

FOR ORAL ADMINISTRATION

Inactive Ingredients

Microcrystalline cellulose, croscarmellose sodium and magnesium stearate. The following are the coloring additives per tablet strength:

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Strength (mcg)	Color additive(s)	
25	FD&C Yellow No. 6 Aluminum Lake	
50	None	
75	FD&C Blue No. 1 Aluminum Lake, D&C Red No. 30 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	FD&C Yellow No. 6 Aluminum Lake, FD&C Red No. 40 Aluminum Lake, D&C Red No. 30 Aluminum Lake	
125	FD&C Red No. 40 Aluminum Lake, D&C Yellow No. 10 Aluminum Lake	
137	FD&C Blue No. 1 Aluminum Lake	
150	FD&C Blue No. 1 Aluminum Lake, D&C Red No. 30 Aluminum Lake	
175	FD&C Blue No. 1 Aluminum Lake, D&C Yellow No. 10 Aluminum Lake	
200	D&C Red No. 30 Aluminum Lake, D&C Yellow No. 10 Aluminum Lake	
300	FD&C Yellow No. 6 Aluminum Lake, FD&C Blue No. 1 Aluminum Lake, D&C Yellow No. 10 Aluminum Lake	

CLINICAL PHARMACOLOGY

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Thyroid hormone synthesis and secretion is regulated by the hypothalamic-pituitarythyroid axis. Thyrotropin-releasing hormone (TRH) released from the hypothalamus
stimulates secretion of thyroid-stimulating hormone, TSH, from the anterior pituitary. TSH,
in turn, is the physiologic stimulus for the synthesis and secretion of thyroid hormones, L
thyroxine (T_a) and L-triidothyronine (T_a), by the thyroid pland. Circulating serum T, and T_a,
levels exert a feedback effect on both TRH and TSH secretion. When serum T, and T_a, levels
increase, TRH and TSH secretion decrease. When thyroid hormone levels decrease, TRH and
TSH secretion increase. TSH secretion increase.

The mechanisms by which thyroid hormones exert their physiologic actions are not completely understood, but it is thought that their principal effects are exerted through control of DNA transcription and protein synthesis. T₃ and T₂ diffuse into the cell nucleus and bind to thyroid receptor proteins attached to DNA. This hormone nuclear receptor complex activates gene transcription and synthesis of messenger RNA and cytoplasmic proteins.

Thyroid hormones regulate multiple metabolic processes and play an essential role in normal growth and development, and normal maturation of the central nervous system and bone. The metabolic actions ofthyroid hormones include augmentation of cellular respiration and thermogenesis, as well as metabolism of proteins, carbohydrates and lipids. The protein

and ulemographics, we were an inequalism of proteins, canoninguists and injunes. The pinch anabolic effects of thyroid hormones are essential to normal growth and development. The physiologic actions of thyroid hormones are produced predominately by T_3 , the majority of which (approximately 80%) is derived from T_4 by deiodination in peripheral

Levothyroxine, at doses individualized according to patient response, is effective as replacement or supplemental therapy in hypothyroidism of any etiology, except transient hypothyroidism during the recovery phase of subacute thyroidits. Levothyroxine is also effective in the suppression of pituitary TSH secretion in the

treatment or prevention of various types of euthyroid goiters, including thyroid nodules, Hashimoto's thyroiditis, multinodular goiter and, as adjunctive therapy in the management of thyrotropin-dependent well-differentiated thyroid cancer (see INDICATIONS AND USAGE, PRECAUTIONS, DOSAGE AND ADMINISTRATION).

Absorption – Absorption of orally administered T_4 from the gastrointestinal (GI) tract ranges from 40% to 80%. The majority of the levothyroxine dose is absorbed from the jejunum and upper ileum. The relative bioavailability of LEVOXYL® tablets, compared to an equal nominal dose of oral levothyroxine sodium solution, is approximately 98%. T_a absorption is increased by fasting, and decreased in malabsorption syndromes and by certain foods such as soybean infant formula. Dietary fiber decreases bioavailability of T4. Absorption may also decrease with age. In addition, many drugs and foods affect T_4 absorption (see PRECAUTIONS, Drug Interactions and Drug-Food Interactions).

PRECAUTIONS, Drug Interactions and Drug-Food Interactions)

Distribution — Circulating thyroid hormones are greater than 99% bound to plasma proteins, linciduing thyroxine-binding globulin (TBG), thyroxine-binding grealburnin (TBPA), and alburnin (TBA), whose capacities and affinities vary for each hormone. The higher sterm levels, slower metabolic clearance, and longer half-life of T₁ compared to T₂. Protein-bound thyroid hormones exist in reverse equilibrium with small amounts of free hormone. Only unbound hormone is metabolically active. Many drugs and physiologic conditions affect the binding of thyroid hormones to serum proteins (see PRECAUTIONS, Drug Interactions and Drug-Laboratory Test Interactions). Thyroid hormones do not readily cross the placental barrier (see PRECAUTIONS, Prepanarov).

Test Interactions). Thyroid hormones do not readily cross the placental barrier (see PRECAUTIONs, Pregnancy).

Metabolism — T₄ is slowly eliminated (see TABLE 1). The major pathway of thyroid hormone metabolism is through sequential deiodination. Approximately eligihy-percent of circulating T₃ is derived from peripheral T₄ by monodeiodination. The liver is the major site of degradation for both T₄ and T₃, with T₄ deiodination also occurring at a number of additional sites, including the kidney and other tissues. Approximately 80% of the daily dose of T₄ is deiodinated to yield equal amounts of T₃ and reverse T₃ (rT₃). T₃ and rT₃ are further deiodinated to diodiothyronic. Thyroid hormones are also metabolized via conjugation will plucuronides and sulfates and excreted directly into the bile and gut where they undergo enteroheacit recriculation. rohepatic recirculation.

