# METHYLPHENIDATE HYDROCHLORIDE - methylph hydrochloride solution Ascend Laboratories, LLC

# HIGHLIGHTS OF PRESCRIBING INFORMATION These highlights do not include all the information needed to use METHYLPHENDATE for METHYLPHENDATE HYDROCHADROBE ORAL SOLUTION. METHYLPHENDATE HYDROCHADROBE ORAL SOLUTION. METHYLPHENDATE HYDROCHADROBE ORAL SOLUTION.

## WARNING: ABUSE AND DEPENDENCE See full prescribing information for complete boxed warning.

CNS stimulants, including methylphenidate hydrochloride oral solution, other methylphenidate- containing products, and amphetamines, have a high potential fo abuse and dependence. (5), 13, 23, 33)
 Assess the risk of abuse prior to prescribing and monitor for signs of abuse and dependence while on therapy. (51, 52)

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# WARNINGS AND PRECAUTIONS Manimes and preductions Manimes and preductions more and p

## ADVERSE REACTIONS Common adverse reactions: tachycardia, papitations, headache, insomnia, anxiety, hyperhidrosis, weight loss, decreased appetite, dry mouth, nausea, and addorniard jani, (0) (6)

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See	17	for PATIENT	COUNSELING	INFORMATION.	
					Revised: 10/2021

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#### FULL PRESCRIBING INFORMATION

## WARNING: ABUSE AND DEPENDENCE

CNS stimulants, including methylphenidate hydrochibride oral solution, other methylphenidate- containing products, and ampletamines, have a hydrocentrylphenidate- containing products, and ampletamines, have a to prescribing, and monitor for signs of abuse and dependence while on therapy (see Warnings and Precautions (5.1), Drug Abuse and Dependence (9.2, 9.3)).

#### 1 INDICATIONS & USAGE

- Methylphenidate hydrochloride oral solution is indicated for the treatment of: Attention Deficit Hyperactivity Disorder (ADHD) in adults and pediatric patients 6 years of age and older Narcolepsy

### 2 DOSAGE AND ADMINISTRATION

2.1 Pretreatment Screening

# Prior to initiating treatment with methylphenidate hydrochloride oral solution, assess for the presence of cardiac disease (i.e., perform a cardful history including family history of sudden death or ventricular arrhythmia, and physical examination) [see Warnings and Precautions (5.2)].

Assess the risk of abuse prior to prescribing, and monitor for signs of abuse and dependence while on therapy. Maintain careful prescription records, educate patients about abuse, monitor for signs of abuse and overdose, and periodically re-evaluate the need for methylphenidiate hydrochloride or al solution use. *[see Boxed Warning, Warnings and Precautions (SJ.), Drug Abuse and Dependence (9)*].

#### 2.2 General Dosing Information Pediatric Patients 6 years of Age and Older

To encommended starting dosage 6 5 mg orally twice daily before breakfast and lunch (preferably 30 to 45 minutes before meak). Increase the dosage gradually, in increments of 5 mg to 10 mg weekly. Daily dosage above 60 mg is not recommended.

#### Adults

Administer orally in divided doses 2 or 3 times daily, preferably 30 to 45 minutes before meals. The maximum recommended daily dose is 60 mg. The average dosage is 20 to 30 mg daily. For adult patients who are unable to sleep if medication is taken late in the day, administer the last dose before 6 p.m.

Pharmacological treatment of ADHD may be needed for extended periods. Periodically re-evaluate the long-term use of methylphenidate hydrochloride oral solution, and adjust the dosage as needed.

#### 2.3 Dosage Reduction and Discontinuation

If paradoxical aggravation of symptoms or other adverse reactions occur, reduce dosage, or, if necessary, discontinue methyphenidate hydrochonide oral solution. Methyphenidate hydrochonide anal solution should be periodically discontinued to assess the pediatric patient's condition. If improvement is not observed after appropriate dosage adjustment over a one-month period, discontinue methyphenidate hydrochonide or al solution.

#### DOSAGE FORMS & STRENGTHS

Methylpheniate hydrochbride oral solution is a clear colorless to pale yellow, grape flavor solution available in a 500 mL bottle in the following strengths: • 5 mg per 5 mL • 10 mg per 5 mL

**4 CONTRAINDICATIONS** 

- 4 CONTRAINDICATIONS Wethybenicate hybricarbindie oral solution is contraindicated in patients: with innown hybricsmetality to methybenetalise or other components of anyonedmust any advantication is any term or the solution of the solution of the methybenetidate (see Adverse Reactions (B)). receiving concentant treatment with monamine oxidase inhibitors (MAOIs), or within 14 days following discontinuation of treatment with an MAOI, because of the risk of hypettensive crises (see Days Interactions (7)).

5 WARNINGS AND PRECAUTIONS

5.1 Potential for Abuse and Depend CS strunds, including methybenidate hydrochloride oral solution, other methybenidate-containing products, and amphetamines, have a high potential for abuse and dependence. Assess ther isk of abuse prior to prescribing, and monitor signs of abuse and dependence while on therapy Isee Boxed Warning, Drug Abuse and Dependence (2, 2, 5, 3).

5.2 Serious Cardiovascular Reactions Subder death, store and mycacital infarction have been reported in adults with CNS studient treatment at recommended doses. Sudden death has been reported in peliabric patients with structural cardiac abnormalities and other serious heart problems taking CNS stimulants at recommended doses for ADHD. Avoid use in patients with hown structural cardiac abnormalities, cardiomyopathy, serious heart hydrim abnormalities, coronary artery disease, and other serious heart problems. Further evaluate patients who develop exertional check patients syncope, or arrhythmias during methythmiate hydrochichties or al solution treatment.

5.3 Blood Pressure and Heart Rate Increases

CNS stimulants cause an increase in blood pressure (mean increase approximately 2 to 4 mmHg) and heart rate (mean increase approximately 3 to 6 bpm). Individuals may have larger increases. Monitor all patients for hypertension and tachycardia. 5.4 Psychiatric Adverse Reactions

Exacerbation of Pre-Existing Psychosis

CNS stimulants may exacerbate symptoms of behavior disturbance and thought disorder in patients with a pre-existing psychotic disorder. Induction of a Manic Episode in Patients with Bipolar Illness

CNS stimulants may induce a manic or mixed mood episode in patients. Prior to initiating treatment, screen patients for risk factors for developing a manic episode (e.g., depressive symptoms or a family history of suicide, bipolar disorder, or depression).

