MICHLIGHTS OF PEESCRIBBING MOTORMATION
HERE highlights for on Included at the information needed to use DICLOFENAC POTASSI
DELECTION OF THE PROPERTY OF THE PR

- See full prescribing information for complete boxed warning

 Non-steroidal anti-inflammatory drugs (IRSADs); cause an increased risk of serious cardiovascular thrombotic events, including myocardal infarction and stroke, which can be fatal. This risk may occur early in treatment and may increase with duration of use [5.1]

- of use (C.)

 Dictionacy postsssium for oral solution is contraindicated in the settling of coronary Dictionacy postssium for oral solution is contraindicated in the settling of coronary NSADs cause an increased risk of serious gastrointestinal (IQ) adverse events coulding blacked), clearation, and perforation of the science of verification, which could be considered to the contraint of the cont

Varnings and Precautions (5.10, 5.12) 4/2021

Weenings and Precautions (5.10, 5.17) 40031

Weenings and Precautions (5.10, 5.17) 40031

BOX (ATRIONS AND USAGE

ADD USAGE CONTROL OF THE ADD USA

DOSAGE AND ADMINISTRATION

Single 50 mg dose; mix single-packet contents with 10 20 ourses or 20 4 bablespoons (30 to 60 mt.) of water prior to adm

• Use the lowest effective dose for obnities disuration consistent with individual patient treatment goals

(2.1)

- DOSAGE FORMS AND STREMONS
 PROME Each containing billmed declarace potentials in a scaleby powder (3)
 Product Each containing billmed declarace potentials in a scaleby powder (3)
 Former Ingeneratively to declarace or Relative or size or specific for the one product (4)
 History or dambnia, unificative, or their allegic-type reactions after taking aspirin or other NSADD (4)
 In no secting or Relative Outpurply (4)

- **DOES DESIGNATION OF THE PROPERTY OF THE PROP

- patients, monitor for signs or worsening renal function (7)

 <u>Diurelies</u>: NADIS can reduce nativirative direct of loop and thiazide diuretics. Monitor patis diuretic efficacy including antihypertensive effects (7)

 <u>Digonin</u>: Concomitant use with dictofenac potassium for oral solution can increase serum concentration and prolong half-life of digonin. Monitor serum digonin levels (7)

USE IN SPECIFIC POPULATIONS Infertility: NSAIDs are associated with reversible infertility. Consider withdrawel of diclofenac potassium for oral solution in women who have difficulties conceiving (8.3)

- FULL PRESCRIBING INFORMATION: CONTENTS*
 WARNING: HISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL
 1 INDICATIONS & USAGE
 2 DOSAGE & ADMINISTRATION
 2 DOSAGE & ADMINISTRATION
 3 DOSAGE FORMS & STREAMTHS
 3 DOSAGE FORMS & STREAMTHS
 3 DOSAGE FORMS & STREAMTHS
 3 LOCAL PROPERTY OF THE STREAMTHS
 5 LOCAL PROPERTY OF THE STREAMTHS
 5 LOCAL PROPERTY OF THE STREAM OF THE STREAMTHS
 5 LOCAL PROPERTY OF THE STREAM OF TH

- Hepatotoxicky
 Hypertexion
 Hypertexion
 Renal Toxicky and Hyperkalem
 Renal Toxicky and Hyperkalem
 Excercised Hyp

- 5.15 Laboratory Monitoring
 ADVERSE REACTIONS
 6.1 Clinical Trials Experience
 6.2 Postmarkers
- 6.2 Postmarketing Experience
 7 DRUG INTERACTIONS
 8 USE IN SPECIFIC POPULATIONS
- Pregnancy Lactation Females and Males of Repr Pediatric Use

- Renal Impairment OVERDOSAGE DESCRIPTION CLINICAL PHARMACOLOGY

- I Bechnism of Action
 I Bechnis
- 13.1 Carchogenesis & Mutagenesis & Impairment un Folksy
 14 CLNICAL STUDIES
 16 HOW SUPPLIED/STORAGE AND HANDLING
 17 PATIENT COUNSELING INFORMATION
 Sections or subsections omitted from the full prescribing information are not listed.

FULL PRESCRIBING INFORMATION

WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS

- ardiovascular Thrombotik Events
 Nonsterodal anti-inflammatory 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 Jee Warnings and Precautions (5.1).
- Diciofenac potassium for oral solution is contraindicated in the setting of coronary artery bypass graft (CABG) surgery [see Contraindications (4) and Warnings and Precautions (5.1)].

Gastrointestinal Bleeding, Ulceration, and Perforation

NSAIDs cause an increased risk of serious spatrointential (c) adverse events
including beeding, userablos, and perforation of the stomach or intentines,
which is the property of the stomach of the stomach of the stomach
warning symptoms. Elderly patients and patients with a prior history of peptic
user disease and/or Gi bleeding are at greater risk for serious GI events (see
Warning and Precautions (2.2).

Dictofenac potassium for oral solution is indicated for the acute treatment of migraine attacks with or without aura in adults (18 years of age or older)

- <u>Limitations of Use</u>
 Dictofenac potassium for oral solution is not indicated for the prophylactic therapy of
- migraine.

 The safety and effectiveness of diclofenac potassium for oral solution have not been established for cluster headache, which is present in an older, predominantly male

2.1 Acute Treatment of Migraine

Administer one packet (50 mg) of dicofenac potassium for oral solution for the acute treatment of migraine. Empty the contents of one packet into a cup containing 1 to 2 ounces or 2 to 4 tablespoons (30 to 60 mL) of water, mix well and drink immediately.

Taking diclofenac potassium for oral solution with food may cause a reduction in effectiveness compared to taking diclofenac potassium for oral solution on an empty stomach [see Clinical Pharmacology (12.3)].

Use the lowest effective dose for the shortest duration consistent with individual patient treatment goals. The safety and effectiveness of a second dose have not been established

2.2 Non-Interchangeability with Other Formulations of Diclofenac

Different formalistics of our dichitense (e.g., dichitense, potassium for oral solution, dichitense sodium enter-coalent abless, di briense sodium enterded-relesse tablets, or dichitense, potassium immediate-relesse tablets, or dichitense, potassium immediate-relesse tablets) appro to be bioquivalent even if the miligram strength is the same. Therefore, it is not possible to convert declaring from any other formalistion of dichitense to dichitense potassium for oral declaring from any other formalistion of dichitense to dichitense potassium for oral

Dictofenac potassium for oral solution, USP is available in individual packets each designed to deliver a 50 mg dose when mixed in water.