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

Table 1: Pharmacokinetic Parameters of Thyroid Hormones in Euthyroid Patients				
Hormone	Ratio in Thyroglobulin	Biologic Potency	11/2(days)	Protein Binding (%) ²
Levothyroxine (T_4) Liothyronine (T_3)	10–20 1	1 4	6–7¹ 2	99.96 99.5
¹ 3 to 4 days in hyperthyroidism, 9 to 10 days in hypothyroidism; ² Includes TBG, TBPA, and TBA				

INDICATIONS AND USAGE

Levothyroxine sodium is used for the following indications:

Levothyroxine sodium is used for the following indications: Hypathyroidism — As replacement or supplemental therapy in congenital or acquired hypothyroidism of any etiology, except transient hypothyroidism during the recovery phase of subacute thyroidils. Specific indications include: primary (thyroidil), secondary (pilutary), and tertiary (hypothalamic) hypothyroidism and subclinical hypothyroidism. Primary hypothyroidism may result from functional deficiency, primary atrophy, partial or total congenital absence of the thyroid gland, or from the effects of surgery, radiation, or drugs, with or without the presence of goiter. Pilutlary 18th Suppression — In the treatment or prevention of various types of euthyroid goiters (see WARNINGS and PRECAUTIONS), including thyroid nodules (see WARNINGS and PRECAUTIONS), subacute or chronic hymphocytic thyroidist (Hashimdots kryvioditis), multinodular goiter (see WARNINGS and PRECAUTIONS) and, as an adjunct to surgery and radiolodine therapy in the management of thyrotropin-dependent well-differentiated thyroid cancer.

CONTRAINDICATIONS

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Levothyroxine is contraindicated in patients with untreated subclinical (suppressed serum TSH level with normal T, and T, levels) or overt thyrotoxicosis of any etiology and in patients with acute myocardial infarction. Levothyroxine is contraindicated in patients with uncorrected adrenal insufficiency since thyroid hormones may precipitate an acute adrenal crisis by increasing the metabolic clearance of glucocorticoids (see PRECAUTIONS). LEVOXIV-® is contraindicated in patients with hyperensitivity to any of the inactive ingredients in LEVOXYC® tablets (see DESCRIPTION, Inactive Ingredients).

WARNING: Thyroid hormones, including LEVOXYL®, either alone or with other rapeutic agents, should not be used for the treatment of obesity or for weight loss inerupeuru ageins, souroul nut eu seus ut inter treatment in ubersity in weight it each in euthyroid patients, dosses within the range of daily hormonal requirements are ineffective for weight reduction. Larger dosses may produce serious or even life threatening manifestations of toxicily, particularly when given in association with sympathonimetic amines such as those used for their anorectic effects.

Levothyroxine sodium should not be used in the treatment of male or female infertility

Levothyroxine sodium should not be used in the treatment of male or female infertility unless this condition is associated with hypothyroxidism. In patients with nontoxic diffuse goiter or nodular thyroid disease, particularly the elderly or those with underlying cardiovascular disease, levothyroxine sodium therapy is contraindicated if the serum TSH level is already suppressed due to the risk of precipitating over thyrotoxicosis (see CONTRAINDICATIONS). If the serum TSH level is not suppressed LEVOXY.1 should be used with caution in conjunction with careful monitoring of thyroid function for evidence of hyperthyroidism and clinical monitoring for potential associated adverse cardiovascular signs and symptoms of hyperthyroidism.

PRECAUTIONS

Levothyroxine has a narrow theraneutic index. Renardless of the indication for use, careful Levothyroxine has a narrow therapeutic index. Regardless of the indication for use, careful dosage titration is necessary to avoid the consequences of over- or under-treatment. These consequences include, among others, effects on growth and development, cardiovascular function, bone metabolism, reproductive function, cognitive function, emotional state, gastrointestinal function, and on glucose and lipid metabolism. Many drugs interact with levothyroxine sodium necessitating adjustments in dosing to maintain therapeutic response (see **Drug Interactions**).

Effects on bone mineral density – In women, long-term levothyroxine sodium therapy has

been associated with decreased bone mineral density, especially in postmenopausal women on greater than replacement doses or in women who are receiving suppressive doses of leveltyproxine soldium. Therefore, it is recommended that patients receiving levoltymas sodium be given the minimum dose necessary to achieve the desired clinical and biochemical

response.

Patients with underlying cardiovascular disease – Exercise caution when administering levothyroxine to patients with cardiovascular disorders and to the elderly in whom there is an increased risk of occult cardiac disease. In these patients, levothyroxine therapy should be initiated at lower doses than those recommended in younger individuals or in patients without cardiac disease (see WARNINGS, PREAUTIONS, Gerainte Use; and DOSAGE AND ADMINISTRATION). If cardiac symptoms develop or worsen, the levothtryoxine doses should be reduced or withheld for one week and then cautiously restarted at a lower dose. Overtreatment with levothyroxine sodium may have adverse cardiovascular effects such as an increase in heart rate, cardiac wall thickness, and cardiac contractility and may precipitate angina or arrhythmias. Patients with coronary ardery disease who are receiving levothyroxine therapy should be monitored closely during surgical procedures, since the possibility of precipitating cardiac arrhythmisas may be greater in those treated with levothyroxine. precipitating cardiac arrhythmias may be greater in those treated with levothyroxine. Concomitant administration of levothyroxine and sympathomimetic agents to patients with coronary artery disease may precipitate coronary insufficiency.

Patients with nontoxic diffuse goiler or nodular thyroid disease – Exercise caution when administering levothyroxine to patients with nontoxic diffuse goiler or nodular thyroid disease in order to prevent precipitation of thyrotoxicosis (see WARNINGS). If the serum TSH is already suppressed, levothyroxine sodium should not be administered (see Contraindications).

Associated endocrine disorders

Hypothalamic/pituitary hormone deficiencies - In patients with secondary or tertiary hypothyroidism, additional hypothalamic/pitutary hormone deficiencies should be considered, and, if diagnosed, treated (see PRECAUTIONS, Autoimmune polyglandular syndrome) for adrenal insufficiency.

syndrome) for adreaal insufficiency.

