DEXAMETHASONE- dexamethasone tablet DEXAMETHASONE INTENSOL- dexamethasone intensol solution, concentrate DEXAMETHASONE- dexamethasone solution Hikma Pharmaceuticals USA, Inc.

Dexamethasone Tablets, USP Dexamethasone Oral Solution, USP and Dexamethasone Oral Solution, USP *Intensol*™ (Concentrate)

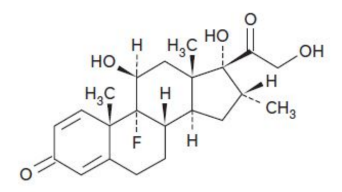
DESCRIPTION

Dexamethasone Tablets, USP are available for oral administration containing either 0.5 mg, 0.75 mg, 1 mg, 1.5 mg, 2 mg, 4 mg or 6 mg of dexamethasone, USP. Each tablet contains the following inactive ingredients: lactose monohydrate, magnesium stearate, starch, sugar, D&C Yellow #10 (0.5 mg and 4 mg), FD&C Blue #1 (0.75 mg and 1.5 mg), FD&C Green #3 (4 mg and 6 mg), FD&C Red #3 (1.5 mg), FD&C Red #40 (1.5 mg), FD&C Yellow #6 (0.5 mg and 4 mg) and Yellow Iron Oxide (1 mg).

Dexamethasone Oral Solution, USP is formulated for oral administration containing 0.5 mg per 5 mL of dexamethasone, USP. The cherry brandy flavored oral solution contains the following inactive ingredients: anhydrous citric acid, cherry brandy flavor, disodium edetate, glycerin, methylparaben, propylene glycol, propylparaben, sorbitol solution and water.

Dexamethasone Oral Solution, USP *Intensol*[™] (Concentrate) is formulated for oral administration containing 1 mg per mL of dexamethasone, USP. In addition, the oral solution contains the following inactive ingredients: alcohol 30% v/v, anhydrous citric acid, benzoic acid, disodium edetate, propylene glycol and water.

Dexamethasone, a synthetic adrenocortical steroid, is a white to practically white, odorless, crystalline powder. It is stable in air. It is practically insoluble in water. The empirical formula is C₂₂H₂₉FO₅. The molecular weight is 392.46. It is designated chemically as 9-fluoro-11 β ,17,21-trihydroxy-16 α -methylpregna-1,4-diene,3,20-dione and the structural formula is:



Glucocorticoids, naturally occurring and synthetic, are adrenocortical steroids that are readily absorbed from the gastrointestinal tract. Glucocorticoids cause varied metabolic effects. In addition, they modify the body's immune responses to diverse stimuli. Naturally occurring glucocorticoids (hydrocortisone and cortisone), which also have sodium-retaining properties, are used as replacement therapy in adrenocortical deficiency states. Their synthetic analogs including dexamethasone are primarily used for their anti-inflammatory effects in disorders of many organ systems.

At equipotent anti-inflammatory doses, dexamethasone almost completely lacks the sodium-retaining property of hydrocortisone and closely related derivatives of hydrocortisone.

INDICATIONS AND USAGE

Allergic States

Control of severe or incapacitating allergic conditions intractable to adequate trials of conventional treatment in asthma, atopic dermatitis, contact dermatitis, drug hypersensitivity reactions, perennial or seasonal allergic rhinitis, and serum sickness.

Dermatologic Diseases

Bullous dermatitis herpetiformis, exfoliative erythroderma, mycosis fungoides, pemphigus, and severe erythema multiforme (Stevens-Johnson syndrome).

Endocrine Disorders

Primary or secondary adrenocortical insufficiency (hydrocortisone or cortisone is the drug of choice; may be used in conjunction with synthetic mineralocorticoid analogs where applicable; in infancy mineralocorticoid supplementation is of particular importance), congenital adrenal hyperplasia, hypercalcemia associated with cancer, and nonsuppurative thyroiditis.

Gastrointestinal Diseases

To tide the patient over a critical period of the disease in regional enteritis and ulcerative colitis.

Hematologic Disorders

Acquired (autoimmune) hemolytic anemia, congenital (erythroid) hypoplastic anemia (Diamond-Blackfan anemia), idiopathic thrombocytopenic purpura in adults, pure red cell aplasia, and selected cases of secondary thrombocytopenia.

Miscellaneous

Diagnostic testing of adrenocortical hyperfunction, trichinosis with neurologic or myocardial involvement, tuberculous meningitis with subarachnoid block or impending block when used with appropriate antituberculous chemotherapy.

Neoplastic Diseases

For the palliative management of leukemias and lymphomas.

Nervous System

Acute exacerbations of multiple sclerosis, cerebral edema associated with primary or metastatic brain tumor, craniotomy, or head injury.

Ophthalmic Diseases

Sympathetic ophthalmia, temporal arteritis, uveitis, and ocular inflammatory conditions unresponsive to topical corticosteroids.

Renal Diseases

To induce a diuresis or remission of proteinuria in idiopathic nephrotic syndrome or that due to lupus erythematosus.

Respiratory Diseases

Berylliosis, fulminating or disseminated pulmonary tuberculosis when used concurrently with appropriate antituberculous chemotherapy, idiopathic eosinophilic pneumonias, symptomatic sarcoidosis.

Rheumatic Disorders

As adjunctive therapy for short-term administration (to tide the patient over an acute episode or exacerbation) in acute gouty arthritis, acute rheumatic carditis, ankylosing spondylitis, psoriatic arthritis, rheumatoid arthritis, including juvenile rheumatoid arthritis (selected cases may require low-dose maintenance therapy). For the treatment of dermatomyositis, polymyositis, and systemic lupus erythematosus.

CONTRAINDICATIONS

Systemic fungal infections (see **WARNINGS: Fungal Infections**) and in patients who are hypersensitive to any components of these products.

WARNINGS

General

Rare instances of anaphylactoid reactions have occurred in patients receiving corticosteroid therapy (see **ADVERSE REACTIONS**).

Increased dosage of rapidly acting corticosteroids is indicated in patients on corticosteroid therapy subjected to any unusual stress before, during, and after the stressful situation.

Immunosuppression and Increased Risk of Infection

Corticosteroids, including dexamethasone, suppress the immune system and increase the risk of infection with any pathogen, including viral, bacterial, fungal, protozoan, or helminthic pathogens. Corticosteroids can:

• Reduce resistance to new infections

- Exacerbate existing infections
- Increase the risk of disseminated infections
- Increase the risk of reactivation or exacerbation of latent infections
- Mask some signs of infection

Corticosteroid-associated infections can be mild but can be severe and at times fatal. The rate of infectious complications increases with increasing corticosteroid dosages.

Monitor for the development of infection and consider dexamethasone withdrawal or dosage reduction as needed.

Do not administer dexamethasone by an intraarticular, intrabursal, intratendinous, or intralesional route in the presence of acute local infection.

Tuberculosis

If dexamethasone is used to treat a condition in patients with latent tuberculosis or tuberculin reactivity, reactivation of tuberculosis may occur. Closely monitor such patients for reactivation. During prolonged dexamethasone therapy, patients with latent tuberculosis or tuberculin reactivity should receive chemoprophylaxis.

Varicella Zoster and Measles Viral Infections

Varicella and measles can have a serious or even fatal course in non-immune patients taking corticosteroids, including dexamethasone. In corticosteroid-treated patients who have not had these diseases or are non-immune, particular care should be taken to avoid exposure to varicella and measles:

- If a dexamethasone-treated patient is exposed to varicella, prophylaxis with varicella zoster immune globulin may be indicated. If varicella develops, treatment with antiviral agents may be considered.
- If a dexamethasone-treated patient is exposed to measles, prophylaxis with immunoglobulin may be indicated.

Hepatitis B Virus Reactivation

Hepatitis B virus reactivation can occur in patients who are hepatitis B carriers treated with immunosuppressive dosages of corticosteroids, including dexamethasone. Reactivation can also occur infrequently in corticosteroid-treated patients who appear to have resolved hepatitis B infection.

Screen patients for hepatitis B infection before initiating immunosuppressive (e.g., prolonged) treatment with dexamethasone. For patients who show evidence of hepatitis B infection, recommend consultation with physicians with expertise in managing hepatitis B regarding monitoring and consideration for hepatitis B antiviral therapy.

Fungal Infections

Corticosteroids, including dexamethasone, may exacerbate systemic fungal infections; therefore, avoid dexamethasone use in the presence of such infections unless dexamethasone is needed to control drug reactions. For patients on chronic dexamethasone therapy who develop systemic fungal infections, dexamethasone withdrawal or dosage reduction is recommended.

Amebiasis

Corticosteroids, including dexamethasone, may activate latent amebiasis. Therefore, it is recommended that latent amebiasis or active amebiasis be ruled out before initiating dexamethasone in patients who have spent time in the tropics or patients with unexplained diarrhea.

Strongyloides Infestation

Corticosteroids, including dexamethasone, should be used with great care in patients with known or suspected Strongyloides (threadworm) infestation. In such patients, corticosteroid-induced immunosuppression may lead to Strongyloides hyperinfection and dissemination with widespread larval migration, often accompanied by severe enterocolitis and potentially fatal gram-negative septicemia.

