HIGHLIGHTS OF PRESCRIBING INFORMATION
These highlights do not include all the information needed to use levothyroxine sodium tablets safely and
effectively. See full prescribing information for levothyroxine sodium tablets.

### LEVOTHYROXINE SODIUM tablets, for oral use

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- absorption. (2.1)

  Starting doos depends on a variety of factors, including age, body weight, cardisvascular status, and concon medications. Peak therapeutic effect may not be attained for 4 to 6 weeks. (2.2)

  See full prescribing information for dooing in specific parient populations. (2.3)

  Adequacy of therapy determined with periodic monotoning of TSH and/or T4 as well as clinical status. (2.4)

# 

- WARNING AND PRECAUTIONS

  Cardiac adverse reactions in the defeaty and in patients with underlying cardiovascular disease. Takine levershyroxine

  Cardiac adverse reactions in the defeaty and in patients with underlying cardiovascular disease. Takine levershyroxine

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  Mayendene come. Do not use earliery and because of the Excess risk of cardiac adverse reactions.

  Mayendene come. Do not use earliery and because of the proceeding come. (5.2)

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ANVEST ETACTIONS

TO ANVEST ETACTIONS

CONTROL

TO PROPER STORY

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\*\*\*DRUG INTERACTIONS\*\*

See full prescribing information for drugs that affect thyoud homomore pharmacolismics and metabolism (e.g., absorption, synthesis, seeverion, catalolism, protein binding, and target tissue responses) and may after the therapeastic response to beverlyvoines confine tables. (7)

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### FULL PRESCRIBING INFORMATION

- WARNING: NOT FOR TREATMENT OF OBESITY OR FOR WEIGHT LOSS
- Thyroid hormones, including levothyroxine sodium tablets, either alone or with other therapeutic agents, should not be used for the treatment of obesity or for weight loss.
- In euthyroid patients, doses within the range of daily hormonal requirem for weight reduction.
- for weight reduction.

  Larger doses may produce serious or even life threatening manifestations of toxicity, particularly when given in association with sympathomimetic amines such as those used for their anorectic effects [see Adverse Reactions (6), Drug Interactions (7.7), and Overdosage (10)]

- HINDA-LOSS

  Hypothyvatian Upportdyvaties odium ubblets are indicated as a replacement therapy in primary (thyroidal), secondary (initiater), and entary (hypothalamic) congenital or acquired hypothyroidism.

  Pluthary Thyroropin (Thyroid-Stimulating Hormane, TSH) Suppression

  Levohyvoxies odium ubblets are indicated as an adjunct to supery and radioidize therapy in the management of thyroropin-dependent well-differentiated thyroid cancer.

- Initiation of Use Joseph and the production of t
- overtreatment with levothyroxine sodium tablets may make the premyrous and the process of the Premions (5.4)]. Levothyroxine sodium tablets are not indicated for treatment of hypothyroidism during the recovery phase of subacute thyroiditis.

### 2 DOSAGE AND ADMINISTRATION

### 2.1 General Administration Information

- Administer levothyroxine sodium tablets as a single daily dose, on an empty stomach, one-half to one hour before breakfast.
- Administer levothyroxine sodium tablets at least 4 hours before or after drugs known to interfere with levothyroxine sodium tablets absorption [see Drug Interactions (7.1)].
- Evaluate the need for dose adjustments when regularly administering within one hour of certain foods that may affect levethyroxine sodium tablets absorption [see Drug Interactions (7:9) and Clinical Pharmacology (12:3)].
- Paramacology (1.2.1). Administer levolutyoxiars sodium tablets to infants and children who carnot swallow intact tablets by crushing the tablet, suspending the freshly crushed tablet in a small amount (5 to 10 mL or 1 to 2 tesponon) of water and immediately administering the suspension by sponon of orderyer. Do not store the suspension. Do not administer in foods that decrease absorption of levolthyroxine sodium tablets, such as solybea-based infant formalia (see Drugh Internitions (7.93).

### 2.2 General Principles of Dosing

- 22 General Principles of Dosing The dose of levolvoyine sodium halves for hypothyroixtan or rininary TSH suppression depends on the most produced as the processing of the produced conditions (residently pregnancy), concontant medications, co-administrated food and the specific name of the condition being treast level Dose post and Administration (2.0), Winnings and Procession (5.0), and Drug hereatment (7.1). Excess great be individually on account for these factors and dose adjustment Dosage and Administration (2.0).

  The peak herapeatic effect of a given dose of levoldysvaries sodium tables may not be antaired for 4 to 6 weeks.