Autoimmune polyalandular syndrome – Occasionally, chronic autoimmune thyroiditis may occur in association with other autoimmune disorders such as adrenal insufficiency, pernicious anemia, and insulin-dependent diabetes mellitus. Patients with concomitant adrenal insufficiency should be treated with replacement glucocorticosis prior to initiation of treatment with levothyroxine sodium. Failure to do so may precipitate an acute adrenal crisis when thyroid hormone therapy is initiated, due to increased metabolic clearance of glucocorticoids by thyroid hormone. Patients with diabetes mellitus may require upward adjustments of their antidiabetic therapeutic regimens when treated with levothyroxine (see PRECAUTIONS, Drug Interactions).

Other associated medical conditions

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Infants with congenital hypothyroidism appear to be at increased risk for other congenital
anomalies, with cardiovascular anomalies (pulmonary stenosis, atrial septal defect, and
ventricular septal defect,) being the most common association.

Information for Patients

Patients should be informed of the following information to aid in the safe and effective use of LEVOXYL®:

- 1. Notify your physician if you are allergic to any foods or medicines, are pregnant or intend to become pregnant, are breast-feeding or are taking any other medications, including prescription and over-the-counter preparations.
- 2. Notify your physician of any other medical conditions you may have, particularly heart Notify your physician of any other medical conditions you may have, particularly heart disease, diabetes, clotting disorders, and adrenal or pituliary gland problems. Your dose of medications used to control these other conditions may need to be adjusted while you are taking LEVDXYL². If you have diabetes, monitor your blood and/or uniary glucose levels as directed by your physician and immediately report any changes to your physician. If you are taking anticoagulants (blood thinners), your clotting status should be cherked trequently.
- physician. I you are aking anicoaguianis (blood thillners), your clotting status shot be checked frequently.

 3. Use LEVOXTL® only as prescribed by your physician. Do not discontinue or change t amount you take or how often you take it, unless directed to do so by your physician.
- 4. The levothyroxine in LEVOXYI.[®] is intended to replace a hormone that is normally produced by your thyroid pland. Generally, replacement therapy is to be taken for life, except in cases of transient hypothyroidism, which is usually associated with an inflammation of the thyroid gland (thyroiditts).
- Take LEVOXYL® in the morning on an empty stomach, at least one-half hour before eating
- 6. LEVOXYL® may rapidly swell and disintegrate resulting in choking, gagging, the tablet getting stuck in your throat or difficulty swallowing. It is very important that you take the tablet with a full glass of water. Most of these problems disappeared when Levoxyl® tablets were taken with water.
- It may take several weeks before you notice an improvement in your symptoms
- 8. Notify your physician if you experience any of the following symptoms: rapid or irregular heartbeat, chest pain, shortness of breath, leg cramps, headache, nervousness, irritability, sleeplessness, tremors, change in appetite, weight gain or loss, vomiting, diarrhea, excessive sweating, heat intolerance, fever, changes in menstrual periods, hives or skin rash, or any other unusual medical event.
- Notify your physician if you become pregnant while taking LEVOXYL®. It is likely that your dose of LEVOXYL® will need to be increased while you are pregnant.
 Notify your physician or dentist that you are taking LEVOXYL® prior to any surgery.
- 11. Partial hair loss may occur rarely during the first few months of LEVOXYL® therapy, but this is usually temporary.

 12. LEVOXYL® should not be used as a primary or adjunctive therapy in a weight control
- program
- 13. Keen | EVOXYI out of the reach of children. Store | EVOXYI away from heat, moisture

Laboratory Tests

The diagnosis of hypothyroidism is confirmed by measuring TSH levels using a sensitive assay (second generation assay sensitivity 0.1 mIU/L or third generation assay sensitivity ≤ 0.01 mIU/L) and measurement of free-T₄.

The adequacy of therapy is determined by periodic assessment of appropriate laboratory

The adequacy of therapy is determined by periodic assessment of appropriate laboratory tests and clinical evaluation. The choice of laboratory tests depends on various factors including the etiology of the underlying thyroid disease, the presence of concomitant medical conditions, including pregnancy, and the use of concomitant medications (see PRECAUTIONS, Drug Interactions and Drug-Laboratory Test Interactions). Persistent clinical and laboratory evidence of hypothyroidism despite an apparent adequate replacement does of LEVDXVIP. any be evidence of inadequate absorption, poor compliance, drug interactions, or decreased T₄ potency of the drug product.

Adults
In adult patients with primary (thyroidal) hypothyroidism, serum TSH levels (using a sensitive assay) alone may be used to monitor therapy. The frequency of TSH monitoring during levothyroxine does titration depends on the clinical situation but it is generally recommended at 6-8 week intervals until normalization. For patients who have recently initiated levothyroxine therapy and whose serum TSH has normalized or in patients who have had their dosage or brand of levothyroxine changed, the serum TSH concentration should be measured after 8-12 weeks. When the optimum replacement dose has been attained, clinical (physical examination) and biochemical monitoring may be performed every 6-12 months, depending on the clinical situation, and whenever there is a change in the patients status. It is recommended that a physical examination and a serum TSH measurement be performed at least annually in patients receiving LEVOXYL® (see WARNINGS, PRECAUTIONS, and DOSAGE AND ADMINISTRATION).