Cerebral Malaria

Avoid corticosteroids, including dexamethasone, in patients with cerebral malaria.

Vaccination

Administration of live or live, attenuated vaccines is contraindicated in patients receiving immunosuppressive doses of corticosteroids. Killed or inactivated vaccines may be administered. However, the response to such vaccines cannot be predicted. Immunization procedures may be undertaken in patients who are receiving corticosteroids as replacement therapy, e.g., for Addison's disease.

Ophthalmic

Use of corticosteroids may produce posterior subcapsular cataracts, glaucoma with possible damage to the optic nerves, and may enhance the establishment of secondary ocular infections due to bacteria, fungi, or viruses. Consider referral to an ophthalmologist for patients who develop ocular symptoms or use corticosteroid-containing products for more than 6 weeks. The use of oral corticosteroids is not recommended in the treatment of optic neuritis and may lead to an increase in the risk of new episodes. Corticosteroids should not be used in active ocular herpes simplex.

Kaposi's Sarcoma

Kaposi's sarcoma has been reported to occur in patients receiving corticosteroid therapy, most often for chronic conditions. Discontinuation of corticosteroids may result in clinical improvement of Kaposi's sarcoma.

Cardio-Renal

Average and large doses of corticosteroids can cause elevation of blood pressure, sodium and water retention, and increased excretion of potassium. These effects are less likely to occur with the synthetic derivatives except when used in large doses. Dietary salt restriction and potassium supplementation may be necessary. All corticosteroids increase calcium excretion.

Literature reports suggest an apparent association between use of corticosteroids and left ventricular free wall rupture after a recent myocardial infarction; therefore, therapy

with corticosteroids should be used with great caution in these patients.

Endocrine

Corticosteroids can produce reversible hypothalamic-pituitary adrenal (HPA) axis suppression with the potential for glucocorticosteroid insufficiency after withdrawal of treatment. Adrenocortical insufficiency may result from too rapid withdrawal of corticosteroids and may be minimized by gradual reduction of dosage. This type of relative insufficiency may persist for months after discontinuation of therapy; therefore, in any situation of stress occurring during that period, hormone therapy should be reinstituted. If the patient is receiving steroids already, dosage may have to be increased.

Metabolic clearance of corticosteroids is decreased in hypothyroid patients and increased in hyperthyroid patients. Changes in thyroid status of the patient may necessitate adjustment in dosage.

PRECAUTIONS

General

The lowest possible dose of corticosteroids should be used to control the condition under treatment. When reduction in dosage is possible, the reduction should be gradual.

Since complications of treatment with corticosteroids are dependent on the size of the dose and the duration of treatment, a risk/benefit decision must be made in each individual case as to dose and duration of treatment and as to whether daily or intermittent therapy should be used.

Cardio-Renal

As sodium retention with resultant edema and potassium loss may occur in patients receiving corticosteroids, these agents should be used with caution in patients with congestive heart failure, hypertension, or renal insufficiency.

Endocrine

Drug-induced secondary adrenocortical insufficiency may be minimized by gradual reduction of dosage. This type of relative insufficiency may persist for months after discontinuation of therapy; therefore, in any situation of stress occurring during that period, hormone therapy should be reinstituted. Since mineralocorticoid secretion may be impaired, salt and/or a mineralocorticoid should be administered concurrently.

Gastrointestinal

Steroids should be used with caution in active or latent peptic ulcers, diverticulitis, fresh intestinal anastomoses, and nonspecific ulcerative colitis, since they may increase the risk of a perforation.

Signs of peritoneal irritation following gastrointestinal perforation in patients receiving corticosteroids may be minimal or absent.

There is an enhanced effect due to decreased metabolism of corticosteroids in patients

with cirrhosis.

Musculoskeletal

Corticosteroids decrease bone formation and increase bone resorption both through their effect on calcium regulation (i.e., decreasing absorption and increasing excretion) and inhibition of osteoblast function. This, together with a decrease in the protein matrix of the bone secondary to an increase in protein catabolism, and reduced sex hormone production, may lead to inhibition of bone growth in pediatric patients and the development of osteoporosis at any age. Special consideration should be given to patients at increased risk of osteoporosis (e.g., postmenopausal women) before initiating corticosteroid therapy.

Neuro-Psychiatric

Although controlled clinical trials have shown corticosteroids to be effective in speeding the resolution of acute exacerbations of multiple sclerosis, they do not show that they affect the ultimate outcome or natural history of the disease. The studies do show that relatively high doses of corticosteroids are necessary to demonstrate a significant effect (see **DOSAGE AND ADMINISTRATION**).

An acute myopathy has been observed with the use of high doses of corticosteroids, most often occurring in patients with disorders of neuromuscular transmission (e.g., myasthenia gravis), or in patients receiving concomitant therapy with neuromuscular blocking drugs (e.g., pancuronium). This acute myopathy is generalized, may involve ocular and respiratory muscles, and may result in quadriparesis. Elevation of creatinine kinase may occur. Clinical improvement or recovery after stopping corticosteroids may require weeks to years.

Psychic derangements may appear when corticosteroids are used, ranging from euphoria, insomnia, mood swings, personality changes, and severe depression, to frank psychotic manifestations. Also, existing emotional instability or psychotic tendencies may be aggravated by corticosteroids.

Ophthalmic

Intraocular pressure may become elevated in some individuals. If steroid therapy is continued for more than 6 weeks, intraocular pressure should be monitored.

Information for Patients

Patients should be warned not to discontinue the use of corticosteroids abruptly or without medical supervision. As prolonged use may cause adrenal insufficiency and make patients dependent on corticosteroids, they should advise any medical attendants that they are taking corticosteroids and they should seek medical advice at once should they develop an acute illness including fever or other signs of infection. Following prolonged therapy, withdrawal of corticosteroids may result in symptoms of the corticosteroid withdrawal syndrome including myalgia, arthralgia, and malaise.

Persons who are on corticosteroids should be warned to avoid exposure to chickenpox or measles. Patients should also be advised that if they are exposed, medical advice should be sought without delay.

Drug Interactions

Aminoglutethimide: Aminoglutethimide may diminish adrenal suppression by corticosteroids.

Amphotericin B injection and potassium-depleting agents: When corticosteroids are administered concomitantly with potassium-depleting agents (e.g., amphotericin B, diuretics), patients should be observed closely for development of hypokalemia. In addition, there have been cases reported in which concomitant use of amphotericin B and hydrocortisone was followed by cardiac enlargement and congestive heart failure.

Antibiotics: Macrolide antibiotics have been reported to cause a significant decrease in corticosteroid clearance (see *Drug Interactions: CYP 3A4 Inducers, CYP 3A4 Inhibitors,* and *CYP 3A4 Substrates*).

Anticholinesterases: Concomitant use of anticholinesterase agents and corticosteroids may produce severe weakness in patients with myasthenia gravis. If possible, anticholinesterase agents should be withdrawn at least 24 hours before initiating corticosteroid therapy.

Anticoagulants, oral: Co-administration of corticosteroids and warfarin usually results in inhibition of response to warfarin, although there have been some conflicting reports. Therefore, coagulation indices should be monitored frequently to maintain the desired anticoagulant effect.

Antidiabetics: Because corticosteroids may increase blood glucose concentrations, dosage adjustments of antidiabetic agents may be required.

Antitubercular drugs: Serum concentrations of isoniazid may be decreased.

Cholestyramine: Cholestyramine may increase the clearance of corticosteroids.

Cyclosporine: Increased activity of both cyclosporine and corticosteroids may occur when the two are used concurrently. Convulsions have been reported with this concurrent use.

Dexamethasone suppression test (DST): False-negative results in the dexamethasone suppression test (DST) in patients being treated with indomethacin have been reported. Thus, results of the DST should be interpreted with caution in these patients.

Digitalis glycosides: Patients on digitalis glycosides may be at increased risk of arrhythmias due to hypokalemia.

Ephedrine: Ephedrine may enhance the metabolic clearance of corticosteroids, resulting in decreased blood levels and lessened physiologic activity, thus requiring an increase in corticosteroid dosage.

Estrogens, including oral contraceptives: Estrogens may decrease the hepatic metabolism of certain corticosteroids, thereby increasing their effect.

CYP 3A4 Inducers: Dexamethasone is metabolized by CYP 3A4. Drugs which induce cytochrome P450 3A4 (CYP 3A4) enzyme activity (*e.g., barbiturates, phenytoin, carbamazepine, rifampin*) may enhance the metabolism of corticosteroids and require that the dosage of the corticosteroid be increased.

CYP 3A4 Inhibitors: Concomitant administration of dexamethasone with erythromycin, a moderate CYP 3A4 inhibitor, has the potential to result in increased plasma

concentrations of dexamethasone. Ketoconazole, a strong CYP3A4 inhibitor, has been reported to decrease the metabolism of certain corticosteroids by up to 60%, leading to increased risk of corticosteroid side effects. In addition, ketoconazole alone can inhibit adrenal corticosteroid synthesis and may cause adrenal insufficiency during corticosteroid withdrawal. Co-administration with other drugs which strongly inhibit CYP 3A4 (e.g., itraconazole, clarithromycin, ritonavir, cobicistat-containing products) may lead to increased plasma concentrations of corticosteroids and potentially increase the risk for systemic corticosteroid side effects. Consider the benefit of co-administration versus the potential risk of systemic corticosteroid effects, in which case patients should be monitored for systemic corticosteroid side effects.

