

AMMONIA INHALANTS- ammonia inhalants inhalant Dynarex Corporation

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

1401 Ammonia Inhalants (Ampule) NDC 67777-251-01

active ingredient each inhalant purpose ammonia 15% inhalant

Uses to prevent or treating fainting

warnings for external use only

do not use if you have breathing problems such as asthma or emphysema

stop use and ask a doctor if condition persists

If swallowed get medical help or contact a Poison Control Center right away

Hold inhalant away from face and crush between thumb and forefinger. carefully approach crushed inhalant to nostrils of affected person.

Store at room temperature away from light

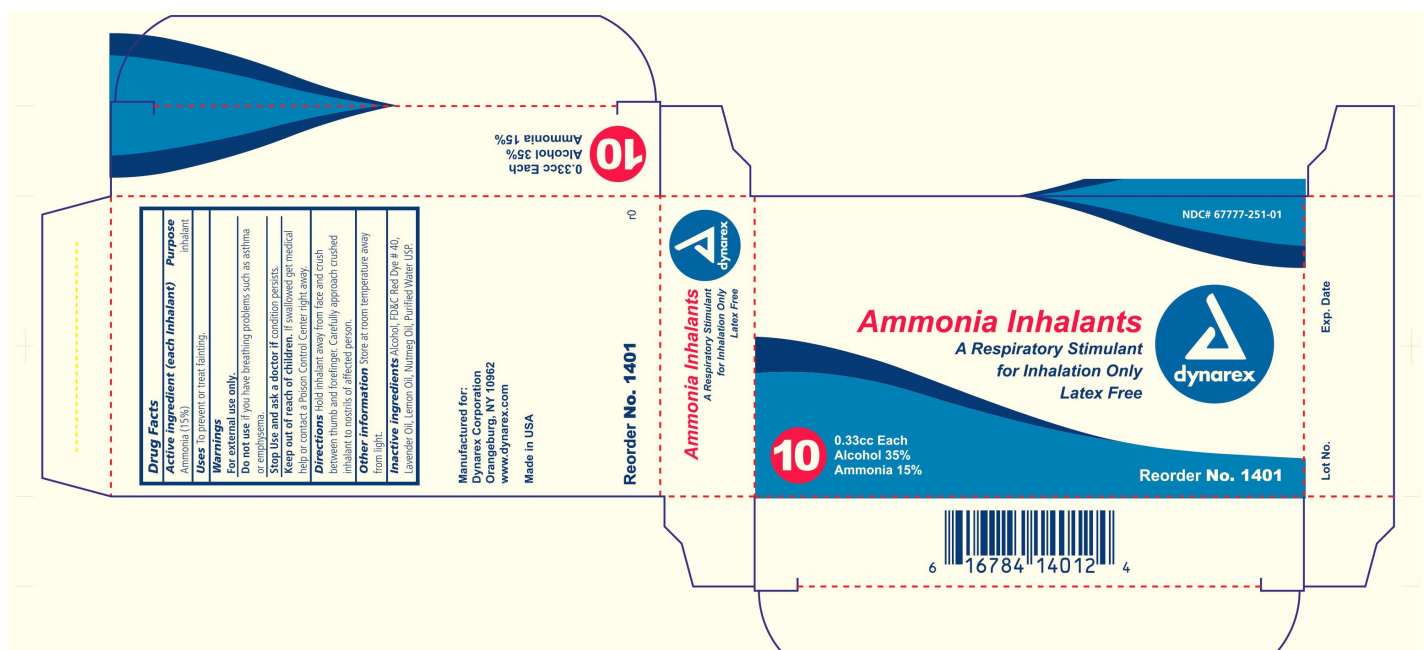
alcohol fdc red dye 40 lavender oil, lemon oil, nutmeg oil, purified water usp

to prevent or treat fainting

for external use only

ammonia inhalants

a respiratory stimulant for inhalation only latex free



AMMONIA INHALANTS

ammonia inhalants inhalant

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:67777-251
Route of Administration	RESPIRATORY (INHALATION)		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
AMMONIA (UNII: 5138Q19F1X) (AMMONIA - UNII:5138Q19F1X)	AMMONIA	0.05 g in 0.33 mL

Inactive Ingredients

Ingredient Name	Strength
ALCOHOL (UNII: 3K9958V90M)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:67777-251-01	0.33 mL in 1 AMPULE; Type 0: Not a Combination Product	02/14/1976	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		02/14/1976	

Labeler - Dynarex Corporation (008124539)

Registrant - Dynarex Corporation (008124529)

Revised: 5/2023

Dynarex Corporation