In patients with congenital hypothyroidism, the adequacy of replacement therapy should be in pacients with congenital hypothyloidsin, the adequacy of replacement metapy should be assessed by measuring both serum TSH (using a sensitive assay) and total- or free-T₄, brould be maintained at all times in the upper half of the normal range. While the aim of therapy is to also normalize the serum TSH level, this is not always possible in a small percentage of patients, particularly in the first few months of therapy. TSH may not normalize due to a resetting of the pituitary-thyroid feedback threshold as a result of in útero hypothyroidism. Failure of the serum T₄ to increase into the upper half of the normal range within 2 weeks of initiation of LEVOXYL® therapy and/or of the serum TSH to decrease below 20 mU/L within 4 weeks should alert the physician to the possibility that the child is not receiving adequate therapy. Careful inquiry should then be made regarding compliance, dose of medication administered, and method of administration prior to raising the dose of LEVOXYL®

The recommended frequency of monitoring of TSH and total or free T₄ in children is as follows; at 2 and 4 weeks after the initiation of treatment; every 1–2 months during the first year of life; every 2–3 months between 1 and 3 years of age; and every 3 to 12 months year of lite; every 2-3 months between 1 and 3 years of age; and every 3 to 12 months thereafter until growth is completed. More frequent intervals of monitoring may be necessary if poor compliance is suspected or abnormal values are obtained. It is recommended that TSM and T₄ levels, and a physical examination, inindicated, be performed 2 weeks after any change in LEVDXT[®] dosage. Routine clinical examination, including assessment of mental and physical growth and development, and bone maturation, schould be performed at regular intervals (see PRECAUTIONS, Pediatric Use and DOSAGE AND ADMINISTRATION).

Secondary (pituitary) and tertiary (hypothalamic) hypothyroidism

Adequacy of therapy should be assessed by measuring serum free-T₄ levels ,which should be maintained in the upper half of the normal range in these patients.

Drug Interactions

Many drugs affect thyroid hormone pharmacokinetics and metabolism (e.g., absorption synthesis, secretion, catabolism, protein binding, and target tissue response) and may after the therapeutic response to LEVOXYL®. In addition, thyroid hormones and thyroid status have varied effects on the pharmacokinetics and action of other drugs. A listing of drug-thyroidal axis interactions is contained in Table 2

The list of drug-thyroidal axis interactions in Table 2 may not be comprehensive due to the introduction of new drugs that interact with the thyroidal axis or the discovery of previously unknown interactions. The prescriber should be aware of this fact and should consult appropriate reference sources. (e.g., package inserts of newly approved drugs, medical literature) for additional information if a drug-drug interaction with levothyroxine is

Table 2: D	Table 2: Drug – Thyroidal Axis Interactions			
Drug or Drug Class	Effect			
	TSH secretion -the reduction is not sustained; , hypothyroidism does not occur			
Dopamine / Dopamine Agonists Glucocorticoids Octreotide	Use of these agents may result in a transient reduction in TSH secretion when administered at the following doses: Dopamine (> 1 mcg/kg/min); Glucocorticoids (hydrocortisone > 100 mg/day or equivalent); Octreotide (> 100 mcg/day).			
Drugs tha	t alter thyroid hormone secretion			

Drugs that may decrease thyroid hormone secretion, which may result in hypothyroidism

Methimazole Propylthiouracil (PTU)

Long-term lithium therapy can result in goit up to 50% of patients, and either subclinical overt hypothyroidism, each in up to 20% of overt hypothyroidism, each in up to 20% patients. The fetus, neonate, elderly and euthyroid patients with underlying thyroidsease (e.g., Hashimoto's thyroiditis or Grave's disease previously treated with additional control of the c urave s oilsease previousy treated with radiolodine or surgery) are among those individuals who are particularly susceptible to iodine-induced hypothyroidism. Oral cholecystographic agents and amiodarone are slowly excreted, producing more prolonged hypothyroidism than parenterally rypunyoudism man parenterally administered iodinated contrast agents. Long-term aminoglutethimide therapy may minimally decrease T₄ and T₅ levels and increase TSH, although all values remain within normal limits in most existent.

Drugs that may increase thyroid hormone secretion, which may result in hyperthyroidism

uniodarone odide (including iodine contain

Indidice and drugs that contain pharmacologic amounts of iodide may cause hyperthyroidism in euthyroid patients with Grave's disease previously treated with antithyroid drugs or in euthyroid patients with thyroid automory (e.g., multimodular goiter or hyperfunctioning thyroid adenoma). Hyperfunctioning thyroid adenoma). Hyperfunctioning thyroid adenoma, is they may discontinuation. Amount of the control of t

Drugs that may decrease T₄ absorption, which may result in hypothyroidism

- Aluminum & Magn
- Simethicone
Bile Acid Sequestrants
- Cholestyramine
- Colestipol
Calcium Carbonate
Cation Exchange Resin:
- Kayexalate
Ferrous Sulfate
Sucralfate

Concurrent use may reduce the efficacy of levothyroxine by binding and delaying or preventing absorption, potentially resulting in hypothyroidism. Calcium carbonate may form ar insoluble chelate with levothyroxine, and ferrous insoluble chelate with levothyroxine, and lerrus sulfate likely forms a ferric-thyroxine complex. Administer levothyroxine at least 4 hours apart from these agents.

Drugs that may alter T_4 and T_3 serum transport – but FT_4 concentration remains normal; and, therefore, the patient remains euthyroid

Drugs that may increase serum TBG concentration	Drugs that may decrease serum TBG concentration	
Clofibrate Estrogen-containing oral contraceptives Estrogens (oral) Heroin / Methadone 5-Fluorouracil Mitotane Tamoxifen	Androgens / Anabolic Steroids Asparaginase Gluccoorticoids Slow-Helease Nicotinic Acid	

Drugs that may cause protein-binding site displacement

Furosemide (> 80 mg IV) Heparin Hydantoins Non Steroidal Anti-Inflammatory Drugs - Fenamates """ "Parone - Phenylbutazone Salicylates (> 2 g/day)

Administration of these agents with levothyro results in an initial transient increase in FT_4 . Continued administration results in a decreas serum T_4 and normal FT_4 and TSH concentral and, therefore, patients are clinically euthyroic Salicylates inhibit binding of T₄ and T₃ to TBG and transthyretin. An initial increase in serum FT, 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%.