CYP 3A4 Substrates: Dexamethasone is a moderate inducer of CYP 3A4. Coadministration with other drugs that are metabolized by CYP 3A4 (*e.g., indinavir, erythromycin*) may increase their clearance, resulting in decreased plasma concentration.

Nonsteroidal Anti-Inflammatory Agents (NSAIDS): Concomitant use of aspirin (or other nonsteroidal anti-inflammatory agents) and corticosteroids increases the risk of gastrointestinal side effects. Aspirin should be used cautiously in conjunction with corticosteroids in hypoprothrombinemia. The clearance of salicylates may be increased with concurrent use of corticosteroids.

Phenytoin: In post-marketing experience, there have been reports of both increases and decreases in phenytoin levels with dexamethasone co-administration, leading to alterations in seizure control.

Skin Tests: Corticosteroids may suppress reactions to skin tests.

Thalidomide: Co-administration with thalidomide should be employed cautiously, as toxic epidermal necrolysis has been reported with concomitant use.

Vaccines: Patients on corticosteroid therapy may exhibit a diminished response to toxoids and live or inactivated vaccines due to inhibition of antibody response. Corticosteroids may also potentiate the replication of some organisms contained in live attenuated vaccines. Routine administration of vaccines or toxoids should be deferred until corticosteroid therapy is discontinued if possible (see **WARNINGS: Infections: Vaccination**).

Carcinogenesis, Mutagenesis, Impairment of Fertility

No adequate studies have been conducted in animals to determine whether corticosteroids have a potential for carcinogenesis or mutagenesis.

Steroids may increase or decrease motility and number of spermatozoa in some patients.

Pregnancy

Teratogenic Effects: Corticosteroids have been shown to be teratogenic in many species when given in doses equivalent to the human dose. Animal studies in which corticosteroids have been given to pregnant mice, rats, and rabbits have yielded an increased incidence of cleft palate in the offspring. There are no adequate and well-controlled studies in pregnant women. Corticosteroids should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus. Infants born to

mothers who have received substantial doses of corticosteroids during pregnancy should be carefully observed for signs of hypoadrenalism.

Nursing Mothers

Systemically administered corticosteroids appear in human milk and could suppress growth, interfere with endogenous corticosteroid production, or cause other untoward effects. Because of the potential for serious adverse reactions in nursing infants from corticosteroids, a decision should be made whether to discontinue nursing or to discontinue the drug, taking into account the importance of the drug to the mother.

Pediatric Use

The efficacy and safety of corticosteroids in the pediatric population are based on the well-established course of effect of corticosteroids, which is similar in pediatric and adult populations. Published studies provide evidence of efficacy and safety in pediatric patients for the treatment of nephrotic syndrome (patients >2 years of age), and aggressive lymphomas and leukemias (patients >1 month of age). Other indications for pediatric use of corticosteroids, e.g., severe asthma and wheezing, are based on adequate and well-controlled trials conducted in adults, on the premises that the course of the diseases and their pathophysiology are considered to be substantially similar in both populations.

The adverse effects of corticosteroids in pediatric patients are similar to those in adults (see **ADVERSE REACTIONS**). Like adults, pediatric patients should be carefully observed with frequent measurements of blood pressure, weight, height, intraocular pressure, and clinical evaluation for the presence of infection, psychosocial disturbances, thromboembolism, peptic ulcers, cataracts, and osteoporosis. Pediatric patients who are treated with corticosteroids by any route, including systemically administered corticosteroids, may experience a decrease in their growth velocity. This negative impact of corticosteroids on growth has been observed at low systemic doses and in the absence of laboratory evidence of hypothalamic-pituitary-adrenal (HPA) axis suppression (i.e., cosyntropin stimulation and basal cortisol plasma levels). Growth velocity may therefore be a more sensitive indicator of systemic corticosteroid exposure in pediatric patients than some commonly used tests of HPA axis function. The linear growth of pediatric patients treated with corticosteroids should be monitored, and the potential growth effects of prolonged treatment should be weighed against clinical benefits obtained and the availability of treatment alternatives. In order to minimize the potential growth effects of corticosteroids, pediatric patients should be *titrated* to the lowest effective dose.

Geriatric Use

Clinical studies did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects. Other reported clinical experience has not identified differences in responses between the elderly and younger patients. In general, dose selection for an elderly patient should be cautious, usually starting at the low end of the dosing range, reflecting the greater frequency of decreased hepatic, renal, or cardiac function, and of concomitant disease or other drug therapy. In particular, the increased risk of diabetes mellitus, fluid retention and hypertension in elderly patients treated with corticosteroids should be considered.

ADVERSE REACTIONS

(Listed alphabetically, under each subsection)

The following adverse reactions have been reported with dexamethasone or other corticosteroids:

Allergic Reactions

Anaphylactoid reaction, anaphylaxis, angioedema.

Cardiovascular

Bradycardia, cardiac arrest, cardiac arrhythmias, cardiac enlargement, circulatory collapse, congestive heart failure, fat embolism, hypertension, hypertrophic cardiomyopathy in premature infants, myocardial rupture following recent myocardial infarction (see **WARNINGS: Cardio-Renal**), edema, pulmonary edema, syncope, tachycardia, thromboembolism, thrombophlebitis, vasculitis.

Dermatologic

Acne, allergic dermatitis, dry scaly skin, ecchymoses and petechiae, erythema, impaired wound healing, increased sweating, rash, striae, suppression of reactions to skin tests, thin fragile skin, thinning scalp hair, urticaria.

Endocrine

Decreased carbohydrate and glucose tolerance, development of cushingoid state, hyperglycemia, glycosuria, hirsutism, hypertrichosis, increased requirements for insulin or oral hypoglycemic agents in diabetes, manifestations of latent diabetes mellitus, menstrual irregularities, secondary adrenocortical and pituitary unresponsiveness (particularly in times of stress, as in trauma, surgery, or illness), suppression of growth in pediatric patients.

Fluid and Electrolyte Disturbances

Congestive heart failure in susceptible patients, fluid retention, hypokalemic alkalosis, potassium loss, sodium retention, tumor lysis syndrome.

Gastrointestinal

Abdominal distention, elevation in serum liver enzyme levels (usually reversible upon discontinuation), hepatomegaly, increased appetite, nausea, pancreatitis, peptic ulcer with possible perforation and hemorrhage, perforation of the small and large intestine (particularly in patients with inflammatory bowel disease), ulcerative esophagitis.

Metabolic

Negative nitrogen balance due to protein catabolism.

Musculoskeletal

Aseptic necrosis of femoral and humeral heads, loss of muscle mass, muscle weakness, osteoporosis, pathologic fracture of long bones, steroid myopathy, tendon rupture,

vertebral compression fractures.

Neurological/Psychiatric

Convulsions, depression, emotional instability, euphoria, headache, increased intracranial pressure with papilledema (pseudotumor cerebri) usually following discontinuation of treatment, insomnia, mood swings, neuritis, neuropathy, paresthesia, personality changes, psychic disorders, vertigo.

Ophthalmic

Exophthalmos, glaucoma, increased intraocular pressure, posterior subcapsular cataracts, vision blurred.

Other

Abnormal fat deposits, decreased resistance to infection, hiccups, increased or decreased motility and number of spermatozoa, malaise, moon face, weight gain.

OVERDOSAGE

Treatment of overdosage is by supportive and symptomatic therapy. In the case of acute overdosage, according to the patient's condition, supportive therapy may include gastric lavage or emesis.

DOSAGE AND ADMINISTRATION

For Oral Administration

The initial dosage varies from 0.75 mg to 9 mg a day depending on the disease being treated.

It Should Be Emphasized That Dosage Requirements Are Variable And Must Be Individualized On The Basis Of The Disease Under Treatment And The Response Of The Patient.

After a favorable response is noted, the proper maintenance dosage should be determined by decreasing the initial drug dosage in small decrements at appropriate time intervals until the lowest dosage that maintains an adequate clinical response is reached.

Situations which may make dosage adjustments necessary are changes in clinical status secondary to remissions or exacerbations in the disease process, the patient's individual drug responsiveness, and the effect of patient exposure to stressful situations not directly related to the disease entity under treatment. In this latter situation it may be necessary to increase the dosage of the corticosteroid for a period of time consistent with the patient's condition. If after long-term therapy the drug is to be stopped, it is recommended that it be withdrawn gradually rather than abruptly.

In the treatment of acute exacerbations of multiple sclerosis, daily doses of 30 mg of dexamethasone for a week followed by 4 mg to 12 mg every other day for one month have been shown to be effective (see **PRECAUTIONS: Neuro-Psychiatric**).

In pediatric patients, the initial dose of dexamethasone may vary depending on the

specific disease entity being treated. The range of initial doses is 0.02 mg to 0.3 mg/kg/day in three or four divided doses (0.6 mg to 9 mg/m²bsa/day).

