HIGHLIGHTS OF PRESCRIBING INFORMATION
TOPICAL SOLUTION safely and effectively. See full prescribing information for DKLOFENAC SODIUM
TOPICAL SOLUTION safely and effectively. See full prescribing information for DKLOFENAC
SODIUM TOPICAL SOLUTION.
DKLOFENAC SODIUM Topical Solution, 1.3% w/w, for topical use initial U.S. Approxit: 1588

- Nonsteroidal 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. (5.1)
- can be feat. The risk may occur withy two values are written and other shadow.

 Can be feat. The risk may occur with your values of the risk of the ri

- nd symptoms of osteoarthritis of the kneets\() (1)

 DOSAGE AND ADMINISTRATION

 the lowest effective dosage for the shortest duration consistent, with individual patient treatment

- Size the Securit delicture diseage for the district administration with includar plants responses to the security of the se

- * Relating in Statistics, of International Conference and International Co

- Note the leaves and find in the finding of the first the
- amerina (5.12, 7)

 <u>Epocyter to (light</u>: Avoid exposure of treated knee(s) to natural or artificial sunlight. (5.15)

 <u>Epoc. Contact</u>: Avoid centact of dicidenac sodium with eyes and mucosa. (5.16)

 <u>Cost Nestertorial Articinal Imministry Druss</u>: Avoid concurrent use with earl MSADs. (5.17)

ADVERSE REACTIONS

Most common adverse reactions with dictofenac sodium topical solution are application site reactions.

(6.1) To report SUSPECTED ADVERSE REACTIONS, contact Lupin Pharmaceuticals Inc. at 1-800-399-2561 or FDA at 1-800-FDA-1088 or www.fda.gov/msdwatch.

DRUG INTERACTIONS

- DRUG INTERACTIONS
 Dougs that Interfere with Hemostasis Se or our fairs, ascient SS-BNS-SNBs; it Monitor patients for bleeding who are concemitantly using dictionate soldium with drugs that interfere with hemostasis. Concemitant

use of dictionac cedam and analysis closes of acpiris in on generally recommended (7).

A Exhibition, designation insecrit foliases of Acpiris in an operandly recommended (7).

A Exhibition, designation insecrit foliases of Acpiris in a second consideration of the activities coloid many affirmabilities and exhipperations we effect of those drugs, Monther Bodo pressure (7) and the activities of the activit

like of digosom. Monitor iserum digosom levels (7)

Interfallay, NSADs are associated with reversible interfallay, Consider withdrawal of dictablinac sodium in watering with the controlling (13).

See 2.7 for PATERIT COUNSELING INFORMATION and Medication Guide.

Beviolet 12/002

FULL PRESCRIBING INFORMATION: CONTENTS* 1 INDICATIONS AND USAGE 2 DOSAGE AND ADMINISTRATION 2.1 General Dosing Instructions 2.2 Sensit Bremarking

- 2 INDICATIONS AND USAGE
 2 DOSAGE AND ADMINISTRATION
 2 2 Special Piez addion.
 3 2 Special Piez addion.
 3 2 Special Piez addion.
 3 2 Special Piez addion.
 4 CONTRAINDICATIONS
 5 VARANINGS AND PERCAUTIONS
 5 VARANINGS AND PERCAUTIONS
 5 VARANINGS AND PERCAUTIONS
 5 1 Repaid Piez Addion.
 5 1 Secretarion of Addinance Addition.
 5 1 Secretarion of Addition Addition.
 5 1 Secretarion of Addition Addition.
 5 1 Secretarion of Addition Redeated to Appirin Sensibility
 5 1 Secretarion of Addition Redeated to Appirin Sensibility
 5 1 Secretarion of Addition Redeated to Appirin Sensibility
 5 1 Secretarion of Addition Redeated to Appirin Sensibility
 5 1 Secretarion of Addition Redeated to Appirin Sensibility
 5 1 Secretarion of Additional Redeated to Appirin Sensibility
 5 1 Secretarion of Additional Redeated to Appiring Sensibility
 5 1 Secretarion of Additional Redeated Secretarion (Red Secretarion Red Se