New Psychotic or Manic Symptoms CAS struitures, at recommended doese, may cluse psychotic or manic symptoms (e.g., halkuchatbons, CAS struitures, at recommended doese, may cluse psychotic or manic symptoms (e.g., halkuchatbons, or mania, if such symptoms occur, consider discontinuing methylphenklate hydrochoride oral solution. In a pooled analysis of multiple short-term, placebo-controlled studies of CNS struitures, psychotic or manic symptoms occurred in approximately 0.1% of CNS stmulant-treated patients, compared to 0 in placebo-treated plants.

5.5 Priapism

3-3 Fraipen
Probinged and painful erections, sometimes requiring surgical intervention, have been reported with methylphenidate products in both pediatric and adult patientis. Praipien dury provide with methylphenidate products in both pediatric and adult patientis. Praipien dury provide dury provide a subscription of the developed after some time on the drug, often subscription adult of the develop after some time on the drug, often subscription adult of the develop after some time on the drug, often subscription adult of the develop after some time on the drug, often subscription to an increase in dose. Priapism has also appeared during a period of drug withdrawal (drug holidays or during discontinuation). Patients who develop after subscription after subscription advection advection and painful erections should seek.

5.6 Peripheral Vasculopathy, Including Raynaud's Phenomeno 5.6 Peripheral Vasculopathy, Including Baynaud's Phenomenon Stimularis used to treat AUP(), nctuding methydyneniadia hydrochloridi oral solution, are associated with peripheral vasculopathy, including Baynaud's phenomenon. Signs and symptoma: are usually intermittent and mith, however, very rar expudse peripheral vasculopathy, including Baynaud's phenomenon, were observed in peripheral vasculopathy, including Baynaud's phenomenon, were observed in postmarketing reports at different lines and at therapeutic doses in all age groups throughout the course of treatment. Signs dipatic Atange's in encessary during treatment with AUPD stimulants, Further clinical evaluation (e.g., rheumatobgy referral) may be appropriate for certain patients.

#### 5.7 Long-Term Suppression of Growth

lants have been associated with weight loss and slowing of growth rate in pediatric patients.

Careful follow-up of weight and height in children ages 7 to 10 years who were randomized to either methylphenidate or non-medication treatment groups over 14 months, so well as it naivables: 35 months to now mere synthesis and the second suggests that consistently medicated children (i.e., treatment for 7 days per wesk throughout the years have a temporary solwing in growth rate (on average, a total of about 2 cm less growth in height and 2.7 kg less growth in weight over 3 years), without evidence of growth rebound during the period of development.

Closely monitor growth (weight and height) in pediatric patients treated with stimulants, including methylphenidate hydrochloride oral solution, Patients who are not growing or gainite hydrochloride as expected may need to have their treatment interrupted. solution that we not been established in pediatric patient less than 6 years of age.

#### 6 ADVERSE REACTIONS

The following adverse reactions are discussed in more detail in other sections of the labeling:

- The following adverse reactions are discussed in more detail in other sectors of the label Abuse and Dependence (see Warnings and Precautions (5.1), Drug Abuse and Dependence (9.2.9.3) Known hypersensitive (see Warnings and Arecautions (4), Hypertensitive crisis when used concentrating with monoamine oxidase inhibitors (see Contraindications (4), Drug Interactions (7)) Serious cardiovacular reactions (see Warnings and Precautions (5.2)) Biodo pressure and heart rate increases (see Warnings and Precautions (5.3)) Psychiatrix adverse reactions (see Warnings and Precautions (5.3)) Psychiatrix adverse reactions (see Warnings and Precautions (5.4)) Prophene (See Warnings and Precautions (5.5)) Precautions (5.1), Nichidam Reynaud: 5 phenomenon (see Warnings and Precautions (5.7)) Long-term suppression of growth (see Warnings and Precautions (5.7))

The following adverse reactions associated with the use of methylphenidate containing products were identified in clinical studies, postmarketing reports, or ilterature. Because some of these reactions were reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

Infections and infestations: nasopharyngitis

Blood and the lymphatic system disorders: leukopenia, thrombocytopenia, anemia, pancytopenia

Immune system disorders: hypersensitivity reactions, including angioedema and anaphylaxis, auricular swelling, bullous conditions, eruntions, exanthemas

Metabolism and nutrition disorders: decreased appetite, reduced weight gain and suppression of growth during prolonged use in pediatric patients

Psychiatric disorders: insomnia, anxiety, restlessness, agitation, psychosis (sometimes with visual and tactile hallucinations), depressed mood, affect lability, mi disorientation, libido changes

Nervous system disorders: headache, dizziness, tremor, dyskinesia including choreoatheetoid movements, drowsiness, convulsions, cerebral arteritis and/or occlusion, serotonin syndrome in combination with serotonergic drugs, migraine

Eye disorders: blurred vision, difficulties in visual accommodation, diplopia, mydriasis

Cardiac disorders: tachycardia, palpitations, increased blood pressure, arrhythmias, angina pectoris, sudden cardiac death, myocardial infarction, bradycardia, extrasystole

Respiratory, thoracic and mediastinal disorders: cough, pharyngolaryngeal pain, dyspnea Gastrointestinal disorders: dry mouth, nausea, vomiting, abdominal pain dyspepsia, diarrhea General disorders: fatigue, hyperpyrexia atobiliary disorders: abnormal liver function, ranging from transaminase elevation to severe hepatic injury

Skin and subcutaneous tissue disorders: hyperhidrosis, pruritus, urticaria, exfoliative dermatitis, scalp hair loss, erythema multiforme rash, thrombocytopenic purpura angioneurotic edema. erythema. fixed drug eruption

Musculoskeletal and connective tissue disorders: arthralqia, muscle cramps, rhabdomyolysis, myalqia, muscle twitching

Renal and urinary disorders: hematuria

Reproductive system and breast disorders: gynecomastia Urogenital disorders: priapism

Vascular disorders: peripheral coldness, Raynaud's phenomenon

Investigations: weight loss

#### 7 DRUG INTERACTIONS

## 7.1 Clinically Important Drug Interactions with methylphenidate hydrochloride oral solution

Table 1 presents clinically important drug interactions with methylphenidate hydrochloride oral solution.