For the purpose of comparison, the following is the equivalent milligram dosage of the various corticosteroids:

Cortisone, 25 mg	Triamcinolone, 4 mg
Hydrocortisone, 20 mg	Paramethasone, 2 mg
Prednisolone, 5 mg	Betamethasone, 0.75 mg
Prednisone, 5 mg	Dexamethasone, 0.75 mg
Methylprednisolone, 4 mg	

These dose relationships apply only to oral or intravenous administration of these compounds. When these substances or their derivatives are injected intramuscularly or into joint spaces, their relative properties may be greatly altered.

In acute, self-limited allergic disorders or acute exacerbations of chronic allergic disorders, the following dosage schedule combining parenteral and oral therapy is suggested:

Dexamethasone sodium phosphate injection, 4 mg per mL

First Day: 1 or 2 mL, intramuscularly

Dexamethasone tablets, 0.75 mg

Second Day: 4 tablets in two divided doses

Third Day: 4 tablets in two divided doses

Fourth Day: 2 tablets in two divided doses

Fifth Day: 1 tablet

Sixth Day: 1 tablet

Seventh Day: No treatment

Eighth Day: Follow-up visit

This schedule is designed to ensure adequate therapy during acute episodes, while minimizing the risk of overdosage in chronic cases.

In *cerebral edema*, dexamethasone sodium phosphate injection is generally administered initially in a dosage of 10 mg intravenously followed by 4 mg every six hours intramuscularly until the symptoms of cerebral edema subside. Response is usually noted within 12 to 24 hours and dosage may be reduced after two to four days and gradually discontinued over a period of five to seven days. For palliative management of patients with recurrent or inoperable brain tumors, maintenance therapy with either dexamethasone sodium phosphate injection or dexamethasone tablets in a dosage of 2 mg two or three times daily may be effective.

Dexamethasone Suppression Tests

1. Tests for Cushing's syndrome

Give 1 mg of dexamethasone orally at 11:00 p.m. Blood is drawn for plasma cortisol determination at 8:00 a.m. the following morning.

For greater accuracy, give 0.5 mg of dexamethasone orally every 6 hours for 48 hours. Twenty-four hour urine collections are made for determination of 17-hydroxycorticosteroid excretion.

 Test to distinguish Cushing's syndrome due to pituitary ACTH excess from Cushing's syndrome due to other causes.
 Give 2 mg of devemethas one orally every 6 hours for 48 hours. Twenty four hours

Give 2 mg of dexamethasone orally every 6 hours for 48 hours. Twenty-four hour urine collections are made for determination of 17-hydroxycorticosteroid excretion.

Proper Use of an *Intensol*™

An Intensol is a concentrated oral solution as compared to standard oral liquid medications. It is recommended that an Intensol be mixed with liquid or semi-solid food such as water, juices, soda or soda-like beverages, applesauce and puddings.

Use only the calibrated oral syringe provided with this product. Draw into the oral syringe the amount prescribed for a single dose. Then empty the contents of the oral syringe into a liquid or semi-solid food, by depressing the plunger until the barrel is empty. Stir the liquid or food gently for a few seconds. The Intensol formulation blends quickly and completely. The entire amount of the mixture, of drug and liquid or drug and food, should be consumed immediately. Do not store for future use.

HOW SUPPLIED

Dexamethasone Tablets, USP

0.5 mg tablets are supplied as a light yellow, flat tablet with beveled edges, scored on one side and product identification "54 299" debossed on the other side.

NDC 0054-8179-25: 10x10 Unit-Dose

NDC 0054-4179-25: Bottle of 100 Tablets

0.75 mg tablets are supplied as a pale blue, flat tablet with beveled edges, scored on one side and product identification "54 960" debossed on the other side.

NDC 0054-8180-25: 10x10 Unit-Dose

NDC 0054-4180-25: Bottle of 100 Tablets

1 mg tablets are supplied as a yellow, flat tablet with beveled edges, scored on one side and product identification "54 489" debossed on the other side.

NDC 0054-8174-25: 10x10 Unit-Dose

NDC 0054-4181-25: Bottle of 100 Tablets

1.5 mg tablets are supplied as a pink, flat tablet with beveled edges, scored on one side and product identification "54 943" debossed on the other side.

NDC 0054-8181-25: 10x10 Unit-Dose

NDC 0054-4182-25: Bottle of 100 Tablets

2 mg tablets are supplied as a white, flat tablet with beveled edges, scored on one side and product identification "54 662" debossed on the other side.

NDC 0054-8176-25: 10x10 Unit-Dose

NDC 0054-4183-25: Bottle of 100 Tablets

4 mg tablets are supplied as a green, flat tablet with beveled edges, scored on one side and product identification "54 892" debossed on the other side.

NDC 0054-8175-25: 10x10 Unit-Dose

NDC 0054-4184-25: Bottle of 100 Tablets

6 mg tablets are supplied as an aqua, flat tablet with beveled edges, scored on one side and product identification "54 769" debossed on the other side.

NDC 0054-8183-25: 10x10 Unit-Dose

NDC 0054-4186-25: Bottle of 100 Tablets

Store and Dispense

Store at 20° to 25°C (68° to 77°F). [See USP Controlled Room Temperature.] Dispense in a tight light-resistant, child-resistant container as defined in the USP/NF.

Dexamethasone Oral Solution, USP

0.5 mg per 5 mL oral solution is supplied as a (cherry brandy flavored) clear colorless solution.

NDC 0054-3177-57: Bottle of 240 mL

NDC 0054-3177-63: Bottle of 500 mL

Store and Dispense

Store at 20° to 25°C (68° to 77°F). [See USP Controlled Room Temperature.] Dispense in a tight light-resistant, child-resistant container as defined in the USP/NF.

Dexamethasone Oral Solution, USP Intensol[™] (Concentrate)

1 mg per mL oral solution is supplied as a clear colorless solution.

NDC 0054-3176-44: Bottle of 30 mL with calibrated oral syringe [graduation of 0.25 mL (0.25 mg), 0.5 mL (0.5 mg), 0.75 mL (0.75 mg) and 1 mL (1 mg) on the oral syringe].

Store and Dispense

Store at 20° to 25°C (68° to 77°F). [See USP Controlled Room Temperature.] Do not freeze. Do not use if solution contains a precipitate. Dispense only in this bottle and only with the calibrated oral syringe provided. Discard opened bottle after 90 days.

Distributed by: **Hikma Pharmaceuticals USA Inc.** Berkeley Heights, NJ 07922

C5000comb/02/k01 Revised February 2024

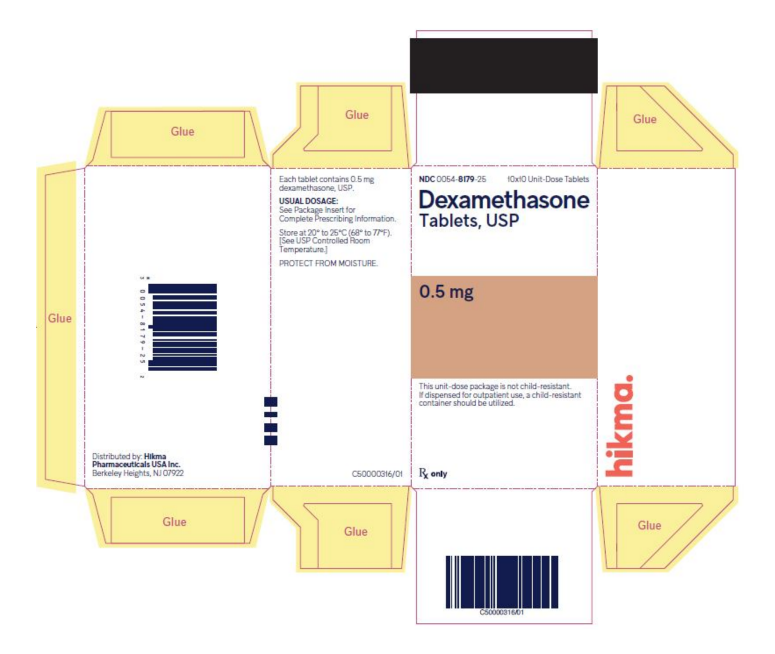
PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 0.5 mg Bottle Label

