BETHANECHOL CHLORIDE- bethanechol chloride tablet Upsher-Smith Laboratories, LLC

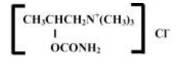
BETHANECHOL CHLORIDE TABLETS, USP 5 mg, 10 mg, 25 mg and 50 mg

Rx only

DESCRIPTION

Bethanechol chloride, USP, a cholinergic agent, is a synthetic ester which is structurally and pharmacologically related to acetylcholine.

It is designated chemically as 2-[(aminocarbonyl)oxy]-N, N, N-trimethyl-1propanaminium chloride. Its molecular formula is $C_7H_{17}CIN_2O_2$ and its structural formula is:



It is a white, hygroscopic crystalline powder having a slight amine-like odor, freely soluble in water, and has a molecular weight of 196.68.

Each tablet for oral administration contains 5 mg, 10 mg, 25 mg or 50 mg bethanechol chloride, USP. Tablets also contain the following inactive ingredients: colloidal silicon dioxide, lactose monohydrate, magnesium stearate, microcrystalline cellulose, sodium starch glycolate, (25 mg and 50 mg) D&C Yellow #10 Lake and FD&C Yellow #6 Lake.

CLINICAL PHARMACOLOGY

Bethanechol chloride acts principally by producing the effects of stimulation of the parasympathetic nervous system. It increases the tone of the detrusor urinae muscle, usually producing a contraction sufficiently strong to initiate micturition and empty the bladder. It stimulates gastric motility, increases gastric tone and often restores impaired rhythmic peristalsis.

Stimulation of the parasympathetic nervous system releases acetylcholine at the nerve endings. When spontaneous stimulation is reduced and therapeutic intervention is required, acetylcholine can be given, but it is rapidly hydrolyzed by cholinesterase and its effects are transient. Bethanechol chloride is not destroyed by cholinesterase and its effects are more prolonged than those of acetylcholine.

Effects on the GI and urinary tracts sometimes appear within 30 minutes after oral administration of bethanechol chloride, but more often 60 to 90 minutes are required to reach maximum effectiveness. Following oral administration, the usual duration of action of bethanechol chloride is one hour, although large doses (300 to 400 mg) have been reported to produce effects for up to six hours. Subcutaneous injection produces a more intense action on bladder muscle than does oral administration of the drug.

Because of the selective action of bethanechol chloride, nicotinic symptoms of cholinergic stimulation are usually absent or minimal when orally or subcutaneously administered in therapeutic doses, while muscarinic effects are prominent. Muscarinic effects usually occur within 5 to 15 minutes after subcutaneous injection, reach a maximum in 15 to 30 minutes, and disappear within two hours. Doses that stimulate micturition and defecation and increase peristalsis do not ordinarily stimulate ganglia or voluntary muscles. Therapeutic test doses in normal human subjects have little effect on heart rate, blood pressure or peripheral circulation.

Bethanechol chloride does not cross the blood-brain barrier because of its charged quaternary amine moiety. The metabolic rate and mode of excretion of the drug have not been elucidated.

A clinical study (Diokno, A.C.; Lapides, J.; *Urol 10*: 23-24, July 1977) was conducted on the relative effectiveness of oral and subcutaneous doses of bethanechol chloride on the stretch response of bladder muscle in patients with urinary retention. Results showed that 5 mg of the drug given subcutaneously stimulated a response that was more rapid in onset and of larger magnitude than an oral dose of 50 mg, 100 mg, or 200 mg. All the oral doses, however, had a longer duration of effect than the subcutaneous dose. Although the 50 mg oral dose caused little change in intravesical pressure in this study, this dose has been found in other studies to be clinically effective in the rehabilitation of patients with decompensated bladders.

INDICATIONS AND USAGE

Bethanechol chloride tablets are indicated for the treatment of acute postoperative and postpartum nonobstructive (functional) urinary retention and for neurogenic atony of the urinary bladder with retention.