CONTRAINDICATIONS

Dictionar polisation for oral solution is contraindicated in the following patients:

• Known hypersensibility (e.g., sinaphylactic reactions and serious shit nections) to

• Known hypersensibility (e.g., sinaphylactic reactions and serious shit nections) to

• Known by Company (e.g., sinaphylactic reactions and serious shit stating apprin or

• Hattory of authma, urticatin, or other alterglo-cippe reactions after taking apprin or

reported in just patients [see Warrings and Precautions (5.7.5.8])

• In the setting of coronary artery bypass graft (CABC) surgery [see Warrings and

Precautions (5.7.4.8)]

5.1 Cardiovascular Thrombotic Events

Clinical trials of several COX-2 selective and nonselective NSAIDs of up to three years duration

duration
have shown an increased risk of serious cardiovascular (CV) thrombotic events, including myocardial infarction (MI) and stroke, which can be fatal. Based on available data, it is unclear that the risk for CV thrombotic events is similar for all INSAIDs. The relative increase in serious CV.

thrombotic events is similar for all KSAIDs. The relative increase in serious CV thrombotic events in the control of the contr

uld remain alert for the development of such events, throughout the entire treatment course, even in the absence of previous CV symple

of treatment. The increase in CV thrombotic risk has been observed most consistently at higher doses. To minimate the potential risk for an adverse CV event in NSADD. treated patients, use the lowest effective dose for the shortest duration possible. Physicians and patients of Patients should be informed about the symptoms of serious CV events and the steps to take if they occur. There is no consistent evidence that concurrent use of asprin mitigates the increased risk of serious CV immotiber events associated with NSADD use. The concurrent use of septim and an NSADD, such as of witherest, processes the risk of serious gastronistical (GI) events [see Warnings and Precautions (S.21)]

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

14 days following CABG surgery found an increased incidence of myocardial infarction and stroke. NSAIDs are contraindicated in the setting of CABG Isee Contraindications (4)1.

Post-MI Patients

Post-ML Distincts.

Observational studies conducted in the Danish National Registry have demonstrated that patients treated with NSAIDs in the postBerd were all interested risk of refaint-cition. CV-related death, and all-cause mortality is produced in the restriction of the relation of the restriction of the rest

up.

Avoid the use of diclofenac potassium for oral solution
in patients with a recent full unless the benefits are expected to outweigh the
risk of recurrent CV thrombotic events. If diclofenac potassium for oral
solution is used in patients with a recent MI, monitor patients for signs of cardiac ischemia

soution is used in patients with a recent M, monitor patients for signs of cardiac ische 5.2 Gastrointestinal Bleeding, Ukceration, and Perforation NSADS, including disclerance, cause serious gastrointestinal (GI) adverse events (GI) and the serious consistency of the serious serious serious serious and intestine, or large intestine, which can be falls. These serious adverse events can corcur at any time, with or without warming symptoms, in patients treated with NSADS. Only one in five patients wind election as et cause upper GI adverse event on the SADS occurred in approximately 3% or patients treated or 3.6 minorities, and in about 2%-4% of patients treated for one year. However, even short-term NSADD therapy is not without these.

Risk Factors for GI Beeding, Ukeration, and Perforation
Patients with a prior history of peptic user disease another GI bleeding who used NSAIDs
had a greater than Developing, a GI belief compared to patients, without these risk factors. Other factors
duration of NSAID these processes of the developing in patients in heads with NSAIDs schule longer
duration of NSAID thesepy concombating use of oral controllectorids, apprint,
anticoagulants, or selective serrorionin respitate inhibitors (SSRI); smoking; use of a
action of the processes of the processes of the developing of the devel

- Strategies to Minimize the GI Bisk in NSAID-treated patients:

 Use the bwest effective dosage for the shortest possible duration.

 Avoid use in patients at higher risk unless benefits are expected to outweigh the bleeding, consists on former than one NSAID at a time.

 Avoid use in patients at higher risk unless benefits are expected to outweigh the bleeding, consists afternate threates on the than NSAID.

 Area and a the time of ti

Elevations of one or more liver tests may occur during therapy with dictofenac potassium for oral solution. These laboratory abnormalities may progress, may persist, or may only be transient with continued therapy. Borderine devotators (less than 3 times the upper limit of the normal (UNI) range) or greater devotators of transammates occurred in about 15% of citotherac. ALT (SGPT) is recommended for the monitoring of liver injury.

In clinical trials, meaningful elevations (i.e., more than 3 times the ULN) of AST (SGGT) occurred in about 2% of approximately 5,700 patients at some time outing treatment of the control of the contro

Abnormal tests occurred during the first 2 months of therapy with diciofenac in 42 of the 51 patients in all trials who developed marked transamiase devalors. In postmarketing reports, cases of draining-induced heaplotable (by heap been reported in the first fundin, and in some cases, the first 21 months of a MSAOL therapy, but can occur at any time during treatment with diciofenac.

Postmarketing surveillance has reported cases of severe hepatic reactions, including liver

sis, jaundice, fulminant hepatitis with and without jaundice, and liver failure. Some of these reported cases resulted in fatalities or liver transp

Inform patients of the warning signs and symptoms of hepatotoxicky (e.g., nauses, fatigue, lethargy, diarrhea, prurtus, jaundice, right upper quadrant tenderness, and Tulke symptoms. It critical signs and symptoms consistent with Net disease develop, or if systems manifestations occur (e.g., existophila, rash, etc.), discontinue dictions of before control of the c

To minimize the potential risk for an adverse liver:

In minimize the potential risk for an adverse liver:

In the property of the property of

5.4 Hypertension

NSAIDs, including dicbfenac potassium for oral solution, can lead to new onset of hypertension or worseling of pre-existing hypertension, either of which may contribute to the increased incidence of CV events. Use NSAIDs, including dicbfenac potassium for oral

oral solution, with caution in patients with hypertension. Monitor blood pressure closely during the initiation of NSAID treatment and throughout the course of therapy.

Patients taking angiotensin converting enzyme (ACE) inhibitors, thiazides, or loop diuretics may have impaired response to these therapies when taking NSAIDs [see Drug Interactions (7)].

reatents stemp arrigorerant converting enzyme (ALL; minitions), intercoloe, or topp autret. S.S. Heart Failure and Edema The Coxts and treatitional MSAID Trialists: Collaboration meta-analysis of randomized controlled trials demonstrated an approximately flow-fold increase in hospitalizations for heart failure in COX2 selective- treated patients, and nonselective MSAID-treated patients, compared to pixecto-treated patients, in a Danish National Registry study of patients with heart failure, NSAID use increased the risk of MJ, hospitalization for heart failure, and detain.

Additionally, fluid retention and edema have been observed in some patients treated with NSAIDs. Use of dicibif may bunt the CV effects of several therapeutic agents used to treat these medical conditions (e.g., duretics, ACE inhibitors, or angiotensin receptor blockers [ARBB] [see Drug Interactions (7)].

Avoid the use of diciofenac potassium for oral solution in patients with severe heart failure unless the benefits are expected to outweigh the risk of worsening heart failure. If diciofenac potassium for oral solution is used in patients with severe heart failure.

