

# Ketorolac Tromethamine Injection, USP

Product Name	NDC Number	Concentration	Fill Volume	Pack Size
Ketorolac Tromethamine Injection, USP 15 mg/l mL	65145-146-25	15mg per mL	1mL	25 vials
Ketorolac Tromethamine Injection, USP 30 mg/l mL	65145-145-25	30mg per mL	lmL	25 vials
Ketorolac Tromethamine Injection, USP 60 mg/2 mL	65145-147-25	30mg per mL	2mL	25 vials

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Strength	Cardinal	Cencora	McKesson
15 mg/1 mL	5960059	10295153	2998763
30 mg/1 mL	5960042	10295001	2998789
60 mg/2 mL	5960067	10294969	2998797

# It's more than a product. It's a promise.



Please see full prescribing information, including boxed warning, for KETOROLAC TROMETHAMINE Injection, USP, enclosed.

## **⊗** AP Rated **⊗** Not Made With Natural Rubber Latex

- For more information visit www.caplin-usa.net
- Caplin Steriles Customer Service: (833) 487-0705

#### FOR INTRAVENOUS/INTRAMUSCULAR USE (15 mg and 30 mg) FOR INTRAMUSCULAR USE ONLY (60 mg)

toroiac tromethamine, a nonsteroidal anti-inflammatory drug (NSAID), is indicated for the short-term (up to 5 days in adults) management of moderately severe acute pain that requires analgesia at the contract of the contra

etorolac tromethamine is not indicated for use in pediatric patients and it is NOT indicated for minor or chronic painful conditions. Increasing the dose of ketorolac tro eyond the label recommendations will not provide better efficacy but will increase the risk of developing serious adverse events.

Ketorolac tromethamine can cause peptic ulcers, gastrointestinal bleeding and/or perforation of the stomach or intestines, which can be fatal. These events can occur at any time during use without warning symptoms. Therefore, ketorolac bromethamine is CONTRAINDICATED in patients with active peptic ulcer disease, in patients with recent gastrointestinal bleeding or perforation, are in patients with a history of peptic ulcer disease or gastrointestinal bleeding. Elderly patients are at greater risk for serious gastrointestinal events (see WARNINGS).

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

Ketorolac tromethamine is CONTRAINDICATED in the setting of coronary artery bypass graft (CABG) surgery (see CONTRAINDICATIONS and WARNINGS)

IAL RISK

Ketorolac tromethamine inhibits platelet function and is, therefore, CONTRAINDICATED in patients with suspected or con incomplete hemostasis and those at high risk of bleeding (see WARNINGS and PRECAUTIONS).

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

Pypersentitivity reactions, ranging from bronchospasm to anaphylactic shock, have occurred and appropriate counteractive measures must be available when administerin betrorists tronsehamine injection (see CONTRAMBICATIONS and WARRINGS), Kostroics tronsehamines is CONTRAMDICATED in patients with previously demonstrated hyperse tronsehamine or allegic manifestations to baptin or other rorestroad aid-inflammatively trugh (SAVIDs).

Ketorolac tromethamine is CONTRAINDICATED for intrathecal or epidural administration due to its alcohol content

SK DURING LABOR AND DELIVERY

MITANT USE WITH NSAIDs

Dosage should be adjusted for patients 65 years or older, for patients under 50 kg (110 lbs.) of body weight (see DOSAGE AND ADMINISTI creatinine (see WARNINGS). Doses of ketorolac tromethamine injection are not to exceed 60 mg (total dose per day) in these patients.

Ketorolac tromethamine tablets are indicated only as continuation therapy to ketorolac tromethamine injection, and the combined duration of use of ket tromethamine tablets is not to exceed 5 (five) days, because of the increased risk of serious adverse events

mended total daily dose of ketorolac tromethamine tablets (maximum 40 mg) is significantly lower than for ketorolac tromethamine injection (maximum 40 mg)

Ketorolac Tromethamine Injection, USP is a member of the pyrrolo-pyrrole group of nonsteroidal anti-inflammatory drugs (NSAIDs). The chemical name for keto dro-1H-pyrrolizine-1-carboxylic acid, compound with 2-amino-2-(nydroxymethyl)-1,3-propanediol (1:1), and the structural formula is presented in Figure 1.

Ketorolac tromethamine is a racemic mixture of [3]S and [-3]R ketorolac tromethamine. Ketorolac tromethamine may exist in three crystal forms. All forms are equally soluble in water. Ketorolac tromethamine and 4.5 and an e-oction/water partition coefficient of 0.25. The methodical weight of ketorolac tromethamine is 378-46. Set of 1.5 miles and 1.5 miles are considered in the consideration of 1.5 miles are considered in the consideration of 1.5 miles are considered in the consideration of 1.5 miles are consideration of 1.

Ketorolac tromethamine possesses no sedative or anxiolytic properties.

The peak analgesic effect of ketorolac tromethamine occurs within 2 to 3 hours and is not statistically significantly different over the recordifference between large and small obsess of ketorolac tromethamine by either route is in the duration of analgesia.