CONTRAINDICATIONS

Hypersensitivity to bethanechol chloride tablets, hyperthyroidism, peptic ulcer, latent or active bronchial asthma, pronounced bradycardia or hypotension, vasomotor instability, coronary artery disease, epilepsy and parkinsonism.

Bethanechol chloride should not be employed when the strength or integrity of the gastrointestinal or bladder wall is in question, or in the presence of mechanical obstruction; when increased muscular activity of the gastrointestinal tract or urinary bladder might prove harmful, as following recent urinary bladder surgery, gastrointestinal resection and anastomosis, or when there is possible gastrointestinal obstruction; in bladder neck obstruction, spastic gastrointestinal disturbances, acute inflammatory lesions of the gastrointestinal tract, or peritonitis; or in marked vagotonia.

PRECAUTIONS

General

In urinary retention, if the sphincter fails to relax as bethanechol chloride contracts the bladder, urine may be forced up the ureter into the kidney pelvis. If there is bacteriuria, this may cause reflux infection.

Information for Patients

Bethanechol chloride tablets should preferably be taken one hour before or two hours after meals to avoid nausea or vomiting. Dizziness, lightheadedness or fainting may occur, especially when getting up from a lying or sitting position.

Drug Interactions

Special care is required if this drug is given to patients receiving ganglion blocking compounds because a critical fall in blood pressure may occur. Usually, severe abdominal symptoms appear before there is such a fall in the blood pressure.

Carcinogenesis, Mutagenesis, Impairment of Fertility

Long-term studies in animals have not been performed to evaluate the effects upon fertility, mutagenic or carcinogenic potential of bethanechol chloride.

Pregnancy

Teratogenic Effects

Animal reproduction studies have not been conducted with bethanechol chloride. It is also not known whether bethanechol chloride can cause fetal harm when administered to a pregnant woman or can affect reproduction capacity. Bethanechol chloride should be given to a pregnant woman only if clearly needed.

Nursing Mothers

It is not known whether this drug is secreted in human milk. Because many drugs are secreted in human milk and because of the potential for serious adverse reactions from bethanechol chloride in nursing infants, a decision should be made whether to discontinue nursing or to discontinue the drug, taking into account the importance of the drug to the mother.

Pediatric Use

Safety and effectiveness in pediatric patients have not been established.

ADVERSE REACTIONS

Adverse reactions are rare following oral administration of bethanechol chloride, but are more common following subcutaneous injection. Adverse reactions are more likely to occur when dosage is increased.

The following adverse reactions have been observed: *Body as a Whole*: malaise; *Digestive*: abdominal cramps or discomfort, colicky pain, nausea and belching, diarrhea, borborygmi, salivation; *Renal*: urinary urgency; *Nervous System*: headache; *Cardiovascular*: a fall in blood pressure with reflex tachycardia, vasomotor response; *Skin*: flushing producing a feeling of warmth, sensation of heat about the face, sweating; *Respiratory*: bronchial constriction, asthmatic attacks; *Special Senses*: lacrimation, miosis.

Causal Relationship Unknown

The following adverse reactions have been reported, and a causal relationship to therapy with bethanechol chloride has not been established: *Body as a Whole*: malaise; *Nervous System*: seizures.

OVERDOSAGE

Early signs of overdosage are abdominal discomfort, salivation, flushing of the skin ("hot feeling"), sweating, nausea, and vomiting.

Atropine Sulfate is a specific antidote. The recommended dose for adults is 0.6 mg. Repeat doses can be given every two hours, according to clinical response. The recommended dosage in infants and children up to 12 years of age is 0.01 mg/kg (to a maximum single dose of 0.4 mg) repeated every two hours as needed until the desired effect is obtained or adverse effects of atropine preclude further usage. Subcutaneous injection of atropine is preferred except in emergencies when the intravenous route may be employed.

The oral LD_{50} of bethanechol chloride is 1510 mg/kg in the mouse.

DOSAGE AND ADMINISTRATION

Dosage must be individualized, depending on the type and severity of the condition to be treated.

Preferably give the drug when the stomach is empty. If taken soon after eating, nausea and vomiting may occur.