5.6 Renal Toxicity and Hyperkalemia

Read Toxicky

Long-term administration of NSAIDs has resulted in renal papallary necrosis and other renal signsy. Renal toxicky has also been seen in patients in whom renal prostaglandins have a compensation of the renal renal prostaglandins have a compensation of the renal performance of renal performance of renal performance of the performance of the

No information is available from controlled clinical studies regarding the use of dicible potassium for oral controlled clinical studies regarding the use of dicible solution in patients with silvaniced renal disease. The renal effects of diciblenac solution in the controlled controlled in the controlled controlled in the controlled may hasten the progression of renal dysfunction in patients with pre-cesting renal disease.

rest valume status in dehydrated or hypovolemic palients prior to initiating dichlemac sastum for oral solution, Norbinz renal function in palents with versal or hepatic airment, heart falure, dehydration, or hypovolemia during sue of dichlemac sastum for oral solution in see Drug inferencies (7)1, Avoid the sue of dichlemac sastum for oral solution.

potassium for oral solution is used in patients with advanced renal disease, monitor patients for signs of worsening renal function.

S.7. Aniaphysecus Necessary Diciofenac has been associated with anaphylactic reactions in patients with and without known hypersensitivey to diciofenac and in patients with aspirinsensitive asthma [see Contraindications (4) and Warnings and Precautions (5.8)].

Seek emergency help if an anaphylactic reaction occurs.

5.8 Exacerbation of Asthma Related to Aspirin Sensitivity

5.8 Exacerbation of Asthma Related to Aspirin Sensithing A subsponding of pointers with asthma may have aspirin-sensible asthma which may include chronic rhinocinustic completed by nasi polyps: severe, potentially fatal brunchospapur, andien for indefense case against and other Isskilds). Because cross-term control of the control o

5.9 Serious Skin Reactions

Diclofenac potassium for oral solution is contraindicated in patients with previous serious skin reactions to NSAIDs [see Contraindications (4)].

Solution (acceptance of the Configuration with Ecological and Systems: Symptoms (DRESS)

Drug Reaction with Ecologicals and Systems: Symptoms (DRESS) has been reported in patients sking (Nation can be configurated production and systems: Symptoms (DRESS) has been reported in patients sking (Nation can desirable production production). Some of these events have been flaid of life threadening, DRESS should be controlled to the configuration of the controlled production. Some of these events have been flaid of life threadening, DRESS should be controlled by the controlled production of the co

Sometimes symptoms of DRESS may resemble an acute viral infection. Eosinophilia is Sometimes symptoms of DRESS may resemble an acute viral infection. Eosinophilia is present. Because this disorder is variable in its presentation, other organ systems not noted here may be involved. It is important too note that early mainfectations of hypersemblyby, such as fever or lymphadenopathy, may be present even though rash is not evident. If such signs or symptoms are present, discording decidence, postsonal more for all solution and evaluate the patient immediately.

and evaluate the pattent mineralizery.

5.11 Medication Oversize Neadache

Diversize of acute migratie drugs (e.g., egglarmine, triptans, oppids, nonsteroidal antiinfilmmatory drugs or combination of these drugs for 10 or more days per month) may

keal to exacerbation of

lead to exacerbation of

gradient of the exacerbation of

gradient of the exacerbation of

leading and treatment of withdrawal symptoms (which often includes a

treatment exacerbation of headached my on the exacerbation

leading of the exacerbation of the exacerbation

leading of the exacerbation of the e

5.12 Fetal Toxicity

ure Closure of Fetal Ductus Arteriosus

Dispolyuframnios.Neonatal Benal Impairment.

Use of NSAIDs, including discidence potassismif for oral solution, at about 20 weeks gestation of late in preparacy may cause preliation or late in preparacy may cause offices and, in some cases, neonatal renal impair are seen, on average, after days to weeks of treatment, although olipsylyramnios has been infrequently reported as soon as 44 hours after NSAID relation.

Opplyadramos is done, but not always, cree-sible with treatment discontinuation. Complications of prohipment or p

If NSAID treatment is necessary between about 2.0 weeks and 3.0 weeks gestation that dischlering potassium for ord solution use to the tweets effective does and solu-darding possible. Consider utrascound monitoring of ammittic fluid if discherine, potassium for ord solution treatment extends beyond 46 hours. Discontinue dicherine, potassium for ord solution folgohydramnios occurs and follow up according to clinical practice (see Line Specific Population (8.1)).

5.13 Hematologic Toxicity

Anemia has occurred in NSAID-treated patients. This may be due to occult or gross blood lass,

fluid retention, or an incompletely described effect upon erythropoiesis. If a patient try potassis od loss, of retention, or an incompletely described effect upon erythropolesis. If a patient treated with diciblenac assium for oral solution has any signs or symptoms of anemia, monitor hemoglobin or hematocrit.

NSAIDs, including dicbfenac potassium for oral solution, may increase the risk of bleeding events. Concomitant use of warfarin and other anticoagulants, antiplatelet agents (e.g., aspirin), and serotonin reuptake hibitors (SSRIs) and serotonin norespications (SRIs) may increase this risk. Monitor these patients and any patient who may be adversely affected by afterations in platelet function for signs of bleeding [see Drug Inferactions (7)].

5.14 Masking of Inflammation and Fever

The pharmacological activity of diclofenac potassium for oral solution in reducing inflammation, and possibly fever, may diminish the utility of diagnostic signs in detecting infections

5.15 Laboratory Monitoring Because serious Gl bleeding, hepatotoxicity, and renal injury can occur without warning symptoms or signs, consider monitoring patients on long-term NSAID treatment with a CBC and a chemistry profile periodically [see Warnings and Precautions (5.2, 5.3, 5.6)].

Discontinue diclofenac potassium for oral solution if abnormal liver tests or renal tests persist or worsen

6 ADVERSE REACTIONS

6 ADVERS RACTIONS
The following sortius alverse reactions are discussed in greater detail in other sections o
Cardiovascular Thrombotic Events (see Warnings and Precautions (5.11)

1 elegations (1) elegation and Perforations (5.21)

1 elegations (1) elegations and Precautions (5.21)

1 elegations (1) eleg

• Hematobytic Toxickly (see Warmings and Precaudition (Laura)

Because chical tribs are conducted under widely varying conditions, adverse reaction rates observed in the circuit stribs are conducted under widely varying conditions, adverse reaction rates observed in the circuit stribs of a fung cannot be directly compared with rates in the chical tribs of another drug and may not reflect the rates observed in practice. In the circuit stribs of a fung cannot be directly contained in for oral solution was evaluated in 2. Perfection controlled risks with a total of 64 furnity are patients and places of the circuit risks with a total of 64 furnity are patients and with circliness potalssism leicher discharge potalssism or consistent in the circuit of the structure of the structure

The most common adverse reactions (i.e., that occurred in 1% or more of dicherace potassium for oral solution, t-readed patients) and more frequent with diciofenac potassium for oral solution. t-readed patients) and more frequent with diciofenac potassium for oral solution than with placebo were naises and deziriess (see Table 1). Table 1: Adverse Reactions With Incidence >1% and Greater Than Placebo in Studies 1 and 2 Combined

Adverse Reactions	Diclofenac Potassium for Oral Solution N=634	Placebo N=646
Gastrointestinal		
Nausea	3%	2%
Nervous System		
Dizziness	1%	0.5%

The most common adverse events resulting in discontinuation of patients following dicofenac potassium for oral solution dosing in controlled clinical trials were urticaria (0.2%) and flushing (0.2%). No withdrawals were due to a serious reaction.