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

intelligation is a cascinic mission or 15° dies ; pr. visionalisation of Intravenous, Intramuscular and Oral Pharmacokinetics
cokinetics of ketorolac tromethamine, following intravenous, intramuscular, and oral doses of ketorolac tromethaon of the oral and intramuscular forms of ketorolac tromethamine was equal to that following an intravenous bolus.

Table 1: Table of Approximate Average Pharmacokinetic Parameters (Mean± Following Oral, Intramuscular and Intravenous Doses of Ketorolac Trome

	Oral <sup>†</sup>	Intramuscular*		Intravenous Bolus ‡		
Pharmacokinetic Parameters (units)	10 mg	15 mg	30 mg	60 mg	15 mg	30 mg
Bioavailability (extent)		100%				
T <sub>max</sub> (min)	44±34	33±21**	44±29	33±21**	1.1±0.7**	2.9±1.8
C <sub>max</sub> (mcg/mL) [Single-dose]	0.87±0.22	1.14±0.32**	2.42±0.68	4.55±1.27**	2.47±0.51**	4.65±0.96
C <sub>max</sub> (mcg/mL) [steady state qid]	1.05±0.26**	1.56±0.44**	3.11±0.87**	N/A <sup>++</sup>	3.09±1.17**	6.85±2.61
C <sub>min</sub> <sup>3</sup> (mcg/mL) [steady state qid]	0.29±0.07**	0.47±0.13**	0.93±0.26**	N/A	0.61±0.21**	1.04±0.35
C <sub>nrq</sub> <sup>4</sup> (mcg/mL) [steady state qid]	0.59±0.2**	0.94±0.29**	1.88±0.59**	N/A	1.09±0.3**	2.17±0.59
V <sub>j</sub> s (L/kg)	0.175±0.039			0.210	±0.044	

adults, following administration of single oral, inframuscular or intravenous doses of kebrolac tromethamine in the recommended dosage ranges, the clearance of the racemate dose not change. This properties that the pharmacokinetics of kebrolac tromethamine in adults, following single or multiple inframuscular, intravenous, or recommended oral doses of kebrolac tromethamine, are linear. At the higher commended doses, there is a proportional increase in the concentrations of free and bound racemate.

JUSTICATION THE MEAN THE PROPRIES AND THE MEAN T Ketorolac tromethamine is excreted in human milk (see PRECAUTIONS - Nursing Mothers).

Reforciac tomelemanine 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.

Accumulation

Kethoricals from elementamine administered as an intravenous bolus, every 6 hours, for 5 days, to healthy subjects (n = 13), showed no significant difference in C<sub>me</sub> on Day 1 and Day 5. Trough levels averaged 0.29 mozymL (SD ± 0.13) on Day 1 and 0.55 mozymL (SD ± 0.23) on Day 6. Steady state was approached after the fourth dose.

cial populations (geriatric, pediatric, renal failure patients, or hepatic disease patients)

Testinetics in Special Populations
Geriatric Patients
Based on single-dose data only, the half-life of the ketorolac fromethamine racemate increased from 5 to 7 hours in the elderly (65 to 78 years) compared with young healthy volunteers (24 to 35 y Table 2). There was little difference in the C\_\_ for the two groups (elderly, 2.52 mog/mt. ± 0.77; young, 2.99 mog/mt. ± 1.03) (see PRECAUTIONS – Geriatric Use).

Politistic Protects
Limited informations available reporting the pharmacolaristics of bosing of leterotace bromelescening in the politistic population. Following a steple intervence books does of 0.5 mp/gs in 1 or inclined information available reporting in 1 or inclined information and inclined inclined information and inclined inclined information and inclined inclined inclined inclined information and inclined inclined

in patients with renal diseases, the ALD\_col each fearitiment increased by approximately 100% compared with telliground visit to a specific process of the patients with renal diseases, the ALD\_col each fearitiment increased by approximately 100% compared with the healthy volunthers. The volume of distribution doubles for the S-enantion 1/3/th for the A-enantioner. The increase in volume of distribution of oubles for the S-enantion 1/3/th for the A-enantioner. The increase in volume of distribution of oubles for the S-enantion 1/3/th for the A-enantioner. The increase in volume of distribution of oubles for the S-enantion 1/3/th for the A-enantioner. The increase in volume of distribution oubles for the S-enantion 1/3/th for the A-enantioner. The increase is volume 1/3/th for the A-enantioner in patient in the A-enantion 1/3/th for the A-enantioner in the A-enantion 1/3/th for the A-enantioner in the A-enantioner

tes of half-life, AUC,, and C,, in 7 patie

cokinetic differences due to race have not been identified.