- Primary Hypothyroidism in Adults and in Adolescents in Whom Growth and Puberty are Complete
- Primary Infrastructions in Adults and in Adolescents in Whom Lirowin and Paberty are Complete Start Techniques (and in Techniques Canality and the security and t

ption, drug interactions, or a combination of these factors

complianer, mulaboroption, drug interactions, or a combination of these factors. For electry patients or patients with underlying cardiac, disease, start with a dose of 12.5 to 25 mcg per day. Increase the dose every 6 to 8 weeks, as needed until the patient its clinically enaltyroid and the searm IST it extrain so normal. The full replacement dose of levolotyrois resolium labelse may be less than 1 mcg per leg per day in eleletrly patients.

In patients with severe longstanding hypothyroidism, start with a dose of 12.5 to 25 mcg per day. Adjust the dose in 12.5 to 25 mcg increments every 2 to 4 weeks until the patient is clinically enaltyroid and the securing 151 level is a formalized.

the dose in 12.50 o.2 mig interment every a saw everes usus as possess common to the common to a normalized.

Secondary or Tertina Phypothyroidism

Surt levolthyroists column tables at the full replacement dose in otherwise healthy, non-elderly individuals, Surt with a lower dose in elderly paintees, patients with underlying cardiovascular disease reasons and the column tables of the column tables of the column tables and the column tables of the

Table 1. Levothyroxine Sodium Tablets Dosing Guidelines for Pediatric Hypothy

AGE	Daily Dose Per Kg Body Weight *		
0 to 3 months	10 to 15 mcg/kg/day		
3 to 6 months	8 to 10 mcg/kg/day		
6 to 12 months	6 to 8 mcg/kg/day		
1 to 5 years	5 to 6 mcg/kg/day		
6 to 12 years	4 to 5 mcg/kg/day		
Greater than 12 years but growth and puberty incomplete	2 to 3 mcg/kg/day		
Growth and puberty complete	1.6 mcg/kg/day		

The dose should be adjusted based on clinical response and laboratory parameters [see Dosage and Adm 2.4] and Use in Specific Populations (8.4)].

Newborns (0 to 3 months) at risk for cardiac failure: Consider a lower starting dose in newborns at risk for cardiac failure. Increase the dose every 4 to 6 weeks as needed based on clinical and laboratory

Children at risk for hyperactivity: To minimize the risk of hyperactivity in children, start at one-fourth the recommended full replacement dose, and increase on a weekly basis by one-fourth the full recommended replacement dose is reached.

Pregnancy

Pre-graining Hypothyroidium: Levothyroxius sodium tables dose requirement may increase during preparety. Measure serum TSH and free-T4 as soon as pregnancy, Measure serum TSH and free-T4 as soon as pregnancy is confirmed and, atminimum, during temperative for the service of the s

### 2.4 Monitoring TSH and/or Thyroxine (T4) Levels

Assess the adequacy of therapy by periodic assessment of laboratory tests and clinical evaluation. 
Persistent clirical and laboratory evidence of hypothyroidism despite an apparent adequate replacement does of levothyroxine sodium tablets may be evidence of inadequate absorption, poor compliance, drug interactions, or a combination of these factors.

nums. In adult patients, with primary hypothyroidism, monitor serum TSH levels after an interval of 6 to weeks after any change in dose. In patients on a stable and appropriate replacement dose, evaluate clinical and biochemical response every 6 to 12 months and whe never there is a change in the pelinical status.

with congrated by ophyroidins, seeses the adequacy of replacement therapy by reasoning determination and one free 24. Motions TSH and under 1 feer 24 in children for 10 feet 20 feet

of development, mental and physical growth, and bone minuration, at regular intervals. While the general aim of heavya is so maritize the seurm 1811 level. TSI may not normalize in some patients due to in stems hypothyroidism causing a resetting of platinary-shyroid feedback. Failure of the level process of the search 1811 and the self-search self-search patients of the search 1811 on the level patients of the search 1811 on the receiving adequate therapy. Assess compliance, does of medication administrated, and method of administration spire to increasing the does of feedback the search of the search 1811 on the search 1811 on the search 1811 of the

Monitor serum free-T4 levels and maintain in the upper half of the normal range in these patients

3 DOSAGE FORMS AND STRENGTHS

Levothyroxine sodium tablets USP are round, colored, scored and debossed with following debossing details on one side and breal-line on other side. They are supplied as follows:

Tablet Strength	Tablet Color/Shape	Debossing Details
25 mcg	Peach/Round	L15
50 mcg	White/Round	L16
75 mcg	Violet/Round	L17
88 mcg	Olive/Round	L19
100 mcg	Yellow/Round	L20
112 mcg	Rose/Round	L21
125 mcg	Tan/Round	L22
137 mcg	Turquoise/Round	L23
150 mcg	Blue/Round	L24
175 mcg	Lilac/Round	L25
200 mcg	Pink/Round	L26
200 more	Cross/Round	1.77

### 4 CONTRAINDICATIONS

traindicated in patients with uncorrected adrenal insufficiency (see

### 5 WARNINGS AND PRECAUTIONS

# 5.1 Cardiac Adverse Reactions in the Elderly and in Patients with Underlying Cardiovascular Disease

Discussion of the Control of the Con

Monitor for cardiac arrhythmias during surgical procedures in patients with coronary artery disease receiving suppressive levothyroxine sodium tablets therapy. Monitor patients receiving concomitant levothyroxine sodium tablets and sympathomimetic agents for signs and sympathoms of coronary insufficiency.