	er T ₄ and T ₃ metabolism bolism, which may result in hypothyroidism
Carbamazepine Hydantolins Phenobarbital Rifampin	Stimulation of hepatic microsomal drug- metabolizing enzyme activity may cause increased hepatic degradation of levothyroxine, resulting in increased levothyroxine requirements. Phenytoin and carbamazepine reduce serum protein binding of levothyroxine, and total- and free-T ₄ may be reduced by 20% to 40%, but most patients have normal serum TSH levels and are clinically euthyroid.
	ase T ₄ 5'-deiodinase activity
Amiodarone Beta-adrenergic antagonists - (e.g., Propranolol > 160 mg/day) Gluccorticolids - (e.g., Dexamethasone > 4 mg/day) Propylthiouracii (PTU)	Administration of these enzyme inhibitors decreases the peripheral conversion of T ₈ to T ₃ , leading to decreased T ₃ levels. However, serum T ₄ levels are usually normal but may occasionally be slightly increased. In patients treated with large doses of propranolol (> 160 mg/day), T ₃ and T ₄ levels change slightly, TSH levels remain normal, and patients are clinically euthyroid. It should be noted that actions of particular beta-adventorigic antigonists may be impaired when the hypothyroid patient is converted to the euthyroid state. Short-term administration of large doses of glucocorticoids may decrease serum T ₂ concentrations by 30% with minimal change in serum T ₄ levels. However, Ino-perm glucocorticoid therapy may result in slightly decreased T ₃ and T ₄ levels due to decreased TBG production (see above).
Anticoagulants (oral)	Thyroid hormones appear to increase the
Coumarin Derivatives Indandione Derivatives	catabolism of vitamin K-dependent clotting factors, thereby increasing the anticoagulant activity of oral anticoagulants. Concomitant use of these agents impairs the compensatory increases in clotting factor synthesis. Prothrombin time should be carefully monitored in patients taking levothyroxine and oral anticoagulants and the dose of anticoagulant therapy adjusted accordingly.
Antidepressants - Tricycliss (e.g., Amtiriptyline) - Tetracycliss (e.g., Maprotiline) - Selective Serotonin Reuptake Inhibitor (SSRIs; e.g., Sertralline)	Concurrent use of tri/letracyclic antidepressants and levothyroxicm may increase the therapeutic and toxic effects of both drugs, possibly due to increased receptor sensitivity to caterholamines. Toxic effects may include increased risk of cardiac arrhythmias and CNS stimulation; onset of action of tricyclics may be accelerated. Administration of sertraline in patients stabilized on levothyroxime may result in increased levothyroxine requirements.
Antidiabetic Agents - Biguanides - Meglitnides - Sulfonylureas - Thiazolidediones - Insulin	Addition of levothyroxine to antidiabetic or insulin therapy may result in increased antidiabetic agent or insulin requirements. Careful monitoring of diabetic control is recommended, especially when thyroid therapy is started, changed, or discontinued.
Cardiac Glycosides	Serum digitalis glycoside levels may be reduced in hyperthyroidism or when the hypothyroid patient is converted to the euthyroid state. Therapeutic effect of digitalis glycosides may be reduced.
Cytokines - Interferon-α - Interfeukin-2	Thereapy with interferon-x has been associated with the development of antithyroid microsomal antibodies in 20% of patients and some have transient hypothyroidism, hyperthyroidism, or both. Patients who have antithyroid antibodies before treatment are at higher risk for thyroid dysfunction during teatment. Interleukin-2 has been associated with transient painless thyroiditis in 20% of patients. Interferon-p and have not been reported to cause thyroid dysfunction.
Growth Hormones - Somatrem - Somatropin	Excessive use of thyroid hormones with growth hormones may accelerate epiphyseal closure. However, untreated hypothyroidism may interfere with growth response to growth hormone.
Ketamine	Concurrent use may produce marked hypertension and tachycardia; cautious administration to patients receiving thyroid hormone therapy is recommended.
Methylxanthine Bronchodilators - (e.g., Theophylline)	Decreased theophylline clearance may occur in hypothyroid patients; clearance returns to normal when the euthyroid state is achieved.
Radiographic Agents	Thyroid hormones may reduce the uptake of ¹²³ I, ¹³¹ I, and ^{99m} Tc.
Sympathomimetics	Concurrent use may increase the effects of sympathomimetics or thyroid hormone. Thyroid hormones may increase the risk of coronary insufficiency when sympathomimetic agents are administered to patients with coronary artery disease.
Chloral Hydrate Diazepam Ethionamide Lovastalin Metoclopramide 6-Mercaptopurine Nitroprusside Para-aminosalicytate sodium Perphenazine Resorionol (excessive topical use) Thiazide Diuretics	These agents have been associated with thyroid hormone and / or TSH level alterations by various mechanisms.

Cal anticoagulants – Levothyroxine increases the response to oral anticoagulant therapy. Therefore, a decrease in the dose of anticoagulant may be warranted with correction of the hypothyroid state or when the LEVOXYL* dose is increased. Prothrombin time should be closely monitored to permit appropriate and timely dosage adulstments (see Table 2).

<u>Distalls obcosides</u> — The therapeutic effects of digitals glycosides may be reduced by levothyroxine. Serum digitals glycoside levels may be decreased when a hypothyroid patient becomes euthyroid, necessitating an increase in the dose of digitals glycosides (see Table 2).

<u>Therap Food Internations</u> — Consumption of certain foods may affect elevothyroxine absorption thereby necessitating an increase in a floreing. Sylvente floor (mich formalls), control on the fall tradition, recessitating an increase in a floreing. Sylvente floor (mich formalls), control on the fall tradition.

<u>Therap Food Internations</u> — Changes in TiES concentration must be considered when interpreting T₄ and T₅ values, which necessitates measurement and evaluation of unbound (free) hornardor determination of the free T₂ incide (Ff.1). Prepanally, increase TBG concentrations. Decreases in TBG concentrations are observed in nephrosis, severe hypoproteimenia, severe liver disease, acromegaly, and after androgen or corticosteroid therapy (see also Table 2). Familia hyper- or hypothyroxine binding plotulinenias have been described, with the incidence of TBG declinency approximating 1 in 9000.