## Table 1: Clinically Important Drug Interactions with methylphenidate

Monoamine Oxidase In	Concomitant use of MAOIs and CNS stimulants, including methylphenidate hydrochlorid
Clinical Impact:	pral solution, can cause hypertensive crisis. Potential outcomes include death, stroke
	myocardial infarction, aortic dissection, ophthalmological complications, eclampsia
	pulmonary edema, and renal failure [see Contraindications (4)].
	Concomitant use of methylphenidate hydrochloride oral solution with monoamine oxidase
Intervention:	inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment is contraindicated.
Antihypertensive Drug	
Clinical Impact:	Methylphenidate hydrochloride oral solution may decrease the effectiveness of drugs used
	to treathypertension [see Warnings and Precautions (5.3)].
	Monitor blood pressure and adjust the dosage of the antihypertensive drug as needed.
Intervention:	
Risperidone	
	Combined use of methylphenidate with risperidone when there is a change, whether a
Clinical Impact:	increase or decrease, in dosage of either or both medications, may increase the risk or extrapyramidal symptoms (EPS).
Intervention:	Monitor for signs of EPS

8 USE IN SPECIFIC POPULATIONS

## 8.1 Pregnancy

ncv Expo Pregna

There is a pregnancy exposure registry that monitors pregnancy outcomes in women exposed to ADHD medications, including methylphenidate hydrochoride oral solution, during pregnancy. Healthcare providers are encouraged to register patients by calling the National Pregnancy Registry for Psychostimulants at 1-866-961-2383.

#### Risk Summary

Published studies and postmarketing reports on methylphenidate use during pregnancy have not identified a drug-associated risk of major birth defects, miscarriage or adverse maternal or fetal outcomes. There may be risks to the fetus associated with the use of CNS stimulants use during pregnancy (see *Clinical Considerations*).

No effects on morphological development were observed in embryo-fetal development studies with oral administration of methylphenidate to pregnant rats and rabbits during organoge 12 and 19 times, respectively, the maximum recommended human dose (MRHD) of 60 mg/dsg view to adults on an mg/m basis. However, spin bifda was observed in rabbits at a dose 65 times the MRHD given to adults. A decrease in pup body weight was observed in a pit-end post-natial development study with orial administration of methylphenidate to rats throughout pregnancy and lactation at doses 7 times the MRHD given to adults ce Datai. sis at doses up to

The estimated background risk of major birth defects and miscarriage for the indicated population is unknown. All pregnancies have a background risk of birth defect, loss, or other adverse outcomes. In the U.S. general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnances 6.2% to 4% and 15% to 20%, respectively.

## Clinical Considerations

Fetal/Neonatal Adverse Reactions CHS stimulants, such as methylphenidate hydrochioride oral solution, can cause vasoconstriction and thereby decrease placental perfusion. No fetal and/or neonala devises reactions have been reported with the use of therapeutic doses of methylphenidate during pregnancy, however, premature delvery and law birth weight nitrains have been reported in amplicatemode dependent mothers.

<u>Data</u> Animal Data

Animal Data In embryo-feal development studies conducted in rats and rabbles, methylphenidate was administered orally at doese of up to 75 and 200 mg/kg/day, respectively, during the period of organogenesis. Mafformations (increased inclence of feal spins Mida) were observed pieve to adults on a mg/mb/basis. The one offect level for embryo-feal development in rabbit was 60 mg/kg/day (19 times the MRHD given to adults on a mg/mb/basis). There was no evidence of morphological development effects in the highest close level (12 times the MRHD on a mg/mb/basis). There was no evidence it welf compared in the offect level for an int advas 25 mg/kg/day (14 times the MRHD on a mg/mb/basis). When methylphenidate was administered to rats throughout pregnancy and licitation at doese of up to 55 mg/kg/day (14 times the MRHD on a mg/mb/basis). Unten methylphenidate was administered to rats throughout pregnancy and licitation at doese of up to 55 mg/kg/day (14 times the MRHD on a mg/mb/basis). Unton other effects on postnaial development methylphenidate to adults on a mg/mb/basis), but no other effects on postnaial development methylphenidate was administered to rats throughout pregnancy and licitation and doese of up to 55 mg/kg/day (14 to adults on a mg/mb/basis), but no other effects on postnaial development methylphenidate to adults on a mg/mb/basis). Unto other effects on postnaial development in rats was 15 mg/kg/day (-2 times the MRHD given to adults on a mg/m<sup>2</sup> basis).

#### 8.2 Lactation Risk Summary

## Clinical Considerations

Monitor breastfeeding infants for adverse reactions, such as agitation, insomnia, anorexia, and reduced weight gain.

#### 8.4 Pediatric Use

The safety and effectiveness of methylphenidate hydrochioride oral solution for the treatment of ADHD have been established in pediatric patients six years of age and older. The safety and effectiveness of methylph hydrochioride oral solution in pediatric patients under six years of age have not been established. The bong-term efficacy of methylphenidate in pediatric patients has not been established.