If cardiac symptoms develop or worsen, reduce the levothyroxine sodium tablets dose or withhold for one week and restart at a lower dose.

### 5.2 Myxedema Coma

3.2. Mysedema Coma Mysedema com is a life-threatering emergency characterized by poor circulation and hypometabolism and may result in unpredictable absorption of levoltyroxine sodium from the gastroinestical tract. Use of oral thyroid hormone drug products is not recommended to treat mysedema coma. Administer thyroid hormone products formulated for intravenous administration to treat mysedema coma.

### 5.3 Acute Adrenal Crisis in Patients with Concomitant Adrenal Insufficiency

3.3 Acture Aurenau Urasis in Patients with Concommant Adrenal Insufficiency. Hypoid hormone increases metabolic clearance of glucocorticoids, Initiation of thy therapy prior to initiating glucocorticoid therapy may precipitate an acture adrenal conflictionsery. Treat patiences with adrenal insufficiency with replacement gluto to initiating treatment with levolthyroxine sodium tabless [see Contraindications (43]].

### 5.4 Prevention of Hyperthyroidism or Incomplete Treatment of Hypothyroidism

3-d Prevention of Hyperthyrodom or Incomplete I realment of Hypothyroidom Levolhyroxies and unable thas an arrow the negatic index. Over-or undertreament with levolhyroxies sodium ablete may have negative reflects on growth and development, cardiovascular function, home methodism, reproductive function, cognisive function, emotional state, gastrointensiand function, and glucose and lipid metabolism. Tirate the dose of levolhyroxies sodium tables carefully and mustior response to trition to avoid these reflects (per Dougon and Administrator) (23). Monits for the presence of drug or food interactions when using levolhyroxies colium tables and adjust the dose as necessary size Puru planterization. 27) and Chinical Pharmacrology (123).

### 5.5 Worsening of Diabetic Control

Addition of levothyroxine therapy in patients with diabetes mellitus may worsen glycemic control and result in increased antidiabetic agent or insulin requirements. Carefully monitor glycemic control after starting, changing, or discontinuing levothyroxine sodium tablets [see Drug Interactions (7.2)].

# 5.6 Decreased Bone Mineral Density Associated with Thyroid Hormone Over-Replacement 2.5 Decreases none sinuario automisty Associated with Infrast Internated both extension and decreased home mirrard density my occur as a result of the objective over-replacement, particularly in post-emappined women. The increased bone resorption may be more application of the contract of the increased bone resorption may be more application of the property of the

### 6 ADVERSE REACTIONS

Adverse reactions associated with levothyroxine sodium tables therapy are primarily those of hyperthyroidism due to therapeutic overdosage (see Warnings and Precontions (5), Overdosage (10)). They include the following:

9 General fusique, increased appetite, weight loss, hea intolerance, sever, excessive sworth of the Control persons opisien: leadache, hyperactivity, nervousness, anticey, triviability, emotional lability,

- incomais

  Mancalosdedent tremos, muscle weskness, muscle spasm

  Cardiovaculor; palpitation, tachycardia, arrhydmias, increased pulse and blood pressure, heart
  failure, angins, mycordiad infarction, cardiac arrest

  Respiratory; obspase

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  Controlomerinal Controlomerinal

  Endocrine: Selectorical Controlomerinal

  Endocrine: Sel

### Reproductive: menstrual irregularities, impaired fertility

Seizures have been reported rarely with the institution of levothyroxine therapy. Adverse Reactions in Children

Pseudotumor ceretori and slipped capital femoral epiphysis have been reported in children receiving levothyroxine therapy. Overtreatment may result in craniosynostosis in infants and premature closure of the epiphyses in children with resultant compromised adult height.

### Hypers ensitivity Reactions

Hypersensitivity reactions to inactive ingredients have occurred in patients reated with thyroid hormone products. These include urticaria, pruritus, skin rash, flushing, angioedema, various gastroinestinal sympioms (abdominal pain, nances, womiting and diarrhea), fever, arthrafgia, serum sickness, and whereign. Hypersensitivity to leveltyrovine itself is not howards occur.

