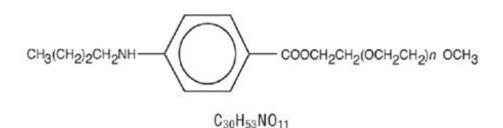
BENZONATATE- benzonatate capsule Proficient Rx LP

Benzonatate Capsules, USP

DESCRIPTION

Benzonatate, a non-narcotic oral antitussive agent, is 2, 5, 8, 11, 14, 17, 20, 23, 26nonaoxaoctacosan- 28-yl p-(butylamino) benzoate; with a molecular weight of 603.0.



Each benzonatate capsule USP, 100 mg contains: Benzonatate, USP 100 mg.

Each benzonatate capsule USP, 200 mg contains: Benzonatate, USP 200 mg.

Benzonatate capsules USP also contain: gelatin 175 bloom bone NF, glycerin 99% USP, methyl/propyl paraben blend (4:1), yellow #10-DC and white ink (shellac glaze in SD-45, titanium dioxide, isopropyl alcohol, n-butyl alcohol, propylene glycol, ammonium hydroxide and simethicone).

CLINICAL PHARMACOLOGY

Benzonatate acts peripherally by anesthetizing the stretch receptors located in the respiratory passages, lungs, and pleura by dampening their activity and thereby reducing the cough reflex at its source. It begins to act within 15 to 20 minutes and its effect lasts for 3 to 8 hours. Benzonatate has no inhibitory effect on the respiratory center in recommended dosage.

INDICATIONS AND USAGE

Benzonatate capsules USP are indicated for the symptomatic relief of cough.

CONTRAINDICATIONS

Hypersensitivity to benzonatate or related compounds.

WARNINGS

Severe hypersensitivity reactions (including bronchospasm, laryngospasm and

cardiovascular collapse) have been reported which are possibly related to local anesthesia from sucking or chewing the capsule instead of swallowing it.

Severe reactions have required intervention with vasopressor agents and supportive measures.

Isolated instances of bizarre behavior, including mental confusion and visual hallucinations, have also been reported in patients taking benzonatate capsules USP in combination with other prescribed drugs.

PRECAUTIONS

Benzonatate is chemically related to anesthetic agents of the para-aminobenzoic acid class (e.g. procaine; tetracaine) and has been associated with adverse CNS effects possibly related to a prior sensitivity to related agents or interaction with concomitant medication.

Information for patients: Release of benzonatate from the capsule in the mouth can produce a temporary local anesthesia of the oral mucosa and choking could occur. Therefore, the capsules should be swallowed without chewing.

Usage in Pregnancy: Pregnancy Category C. Animal reproduction studies have not been conducted with benzonatate capsules USP. It is also not known whether benzonatate capsules USP can cause fetal harm when administered to a pregnant woman or can affect reproduction capacity. Benzonatate capsules USP should be given to a pregnant woman only if clearly needed.

Nursing mothers: It is not known whether this drug is excreted in human milk. Because many drugs are excreted in human milk caution should be exercised when benzonatate capsules USP are administered to a nursing woman.

Carcinogenesis, mutagenesis, impairment of fertility: Carcinogenicity, mutagenicity, and reproduction studies have not been conducted with benzonatate capsules USP.

Pediatric Use: Safety and effectiveness in children below the age of 10 have not been established.

ADVERSE REACTIONS

Potential Adverse Reactions to benzonatate capsules USP may include: Hypersensitivity reactions including bronchospasm, laryngospasm, and cardiovascular collapse possibly related to local anesthesia from chewing or sucking the capsule.

CNS: sedation; headache; dizziness; mental confusion; visual hallucinations.

GI: constipation; nausea; GI upset.

Dermatologic: pruritus; skin eruptions.

Other: nasal congestion; sensation of burning in the eyes; vague "chilly" sensation; numbness of the chest; hypersensitivity.

Rare instances of deliberate or accidental overdose have resulted in death.

OVERDOSAGE

Overdose may result in death.

The drug is chemically related to tetracaine and other topical anesthetics and shares various aspects of their pharmacology and toxicology. Drugs of this type are generally well absorbed after ingestion.

Signs and Symptoms:

If capsules are chewed or dissolved in the mouth, oropharyngeal anesthesia will develop rapidly. CNS stimulation may cause restlessness and tremors which may proceed to clonic convulsions followed by profound CNS depression.

Treatment:

Evacuate gastric contents and administer copious amounts of activated charcoal slurry. Even in the conscious patient, cough and gag reflexes may be so depressed as to necessitate special attention to protection against aspiration of gastric contents and orally administered materials. Convulsions should be treated with a short-acting barbiturate given intravenously and carefully titrated for the smallest effective dosage. Intensive support of respiration and cardiovascular-renal function is an essential feature of the treatment of severe intoxication from overdosage.