Long-Term Suppression of Growth Growth should be monitored during treatment with stimulants, including methylphenidate hydrochbride oral solution. Pediatric patients who are not growing or gaining weight as expected may need to have their treatment interrupted [see Warnings and Precautions (5.6)].

n a study conducted in young rats, methylphomidate was administered orably at dooses of up to 100 mg/gding/mg for 9 sevels, starting early in the postnatial period (bootshuld Day 7) and continuing through sexual maturky (postnatal Week 10). When these animals were tested as adults (postnatal Week 10). When these animals were tested as adults (postnatal Week 10). When these animals were tested as adults (postnatal Week 10). When these animals were tested as adults (postnatal Week 10). The makes and females previously treated with 50 mg/kg/day (approximately 4 times the MRHD of 60 mg/day given to children on a mg/m<sup>2</sup> basis) or greater, and a defici in the acquickshort of a specific learning task was seen in females. greater, and a deficit in the acquisition of a specific learning task was seen in females exposed to the highest doss (8 times the MRHD given to children on a mg/m<sup>2</sup> basis). The no effect level for juvenile neurobehavioral development in rats (5 mg/kg/day) is less than the MRHD given to children on a mg/m<sup>2</sup> basis. The clinical significance of the long-term behavioral effects doserved in rats is unknown.

8.5 Geriatric Use Methylphenidate hydrochloride oral solution has not been studied in the geriatric population

#### 9 DRUG ABUSE AND DEPENDENCE

9.1 Controlled Substance

Methylphenidate hydrochloride oral solution contains methylphenidate hydrochloride, a Schedule II controlled substance.

9.2 Abus 9.2 Abuile CNS struinulus, including methylphenidate hydrochloride oral solution, other methylphenidate, containing products, and ampletamines have a high potential for abuse. Abuse is the interthomal non-therapeutic use of a wing, wenn once, use a bacheve a behaviorial, cognitive, and physiological phenomena that may include a strong desire to take the drug. difficulties in controlling drug use (e.g., continuing drug use despite harmful consequences, giving a higher priority to drug use than other activities and obligations), and possible tolerance or physical dependence. Both abuse and misuse may lead to addiction, and some individuals may develop addiction even when taking methylphenidate hylprichicities or activities in controlling drug use (e.g., continuing drug use despite harmful consequences, giving a higher priority to drug use than other activities and obligations), and possible tolerance or physical dependence. Both abuse and misuse may lead to addiction, and some individuals may develop addiction even when taking methylphenidate hylprichicities other contended.

Signs and symptoms of CNS stimulant abuse include increased heart rate, respiratory rate, blood pressure, and/or sweating, dileted pupils, hyperactivity, restlessness, insomnia, decreased appetite, loss of coordination, tremores, fluched dski, vomiting, and/or abdominal pain. Anxiety, psychosis, hostilty, aggression, and suicidal or homicial ideabits have also been observed. Individuals who abuse CNS stimulants may chew, snort, inject, or use other unapproved routes of administration which may result in overdose and death [see Overdosage (10)].

To reduce the abuse of methylphenidate hydrochloride oral solution, assess the risk of abuse prior to prescribing. After prescribing, the device studies and abuse the studies and their families about abuse and on proper storage and disposal abuse while on therapy, and re-evaluate the need for methylphenidate hydrochloride oral solution use.

### 9.3 Dependence

Physical Dependence Provsati Desendence Methylphendate hydrochloride oral solution may produce physical dependence from continued therapy. Physical dependence is a li yndrome produced by abrupt essation, rapid dose reduction, or administration of an antagonst. Whithrawal symptoms after abrupt cessation following probaged high-dosage administration of CMS stimulas include dysphork model, depression; fatigue, vivid, urpleasant dreams; insomnia or hypersonnia; increased appetite; and psychomotor relarations or against.

Tolerance

Methylphenidate hydrochloride oral solution may produce tolerance from continued therapy. Tolerance is a state of adaptation in which exposure to a drug results in a reduction of the drug's desired and/or undesired effects over time).

## 10 OVERDOSAGE

Human Experience Signs and symptoms of acute methylphenidate overdosage, resulting principally from overstimulation of the central nervous system and from excessive sympathomimetic effects, may include the following nausea, vomking, diarrhea, restlessness, anxiety, agitation, tremors, hyperreflexia, muscle twitching,

convulsions (which may be followed by coma), euphoria, confusion, hallucinations, delirium, sweating, flushing, headache, hyperpyrexia, tachycardia, palpitations, cardiac arrhythmias, hypertension, hypotension, tachypnea, mydriasis, dryness of mucous membranes, and rhabdomyolysis.

Overdose Management Concut with a Certified Polion Control Center (1.400-222. 1222) for the bitest recommendations on the management of overdosage with methylphenidate. Provide supportive care, including close medical supervision and monitoring. Transment should consist of those general measures employed in the management of overdosage with any drug. Consider the possibility of multiple drug overdosage. Ensure an adequate airway, oxygenation, and ventilation. Monitor cardiac rhythm and vital signs. Use supportive and symptomatic measures.

### 11 DESCRIPTION

Methylphenidate hydrochloride oral solution contains methylphenidate hydrochloride a CMS struulant. It is available as an oral solution in 5 mg/5 mi, and 10 mg/5 mi, strengths for oral administration. Chemically, methylbhenidate hydrochloride is (*i*, (racemic), methyl α-phenyl-2-piperdineacetate hydrochloride and its structural formula is:

~#~ HCI Ì

Methylphenidate Hydrochloride C<sub>14</sub>H<sub>19</sub>NO<sub>2</sub> • HCI MW = 269.77 Clampino2 \* nci nive = 209.77 Methylphenidate hydrochloride USP is a white, odorless, fine crystalline powder. Its solutions are acid to fitmus. It is freely soluble in water and in methanol, soluble in alcohol, and slightly soluble in chloroform and in acetone.

Each mL of methylphenidate hydrochloride oral solution 5 mg/5 mL contains 1 mg of methylphenidate hydrochloride USP.

Each mL of methylphenidate hydrochloride oral solution 10 mg/5 mL contains 2 mg of methylphenidate hydrochloride USP.

Methylphenidate hydrochloride oral solution also contains the following inactive ingredients: glycerin, polyethylene glycol 1450, hydrochloric acid, grape 501417C and purified water.

### 12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Methylphenidate hydrochloride is a central nervous system (CNS) stimulant. The mode of therapeutic not known.

## 12.2 Pharmacodynamics

Active transmostgrammus Methylpheniata is a nacemic multiple of the d- and Ethree enantomers. The d-three enantomer is more pharmacologically active than the fitnee enantomer. Methylpheniata belocks the reuptake of norpeinpehrine and dopamne into the presynaptic neuron and increases the release of these monoannies into the extransmortal page.