- 6 Eye Exposure
 7 Oral Nonsteroidal Anti-Inflammatory Drugs
 WERSE REACTIONS
 Clinical Trials Experience
 Postmarketing Experience
- 6.2 Postmarketing Experience DRUG INTERACTIONS USE IN SPECIFIC POPULATIONS

- 8.4 Pediatric Use
 3.5 Geristri Use
 10 VENDOSAGE
 10 VENDOSAGE
 11 CLINICAL PHARMACOLOGY
 12.1 Mechanism Of Action
 12.3 Pharmacolostets
 13 NONCHINICAL TOXICOLOGY
 13.4 Carringeress, Matageness, Impairment of Fertility
 13.4 Arinam Eloxicology and or Pharmacology
 13.4 Arinam Eloxicology and or Pharmacology
 13.4 Arinam Eloxicology and or Pharmacology
 13.4 Arinam Eloxicology and the New York Carring State Office of Pharmacology
 14.4 Studies no Descentritis of the Knee
- 14 CLINICAL STUDIES
 14.1 Studies in Osteoarthritis of the Knee
 16 HOW SUPPLIED/STORAGE AND HANDLING
 17 PATIENT COUNSELING INFORMATION
 ** Sertions or subsertions omitted from the full prescribing

WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS

Centiovascular Thrombotic Events

Nonsteroidal anti-inflammatory drugs (NEAIDs) cause an increased

Nonsteroidal anti-inflammatory drugs (NEAIDs) cause an increased

infraction and stroke, which can be fast. This risk may occur early in
treatment and may increase with duration of use [see WARNINGS

AND PRECAUTIONS (5.1)].

AND PRECAUTIONS (5.1)].

Diclofenac sodium is contraindicated in the setting of coronary artery bypass graft (CABG) surgery [see CONTRAINDICATIONS (4) and WARNINGS AND PRECAUTIONS (5.1)].

WARNINGS AND PRECAUTIONS 7.3.

Seatonitas than Biseston. Elecanta, and Parforation

MAMD and Parforation

MAMD and Parforation

MAMD and part of the seaton of the stomest including bleeding, uceration, and perforation of the stomest or intestines, which can be falsh. These events can occur a seaton of the sea

2 DOSAGE AND ADMINISTRATION

2.1 General Dosing Instructions
Use the lowest effective desage for the shortest duration consistent with independent retentines goods few MARAIMOR AND PRECAUTIONS (5.3))
For the relief of the signs and symptoms of estematrities of the knee(s), the recommended does 6.0 drops per five. 4 times a day.
Apply dictionars sodium topical solution to clean, dry skin.

To avoid spilege, dispense dichrimes sodium topical solution 10 drops as a time either To avoid spilege, dispense dichrimes sodium topical solution 10 drops as a time either sodium topical solution evenly around front, back and sides of the kines. Repeat this procedure until 40 drops have been applied and the kines is completely overed with solution. To treat the other kines, if symptomatic, repeat the procedure.

Application of diciofense sodium topical solution in an amount exceeding or less than the recommended dose has not been studied and is therefore not recommended.

2. Special Precautions

Avoid shower/splashing for a least 30 minutes after the application of discrimations

**Avoid shower/splashing for a least 30 minutes after the application of discrimations of the production to the treated kines.

**Avoid contact of discrimation shower and the splash of the spl

Little the treated knee(s) is completely dry, avoid skin-to-skin contact between other people and the treated knee(s).

**Do not use combrasted therapy with diclofenac sodium topical solution and an oral exclusions the benefit sufweight the risk and conduct periodic abbratory evaluations.