Do not use CNS stimulants.

DOSAGE AND ADMINISTRATION

Adults and Children over 10: Usual dose is one 100 mg or 200 mg capsule t.i.d. as required. If necessary, up to 600 mg daily may be given.

HOW SUPPLIED

Benzonatate capsules USP 100 mg are available for oral administration as a clear, colorless to pale yellow oil in a clear, yellow softgel shell imprinted "A" in white ink.

They are supplied as follows:

Bottles of 10 (NDC 71205-885-10)

Bottles of 14 (NDC 71205-885-14)

Bottles of 15 (NDC 71205-885-15)

Bottles of 20 (NDC 71205-885-20)

Bottles of 21 (NDC 71205-885-21)

Bottles of 30 (NDC 71205-885-30)

Bottles of 100 (NDC 71205-885-00)

Bottles of 500 (NDC 71205-885-55)

Benzonatate capsules USP 200 mg are available for oral administration as a clear, colorless to pale yellow oil in a clear, yellow softgel shell imprinted "A2" in white ink.

They are supplied as follows:

Bottles of 10 (NDC 71205-886-10)

Bottles of 14 (NDC 71205-886-14)

Bottles of 15 (NDC 71205-886-15)

Bottles of 20 (NDC 71205-886-20)

Bottles of 21 (NDC 71205-886-21)

Bottles of 30 (NDC 71205-886-30)

Bottles of 100 (NDC 71205-886-00)

Bottles of 500 (NDC 71205-886-55)

Storage

Store at 20° to 25°C (68° to 77° F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tight, light-resistant container [see USP].

Protect from moisture.

Manufactured by:

Catalent Pharma Solutions Windsor, ON Canada N9C 3R5

Manufactured for:

Acella Pharmaceuticals, LLC Alpharetta, GA 30005

Repackaged and Relabeled by:

Proficient Rx LP Thousand Oaks, CA 91320

L-0273 Rev 1020-01

PACKAGE LABEL.PRINCIPAL DISPLAY PANEL





NDC 71205-885-00

RX Only

Packaged By: Proficient Rx LP Thousand Oaks, CA 91320

Benzonatate 100mg

#100 Capsules

Each capsule contains: 100 mg of benzonatate.

Clear, colorless to pale yellow oil in a clear, yellow softgel shell imprinted "A" in white ink.

Usual dosage: See package insert.

Product ID: QB088500

Mfr. By: Catalent Pharma Solutions Windsor, ON Canada N9C 3R5

Store at 20°-25°C (68°-77°F). Dispense in a tight, light-resistant container [see USP]. Protect from moisture.

Keep medication out of the reach of children





NDC 71205-886-14

RX Only

Packaged By: Proficient Rx LP Thousand Oaks, CA 91320

Benzonatate 200mg

#14 Capsules

Each capsule contains: 200 mg of benzonatate.

Clear, colorless to pale yellow oil in a clear, yellow softgel shell imprinted "A2" in white ink.

Product ID: QB088614

Mfr. By: Catalent Pharma Solutions Windsor, ON Canada N9C 3R5 Store at 20°-25°C (68°-77°F)

Keep medication out of the reach of children

BENZONATATE benzonatate capsule					
Product Information					
Product Type	HUMAN PRESCRIPTION DRUG	ltem Code (Source)	NDC:71205-885(NDC:42192- 617)		
Route of Administration	ORAL				
Active Ingredient/Active Moiety					

GTIN: 00371205885007 SN# MASTER Exp. 00/00/00 Lot #.00000



GTIN: 00371205886141 SN# MASTER Exp. 00/00/00 Lot #:00000



	Ingredient Name		Basis of S	trengt	h Strength
BENZONATATE (UN	NII: 5P4DHS6ENR) (BENZONATATE - UNII:5P4DH	56ENR)	BENZONATAT	E	100 mg
Inactive Ingre	dients				
-	Ingredient Name			S	trength
GELATIN, UNSPEC	IFIED (UNII: 2G86QN327L)				
GLYCERIN (UNII: PC	DC6A3C0OX)				
METHYLPARABEN	(UNII: A2I8C7HI9T)				
PROPYLPARABEN	(UNII: Z8IX2SC1OH)				
D&C YELLOW NO.	10 (UNII: 35SW5USQ3G)				
SHELLAC (UNII: 46N	J107B71O)				
TITANIUM DIOXIDE	E (UNII: 15FIX9V2JP)				
ISOPROPYL ALCO	HOL (UNII: ND2M416302)				
BUTYL ALCOHOL (UNII: 8PJ61P6TS3)				
PROPYLENE GLYC	OL (UNII: 6DC9Q167V3)				
AMMONIA (UNII: 51	38Q19F1X)				
DIMETHICONE (UN	II: 92RU3N3Y1O)				
SILICON DIOXIDE	(UNII: ETJ7Z6XBU4)				
Product Chara					
Color	YELLOW (clear; yellow)	Score no score			score
Shape	CAPSULE	Size 6mm		nm	
Flavor		Imprint Code A			
Contains					
Packaging					
# Item Code	Package Description		ting Start Date	Marl	ceting End Date
1 NDC:71205-885- 10	10 in 1 BOTTLE; Type 0: Not a Combination Product	09/17/202	1	08/31/2025	
2 NDC:71205-885- 14	14 in 1 BOTTLE; Type 0: Not a Combination Product	09/17/2021 08/31/2025		025	
3 NDC:71205-885- 15	15 in 1 BOTTLE; Type 0: Not a Combination Product	09/17/2021 03/31/2024		024	
4 NDC:71205-885- 20	20 in 1 BOTTLE; Type 0: Not a Combination Product	09/17/202	1	08/31/2	025