<u>Carrinopenests</u>, Mutagenessis, and Impairment of Fertility — Animal studies have not been

In 9000.

Carcinogenesis, Mutagenesis, and Impairment of Fertility — Animal studies have not been performed to evaluate the carcinogenic potential, mutagenic potential or effects on fertility of leothyroxice. The symbietic r. ja IEVXVII "si is directal to that produced naturally by the human formation of the produced study by the human formation of the produced study in the produced study. The study claim of the produced study is the produced study in the produced study for appropriate clinical indications should be littrated to the lowest effective replacement does.

Prepagacy — Expegory A — Studies in women taking betwhyroxine sodium during pregnancy have not shown an increased risk of congenital abnormalities. Therefore, the possibility of fetal harm appears remote LEVXVII" should not be discontinued during pregnancy and hypothyroidism diagnosed during pregnancy should be promptly treated.

Hypothyroidism during pregnancy is associated with a higher rate of complications, including spontaneous abortion, pre-ctampsia, stillibrith and premature delivery. Maternal hypothyroidism may have an adverse effect on fetal and childhood growth and development. During pregnancy, serum T₁ levels may decrease and serum TSH levels increase to values outside the normal range. Since elevations is serum TSH may occur as early as 4 weeks gestation, pregnant women taking LEVOXY! Should have their TSH measured during each trimester. An elevated serum TSH level should be corrected by an increase in the dose of LEVOXY! Since postpartum TSH levels as risinitar to preconception values, the LEVOXY! dosage should return to the pre-pregnancy dose immediately after delivery. A serum TSH level should be obtained 6-8 weeks postpartum.

Thyroid hormones do not readily cross the placental barrier, however, some transfer does occur as evidenced by levels in cord blood of athyreotic teluses being approximately one-third maternal levels. Transfer of thyroid hormone from the mother to the fetus, however, may not be adequate to prevent in utero hypothyroidism.

ero hypothyroidism.

Nursing Mothers – Although thyroid hormones are excreted only minimally in human milk, caution nould be exercised when LEVOXYL® is administered to a nursing woman. However, adequate placement doses of levothyroxine are generally needed to maintain normal lactation.

Pediatric Use

General
The goal of treatment in pediatric patients with hypothyroidism is to achieve and maintain normal intellectual and physical growth and development.

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The initial dose of levothyrozine varies with age and body weight (see DOSAGE AND ADMINISTRATION) Table 3). Dosing aguistments are used on an assessment of the individual patient's clinical and laboratory parameters (see PRECAUTIONS, Laboratory Tests).

In children in whom a diagnosis of permanent hypothyroidism has not been established, it is recommended that levothyroxine administration be discontinued for a 30-day trial period, but only after the tRH is a state of the seed of the seed of the seed of the seed of the tRH is a state of the seed of the seed of the tRH is a state of the seed of the seed of the tRH high the diagnosis of permanent hypothyroidism is established, and levothyroxine therapy should be reinstituded. If the 1_g and TSH levels are normal, euthyroidism may be assumed and, therefore, the hypothyroidism can be considered to have been transient in this instance, however, the physician should cartully monitor the child and repeat the thyroid function tests if any signs or symptoms of the results of the levothyroxine withdrawal test are incondusive, careful follow-up and subsequent testing will be necessary.

Since some more severely affected children may become clinically hypothyroid when treatment is discontinued for 30 days, an alternate approach is to reduce the replacement dose of levothyroxine by half during the 30-day trial period. If, after 30 days, the serum TSH is elevated above 20 mU/L, the diagnosis of permanent hypothyroidism is confirmed, and full replacement therapy should be resumed. However, if the serum TSH has not risen to greater than 20 mU/L, levothyroxine treatment is discontinued for another 30-day trial period followed by repeat serum TSH and TSH.

The presence of concomitant medical conditions should be considered in certain clinical crimustances and if present, appropriately treated (see PRECAUTIONS).

Congenital hypothyro

maturation. Therefore, LEVOXYL* therapy should be initiated immediately upon diagnosis and is generally continued for life.

During the first 2 weeks of LEVOXYL* therapy, infants should be closely monitored for cardiac overload, arriythmias, and aspiration from avid suckling.

The patient should be monitored closely to avoid undertreatment or overtreatment. Undertreatment may have deleterious effects on intellectual development and linear growth. Overtreatment has been associated with cranicosynosiosis in infants, and may adversely affect the tempo of brain maturation and accelerate the lone age with resultant premature closure of the epophyses and compromised adult stature. Reaction Happith or an Intellectual Collectual and the stature of the control of th

Geratric Use

Because of the increased prevalence of cardiovascular disease among the elderly, levothyroxine therapy should not be initiated at the full replacement dose (see WARNINGS, PRECAUTIONS, and DOSAGE AND ADMINISTRATION).

Adverse Reactions

Adverse reactions associated with levothyroxine therapy are primarily those of hyperthyroidism due to therapeutic overdosage. They include the following:

General: failing, increased applict, weight loss, heat intolerance, fever, excessive sweating;

Central nervous system: headache, hyperactivity, nervousness, anxiety, irritability, emotional lability,

somnia; Musculoskeletal: tremors, muscle weakness; Cardiac: palpitations, tachycardia, arrhythmias, increased pulse and blood pressure, heart failure gina, myocardial infarction, cardiac arrest;

angina, myocardial infarction, cardice arrest;

*Pulmanary,*Ospinas;

*Git diarrhea, vomiting, abdominal cramps;

*Bernatologis: hair loss, flushing;

*Repraductive: menistrual irregularities, impaired fertility.

*Pseudotumor cerebi and slipped capital femoral epiphysis have been reported in children receiving evoltyroxine therapy. Overtreatment may result in cranicognostics in infants and prenature closure of the epiphyses in children with resultant compromised adult height.