The usual adult oral dose ranges from 10 to 50 mg three or four times a day. The minimum effective dose is determined by giving 5 to 10 mg initially and repeating the same amount at hourly intervals until satisfactory response occurs, or until a maximum of 50 mg has been given. The effects of the drug sometimes appear within 30 minutes and are usually maximal within 60 to 90 minutes. The drug effects persist for about one hour.

If necessary, the effects of the drug can be abolished promptly by atropine [see **OVERDOSAGE**].

HOW SUPPLIED

Bethanechol Chloride Tablets, USP

5 mg tablets are white, 7/16" diameter, round flat faced bevel edge tablets; one side scored and debossed BCL bisect 5, one side debossed 832. They are supplied as follows:

Bottles of 100 tabletsNDC 0832-0510-00Bottles of 1,000 tabletsNDC 0832-0510-10Unit-dose cartons of 100
tabletsNDC 0832-0510-01

10 mg tablets are white, 7/16" diameter, round flat faced bevel edge tablets; one side scored and debossed BCL bisect 10, one side debossed 832. They are supplied as follows:

Bottles of 100 tabletsNDC 0832-0511-00Bottles of 250 tabletsNDC 0832-0511-25Bottles of 1,000 tabletsNDC 0832-0511-10Unit-dose cartons of 100
tabletsNDC 0832-0511-01

25 mg tablets are yellow, 7/16" diameter, round flat faced bevel edge tablets; one side scored and debossed BCL bisect 25, one side debossed 832. They are supplied as follows:

Bottles of 100 tablets	NDC 0832-0512-00
Bottles of 250 tablets	NDC 0832-0512-25
Bottles of 500 tablets	NDC 0832-0512-50
Bottles of 1,000 tablets	NDC 0832-0512-10
Unit-dose cartons of 100 tablets	NDC 0832-0512-01

50 mg tablets are yellow, 7/16" diameter, round flat faced bevel edge tablets; one side scored and debossed BCL bisect 50, one side debossed 832. They are supplied as follows:

Bottles of 100 tablets	NDC 0832-0513-00
Bottles of 500 tablets	NDC 0832-0513-50
,	NDC 0832-0513-10
Unit-dose cartons of 100 tablets	NDC 0832-0513-01

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 container as defined in the USP.

Keep out of reach of children.

Manufactured by UPSHER-SMITH LABORATORIES, LLC Maple Grove, MN 55369

Revised: 5/2020

PRINCIPAL DISPLAY PANEL - 5 mg Tablet Bottle Label

NDC 0832-0510-00

Bethanechol Chloride Tablets, USP

5 mg

100 Tablets **Rx only**

UPSHER-SMITH



PRINCIPAL DISPLAY PANEL - 10 mg Tablet Bottle Label

NDC 0832-0511-00

Bethanechol Chloride Tablets, USP

10 mg

100 Tablets **Rx only**

UPSHER-SMITH

NDC 0832-0511-00 Bethanechol Chloride Tablets, USP	Each tablet contains: Bethanechol Chloride, USP 10 mg Usual Dosage: See package insert for full prescribing information. 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 container as defined in the USP.
10 mg	Keep out of reach of children.
100 Tablets Rx only	Manufactured by UPSHER-SMITH LABORATORIES, LLC Maple Grove, MN 55369
UPSHER-SMITH	© 2017 Upsher-Smith Laboratories, LLC 112980-01 R0917 =>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>

PRINCIPAL DISPLAY PANEL - 25 mg Tablet Bottle Label

NDC 0832-0512-00

Bethanechol Chloride Tablets, USP

25 mg

100 Tablets **Rx only**

UPSHER-SMITH

NDC 0832-0512-00 Bethanechol Chloride Tablets, USP	Each tablet contains: Bethanechol Chloride, USP 25 mg Usual Dosage: See package insert for full prescribing information. 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 container as defined in the USP.
25 mg	Keep out of reach of children.
100 Tablets Rx only	Manufactured by UPSHER-SMITH LABORATORIES, LLC Maple Grove, MN 55369
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PRINCIPAL DISPLAY PANEL - 50 mg Tablet Bottle Label

