# HALOPERIDOL- haloperidol tablet Clinical Solutions Wholesale, LLC

Haloperidol Tablets, USP

#### WARNING

sed Mortality in Elderly Patients with Dementia-Related Psychosis Increased Moralisy in Elderly Platients with Demendia Related Psychosis Elderly patients with demension-related psychosis treased with using whether the second second second second second second second second second treased prior that the second second second second second second second treased patients using asymptotic alterpsychosis treased with the second varied, most of the deads appared to be either catallowscular (e.g., heart failure, sudeen deads) infections (e.g., promormal) in nature. Determination second second second second sector which the trengency of comparison of the second second second second sector which the trengency of comparison of the second second second second sector which the trengency of patients with demension of the second second sector which the trengency of patients with demension of the second second second second sector which the trengency of patients with demension of cond second second second second sector which the trengency of patients with demension of cond second s

### DESCRIPTION

Haloperidol is the first of the butyrophenore series of major tranquilizers. The chemical designation is 4-[4-(.p-chlorophenyl)-4-hydroxypiperidim)-4'-fluorobutyrophenone. It has the following structural

Each haloperidol tablet, USP intended for oral administration contains haloperidol, USP 5 mg or 10 mg, or 20 mg, in addition each haltet contains the following institute ingredients: calcium intentra, dibasit Viellow 101 Administration and the State State (State 1) and the State State (State 1) and the State State State Viellow 101 Administration Laboration 2016 and Administration and the State State (State 1) and the State State Laboration and the State State State (State 1) and the State State State (State 1) and the State Sta

CLINICAL PHARMACOLOGY -has not been clearly established.

### INDICATIONS AND USAGE

Haloperidol is indicated for use in the management of manifestations of psychotic disorders Haloperiols is indicated for use in the management of multi-stations of psychoic disorders. Haloperiols is indicated for the corrol of is can vocal ultraters of Tourner's Disorder is children of containing exploration is effective for the treatment of severes behavior problems in children of containing exploration of the proceeding of the state of the symmetry proceedings of motor activity with accompanying context Gisorders constraining of source and proceeding symptoms: implovity, officially sustaining anterion, aggressivity, mood hality, and poor forstration tolerance. Haloperiod is should be reserved for these two groups of children outly after failure to respond to psychotrapy or medication of the antantipsychotics.

### CONTRAINDICATIONS

Haloperidol is contraindicated in severe toxic central nervous system depression or comatose states from any cause and in individuals who are hypersensitive to this drug or have Parkinson's disease.

# WARNINGS

Increased Mortality in Elderly Patients with Dementia-Related Psychosis Elderly patients with dementia-related psychosis treated with antipsychotic drugs are at an incre risk of death. Haloperidol is not approved for the treatment of patients with dementia-related ps (see BOXED WARNING).

Cardiovas cular Effects Caronwacuur Linets Cases of suddendarding, GT-prolongation, and Torsades de Pointes have been reported in patients receiving hia/peritôn. Higher than recommended donss of any formaliation of haloperitôn lapore trô associated with a higher risk of QT-prolongation and Torsades de Pointes. Althoug classes have been reported even in the absence of predisposing facture, particular cataoni is advised in treating patients with other QT-prolonging conditions (netricad) experison particular cataoni is advised in treating patients with other QT-prolonging conditions (netricad) experison particular cataoni sa divised in treating patients with other QT-prolonging conditions (netricad) experison patients (particular) physicalients and hypomagneemia (harge short prolong QT, underlying cardiar absormatilies, hypodysolidsm and familia long QT-yardowe).

### Tardive Dyskinesia

A surver crystancial A syndrome consisting of potentially irreversible, involuntary, dyskinetic movemens may develop in patients treated with antipsychotic drugs. Although the prevalence of the syndrome appears to be indigets atom gate dealery, sepecially develop's women, it is insolite to rety quore prevalence estima-to predict, at the integriton of antipsychotic treatment, which patients are likely to develop the syndrome. Whether antipsychotic drug predicts differ in their portential coase turbide opskyttenia is subatown. When an amplycholic ang produces mitter in mer potential to cause arraye opsanesia is unanown Both the risk of welpoping tardrey dysknesia and the likelihood that it will become irreversible are believed to increase as the duration of treatment and the total cumulative dose of antipsychotic drugg administered to the patient increase. However, the syndrome can develop, although much less commonly, after relatively brief treatment periods at low doses.