NDC 0054-4179-25 100 Tablets Dexamethasone Tablets, USP 0.5 mg



PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 0.5 mg Unit-Dose Carton

NDC 0054-8179-25 10x10 Unit-Dose Tablets Dexamethasone Tablets, USP 0.5 mg



PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 0.75 mg Bottle Label

NDC 0054-4180-25 100 Tablets Dexamethasone Tablets, USP 0.75 mg



PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 0.75 mg Unit-Dose Carton

NDC 0054-8180-25 10x10 Unit-Dose Tablets Dexamethasone Tablets, USP 0.75 mg



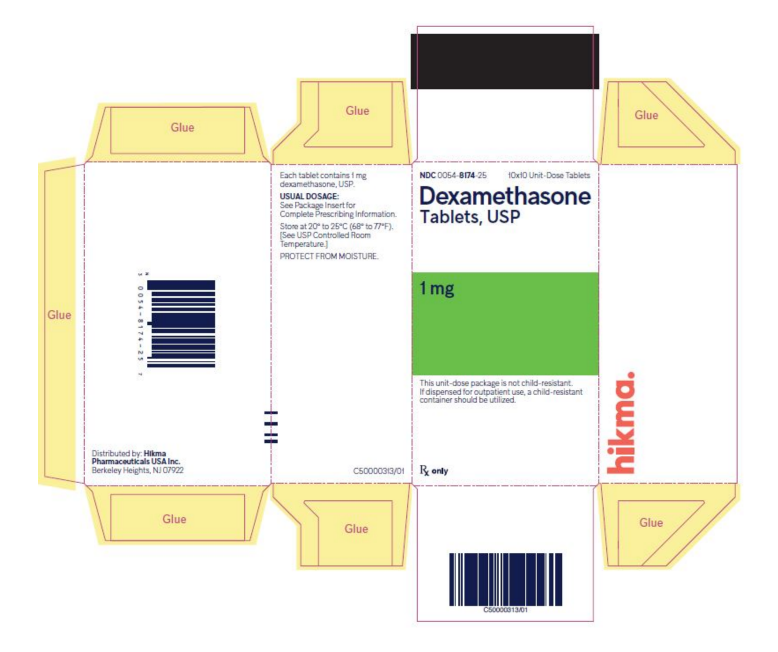
PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 1 mg Bottle Label

NDC 0054-4181-25 100 Tablets Dexamethasone Tablets, USP 1 mg



PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 1 mg Unit-Dose Carton

NDC 0054-8174-25 10x10 Unit-Dose Tablets Dexamethasone Tablets, USP 1 mg



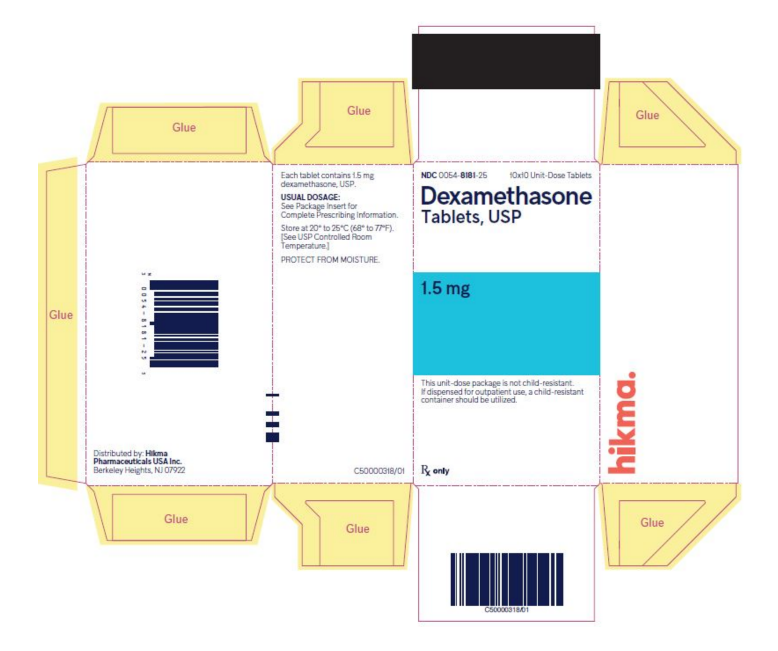
PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 1.5 mg Bottle Label

NDC 0054-4182-25 100 Tablets Dexamethasone Tablets, USP 1.5 mg



PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 1.5 mg Unit-Dose Carton

NDC 0054-8181-25 10x10 Unit-Dose Tablets Dexamethasone Tablets, USP 1.5 mg



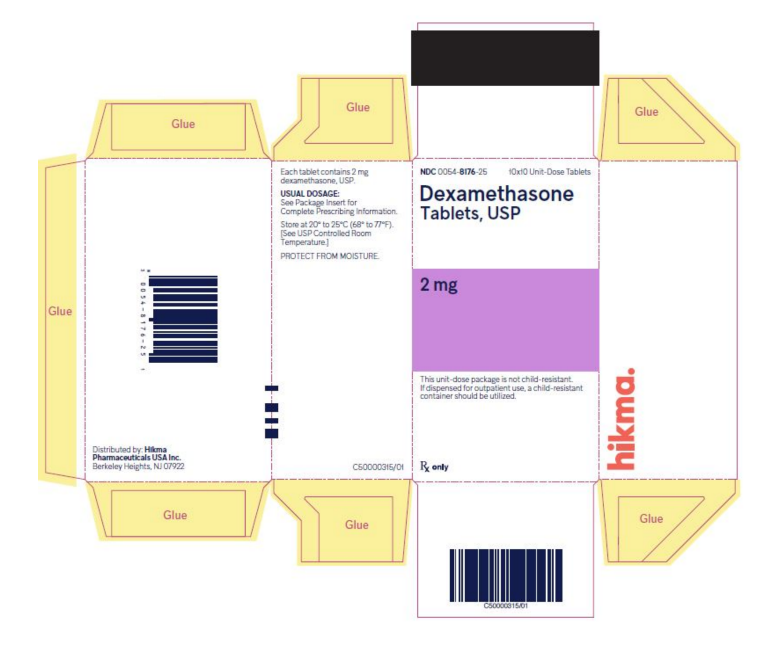
PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 2 mg Bottle Label

NDC 0054-4183-25 100 Tablets Dexamethasone Tablets, USP 2 mg



PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 2 mg Unit-Dose Carton

NDC 0054-8176-25 10x10 Unit-Dose Tablets Dexamethasone Tablets, USP 2 mg



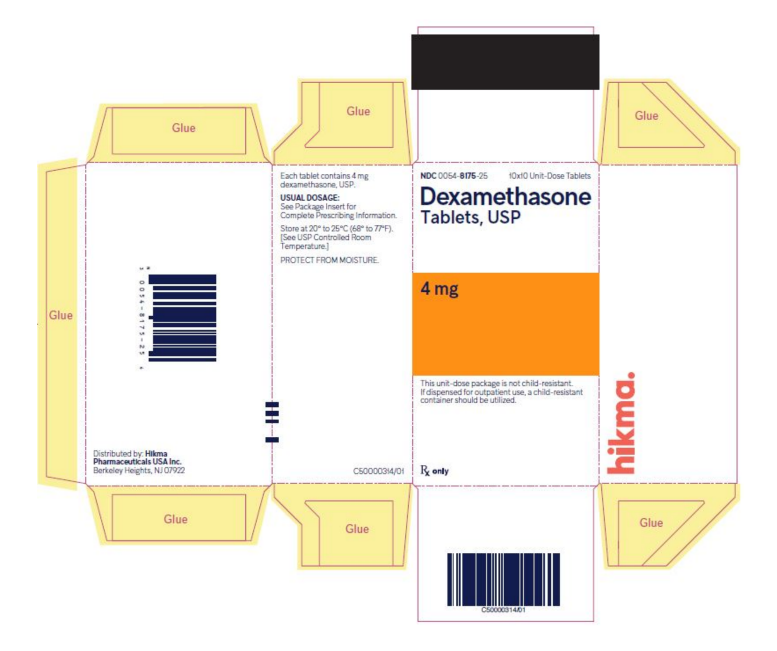
PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 4 mg Bottle Label

NDC 0054-4184-25 100 Tablets Dexamethasone Tablets, USP 4 mg



PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 4 mg Unit-Dose Carton

NDC 0054-8175-25 10x10 Unit-Dose Tablets Dexamethasone Tablets, USP 4 mg



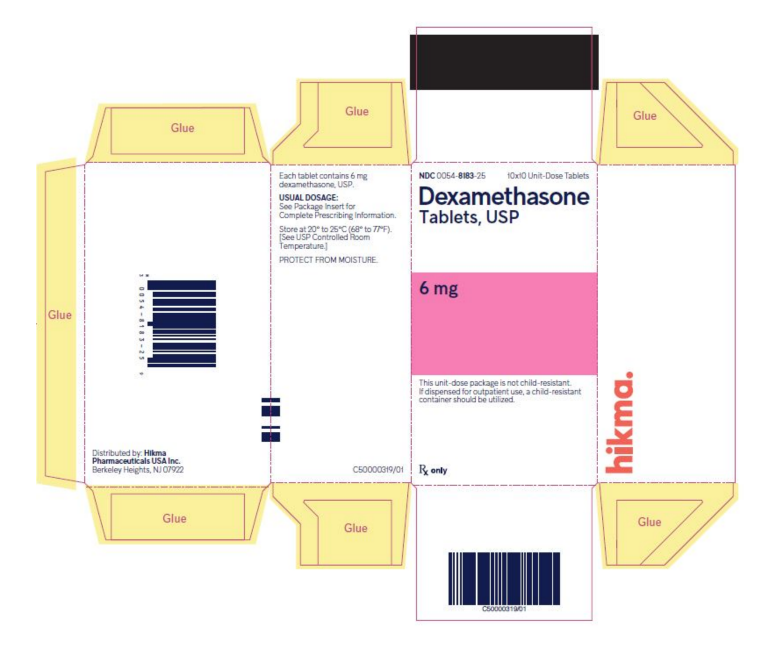
PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 6 mg Bottle Label