NDC 0832-0513-00

Bethanechol Chloride Tablets, USP

50 mg

100 Tablets **Rx only**

UPSHER-SMITH



BETHANECHOL CHLORIDE

bethanechol chloride tablet

Product Information					
Product Type	HUMAN PRESCRIPTION DRUG	ltem Cod	e (Source)	NDC:0	832-0510
Route of Administration	ORAL				
Active Ingredient/Active	Moiety				
-	redient Name		Basis of Stre	ength	Strength
Bethanechol Chloride (UNII: H4	QBZ2LO84) (Bethanechol - UNII:004	F72P8F4)	Bethanechol Chl	oride	5 mg
Inactive Ingredients					
	Ingredient Name			S	trength
	3U4)				
SILICON DIOXIDE (UNII: ETJ7Z6X					
lactose monohydrate (UNII: EU/26X					
	Q57Q8I5X)				

P	roduct Chara	acteris	tics					
	olor		WHITE	Score			2 piece	25
	nape		ROUND	Size			11mm	
	avor			Imprint Co	de		BCL;5;	832
Co	ontains							
Pa	ackaging							
#	ltem Code		Package	Description	Ма	rketing Star Date	rt I	Marketing End Date
1	NDC:0832-0510- 00	100 in 1 Product	BOTTLE; Type	e 0: Not a Combinatio	on 01/24	/2010		
2	NDC:0832-0510- 10	1000 in 1 Product	BOTTLE; Typ	e 0: Not a Combinat	ion 01/24	/2010		
3	NDC:0832-0510- 01	100 in 1	CARTON		01/24	/2010		
3	NDC:0832-0510- 89	1 in 1 BL Product	ISTER PACK; ⁻	Type 0: Not a Combin	ation			
R/	larkoting	Inform	nation					
Iv	larketing						-	
	Marketing Category	Ap		umber or Monog Citation	aph Ma	arketing Sta Date	rt	Marketing End Date
	IDA		040633		01/2	4/2010		

BETHANECHOL CHLC bethanechol chloride tablet	DRIDE				
Product Information					
Product Type	HUMAN PRESCRIPTION DRUG	Item Cod	e (Source)		832-0511
Route of Administration	ORAL	item cou	e (Source)	NDC.0	052 0511
Active Ingredient/Active	Moiety				
Ingr	edient Name		Basis of Stre	ngth	Strength
Bethanechol Chloride (UNII: H4Q	BZ2LO84) (Bethanechol - UNII:004	4F72P8F4)	Bethanechol Chlo	oride	10 mg
Inactive Ingredients					
	Ingredient Name			S	Strength
SILICON DIOXIDE (UNII: ETJ7Z6XB	U4)				
lactose monohydrate (UNII: EWQ	57Q8I5X)				
magnesium stearate (UNII: 7009	7M6I30)				
MICROCRYSTALLINE CELLULOSI	(UNII: OP1R32D61U)				

sc	DIUM STARCH	GLYCOLA		ro (UNII: 5856J3G2A2)			
Pı	roduct Chara	acteris	tics				
Сс	olor		WHITE	Score		2 piec	es
Sł	nape		ROUND	Size		11mm	l
Fla	avor			Imprint Code		BCL;1	0;832
Co	ontains						
Pa	ackaging						
#	ltem Code		Package Des	scription	Marketing St Date	tart	Marketing End Date
1	NDC:0832-0511- 00	100 in 1 Product	BOTTLE; Type 0: N	ot a Combination	01/24/2010		
2	NDC:0832-0511- 25	250 in 1 Product	BOTTLE; Type 0: No	ot a Combination	01/24/2010		
3	NDC:0832-0511- 10	1000 in 2 Product	1 BOTTLE; Type 0: I	Not a Combination	01/24/2010		
4	NDC:0832-0511- 01	100 in 1	CARTON		01/24/2010		
4	NDC:0832-0511- 89	1 in 1 BL Product	ISTER PACK; Type (): Not a Combination			
R.A	larkotina	Inform	mation				
IV	larketing						
	Marketing Category	Ар	plication Numbe Citat	er or Monograph ion	Marketing S Date	tart	Marketing End Date
AN	IDA	ANDA	040634		01/24/2010		