Table 2:
The Influence of Age, Liver and Kidney Function, on the Clearance and Terminal Half-life of Ketorolac Trometh
(Inframuscular' and Oral') in Adult Populations

	[in L/h/kg] <sup>3</sup>		[in hours]	
Type of Subjects	Intramuscular Mean (range)	Oral Mean (range)	Intramuscular Mean (range)	Oral Mean (range)
Normal Subjects Inframuscular (n = 54) mean age = 32, range = 18-60 Oral (n = 77) mean age = 32, range = 20-60	0.023 (0.010-0.046)	0.025 (0.013-0.050)	5.3 (3.5-9.2)	5.3 (2.4-9)
Healthy Elderly Subjects Intramuscular (n = 13), Oral (n = 12) mean age = 72, range = 65-78	0.019 (0.013-0.034)	0.024 (0.018-0.034)	7 (4.7-8.6)	6.1 (4.3-7.6)
Patients with Hepatic Dysfunction Intramuscular and Oral (n = 7) mean age = 51, range = 43-64	0.029 (0.013-0.066)	0.033 (0.019-0.051)	5.4 (2.2-6.9)	4.5 (1.6-7.6)
Patients with Renal Impairment Intramuscular (n = 25), 0ral (n = 9) serum creatinine = 1.9-5.0 mg/dL, mean age (intramuscular) = 54, range = 35-71 mean age (0ral) = 57, range = 3970	0.015 (0.005-0.043)	0.016 (0.007-0.052)	10.3 (5.9-19 <i>2</i> )	10.8 (3.4-18.9)
Renal Dialysis Patients Intramuscular and Oral (n = 9) mean age = 40, range = 27-63	0.016 (0.003-0.036)	-	13.6 (8.0-39.1)	-

Intravenous-Administration: In normal subjects (n=27), the total clearance of 30 mg intravenous-administered ketorolac tromethamine was 0.030 (0.017-0.051) L/h/kg. The te (4.0-7.9) hours, (see Kinetics in Special Populations for use of intravenous dosing of ketorolac tromethamine in pediatric patients.)

re all patients received morphine by a PCA device, patients treated with ketorolac tromethamine intravenous as fixed intermittent boluses (e.g., 30 mg initial dose followed by 15 significantly less morphine (26%) than the placebo group. Analysisia was significantly superior, at various postdosing pain assessment times, in the patients receiving ketorolac us PCA morphine accompared to patients receiving Nationalisated morphine alone.

INDICATIONS AND USAGE.
Carefully consider the optimization performs the form of the property o

Acute Pain in Adult Patients
Kotronics Tromethamine is indicated for the short-term (\_55 days) management of moderately severe acute pain that requires analgesia at the opioid level, usually in a postoperative
always be initiated with intravenous or inframuscular dosing of ketorolac tromethamine, and oral ketorolac tromethamine is to be used only as continuation treatment, if necessary. The total combined duration of use of keterolac tromethamine injection and oral keterolac tromethamine is not to exceed 5 days of use because of the potential of increasing the frequency and severity of adverse reactions associated with the recommended doses (see WARRINGS, PRECAUTIONS, DOSAGE AND ADMINISTRATION), and ADVERSE REACTIONS). Patients should be switched to alternative analogies as soon as possible, but keterolac incremelamine therapy is not to exceed 5 days.

CONTRAINDICATIONS
(see also Boxed WARNING)

Ketorolac Tromethamine is contraindicated in patients with previously demonstrated hypers

Ketorolac tromethamine is contraindicated in patients with active peptic ulcer disease, in patients with recent gastrointestinal blee aastrointestinal bleeding.

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

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

mine is contraindicated in patients currently receiving aspirin or NSAIDs be

The concomitant use of ketorolac tromethamine and probenecid is contraindicated.

The concomitant use of ketorolac tromethamine and pentoxifylline is contraindicated.

The total combined duration of use of oral ketorolac tromethamine and intravenous or intramuscular dosing of ketorolac tromethamine is not to exceed 5 days in adults. Ket for use in pediatric patients. The most serious risks associated with ketorolac tromethamine are

Gastrointestinal Effects - Risk of Wiceration, Bleeding and Perforation:
Ketorolac tromelhamine is contraindicated in patients with previously documented peptic utions and/or gastrointestinal (G) bleeding, Ketorolac tromelhamine can cause serious GI advantages and the contraint of the storolach, small intestine, or large intestine, which can be falsal. These serious adverse events can occur at any time, with or without symploms, in patients treated with learnonic bromelhamine.

The incidence and severity of gastro days.

In postmarketing experience, postoperative hemalomas and other signs of wound bleeding have been reported in association with the peri-operative use of intravenous or intramuscular dosing of lot tromethamina. Therefore, peri-operative use of keterolac tromethamine should be avoided and postoperative use be undertaken with caution when hemostasis is critical (see PRECAUTIONS).

Recal Effects of Text Section 1 (Text Section 1) to receive the received and other result signs, Result solicity has also been seen in patients in whom most prostaglanders have a compressable or the result signs of the result

Referoise fromethamine and its metabolites are eliminated primarily by the kickneys, which, in patients with reduced creatinine clearance, will result in deministed clearance of the drug (see CAINECAL PRAR-MADOLOGY). Therefore, ketoricals cornelmense should be used with causion in patients with impaired rured function (see DOSAGE AND ADMINISTRATION) and such patients should be followed closely With the use of selectric comerbamies, they have been reports of calcer resultable, inclinital and instruction condrows.

Impaired Renal Function Ketorolac tromethamine is con caution in patients with impairisk of developing acute renal

(see Curil INAMUNIATION SINC TRANSPORT TO THE CONTRIBUTION OF THE

To minimize the potential risk for an adverse CV event in INSAID-treated patients, use the lowest effective dose for the shortest duration possible. Physicians and patients should remain alert for the opened 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 if they occur.