commutsy, after retainvey hiel resument perioda al too doses. There is no known stament for estabilistic cases of and/or dopkinesia, adfounds the syndrome may rentis, partially or completely, if antipy-choic restament is withdrawn. Antipy-thoic resument, iself, however, may suppress (or partially suppress) he signs and symposon of the syndrome and dheredy may possibly musk the underlying process. The effect that symptomatic suppression has upon the long-term comise of the syndrome is unknown.

course of the syndrome is unknown. Given-base conditions, misrycychodic drugs should be prescribed in a muner shu is mort likely to minimize the occurrence of tardive dyskiresia. Chronic antipyscholic resumes should generally be reserved for patients who suffer from a chronic illensis shu, i) is shown to respond a patient drugs, and, 2) for whom alternative, equally effective, har potentially less harmful resumers are not shuilble er appropriate. In patients who or regime chronic treatment, the smalles charge and the submets duration of treatment producing a statisfactory clinical response should be tought. The rest for statisfactor of the statistical statistical statistical statistical statistical statistical statistical distantion of tardive dyskinesis appear in a patient on antipsycholics, drug discontinuinal matchields the constructive producing a statistic statistical statistical statistical statistical distantiation of the statistical s

### Neuroleptic Malignant Syndrome (NMS)

A potentially late symptom compete sometime reference and a so Neurologic Multiplean Spechrone (NMS) hybersensisted and the source of the sour

(thabdomylshi) and care renal failure. The diagnostic evaluation of patterns with this synchrone is complicated. In arriving at a diagnosti, it is important to identify cases where the clinical presentation includes both serious medical litess (e.g., parmonia, systemic relevance) in teredo or inadequarks (trended varagamatial sign and symptom (TFS). Other important considerations in the differential diagnosis include central symptom (TFS). There important considerations in the differential diagnosis include central trends and the symptom of the symptom of the symptomic or any systemic diagnosis (trends and the drags not essential to concurrent therapy, 2) intensity symptomic treatment and medical monitoring, there is no general agreement about specific pharmacological reanners regimes for uncomplicated NMS.

If a patient requires sutpsychoid dug treatment after recovery from NNS, the potent introduction of drug therapy should be carefully considered. The patient should be carefully movitored, since returnerses of NAS have been reported. Hyperprysexia and heat strole, not associated with the above symptom complex, have also been reporte with hadperichi.

### Falls

r an Haloperidol may cause somnolence, postural hypotension, motor and sensory instability, which may lead to falls and, consequently, fractures or other injuries. For patients with diseases, conditions, or medications that could exacerbate these effects, complete fall risk assessments when initiating anipsychotic treatment and recurrently for patients on long-term anipsychotic therapy.

Usage In Pregnancy Non-teratogenic Effects

Non-terangeme EHecs Nonnes equose do antipox/holic drugs, during the third trimester of pregnancy are at risk for extrapyramidal and/or withdrawal symptoms following delivery. There have been reports of agitation, hypertonia, hypotra terror, somolecer, respiratory distrust sand leeding distorted in these renorates. These complications have varied in severity, while in some cases symptoms have been soft-limited, in other cases renorates have required intensive care unit support and prolonged boogalitation. ridol should be used during pregnancy only if the potential benefit justifies the potential risk to

the feats. Redens gives 2 to 20 times the usual maximum human dose of haloperiddol by oral or parenteral routes aboved an increase in incidence of resorption, reduced ferling, delayed delivery and page montality. No-haber observed in the circle view 15 into with the usual maximum human dose. Cleft palae in nice agness in to be anonpecific response to stress or marinoal inhalance as well as to a variety of drugs, and here is nevidence to relate his phenomenous predictable human factor must obtain agges. is ne vidence to relate his phenomenon to predicable humanrisk for most of these agrees. There are no well concolled studies with huboprido lin preguato womes. There are reports, however, of cases of link malformations observed following metarula are of haloperiola along with other droggest witch have supercised transgering phenetia during the first integree of preguators. Casand are alondowing many de host haloperiols, his dug shadoh be used during generatory or in worme likely to become preguatory if the bacefit clearly patifies a patiental risk to the fease. Indicas should not be nareed during during the similar to the second strategest preduction of the second strat