BETHANECHOL CHLC bethanechol chloride tablet	DRIDE				
Product Information					
Product Type	HUMAN PRESCRIPTION DRUG	ltem Cod	e (Source)	NDC:0	832-0512
Route of Administration	ORAL				
Active Ingredient/Active	Moiety				
Ingr	edient Name		Basis of Stre	ngth	Strength
Bethanechol Chloride (UNII: H4Q	BZ2LO84) (Bethanechol - UNII:004	F72P8F4)	Bethanechol Chlo	oride	25 mg
Inactive Ingredients					
	Ingredient Name			S	trength
SILICON DIOXIDE (UNII: ETJ7Z6XB	U4)				
lactose monohydrate (UNII: EWQ	57Q8I5X)				
magnesium stearate (UNII: 7009	7M6I30)				

sc	DIUM STARCH	GLYCOLATE TYPE A P	OTATO (UNII: 5856J3G2A2	2)		
Dð	C YELLOW NO.	10 (UNII: 355W5USQ30	G)			
FC	D&C YELLOW NO	D. 6 (UNII: H77VEI93A8)				
P	roduct Chara	acteristics				
	olor	YELLOW	Score		2 piece	25
	nape	ROUND	Size		11mm	
	avor		Imprint Code		BCL;25	;832
	ontains					
Pa	ackaging					
#	ltem Code	Package	Description	Marketing St Date	art	Marketing End Date
1	NDC:0832-0512- 00	100 in 1 BOTTLE; Type Product	0: Not a Combination	01/24/2010		
2	NDC:0832-0512- 25	250 in 1 BOTTLE; Type Product	0: Not a Combination	01/24/2010		
3	NDC:0832-0512- 50	500 in 1 BOTTLE; Type Product	0: Not a Combination	01/24/2010		
4	NDC:0832-0512- 10	1000 in 1 BOTTLE; Typ Product	e 0: Not a Combination	01/24/2010		
5	NDC:0832-0512- 01	100 in 1 CARTON		01/24/2010		
5	NDC:0832-0512- 89	1 in 1 BLISTER PACK; T Product	ype 0: Not a Combination			
Μ	larketing	Information				
	Marketing Category	Application Nu	Imber or Monograph Citation	Marketing S Date	tart	Marketing End Date
	IDA	ANDA040635		01/24/2010		

BETHANECHOL CHLC bethanechol chloride tablet	DRIDE				
Product Information					
Product Type	HUMAN PRESCRIPTION DRUG	ltem Cod	e (Source)	NDC:0	832-0513
Route of Administration	ORAL				
Active Ingredient/Active	Moiety				
Ingr	edient Name		Basis of Stre	ngth	Strength
Bethanechol Chloride (UNII: H4C	BZ2LO84) (Bethanechol - UNII:004	F72P8F4)	Bethanechol Chlo	oride	50 mg