6.2 Postmarketing Experience

The following adverse reactions have been identified during post approval use of disofense. or disofense or d Adverse Reactions Reported With Diclofenac and Other NSAIDs

Adverse Reactions Reported WRb Dichfenic and Other NSAIDs
In patients blank globines or other NSAIDs, the most frequently reported adverse reactions occurring in approximately
13-to 10% of patients are: Gir escritors, (including plenforation, healthour, constipation,
13-to 10% of patients are: Gir escritors, (including perforation, healthour, constipation,
14-to 10% of patients are: Gir escritors, (including perforation, healthour, constitution, and constitution, and the constitution of the constitution and the constitution of the constitut

Body as a Whole: Fever, infection, sepsis

Bady is a Whitel: Fever, infection, sepss.

Cardiovascular System; Congestive heart failure, hypertension, tachycardia, syncope

Destive System; Dry mouth, esophapità, gastricipeptic uters, gastrioti, gastriotinestinal bleeding, glossitis, hematemesis, hepatitis, jaundice

Hemic and Lymphatic System; Ecchymosis, eosinophilia, leutopenia, melena, purpura, rectal bleeding, stomatitis, thrombocytopenia

Harmic and Jumphatic Systems (Ecrlymosis, ecsiophial, leukopeniu, melena, purpuru, rectal bleeding, stomatilis, thrombocytopenia Medhabic and Mukrimosi Weight Change.

Nervous System, Anviety, asthemia, confusion, depression, dream abnormables, drowsiness, insomnia, malabie, nervousness, paresthesia, somnolence, tremors, vertigo Respiratory System, Anthrus, dyspene.

Sikh and Appendages, Alopecia, photosensikky, sweating increased

Social States; Birved Vision

Linggenital System; Cystilla, dysuria, hematuria, interstitäi nephrillis, oliguria/polyuria, proteinuria, renal failure

Other adverse reactions in patients taking NSAIDs, which occur rarely, are

Rody as a Whole: Anaphylactic reactions, appetite changes, death

<u>Cardiovascular System</u>: Arrhythmia, hypotension, myocardial infarction, paipitations, vasculitis

<u>Dipestive System</u>: Colitis, eructation, liver failure, pancreabilis

<u>Hemic and Lymphatic System</u>: Agranulocytosis, hemolytic anemia, aplastic anemia, lymphadenopathy, pancytopenia

Metabolic and Nutritional: Hyperglycemia Nervous System: Convulsions, coma, hallucinations, meningitis Respiratory System: Respiratory depression, pneumonia

Skin and Appendages: Angioedema, toxic epidermal necrolysis, erythema multiforme, exfoliative dermalitis, Stevens-johnson syndrome, utilization. Second Senses: Conjunctivitis, hearing impairment

7 DRUG INTERACTIONS

See Table 2 for clinically significant drug interactions with diclofenac
Table 2: Clinically Significant Drug Interactions with Diclofenac

Drugs That Interfere	with Hemostasis
Clinical Impact:	 Dicipfenac and anticoagulants such as warfarin have a synergistic effect or bleeding. The concomitant use of dicipfenac and anticoagulants have an increased risk of serious bleeding compared to the use of either drug alone.
	 Serotonin release by platelets plays an important role in hemostasis. Case- control and cohort epidemiological studies showed that concomitant use o drugs that interfere with serotonin reuptake and an NSAID may potentiate the risk of bleeding more than an NSAID alone.
intervention:	Monitor patients with concomitant use of diciplenac potassium for or solution with anticogulants (e.g., warfarin), antiplatelet agents (e.g., aspirin selective serotonir reuptake inhibitors (SRISIs), and serotonin nore
Aspirin Clinical Impact:	
	Controlled clinical studies showed that the concomitant use of NSAIDs an analysisk does of aspirin does not produce any greater theraportice than the use of NSAIDs alone. In a clinical study, the concomitant use of all NSAID and aspirin was associated with a significantly nereased incidence to the control of the con
Intervention: ACE Inhibitors, Angio	of aspirin is not generally recommended because of the increased risk of bleeding [see Warnings and Precautions (5.13)].
Clinical Impact:	NSAIDs may diminish the antihypertensive effect of angiotensin converting
,	enzyme (ACE) nibibors, angiotensin receptor blockers (ARBs), or beta- blockers (including propra anoiba). In patients who are elderly, volume-depleted (including those on diuretic therapy), or have renal impairment, co-administration of an INSAID with ACI inhibitors or ARBs may result in deterioration of renal function, including possible acute renal failure. These effects are susually reversable.
Intervention:	 During concomitant use of disoferace potessium for oral solution and ACE- inhibitors, ARRS, or beta-blockers, monitor blood pressure to ensure that the desired blood pressure is obtained. During concomitant use of disclorate, potassium for oral solution and ACE- inhibitors or ARRS in patients who are elderly, volume-depiteds, or have the property of the property of the property of the property of the Warnious and Precautions (S. Oir "sign of vivorening renaf Inniction [see Warnious and Precautions (S. Oir "sign of vivorening renaf Inniction [see
Diuretics	
Clinical Impact:	Clinical studies, as well as post-marketing observations, showed that NSAID reduced the natriuretic effect of loop diuretics (e.g., furosemide) and thiazid diuretics in some patients. This effect has been attributed to the NSAII highlition of repail prostatulation synthesis.
Intervention:	inhibition of renal prostaquandin synthesis. During concomitant use of diclofenac potassium for oral solution wit diuretics, observe patients for signs of worsening renal function, in addition to assuring diuretic efficacy including antihypertensive effects see Warnings and Precautions (5.6).
Digoxin Clinical Impact:	The concomitant use of diclofenac with digoxin has been reported to increas
Intervention:	the serum concentration and prolong the half-life of digoxin. During concomitant use of diclofenac potassium for oral solution and digoxir monitor serum digoxin levels.
Lithium Clinical Impact:	NSAIDs have produced elevations in plasma lithium levels and reductions is
Intervention:	renal Rhium clearance. The mean minimum Rhium concentration increase 15%, and the renal clearance decreased by approximately 20%. This effect has been attributed to NSAID inhibition of renal prostaglandin synthesis. During concomitant use of diciplenac potassium for oral solution and Rhium
	monitor patients for signs of Rhium toxicity.
Methotrexate Clinical Impact:	Concomitant use of NSAIDs and methotrexate may increase the risk for
Intervention:	Concomitant use of NSAIDs and methotrexate may increase the risk for methotrexate toxicity (e.g., neutropenia, thrombocytopenia, reni dysfunction). During concomitant use of distributions on the said of the concomitant use of distributions on the said of the concomitant use of distributions on the said of the concomitant use of distributions on the said of the concomitant use of distributions on the said of the concomitant use of distributions on the said of the concomitant use of distributions on the concomitant use of distributions of the concomitant use of the concomitant use of the concomitant use of distributions of the concomitant use of the conc
	During concomitant use of diclofenac potassium for oral solution an methotrexate, monitor patients for methotrexate toxicity.
Cyclosporine Clinical Impact: Intervention:	Concomitant use of diciofenac potassium for oral solution and cyclosporin may increase cyclosporine's nephrotoxicky. During concomitant use of diciofenac potassium for oral solution an cyclosporine, monitor patients for signs of worsening renal function.
NSAIDs and Salicylat	
Clinical Impact:	Concomitant use of diciofenac with other NSAIDs or salicylates (e.g., diffunisa salsalate) increases the risk of GI toxicity, with little or no increase in efficact see Warnings and Precautions (5.21).
Intervention:	The concomitant use of diclofenac with other NSAIDs or salicylates is no recommended.
Pemetrexed	
Clinical Impact:	Concomitant use of diciofenac potassium for oral solution and pemetrexe may increase the risk of pemetrexed-associated myelosuppression, renal, an Gl toxicky (see the pemetrexed prescribing information).
Intervention:	During concomitant use of NSAIDs and pemetrexed, in patients with ren- impairment whose creatinine clearance ranges from 45 to 79 mL/min, monitor for myelosuppression, renal and GI toxicity.
	NSAIDs with short elimination half-lives (e.g., diclofenac, indomethacin) shoul be avoided for a period of two days before, the day of, and two days followin administration of pemetrexed.
	In the absence of data regarding potential interaction between pemetrexe and NSAIDs with longer half-lives (e.g., meloxicam, nabumetone), patient taking these NSAIDs should interrupt dosing for at least five days before, th day of, and two days following pemetrexed administration.
Inhibitors of Cytochi Clinical Impact:	Dictofenac is metabolized predominantly by Cytochrome P-450 CYP2C9. Co administration of medications that inhibit CYP2C9 may affect the pharmacokinetics of dictofenac [see Clinical Pharmacology (12-3)] During concomitant use of dictofenac potassium for oral solution and drug that inhibit CYP2C9, an increase in the duration between dictofenac potassium