### Combined Use of Haloperidol and Lithium

Combined Use of Halperidal and Lihhum Amerophologuida polymore themacrined by weahness, lethargy, fever, trendomstess and condusion, estrapycatifiel symptome, letharcytrisis, elevands terum enzymes, BUN, and BEBS followed by inversible beard damage has occurred in a few gattern treatwork with libitant gales halperidal. A casaal reliationship between here events and the concontant administration of linham and halperidal has no bene stabilished, howevere, gattern receiving such contained drangs should be manistered closely for early evidence of neurological toxicity and restment discontinued promptly if such signs appear.

### General

Haloperidol may impair the mental and/or physical abilities required for the performance of hazardous takes such as operating machinery or driving a motor vehicle. The ambulatory patient should be warned accordingly.

The use of alcohol with this drug should be avoided due to possible additive effects and hyp

#### PRECAUTIONS

Leukopenia, Neutropenia and Agranulocytosis In clinical trial and postmarketing experience, events of leukopenia/neutropenia have been reported temporally related to antipsychotic agents, including haloperidol tablets USP. Agranulocytosis (including fatal cases) has also been reported.

unauong tami cases) tan also been reported. Second and the second

Loursee dual resolvery indicated quadrandy in gaterine: Happenbolin, and happenbolin, an

should be used. receiving anticonsubant medications, with a history of seizures, or with EEG abnormalities, because hadoperiols imp lower the counsilive threshold. If indicated, adequae anticonsubant with hows and lenger, or with history of all lenger traceations to dugs. receiving anticongulanes, since an isolated instance of interference occurred with the effects of one anticongulant (phenelindore).

If concomiant antiparkinon medication is required, it may have to be continued after haloperidol-discontinued because of the difference in neuration rates. If both are discontinued simultaneously, estimation and the second concountering with haloperidol.

As with other artipsychotic agents, it should be noted that haloperidol may be capable of potentiating CNS depressants such as anesthetics, opiates, and alcohol.

Use support the standard structure of the structure of th

When haloperidol is used to control mania in cyclic disorders, there may be a rapid mood swing to depression. aepression. Severe neurotoxicity (rigidity, inability to walk or talk) may occur in patients with thyrotoxicosis who are also receiving antipsychotic medication, including haloperidol.

Server ensurancies (registant), manimulty is wain, de tala jung decidi in planettes with intyrotaticotas won hormangenic generational de hologorial do area. Note in the Area Schandell in retoronal activation assays, Negative or inconsistent positive findings, have bere no handed in in vitro adia invois studies of effects of hologorial on chromosome structure and annubers. The available cryogenetic evidence is considered too inconsistent to be conclusive at this time. Carcinogencity studies using on al hologorialo were conduced in Wister rans (doned at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and in Allans Sviss mice (dosed at up to 5 mg/kg daily for 24 months) and the fore for theorem (at the dosed at up to 20 times the exaual duily significant increase in interactive of a realistical mice (dose at the dose at a statistical dosed at the statistical dosed at the statistical significant increase in interactive) and alla duily dose for chronic or revistant patients at the month duily significant differences in increasers of a dosed at the statistical significant differences in increasers in the evolution statistical traditional to mate realistical statisticantic differences in increduces or stati

and not, ware user was assuming any significant intractor in philing glast regulation. Build of the second seco

Pregnancy: Non-teratogenic Effects

Non-transgenic Lifects Nonatise exposed a antipsycholic drugs, during the third trimester of pregnancy are at risk for extrapyrantial and/or withdrawal symptoms following delivery. There have been reports of agitat hypertonia, hypotonia, tremori, somolecer, respiratory distress and feeding disorder in these reom These complications have varied in severity; while in some cases symptoms have been self-limite other cases moments have required intensive care unit support and prolonged thospital lazion. Haloperidol should be used during pregnancy only if the potential benefit justifies the potential risk to the feus.

Pediatric Use Safety and effect ness in pediatric patients have not been established.

