KETOROLAC TROMETHAMINE TABLETS USP, 10 mg

0314

Rx only

WARNING

Ketorolac tromethamine tablets, a nonsteroidal anti-inflammatory drug (NSAID), are indicated for the short-term (up to 5 days in adults), management of moderately severe acute pain that requires analgesia at the opioid level and only as continuation treatment following IV or IM dosing of ketorolac tromethamine, if necessary. The total combined duration of use of ketorolac tromethamine tablets and ketorolac tromethamine tablets and ketorolac tromethamine tablets.

Ketorolac tromethamine tablets are not indicated for use in pediatric ketorolac tromethamine tablets are not indicated for use in pediatric patients and they are NOT indicated for minor or chronic painful conditions. Increasing the dose of ketorolac tromethamine tablets beyond a dally maximum of 40 mg in adults will not provide better efficacy but will increase the risk of developing serious adverse even

AGSTROINTESTINAL RISK

• Ketorolac tromethanine, including ketorolac tromethanine tablets can cause peptic ulers, gastrointestinal bleeding and/or perforation of the stomach or intestines, which can be fatal. These events can occur at any time during use and without warming symptoms. Therefore, ketorolac tromethanine is CONITRAINDICATED in patients with active peptic ulere disease, in patients with a history of peptic user disease or gastrointestinal beleding. Existery patients are at greater risk for serious gastrointestinal events (see WARNINGS).

- CARDIOVASCULAR THOMBOTIC EVENTS

 Nonsteroidal anti-riflammatory drugs (NSAIDs) cause an increased risk of serious cardiovascular thrombotic events, including myocardial infarction and stroke, which can be fatal. This risk may occur early in treatment and may increase with duration of use (see WARNINGS and PRECAUTION).

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RENAL RISK • Ketorolac t

ENAL NISK

Ketorolac tromethamine is CONTRAINDICATED in patients with advanced renal impairment and in patients at risk for renal failure due to volume depletion (see WARNINGS).

RISK OF BLEEDING

SK OF BLEEDING
Ketorolac tromethamine inhibits platelet function and is, therefore,
CONTRAINDICATED in patients with suspected or confirmed cerebrovascular
bleeding, patients with hemorrhagic idathesis, incomplete hemostasis and those
at high risk of bleeding (see WARNINGS and PRECAUTIONS).

Ketorolac tromethamine is CONTRAINDICATED as prophylactic analgesic before any major surgery.

RISK DURING LABOR AND DELIVERY

The use of ketorolac tromethamine in labor and delivery is contraindicated because it may adversely affect fetal circulation and inhibit uterine contractions

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SPECIAL POPULATIONS

• Dosage should be adjusted for patients 65 years or older, for patients under it y(1010 ib) of body weight (see DOSAGE AND ADMINISTRATION) and for patients with moderately elevated serum creatinine (see WARNINGS).

DESCRIPTION

Ketorola: Tromethamine Tablets USP are a member of the pyrrols-pyrrole group of nonsterodial anti-inflammatory drugs (INSAIDs). The chemical name for ketorolac tromethamine. [95 Fs (±-5-bersoy2-3-dilydro-1-4-pyrrolizen-1-carboy/sc act, compound with 2-amino-2-(hydroxymethyl)-1,3-propanediol (1:1). The structural formula 6:

C ₁₅H ₁₃NO ₃·C ₄H ₁₁NO ₃ M.W. 376.40

Ketorolac tromethamine, USP is a racemic mixture of [-]S and [+]R ketorolac tromethamine, USP. Ketorolac tromethamine, USP may exist in three crystal forms. All forms are equally soluble in water. Ketorolac tromethamine, USP has a pKa of 3.5 and an n-octanol/water partition coefficient of 0.26.

Ketorolac Tromethamine Tablets USP are white, round, convex, unscored, film-coated tablets. Each tablet, for oral administration, contains 10 mg leetorolac tromethamine, USP, the active ingredient in addition, each tablet contains the following inactive ingredients: hydroxypropyl cellulose, hypromelose, lactose monohydrate, magnesium stearate, microrostation cellulose, polythyline gloco, and titanium dioxide.

Pharmacodynamics

Final maccoynamics Ketoroisc tromethamine is a nonsteroidal anti-inflammatory drug (NSAID) that exhibits analgesis activity in animal models. The mechanism of action of ketoroisc, like that of other NSAIDs, is not completely understood but may be related to prostaginadin synthetise inhibition. The biological activity of ketoroisc tromethamine is associated with the Sform. Ketoroisc tromethamine possesses no sociative or anothery properties. The peak analgesis effect of retoroisc tromethamine occurs within 2 to 3 hours and is not statistically soprificantly different over the recommended dosage range of ketoroisc tromethamine is in the duration of analgesia.

Ketorolac tromethamine is a racemic mixture of [-]S- and [+]R-enantiomeric forms, with the S-form having analgesic activity.

Comparison of IV, IM and Oral Pharmacokinetics

The pharmacoknetics of ketorolac tromethamine, following IV and IM doses of ketorolac tromethamine and oral doses of ketorolac tromethamine, are compared in **Table** 1. In adults, the extent of bioavaliability following administration of the ORAL form of ketorolac tromethamine and the IM form of ketorolac tromethamine was equal to that following an IV bolus.

Linear Kinetics

Linear Knetics In adults, flollwing administration of single ORAL doses of ketorolac tromethamine or IM or IV doses of ketorolac tromethamine in the recommended dosage ranges, the clearance of the racemate does not change. This implies that the pharmacokinetics of ketorolac tromethamine in adults, following single or multiple IM or IV doses of ketorolac tromethamine or recommended ord doses of ketorolac tromethamine in a relinear. At the higher recommended doses, there is a proportional increase in the concentrations of free and bound racemate.

Absorption

Kotorolac tromethamine is 100% absorbed after oral administration (see **Table 1**). Oral administration of ketorolac tromethamine after a high-fat meal resulted in decreased peak and delayed time-to-peak concentrations of ketorolac tromethamine by about 1 hour. Antacids did not affect the extent of absorption.

Distribution

Distribution

The mean apparent volume ($V_{\rm a}$) of ketorolac tromethamine following complete distribution was approximately 13 iters. This parameter was determined from single-dose data. The ketorolac tromethamine racemate has been shown to be highly protein bound (99%). Nevertheless, plasma concentrations as high as 10 mcg/mL will only occupy approximately 5% of the abumin binding sites. Thus, the unbound fraction for each enanthomer will be constant over the therapeutic range. A decrease in serum abumin, however, will result in increased free drug concentrations.

Ketorolac tromethamine is excreted in human milk (see **PRECAUTIONS, Nursing Mothers**).

Metabolism

Retorolac tromethamine is largely metabolized in the liver. The metabolic products are hydroxylated and conjugated forms of the parent drug. The products of metabolism, and some unchanged drug, are excreted in the urine.