Cardiac Electrophysiology

A formal QT study has not been conducted in subjects taking methylphenidate hydrochloride oral solution.

The effect of dexmethylphenidate, the pharmacologically active d-enartomer of methylphenidate hydrochiorde oral solution, on the QT interval was evaluated in a double-bind, placedo, and open-label active (mosfloxachi) and the solution of the solution of the solution of the solution of the solution extended-release capsule in 75 healthy volunteers. Electrocardiograms were collected up to 12 hours positiose. Frederica's method for health rate correction was employed to derive the corrected QT interval (QTcF). The maximum mean prolongation of QTcF intervals was less than 5 ms, and the upper link of the 90% confidence interval was below the twoshold of chinaci concern and there was no exident exposure response relationship.

## 12.3 Pharmacokinetic

Absorption Following a single dose administration of 20 mg methylphenidate hydrochloride oral solution and 20 mg tablet of methylphenidate hydrochloride in healthy volunteers under fastet conditions, time to peak plasma concentration ( $T_{max}$ ) of methylphenidate was at 1 to 2 hours after dosing, and:

 The mean peak plasma concentration (C<sub>max</sub>) of methylphenidate was 9.1 ng/mL and 9.8 ng/mL, respectively. 9.8 ng/mL, respectively.
 The mean area under concentration curve (AUC) of methylphenidate was 46.7 hour\*ng/mL and 50.0 hour\*ng/mL, respectively.

Effect of Food Ingestion of a high-fat meal with methylphenidate hydrochloride oral solution increased methylphenidate mean  $C_{\rm max}$  and AUC by about 13% and 25%, respectively. Time to  $C_{\rm max}(T_{\rm max})$  was delayed by approximately 1 hour

Distribution Plasma protein binding is 10% to 33%. The volume of distribution was 2.65  $\pm$  1.11 L/kg for d-methylphenidate and 1.80  $\pm$  0.91 L/kg for I-methylphenidate.

Elimination

The mean terminal half-life (t\_{b/2}) of methylphenidate was 2.7 hours following administration of 20 mg  $\,$ methylphenidate hydrochloride oral solution. The systemic clearance is  $0.40 \pm 0.12$  L/h/kg for d-methylohenidate and  $0.73 \pm 0.28$  L/h/kg for I-methylohenidate.

Metabolism

Methylphenidate is metabolized primarily by deesterification to alpha-phenyl-piperidine acetic acid (ritalinic acid), which has little or no pharmacologic activity.

Excretion After oral dosing of radiolabeled methylphenidate in humans, about 90% of the radioactivity was recovered in urine. The main urinary metabolite was ritalinic acid, accounting for approximately 80% of the dose.

#### Specific Populations Male and Eemale Patients, Racial Groups, and Age

The effect of gender, race, and age on the pharmacokinetics of methylphenidate after methylphenidate hydrochloride oral solution administration have not been studied.

Patients with Renal Impairment

Methyphenidate hydrochloride oral solution has not been studied in in patients with renal impairment. Since renal clearance is not an important route of methyphenidate clearance, renal impairment is expected to have little effect on the pharmacokinetics of methylphenidate hydrochloride oral solution.

### Patients with Hepatic Impairment

Methylphenidate hydrochiotisko oral sociation has not been studied in patients with hepatic impairment. Since methylphenidate is metabolized primarily to ritalinic acid by nonmicrosomal hydrolytic esterases that are widely distributed throughout the body, hepatic impairment is expected to have minimal effect on the pharmacokinetics of methylphenidate hydrochiotie oral solution.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis & Mutagenesis & Impairment Of Fertility

Carcinogenesis In a lifetime carcinogenicity study carried out in B6C3F1 mice, methylphenidate caused an increase in hepatocellular adenomas and, in males only, an increase in hepatoblastomas, at a daily dose of approximately 60 mg/kg/day. This dose is approximately 5 times the maximum recommended human dose (MHMD) of 60 molg dgs who baduks on a mg/m² basis. Henatoblastoma is a relatively rare rodent malignant tumor type. There was no increase in total malignant hepatic tumors. The mouse strain used is sensible to the development of hepatic tumors, and the significance of these results to humans is unknown.

ylphenidate did not cause any increase in tumors in a lifetime carcinogenicity study

Metryphynetiuate du not ouse any second a manufacture of a second s

## In a 24-week carcinogenicity study in the transgenic mouse strain p53+/-, which is sensitive to genotoxic carcinogens, there was no evidence of carcinogenicity. Male and female mice were fed diets containing the same concentration of methylphenidate as in the lifetime carcinogenicity study; the high-dose groups were exposed to 60 to 24 mg/kg/day of methylphenidate.

#### Mutagenes is

Introduction Methylphendale was not mutagenic in the *in vitro* Arms; reverse mutation assay, in the methylphendale was not mutagenic in the *in vitro* Arms; reverse mutation assay and an another chromosome beardings were increased, indicative of a wark clastoger is and an *in vitro* assay in cultured Chinese Hamster Ovary (CHO) cells. Methylphendate was negative *in vive* in males and females in the mouse bone marrow microructuus assay.

#### Impairment of Fertility

Immediates of realisty No human data on the effect of methylphenidate on fertility are available. Methylphenid in male or female mice that were fed diets containing the drug in an 18-week continuous breeding study. The study was conducted at doses up to 160 mg/kg/day, approximately 13 times the maximum recommended human dose of 60 mg/kg/gupte to adults on a ng/m<sup>2</sup> basis. Methylphenidate did not impair fertility

#### 16 HOW SUPPLIED/STORAGE AND HANDLING

How Supplied

## Methylphenidate hydrochloride oral solution is a clear colorless to pale yellow, grape flavor solution available in the following strengths:

5 mg per 5 ml Bottles of 500 mL.....NDC 67877-602-91

- 10 mg per 5 mL
- Bottles of 500 mL.....NDC 67877-603-91

Storage and Handling

Store at 20° to 25°C (68° to 77°F) [see USP Controlled Room Temperature]. Dispense in tight container with child-resistant closure

#### Disposal

Databasi Databasi Comply with local laws and regulations on drug disposal of CNS stimulants. Dispose of remaining, unused, or expired methylphendiate hydrochoride oral solution by a medicine take-back program or by an authorized collector back program or authorized collector is available, mix methylphenidate hydrochorized oral solution with an undesirable, nortoxix substance to make R less appealing to children and pets. Place the mixture in a container such as a sealed plastic bag and discard methylphenidate hydrochorized oral solution in the household trash.