Status Post Ceronary Artery Bypass Graft (CABS) Surgery
Two large, controlled clinical trials of a CDC-2 selective IRSIDI for the treatment of pain in the first 10 to 14 days following CABS surgery found an increased incidence of myor.
INSIGH zero-controlled on the setting of CABS surgery (see CONTRAMIDICATIONIS).

The control of the conducted in the Daniel National Registry have demonstrated that patients braided with NSADs in the post-All period were all recreased risk of reintertion. Oxived counter mortal programs in the first weet of treatment, in this same control, the incidence of odeshi in the first year post Milw was 20 per 100 proses years in RSADD opposed patients. Although the absolute rate of death declined somewhat after the first year post-MI, the increased relative risk of death in KSADD users per the next four years 1000-vsp.
Avoid the use of kstorotac fromethatismine in patients with a recent MI united somewhat after the first year post-MI, the increased relative risk of death in KSADD users per Avoid the use of kstorotac fromethatine in patients with a recent MI united somewhat after the first year post-MI, the increased relative risk of death in KSADD users per Avoid the use of kstorotac fromethatine risk years and the patients with a recent MI united patients for grant and the patients of the patients and the patients with a recent MI united patients for grant and the patients of the patients of the patients with a recent MI united patients for grant and the patients are patients and the patients and the patients are patients are patients are patients are patients are patients and the patients are patie

Hyperfension

ISSUE, including lettorate tromethamine, can lead to creat of new hyperfension or worsening of pre-existing hyperfension, either of which may contribute to the increased incidence of CV events. Patients thating histories or topo disvelocitic may have impaired response to these therapies when taking ISSUES. ISSUE), including hetroteic transferamine, should be used with caution in patients with hyperfension Blood pressure (RF) should be monitored closely during the initiation of ISSUE treatment and throughout the course of therapy.

Next Fallurs and Edenia
The Cobb and traditional SKRID Trialists' Collaboration meta-analysis of randomized controlled trials demonstrated an approximately two-fold two-treated patients and nonselective NSAID-treated patients compared to placebo-treated patients. In a Danish National Registry study of path 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 agmedical conditions (e.g., diuretics, ACE inhibitors, or angiotensin receptor blockers (ARBs) (see PRECAUTIONS – Drug Interactions).

Avoid the use of ketorolac tromethamine in patients with severe heart failure unless the benefits are expected to outweigh the risk of worsening heart failure. If ketorolac tromethe with severe heart failure, monitor patients for signs of worsening heart failure.

Drug Reaction with Existinghilla and Systemic Symptoms (DRESS) to the production of the production of

Disphydramnion-Neonatal Renal Impairment:
Use of NSAND, including ledorolac Comethamine, at about 20 weeks gestation or later in pregnancy may cause fetal renal dysfunction leading to oligophydramnios and, in some cases, neonatal renal men. These adverse conciones are seen, on average, after days to weeks of treatment, although oligophydramnion has been infrequently reported as soon as 48 hours after NSAD initiation. Oligophydramnion for the contractive reported as soon as 48 hours after NSAD initiation. Oligophydramnion for the contractive reported as soon as 48 hours after NSAD initiation. Oligophydramnion for the contractive and delayed lung maturation. In some postnesses of impaired monation after national renal function, indexes procedures such as exacting partnershown of algebys were required in delayed to report and an activity of the contractives and delayed lung maturation. In some postnesses of impaired monation after national renal function, indexes procedures such as exacting partnershown of algebys were required.

If INCAID treatment is necessary between about 20 weeks and 30 weeks gestation, limit ketorolar tromethamine use to the lowest effective dose and shortest duration possible. Consider ultrasour of ammidic fluid if ketorolar tromethamine freatment extends beyond 48 hours. Discontinue ketorolar bromethamine if oligohydramnios occurs and follow up according to clinical practice (see PR Preparancy).

cal activity of ketorolac tromethamine in reducing inflammation may diminish the utility of this diagnostic sign in detecting co

Hepatic Effects

the control of the

Hematologic Effects
contains seen in patients receiving NSADs, including betroites bornelhamine. This may be due to fluid intention, occult or gross GI blood loss, or an incompletely described effect upon exploracontains seen in patients receiving NSADs, including betroites bromethamine, should have their hemoglobin or humatoric checked if they eithed any signs or symptom of nemia ISADs intelligent
patient aggregation and have been shown to protong beterding them some patients. Using applicit transport or patient structure patients aggregation and have been shown to protong beterding their some patients. Using applicit transport or patient structure patients with the patients of the patients of the patient structure patients and patients with the patients with the patients with the patients and the patients of the patients are patients. Using applicit furnition is quantitatively less of administratively seed and patients are administratively seed and patients are administratively and the patients are administratively seed and patients are administratively seen and patients are administratively seen administratively seed and patients are administratively seed and patients are administratively seen and patie