Geriatric Use 

### ADVERSE REACTIONS

Cardiovascular Effects

Cardiovarcular Effects Tachycardia, hypotension, and hypertension have been reported. QT prolongation and/or ventricular anhydmins have also been reported, in addition to ECG pattern changes compatible with the polymorphons configuration of transa de poinses, and may occurs unner frequently with high does and in preclaposed patients (see WARNINGS and PRECAUTIONS). Cases of sudden and unseptend deals have been reported in association with the administration of haloperiodi. The nature of the evidence makes it impossible to determine definitively what role, II any haloperiodi played in the outcome of the reported cases. The possibility that haloperiodic cased deals canner, of course, be excluded, that it is to be lapt in mind that sudden and unsequend ded may occur in pycholic patterns when they go surreade of when they are tested with other antipychoic drages.

(P) toting neural and the second seco

Up yours. Closs effect: Symptoms of dystonia, prolonged abnormal contractions of muscle groups, may occur in susceptible individuals during the first few days of remarker. Dystonic symptoms include: space of the benchmark, and/or protonion of the mogark. While these symptoms can concur at level works, they occur more frequently and write greater severity with high pottery and a higher doses of first generation imprychoic drugs. A deviated first deviate dystonia is dones or first generation imprychoic drugs. A deviated first deviate dystonia is observed in mulei and younger egg troups.

### WITHDRAWAL EMERGENT NEUROLOGICAL SIGNS

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#### TARDIVE DYSKINESIA

LARGUYE DISARDESIA As with all analysochoic agents, haloperidol has been associated with persistent dyskinesias. Tardi dyskinesia, a synchrone consisting of potentially irrerestible, incolotatory, dyskinetic answerners, mu appear is income painteen soling are methodary or may occur and ring theory has been discontinued, and the synchronized and the hybridical involutionary movements of logance, face, month origo in equile, patient checks, nucleiring of meant, cheving movements of logance and the synchronized and the synchronized and unreterment of extractionized and the truth.

novement of extension and the trunk. There is no haven effective reasonmer for tardive dyskinestic, antiparkinon agents usually do not alleviate the symptoms of this symptome. It is suggested that all antipycoind agents the discontinger degrees of which consistent and alleviate agents of the symptome target of the symptome degrees or which constitutions agents alleviate agents the symbolic may be maded. It has been reported that first venturical antiverse of the tongor may be maded.

TARDIVE DYSTONIA

# Tardive dystoria, not associated with the above syndrome, has also been reported. Tardive dysto characterized by delayed onset of choreic or dystoric movemens, is often persistent, and has the potential of becoming irreversible.

### OTHER CNS EFFECTS

normia, reviso ar terior network and the second sec

#### Body as a Whole

Neuroleptic malignant syndrome (NMS), hyperpyrexia and heat stroke have been reported with haloperidol (see WARNINGS for further information concerning NMS).

### Hematologic Effects

### Liver Effects

aired liver function and/or jaundice have been reported. Dermatologic Reactions

Maculopapular and acneiform skin reactions and isolated cases of photosensitivity and loss of hair.

### Endocrine Disorders

Lactation, breast engorgement, mastalgia, menstrual irregularities, gynecomastia, impotence, increased libido, hyperglycemia, hypoglycemia and hyponatremia.

#### Gas trointes tinal Effects

Anorexia, constipation, diarrhea, hypersalivation, dyspensia, nausea and yomiting

# Autonomic Reactions Dry mouth, blurred visio

vision, urinary retention, diaphoresis and priapism Respiratory Effects Laryngospasm, bronc

, bronchospasm and increased depth of respiration. Special Senses Cataracts, retinopathy and visual disturbances.

Postmarketing Events

Hyperanmonemia has been reported in a 5½ year old child with citrullinemia, an inherited disorder of ammoria excretion, following treatment with haloperidol.