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy Risk Summary

Interesting Conference potassium for red to the fact of the conference of the confer

Premature Closure of Fetal Ductus Arteriosus

Use of MSAIDs, including dicbfenac potassium for oral solution, at about 30 weeks gestation or later in pregnancy increases the risk of premature closure of the fetal ductus arteriosus.

Olgohydramnios/Weonatal 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 olgohydramnios, and in some cases, neonatal renal impairment

Data from observational studies regarding other potential enterproprietal risks of IRSAID use in women in the first or second trimesters of pregnancy are inconclusive. In animal studies, or all animal studies of a studies of

All pregnancies have a background risk of birth defects, loss, or other adverse outcomes, in the U.S. general population, the estimated background risk of major birth most provided by the provided of the provided provided by the provid

Clinical Considerations.

Disease-Associated Maternal ander Embryo/Ental Risk

Several studies have suggested that women with migrane may be at increased risk of preeclampsia and gestational hypertension during pregnancy.

Seema to Judies have suggested that women with impaired may be at increased has of precumpsia and gestatorial hypertension during pregnant Fetal/Recordal Adverse Reactions

Premature Closure of Fetal Ductus Arteriosus:

Avoid use of NSAIDs in women at about 30 weeks gestation and later in pregnancy, because NSAIDs, including dicofenac solution, can cause premature closure of the fetal ductus arteriosus (see Data).

Oligohydramnics/Neconatal Renal Impairment:

If an ISAID in necessary at about 20 weeks gestation or later in pregnancy, limit the use to the lowest effective dose and shortest duration possible. If dicoference potassium for oral solution, can consider most possible most participation or significant potassium for oral solution. Once placed the fours, consider monitoring with uttrasourd for eligiphydramnios. If oligohydramnios occurs, discentinue dicioferac potassium for oral solution and follow up according to clinical practice (see Data).

Labor or Delivery

The effects of diobfenac potassium for oral solution on labor and delivery in pregnant women are unknown. In ral studies, material exposure to KSAIDs, as with other drugs known to inhibit prostaglandin synthesis, increased the incidence of dystocia, delayed parturktion, and decreased pup survival material exposure to KSAIDs, as with other drugs known to inhibit prostaglandin synthesis, increased the incidence of dystocia, delayed parturktion, and decreased pup survival

Fremature Closure of Fetal Ductus Arteriosus:
Published Renature 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.

Oligohydramnios, Neonatal Renal Impairment:
Published studies and postmarketing reports describe maternal NSAID use at about 20 weeks gestation or later in pregnancy associated with fetal renal dysfunction leading to oligohydramnios, and in

some cause, monital renal impairment. These adverse outcomes are seen, on average, after days to weeks of treatment, abh violates in many cases, but not all, the decrease is ammotic fluid was transient and reversible with cases of the drug. There have been aimfact anumber of case reports of material KIAMI use and sevential renal dysfunction without objective remains, some cases of material KIAMI use and sevential renal dysfunction without objective remains, some remains of the decrease of the second dysfunction without objective remains, some cases of the decrease of the decrease of the second dysfunction remains of the decrease o

rectionable and a second of the rection of the rect

neonatal outcomes with maternal MSIDs use Beause the published safety data on neonatal outcomes with maternal MSIDs use Beause the published safety data on neonatal outcomes involved mostly preterm infants, the generalizability of certain reported risks to the full-term infant exposed to NSAIDs through maternal use is uncertain.

Animal Data

Animal Data

Oral administration of dicitienes, sodium to preprient mice and rabbits during originogenesis resulted in embryofteal toxicity at oral doses of up to 20 and 10 magliciday, to to approximately 2 and 4 times, respectively, where there, respectively, the recommended human dose [Rifell] of 50 mg/day, but seed on body surface area [mg/m In ras, or all administration of dicitienes at doses of up to 10 mg/kg/day (up to approximately 2 times the RHI on a mg/m? basis) during organogenesis resulted in increased enhyoridant mortally and resulted flat body weight.

8.2 Lactation

Back Summay.

Data from published Rerature reports with oral preparations of diciplenae indicate the presence of small amounts of diciplenae in human milk. There are no data on the effects on the breadfed infart, or the effects on milk production.

The production of the production

8.3 Females and Males of Reproductive Potential

Fements

Managed in Managed in the number of prostaglands

Managed in Managed

interitation for oral solution, in women who have difficulties conceiving or who are undergoing investigation of infertility.

8.5 Geriatric Use Elderly patients, com

B.3. Gerantire Use

Ether) patients, compared to younger patients, are at greater risk for NSAID-associated serious cardiovascular, gastrointestinal, and/or renal adverse reactions. If the anticipated benefit for the dependent of the patient outweights these potential risks, monitor patients for adverse effects [see Warnings and Precautions (51, 52, 53, 56, 515]].

Clinical studies of diclofenac potassium for oral solution did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects.