The principal route of elimination of ketorolac and its metabolites is renal. About 92% of a given dose is found in the urrier, approximately 40% as metabolites and 60% as of a given dose is found in the urrier, approximately 40% as metabolites and 60% as of the principal dose study with 10 mg ketorolac tromethamine (n = 9) demonstrated that the Sienatose study with 10 mg ketorolac tromethamine (n = 9) demonstrated that the Sienatose is classed approximately two times faster than the R-enantiomer and that the clearance was independent of the route of administration. This means that the ratio of SiR plasma concentrations decreases with time after each dose. There is title or no inversion of the R- to S- form in humans. The clearance of the racermate in normal of the Table 2 (see CLINICAL PHARMACOLOGY. Kinetics in Special Populations).

The half-life of the ketorolac tromethamine S-enantiomer was approximately 2.5 hours ($5D \pm 0.4$) compared with 5 hours ($5D \pm 1.7$) for the R-enantiomer. In other studies, the half-life for the racemate has been reported to lie within the range of 5 to 6 hours

Ketorolac tromethamine administered as an IV bolus every 6 hours for 5 days to healthy

subjects (n = 13), showed no significant difference in C $_{max}$ on Day 1 and Day 5. Trough levels averaged 0.29 mcg/mL (SD \pm 0.13) on Day 1 and 0.55 mcg/mL (SD \pm 0.23) on Day 6. Steady state was approached after the fourth dose.

Accumulation of ketorolac tromethamine has not been studied in special populations (geriatric, pediatric, renal failure or hepatic disease patients).

Kinetics in Special Populations

Goriatric Patients

Based on single-dose data only, the half-life of the ketorolac tromethamine racemate increased from 5 to 7 hours in the edety (56 to 78 years) compared with young healthy volunteers (24 to 35 years) (see Table 2). There was little difference in the C max for the two groups (edety), 2.52 mcg/ml. ± 0.77; young, 2.99 mcg/ml. ± 1.03) (see PRECAUTIONS, Gerfatric Use (c. 65 Years of Ago).

Pediatric Patients

Pedditr.' Patients.

Limited Information is available regarding the pharmacokinetics of dosing of ketorolac tromethamine in the pedditric population. Following a single intravenous bolus dose of 0.5 mg/kg in 10 children 4 to 8 years old, the half few as 5.8 ± 1.6 hours, the average clearance was 0.042 ± 0.01 L/hr/kg, the volume of distribution during the terminal phase (V_g) was 0.34 ± 0.01 L/kg and the volume of distribution at steady state (vs)s was 0.26 ± 0.08 L/kg. The volume of distribution at steady state (vs)s was 0.26 ± 0.08 L/kg. The volume of distribution at steady state (vs)s was 0.26 ± 0.08 L/kg. The volume of distribution and clearance of ketactock in pedditric patients was higher than those observed in adult subjects (see Table 1.1) There are no pharmacochretic data vasible for administration of ketorolac tromethamine by the IM roude in pedditric patients.

Renal Insufficiency

Based on single-dose data only, the mean half-life of ketorolac tromethamine in renally impaired patients is between 6 and 19 hours and is dependent on the extent of the impairment. There is poor correlation between creatinne learance and total ketorolac tromethamine clearance in the elderly and populations with renal impairment (r = 0.5).

In patients with renal disease, the AUC $_{\rm e}$ of each enantiomer increased by approxima 100% compared with healthy volunteers. The volume of distribution doubles for the S enantiomer and increases by 1/5th for the R-enantiomer. The increase in volume of distribution of ketorolac tromethamine implies an increase in unbound fraction.

The AUC a-ratio of the ketorolac tromethamine enantiomers in healthy subjects and patients remained similar, indicating there was no selective excretion of either enantiomer in patients compared to healthy subjects (see WARNINGS, Renal Effects)

Hepatic Insufficiency

There was no significant difference in estimates of half-life, AUC $_{\infty}$ and C $_{\rm max}$ in 7 patients with liver disease compared to healthy volunteers (see **PRECAUTIONS**, **Hepatic Effect** and **Table 2**).

Pharmacokinetic differences due to race have not been identified.

Table 1: Table of Approximate Average Pharm cokinetic Parameters (Mean ± SD) Following Oral, Intramuscular and Intra

Pharmacokinetic Parameters (units)			Bol	us [‡]		mg	mg	mg	mg	mg	mg	Bioavailability (extent)			max [§] (min)	±)34	± ±	± ±	1 C _{ma} : (mcg 7 [sing dose	ı/mL) ile- (±	±	±	±	±	± 0.96	(mcg/mL)	± 0.26	±	3.11N ± 0.87	· ±	096. ± ± 172.	±
C _{min⁸} (mcg/mL) [steady state qid]	0.13 1	0.93 N/A ± 0.26	0.21 1	± 0.35	(mcg/mL)	± 0.20	0.94 ± 0.29	±		±	2.17 ± 0.59	F 1.1 3/	0.175 ± 0.039	±	meta	boliz	edex	creteo feces	% Do d in uri				% Pla bindir			ein				•			_

- fasted volunteers
 † Derived from IM pharmacokinetic studies in 54 normal
- volunteers Derived from IV pharmacokinetic studies in 24 normal

- volunteers

 5 Time-to-peak plasma concentration

 ¶ Mean value was simulated from observed plasma
 concentration data and standard deviation was simulated
 from percent coefficient of variation for observed C max and
 T max data
- # Peak plasma concentration p Not applicable because 60 mg is only recommended as a single dose
- 8 Trough plasma concentration à Average plasma concentration

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L/h/kg] [‡] hours] 54)	IM (n =	Dysfunction	IM (n =	IM an	nd
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= 77)	range =		mg/dL,	= 27	to
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range			to 71		
20 to	50		mean age		
T T			(Oral) =		
			57, range		
			= 39 to 70	(I	
a February draw 20 mg					

- Estimated from 30 mg single IM doses of ketorolac
- ketorolac tromethamine † Estimated from 10 mg single oral doses of ketorolac tromethamine ‡ Liters/hour/kilogram

In normal adult subjects (n = 37), the total clearance of 30 mg IV-administered ketorolac tromethamine was 0.030 (0.017 to 0.051) L/h/kg. The terminal half-life was 5.6 (4 to 7.9) hours (see Kinetics in Special Populations for use of IV dosing of ketorolac tromethamine in pediatric patients).

CLINICAL STUDIES

Adult Patients

Audit Patients
In a postoperative study, where all patients received morphine by a PCA device, patients treated with ketorolac tromethamine. ** as fixed intermittent boluses (e.g., 30 mg initial placedo group, Analgesia was significantly superior, a various; postodorispi pain assessment times, in the patients receiving ketorolac tromethamine. ** plus PCA morphine as compared to patients receiving PCA-administered morphine alone.

Padiatric Patients

There are no data available to support the use of ketorolac tromethamine tablets in pediatric patients.

INDICATIONS AND USAGE

Carefully consider the potential benefits and risks of Ketorolac Tromethamine Tablets USP and other treatment options before deciding to use Ketorolac Tromethamine Tablets USP. Use the lowest effective dose for the shortest duration consistent with individual patient treatment goals.