## OVERDOSAGE

OVERDOSALE Maintenation In general, the symptoms of overdosage would be an esuggradion of howen plarmacologic effects and adverse reactions, be most prominent of which would be: 1) severe entrapyramidal reactions, 2) hypotension, or 3) sedands. The patient would appear commons with respiratory depression and software and the provide setable and the second setable and the second setable and by the standard by the adjust of the second setable and the second setable and the demonstrated by the adjust of the second setable considered and the second setable and the demonstrated by the adjust of the second setable and the second setable and the second setable and the most of the second setable considered. (for further information regarding useds de pointer, please refer to ADVERSE REACTIONS).

refer to ADVERSE REACTONS). **Treasment** Gauric Houge or induction of events should be carried out immediately followed by admistrations divident chronical. Since there is no specific andobu, resament is primarily supportive. A patent airway must be established by use of an enopharyngel airway or enduratable able or, in prolonged cases of come, by trachesomy. Respiratory depression may be contracted by stifficial respiration; methods and respirators. Hypotension and circulatory collapse may be contracted by stifficial respirators. Hypotension and the statement of the statement of the statement of the statement of the statement inversion flush, Bason, or core entered babant; and supports agents are its a metamining intervents flush, Bason, or core entered babant; and supports agents are its an enterement. Hypotensis and the statement of the statement of

#### DOSAGE AND ADMINISTRATION

DOSAGE AND ADMINISTRATION There is considered variation from guider to pattern in the amount of medication required for treatment. As with all antipyychoic drugs, shouge should be individualized according to the redsh and required the antipyychoic drugs. Shouge should be individualized according to the redsh and required the antipyychoic drugs. The antipyychoic drugs and the should be antipyychoic drugs and the antipyychoic drugs and antipyychoic drugs and antipyychoic drugs and Children debilismed or geniatic gathers, as well as those with a history of adverse tractions to children debilismed or geniatic gathers, and well as those with a history of adverse tractions to children debilismed or geniatic gathers, and well as those with a history of adverse tractions to children debilismed or geniatic gathers, and well as those with a history of adverse tractions to children debilismed or geniatic gathers, and well as those with a history of adverse tractions to children debilismed or geniatic gathers, and well as those with a history of adverse tractions to children debilismed or geniatic gathers, and a children debilismed or history and children debilismed or geniatic gathers, and a children debilismed or history and children debilismed or geniatic gathers and a liver debilismed or history and children debilismed or geniatic gathers and a liver debilismed or history and children debilismed or geniatic gathers and a liver debilismed or history and children debilismed or geniatic gathers and a liver debilismed or history and children debilismed or geniatic gathers and a liver debilismed or history and children debilismed or geniatic gathers and a liver debilismed or history and children debilismed or geniatic gathers and a liver debilismed or history and children debilismed or geniatic gathers and a liver debilismed or history and children debilismed or history and adverse trank and history adverse trank and

# Oral Administration INITIAL DOSAGE RANGE

Adults

 Moderate Symptomatology
 0.5 mg to 2 mg b.i.d. or t.i.d.

 Severe Symptomatology
 3 mg to 5 mg b.i.d. or t.i.d.

# To achieve prompt control, higher doses may be required in some cases.

Geriatric or Debilitated Patients

0.5 mg to 2 mg b.i.d. or t.i.d. 3 mg to 5 mg b i d Patients who auens verely disturbed or inadequately controlled may require dosage adjustment. Daily do

Children The following recommendations apply to children between the ages of 3 and 12 years (weight range 15 to 40 Ag). Malaportialo is not intended for children under 3 years old. Therapy should begin at the lowest does possible (0.5 mg per day). It required, the does should be increased by an increment of 0.5 and a 50 T-20 junctuals undit the desire the demengencie effect is obtained, (see clam below).

The total dose may be divided, to be given b.i.d. or t.i.d.

 Bycknik: Disorders
 Bo35 mg/kg/day to 0.15 mg/kg/day

 Non-Psychiatic Relation Disorders and Tourner's Disorder
 Biosofter (Section 2010)
 Biosofter (Section 2010)

Maintenance Dosage Upon achieving a satisfactory therapeutic response, dosage should then be gradually reduced to the lowest effective maintenance level.

Switchover Procedure

Switchover Procedure The cord form studied supplort the injectable as soon as practicable. In the absence of binavailability studies establishing likeoquivalence briveen these two dosage forms, the following guidelines for administered in the preceding 24 shows may be used. Since this dose is only animital estimate, it is recommended that careful monitoring of clinical sign and symposin, including clinical efficiency, sedation, and adverse effects, be carried outperiodically for the first several days following the accomplicated. Depending on the pattern's clinical size of symposite of days of a low equivalence accomplicated. Depending on the pattern's clinical size of symposite of adverse of the pattern's accomplicated. Depending on the pattern's clinical stands, the first oral dose should be given within 12-24 bars following the interpretent dose.