3 DOSAGE FORMS AND STRENGTHS
Diclofenac Sodium Topical Solution USP, 1.5% w/w

CONTRANSMENTATIONS
 Disclores sodium is contraindicated in the following patients:
 Known hypersensibity (e.g., anaphylactic reactions and serious skin reactions) to disclores, or any components of the drug product, less WARMINGS AND
 History of asthma, urticaria, or other allergic-type reactions after taking aspirin or

other NSAIDs. Severe, sometimes fatal, anaphylactic reactions to NSAIDs have been reported in such patients (see WARNINGS AND PRECAUTIONS (5.7.5.8)).

In the setting of coronary artery bypass graft (CABG) surgery [see WARNINGS AND PRECAUTIONS (5.1)].

5 WARNINGS AND PRECAUTIONS

S WANNINGS AND PRECAUTIONS

5.1. Cardiovascular Thrombotic Events

Clinical train of several COX.2 selective and nonselective NSAIDs of up to three years developed to the control of the

Chevrationist acutes conducted in the Danish National Registry have demonstrated that Christophia and Christophia and Christophia and Christophia and Christophia and Christophia and a Cause mortally beginning in the first week of treatment. In this same cohort, the includes of elabor in the first year post-M was 20 pc 100 person compared patients. Almosphia the abouts are of death clocking beginning that year post-Milk, the increased relative risk of each in NSAID scens persisted over at least Anoth the use of divinerance.

Another used not dependent on the control in Soul users persisted over at east Another used officience accolum in petitions with a recent Multiple size the persistence of the expectation of the size of dependent control in Soulimb is used in patients with a recent Multiple most persistent for signs of carbon chemical flowledges collumb is used in patients with a recent Multiple most persistent for signs of carbon chemical flowledges. Soulimb is used in patients with a recent Multiple signs in the signs of the soulimb is significant to the soulimb in the signs of the signs of the soulimb in the signs of the signs of the signs of the soulimb is significant to the signs of t

Risk Factors for GI Bleeding, Ulceration, and Perforation

Risk Factors for GI Bleeding, Ulcration, and Perforation

Paleirst such a print history of perfix cert Gensea and/or GI bleeding who used NSADs had a greater than 10-foot increased risk for devolvage, a GI bleed compared to patients that the print of the GI bleeding with the GI bleeding of GI bleeding or GI bleeding or

- risk for Gi bleeding.

 Strategies to Minimize the Gi Risks in MSAID-treated Patients

 Use the lowest effective disage for the shortest possible duration.

 Avoid use in patients of more than one MSAID at a time.

 Avoid use in patients at higher risk unless benefits are expected to outweigh the increased risk of beleding. For such patients, as well as those with active Gi bleedin consider alternate therapies other than MSAID.

 Remarks alter for agrin and symptoms of of liceration and bleeding during MSAID.

In clinical trials, of oral dicibfenac-containing products, meaningful elevations (i.e., more than 3 times the ULM) of AST (SGOT) were observed in about 2% of approximately 5,700 patients at some time during dicibfenac treatment (ALT was not measured in all studies).

5.100 petition at 500 er une surrig subsense, seasones que la merca participato de la Segui popular de 100 er une surrigio de la Segui popular del Segui

when compared to other ISSADE. Elevations in transmissies were seen more requested in patients with activativities have in those with the manded stricts.

Amout all international elevations in transmissies were detected before patients became decidence in 42 of 185 patients in all this wish dovelveber manded stransmissies elevations.

In the contractivity reports, case of drug included inspatiolacity, have been reported by the internativity reports, case of drug included in patients between the patients of the patients of the patients and the patients of the patients

immeasure/.

Inform patients of the warning signs and symptoms of hepatotoxicity (e.g., nausea, fatgue, lethargy, diarriea, prurfux, joundice, right upper quadrant tenderness, and fluide symptoms. If chical signs and symptoms consistent with her disease develop, or if systems: manifestations occur (e.g., easinophia, rash, etc.), diacordinue diciblenac socialm immediately, and perform a facical revaluation of the patient.