Acute Pain in Adult Patients

Acute Pain in Adult Patients

Ketorola: Tromethamine Tablets USP are indicated for the short-term (s 5 days) management of moderately severe acute pain that requires analyses at the opioid level, usually in a postoperative setting. Thereipy should wavey be initiated with 10 or 1M dosing of ketorola: tromethamine and Ketorola: Tromethamine Tablets USP are to be used only The total coming of the control of the

CONTRAINDICATIONS

(See also Boxed WARNING.)

Ketorolac tromethamine is contraindicated in patients with previously demonstrated hypersensitivity to ketorolac tromethamine.

inputs sensaway to Rectificate continuations. Retorolac tromethamine is contraindicated in patients with active peptic ulcer disease, in patients with recent gastrointestinal bleeding or perforation and in patients with a history of peptic ulcer disease or gastrointestinal bleeding.

Ketorolac tromethamine should not be given to patients who have experienced asthma, urticaria, or alergic-type reactions after taking aspirin or other NSAIDs. Severe, rarely fatal, anaphylactic-like reactions to NSAIDs have been reported in such patients (see WARNINGS, Anaphylactoid Reactions and PRECAUTIONS, Preexisting Asthma).

Ketorolac tromethamine is contraindicated as prophylactic analgesic before any major surgery.

Ketorolac tromethamine is contraindicated in the setting of coronary artery bypass graft (CABG) surgery (see **WARNINGS**).

Ketorolac tromethamine is contraindicated in patients with advanced renal impairment or in patients at risk for renal failure due to volume depletion (see **WARNINGS** for correction of volume depletion).

Retorolac tromethamine is contraindicated in labor and delivery because, through its prostaglandin synthesis inhibitory effect, it may adversely affect fetal circulation and inhibit uterine contractions, thus increasing the risk of uterine hemorrhage.

Ketorolac tromethamine inhibits platelet function and is, therefore, contraindicated in

patients with suspected or confirmed cerebrovascular bleeding, hemorrhagic diathesis, incomplete hemostasis and those at high risk of bleeding (see **WARNINGS** and **PRECAUTIONS**).

Ketorolac tromethamine is contraindicated in patients currently receiving aspirin or NSAIDs because of the cumulative risks of inducing serious NSAID-related adverse

The concomitant use of ketorolac tromethamine and probenecid is contraindicated

The concomitant use of ketorolac tromethamine and pentoxifylline is contraindicated

(See also Boxed WARNING.)

The total combined duration of use of ketorolac tromethamine tablets and IV or IM dosing of ketorolac tromethamine is not to exceed 5 days in adults. Ketorolac tromethamine tablets are not indicated for use in pediatric patients. The most serious risks associated with ketorolac tromethamine are:

Gastrointestinal Effects - Risk of Ulceration, Bleeding, and Perforation

Ketorola: tromethamine is contraindicated in patients with previously documented peptit ulcers and/or GI bleeding. Ketorola: cromethamine can cause serious gastronitestimal (GI) adverse events including bleeding, ulceration and perforation, of the stomach, small intestine, or large intestine, which can be fatal. These serious adverse events can occur at any time, with or without warning symptoms, in galestics treated with ketorolac.

tromethamine.

Only one in five patients who develop a serious upper GI adverse event on MSAID therapy is symptomatic. More upper gestrointestinal problems, such as dyspepsia, a common and may also occur at any time during MSAID therapy, The inclence and severity of gastrointestinal complications increases with increasing dose of, and durat or treatment with, ketoroisc tromethamine. Do not use ketoroisc tromethamine for more than five days. However, even short-term therapy is not without risk. In addition patients treated with MSAIDs include connomitant use of oral corticosteroids, or anticoagulants, longer duration of NSAID therapy, smoking, use of alcohol older age, and poor general health status. Most spontaneous reports of risk GI Generals are delety or debilitated patients and therefore, special care should be taken in treating the population.

population.

To minimize the potential risk for an adverse GI event, the lowest effective dose should be used for the shortest possible duration. Patients and physicians should remain alert for signs and symptoms of GI uteration and bleading during NSAID therapy and promptly initiate additional evaluation and treatment if a serious GI adverse event is suspected. This should include discontinuation of leteroise tromethamine until serious GI adverse event is uspected in the subject of the control o

NSAIDs should be given with care to patients with a history of inflammatory bowel disease (ulcerative colitis, Crohn's disease) as their condition may be exacerbated.

Hemorrhage

Because prostaginariins play an important role in hemostasis and NSAIDs affect platelet aggregation as wel, use of ketoroluc tromethamine in patients who have coaguitation disorders should be undertaken very cautiously, and those patients should be carefully monitored. Patients on therapeutic doses of anticoaguitants (e.g., heparin or dicumarol derivatives) have an increased risk of bleeding completations if given ketoroluc tromethamine concurrently, therefore, physicians should antimister such concomitation therapy only extremely cautiously. The concurrent use of ketoroluc tromethamine concurrently. The concurrent use of ketoroluc tromethamine concurrently. The concurrent use of ketoroluc tromethamine concurrently. The concurrent use of ketoroluc tromethamine concurrently and destrains have not been studied extensively, but may also be associated with an increased risk of beeding. Until data from such studies are available, physicians should carefully weigh the benefits against the risks and use such concomitant therapy in these patients only extremely cautiously. Patients receiving therapy that affects hemostase should be monitored closely.

In postmarketing experience, postoperative hematomas and other signs of wound bleeding have been reported in association with the per-loperative use of N or IM dosing of ketorols: tromethamine. Therefore, per-loperative use of ketorols: tromethamine should be avoided and postoperative use be undertaken with caution when hemostasis is critical (see PRECAUTIONS).

Renal Effects

xenal Effects
Long-term administration of NSAIDs has resulted in renal papillary necrosis and other renal injury, Renal toxicity has also been seen in patients in whom renal prostaglandins administration of a NSAID may cause a fonce-dependent reduction in prostaglandin formation and secondary, in renal blood flow, which may precipitate over tenal decompensation. Patients at greatest risk of this reaction are those with impared renal function, heart failure, live of sylinticion, hose taking duriects and ACE inhibitors, and the diderly. Discontinuation of NSAID therapy is usually followed by recovery to the pretreatment state.

pretreatment state. Ketorola tromethamine and its metabolites are eliminated primarily by the kidneys, which, in patients with reduced creatinine clearance, will result in diminished clearance of the drug (see CLINICAL PHARMACOLOGY). Therefore, lestoriac tromethamine should be used with caution in patients with impaired renal function (see DOSAGE AND ADMINISTRATION) and such patients should be followed closely. With the use of leatorolac tromethamine, there have been reports of acute renal failure, interstitial neightists and neightotic syndrome.

Impaired Renal Function

Impaired Kenal Function

Ketrolac trombamine is contraindicated in patients with serum creatinine concentrations indicating advanced renal impairment (see CONTRAINDICATIONS). Ketrolac trombamine should be used with caution in patients with impaired renal function or a history of kidney disease because it is a potent inhibitor of prostaglanding synthesis. Because patients with underlying renal insufficiency are at Increased risk of developing acute renal decompensation or failure, the risks and benefits should be assessed prior to glying ketrolac trombamine to these patients.