### HOW SUPPLIED

Haloperidol Tablets USP, 5 mg are green, capsule-shaped, flat-faced, beveled-edge tablets debossed with the logo of ZC', '07' and partial bisect, on one side and plain on the other side and are supplied as follows:

follows: NOC 63822-079-10 in houtes of 100 tables NDC 63822-079-10 in houtes of 1000 tables Habyeriold Tables USP, 100 are tables press, granule-shaped, flat faced, heveled-edge tables debosted with the logo of 732, 000 and partial bisect, on one side and plain on the other side and are signified as follows: NDC 63822-080-06 in houtes of 30 tables NDC 63822-080-06 in houtes of 1000 tables NDC 63822-080-10 in houtes of 1000 tables NDC 63822-080-10 in houtes of 1000 tables

Haloperidol Tablets USP, 20 mg are coral, capsule-shaped, flat-faced, beveled-edge tablets debossed with the logo of 'ZC', '09' and bisect on one side and plain on the other side and are supplied as follows:

with the log of 72C, 109 and black or none side and plain on the other side and are supplied as follow NDC 683372-018-01 in bottlers of 1040 tablets Store at 30° to 25°C (68° to 77°F) [Sec USP Controlled Room Temperature]. Disprese in a tigk hip (hip-resistant constrainter). All trademarks are the property of Zyloas group. Call your deters for medical advice about side effects. You may report side effects to FDA at 1-800+104.

Manufactured by: Cadila Healthcare Ltd.

### Ahmedabad, India Distributed by: Zvdus Pharr ceuticals USA Inc. ton, NJ 08534 Pennington, NJ 08534 Rev.: 11/16 Revision Date : 2016/11/12 Repackaged by: Clinical Solutions Wholesale Franklin, TN 37067

Package/Label Display Panel

CLINICAL Solutions WHOLESALE Haloperido 5mg Tables NDC: 58118-1079-8 Qty: 30 OTR: 00308118107476 
 Mg, NDC: 68332-3077-01
 GTW: 02358118107863

 Mg, Lot #: SAMPLE123
 Betch #: SAMPLE436

 Mg, Joy Arban Pramaceedicals (JSA) Inc.
 Eps: 2020 Jul

 Penningfore, NL, USA, 68554
 SN: "YMMERCOD0000

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HALOPERIDOL

Product Information HUMAN FRESCRIPTION DRUG Irem Code (Source) NDC-581 ration ORAL Anute of Art

Active Ingredient/Active Molety
Ingredient Name Basis of Strength
Strength
Mai OPERIDOL (UNE 2522781.DD) (MALOPERIDAL 5 mg Inactive Ingredients Ingredient Name Strength

CALCIUM STEARATE (UNE: 776XM7047L)	
D&C YELLOW NO. 10 (UNIX 355W5USQ3G)	
DIBASIC CALCIUM PHOSPHATE DIHYDRATE (UNIt: O7TSZ97GEP)	
FINC BLUE NO. 1 (UNE HIR47K3TBD)	
PO VIDO NE K30 (UNIE U725QWY32X)	
SOBUM STARCH GLYCOLATE TYPE A POTATO (UNE 585633G2A2)	
STARCH, CORN (UNE 08232NY35J)	

Product Character	istics		
Color	green (GREEN) Sco	re	2 pieces
Shape	OVAL (CAPSULE) Size		10 mm
Flavor	Imp	rint Code	ZC;07
Contains			
Packaging			
# Item Code	Package Description	Marketing Start Date	Marketing End Date
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Marketing Info	rmation		1
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Marketing Info Marketing Category ANDA Labeler - Clinical S	rmation Application Number or Menograph Citation ANDA077580	Marketing Start Date	Marketing End Date

Establishment New Address ID913 Business Operations
Charls Salaina Models, LLC (2017)2542 Preprint 2017 12542 Preprint 2017 2017
Revine: 72013 Chird Scharms Webriek, LLC