NDC 0054-4186-25 100 Tablets Dexamethasone Tablets, USP 6 mg



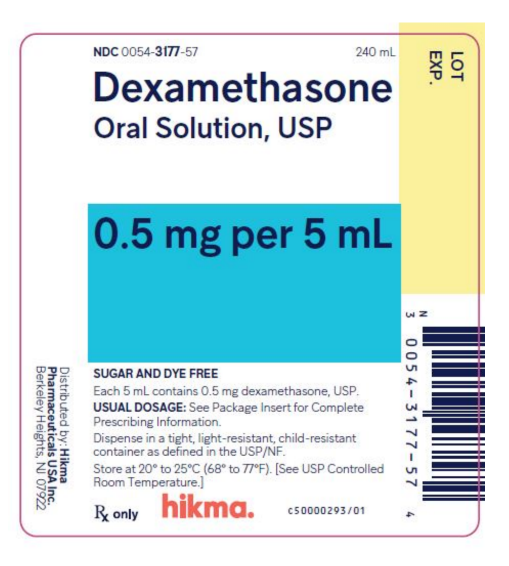
PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 6 mg Unit-Dose Carton

NDC 0054-8183-25 10x10 Unit-Dose Tablets Dexamethasone Tablets, USP 6 mg



PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 0.5 mg per 5 mL Oral Solution

NDC 0054-3177-57 240 mL Dexamethasone Oral Solution, USP 0.5 mg per 5 mL



PACKAGE/LABEL PRINCIPAL DISPLAY PANEL - 1 mg per mL Oral Solution Intensol[™] (Concentrate)

NDC 0054-3176-44 30 mL Dexamethasone Oral Solution, USP Intensol[™] (Concentrate) 1 mg per mL



DEXAMETH	SONE						
dexamethasone t	ablet						
Product Inform	nation						
Product Type		HUMAN PRESCRI	PTION DRUG	ltem Cod	le (Source)	NDC:0	054-4179
Route of Adminis	stration	ORAL					
Active Ingredi	ent/Active	Moiety					
Ingredient Name Basis of S						trength	Strengt
DEXAMETHASONE	(UNII: 7S5I7G3	JQL) (DEXAMETHA	SONE - UNII:7S5I7	G3JQL)	DEXAMETHAS	ONE	0.5 mg
Inactive Ingre	ulents	Ingredient N	lame			Strength	
mactive myre	ulents	In a ve die nt N				Chu	+ l a
D&C YELLOW NO.	10 (UNII: 355V	-					
FD&C YELLOW NO							
LACTOSE MONOH	(DRATE (UNII:	EWQ57Q8I5X)					
MAGNESIUM STEA	RATE (UNII: 70	097M6I30)					
STARCH, CORN (UN	III: 08232NY35	ij)					
SUCROSE (UNII: C1	51H8M554)						
Product Chara	cteristics						
	YEL	LOW	Score			2 pieces	
Color							
	ROL	IND	Size			6mm	
Shape		IND	Size Imprint Code			6mm 54;299	
Shape		IND					
Flavor		IND					
Shape Flavor		IND					

	1						
1 NDC:0054- 4179-25	100 in 1 BOTTL Combination Pr	E, PLASTIC; Type 0: Not a oduct		07/25/19	975		
Maxication	Informat	1					
Marketing							
Marketing Category	Applica	tion Number or Monog Citation	graph		eting Start Date		eting End Date
ANDA	ANDA08461	1		07/25/19	75		
DEXAMETH	ASONE						
dexamethasone							
Product Info	rmation						
Product Type		HUMAN PRESCRIPTION DRI	UG	ltem Cod	le (Source)	NDC:0	054-4180
Route of Admin	istration	ORAL					
Active Ingred	iont/Activo	Majaty					
Active Ingred		•			Pacia of St	ronath	Strongt
DEVAMETUACON	-	edient Name JQL) (DEXAMETHASONE - UN			Basis of St DEXAMETHASO	-	
DEXAMETRASUN		JQL) (DEXAMETHASONE - OF	11.735170	33JQL)	DEAMETHASO		0.75 mg
Inactive Ingre	edients						
		Ingredient Name				Str	ength
FD&C BLUE NO.							
LACTOSE MONOH							
MAGNESIUM STE							
STARCH, CORN (L		J)					
SUCROSE (UNII: C	151H8M554)						
Product Char	acteristics						
Color	BLUE (Pa	ale)	Score			2 piec	es
Shape	ROUND (ROUND)	Size			6mm	
Flavor			Imprin	t Code		54;960)
Contains							
Packaging							
# Item Code	Pa	ackage Description			eting Start Date		eting End Date
1 NDC:0054-		E, PLASTIC; Type 0: Not a		06/03/19		L	Jale
4180-25	Combination Pr	oduct					
Marketing	Informat	ion					
marketing	mormal						

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
ANDA	ANDA084613	06/03/1975			
DEXAMETHASONE					
dexamethasone ta	ablet				

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	ltem Code (Source)	NDC:0054-4181
Route of Administration	ORAL		

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
DEXAMETHASONE (UNII: 7S5I7G3JQL) (DEXAMETHASONE - UNII:7S5I7G3JQL)	DEXAMETHASONE	1 mg

Inactive Ingredients					
Ingredient Name	Strength				
FERRIC OXIDE YELLOW (UNII: EX43802MRT)					
LACTOSE MONOHYDRATE (UNII: EWQ57Q8I5X)					
MAGNESIUM STEARATE (UNII: 70097M6I30)					
STARCH, CORN (UNII: 08232NY3SJ)					
SUCROSE (UNII: C151H8M554)					

Product Characteristics					
Color	YELLOW	Score	2 pieces		
Shape	ROUND	Size	6mm		
Flavor		Imprint Code	54;489		
Contains					

P	ackaging			
#	ltem Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0054- 4181-25	100 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	09/15/1983	

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
ANDA	ANDA088306	09/15/1983			

DEXAMETH dexamethasone								
Product Info	rmation							
Product Type		HUMAN PRESC	RIPTION DRUG	ltem Cod	le (Source)	e) NDC:0054-4182		
Route of Admir	nistration	ORAL						
Active Ingred	lient/Active	Moiety						
	Ingr	edient Name	2		Basis of St	trength	Strength	
DEXAMETHASON	E (UNII: 7S5I7G3	JQL) (DEXAMETH	ASONE - UNII:7S5I7	G3JQL)	DEXAMETHAS	ONE	1.5 mg	
Inactive Ingr	edients							
		Ingredient	Name			Str	ength	
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)								
FD&C RED NO. 3								
FD&C RED NO. 4								
LACTOSE MONO								
MAGNESIUM STE STARCH, CORN (I								
SUCROSE (UNII: C		J <i>1</i>						
Product Chai	acteristics							
Color	PIN		Score			2 pieces		
Shape	ROU	JND			6mm			
Flavor			Imprint Code		54;943			
Contains								
Packaging								
# Item Code	Pa	ackage Desc	ription	Mark	eting Start Date		eting End Date	
1 NDC:0054- 4182-25	Combination Pr			05/19/19	975			
2 NDC:0054- 4182-31	1000 in 1 BOTT Combination Pr	LE, PLASTIC; Ty oduct	pe 0: Not a	07/17/20	006	06/18/20	18	
Marketing	Informat	ion						
Marketing Category	Applica	tion Number Citatio	or Monograph n		ting Start Date		eting End Date	
		-						
ANDA	ANDA08461	0		05/19/19	/5			

DEXAMETHASONE

Product Info	rmation							
Product Type		HUMAN PRESCR	IPTION DRUG	ltem Cod	le (Source)	NDC:0054-4183		
Route of Admir	nistration	ORAL						
Active Ingred	lient/Active	Moiety						
	Ingi	redient Name			Basis of S	trength	Strengt	
DEXAMETHASON	E (UNII: 7S5I7G3	BJQL) (DEXAMETHA	SONE - UNII:7S5I7	G3JQL)	DEXAMETHAS	ONE	2 mg	
nactive Ingr	edients							
		Ingredient I	Name			Stre	ength	
LACTOSE MONOI								
MAGNESIUM STE								
STARCH, CORN (USUCROSE (UNII: C		oJ)						
	·							
Product Char	acteristics							
Color		ITE	Score			2 pieces		
Shape 	RO	UND	Size			6mm		
Flavor			Imprint Code			54;662		
Contains								
Packaging				Marke	eting Start	Marke	ting End	
	P	ackage Descr	iption		Date	C	Date	
# Item Code		LE, PLASTIC; Type			Date		Date	
<pre># Item Code NDC:0054-</pre>	100 in 1 BOTT	LE, PLASTIC; Type			Date		Date	
NDC:0054-	100 in 1 BOTTI Combination P	LE, PLASTIC; Type roduct			Date		Date	
 item Code NDC:0054- 4183-25 	100 in 1 BOTTI Combination P	LE, PLASTIC; Type roduct	e 0: Not a	08/26/19	Date	Marke	Date eting End Date	

dexamethasone tablet

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0054-4184
Route of Administration	ORAL		