Anaphylactoid Reactions

As with other MSAIDs, anaphylactoid reactions may occur in patients without a known previous exposure or hypersensitivity to ketoroids: tromethamine. Ketoroids: complex typically occurs in asthmatic patients with one open entering the or without complex typically occurs in asthmatic patients with one open enter infinite with or without nasal polypis, or who exhibit sewere, potentially fatal bronchospasm after taking sprint or other MSAIDs (see CONTRAINDIACTIONS and PREAZUTIONS, Precessiting

Cardiovascular Effects

Cardiovascular Thrombotic Events

Cardiovascular Thrombotic Events

Cinical risks of several COX.2 selective and nonselective NSAIDs of up to three years duration have shown an increased risk of serious cardiovascular (CV) thrombotic events, including myocardial infaction (MR) and strike, which can be fatal. Based on available data, it is unclear that the risk for CV thrombotic events is similar for all NSAIDs. The relative increase in serious CV thrombotic events over baseline conferred by NSAID use appears to be similar in those with and without known CV disease. However, patients with known CV disease or risk factors for CV disease. However, patients with known CV disease or risk factors had nighter than the control of the control o

To minimize the potential risk for an adverse CV event in NSAID-treated patients, use the lowest effective dose for the shortest duration possible. Physicians and patients should lowest effective dose for the shortest duration possible. Physicians and patients sho remain alert for the development of such events, throughout the entire treatment course, even in the absence of previous CV symptoms. Patients should be informed about the symptoms of serious CV events and the steps to take if they occur.

here is no consistent evidence that concurrent use of aspirin mitigates the increase risk of serious CV thrombotic events associated with NSAID use. The concurrent use aspirin and an NSAID, such as keterolac tromethamine, increases the risk of serious gastrointestinal (GI) events (see WARNINGS).

Status Post Coronary Artery Bypass Graft (CABG) Surgery

Two large, controlled clinical trials of a COX-2 selective NSAID for the treatment of pain in the first 10 to 14 days following AGB surgery found an increased incidence of impocardial infarction and stroke. NSAIDs are contrandicated in the setting of CABG (see CONTRAINDICATIONS).

Post-MI Patients

POSCHE TABERIS.

Observational studies conducted in the Danish National Registry have demonstrated that patients treated with NSAIDs in the post-Mip period were at increased risk of reinfarction, CV-related death, and all-cause mortally beginning in the first week of treatment. In this years in NSAID-treated patients compared to 12 per 100 person years in non-NSAID exposed patients. Athough the absolute rate of death decined somewhat after the first year post-MI, the increased relative risk of death in NSAID users persisted over at least the next four years of follow-up.

Avoid the use of ketorolac tromethamine tablets in patients with a recent MI unless the benefits are expected to outweigh the risk of recurrent CV thrombotic events. If ketorolac tromethamine tablets are used in patients with a recent MI, monitor patients for signs of cardiac ischemia.

Hypertension

Hypertension NSAIDs, including ketorolac tromethamine, can lead to onset of new hypertension or worsening of preexisting hypertension, either of which may contribute to the increased incidence of CV events. Patients taking thizacids or loop duretics may have impaired response to these therapies when taking NSAIDs. NSAIDs, including ketorolac tromethamine, should be used with cauthor in patients with hypertension. Blood pressure IOP should be nonkored closely during the intaktion of NSAID treatment and throughout the course of therapy.

Heart Failure and Edema

neart nature and Edema

The Coxib and traditional NSAID Trialists' Collaboration meta-analysis of randomized controlled trials demonstrated an approximately two-fold increase in hospitalizations for heart failure in COX-2 selective-treated patients and nonselective NSAID-treated patients compared to pixe-bet-treated patients. In a Danish National Registry study of patients with heart failure, NSAID use increased the risk of MI, hospitalization for heart failure, and death.

Additionally, fluid retention and edema have been observed in some patients treated with NSAIDs. Use of ketorolac tromethamine may blunt the CV effects of several therapeutic agents used to treat these medical conditions [e.g., dutnetics, ACE inhibots, or

angiotensin receptor blockers (ARBs)] (see DRUG INTERACTIONS)

Avoid the use of ketorolac tromethamine tablets in patients with severe heart failure. Avoid the use of ketorolac tromethamine tablets in patients with severe heart failure. ketorolac tromethamine tablets are used in patients with severe heart failure, morpatients for signs of worsening heart failure.

Skin Reactions

NSAIDs, including ketoroic tromethamme, can cause serous skin adverse events such as exfoliative dematiks, Selvens-johnson syndrome (SJS), and toxic geldermal necrolysis (TEN), which can be fatal. These serious events may occur without warning. Paleiters should be informed about the signs and symptoms of serous skin manifestations and use of the rule you should be discontinued at the first appearance of skin rash, muccost lessons, or any other sign of hypersensitivity.

In late pregnancy, as with other NSAIDs, ketorolac tromethamine should be avoided because it may cause premature closure of the ductus arteriosus.

General

Ketorolac tromethamine cannot be expected to substitute for corticosteroids or to tre corticosteroid insufficiency. Abrupt discontinuation of corticosteroids may lead to disease exacerbation. Patients on prolonged corticosteroid therapy should have their therapy tapered slowly if a decision is made to discontinue corticosteroids.

the rapy capered sown in a decision is made to discontinue controsterious.

The pharmacological activity of ketorolac tromethamine in reducing inflammation diminish the utility of this diagnostic sign in detecting complications of presumed noninfectious, painful conditions.

repatic errect. Ketorolac tromethamine should be used with caution in patients with impaired hepatic function or a history of liver disease. Borderline elevations of one or more liver tests may occur in up to 15% of patients taking NSAIDs including ketorolac tromethamine. These laboratory abnormalities may progress, may remain unchanged, or may be transient with continuing thereapy. Notable ketworation of ALT or ST (approximately 15% or patients in the continuing the patients of the continuing t

A patient with symptoms and/or signs suggesting iver dysfunction, or in whom an abnormal liver test has occurred, should be evaluated for evidence of the development of a most severe hapatic reaction within on the gay with seed loc tronsferamme. If manifestations occur (e.g., essinophila, rash, etc.), ketorolac tromethamine should be discontinued.

Anemia is sometimes seen in patients receiving NSAIDs, including ketorolac tromethamine. This may be due to fluid retention, occult or gross Gi blood loss, or an incompletely described effect upon erythropiess. Platents on long semme treatment with necompletely described effect upon erythropiess. Platents on long semme treatment with checked if they exhibit any signs or symptoms of anemia. NSAIDs inhibit platelet dapgregation and have been shown to profulop bleeding time in some patients. Unlike aspirin, their effect on platelet function is quantitatively less, of shorter duration, and reversible. Patients receiving lextrools ctromethamine who may be adversely affected by alterations in platelet function, such as those with cogulation disorders or patients receiving airticogallutics, should be carefully manifest.