### 7 DRUG INTERACTIONS

7.1 Drugs Known to Affect Thyroid Hormone Pharmacokinetics

Many drugs can exert effects on thyroid hormone pharmacokinetics and metabolism (e.g., absorption, synthesis, secretion, catabolism, protein binding, and target tissue response) and may after the therapeutic response to levolvhyroits endium tables (see Tables 2 to 5 beet Tables 2).

	Table 2. Drugs That May Decrease T4 Absorption (Hypothyroidism)\
Potential impact: Concurrent use may	reduce the efficacy of levothyroxine sodium tablets by binding and delaying or preventing absorption, potentially resulting in hypothyroidism.
Drug or Drug Class	Effect
Calcium Carbonate Ferrous Sulfate	Calcium carbonate may form an insoluble chelate with levothyroxine, and ferrous sulfate likely forms a ferric-thyroxine complex. Administer levothyroxine sodium tablets at least 4 hours apart from these agents.
Orlistat	Monitor patients treated concomitantly with orlistat and levothyroxine sodium tablets for changes in thyroid function.
Bile Acid Sequestrants -Colesevelam -Cholestyramine -Colestipol Ion Exchange Resins -Kayexalate -Sevelamer	Bile acid sequestrams and ion exchange resins are known to decrease levothyroxine absorption. Administer levothyroxine sodium tablets at least 4 hours prior to these drugs or monitor TSH levels.
Other drugs: Proton Pump Inhibitors Sucral fate Artacids - Aluminum & Magnesium Hydroxides - Simethicone	Gastric acidity is an essential requirement for adequate absorption of levothyroxine. Sucraffate, attacids and proton pump inhibitors may cause hypochlorhydria, affect intragastric pH, and reduce levothyroxine absorption. Monitor patients appropriately.

	Table 3. Drugs: That May Alter T4 and Triiodothyronine (T3) Serum Transport Without Affecting Free Thyroxine (T74) Concentration (Euthyroidis m)			
Drug or Drug Class	Effect			
Clofibrate	These drugs may increase serum thyroxine-binding globulin (TBG) concentration.			
Estrogen-				
containing oral contraceptives				
Estrogens (oral)				
Heroin / Methadone				
5-Fluorouracil				
Mitotane				
Tamoxifen				
Androgens / Anabolic Steroids	These drugs may decrease serum TBG concentration.			
Asparaginase				
Glucocorticoids				
Slow-Release Nicotinic Acid				
Potential impact (below): Admi	inistration of these agents with levothyroxine sodium tablets results in an initial transient increase in FT4. Continued administration results in a decrease in serum T4 and normal FT4 and T5H concentrations.			
Salicylates (> 2 g/day)	Salicylates inhibit binding of T4 and T3 to TBG and transthyretin. An initial increase in serum FT4 is followed by return of FT4 to normal levels with sustained therapeutic serum salicylate concentrations, although total T4 levels may decrease by as much as 30%.			
Other drugs:	These drugs may cause protein-binding site displacement. Furosemide has been shown to inhibit the protein binding of T4 to TBG and albumin, causing an increase free T4 fraction in serum. Furosemide competes for T4-			
Carbamazepine	binding sites on TBG, prealbumin, and albumin, and albumin so that a single high dose can acutely lower the total T4 level. Phenytoin and carbamazeoine reduce serum protein binding of levothyroxine, and total and free T4 may be reduced by 20% to 40%, but most patients have normal serum TSH levels and are clinically euthyroid. Closely monitor thyroid hormone parameters.			
Furosemide (> 80 mg IV)				
Heparin				
Hydantoins				
Non-Steroidal Anti-				
inflammatory Drugs				
-Fenamates				
1				

### Table 4. Drugs That May Alter Hepatic Metabolism of T4 (Hypothyroidism)

Potential impact: Stirr	Potential impact: Stimulation of hepatic microsomal drug-metabolizing enzyme activity may cause increased hepatic degradation of levothyroxine, resulting in increased levothyroxine sodium tablets requirements.			
Drug or Drug Class	Effect			
Phenobarbital	Phenobarbital has been shown to reduce the response to thyroxine. Phenobarbital increases L-thyroxine metabolism by inducing uridine 5'-diphospho-			
Rifampin	glucuromosyltransferase (UGT) and leads to a lower T4 serum levels. Changes in thyroid status may occur if barbiturates are added or withdrawn from patients being treated for hypothyroidism. Rifampin has been shown to accelerate the metabolism of levothyroxine.			