To minimize the potential risk for an adverse liver-related event in patients treated with diclofenac sodium, use the lowest effective dose for the shortest duration possible. Exercise caution when prescribing diclofenac sodium with concomitant drugs that are known to be potentially hepatotoxic (e.g., acetaminophen, antibiotics, antiepteptics).

5.4 Hypertension
NSAIOs, including dicidence sodium, can lead to new onset of hypertension, or
worseing of predicting hypertension, either of which may contribute to the increased
worseing of predicting hypertension, either of which may contribute to the increased
that of duretic, or loop duretics may have impaired response to these therapies when
taking ISSAIOs (see DOIG INTERACTIONS (7)).

Monitor blood pressure (BP) closely during the initiation of NSAID treat throughout the course of therapy.

5.5 Heart Failure and Edema

The Coxib and traditional MSAID Trislists' Collaboration meta-analysis of randomized controlled trials demonstrated an approximately two-fold iscrease in hospitalizations for heart failure in CCV2 selective-breader platents and numberate two MSAID tradestarg platents with the control of the Company of the

Additionally, fluid retention and selema have been observed in some patients treated with NSAIDs. Use of dictofenox may blust the CV effects of soveral therapeutic agents used to treat these medical conditions (e.g., duretics, ACE inhibitors, or angistensin receptor blockers (ARBs)) [see DRUG INTERACTIONS (7)].

Avoid the use of dicfofenac sodium in patients with severe heart failure unless th benefits are expected to outweigh the risk of worsening heart failure. If dicfofen sodium is used in patients with severe heart failure, monitor patients for signs of worsening heart failure.

wersening heart failure.

5. Eneard Toxicky and Hyperkalemia
Renal Toxicky and Hyperkalemia is resulted in result papillary necrosis and other
renal ejary.

Renal toxicky has also been seen in patients in whom renal protosplandins have a
compensatory role in the maintenance of renal perfusion. In these patients,
renal toxicky has also been seen in patients in whom renal protosplandins have a
compensatory role in the maintenance of renal perfusion. In these patients,
represents and, secondarily, in real blood live, which may preside over it renal
discompensation. Patients all greatest risk of this reaction are those with impaired renal
discompensation. Patients all represents related to this reaction are those with impaired renal
discorptions. The results of the reaction are those with impaired renal
discorptions. The results are related to the reaction are those with impaired renal
discorptions. The results are related to the reaction are those with impaired renal
discorptions. The results are related to the reaction of MSADD therapy
was usually followed by recovery to the preferentment state.

usuary rusowed by recovery to the pretreatment state.

No information is available from controlled clinical studies regarding the use of disclerasodium in patients with advanced renal disease. The renal effects of dictofenas sodium
may hasten the progression of renal dysfunction in patients with preexisting renal
disease.

disease. Correct volume status in dehydrated or hypovolemic patients prior to initiating diciblem sodium. Monitor renal function in patients with renal or hepaic impairment, heart fails dehydration, or hypovolemia during use of deficient actioning feed (in IPRSACT/IOI) dehydration are supported action and the second prior of the second prior to the the benefits are expected to authority the risk of wonsemp renal function. If diciblems socialism is used in plates with advanced renal disease, monitor patients for signs of worsering renal function.

5.7 Anaphylactic Reactions

3.7 Anaphysicitx Reactions
Dickfene, has been associated with anaphylactic reactions in patients with and without known hypersensibility to dicidence and in patients with aspirin-sensitive asthma [see COMPRAINICIATIONS (4) and WARNINGS AID PREACTIONS (5.8)].
Seek emergency help if an anaphylactic reaction occurs.

5.8 Exacerbation of Asthma Related to Aspirin Sensitivity

A subpopulation of platents with safthm many have appirs sensible actima which may include crome riminations of platents with safthm many have appirs severe, potentially final include crome riminations. The platent platent is platent plat

offer pre-wardrage in the first and symptoms or examina3.5 scripts. She Reschedular
3.5 scripts. She Reschedular
3.5 scripts. She Reschedular
4.5 scripts. She Reschedular
5.5 scripts. She Reschedular
6.5 scripts. Sh

skin reactions to NSAIDs [see CONTRAINDICATIONS (4)].