*Sectures have been reported rarely with the institution of levolthyroxine therapy.

*Inadequate levolthyroxine dosage will produce or fail to ameliorate the signs and symptoms of hypothyroxine.

hypothypoidism. Hypersensitivity reactions to inactive ingredients have occurred in patients treated with thyroid hypersensitivity reactions to inactive ingredients have occurred in patients treated with thyroid homone products. These include urticaria, purufus, skin rash, flushing, angiodetena, various Gi symptoms (abdominal pain, nausea, owniting and diarrhea), fever, arthrafagia, serum sickness and wheezing, Hypersensitivity to levothyroxine itself is not known to occur. In addition to the above events, the following have been reported, predominately when Levoxyl* tablets were not taken with water. choking, gagging, tablet stuck in throat and dysphagia (see Information for Patients).

OVERDOSAGE

OVERDOSAGE
The signs and symptoms of overdosage are those of hyperthyroidism (see PRECAUTIONS and ADVERSE REACTIONS), in addition, confusion and disorientation may occur. Gerebral embolism, shock, coma, and death have been reported. Seizures have occurred in a child ingesting approximately 20 mg of levothyroxine. Symptoms may not necessarily be evident or may not appear until several days after ingestion of levothyroxine sodium. Treatment of Overdosarilo, may not appear until several days after ingestion of levothyroxine sodium. Treatment of Overdosarilo, and the control of the control over the control of the control of the control over the control of the control over the contr

Action Massive Overfosage — This may be a life-threatening emergency, therefore, symptomatic and supportive therapy should be instituted immediately. If not contraindicated (e.g., by secures, compared to loss of the gap reflexy), the stomach should be empidied by emests or gastric lavage to decrease gastrointestinal absorption. Activated charcoal or choicetyramine may also be used to decrease gastrointestinal absorption. Activated charcoal or choicetyramine may also be used to decrease absorption. Central and peripheral increased sympathetic activity may be treated by administrated to the contract of the contract o

DOSAGE AND ADMINISTRATION

DOSAGE AND ADMINISTRATION

General Principles:

The goal of replacement therapy is to achieve and maintain a clinical and biochemical euthyroid state. The goal of supersessive therapy is to inhibit growth and/or function of abnormal thyroid tissue. The dose of LEVOXYL* that is adequate to achieve these goals depends on a variety of factors including the patient's age, body weight, cardiovascular status, concomitant medical conditions, including pregnancy, concomitant medications, and the specific nature of the condition being treated (see WARNINGS and PRECAUTIONS, 1-there, the following recommendations serve only as dosing guidelines. Dosing must be individualized and adjustments made based on periodic assessment of the patient's clinical response and bloratory parameters (see PRECAUTIONS, Laboratory 17 est) and Latest one-half browt before any food is eaten. LEVOXYL* should be taken might and only a from drugs that are known to interfere with its absorption (see PRECAUTIONS, Dury Interactions).

LEVOXYL* should be taken with water (see Informations for Patients and ADVERSE REACTIONS).

Due to the long half-life of levolthyroxine, the peak therapeutic effect at a given dose of levothyroxine sodium may not be attained for 4-6 weeks.

sodium may not be attained for 4–6 weeks.
Caution should be exercised when administering LEVOXYL® to patients with underlying cardiovascular disease, to the elderly, and to those with concomitant adrenal insufficiency (see PRECAUTIONS)
Specific Patient Populations:
Hypothroxidism in Adults and in Children in Whom Growth and Puberty are Complete (see WARNINGS and PRECAUTIONS). Laboratory Tests)
Therappy may begin at 4-fill amount of the patient of the p

and PRECAUTIONS. Laboratory Tests!
Therapy may begin at full replacement doses in otherwise healthy individuals less than 50 years old and in those older than 50 years who have been recently treated for hyperthyroidism or who have been reportlyorid for only a short time (such as a few months). The average full replacement dose of levothyroine sodium is approximately 1.7 mcg/kg/day (e.g. 100-125 mcg/day for a 70 kg adult). Older patients may require less than 1 mcg/kg/day. Levothyroines sodium doses greater than 200 mcg/day seldom required. An inadequable response to daily doses 300 mcg/day is rare and may indicate poor compliance, malabeoprofile, andiof ording interactions.

compliance, malabsorption, and/or drug interactions.

For most patients older than 30 years of for patients under 50 years of age with underlying cardiac.

For most patients older than 30 years of for patients under 50 years of age with underlying cardiac for most patients older than 30 years of for patients with patients with patients of the patients with cardiac diseases is 12.5-25 mog/day, with pradual dose increments at 4-6 week intervals. The breithyroxine sodium in elderly patients with cardiac diseases is 12.5-25 mog/day, with pradual dose increments at 4-6 week intervals. The breithyroxine sodium dose is generally adjusted in 12.5-25 mog increments until the patient with primary hypothyroidism, the circummended initial evoltyroxine sodium dose is 12.5-25 mog/day with increases of 25 mog/day every 2-4 weeks, accompanied by clinical and laboratory assessment, until the TSH level is normalized.

In patients with secondary (pitultary) or terfary (hypothalamic) hypothyroidism, the levothyroxine sodium dose should be titrated until the patient is clinically euthyroid and the serum free-14 level is restored to the upper half of the normal range.

For most patients with secondary (accounted Hypothyroidism (see PRECAUTIONS, Laboratory Tests)

Pediatric Dosage – Congenital or Acquired Hypothyroidism (see PRECAUTIONS, Laboratory Tests).

Il Principles eneral, levothyroxine therapy should be instituted at full replacement doses as soon as poss in diagnosis and institution of therapy may have deleterious effects on the child's intellectua

Physical growth and development.

Undertreatment and overtreatment should be avoided (see PRECAUTIONS, Pediatric Use).