Pre-existing fathma Platients with ashiman may have aspirin-sensitive ashima. The use of aspirin in patients with aspirin-sensitive ashima has been associated with severe bronchospasm which can be fatal. Since cross reactively, including bronchospasm, between aspirin and other nonstensicial anti-inflammatory drugs has been reported in such aspirin-sensitive patients, lestorolac bronethamine should not be administered to patients with his form of aspirin-sensitivity and should be used with caudin, in patients with pre-existing asthma.

and/or signs suggesting liver dysfunction, or in whom an abnormal liver test has occurred, should be evaluated for evidence of the development of a more severe hepatic re rolac tromethamine. If clinical signs and symptoms consistent with liver disease develop, or if systemic manifestations occur (e.g., eosinophilia, rash, etc.), ketorolac trometh

Information for Patients
Reformation for Patients

Physicians, when prescribing interrolar branching i

when observing any indicative sign or symptoms including exposition pages and express mediens, and hemalemenia. Patients should be apprised of the importance of this follow-up (see WARNINGS, Section.State Reactions. including DRISS Andreap Extens. State Reactions. including DRISS Andreap Extens. State States Reactions. Including DRISS Andreap Extens. States Reactions are considered for the States Reactions. Including DRISS Andreap Extens. States Reactions. Including DRISS Andreap Extens. States Reaction States Reac

WARMINGS.

It is instructed of the warring signs and symptoms of hepatotoxicity (e.g., nausea, faligue, lethargy, pruffus, jaundice, right upper quadrant tenderness, and "file-like" symptoms). If these exists chold be instructed to stop threship and seek immediate medical therapy.

Only the instructed for a large legal of an anaphylotycation (e.g., efficiently treating, swelling off the face or through.) If these occur, patients should be instructed to seek immediate emergency of the face or through. If these occur, patients should be instructed to seek immediate emergency

and women to avoid use of keterolac tromethamine and other NSAIDs starting at 30 weeks gestation because of the risk of the premature closing of the fetal ductus arteriosus. If historicals tromethamine is needed for a preparant woman between about 20 to 30 weeks gestation, advise her that she may need to be monitored for oligohydramnios, if treat longer than 48 hours (see WARHINGS: Fetal Toxicity, PRECAUTIONS, Pregnancy).

Drug Interactions
Ketrockis 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 metabolizing there

Laboratory Tests
Because serious Gifract ulcerations and bleeding can occur without warning symptoms, physicians should monitor for signs or symptoms of Gi bleeding. Patients on long-term treatment with INSADs, should be the their CRG and a chemistry profile checked periodically. If clinical signs and symptoms consistent with liver or renal disease develop, systemic manifestations occur (e.g., eximpoliia, rash etc.) or if abnormal liver tests persist or worsen, ketorolac tronethamine should be discontinued.

The in vito barding of warfarin to plasms proteins is only slightly reduced by laterolar bronthamine (96.5% control in 99.3%) when leterolar plasms encentralized on a control protein barding in the dutudus indicated that at therepastic connectations of salicylate (200 mog/mit,) the binding of leterolar was reduced from the control protein barding in the dutudus indicated that at therepastic connectations of salicylate (200 mog/mit,) the binding of leterolar was reduced from the control protein barding in the control protein barding in the deteroids the control increase in unbound deteroids, plasma levels. Therepastic concertations of digestin, warfarin, theyerefor, napresen, plasma levels. Therepastic concertations of digestin, warfarin, theyerefor, napresen, plasma levels.

In a study involving 12 adult volunteers, oral ketorolac tromethamine was coadministered with a single dose of 25 mg warfarin, causing no significant changes in pharmacokinetics or pharmacodynamics warfarin in another study, ketorolac bromethamine doced intravenous or intramuscular was given with two doses of 5000 U of Appartis 11 healthy volunteers, resulting in an eman emplaie beforeing fine intermed and 60 mitized (64 or 25 mill not pleage into an ead 5. mitized). 50 to 8.m mill by relaxed, on Although these results on or indicate a significant interact between ketorolac tromethamine and variation or begain, the administration of storolac bromethamine is patients taking anticoagularits should be dose extremely cautiously and patients should be dose montred (see Waddiness our Precautions). — Healtablegic Effects).

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

Aspairs
When heterotics bromethamine is administered with appire, its protein briding is reduced, although the clearance of free heterotics tromethamine is not altered. The clinical significance of this interaction has been been also also although the clearance of free heterotics tromethamine is not altered. The clinical significance of this interaction has been been also although the clearance of free heterotics tromethamine is not appreciately recommended because of the potential of increased adverse effects.

Districtions. Districtions are selected as a postmarketing observations, have shown that kelorolac tromethamine can reduce the natifuratic effect of furosemide and thiazides in some patients. This response has been attributed to inhibition of renal prostagation synthesis. 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 during efficacy.

Probenecid

Concomitant administration of oral lebtorolac tromethamine and probenecid resulted in decreased clearance and volume of distribution of Nettorolac and significant increases in Nettorolac pissma levels (bital ML increased approximately three-lold from 5.6 to 15.1 hours. Therefore, concomitant use of Nettorolac tromethamine and probenecid is contraindicated.