## 17 PATIENT COUNSELING INFORMATION

Advise the patient to read the FDA-approved patient labeling (Medication Guide)

Controlled Substance Status/High Potential for Abuse and Dependence Lantrollers Justistance Statisting Protectional for Adults and Lependencke Advise patients that methylybenitatis hydrochroliter oraci solution is a federally controlled substance, and it can be abused and lead to dependence [see Drug Abuse and Dependence (3), 19, 2, and 3)]. Instruct patients that they should not give methylphenidate hydrochrolite oral solution to anyone else. Advise patients to store methylphenidate hydrochroliter oral solution in a safe pace, preferably locked, to prevent abuse. Advise patients to comply with laws and regulations on drug disposal. Advise patients to dispose of remaining, unused, or expired methylphenidate hydrochroliter oral solution to a methyle back, preferably locked, to prevent abuse. Advise patients to comply with laws and regulations on drug disposal. Advise patients to dispose of remaining, unused, or expired methylphenidate hydrochroliter oral solution to a methyle advision take and pendence (9), How Supplied/Storage and Handling (16)].

#### Serious Cardiovascular Risks

Advise patients that there is a potential serious cardiovascular risk including sudden death, myocardial infarction, and stroke with methylpieneidate hydrochioride oral solution. Instruct patients to contact a healthcare provider immediately if the develop symptoms such as exercisional chest pain, unexplained syncope, or other symptoms suggestive of cardiac disease [see Warnings and Pre-automs (5.2)].

#### Blood Pressure and Heart Rate Increases

Instruct patients that methylphenidate hydrochloride oral solution can elevate blood pressure and heart rate (see Warnings and Precautions (5.3)).

Psychiatric Risks EXCLUDEL.ENDE Advise patients that methylphenidate hydrochloride oral solution, at recommended dosse, can cause psychotic or main: symptoms. even in patients without prior history of psychotic symptoms or mania [see Warnings and Precautions (5.4)].

Priapism Advise patients of the possibility of painful or prolonged penile erections (priapism). Instruct the patient to seek immediate medical attention in the event of priapism [see Warnings and Precautions (5.5)].

Circulation Problems in Fingers and Toes (Peripheral Vasculopathy, Including Raynaud's Phenomenon) Crcuation Troberns n. Fngers and Tees. IPerported Vascubpathy. Including Raymad's Theometional Instruct patients about the risk of peripheral vascubpathy, including Raymad's phenomenon, and associated signs and symptoms: fingers or toes may feel numb, cool, painful, and/or may change color from pale, to blue, to red.

Instruct patients to report to their physician any new numbness, pain, skin color change, or sensitivity to temperature in fingers or toes. Instruct patients to call their physician immediately with any signs of unexplained wounds appearing on fingers or toes while taking methylphenidate hydrochloride oral solution. Further clinical evaluation (e.g., rheumatology referral) may be appropriate for certain patients (see Warnings and Precautions (5.6)).

Suppression of Growth Advise patients that methylphenidate hydrochloride oral solution may cause slowing of growth and weight loss in pediatric patients (see Warnings and Precautions (5.7)).

Pregnancy Exposure Registry Inform patients that there is a pregnancy exposure registry that monitors pregnancy outcomes in patients exposed to methylphenidate hydrochloride oral solution during pregnancy (see Use in Specific Populations (8.1)).

#### Manufactured in India by: Alkem Laboratories Limited

H.O.: ALKEM HOUSE Senapati Bapat Marg, Lower Parel Mumbai - 400 013, INDIA

Distributed by:

Ascend Laboratories, LLC Parsippany, NJ 07054 Revised: October, 2021

MEDICATION GUIDE

Methylphenidate Hydrochloride Oral Solution, CII (METH il FEN i date)

What is the most important information I should know about methylphenidate hydrochloride oral solution?

what is the most important mormation i should know about mathylphenidate hydrochoride or al solution can cause serious side effects, Bethylphenidate hydrochoride or al solution can cause serious side effects, is a solution of the phenident of the phenide hydrochoride or al solution, other emphylphenidate containing medicines, and amphetamines have a high chance for abuse and can cause physical and psychobical dependence. Your healthcare provider should check you or your child for signs of abuse and dependence before and curing transmit with methylphenidate hydrochoride oral solution. or dependent with methylphenidate hydrochoride oral solution and curing transmit with methylphenidate hydrochoride oral solution. Or the solution of the dependence and forg addiction. Heart-celled problems, including: e souden death, nichtigen with use differences between physical and psychobical dependence and drug addiction. Heart-celled problems, including: e souden death, nichtigen with use heart attack in adults e souden death, nichtigen with use heart problems or heart defects. e souden death, nichtigen with use heart problems or heart defects. With methylphenidate hydrochoride or al solution. Taylor unbakarse provider for or your child have any heart problems, heart defects, high blood pressure, or have a family heatory of these providers in solution. Callyour healthcare provider or og to the nearest hospital emergency room right away if you or your child have any spison of heart forbedms. Callyour healthcare or colles have heaty throbems, heart defects, high blood pressure, and heart and the death with methylphenidate for allyour healthcare provider or og to the nearest hospital emergency room right away if you or your child have any spison of heart tracker and your or your child have any heart problems. Nort methance provider should check your or your child's blood pressure and heart rate regularly during treatment with methylphenidate for allyour healthcare or child have any hearts are provid

awe a family history of these problems. Your healthcare provider should check colchride oral solution. Il your healthcare provider or go to the nearest hospital emergency in right away if you or your child have any signs of heart problems h as chest pain, shortness of breath, righting during treatment with methylphenidate hydrochloride oral Call your hea room right av