8.6 Hepatic Impairment

Execute Inpact metabolism recourst for almost 100% of disofense einhaldon. Berushe helpack metabolism recourst for almost 100% of disofense einhaldon Berushe helpack for control and the considered for returnent with disofense potassium for oral solution only the benefits outwarph the risks. There is insufficient information washable to support dosing recommendations for disofense potassium for oral solution in patients with prefer solution.

In placement with replace insurincentry (see Lunica enamenacings) (12.7)).

No information is available from controlled clinical studies regarding the use of diciofenacy obstassism for oral postessism for oral solution in patients with advanced renal disease. Therefore, treatment with diciofenacy obstassism for oral solution is not recommended in patients with advanced renal disease. If dicidenacy potassism for oral solution is not recommended in patients with advanced renal disease. If dicidenacy potassism for oral solution therapy must be initiated, close monitoring of the patients' renaf infunction is advisable.

10 OVERDOSAGE

Symptoms following acute NSAID overdoses have been typically imited to lethargy, drowsiness, nausea, vomiting, and epigastric pain, which have been generally reversible with supportive care. Gastrointestinal bleeding has occurred. Hypertension, acute renal failure, respiratory depression and, coma have occurred, but were rare [see Warnings and Precautions (5.1, 5.2, 5.4, 5.6].

Manage patients with symptomatic and supportive care following an INSAID overdosage.

There are,

miles are,

mile

For additional information about overdosage treatment contact a poison control center (1-800-222-1222).

Anaphylactic reactions have been reported with therapeutic ingestion of NSAIDs, and may occur following an overdose

Dictofenac potassium for oral solution, USP is a nonsteroidal anti-inflammatory drug, available as a buffer doshibe powder, designed to be mixed with water prort to real administration. Dictofenac potassium for oral solution, USP is a white to off-white, peppermit flavored powder for oral solution packaged in individual unit dose packets.

The chemical name is 2-f(2.6-dichlorophenyl)aminol benzeneacetic acid mo

sait.

The molecular weight is 334.25. Its molecular formula is C₁₄H₁₀Cl₂NKO₂, and it has the following structure.

The inactive ingredients in diclofenac potassium for oral solution, USP include: Glyceryl behenate, mannitol, peppermint flavor, potassium bicarbonate, and sucralose.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Diclofenac potassium for oral solution has analgesic, anti-inflammatory, and antipyretic properties.

The mechanism of action of diclofenac potassium for or al solution, like that of other NSAIDs, is not completely understood but involves inhibition of cyclooxygenase (COX-1 and COX-2).

1 and COX 2). Dictionax is a potent hibitor of prostaglandin synthesis in vtro. Dictionax concentrations reached during therapy have produced in vito effects. Prostaglandins sensitize afferent nerves and potentiate the action of braykinn in inducing pain a minial modes. Prostaglandins are mediators of inflammation. Because dictionax is an inhibitor of prostaglandin synthesis, its mode of action may be due to a decrease of prostaglandins profibered tissues.

12.3 Pharmacokinetics

Ablacticition.

Dictorieus: is 100% absorbed after oral administration compared to intravenous Dictorieus: is 100% absorbed after oral administration to measured by uniter ecovery. However, due to this types metabolism, administration is measured by uniter the experiment of the property of the property of the experiment of the experim

<u>Distribution</u> The apparent volume of distribution (V/F) of diclofenac potassium is 1.3 L/kg.

Diclofenac is more than 99% bound to human serum proteins, primarily to albumin. Serum protein binding is constant over the concentration range (0.15-105 µg/mL) achieved with recommended doses.

Elimination Metabolism

Metabolism

The dischers, metabolites have been identified in human plasma and urine. The metabolites include 4*hydroxy, 3-hydroxy, 4*-5-dhydroxy and 3-hydroxy 4*hydroxy, 5-hydroxy, 4*-5-dhydroxy and 3-hydroxy 4*hydroxy, 5-hydroxy, 4*-5-dhydroxy and 3-hydroxy 4*hydroxy, 5-hydroxy, 4*hydroxy, 5-hydroxy, 4*hydroxy, 4*-

Diclofenac is eliminated through metabolism and subsequent urinary and bilary excretion of the glucuronide and the sulfate conjugates of the metabolites. Little or no

excretion of the glucuronide and the sulfate conjugates of the metabolites. Little or no free unchanged discloriens.

Beccreted in the urine. Approximately 65% of the dose is excreted in the urine and approximately 35% in the bile as conjugates of unchanged discloriens; approximately 56% of the dose is excreted in the urine. Approximately 76% of the dose is excreted in the urine and approximately 35% in the bile as conjugates of unchanged discloriens. ation is not a significant pathway of elimination for unchanged diclofenac, dosing adjustment in patients with mild to mo

nt: The liver metabolizes almost 100% of diclofenac; there is ation available to support dosing recommendations for diclofenac

tick Impairment: The liver metabolizes almost 100% of diclofenac; there is "Increast information available to support dissing recommendations for diclofenac salam for orda son in patients with hepatic insufficiency (see Warnings and Precautions (5.3) and Use in Specific Populations (8.6)].

Renal Impairment: In patients with renal impairment (inulin clearance 60-90, 30-60, and <30 ml, lmin; N=6 in each group), AUC values and elimination rate were comparable to those in healthy subjects [see Warnings and Precautions (5.0) and Use in Specific Populations (8.7)].

Drug Interaction Studies

en NSAIDs were administered with aspirin, the protein binding of NSAIDs

we'r reduced,
although the clearance of free NSAID was not altered. The clinical significance of this interaction is not known. See Table 2 for clinically significant drug interactions of NSAIDs with aspirin [see Drug Interactions (7)].

13.1 Carcinogenesis & Mutagenesis & Impairment Of Fertility

2-year carcinogenicity study conducted in mice employing dicbfenac sodium at doses up to 0.3 mg/kg/day (less than the RHD on a mg/m² basis) in males and 1 m/kg/day (less than the RHD on a mg/m² basis) in females did not reveal any noncoperic potential.

Diciofenac sodium was not genotoxic in in vitro (reverse mutation in bacteria [Ames], mouse lymphoma tk) or in in vivo (including dominant lethal and male germinal epithelial chromosomal aberration in Chinese hamster Jassays.

Impairment of Fertility

Diclofenac sodium administered to male and female rats at 4 mg/kg/day (less than the RHD on a mg/m² basis) did not affect fertility.

The efficacy of diciofenac potassium for oral solution in the acute treatment of migraine headache was demonstrated in two randomized, double blind, placebo-controlled trials.