	ive Ingred	lient/Act	ive Moiety	у					
		lı	ngredient	Name			Basis of St	trength	Strength
DEX	AMETHASONI	e (UNII: 755	I7G3JQL) (DE>	KAMETHASON	E - UNII:7S5I7G	i3JQL)	DEXAMETHAS	ONE	4 mg
Ina	ctive Ingre	edients							
			Ingre	dient Nam	e			Stre	ength
D&C	YELLOW NO	. 10 (UNII: 3	35SW5USQ3G	i)					
FD&	C GREEN NO	. 3 (UNII: 3P	30NR601S)						
LAC.	TOSE MONOF	HYDRATE (U	JNII: EWQ57Q	8I5X)					
MAG	SNESIUM STE	ARATE (UNI	I: 70097M6I30)					
	RCH, CORN (L		-						
SUC	ROSE (UNII: C	151H8M554)						
_	oduct Char	acteristi							
Cold	or		GREEN	Sco	re		:	2 pieces	
		uccense		Sco Size	-			2 pieces 6mm	
Sha	pe		GREEN	Size	-				
Sha Fla v	pe		GREEN	Size	•			6mm	
Sha Fla v	ipe vor		GREEN	Size	•			6mm	
Sha Flav Con	ipe vor		GREEN	Size	•			6mm	
Sha Flav Con Pac	ipe /or itains		GREEN ROUND	Size	rint Code			6mm 54;892 Marke	eting End Date
Sha Flav Con Pac # I	ope vor otains ckaging	100 in 1 BC	GREEN ROUND	Size	rint Code		eting Start Date	6mm 54;892 Marke	
Sha Flav Con Pac # I	ope vor stains ckaging Item Code DC:0054-	100 in 1 BC	GREEN ROUND Package	Descriptio	rint Code		eting Start Date	6mm 54;892 Marke	
Sha Flav Con Pac # I 1 N 4	ope vor stains ckaging Item Code DC:0054-	100 in 1 BC Combinatio	GREEN ROUND Package	Descriptio	rint Code		eting Start Date	6mm 54;892 Marke	
Pac # 1 1 ^N 4 Ma	kaging tem Code DC:0054- 184-25	100 in 1 BC Combinatio	GREEN ROUND Package OTTLE, PLAST on Product	Descriptio	rint Code on ot a	07/19/19	eting Start Date	6mm 54;892 Marke Marke	

DEXAMETHASONE			
dexamethasone tablet			
Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0054-4186
Route of Administration	ORAL		
Active Ingredient/Active	Moiety		

		Ingr	edient Name			Basis of St	-	Strength
DE	XAMETHASON	E (UNII: 75517G3	BJQL) (DEXAMETHASONE - UNII:	7S5I7G3	BJQL)	DEXAMETHASO	NE	6 mg
n	active Ingro	edients						
			Ingredient Name				St	rength
	&C GREEN NO							
	CTOSE MONOR							
	AGNESIUM STE							
	ARCH, CORN (L CROSE (UNII: C		oJ)					
30		1311000334)						
Pı	oduct Char	acteristics						
	lor	TURQUOIS	SE (Aqua)	Score			2 pie	res
	ape	ROUND	JE (Aquu)	Size			2 pie	
	avor	NOUND		Imprin	nt Cod		54;7	•
	ontains			mpm			54,7	
	intamis							
Pa	ackaging							
		_			Mar	keting Start	Mark	eting End
#	ltem Code	Pa	ackage Description		man	Date		Date
	NDC:0054- 4186-25	100 in 1 BOTTL Combination Pr	E, PLASTIC; Type 0: Not a		09/15/	1983		
Μ	arketing	Informat	ion					
	Marketing Category	Applica	tion Number or Monogra Citation	ph	Marl	keting Start Date		eting End Date
AN	DA	ANDA08831	.6		09/15/1	.983		
		ACONE						
_	EXAMETH							
le	xamethasone	intensol soll	ition, concentrate					
Ρ	roduct Info	rmation						
Pr	oduct Type		HUMAN PRESCRIPTION DRUG	It	em Co	ode (Source)	NDC	0054-3176
	oute of Admin	istration	ORAL		em et		n b c	00010170
n		istration						
A	ctive Ingred	ient/Active	Moietv					
			edient Name			Basis of Stre	nath	Strength
DF	XAMETHASON	•	BJQL) (DEXAMETHASONE - UNII:	7551763		DEXAMETHASON	-	1 mg in 1 m
~ -		_ (0 / 00// 00			, ~ - /		_	
	active Ingro	adionts						

		Ingredient Name			Strength
AL	COHOL (UNII: 3	3K9958V90M)			
BE	NZOIC ACID (L	JNII: 85KN0B0MIM)			
CI	TRIC ACID MO	NOHYDRATE (UNII: 2968PHW8QP)			
ED	ETATE DISOD	IUM (UNII: 7FLD91C86K)			
PR	OPYLENE GLY	COL (UNII: 6DC9Q167V3)			
w	ATER (UNII: 059	QF0KO0R)			
Pa	ackaging				
	ackaging Item Code	Package Description		Marketing Start Date	Marketing End Date
#		Package Description 30 mL in 1 BOTTLE, GLASS; Type 1: Convenience of Co-Package	Kit	-	-
#	Item Code NDC:0054-	30 mL in 1 BOTTLE, GLASS; Type 1: Convenience	Kit	Date	-
#	Item Code NDC:0054-	30 mL in 1 BOTTLE, GLASS; Type 1: Convenience	Kit	Date	-
# 1	Item Code NDC:0054- 3176-44	30 mL in 1 BOTTLE, GLASS; Type 1: Convenience	Kit	Date	-
#	Item Code NDC:0054- 3176-44	30 mL in 1 BOTTLE, GLASS; Type 1: Convenience of Co-Package	Kit	Date	-

DEXAMETHASONE				
dexamethasone solution				
Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	ltem C	Code (Source)	NDC:0054-3177
Route of Administration	ORAL			
Active Ingredient/Active	Moiety			
Ingree	dient Name		Basis of Strength	Strength
DEXAMETHASONE (UNII: 75517G3)	QL) (DEXAMETHASONE - UNII:7S5I7	G3JQL)	DEXAMETHASONE	0.5 mg in 5 mL
Inactive Ingredients				
	Ingredient Name			Strength
CITRIC ACID MONOHYDRATE (UN	III: 2968PHW8QP)			
EDETATE DISODIUM (UNII: 7FLD9)	1С86К)			
CHERRY (UNII: BUC5I9595W)				
GLYCERIN (UNII: PDC6A3C0OX)				
METHYLPARABEN (UNII: A218C7HI	•			
PROPYLENE GLYCOL (UNII: 6DC90				
PROPYLPARABEN (UNII: Z8IX2SC1	.OH)			
WATER (UNII: 059QF0KO0R)				
SORBITOL (UNII: 506T60A25R)				

Packaging							
# Item Coc	le	Package	Description	Mar	keting Start Date		eting Enc Date
1 NDC:0054- 3177-57		1 BOTTLE, PLA ion Product	ASTIC; Type 0: Not a	11/01/	2007		
2 NDC:0054- 3177-63		1 BOTTLE, PLA ion Product	ASTIC; Type 0: Not a	09/01/	1983		
Marketiı	ng Infor	mation					
Marketi Catego			mber or Monograp itation	h Mark	eting Start Date		ting End ate
ANDA	ANDA	088248		09/01/19	983		
		F					
DEXAME lexamethas							
Product Ir	oformation	า					
Product Typ		HUMAN	PRESCRIPTION DRUG	ltem Co	de (Source)	NDC:0	054-8179
	ministratio					noord	0010175
Active Ing	redient/Ac	tive Moiet	V				
		Ingredient	Name		Basis of St	rength	Strengt
DEXAMETHAS	ONE (UNII: 75	517G3JQL) (DE)	AMETHASONE - UNII:7S	5I7G3JQL)	DEXAMETHASO	NE	0.5 mg
Inactive In	igredients						
		-	dient Name			Stre	ength
		: 35SW5USQ3G)				
FD&C YELLO	-						
		(UNII: EWQ57Q					
STARCH, COR		NII: 70097M6I3())				
SUCROSE (UN							
		, , ,					
	haracteris	tics					
Product C		VELLOW!	Score		2	2 pieces	
Color		YELLOW	Beere				
Color Shape		ROUND	Size			Smm	
Color Shape Flavor				le		5mm 54;299	
Product C l Color Shape Flavor Contains			Size	le			
Color Shape Flavor			Size	le			

# Item Code	Package Description	Marketing Start Date	Marketing End Date
1 NDC:0054- 8179-25	10 in 1 CARTON	07/25/1975	
1	10 in 1 BLISTER PACK; Type 0: Not a Combination Product		
Marketing	Information		
Marketing Marketing Category	Information Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
Marketing	Application Number or Monograph	-	-
Marketing Category	Application Number or Monograph Citation	Date	-