5 NDC:71205-885-21 in 1 BOTTLE; Type 0: Not a Combination Product 6 NDC:71205-885-30 NDC:71205-885-9 Product Not a Combination 09/17/2021 08/31/2025 7 NDC:71205-885- 100 in 1 BOTTLE; Type 0: Not a Combination 09/17/2021 03/31/2025 00 Product 8 NDC:71205-885-55 500 in 1 BOTTLE; Type 0: Not a Combination Product 09/17/2021 03/31/2025

Marketing Information

Marketing

Application Number or Monograph Marketing Start Marketing End

09/17/2021

08/31/2025

Category	Citation	Date	Date
ANDA	ANDA091310	08/02/2021	08/31/2025

BENZONAT	ΆΤΕ					
enzonatate ca	psule					
Product Info	rmation					
Product Type		HUMAN PRESCRIPTION DRUG	ltem Code (Source)	•		
Route of Admir	nistration	ORAL				
Active Ingred	lient/Active	Moiety				
		dient Name		Basi	s of Strengt	h Strength
BENZONATATE (U	-	R) (BENZONATATE - UNII:5P	4DHS6ENR)			200 mg
Inactive Ingr	edients					
		Ingredient Name			S	trength
GELATIN, UNSPE	CIFIED (UNII: 2G	86QN327L)				
GLYCERIN (UNII: P	DC6A3C0OX)					
METHYLPARABEN	(UNII: A2I8C7HI	ЭТ)				
PROPYLPARABEN	I (UNII: Z8IX2SC1	.OH)				
D&C YELLOW NO	. 10 (UNII: 355V	/5USQ3G)				
SHELLAC (UNII: 46	5N107B71O)					
TITANIUM DIOXIC	DE (UNII: 15FIX9V	2JP)				
ISOPROPYL ALCO	HOL (UNII: ND2I	M416302)				
BUTYL ALCOHOL	(UNII: 8PJ61P6T9	53)				
PROPYLENE GLY	COL (UNII: 6DC9	Q167V3)				
AMMONIA (UNII: 5	138Q19F1X)					
DIMETHICONE (U	NII: 92RU3N3Y1C)				
SILICON DIOXIDE	(UNII: ETJ7Z6XB	U4)				
Product Char	acteristics					
Color	YELLOW (cle	ar; yellow)	Score		no	score
Shape	CAPSULE		Size		11	mm
Flavor Imprint Code A2						
Contains			•			
Packaging						
# Item Code	Pa	kage Description	Marke	ting S [.] Date	tart Marl	ceting End Date
1 NDC:71205-886	- 10 in 1 BOTTL Product	E; Type 0: Not a Combinatio			11/30/2	

2	14	Product 09/1//2021		11/20/2023		
3	NDC:71205-886- 15	15 in 1 BOTTLE; Type 0: Not a Combination Product	09/17/2021	11/30/2025		
4	NDC:71205-886- 20	20 in 1 BOTTLE; Type 0: Not a Combination Product	09/17/2021	11/30/2025		
5	NDC:71205-886- 21	21 in 1 BOTTLE; Type 0: Not a Combination Product	09/17/2021	04/30/2025		
6	NDC:71205-886- 30	30 in 1 BOTTLE; Type 0: Not a Combination Product	09/17/2021	04/30/2025		
7	NDC:71205-886- 00	100 in 1 BOTTLE; Type 0: Not a Combination Product	09/17/2021	11/30/2025		
8	NDC:71205-886- 55	500 in 1 BOTTLE; Type 0: Not a Combination Product	09/17/2021	11/30/2025		
Marketing Information						
	Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
٨N	IDA	ANDA091310	08/02/2021	11/30/2025		

Labeler - Proficient Rx LP (079196022)

Establishment				
Name	Address	ID/FEI	Business Operations	
Proficient Rx LP		079196022	REPACK(71205-885, 71205-886), RELABEL(71205-885, 71205-886)	

Revised: 5/2024

Proficient Rx LP