	active Ingre	dients	5					
			Ir	ngredie	ent Name			Strength
SII	ICON DIOXIDE	(UNII: ET)	J7Z6XBU4)					
lad	tose monohydi	rate (UN	II: EWQ57Q8I	5X)				
ma	gnesium stear	ate (UNII	: 70097M6I30	0)				
MI	CROCRYSTALLI	NE CELL	ULOSE (UNII	I: OP1R32	2D61U)			
so	DIUM STARCH	GLYCOL	ATE TYPE A	ΡΟΤΑΤΟ) (UNII: 5856J3G2A2)	1		
D۵	C YELLOW NO.	10 (UNI	I: 35SW5USQ	(3G)				
FD	&C YELLOW NO	D. 6 (UNI	I: H77VEI93A8	8)				
_			_					
Pı	oduct Chara	acteris	stics					
Со	lor		YELLOW		Score		2 pieces	
Sh	аре		ROUND		Size		11mm	
Fla	vor				Imprint Code		BCL;50;8	832
Co	ntains							
_								
Pa	ckaging							
Pa #	ickaging Item Code		Packag	je Desc	ription	Marketing St Date	art	Marketing End Date
#		100 in 1 Product	BOTTLE; Typ		•		art	Marketing End Date
# 1 2	Item Code	Product	BOTTLE; Typ BOTTLE; Typ	pe 0: Not	a Combination	Date	art	_
# 1 2	Item Code NDC:0832-0513- 00 NDC:0832-0513- 50	Product 500 in 1 Product	BOTTLE; Typ BOTTLE; Typ 1 BOTTLE; Typ	pe 0: Not pe 0: Not	a Combination	Date 01/24/2010	art	
# 1 2 3	Item Code NDC:0832-0513- 00 NDC:0832-0513- 50 NDC:0832-0513-	Product 500 in 1 Product 1000 in Product	BOTTLE; Typ BOTTLE; Typ 1 BOTTLE; Typ	pe 0: Not pe 0: Not	a Combination a Combination	Date 01/24/2010 01/24/2010	art	
# 1 2 3 4	Item Code NDC:0832-0513- 00 NDC:0832-0513- 50 NDC:0832-0513- 10 NDC:0832-0513- 01	Product 500 in 1 Product 1000 in Product 100 in 1	BOTTLE; Typ BOTTLE; Typ 1 BOTTLE; Typ CARTON LISTER PACK;	pe 0: Not pe 0: Not ype 0: No	a Combination a Combination	Date 01/24/2010 01/24/2010 01/24/2010	art	
# 1 2 3 4	Item Code NDC:0832-0513- 00 NDC:0832-0513- 50 NDC:0832-0513- 10 NDC:0832-0513- 01 NDC:0832-0513-	Product 500 in 1 Product 1000 in Product 100 in 1 1 in 1 B	BOTTLE; Typ BOTTLE; Typ 1 BOTTLE; Typ CARTON LISTER PACK;	pe 0: Not pe 0: Not ype 0: No	a Combination a Combination ot a Combination	Date 01/24/2010 01/24/2010 01/24/2010	art	_
# 1 2 3 4 4	Item Code NDC:0832-0513- 00 NDC:0832-0513- 50 NDC:0832-0513- 10 NDC:0832-0513- 01 NDC:0832-0513-	Product 500 in 1 Product 1000 in Product 100 in 1 1 in 1 B Product	BOTTLE; Typ BOTTLE; Typ 1 BOTTLE; Typ CARTON LISTER PACK;	pe 0: Not pe 0: Not ype 0: No	a Combination a Combination ot a Combination	Date 01/24/2010 01/24/2010 01/24/2010	art	
# 1 2 3 4 4	Item Code NDC:0832-0513- 00 NDC:0832-0513- 50 NDC:0832-0513- 10 NDC:0832-0513- 01 NDC:0832-0513- 89	Product 500 in 1 Product 1000 in 1 1 in 1 B Product	BOTTLE; Typ BOTTLE; Typ 1 BOTTLE; Typ CARTON LISTER PACK; Mation	ре 0: Not pe 0: Not ype 0: No ; Туре 0:	a Combination a Combination ot a Combination Not a Combination	Date 01/24/2010 01/24/2010 01/24/2010		

Labeler - Upsher-Smith Laboratories, LLC (047251004)

Establishment					
Name			dress	ID/FEI	Business Operations
Upsher-Smith Laboratories, LLC				047251004	ANALYSIS(0832-0510, 0832-0511, 0832-0512, 0832-0513)
Establishment					
Name	Address	ID/FEI		Business Operations	
Upsher-Smith Laboratories , LLC		07911182	0 0511	• •	0832-0511, 0832-0512, 0832-0513) , PACK(0832-0510, 0832- 0832-0513) , manufacture(0832-0510, 0832-0511, 0832-

Revised: 5/2023