Preexisting Asthma

Preexisting Asthma

Pletins with statum any have aspirin-sensitive asthma. The use of aspirin in patients with aspirin-sensitive asthma has been associated with severe bronchespasm which ran falls. Since rores reactivity, including promotiopagms between a piper and whe nonsteroidal anti-inflammatory drugs has been reported in such aspirin-sensitive patients, ketoroide to more than a point of the administer do to patients with this form of aspirin sensitivity and should be used with caution in patients with preexisting asthma.

Ketorolac tromethamine is a potent NSAID and may cause serious side effects such as gastrointestinal bleeding or kidney failure, which may result in hospitalization and even fatal outcome.

reactivization.

Physicians, when prescribing ketorolac tromethamine, should inform their patients of their guardians of the potential risks of ketorolac tromethamine treatment (see Boxe transport of the potential risks). The properties of their patients of their

- events, and advesse patients not to give ketorolic tromethamine tablets to other family members and to discard any unused drays. Remember that the total combined duration of use of ketorolic tromethamine tablets. Remember that the total combined duration of use of ketorolic tromethamine tablets. Remember that the total combined duration of use of ketorolic tromethamine tablets are not indicated for use in pediatric patients. Returned the tromethamine tablets are not indicated for use in pediatric patients. Patients should also be encouraged to read the NSAID and periodically during the course of ongoing therapy, Patients should also be encouraged to read the NSAID and accompanies each prescription dispersed.

 Cardiovascular Thrombotic Events
 Advise patients to be selet for the symptoms of cardiovascular thrombotic events, Advise patients to be selet for the symptoms of cardiovascular thrombotic events, Advise patients to be selet for the symptoms of cardiovascular thrombotic events, Advise patients to be selet for the symptoms of cardiovascular thrombotic events, Advise patients to be select for the symptoms of cardiovascular thrombotic events, Advise patients to be selected for the symptoms of cardiovascular thrombotic events, Advise patients to be selected for the symptoms of the selected for the symptoms of cardiovascular thrombotic events, School, and the selected for t

- Immediately it they develop any type or reast and such as the prossible prossible.

 An Edema And Edema And

Laboratory Tests

Recause serious GI tract ulcerations and bleeding can occur without warning symptoms, physicians should monitor for signs or symptoms of GI bleeding. Patients on long-term proceedings of the process of

Ketorolac is highly bound to human plasma protein (mean 99.2%). There is no evidence in animal or human studies that ketorolac tromethamine induces or inhibits hepatic enzymes capable of metabolicing itself or other drugs.

Warfarin, Digoxin, Salicylate, and Heparin

Warfarin, Digoxin, Salicylake, and Heparn
The in vitor binding of warfarin to plasma proteins is only slightly reduced by ketorolac tromethamine (99.5% control vs 99.3%) when ketorolac plasma concentrations reach 5 to 10 mcg/ml. Ketorolac does not alter dilgozing protein binding, in vitor studies indicate that, at therapeutic concentrations of salicylate (300 mcg/ml.), the binding of ketorolac was reduced from approximately 92.9% to 91.5%, representing a potential twofold increase in unbound ketorolac plasma levels. Therapeutic concentrations of digoxin, warfarin, bupperfore, nagrozene, plroxixcam, acteralminophen, phenytolin and warfarin, ibuprofen, naproxen, piroxicam, acetaminophen, phenyto tolbutamide did not alter ketorolac tromethamine protein binding.

tabutamide did not aler ketoroku tromethamine protein binding. In a study involving 12 adut voluntees, ketoroku tromethamine tablets were coadiministered with a single dose of 25 mg warfarin, causing no significant changes in pharmacokinetics or pharmacokinetics. In another study, ketoroku tromethamine dosed IV or IM was given with two doses of 5000 U of heparin to 11 healthy volunteers, resulting in a mean template bleeding time of 6. minutes (3.2 to 11.4 min) compared to a mean of 6 minutes (3.4 to 7.5 min) for heparin alone and 5.1 minutes (3.2 to 8.5 min) for pischeo. Although these results do not indicate a significant interaction between ketoroku tromethamine and warfarin or heparin, the administration of ketoroku tromethamine to palents taking airt coagulants should be done extremely PRECAUTIONS, Hematologic Effect), monitored (see WARNINGS and

The effects of warfarin and NSAIDs, in general, on GI bleeding are synergistic, such that the users of both drugs together have a risk of serious GI bleeding higher than the users of either drug alone.

Aspirin

When ketorolac tromethamine is administered with aspirin, its protein binding i reduced, although the clearance of free ketorolac tromethamine is not aftered clinical significance of this interaction is not known, however, as with other NS concomitant administration of ketorolac tromethamine and aspirin is not gener recommended because of the potential of increased adverse effects.

Clinical studies, as well as postmarketing observations, have shown that ketorolad

tromethamine can reduce the natriuretic effect of furosemide and thiazides in some patients. This response has been attributed to inhibition of renal prostaglandin synthe During concomitant therapy with NSAIDs, the patient should be observed closely for signs of renal failure (see WARNINGS, Renal Effects), as well as to assure duretic efficacy.

Prohenecid

Concombant administration of leterolac tramethemine tablets and probeneoid resulted in decreased clearance and volume of distribution of leterolac and significant icrosses in leterolac plasma levels (total AUC increased approximately threefold from 5.4 to 17.8 morghmin) and reminal half-life increased approximately throrid for mo.6.6 to 15.1 hours. Therefore, concombant use of ketorolac tromethamine and probeneoid is contrainficated.

NSAIDs have produced an elevation of plasma lithium levels and a reduction in renal lithium clearance. The mean minimum thatum concentration increased 15% and the renal clearance was decreased by approximately 20%. These effects have been attributed to inhibition of renal prostaglandin synthesis by the NSAID. Thus, when NSAIDs and lithium are administered concurrently, subjects should be observed carefully for signs of lithium toxicity.

Methotrexate

NSAIDs have been reported to competitively inhibit methotrexate accumulation in rabbit kidney slices. This may indicate that they could enhance the toxicity of methotrexate. Caution should be used when NSAIDs are administered concomitantly with methotrexate.

ACE Inhibitors/Angiotensin II Receptor Antagonists

Concomitant use of **ACE inhibitors and/or angiotensin II receptor antagonists** may increase the risk of renal impairment, particularly in volume-depleted patients.

Reports suggest that NSAIDs may diminish the anthypertensive effect of ACE inhibitors and/or angiotensin II receptor antagonists. This interaction should be given consideration in patients taking NSAIDs concomitantly with ACE inhibitors and/or angiotensin II receptor antagonists.

Antiepileptic Drugs

Sporadic cases of seizures have been reported during concomitant use of ketorolac tromethamine and **antiepileptic drugs** (phenytoin, carbamazepine).

Psychoactive Drugs

Hallucinations have been reported when ketorolac tromethamine was used in patients taking **psychoactive drugs** (fluoxetine, thiothixene, alprazolam).

Pentoxifulline When ketorolac tromethamine is administered concurrently with pentoxifylline, there is an increased tendency to bleeding.

Nondepolarizing Muscle Relaxants

In postmarketing experience there have been reports of a possible interaction betwee ketorolac tromethamine ^{IV/M} and **nondepolarizing muscle relaxants** that resulte apnea. The concurrent use of ketorolac tromethamine with muscle relaxants has no been formally studied.