 Mental (psychiatric) problems, including:
 new or worse behavior and thought problems
 new or worse bipolar lines:
 new psychotic symptoms (such as hearing voices, or seeing or believing things that are not real or new main csymptoms that are not real or new main: symptoms Ful your healthcare provider about any mental problems you or your child have, or about a family history of suicide, bipolar illness, or depression. Call your healthcare provider right away if you or your child have Marchonize oral solution is processing marchine used for the treatment of Attention Defict Hyperactivity Disorder (ADHD) in people 6 years of age and older. Methylphenidate hydrochloride oral solution may help increase attention and decrase impulsiveness and hyperactivity in people with ADHD. It is not known if methylphenidate hydrochloride oral under 6 years of age and to known if methylphenidate hydrochloride oral under 6 years of age Attentylphenidate hydrochloride oral solution is a federally controlled substance (CII) because it contains methylphenidate hydrochloride oral solution in a safe pixet to protect it from thet. Never give hydrochloride oral solution in a safe pixet to protect it from thet. Never give solution may harm others and is against the isw Do not take methylphenidate hydrochloride oral solution if you or your child are: a lærgs to methylphenidate hydrochloride or any of the ingredients in methylphenidate hydrochloride oral solution. See the end of this Medication Guide for a complete list of ingredients in methylphenidate hydrochloride oral soution. taking, or have stopped taking within the past 14 days, a medicine called a monoamir efore taking methylphenidate hydrochloride oral solution tell your ealthcare provider about all your medical conditions, including if you amine oxidase inhibitor (MAOI). healthcare provider about all your medical conditions, in or your child: • have heart problems, heart defects, or high blood pressure • have heat problems, heard ordersts, or high blood pressure have neat problems including psychosis, main, blood pressure have charalized problems including psychosis, main, bloopt inless, or depression, or have can apply bloop or blooms in fingers and toos are pregnant or plan to become pregnant. It is not hown if methybhenidate hydrochoride oral solution during pregnancy. The purpose of the registry is to collect information about the headh of framalise exposed to methybhenidate hydrochoride oral solution during pregnancy. The purpose of the registry is to collect information about the headh of framalise exposed to methybhenidate hydrochoride oral solution during pregnancy. The purpose of the registry is to collect out registreng with the headh of framalise exposed to methybhenidate about registreng with the haldhow of regnancy Registry for Srychostimulants at 1-866-961-2388. are breastleding or plan to breastfeed. Methybhenidate hydrochoride oral solution. solution. Tell your healthcare provider about all the medicines that you take or your child take, including prescription and over the counter medicines, vtamins, and herbai supplements. Methylehendale cause serious size effects. Sometimes the dose of other medicines with need to be changed during treatment with methylehendale hydrochboride oral solution. Your healthcare provider with cettrylehendale hydrochboride solution. Your healthcare provider with cettrylehendale hydrochboride oral souton can be taken with other medicines. Especially tell your healthcare provider if you or your child take a medicine used to treat depression called a monoamine oxidase inhibitor (MAO). Know the medicines that you take or your child take. Keep a list of your medicines with you to show your healthcare provider and pharmacit. Do not start any new medicine during treatment with methylphenidate hydrochloride oral solution without taking to your healthcare provider first. taking to your healthCare provider trist.
How should methylphenidate hydrochloride oral solution exactly as prescribed by your healthcare provider.
Your healthcare provider may change the dose if needed.
Children 6 years of age and older:
Store breakfast and twick. 30 to 45 before a meal, as prescribed by your healthcare provider.
Aduts:
Take methylphenidate hydrochloride oral solution by mouth 2 or 3 times a day, 30 to 45 before a meal, as prescribed by your healthcare provider.
Vaduts:
Take methylphenidate hydrochloride oral solution by mouth 2 or 3 times a day, 30 to 45 before a meal, as prescribed by your healthcare provider.
Your, habiticare provider you kale your tisk dave of methylphenidate hydrochloride oral solution before 6 p.m.
Your, habiticare provider you sentimes, stop methylphenidate hydrochloride oral solution is taken taken in the day, take your last dase of methyphenidate with hydrochorbide and solution better & Jun. • Your healthcare provider may sometimes stop methyphenidate hydrochoride oral solution itsufficient for a white or levek ADD0 symptoms as hydrochoride oral solution itsufficient for a white or levek ADD0 symptoms as hydrochoride oral solution itsufficient for a white oral solution itsufficient of a white oral solution itsufficient and solution oral solution, call your poison control center at 1-800-222. 222 or go to the nearest hospital emergency your oral fait away. What are the possible side effects of methyphenidate hydrochoride oral solution can cause serious side effects, including: Painful and prolonged erecticates (graphem). Project has happened in males projecting and the projection of the solution of the solution of the solution rectify benicidate hydrochoride oral solution? • Circulation problems in fingers and toos (peripheral vasculopathy, including Raymaud's phenomenon). Signs and symptoms may include: • fregers or toos may feel numb, coch, painful • redify your healthcare provider if you or your child have numberes, pain, sinc coor change, or sensitivity to temperature in the fingers or toos, or your our child have muthyphenidate hydrochorbide eral solution.

# of unexplained wounds appearing on the second second

nyurucniorde oral solution. Methylphenidate hydrochloride oral solution treatm may be stopped if your child is not growing or gaining weight. The most common side effects of methylphenidate hydrochloride oral solution include:

<ul> <li>increased heart rate</li> </ul>	<ul> <li>irregular heart beat (palpitations)</li> </ul>
<ul> <li>headache</li> </ul>	<ul> <li>trouble sleeping</li> </ul>
<ul> <li>anxiety</li> </ul>	<ul> <li>sweating</li> </ul>
<ul> <li>weight loss</li> </ul>	<ul> <li>decreased appetite</li> </ul>
<ul> <li>dry mouth</li> </ul>	<ul> <li>nausea</li> </ul>
<ul> <li>stomach pain</li> </ul>	

# These are not all the possible side effects of methylphenidate hydrochloride oral solution.Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800- FDA-1088.