### Table 5. Drugs That May Decrease Conversion of T4 to T3

Potential impact: Administration of these enzyme inhibitors decreases the peripheral conversion of T4 to T3, leading to decreased T3 levels. However, serum T4 levels are usually normal but may occasionally be slightly increased.				
Drug or Drug Class	Effect			
	in patients treated with large doses of propramolol (> 160 mg/day), T3 and T4 levels change, TSH levels remain normal, and patients are clinically euthyroid. Actions of particular beta-			
	adrenergic antagonists may be impaired when a hypothyroid patient is converted to the euthyroid state.			
	Short-term administration of large doses of glucocorticoids may decrease serum T3 concentrations by 30% with minimal change in serum T4 levels. However, long-			
	term glucocorticoid therapy may result in slightly decreased T3 and T4 levels due to decreased TBG production (See above).			
	Amiodarone inhibits peripheral conversion of levothyroxine (T4) to triiodothyronine (T3) and may cause isolated biochemical changes (increase in serum free-			
Amindarone	T4 and decreased or normal free. T3) in clinically enthyroid nations			

### 7.2 Antidiabetic Therapy

Addition of levothyroxine sodium tablets therapy in patients with diabetes mellitus may worsen glycenic control and result in increased antidiabetic agent or insulin requirements. Carefully monitor glycenic control, especially when thyroid therapy is started, changed, or discontinued [see Wornings and Prenautions (5.5)].

7.3 Oral Anticoagulants
Levodhyoxine sodium tablet increases the response to oral anticoagulant therapy. Therefore, a
decrease in the dose of anticoagulant may be warranted with correction of the hypothyroid sate or
when the levolyhyoxine sodium tables dose is increased. Closely monitor coagulation tests to permit
appropriate and interploy-dosage adjunters.

7.4 Digitalis Glycosides

Nu yesloc the therapeutic effects of digitalis glycosides. Serum digitalis glycoside selection of digitalis glycoside levels may decrease when a hypothyroid patient becomes eathyroid, necessitating an increase in the dose of digitalis glycosides.

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### 7.6 Ketamine

Concurrent use of ketamine and levothyroxine sodium tablets may produce marked hypertension and tachycardia. Closely monitor blood pressure and heart rate in these patients.

### 7.7 Sympathomimetics

Concurrent use of sympathonimetics and levothyroxine sodium tablets may increase the effects of sympathonimetics or thyroid hormone. Thyroid hormones may increase the risk of coronary insufficiency when sympathonimetic agents are administered to patients with coronary artery disease.

7.8 Tyrosine-Kinase Inhibitors
Concurrent use of tyrosine-kinase inhibitors such as imutinib may cause hypothyroidism. Closely monitor TSH levels in such patients.

### 7.9 Drug-Food Interactions

Commission of certain looks my affect levolsynasius sofiam tablest absorption thereby recessisting subjectives in doising look Department of the Commission of certain looks and post Dongs and Administration (2.1). Subjecta flour, continued areal, walnum, and descript flow in plant and descript the responsive sofiam tablest commende gastorine-stall tract. Grapefruit juice my delay the absorption of levolsynasius and arealors and reduce in biovariability.

### 7.10 Drug-Laboratory Test Interactions

Consider-funges in TBG concentrations when interpreting T4 and T3 values. Measure and evaluate unbound (free) hormone and/or determine her fee-T4 false, (TT4)) in this circ unsance. Regulary, the fee-T4 false, (TT4) in this circ unsance. Regulary, the fee-T4 false, (TT4) in this circ unsance. Regulary, the fee-T4 false (T4) in the circ unsance. Regulary (T4) in the circ unsance. TBG concentration, Nephronis, severe hypoprolicentais, severe liver disease, scrome gly, androgens, and corticosteroids decrease TBG concentration. Familial hyper- or hypo-thyrotic baining plouliurinals have been described, with the incidence of TBG deficiency approximating 1 in 9000.

### 8 USE IN SPECIFIC POPULATIONS

B USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Risk Summay

Experience with levolhyroxian use in pregnant wornes, including data from post-marketing studies, have not reported increased rates of major birth defects or miscarriages feer Dauly. There are risks to the mother and feen associated with unreased prophyroxiation in pregnancy. Size Tell levels may increase during pregnancy, Tell should be monitored and levoltyroxians codium tabless dosage adjusted levoltyroxian resolum tabless dosage adjusted levoltyroxian resolum tabless dosage adjusted levoltyroxian resolum tables should not be discontinued during pregnancy, and hypothyroidism diagnosed during pregnancy should be promptly treated.

The estimates background risk of major brith defects and six-carriage for the infectod population is unknown in the U.S. general population, the estimates background risk of major brith defects and stackersinge in clinically recognized pregnance is 2 to 4 the art 15 to 20%, respectively.

## Clinical Considerations Disease-Associated Maternal and/or Embryo/Fetal Risk

Maternal hypothyroidism during pregnancy is associated with a higher rate of complications, includin spontaneous abortion, gestational hypertension, pre-eclampsia, stillbirth, and premature delivery. Untreated maternal hypothyroidism may have an adverse effect on fetal neurocognitive development.

umerase unternal typothyrotisus may have an adverse effect on feal neurocogative development. Done Adjantense Dirigin Pryampuro and the Propatrum Period.