Do not apply dictofemac sodium topical solution to open skin wounds, infections, inflammations, or exfoliative dermatitis, as it may affect absorption and tolerability of the drue.

drug.

5.10 Drug Reaction with Eosinophila and Systemic Symptoms (DRESS)
Drug Reaction with Eosinophila and Systemic Symptoms (DRESS) has been regorted in pellerest salving (RSBS) and in dictional cooling. Some of these events have been reported in pellerest stalving (RSBS) so which collidates cooling. Some of these events have been reported in the pellerest specific pellerest specific pellerest pellere

5.11 Fetal Toxicity

Avoid use of NSAIDs, including diclofenac, in pregnant women at about 30 weeks gestation and later. NSAIDs, including diclofenac, increase the risk of premature closure of the fetal ductus arterious at approximately this gestational age.

Olgohydramnios/Neonatal Renal Impairment:

Opinity distances altered at Bread Impairment.

We of NADIAs, Furthy disclorines adout 20 weeks gestation or later in pregnancy may cause feat mend dysfunction leading to oploylydramios and, is some cause, recorded rend impairment. These observations are seen as manage, after the cause of the control of t

50.24 Rematology in Taxicity per oder in Section. Productions (sel.);
Anemie has occurred in NSAID treated patients. This may be due to occult or gross belood toss. But intention, or an incompletely described effect on erythropoies. If a patient treated with dischleres soulim has any sign or symptoms of anemis, monitor NSAIDs, reckling plotforms coulim, may have been souliment of anemis monitor NSAIDs, reckling plotforms coulim, may have been seen from the ordering control of the complete conditions such as compulsion disorders, concentrate use of warfars, other arcinosoguistics, relinited experts (se.), participate experts (se.) participate in the complete conditions such as compulsion of sections in ordering the conditions of the conditio

The effects of dicbfenac sodium on platelet function were studied in 10 healthy subjects administered 80 drops four times a day for 7 days. There was no significant change in platelet aggregation following one week of treatment [see CLINICAL PHARMACOLOGY (12-4)].

5.13 Masking of Inflammation and Fever

The pharmacological activity of diclofenac sodium in reducing inflammation, and possibly fever, may diminish the utility of diagnostic signs in detecting infections.

S-14 Laboratory Monitoring

Because serious Gi bleeding, hepotoxicity, 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 (\$.2, 5.3,
5.0].

5.15 Sun Exposure
Instruct patients to avoid exposure to natural or artificial sunlight on treated knee(s)
because studies in animals indicated topical dictoferac treatment resulted in an earlier
onset of ultraviolet light-induced skin tumors. The potential effects of dictoferac sodium
on skin response to utraviolet damps in humans are not known.

S.16 Eye Exposure
Avoid contact of diciofenac sodium with eyes and mucosa. Advise patients that if eye contact occurs, immediately wash out the eye with water or salme and consult a physicani if irration presists for more than an hour.

A.3.7 or in Nonsteroidal Ant-Inflammatory Drugs
Concembar use of oral NSAIDs with disofense sodium resulted in a higher rate of
rectal hemorrhage, more frequent absormading catastine, use and hemoglobin. Therefore,
do not use combination therapy with disofense sodium and an oral NSAID unless the
benefit outledge in the risk and conduct periodic biotratory productions.

6 ADVERSE REACTIONS

- 6 ADVERSE REACTIONS
 The following adverse reactions are discussed in greater detail in other sections of the libering.
 The following adverse reactions are discussed in greater detail in other sections of the libering.

