Corp.

17211 Valley View Ave.

Cerritos, CA 90703 USA



## **IDODENT FLOURIDE MINT**

fluoride paste

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:65923-181
Route of Administration	DENTAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
<b>SODIUM MONOFLUOROPHOSPHATE</b> (UNII: C810JCZ56Q) (FLUORIDE ION - UNII:Q80VPU408O)	FLUORIDE ION	0.76 g in 100 g	

Inactive Ingredients		
Ingredient Name	Strength	
PROPYLPARABEN (UNII: Z8IX2SC1OH)		
METHYLPARABEN (UNII: A218C7H19T)		
POLYETHYLENE GLYCOL 400 (UNII: B697894SGQ)		
SODIUM LAURYL SULFATE (UNII: 368GB5141J)		
CALCIUM CARBONATE (UNII: H0G9379FGK)		
WATER (UNII: 059QF0KO0R)		

CARBOXYMETHYLCELLULOSE SODIUM, UNSPECIFIED FORM (UNII: K6790BS311)	
SORBITOL (UNII: 506T60A25R)	
HYDRATED SILICA (UNII: Y6O7T4G8P9)	
SACCHARIN SODIUM (UNII: SB8ZUX40TY)	
SODIUM PYROPHOSPHATE (UNII: O352864B8Z)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
SODIUM SILICATE (UNII: IJF18F77L3)	

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:65923-181- 18	1 in 1 CARTON	12/21/2016	10/13/2021	
1		181 g in 1 TUBE; Type 1: Convenience Kit of Co-Package			

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	part355	10/23/2014	

## Labeler - United Exchange Corp. (840130579)

Revised: 11/2021 United Exchange Corp.