Pregnancy may increase levoltyroxine sodium balbes requirements. Serum TSH levels should demonitored and the levoltyroxine sodium thales dosage adjanted during pregnancy. Since postpartum TSH levels are similar to preconceptions values, the levoltyroxine sodium tables dosage about return to the per-pregnancy flow immediately after delivery (net Dosage and Administrator) CSM on immediately after delivery (net Dosage and Administrator) CSM on immediate aly after delivery (net Dosage and Administrator) CSM.

# Data Human Data

Levothyroxine is approved for use as a replacement therapy for hypothyroidism. There is a long experience of levothyroxine use in pregnant women, including data from post-marketing studies that have not reported increased rates of fetal malformations, miscarriages or other adverse maternal or fetal outcomes associated with levothyroxine use in pregnant women.

### 8.2 Lactation

B.2 Lactation

Risk Summay

Limited published studies report that levothyroxine is present in human milk. However, there is insufficient information to determine the effects of levothyroxine on the breastfed infant and no available information on the effects of levothyroxine on milk production. Adequate levothyroxine transment during lactation may normalize milk production in sphopyroid lactuating mothers. The developmental and health teerfits of breastfeeding should be considered along with the mother's clinical need for levothyroxine sodium thiest and any potential adverse effects on the breastfed infant from levothyroxine sodium thiest and any potential adverse effects on the breastfed infant from levothyroxine sodium tables or from the underlying maternal condition.

The initial dose of levothyroxine sodium tablets varies with age and body weight. Dosing adjustments are based on an assessment of the individual patient's clinical and laboratory parameters [see Dosage and Administration (2.3, 2.4)].

Administration (2.3, 2.41).
In children in whom adignosis of permasers hypothyroidism has not been established, discontinue levolthyroxine sodium ablets administration for a rial period, but only after the child is at least 3 years of age. Others rewrit 4 and TSH levels at the end of the trial period, and use laboratory test results and clinical assessment to guide diagnosis and reatment, if warranted.

Camerical Hypothyroidism five Poospos and Administration (2.3, 2.4)
Rapid restoration of normal serum T4 concentrations is essential for preventing the adverse effects of congestial hypothyroidism no interlectual development as well as on overall physical growth and manutation. Therefore, intime levoltyroxine sodium ablets therapy immediately upon diagnosis. Levelshyrosites generally canable of the It is always platters.

Closely monitor infants during the first 2 weeks of levoltyroxine sodium tablets therapy for cardiac overtical, armylotions, and application forms and scaleling.

overload, arrhydmias, and aspiration from avid suckling.

Closely monitor puriests to avoid undertreatment or overtreatment. Undertreatment may have deleterious effects on intellectual development and linear growth. Overtreatment is susceilared with cranicosyssosists in infiname, may ackerely effect the tempo of brain naturation, and may accelerate the bone age and result in premature epiphyseal closure and compromised adult stature.

Acquired Hypothymidian Intellatatic Fidencies

Closely monitor patients to avoid undertreatment and overtreatment. Undertreatment may result in poor school performment due to impaired concernation and slowed mentation and in reduced adult height.

Overtreatment may accelerate the bone age and result in premature epiphyseal closure and compromised adult stature.

Treated children may manifest a period of catch-up growth, which may be adequate in some cases to normalize adult height. In children with severe or prolonged hypothyroidism, catch-up growth may not be adequate to normalize adult height.

The Because of the interest of

### 10 OVERDOSAGE

10 OVERDOSAGE:
The signs and symptoms of overdosage are those of hyperthyroidism (see Warnings and Perconsions (5) and Adverse Roctions (6)). In addition, confusion and disoriemation may occur. Cerebral entholism, shock, com, and denth have been reported. Scienzes occurred in a 3-year of delithing gesting 3.6 mg of levultyroxine. Symptoms may not necessarily be evident or may not appear until several days after ingestion of levolutyroxine sodium. Reduce the levolutyroxine sodium. Reduce the levolutyroxine sodium like appearance of the source of the source

For current information on the management of poisoning or overdosage, contact the National Poison Control Center at 1-800-222-1222 or www.poison.org.

### 11 DESCRIPTION

Levohyvoxine sodium tablest USP contain symbetic crystalline L-3,3,5,5 tetraiodothyvoxine sodium salt [levohyvoxine (T4) sodium]. Symbetic T4 is chemically identical to that produced in the human thyroid gland. Levohyvoxine (T4) sodium has an empirical formula of C 12H 101 4N NaO 4\*H 2O, molecular weight of 798.85 (unhydrous), and structural formula as shower.