How should I store methylphenidate hydrochloride oral solution? • Store methylphenidate hydrochloride oral solution at room temperature betw

- Store methylphendidate hydrochhoride oral solution at noom temperature between 68°F to 37°F (20°C to 25°C). Store methylphenidate hydrochhoride oral solution in a safe place, like a locked cabinet.
  Protect rom high and mosture.
  Dispose of remaining, mused, or expired methylphenidate hydrochhoride oral solution in the methylphenidate hydrochhoride oral solution by the methylphenidate hydrochhoride oral solution in the table of the solution in the safe solution by a methylphenidate hydrochhoride oral solution by a methylphenidate hydrochhoride oral solution by a methylphenidate hydrochhoride oral solution with an understable, montoxic subctance such as dirtylphenidate hydrochhoride oral solution with an understable, montoxic subctance such as dirtylphenidate hydrochhoride oral solution in the household trash.
  Keen pethylphenidate hydrochhoride oral solution in the household trash.

## methylphenidate nydrochloride oral solution in the nousehol Keep methylphenidate hydrochloride oral solution and all medicines out of the reach of children.

Conversion information about the safe and effective use of methylphenidate hydrochoride and isolation. Netdicines are sometimes prescribed for purposes other than those letted in a Medication Guide. Do not use methylphenidate hydrochoride or al solution for a choriden for which it was not prescribed. Do not give methylphenidate hydrochoride oral solution to a chore people, went f they have the same symptoms. It may have mean at its against the law.

You can ask your healthcare provider or pharmacist for information about methylphe hydrochloride oral solution that was written for healthcare professionals.

What are the ingredients in methylphenidate hydrochloride oral solution/Active Ingredients: methylphenidate hydrochloride USP Inactive and purfield valer: Manufactured in India byAtken Laboratorise LimitedH 0. ALKEM HOUSE.Senapail Bapat Marg, Lower Parel,Mumbai – 400 013, INDIA Distributed byAscerd Laboratorise, LLCParel,Spany, N (07054

For more information, call 1-877-ASC-RX01 (877-272-7901). This Medication Guide has been approved by the U.S. Food and Drug Administration. Revised: October, 2021 PT 2994

#### PACKAGE LABEL.PRINCIPAL DISPLAY PANEL

NDC 67877-602-91 Methylphenidate Hydrochloride Oral Solution 5 mg/5 mL Rx Only





NDC 67877-603-91 Methylphenidate Hydrochloride Oral Solution 10 mg/5 mL Rx Only



## METHYLPHENIDATE HYDROCHLORIDE

methylphenidati	a hydrochlori	de solution				
Product Info	rmation					
Product Type		HUMAN PRESCRIPTION DRUG	Item Cod	e (Source)	ND	:67877-602
Route of Admin	istration	ORAL	DEA Sche		CI	
Note of Admin	in the test of tes		DER Sein	- une		
Active Ingred	ient/Active	Moiety				
		edient Name		Basis of St		
METHYLPHENIDATI	TE HYDROCHI	ORIDE (UNI: 4835C438HI)		METHYLPHENIDA	TE	5 mg in 5 mL
PIETTED TEMERIN	041.207223	(1-49)		monocheomor		in 5 mc
Inactive Ingr	edients					
GLYCERIN (UNII: P	DC6A3C0OX)	Ingredient Name			Str	ength
HYDROCHLORIC		17582CB)				
		(UNII: O)4Z5Z32L4)				
WATER (UNII: 059	QF0KO0R)					
Product Char	acteristics					
Color	WHITE (Clear	coloriess to pale yellow)		Score		
Shape				Size		
Flavor	GRAPE			Imprint (	Code	
Contains						
Packaging						
t Item Code		ackage Description	Mar	keting Start	Marke	ting End
				Date	D	ate
1 NDC:67877-602	500 mL in 1 B	OTTLE; Type 0: Not a Combinat	ion 04/01/	2020		
Marketing Category	Applica AND42116	ation Number or Monogra Citation		rketing Start Date	Marke	ting End ate
ANDA	ANDA2116	47	04/03	/2020		
METHYLPH methylphenidati Product Info	e hydrochlori	de solution	•			
Product Type	mation	HUMAN PRESCRIPTION DRUG		- ( <b>F</b>	ND	2:67877-603
Route of Admir		OBAI	DFA Sche		CI	
Route of Admin	istration	UNAL	DEA SCH	raule	C.	
Active Ingred	ient/Active	Moiety				
	Ingre	edient Name		Basis of St	rength	Strength
METHYLPHENIDATI	TE HYDROCHI	ORIDE (UNI: 4835C438HI)		METHYLPHENIDA	TE	10 mg in 5 mL
METHTOPHENIUKI	: - UNII:2072.23	(42.49)		HTDNOC HLDNDE		in 5 mc
Inactive Ingr	adionto					
mactive mgr	culencs	Ingredient Name			C++	ength
GLYCERIN (UNII: P	DC6A3C0OX)					
HYDROCHLORIC						
POLYETHYLENE	ILYCOL 1450	(UNII: O)4Z5Z32L4)				
WATER (UNI: 059	(FORCOR)					
Product Char	acteristics					
Color		coloriess to pale yellow)		Score		
Shape				Size		
Flavor	GRAPE			Imprint (	Code	
Contains						
Packaging						
# Item Code	P	ackage Description	Mar	keting Start	Marke	ting End
NDC-67877 602		OTTLE; Type 0: Not a Combinat	ion			ate
		orrec, type u: not a combinat	oon 04/01/	2020		

2	ackaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
L	NDC:67877-603- 91	500 mL in 1 BOTTLE; Type 0: Not a Combination Product	04/01/2020	
M	larketing	Information		
M	larketing Marketing Category	Information Application Number or Monograph Citation	Marketing Start Date	Marketing End Date

### Labeler - Ascend Laboratories, LLC (141250469) Eet

Name	Address	ID/FEI	Business Operations
Nem Laboratories Limited		677605851	MANUFACTURE(67877-602, 67877-603)

Revised: 12/2021