Selective Serotonin Reuptake Inhibitors (SSRIs)

There is an increased risk of gastrointestinal bleeding when selective serotonin reuptake inhibitors (SSRIs) are combined with NSAIDs. Caution should be used when NSAIDs are administered concomitantly with SSRIs.

Carcinogenesis, Mutagenesis, Impairment of Fertility

An 18 month study in nice with ord does of lettorotac tronethamine at 2 mg/kg/day (0.9 times the human systemic exposure at the recommended IM or IV dose of 30 mg (d), based on are-under-the-plasma-concentration curve [AUC]), and a 24 month study in rats at 5 mg/kg/day (0.5 times the human AUC) showed no evidence of tumorigenicky.

Ketrolac tromethamine was not mutagenic in the Ames test, unscheduled DNA synthesis and repair, and in forward mutation assays. Ketrolac tromethamine did not cause chromosome breakage in the in vivo mouse micronucleus assay. At 1590 mcg/mL and at higher concentrations, ketrolac tromethamine increased the incidence of chromosomal aberrations in Chiese hamster ovariant cells.

Impairment of fertility did not occur in male or female rats at oral doses of 9 mg/kg (0.9 times the human AUC) and 16 mg/kg (1.6 times the human AUC) of ketorolac tromethamine, respectively.

Pregnancy

Teratogenic Effects

Preanancy Category C

regulars, v.ausgury V. Reproduction studies have been performed during organogenesis using daly oral dose of ketorolec tromethamine at 2.6 mg/kg (0.37 times the human AUC) in rabbits and at 10 mg/kg (1 times the human AUC) in rats. Results of these studies did not reveal evidence of teratogenicky to the fetus. However, animal reproduction studies are not always predictive of human response.

Nonteratogenic Effects

Because of the known effects of nonsteroidal anti-inflammatory drugs on the fetal cardiovascular system (closure of ductus arteriosus), use during pregnancy (particular) late pregnancy) should be avoided. Oral doses of ketorolac tromethamine at 1.5 mg/kg (0.14 times the human AUC), administered after gestation Day 17, caused dystocia and higher pun mortally in rats.

There are no adequate and well-controlled studies of ketorolac tromethamine in pregnant women. Ketorolac tromethamine should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Labor and Delivery

The use of ketorolac tromethamine is contraindicated in labor and delivery because through its prostaglandin synthesis inhibitory effect, it may adversely affect fetal circulation and inhibit uterine contractions, thus increasing the risk of uterine hemorrhage (see CONTRAINDICATIONS).

Effects on Fertility

The use of ketorolac tromethamine, as with any drug known to inhibit cyclooxygenase/prostaglandin synthesis, may impair fertifily and is not recommended in women attempting to concelve. In women who have difficulty conceiving or are undergoing investigation of infertifily, withdrawal of ketorolac tromethamine should be considered.

Nursing Mothers

Nursing Mothers

Thirded data from one published study involving 10 breastfeeding women 2 to 6 days postpartum showed low levels of ketorobic in breast milk. Levels were undetectable (less than 5 ng/ml.) 1 d of the patients. After a single administration of 10 mg of ketorobic tromethamine tablets, the maximum milk concentration observed was 7.3 ng/ml., and the maximum milk-to-plasma ratio was 0.037. After 1 day of dosing 10 mg every 6 hours), the maximum milk concentration was 7.3 ng/ml., and the maximum milk-to-plasma ratio was 0.037. After 1 day of dosing 10 mg every 6 hours), the maximum milk-to-oncentration was 7.3 ng/ml., and the maximum milk-to-query 6 mg milk-to-plasma ratio 6 lag milk-to-plasma ratio 6 lag milk-to-plasma ratio 6 lag milk-to-plasma for 10 kg milk-to-plasma for 1

Exercise caution when ketorolac is administered to a nursing woman. Available information has not shown any specific adverse events in nursing infants; however, instruct patients to contact their infant's health care provider if they note any adverse

Pediatric Use

Ketorolac tromethamine tablets are not indicated for use in pediatric patients. The safety and effectiveness of ketorolac tromethamine tablets in pediatric patients below the age of 17 have not been established.

Geriatric Use (≥ 65 Years of Age)

Because ketrorbic tromethamine may be cleared more slowly by the elderly (see CLINICAL PHARMACOLOGY) who are also more sensitive to the dose-related adverseffect of MSAIDs (see MARMINGS, Gastrointestimal Effects – Risk of Ulceration, ADMINISTRATION), and careful clinical monitoring must be used when treating the elderly with ketorolac tromethamine.

ADVERSE REACTIONS

ADVERSE REACTIONS

Adverse reaction rates increase with higher doses of ketorolac tromethamine.
Practitioners should be alert for the severe complications of treatment with ketorolac
tromethamine, such as GI ulceration, bededing and perforation, postoperative bleeding
acute reral failure, anaphylactic and anaphylactod reactions and laver failure (see Box
WARNING, WARNINGS, PREACTIONS, and DOSAGE AND ADMINISTRATION).
These INSAID-related complications can be serious in certain patients for whom keture
tromethamine is indicated, specified when the drug is used mappropriets.

In patients taking ketorolac tromethamine or other NSAIDs in clinical trials, the most frequently reported adverse experiences in approximately 1% to 10% of patients are

Gastrointestinal (GI) experier		
abdominal pain *	constipation/diarrhea	dyspepsia *
flatulence	GI fullness	GI ulcers (gastric/duodenal)
gross bleeding/perforation	heartburn	nausea *
stomatitis	vomiting	
Other experiences:		
abnormal renal function	anemia	dizziness
drowsiness	edema	elevated liver enzymes
headaches *	hypertension	increased bleeding time
injection site pain	pruritus	purpura
rashes	tinnitus	sweating
* Incidence greater than 10%	5	

Additional adverse experiences reported occasionally (< 1% in patients taking ketorolar tromethamine or other NSAIDs in clinical trials) include:

Body as a Whole: fever, infections, sepsis

Cardiovascular: congestive heart failure, palpitation, pallor, tachycardia, syncope Dermatologic: alopecia, photosensitivity, urticaria

Gastrointestinal: anorexia, dry mouth, eructation, esophagitis, excessive thirst, gastritis, glossitis, hematemesis, hepatitis, increased appetite, jaundice, melena, rectal bleeding.