Levothyroxine sodium tables USF for oral administration are supplied in the following strengths: 25 mg, 50 mg, 75 mg, 88 mg, 100 mg, 112 mg, 125 mg, 137 mg, 150 mg, 175 mg, 200 mg, and 300 mg, Each levothyroxine sodiumables USF contains the incurve ingredients corn starch, croscarmellose sodium, magnesium stearte, maritiol and sodium bicarbonate. Table 6 provides a listing of the color additives by tablest strength:

Strength (mcg)	Color additive(s)
25	FD&C Yellow No. 6 Aluminum Lake *
50	FD&C Blue 1 Aluminum Lake
75	FD&C Red No. 40 Aluminum Lake, FD&C Blue No. 2 Aluminum Lake
88	FD&C Yellow No. 6 Aluminum Lake *, FD&C Blue No. 1 Aluminum Lake, D&C Yellow No. 10 Aluminum Lake
100	FD&C Yellow No. 6 Aluminum Lake *, D&C Yellow No. 10 Aluminum Lake
112	D&C Red No 27 Aluminum Lake
125	FD&C Yellow No. 6 Aluminum Lake *, FD&C Blue No. 1 Aluminum Lake, FD&C Red No. 40 Aluminum Lake, FD&C Blue No. 2 Aluminum Lake
137	FD&C Blue No. 1 Aluminum Lake
150	FD&C Blue No. 2 Aluminum Lake
175	FD&C Blue No. 1 Aluminum Lake, D&C Red No. 27 Aluminum Lake
200	FD&C Red No. 40 Aluminum Lake
300	FD&C Yellow No. 6 Aluminum Lake *, FD&C Blue No. 1 Aluminum Lake, D&C
	Yellow No. 10 Aluminum Lake

Levothyroxine sodium tablet USP meets USP Dissolution Test 2.

12.1 Mechanism of Action
Thyroid hormones eart their physiologic actions through control of DNA transcription and protein synthesis. Triolodorismic (3) and 1-dayoxxine (74) diffuse into the cell nucleus and hard to thyroid receptor protein attached to DNA. This hormore nucleur receptor complex activates gene transcription and synthesis of nessering NNA and expolation groutins.

The physiological actions of thyroid hormones are produced predominantly by T3, the majority of which (approximately 80%) is derived from T4 by deiodination in peripheral tissues.

Oral levothyroxine sodium is a synthetic T4 hormone that exerts the same physiologic effect as endogenous T4, thereby maintaining normal T4 levels when a deficiency is present.

### 12.3 Pharmacokinetics

### Absorption

Absorption of orally administered T4 from the gastrointestinal tract ranges from 40% to 80%. The majority of the levolstyroizer solium ables does is absorbed from the jajuma and upper ileum. The relative bioavailables of 10 evolstyroizer solium ables, compared to an equal norminal does of oral levolstyroizer solium soliution, is approximately 9.5%, 17 absorption is increased by fasting, and decreased in malkanilly of 15%. Absorption systems and up certain foods such as spyleams, Dietary fifter decreases from the control of th

### Distribution

Distribution
Circulaing thyroid hormones are greater than 99% bound to plasma proteins, including thyroxine-binding globulin (TBG), thyroxine-binding globulin (TBA), the protein binding globulin (TBA), the protein binding globulin (TBA) that the protein globulin (TBA) that the prot

Mentodosa. 
Tal is slowly eliminated (see Table 7). The major pathway of thyroid hormone metabolism is through sequential desionation. Approximately 80% of circulating T3 is derived from peripheral T4 by monodecidination. The lever is the major size of degradation for both T4 and T3, with T4 decidination monodecidination. The lever is the major size of degradation for both T4 and T3, with T4 decidination of 80% of the daily dose of T4 is decidinated to yield regula amounts of T3 and reverse T3 (FT3). T3 and Further decidination do to includy prices. Thyroid hormone are also metabolized via configuration with glucuronides and sulfates and excreted directly into the bile and gut where they undergo enterohyptic recruication.

### Excretion

Thyroid hormones are primarily eliminated by the kidneys. A portion of the conjugated hormone reaches the colon unchanged and is eliminated in the feecs. Approximately 20% of T4 is eliminated in the stool. Urinary excretion of T4 decreases with age.

Tube 7.1 minutokancuc 1 municitis vi 1 myrota 110 mones an 2 utanyrota 1 utents							
Hormone	Ratio in Thyroglobulin	Biologic Potency	t <sub>1/2</sub> (days)	Protein Binding (%)*			
Levothyroxine (T4)	10 to 20	1	6 to 7 †	99.96			
Linthrangino (T2)	1		< 2	99.5			

\* Includes TBG, TBPA, and TBA
† 3 to 4 days in hyperthyroidism, 9 to 10 days in hypothyroidism

### 13 NONCLINICAL TOXICOLOGY

### 13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Standard animal studies have not been performed to evaluate the carcinogenic potential, mutagenic potential or effects on fertility of levothyroxine.