Lithium

INSUDs have produced an elevation of plasma lithium levels and a reduction in renal lithium dearance. The mean minimum lithium 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 INSUD. Thus, when INSUDs and lithium are administered concurrently, subjects should be observed carefully for signs of lithium busicity.

Methotrexate
NSAUs 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 NSAUs are administered concomitantly with methotrexate.

ACE Inhibitors/Angiotensin II Receptor Antagonists
Corcomitant 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 INSAIDs may diminish the antihypertensive effect of ACE inhibitors and/or angiotensin II receptor antagonists. This interaction should be given consideration in patients taking INSAIDs concomitantly with ACE inhibitors and/or angiotensin II receptor antagonists. This interaction should be given consideration in patients taking INSAIDs concomitantly with ACE inhibitors and/or angiotensin II receptor antagonists.

Antiepilieptic Drugs

Sporadic cases of seizures have been reported during concomitant use of ketorolac tromethamine and antiepileptic drugs (phenytoin, carbam Psychoactive Drugs
Hallucinations have been reported when ketorolac tromethamine was used in patients taking psychoactive drugs (fluoxetine, thiothixene, alprazolam).

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

Hondepolarizing Muscle Relaxants
In postmatching experience them have been reports of a possible interaction between velorotics transitioning intravenous/intramuscular and nondepolarizing muscle relaxants that resulted in apnea.
The concurrent use of behavior it branchismine with muscle relaxants has not been formally studied.

Selective Serotenin Reuptake Inhibitors (SSRIs)
There is an increased risk of gastrointestinal bleeding when selective serotenin reuptake inhibitors (SSRIs) are combined with NSAIDs. Caudion should be used when NSAIDs are administrative with SSRIs.

Carcinogenesis, Mutagenesis, and Impairment of Fertility
An 18-month study in mice with oral doses of National consellamine tablets at 2 mg/kg/day (0.9 times the human systemic exposure at the recommended intramuscular or intravenous dose of 30 mg qid, based or are-under-the-plasmes-concentration curve (NUC), and a 24-month study in rate at 5 mg/kg/day (0.5 times the human AUC) showed no evidence of humorigenicity.

Ketorolac tromethamine was not mutagenic in the Ames test, unscheduled DNA synthesis and repair, and in forward mutation assays. Ketorolac tromethamine did not cause chromosome breakage in the in vivo mouse micronucleus assay. At 1590 mog/mL and at higher concentrations, ketorolac tromethamine increased the incidence of chromosomal aberrations in Chinese hamster ovarian 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 trometha

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Premature Closure of Fetal Ductus Arteriosus
Use of NSAIDs, including ketorolac tromethamine, at about 30 weeks gestation or later in pregnancy increases the risk of premature closure of the fetal ductus arteriosus.

Oligohydramnios/Neonatal Renal Impairment
Use of NSAIDs at about 20 weeks gestation or later in pregnancy has been associated with cases of fetal renal dysfunction leading to oligohydramnios, and in some cases, neonatal renal impairment.

Data from observational shadies regarding other potential embryofetal risks of NSAID use in women in the first or second trimesters of preguancy are inconclusive. Reproduction studies have been preformed dura organogenesis using daily and sloses of sebrosiac bromehamine at 3.5 mg/kg (0.37 films the human AUL) in abilitias and at 10 mg/kg (1.07 films the human AUL) in install, as the human AUL prints. Results of these satisfies dislorate reveal relative of characteristic studies and the contraction of characteristic studies and the contraction of the contract

Clinical Considerations

Fetal/Neonatal Adverse Rea

Premature Closure of Fetal Ductus Arteriosus

Avoid use of NSAIDs in women at about 30 weeks gestation and later in pregnancy, because NSAIDs, including ketorolac tromethamine, can cause premature closure of the fetal ductus arts WARNINGS, Fetal Toxicity.

If an NSAID is necessary at about 20 weeks gestation or later in pregnancy, limit the use to the lowest effective dose and shortest duration possible. If ketorolac tromethamine treatment extends beyond 48 hours, consider monitoring with ultrasound for oligohydramnios. If oligohydramnios occurs, discontinue ketorolac tromethamine and follow up according to clinical practice (see WARNINGS; Fetal Toxicity).

Data
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Himano Data
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

Fremature Closure of Fetal Ductus Arteriosus:
Published illerature reports that the use of NSAIDs at about 30 weeks of gestation and later in pregnancy may cause premature closure of the fetal ductus arteriosus.

Oligohydramnico Necontal Renal Impairment:
Publish of Studies and postmarking reports describe maternal NSAID use at about 20 weeks gestation or later in pregnancy associated with fetal renal dysfunction leading to oligohydramnics, and in some success, necontail renal impairment. These adverse outcomes are seen, on average, after days to weeks of treatment, although dispinydramnics has been infrequently reported as soon as 45 hours after NSAID initiation. In many cases, but not all, the decrease in amnioch fluid was transient and reversible with cessation of the drug. There have been a limited number of case reports of maternal INSAID use and necessation are recorded in the control without displayments, somed of which were inversible. Some cases of necessating of students or expert of teachers with invester procedure, such as exchange frantisation.