LEVOXY.® may be administered to infants and children who cannot swallow intact tablets by

the tablet and suspending the freshly crushed tablet in a small amount (5–10 mL or 1–2 teaspoons) or water. This suspension can be administered by spoon or dropper. **DO NOT STORE THE SUSPENSION** Foods that decrease absorption of eventyroxine, such as oxylean intant fromula, should not be used to administering levothyroxine sodium tablets. (see **PRECAUTIONS**, **Drug-Food Interactions**).

Newborns
The recommended starting dose of levothyroxine sodium in newborn infants is 19–15 mcg/kg/day. The recommended starting dose of levothyroxine sodium in newborn infants is 16–15 mcg/kg/day. An observation of levothyroxine sodium infants at risk for cardiac failure, and the dose should be increased in 4-5 weeks as needed based on clinical and laboratory response to treatment. In infants with very love (< 5 mcg/dc) or undetectable serum T₄ concentrations, the recommended infinal starting dose is 50 mcg/day of levothyroxine sodium.

Intains and Children
Levothyroxine therapy is usually initiated at full replacement doses, with the recommended dose per body weight decreasing with age (see TABLE 3). However, in children with chronic or severe hypothyroidism, an initial dose of 25 mag(day of levothyroxine sodium is recommended with increments of 25 mag every 2-4 weeks until the desired effect is achieved.
Hyperactivity in an older child can be minimized if the starting dose is one-fourth of the recommended ill replacement dose, and the dose is then increased on a weekly basis by an amount equal to one-fourth the full recommended replacement dose until the full recommended replacement dose is reached.

AGE Daily Dose Per Ko Body Weight		
0-3 months	10-15 mcg/kg/day	
3–6 months	8-10 mcg/kg/day	
6-12 months	6–8 mcg/kg/day 5–6 mcg/kg/day 4–5 mcg/kg/day	
1–5 years		
6-12 years		
>12 years 2-3 mcg/kg/day		
Growth and puberty complete 1.7 mcg/kg/day		

Pregnancy—Pregnancy may increase levothyroxine requirements (see **PREGNANCY**).

Subclinical Hypothyroidism— If this condition is treated, a lower levothyroxine sodium dose (e.g. 1 neg/kg/day) than that used for full replacement may be adequate to normalize the serum TSH level admits who are not treated should be monitored yearly for changes in clinical status and thyroid laboratory parameters.

laboratory parameters.

75H Suppression in Welf-differentiated Thyroid Cancer and Thyroid Modules — The target level for TSH suppression in Welf-differentiated Thyroid Cancer and Thyroid Modules — The target level for TSH suppression in these conditions has not been established with controlled studies. In addition, the efficacy of TSH suppression for benign nodular disease is controlled studies. In addition, the efficacy of TSH suppression should be individualized based on the specific disease and the patient being treated. In the treatment of vell differentiated (papillary and folicular) thyroid cancer, levothyroids is used as an adjunct to surgery and radioiodine therapy. Generally, TSH is suppressed to -0.1 mU/L, and this susually requires a levothyroxies obdimin dose of greater than 2 meg/agi/agi. However, in patients with high-risk tumors, the target level for TSH suppression may be -0.01 mU/L. In multimodular gother, TSH is generally suppressed to a higher target (e.g., 0.1-0.5 mU/L for modules and 0.5-1.0 mU/L for multimodular gother. TSH is generally suppressed to a higher target (e.g., 0.1-0.5 mU/L for modules and 0.5-1.0 mU/L for multimodular gother) than that used a higher target (e.g., 0.1-0.5 mU/L for modules and 0.5-1.0 mU/L for multimodular gother) than that used suppressed the to the risk of precipitating overt thyrotoxicoxis (see CONTRAINDICATIONS, WARNINGS and PRECAUTIONS).

America (Nova University). Myzedema coma is a life-threatening emergency characterized by poor circulation and hypometabolism, and may result in unpredictable absorption of levothyroxine sodium from the distribution of the commended to treat this condition. Thyroid hormone drug products are not recommended to treat this condition. Thyroid hormone products formulated for intravenous administration should be administered.

HOW SUPPLIED

Strength (mcg)	Color	NDC # for bottles of 100	NDC # for bottles of 1000	NDC # for Unit Dose Cartons of 100
25	Orange	NDC 52604-5025-1	NDC 52604-5025-2	NDC 52604-5025-5
50	White	NDC 52604-5050-1	NDC 52604-5050-2	NDC 52604-5050-5
75	Purple	NDC 52604-5075-1	NDC 52604-5075-2	NDC 52604-5075-5
88	Olive	NDC 52604-5088-1	NDC 52604-5088-2	NDC 52604-5088-5
100	Yellow	NDC 52604-5100-1	NDC 52604-5100-2	NDC 52604-5100-5
112	Rose	NDC 52604-5112-1	NDC 52604-5112-2	NDC 52604-5112-5
125	Brown	NDC 52604-5125-1	NDC 52604-5125-2	NDC 52604-5125-5
137	Dark Blue	NDC 52604-5137-1	NDC 52604-5137-2	NDC 52604-5137-5
150	Blue	NDC 52604-5150-1	NDC 52604-5150-2	NDC 52604-5150-5
175	Turquoise	NDC 52604-5175-1	NDC 52604-5175-2	NDC 52604-5175-5
200	Pink	NDC 52604-5200-1	NDC 52604-5200-2	NDC 52604-5200-5
300	Green	NDC 52604-5300-1	NDC 52604-5300-2	NDC 52604-5300-5

LEVOXYL® (levothyroxine sodium tablets, USP) are supplied as oval, color-coded, potency marked tablets in 12 strengths:

STORAGE CONDITIONS
20°-25°C (68°-77°F) with excursions permitted between 15°-30°C (59°-86°F).
Meets USP Dissolution Tests 1 and 2.

Rx ONLY

MANUFACTURER
JONES PHARMA INCORPORATED
(A wholly owned subsidiary of King Pharmaceuticals, Inc.)
Bristol, VA 24201