### 16 HOW SUPPLIED/STORAGE AND HANDLING

Levothyroxine sodium tablets USP are round, colored, scored and debossed with following debossing details on one side and break-line on other side. They are supplied as follows:

Strength (mcg)	Color/Shape	Debossing Details	NDC# for bottles of 30
137	Turquoise/Round	L23	70934-590-30

Store at 25°C (77°F); excursions permitted to 15° to 30° C (59° to 86° F) [see USP Controlled Roon Temperature]. Levothyroxine sodium tablets USP should be protected from light and moisture.

17 PATIENT COUNSELING INFORMATION
Institute patient of the following information to aid in the safe and effective use of levoltyroxins softum tables:

- sodium nables:

  Dosing and Administration

  Dosing and Administration

  Instruct patients to take levels/processes sodium nabless only as directed by their healthcare provider, stames, one-half to one host before breakfast, the satisfies dose, perfectably on an empty stames, one-half to one host before breakfast.

  Inform patients has agrees such as inon and calcium supplements and tractice and observed and about professor in the satisfies of the
- of these agents.

  Instruct patients to notify their healthcare provider if they are pregnant or breastfeeding or are thinking of becoming pregnant while taking levothyroxine sodium tablets.

- Inform patients that it may take several weeks before they notice as improvement in symptoms.
   Inform patients that the levolytroxine in levolytroxine sodium ablet is intended to replace a hormoze that is normally produced by the thyroid gland. Generally, replacement therapy is to be taken for life.
   Inform patients that levolytroxine sodium tablets should not be used as a primary or adjunctive therapy in a weight control program.
   Inform patients that levolytroxine sodium tablets should not be used as a primary or adjunctive therapy in a weight control program.
   Inform patients of the program of the properties of the program of the pro

prior to any surgery.

Adverse Recention

• Instruct patients to moify their healthcare provider if they experience any of the following symptoms: rapid or irregular hearthcat, cheat pain, thormess of broath, leg cramps, headache, symptoms rapid or irregular hearthcat, cheat pain, thormess of broath, leg cramps, headache, accressive swearing, host intolerance, fewer, changes in memmual periods, lives or sidn rash, or any other musual medical event.

• Inform patients that parial had rios may occur rarely during the first few months of levothyroxine sodium tables the bency, but this is usually superpary.

sodium tablets therapy, but di Manufactured for: Lupin Pharmaceuticals, Inc. Baltimore, Maryland 21202 United States Manufactured by: Lupin Limited Pithampur (M.P.) + 454 775 INDIA Revised: June 2020

ID#:264654

# Principal Display Panel NDC: 70934-590-30



Pre	oduct Informa	tion							
Pro	duct Type		HUMAN PRESCRIP	TION DRUG	Item Coo	le (Source)	NDC:709	34-590(NDC	:68180-97
Ros	ate of Administr	ation	ORAL						
Act	tive Ingredier	t/Active Mo	iety						
		Ingre	lient Name			Basi	is of Stree	ngth	Streng
	OTHYROXINE S EQ51BO43MG4)	EXINE SO DIUM (UNE-91765S329G) (LEVOTIFITOXINE - LEVOTIFITOXINE SO ANIFITEROUS				DEUM 0.137 m			
Ina	ctive Ingredi	ents							
			Ingredient N	lame				St	rength
CRC	SCARMILLOSI	SODIUM (UNI	M28OLHBH8)						
FDA	C BLUE NO. 1 (U	IND HSR47K3TB	D)						
MAG	SNESIUM STEAR	ATE (UNI: 7005	7MSE30)						
MAZ	NNITOL (UNR 30	WL53L36A)							
sor	NUM BICARBON	ATE (UNE: 8 MD	F5V39QO)						
Pro	nduct Charact	eristics							
Col	or	turquo	ise	Score				2 pieces	
Sha	ne .	POUN	n	Size				5mm	
Flav	er			Imprint C	ede			L23	
Con	tains								
Pac	kaging		Package Description						
z	kaging Item Code		Package Descrip	ption		Marketii Da			rting End Date
z	Item Code DC:70934-590-	30 in 1 BOTTLE Product	Package Descrip		atio n				
1 N	Item Code DC:70934-590-	Product			atio n	Da			
ı N	Item Code DC:70934-590-	product		iot a Combin		Da	te		Date

Labeler - Denton Pharma, Inc. dba Northwind Pharmaceuticals (080355546) Registrant - Denton Pharma, Inc. dba Northwind Pharmaceuticals (080355546)

Revised: 2/2020

Denton Pharma, Inc. dba Northwind Pharmaceuticals