Methodological limitations of these postmarketing studies and reports include lack of a control group; limited information regarding dose, duration, and timing of drug exposure; and concomitant use of other medications. These limitations proclude establishing a reliable estimate of the risk of adverse field and necestal outcomes with maternal NSAD use. Because the published safety data on necestal outcomes involved mostly preferring infants, the generalizability of certain report of risks to the full-terminiant exposed to NSADs through maternal uses uncertain.

Labor and Belivery
The use of substrate brenshamine is contraindicated in labor and delivery because, through its prostaglandin synthesis inhibitory effect, it may adversely affect fetal circulation and inhibit use the increasing the rick of uterior immortage (see CONTINUMENTINES).

Effects on Fulfilly
The use of ketorolac transitions, as with any drug known to inhibit cyclopogenase/ prostaglandin synthesis, may impair fertility and is not recommended in women attempting to conceive. In won tave difficulty conceiving or are undergoing investigation of infertility, withdrawal of ketorolac transitionaries about the considered.

Nursing Mothers

Limited data from one published dately that included 10 brassification growner 2-6 days postparfum aboved low levels of indecidac in brassification and under the control of the control

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 healthcare provider if they note any adverse events. Pediatric Use
Retorolac tromethamine is not indicated for use in pediatric patients. The safety and effectiveness of ketorolac tromethamine in pediatric patients below the age of 17 years have not been established.

istatic Use, CleST Years of Ago)
axis determine many be desered more slowly by the elderly (see CLINICAL PHARMACOLOGY) who are also more sensitive to the dose-related adverse effects of ISAIDs (see WARMINGS— troinisestated Effects—Risk of Ulceration, Bleeding, and Perforation), extreme caution and reduced dosages (see DOSAGE AND ADMINISTRATION) and careful clinical monitoring must be used in restaint the elderly with lactings to tenderly and prefer and the prefer and

ADVERSE REACTIONS

were maction rates increase with higher dosse of leterorise tromethamine. Practitioners should be start for the severe complications of treatment with leterorise, tromethamine, such as G.I. ulcoration, endering and perfortantly complement producers and a result failure, manaphylactic and anaphylactical reactions and liver failure (see Beade Walmille, Walmille, Walmille, Specially), and DSAGE AND AD-WISTRATION, These ItS-AD-related complications can be serious in certain patients for whom relatorise tromethamine is indicated, especially when the drug is used inappropriately.

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) experiences including:			
abdominal pain	abdominal pain	abdominal pain	
flatulence	GI fullness	Gl ulcers (gastric/duodenal)	
gross bleeding/perforation	heartburn	nausea*	
stomatitis	vomiting		

abnormal renal function	anemia	dizziness
drowsiness	edema	elevated liver enzymes
headaches*	hypertension	increased bleeding time
injection site pain	pruritus	purpura
rashes	tinnitus	sweating

Additional adverse experiences reported occasionally (<1% in patients taking ketroriac tromethamine or other NSADs in clinical triats) include:

Body as a Whote: Ever, infection, sepsis

Cardiovasculara competive heart failure, palpitation, pallor, tuchycardia, syncope

Dermatologie: alopoca, photosensitivity, uriticaria

Castrollandania erroria, dy nonder, uncutation, esphagitis, excessive thirst, gashifis, glossifis, hematemesis, hepatitis, increased appetite, jaundice, melena, rectal bleeding

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Table 3: Incidence of Clinically Serious G.I. Bleeding as Related to Age, Total Daily Dose, and History of G.I. Perforation, Ulcer, Bleeding (PUB) after up to 5 Days of Treatment with Ketorolac Tromethamine Injection

Total Baily Bose of Keterolae Tremethamine Injection

Age of Patients					
age of Patients	≤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%	

	Total Daily Dose of Ketorolac Tromethamine Injection					
	Age of Patients	≤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%	25.0%	

## To report SUSPECTED ADVERSE REACTIONS, contact Caplin Steriles Limited at 1-866-978-6111 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

OVERIOSACE Symptoms and Signs Symptoms following acute NSADs overdoses are usually limited to lethargy, drowsiness, nausea, ventiling, and epigastric pain, which are generally reversible with supportive care. Gastrointestinal bleeding can occur. Hyperference, acute renal failure, respiratory depression and coma may occur, but are rare. Anaphylactoid resctions have been reported with therapeutic ingestion of NSADs, and may occur following an evention.

Patients should be managed by symptomatic and supportive care following a NSADs overdose. There are no specific antidotes. Forced diuresis, alkalization of urine, hemodialysis or hemoperfusion may not be useful due to high protein binding. Single overdoses of ketorolac branethamine have been variously associated with abdominal pain, nausea, vomiting, hyperventitation, peptic ulcers and/or erosive gastritis and renal dysfunction which have resolved after discontinuation of dozing.

Total duration of treatment in adult patients: the comb ined duration of use of intravenous or intramuscular dosing of ketorolac tro nine and oral ketorolac tron

ENTRODUCE TROMBET RAMME BLECTION

Ketronica tromethramine injection may be used as a single or multiple dose on a regular or "as needed" schedule for the management of moderately severe, acute pain that requires analgesia at the opioid very usually in a postportaine stricture, leverorisms should be corrected prior to the administration of ketronic tromethramine (see WARNINGS - Benat Effects). Patients should be switched to alternative analgesics as soon as possible, but ketronics tromethramine therapy is not to exceed 5 days.