Hemic and Lymphatic: ecchymosis, eosinophilia, epistaxis, leukopenia
thrombocytopenia

Metabolic and Nutritional: weight change

Nervous System: abnormal dreams, abnormal thinking, anxiety, asthenia, confusion, depression, euphoria, extrapramial symptoms, hallucinations, hyperkinesis, nability to concentrate, insommia, nervousness, paresthesia, sommolence, stupor, tremors, vertigo, malaise

Reproductive, female: infertility

Respiratory: asthma, cough, dyspnea, pulmonary edema, rhinitis

Special Senses: abnormal taste, abnormal vision, blurred vision, hearing loss

Urogenital: cystitis, dysuria, hematuria, increased urinary frequency, interstitial nephritis, oliguria/polyuria, proteinuria, renal failure, urinary retention

Other rarely observed reactions (reported from postmarketing experience in patients taking ketorolac tromethamine or other NSAIDs) are:

Body as a Whole: angioedema, death, hypersensitivity reactions such as anaphylaxis, anaphylactoid reaction, laryngeal edema, tongue edema (see WARNINGS), myalgia

Cardiovascular: arrhythmia, bradycardia, chest pain, flushing, hypotension, myocardial infarction, vasculitis

marcton, vascuuts

Dermatologic: exfolative dermatitis, erythema multiforme, Lyell's syndrome, bullous reactions including Stevens-Johnson syndrome and toxic epidermal necrolysis

Gastrointestinal: acute pancreatitis, liver failure, uterative stomatitis, exacerbation of inflammatory bowel disease (uterative colits, Crohn's disease)

Hemic and Lymphatic: agranulocytosis, aplastic anemia, hemolytic anemia, lymphadenopathy, pancytopenia, postoperative wound hemorrhage (rarely req blood transfusion - see Boxed WARNING, WARNINGS, and PRECAUTIONS;

Metabolic and Nutritional: hyperglycemia, hyperkalemia, hyponatre

Nervous System: aseptic meningitis, convulsions, coma, psychosis

Respiratory: bronchospasm, respiratory depression, pneumonia Special Senses: conjunctivitis

Urogenital: flank pain with or without hematuria and/or azotemia, hemolytic uremic syndrome

Postmarketing Surveillance Study

A large postmarketing observational, nonrandomized study, involving approximately 10,000 patients receiving leteroise tromethamine ^{MRI}, demonstrated that the risk of cinically serious pastromitectural (of) bleeding was obse-dependent (see Tables 3A a 3B). This was particularly true in etierly patients who received an average daily dose greater than 60 mightay of reference tromethamine ^{MRI} (see Tables 3A).

Table 3: Incidence of Clinically Serious GI Bleeding as Related to Age, Total Daily Dose, and History of GI Perforation, Ulcer, Bleeding (PUB) After up to 5

,					
		ents Without His			
Age of Patients	Tot	al Daily Dose of Ket	orolac Tromethamin	e IV/IM	
	≤ 60 mg	> 60 to 90 mg	> 90 to 120 mg	> 120 mg	
< 65 years of age	0.4%	0.4%	0.9%	4.6%	
≥ 65 years of age	1.2%	2.8%	2.2%	7.7%	
	B. Adult Pa	tients With Histo	ry of PUB		
Age of Patients	Tot	al Daily Dose of Ket	orolac Tromethamin	e IV/IM	
	≤ 60 mg	> 60 to 90 mg	> 90 to 120 mg	> 120 mg	
< 65 years of age	2.1%	4.6%	7.8%	15.4%	
≥ 65 years of age	4.7%	3.7%	2.8%	25%	

Symptoms following acute NSAID overdoses are usually limited to lethargy, drowshess, nausea, vomiting, and epigastric pain, which are generally reversible with supportive care. Gastronitestrain bleeding can occur. Hypertension, acute renaf faiture, respiratory depression and coma my occur, but are rare. Anaphyloctoid reactions have been reported with therapeutic nigestion of NSAIDs, and my occur flowing an overdose.

Treatment

Treatment
Patients should be managed by symptomatic and supportive care following a NSAIDs overdose. There are no specific antidotes. Emesis and/or activated charcoal (60 g to 10 g in adults, 1 glob to 2 g/kg in children) and/or osmotic catharite may be indicated in patients seen within 4 hours of ingestion with symptoms or following a large oral patients seen within 4 hours of ingestion with symptoms or following a large oral hemodalysis or themoperfusion may not be useful due to high protein binding.
Single overdoses of ketonois transchamine have been variously associated with abdomain pain, nutures, owniting, hyperventiation, peptit uckers and/or erosive gastritis and renal dysfunction which have resolved after discontinuation of dosing.

DOSAGE AND ADMINISTRATION

Carefully consider the potential benefits and risks of ketorolac tromethamin tablets and other treatment options before deciding to use ketorolac tromethamine tablets. Use the lowest effective does for the shortest duration consistent with individual patient treatment goals. In adults, the combined duration of use of IV or IM dosing of ketorolac tromethamine tablets is not to exceed 5 days. In adults, the use of ketorolac tromethamine tablets is only indicated as continuation therapy to IV or IM dosing of ketorolac tromethamine tablets.

Transition from IV or IM dosing of ketorolac tromethamine (single- or multiple-dose) to multiple-dose ketorolac tromethamine tablets:

Patients age 17 to 64: 20 mg PO once followed by 10 mg q4 to 6 hours prn **not > 40** mg/day

Patients age \geq 65, renally impaired, and/or weight < 50 kg (110 lbs): 10 mg PO once followed by 10 mg q4 to 6 hours prn <code>not > 40 mg/day</code>

Oral formulation should not be given as an initial dose

Do not shorten dosing interval of 4 to 6 hours.

Total duration of treatment in adult patients: the combined duration of use of IV or IM dosing of ketorolac tromethamine and ketorolac tromethamine tablets is not to exceed 5 days.

The following table summarizes ketorolac tromethamine tablet dosing instructions in terms of age group:

Table 4: Summary of Dosing Instructions

Patient Population	Ketorolac Tromethamine Tablets (following IV or IM dosing of ketorolac tromethamine)
Age < 17 years	Oral not approved
Adult Age 17 to 64 years	20 mg once, then 10 mg q4 to 6 hours prn not > 40 mg/day
Adult Age ≥ 65 years, renally impaired, and/or weight < 50 kg	10 mg once, then 10 mg q4 to 6 hours prn not > 40 mg/day

HOW SUPPLIED

Ketorolac Tromethamine Tablets USP are available as follows:

To mg: White, round, convex, unscored, film-coated tablets, debossed "93" on one side and "314" on the other side. They are available in bottles of 4 tablets (NDC 66267-832-04) and 8 tablets (NDC 66267-832-08) Store at 20° to 25°C (68° to 77°F) [See USP Controlled Room Temperature].

Dispense in a tight, light-resistant container as defined in the USP, with a child-resistant closure (as required).

PROTECT FROM LIGHT AND EXCESSIVE HUMIDITY

KEEP THIS AND ALL MEDICATIONS OUT OF THE REACH OF CHILDREN.

Distributed By

TEVA PHARMACEUTICALS USA, INC.