Patients errolled in these two trials were predominantly female (85%), and white (86%), with a mean age of 40 years (range: 18 to 65). Patients were instructed to treat a migraine of moderate to severe pain with 1 dose of study medication. Patients evaluated their headache pain 2 hours later. Associated symptoms of nauses photophobia, and phonophobia were also evaluated. In addition, the proportion of Photosphobia, and phonophobia were also evaluated. In adultant, une property photosphobia, and phonophobia were also evaluated. In adultant, une property photosphobia patients who were "sustained pain free", defined as a reduction in headache severity from moderate or severe pain to

no pain at 2 hours post- dose without a return of mild, moderate, or severe pain and no use of rescue medication for 24 hours post-dose, was also evaluated. In these studies,

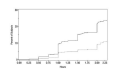
this gold is a consideration for 2.4 hours post-doce, was also evaluated. In these susues, the percentage of patients schewing pain freedom 2 hours, after treatment and sustained pain freedom 7 to 2.4 hours post-solution compared with those who received piaceto (see Table 3). The percentage of patients schewing pain refed 2 hours after treatment (defined as a reduction in headsche spatients schewing pain refed 2 hours after treatment (defined as a reduction in headsche spatients schewing pain refed 2 hours after treatment (defined as a reduction in headsche spatients) and the spatients who received piaceto (see Table 3).

Septimized with those who received piaceto (see Table 3).

Table 3: Percentage of Patients with 2-Hour Pain Freedom, Sustained Pain Freedom 2-24 Hours, and 2-Hour Pain Relief Following Treatment

Study 1 Hour Pain Free 2-Hour Pain Free 2-24h Sustained Pain F 2-Hour Pain Relief 48% Diclofenac Potassium for Oras Solution (n=343) 25% 19% 65% Study 2 lacebo (n=347) 2-Hour Pain Free 2-24h Sustained Pain Free 2-Hour Pain Relief

The estimated probability of achieving migraine headache pain freedom within 2 hours following treatment with dicbfenac potassium for oral solution is shown in Figure 1. Figure 1: Percentage of Patients with Inikial Headache Pain Freedom within 2 Hours



There was a decreased incidence of nausea, photophobia and phonophobia following administration of dicbfenac potassium for or al solution, compared to placebo. The efficacy and safety of dicbfenac potassium for oral solution was unaffected by age or gender of the patient.

Diciofenac potassium for oral solution, USP 50 mg, is a white to off-white, peppermint flavored powder for oral solution, supplied as individual dose packets. Each individual packet is designed to deliver a dose of 50 mg diciofenac potassium when mixed in water.

water.

NDC 67877-772- 49 Individual diclofenac potassium for oral solution, USP packets

NDC 67877-772- 58 Boxes of nine (9) diclofenac potassium for oral solution, USP packets

Store at 25°C (77°F). Excursions permitted from 15°C to 30°C (59°F to 86°F). [See USP Controlled Room Temperature]

Advise the patient to read the FDA-approved patient labeling (Medication Guide) that accompanies each prescription dispersed. Inform patients, families, or their caregivers of the following informatic before initiating therapy with dictofenac potassium for oral solution and periodically during the course of ongoing therapy.

Advise patients to be alert for the symp scular thrombotic events, including chest pain, shortness of breath, weakness, or slurring of speech, and to report any of these symptoms to their health care provider immediately [see Warnings and Precautions (5.1)].

Debtemen, postable and the Committee of the Committee of

Hepatotoxisty Inform patients of the warning signs and symptoms of hepatotoxickly (e.g., nausea, Tatique, etharqy, prurkus, diarrhea, jaundice, right upper quadrant tenderness, and "Tu-like" symptoms). If these occur, instruct patients to stop diciofens potassium for oral solution and seek immediate medical therapy (see Warnings and Precautions (5.3).

Heart Fahire and Edema
Advise patients to be alert for the symptoms of congestive heart failure including
shortness of
brindings.

The advise patients are supported by the state of the symptoms occur [see Warnings and
Precautions (5.2)].

ents of the signs of an anaphylactic reaction (e.g., difficulty breathing, sweling of the face or throat). Instruct patients to seek immediate emergency help if these occur [see Contraindications (4) and Warnings and Precautions (5.71).

Serious Skin Reactions, Including DRESS
Advise patients to stop taking dichorance plants for oral solution immediately if they
deebeg any type of risth, blatters, fewer or other signs of hypersensibility such as
tching and to contact their healthcare provider as soon as possible. Dicherance
solution, like other MSAIDs, can cause serious skin reactions such as enfoliately deemaktis.
Sleever-johnson syndrome (SSS), took expediermal necrosis (TBIs), and DRESS, which may
carried it hospitablestions and even death the Winnings and Procastions (5.6, 5.10).

Medication Overuse Headache
Inform patients that use of acute migraine drugs for 10 or more days per month may
lead to
are excernation of headache and encourage patients to record headache frequency and drug use (e.g., by keeping a headache diary) [see Warnings and Precautions (5.11)].

Fetal Toxicity

International International Processing of the Pr

Advise patients to notify their heathcare provider if they are breastfeeding or plan to breastfeed [see Use in specific Populations (8.2)].

Female Fertility

Advise females of reproductive potential who desire pregnancy that NSAIDs, including dicidence potassium for oral solution, may delay or prevent rupture of ovarian follicles, which has been associated with reversible infertility in some women [see Use in Specific Populations (8.3)].

Avoid Concomitant Use of NSAIDs

Inform patients that the concomitant use of dichfense potassium for oral solution with other MSAIDs or saks/dates (e.g., offlunisk) sakslated is not recommended due to with other MSAIDs or saks/dates (e.g., offlunisk) sakslated is not recommended due to the increased risk of gastrointestinal subcky, and title or no increase in efficacy [see Warmings and Precautions (7.2) and Dug Interactions (7)). Alert patients that MSAIDs may be precent in view the counter* motic alons for treatment of colds, feeer, or insommit.

Use of NSAIDS and Low-Dose Aspirin

Inform patients not to use low-dose aspirin concomitantly with diclofenac potassium for oral solution until they talk to their healthcare provider [see Drug Interactions (7)].

Distributed by: Ascend Laboratories, LLC

Parsippany, NJ 07054

Diclofenac Potassium (dye-KLOE-fen-ak poe-TAS-ee-um) for oral solution

What is the most important information I should know about diciofenac potassium for oral solution? Diclofenac potassium for oral solution contains diclofenac (a non-steroidal anti-inflammatory drug or NSAID).

Dickbenac potassium for oral solution contains dickbenac (a non-steroidal anti-inflammatory drig or MSAID).

NSAIDs, including dickbenac potassium for oral solution, can cause serious side effects, including with the contained of the contained

What is diclofenac potassium for oral solution?

What is disclofense potassium for oral solution?

Discloress potassium for oral solution is a prescription medicine used to treat migraine attacks in adults. It does not prevent or lessen the number of migraines you have, and it is not for other types of headaches. Disclorenze potassium for oral solution contains disclores potassium for noral service in the disclorest potassium for noral solution.

When should It take disclorenze potassium for oral solution.

Take disclorenze potassium for oral solution exactly as your headache per provider tells you to take it.

Take 1 dose of diciblenac potassium for oral solution to treat your migraine headache:

• take a single dose packet
• open packet only when you are ready to use it.
• open packet only when you are ready to use it.
• or contents of packet into 1 to 2 outness or 2 to 4 tablespoons (30 to 60 mL) of
• mix well and mixth the water and power mixture
• throw away empty packet in a safe place and out of the reach of children.
• throw away empty packet in a safe place and out of the reach of children.
• effectubriens compared to taking dicionac potassium for oral solution on an empty stomach.
• on the children or dicionaction potassium for oral solution.
• of not take more dicionaction potassium for oral solution.
• of not take more dicionaction contents are of overdoos, get medical help or contents a Posion Control Center right away.