D	EXAMETH	ASONE						
de	xamethasone	tablet						
Ρ	roduct Infor	mation						
Р	roduct Type		HUMAN PRESCRIPTION DRU	JG	ltem Cod	e (Source)	NDC	:0054-8180
Re	oute of Admin	istration	ORAL					
A	ctive Ingred	ient/Active	Moiety					
		Ingr	edient Name			Basis of S	trengtł	n Strength
DE	XAMETHASONE	(UNII: 7S5I7G3	JQL) (DEXAMETHASONE - UN	III:7S5I7	G3JQL)	DEXAMETHAS	ONE	0.75 mg
In	active Ingre	edients						
			Ingredient Name				St	rength
FD	&C BLUE NO. 1	. (UNII: H3R47K3	TBD)					
LA	стоѕе монон	YDRATE (UNII:	EWQ57Q8I5X)					
M	AGNESIUM STEA	RATE (UNII: 70	097M6I30)					
	ARCH, CORN (U		J)					
รบ	JCROSE (UNII: C1	L51H8M554)						
P	roduct Chara	acteristics						
Сс	olor	BLUE (Pa	le)	Score			2 pie	ces
Sł	nape	ROUND (ROUND)	Size			6mm	1
Fla	avor			Imprin	nt Code		54;9	60
Co	ontains							
Pa	ackaging							
#	ltem Code	Ра	ckage Description			ting Start Date	Mark	eting End Date
1	NDC:0054- 8180-25	10 in 1 CARTON	J		06/03/197	5		

1	10 in 1 BLI Product	STER PACK; Ty	pe 0: Not a Combinatior				
Marketing	Inform	ation					
Marketing		lication Num	ber or Monograph		eting Start		ting End
Category ANDA	ANDA08		ation	06/03/19	Date 75	L	Date
						1	
DEXAMETH	ASONF	:					
lexamethasone		•					
Product Infor	mation						
Product Type		HUMAN P	RESCRIPTION DRUG	Item Cod	le (Source)	NDC:0	054-8174
Route of Admini	istration	ORAL					
Active Ingredi	ient/Acti	ive Moiety					
	lı	ngredient N	ame		Basis of St	rength	Strengt
DEXAMETHASONE	(UNII: 7S5	I7G3JQL) (DEXA	METHASONE - UNII:7S5I	7G3JQL)	DEXAMETHAS	DNE	1 mg
nactive Ingre		-	ent Name			Stro	ength
LACTOSE MONOH MAGNESIUM STEA			DA)				
STARCH, CORN (UI		•					
SUCROSE (UNII: C1							
Product Chara	acteristi	cs					
Color		YELLOW	Score			2 pieces	
Shape		ROUND	Size			6mm	
Flavor			Imprint Code			54;489	
Contains							
Packaging				Marke	ting Start		
		Package D	escription		Date		ting End ate
# Item Code	10 in 1 CA	-	escription		Date		
1 NDC:0054- 8174-25		N	pescription	C 09/15/198	Date		

Marketing Category	Applic	ation Number Citatio	or Monograph n		eting Start Date		eting End Date
ANDA	ANDA0883	06		09/15/19	83		
DEXAMETH	ASONE						
lexamethasone	tablet						
Product Info	mation						
Product Type		HUMAN PRESC	RIPTION DRUG	ltem Coo	le (Source)	NDC:0	054-8181
Route of Admin	istration	ORAL					
Active Ingred	ient/Active	e Moiety					
	Ing	redient Name	9		Basis of S	trength	Strengt
DEXAMETHASONI	(UNII: 7S5I7G	3JQL) (DEXAMETH	ASONE - UNII:7S5I7	G3JQL)	DEXAMETHAS	ONE	1.5 mg
Inactive Ingre	edients						
		Ingredient	Name			Stre	ength
FD&C BLUE NO. 1							
FD&C RED NO. 3							
FD&C RED NO. 40 LACTOSE MONOF							
MAGNESIUM STE							
STARCH, CORN (L	-						
SUCROSE (UNII: C		-					
Product Char	acteristics	5					
Color	PI	NK	Score			2 pieces	
Shape	RC	DUND	Size			6mm	
Flavor			Imprint Code			54;943	
Contains							
Packaging							
# Item Code	Р	ackage Desc	ription		ting Start Date		ting End ate
1 NDC:0054- 8181-25	10 in 1 CARTO	N		05/19/197	5		
1	10 in 1 BLIST Product	ER PACK; Type 0:	Not a Combination				

ANDA ANDA084610	05/19/1975	

DEXAMETH dexamethasone								
Product Infor	rmation							
Product Type		HUMAN PRESCR	IPTION DRUG	ltem Cod	le (Source)	NDC:0	0054-8176	
Route of Admin	istration	ORAL						
Active Ingred	ient/Active	Moiety						
	-	redient Name				-	Strength	
DEXAMETHASONE	(UNII: 7S5I7G	3JQL) (DEXAMETHA	SONE - UNII:7S5I7	G3JQL)	DEXAMETHAS	ONE	2 mg	
Inactive Ingre	edients							
, i i i i i i i i i i i i i i i i i i i		Ingredient I	Name			Strength		
LACTOSE MONOH	IYDRATE (UNII:	-				ou ong u		
MAGNESIUM STEA	ARATE (UNII: 70	0097M6I30)						
STARCH, CORN (U	INII: 08232NY3	SJ)						
SUCROSE (UNII: C	151H8M554)							
Product Characteristics								
Color	WH	IITE Score 2 pie					pieces	
Shape		UND Size			6mm			
Flavor			Imprint Code			54;662		
Contains								
Packaging								
# Item Code	Pa	Package Description		Marketing Start Date		Marketing End Date		
1 NDC:0054-	10 in 1 CARTO	N		08/26/198				
▲ 8176-25	10 in 1 BLISTER PACK; Type 0: Not a Combination							
1 Product								
Marketing Information								
Marketing Category		ation Number o Citation			ting Start Date		eting End Date	
ANDA	ANDA08791	16		08/26/19	82			

dexamethaso	ne tablet						
Product Inf	ormation						
Product Type	•	HUMAN PRE	SCRIPTION DRUG	ltem Cod	le (Source)	NDC:0	0054-8175
Route of Adn	ninistration	ORAL					
Active Ingre	edient/Activ	e Moiety					
	Ing	gredient Na	me		Basis of S	trength	Strengt
DEXAMETHASO	NE (UNII: 75517	G3JQL) (DEXAME	THASONE - UNII:7S5I7	G3JQL)	DEXAMETHAS	ONE	4 mg
nactive Ing	gredients						
		Ingredie	nt Name			Str	ength
	NO. 10 (UNII: 35						
	IO. 3 (UNII: 3P3)						
	OHYDRATE (UN)				
	TEARATE (UNII:						
	I (UNII: 08232NY	'3SJ)					
SUCROSE (UNII							
FD&C YELLOW	NO. 6 (UNII: H7	7VEI93A8)					
Product Ch	aracteristic	S					
						2 pieces	
Shape ROUND Size					6mm		
Flavor			Imprint Code			54;892	
Contains			•				
Packaging							
# Item Cod	e I	Package Description Marketing Start Date		-	Marketing End Date		
1 NDC:0054- 8175-25	10 in 1 CAR	10 in 1 CARTON		07/19/1978			
1	10 in 1 BLIS Product	10 in 1 BLISTER PACK; Type 0: Not a Combination Product					
Marketin	g Informa	ation					
		cation Numb Citat	er or Monograph		eting Start Date		eting End Date
Marketin Category	/	Citai					Juic
	ANDA084			07/19/19			Jute

DEXAMETHASONE

	xamethasone	tablet							
P۱	roduct Infor	mation							
Pr	oduct Type		HUMAN PRESCRIPTION DRUG	G	ltem Cod	e (Source)	NDC:0	054-8183	
	oute of Admin	istration	ORAL			- (,			
40	tive Ingred	ient/Active	Moiety						
		Ingre	edient Name			Basis of St	trength	Strengt	
DE	XAMETHASONE	: (UNII: 7S5I7G3)	QL) (DEXAMETHASONE - UNII	1:75517	G3JQL)	DEXAMETHAS	ONE	6 mg	
n	active Ingre	edients							
			Ingredient Name				Strength		
	&C GREEN NO.								
	стоѕе монон								
	GNESIUM STEA								
	ARCH, CORN (U))						
50	CROSE (UNII: C	L51H8M554)							
Pr	oduct Char	acteristics							
Co	lor	TURQUOISE (Aqua) S			re		2 piec	ces	
Shape ROUND			Size	•		6mm			
Flavor				Imprint Code			54;76	9	
Co	ntains								
_									
Pa	ackaging					_			
#	ltem Code	Ра	ckage Description			ting Start Date		ting End Date	
	NDC:0054- 8183-25	10 in 1 CARTON	1		09/15/198	3			
L		10 in 1 BLISTER Product	R PACK; Type 0: Not a Combi	nation					
			• • • • • • • • • • • • • • • • • • •						
Μ	arketing	Informat	ion						
M	arketing Marketing Category		ION tion Number or Monogr Citation	aph		ting Start Date		eting End Date	

Labeler - Hikma Pharmaceuticals USA, Inc. (080189610)

Establ	ishmen	t	
Name	Address	ID/FEI	Business Operations

West-Ward Columbus Inc.

Revised: 2/2024

Hikma Pharmaceuticals USA, Inc.