North Wales, PA 19454 Rev. K 7/2015

Medication Guide for Nonsteroidal Anti-inflammatory Drugs (NSAIDs)
What is the most important information I should know about medicines
called Nonsteroidal Anti-inflammatory Drugs (NSAIDs)?
NSAIDs can cause serious side effects, Including:

Increased risk of a heart attack or stroke that can lead to death. This risk may happen early in treatment and may increase:

- with increasing doses of NSAIDs
- with longer use of NSAIDs
- Do not take NSAIDs right before or after a heart surgery called a "coronary artery bypass graft (CABG)."
- Avoid taking NSAIDs after a recent heart attack, unless your healthcare provider tells you to. You may have an increased risk of another heart attack if you take NSAIDs after a recent heart attack.
- Increased risk of bleeding, ulcers, and tears (perforation) of the esophagus (tube leading from the mouth to the stomach), stomach and

intestines: anytime during use

The risk of getting an ulcer or bleeding increases with:

past history of stomach ulcers, or stomach or intestinal bleeding with use of NSAIDs

older age

taking medicines called "corticosteroids", "anticoagulants", "SSRIs", or "SNRIs"

increasing doses of NSAIDs

longer use of NSAIDs

 poor health advanced liver disease

 smoking drinking alcohol

bleeding problems

NSAIDs should only be used: • exactly as prescribed

at the lowest dose possible for your treatment

• for the shortest time needed

What are NSAIDs?

NSAIDs are used to treat pain and redness, swelling, and heat (inflammation) from medical conditions such as different types of arthritis, menstrual cramps, and other

types of short-term pain. Who should not take NSAIDs? Do not take NSAIDs: if you have had an asthma attack, hives, or other allergic reaction with aspirin or any other NSAIDs.

right before or after heart bypass surgery

fore taking NSAIDs, tell your healthcare provider about all of your medical nditions, including if you: have liver or kidney problems

- are pregnant or plan to become pregnant. Talk to your healthcare provider if you are considering taking NSAIDs during pregnancy. You should not take NSAIDs after 29 weeks of pregnancy.
- are breastfeeding or plan to breast feed.

Tell your healthcare provider about all of the medicines you take, including prescription or over-the-counter medicines, vitamins or herbal supplements. NSAIDs and some other medicines can interact with each other and cause serious side effects. Do not start taking any new medicine without talking to your

healthcare provider first.
What are the possible side effects of NSAIDs?

new or worse high blood pressure heart failure

- life-threatening skin reactions
- low red blood cells (anemia)
- Ife-threatening allergic reactions
- Other side effects of NSAIDs include: stomach pain, constipation, diarrhea, gas heartburn, nausea, vomiting, and dizziness.

Get emergency help right away if you get any of the following symptoms: • shortness of breath or trouble breathing • slurred speech

chest pain

swelling of the face or throat

weakness in one part or side of your body

Stop taking your NSAID and call your healthcare provider right away if you get any of the following symptoms:

 vomit blood

diarrhea

is black and sticky like tar

itchina

unusual weight gain

your skin or eyes look yellow

skin rash or blisters with fever

swelling of the arms, legs, hands and feet

flu-like symptoms

If you take too much of your NSAID, call your healthcare provider or get medical help right away. These are not all the possible side effects of NSAIDs. For more information, ask your healthcare provider or pharmacet about NSAIDs. Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-000-FDA-1008. Other information about NSAIDs

Aspirin is an NSAID but k does not increase the chance of a heart attack. Aspirin can cause bleeding in the brain, stomach, and intestines. Aspirin can also cause ulcers in the stomach and intestines.

Some NSAIDs are sold in lower doses without a prescription (over-the-counter). Tak to your healthcare provider before using over-the-counter NSAIDs for more than 10 days.

General information about the safe and effective use of NSAIDs

Medicines are sometimes prescribed for purposes other than those listed in a dedication Guide. Do not use NSAIDs for a condition for which it was not prescribed. D tot give NSAIDs to other people, even if they have the same symptoms that you have.

It may harm them.

If you would like more information about NSAIDs, talk with your healthcare provider. You are sky your pharmacist or healthcare provider for information about NSAIDs that is

written for health professionals. This Medication Guide has been approved by the U.S. Food and Drug Administration.

Package/Label Display Panel



Product Information						
Product Type	HUMAN PRI DRUG	ESCRIPTION	Item Code (Source)		NDC:66267-832(0314)	NDC:0093-
Route of Administration	ORAL					
Active Ingredient/Active						
	gredient Na				is of Strength	Strengt
KETOROLAC TROMETHAMINI	(UNII: 4EVE59-	46BQ) (KETORI	DLAC -	KETOR	OLAC THAMINE	10 mg
Inactive Ingredients						
Inactive Ingredients	Ingre	dient Name				Strength
•						Strength
HYDROXYPROPYL CELLULOS	E (1600000 W	AMW) (UNII: F				Strength
HYDROXYPROPYL CELLULOS HYPROMELLOSE 2910 (3 MP	E (1600000 W A.S.) (UNII: 0VU	YAMW) (UNII: F T3PMY82)				Strength
HYDROXYPROPYL CELLULOS HYPROMELLOSE 2910 (3 MP HYPROMELLOSE 2910 (50 M	E (1600000 W A.S.) (UNII: 0VU PA.S.) (UNII: 1IV	/AMW) (UNII: F T3PMY82) /H67816N)				Strength
HYDROXYPROPYL CELLULOS HYPROMELLOSE 2910 (3 MP HYPROMELLOSE 2910 (50 M LACTOSE MONOHYDRATE (UI	E (1600000 W A.S) (UNII: 0VU PA.S) (UNII: 1IV NII: EWQ57Q8IS:	/AMW) (UNII: F T3PMY82) /H67816N)				Strength
HYDROXYPROPYL CELLULOS HYPROMELLOSE 2910 (3 MP HYPROMELLOSE 2910 (50 M LACTOSE MONOHYDRATE (UM MAGNESIUM STEARATE (UMI:	E (1600000 W A.S) (UNII: 0VU PA.S) (UNII: 1IV NII: EWQ57Q8I5: 70097M6I30)	/AMW) (UNI: 8 T3PMY82) /H67816N) X)				Strength
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Inactive Ingredients HYDROXYPROPYL CELLULOS HYPROMELLOSE 2310 (3 MP HYPROMELLOSE 2310 (50 M LACTOSE MONOHYDATE (UM MAGNESIUM STEARATE (UMI) CELLULOSE, MICROCHYSTAL POLYETHYLENE GLYCOL 400 TITTANIUM DIOXIDE (UMI): 1581	E (1600000 W A.S) (UNII: OVU PA.S) (UNII: 1IV NII: EWQ57Q8I5. 70097M6I30) LINE (UNII: OP1 (UNII: B697894	VAMW) (UNII: 8 T3PMY82) (H67816N) X) LR32D61U)				Strength
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SI	nape	ROUND	Size		8mm
FI	avor		Imprint Code		93;314
C	ontains				
P	ackaging				
#	Item Code	Package D	escription	Marketing Start Date	Marketing End Date
1	NDC:66267-832- 04	4 in 1 BOTTLE; Type 0: Product	Not a Combination 01	1/24/2017	
2	NDC:66267-832- 08	8 in 1 BOTTLE; Type 0: Product	Not a Combination 01	1/24/2017	
N	larketing	Information			
		Annilosation Mus	nber or Monograph	Marketing Start	Marketing En
	Marketing Category	Application Null	tation	Date	Date

Labeler - NuCare Pharmaceuticals, Inc. (010632300)

Establishment

Name Address IDFEI Business Operations

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