Who should not take diclofenac potassium for oral solution?

Do not take diclofenac potassium for oral solution:

• if you have had an asthma attack, hives, or other allergic reaction with aspirin, diclofenac, or any other NSAIDs.

right before or after heart bypass surgery.

Before taking diclofenac potassium for oral solution, tell your healthcare provider about all of your medical conditions, including if

You:

* have lex or kidiney problems

* have a history of stomach uker or bleeding in your stomach or intestines

* have any allerges to any medicines

* have othest pain, shortness of breath, irregular heartbeats

* have chest pain pressure

* have appressure

• have high blood pressure

• have eithing
• have eithing
• are pregnant, that's you might be pregnant, or are trying to become pregnant. Taking
• are pregnant, that's you might be pregnant, or are trying to become pregnant. Taking
• pregnant, or little may harm your unborn babb, if you need to take IKSIDs for
more than 2 days when you are between 20 and 30 weeks of pregnancy, your
heathcare provider and to the amount of talk in your words around
pregnancy as should not take IKSIDs after about 30 weeks of
pregnancy as hould not take IKSIDs after about 30 weeks of
pregnancy are provider about a of it the modelines you take, including prescription or overtife your heathcare provider about at of the modelines you take, including prescription or overtife your heathcare provider about and of the modelines you take, including prescription or overdied before produced to the provider about 30 weeks of
the discovery providers about 50 or the provider of the provider of the order of the provider of the provider of the order of the provider o

Know the medicines you take. Keep a list of your medicines and show it to your doctor and pharmacist when you get a new medicine

What are the possible side effects of diclofenac potassium for oral solution?

Diclofenac potassium for oral solution can cause serious side effects, including:

See "What is the most important information I should know about diclefense potassium for oral solution?"

- New York of high blood pressure
- heart failure
- bleeding and uiters in the stomach and intestine
- bleeding and uiters in the stomach and intestine
- bleeding and uiters in the stomach and intestine
- but ore blood coff, identina)
- bleeding and uiters in the stomach and intestine
- bleeding and uiters in the stomach and intestine
- but or the blood coff, identinal
- interest in the stomach and intestine the stomach and intestine the stomach and intestine the stomach and interest in the stomach and interest interest

Other side effects of NSAIDs include: stomach pain, constipation, diarrhea, gas, hearthurn, nausea, vomiting and dizziness.

Get emergency help right away if you get any of the following symptoms:

	breathing	slurred speech
٠	chest pain	swelling of the face or throat
•	weakness in one part or side of your body	

Stop taking diclofenac potassium for oral solution and call your health provider right away if you get any of the following symptoms

 nausea that seems out of proportion to your migraine 	
sudden or severe pain in your belly	 there is blood in your bowel movement or it is black and sticky like tar
more tired or weaker than usual	unusual weight gain
diarrhea	more tired or weaker than usual
itching	 skin rash or blisters with fever
your skin or eyes look yellow	 swelling of the arms, legs, hands and feet
indigestion or stomach pain	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 pharmacist about NSAIDs.

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
 Aspirin is an NSAID but it 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.

the stomach and intestines.

Some NSAIDs are sold in lower doses without a prescription (over-the counter). Take
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.

salite symptotis using you make it may main treat.

If you would be more information about MSAIDs, the most of relating to provider for information about NSAIDs that is written for health professionals.

Manufactured by:

Altern Laboratories Ltd.,

Distributed by: Ascend Laboratories, LLC Parsippany, NJ 07054

This Medication Guide has been approved by the U.S. Food and Drug Administration. Revised 7/2022 PT-9123





F	roduct Infon	mation					
p	roduct Type		HUMAN PRESCRIPTION DRUG	Item Cod	ie (Source)	NDC:6	7877-772
R	oute of Admini	stration	ORAL				
A	ctive Ingredi	ent/Active	Moiety				
		Ingr	edient Name		Basis of S	trenath	Strengt
	NI:14408QL0L1)	ASSIUM (UNII:	L4DSUA6CB4) (DICLOFENAC -		DICLOFENAC POTASSIUM		50 mg
li	nactive Ingre	dients					
			Ingredient Name			Stre	ength
	LYCERYL DIBENE		WTH25Y52)				
	ANNITOL (UNIT 3						
			HM5Z15LEBN)				
s	UCRALOSE (UNI: EPPERMINT (UNI:	96X8UQ3Z D4)	: HMSZ1SLEBN)				
P	UCRALOSE (UNI: EPPERMINT (UNI: Troduct Chara olor	96KBUQ3ZD4) V95R5KMY2B)	hite Powder)	Score			
P	UCRALOSE (UNI: EPPERMINT (UNI: Troduct Chara olor hape	VESTSKMYZE) VESTSKMYZE) ACTORISE (VALUE (V	ibite Powder)	Size			
PCSF	ucratose (uni: eppermint (uni: roduct Chara olor hape lavor	96KBUQ3ZD4) V95R5KMY2B)	ibite Powder)	Size	e int Code		
PCSF	UCRALOSE (UNI: EPPERMINT (UNI: Troduct Chara olor hape	VESTSKMYZE) VESTSKMYZE) ACTORISE (VALUE (V	ibite Powder)	Size			
PCSFC	ucratose (uni: eppermint (uni: roduct Chara olor hape lavor	VESTSKMYZE) VESTSKMYZE) ACTORISE (VALUE (V	ibite Powder)	Size Impri	int Code		
PCSFC	roduct Charz olor hape lavor ontains	PEPPER	ibite Powder)	Size Impri	int Code	Market Da	ing End
PCSFC	UCRALOSE (UNI: EPPERMINT (UNI: Product Chara olior hape la vor ontains Tackaging Item Code NDC:67877-772-755	PEPPER Pain 1 CARTO	inhie Powder) INIT kkage Description	Size Impri	int Code	Market De	ing End
PCSFC	roduct Chara olor ontains item Code NDC-67877-772-	PEPPER Pain 1 CARTO	inhe Powder) IENT Kkage Description	Size Impri	int Code	Market Da	ing End
P C S F C	ucratoss (uni- roduct Chara oler hape laver ontains tackaging ltem Code NDC-87877-772- 59 NDC-87877-772- 49	PAPPER Part 1 1 1 PACKET Product	this Powder) INT Lkage Description (1) Type 0: Not a Combination	Size Impri	int Code	Market Da	ing End
P C S F C	UCRALOSE (UNI- EPPERMINT (UNI- Product Chara ofor hape lavor ontains ackaging Rem Code NDC-67877-772- 58 NDC-67877-772-	Perpere Part 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	this Powder) INT Lkage Description (1) Type 0: Not a Combination	Size Impri	int Code	Da	ing End ite

Labeler - Ascend Laboratories, LLC (141250469)