When administering ketorolac tromethamine injection, the intravenous bobs must be given over no less than 15 seconds. The intramuscular administration should be given slowly and deeply into the The analgesic effect begins in ~30 minutes with maximum effect in 1 to 2 hours after dosing intravenous or intramuscular. Duration of analgesic effect is usually 4 to 6 hours.

### Single-Dose Treatment: The following regimen should be limited to single administration use only Intramuscular Dosino

Patients <65 years of age: One dose of 60 mg.
 Patients ≥65 years of age, renally impaired and/or less than 50 kg (110 lbs) of body weight: One dose of 30 mg.

Patients <65 years of age: One dose of 30 mg.
 Patients ≥65 years of age, renally impaired and/or less than 50 kg (110 lbs) of body weight. One dose of 15 mg.

the Pape Transferred (Indicements or Informatical Professor Section 2014). The Contract of Contract Co

For breakfirrough pain, do not increase the dose or the frequency of ketorolac tromethamine. Consideration should be given to supplementing these regimens with low doses of opioids "as needed" urtherwise contraindicated.

eutical Information for Ketorolac Tromethamine Injection fromethamine injection should not be mixed in a small volume (e.g., in a syringe) with morphine sulfate, meperidine hydrochloride, pro in precipitation of betworks from solving.

teral drug products should be inspected visually for particulate matter and discoloration prior to administration, whenever solo

NDC	Strength of product	Package Factor
65145- 146-25	15 mg/mL Single-dose vial	25 Vials per Carton
65145- 145-25	30 mg/mL Single-dose vial	25 Vials per Carton
65145-147-25	60 mg/2 mL (30 mg/mL) Single-dose-vial	25 Vials per Carton

Store at 20 to 25°C (68 to 77°F). [See USP Controlled Room Temperature.]

Protect from light.

Retain in carton until time of use

Made in India Distributed by : Caplin Steriles USA Inc,

Hamilton, NJ 08619.

Revised: 05/2024 Medication Guide for Non-Steroidal Anti-Infla

What is the most important information I should know about medicines called Non-Steroidal 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 in

with increasing doses of NSAIDs
 with longer use of NSAIDs

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 HSAIDs after heart attack.

Increased risk of bleeding, ulcers, and tears (perforation) of the esophagus (tube leading from the mouth to the stomach), stomach and intesti

Do not take NSAIDs right before or after a heart surgery called a "coronary artery bypass graft (CABG)."

The risk of getting an ulcer or bleeding increases with: rssk of getting an ulcor or bleeding increases with:

past history of stomach ulcars, or stomach or intestinal bleeding with the use of NSAIDs staking medicines called "confcosteroids", "anticoagulants", "SSRIs", or "SNRIs" increasing doses of NSAIDs increasing doses of NSAIDs smoking and the stake of t

exactly as prescribed
 at the lowest dose possible for your treatm
 for the shortest time needed

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?

if you had an asthma attack, hives, or other allergic reaction with aspirin or any other NSAIDs.
 ight before or after heart bypass surgery.

Before taking NSAIDs, tell your healthcare provider about all of your medical conditions, including if you

have liver or kidney problems have high blood pressure have asthma an programed or gian to become programt. Taking ISAIDs at about 20 seeks of programory or later may harm your unborn baby. If you need to babe ISAIDs for more than 2 days when you are between 20 and 30 seeks of programory, your healthcare provider may need to monitor the amount of fluid in your womb around your baby. **You should not take ISAIDs after about 30 weeks** of **programory**.

are brantlesteding 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 supp 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? NSAIDs can cause serious side effects, includir

See "What is the most important information is about know about medicines called Nonsteroidal Anti-Inflammatory Drugs (NSAIDs)?"

new or worse high blood pressure
heart failure
less reproblems including kiner failure
less reproblems reproblems including kiner failure
less reproblems mergency help right away if you get any of the following symptoms: shortness of breath or toucle breathing chest pain chest pain could be reading chest pain weakness in one part or side of your body storned greath swelling of the face or throat

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

nausea more tired or weaker than usual diarrhea itching

diamness

your sikin or eyes look yellow
indigestion or stomach pain
the-like eyentions
want blood

of like year blook yellow
indigestion or stomach pain
twent blood

of like year bowel movement or it is black and sticky like tar
unusual weight glant
sikin rash or blistens with fever

overling of the arms and legs, hands and feet

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 phar Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

Other information about NSAIDs

Appire is an NSAID medicine but it does not increase the chance of a heart attack. Appire can cause bleeding in the brain, stomach, and intestines. Appire can also cause ultimistations.
Some ISAIDs are sold in lower doses without a prescription (over-the-counter). Talk 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 Medication Guide. Do not use NSAIDs for a condition for which it was not prescribed. Do not 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 can ask your pharmacist or healthcare provider for information about NSAIDs that is written for health pro

This Medication Guide has been approved by the U.S. Food and Drug Administration Made in India

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