 10 Electing, Userbara and Perioration (E. S.)
 11 Hepationscity [see WARRINGS AND PRECAUTIONS (S. 3)]
 12 Hepationscity [see WARRINGS AND PRECAUTIONS (S. 3)]
 13 Hepationscity [see WARRINGS AND PRECAUTIONS (S. 3)]
 14 Renal Toxicky and Hyperfacients [see WARRINGS AND PRECAUTIONS (S. 6)]
 15 Renal Toxicky and Hyperfacients [see WARRINGS AND PRECAUTIONS (S. 6)]
 16 Serious Sin Reactions [see WARRINGS AND PRECAUTIONS (S. 6)]
 17 Serious Sin Reactions [see WARRINGS AND PRECAUTIONS (S. 6)]
 18 Hemanicing Toxicky and WARRINGS AND PRECAUTIONS (S. 6)]
 18 Serious Sin Reactions [see WARRINGS AND PRECAUTIONS (S. 6)]
 18 Serious Sin Reactions [see WARRINGS AND PRECAUTIONS (S. 6)]

• Services Shin Reactions: [see WARNINGS AND PRECAUTIONS (5.9)]
• Intendispice; Tooks, [see WARNINGS AND PRECAUTIONS (5.9)]
• Intendispice; Tooks, [see WARNINGS AND PRECAUTIONS (5.9)]
• 1. Control of the Chical trials of a drug cannot be directly compared to rates in the clinical trials of the size of a drug cannot be directly compared to rates in the clinical trials of an drug and range in the clinical trials of the rates. See which the clinical trials are designed in the clinical trials of the rates and the clinical trials and the drug and range in the clinical trials and see which the clinical trials are conducted to the clinical trials and the clinical trials are clinically as the clinical trials are clinical trials. The clinical trials are clinically as the clinical trials. The clinical trials are clinically as the clinical trials are clinically as the clinical trials. The clinical trials are clinically as the clinical trials are clinically as the clinical trials are clinically as the clinical trials. The clinical trials are clinically as the clinical trials are clinically as the clinical trials. The clinical trials are clinically as the clinical trials are clinically as the clinical trials are clinically as the clinical trials. The clinical trials are considered as the clinical trials are clinically as the clinical trials are clinically as the clinical trials. The clinical trials are clinically as the clinical trials are clinically as the clinical trials are clinically as the clinical trials. The clinical trials are clinically as the clinical trials are clinically as the clinical trials are clinically as the clinical trials. The clinical trials are clinically as the clinical trials are clinically as the clinical trials are clinically as the cl

Table 1: Adverse Reactions occurring in ≥1% of patients treated with Diciofenac Sodium Topical

Treatment Group:	Diclofenac Sodium Topical Solution, 1.5% w/w Topical Placeb			
	N=911	N=332		
Adverse Reaction"	N (%)	N (%)		
Dry Skin (Application Site)	292 (32)	17 (5)		
Contact Dermatitis (Application Site)	83 (9)	6 (2)		
Dyspepsia	72 (8)	13 (4)		
Abdominal Pain	54 (6)	10 (3)		
Flatulence	35 (4)	1 (<1)		
Pruritus (Application Site)	34 (4)	7 (2)		
Diarrhea	33 (4)	7 (2)		
Nausea	33 (4)	3 (1)		
Pharyngitis	40 (4)	13 (4)		
Constipation	29 (3)	1 (<1)		
Edema	26 (3)	0		
Rash (Non-Application Site)	25 (3)	5 (2)		
Infection	25 (3)	8 (2)		
Ecchymosis	19 (2)	1 (<1)		
Dry Skin (Non-Application Site)	19 (2)	1 (<1)		
Contact Dermatitis, vesicles (Application Site)	18 (2)	0		
Paresthesia (Non-Application Site)	14 (2)	3 (<1)		
Accidental Injury	22 (2)	7 (2)		
Pruritus (Non-Application Site)	15 (2)	2 (<1)		
Sinusitis	10 (1)	2 (<1)		
Haltosis	11(1)	1 (<1)		
Application Site Reaction (not otherwise specified)	11(1)	3 (<1)		

6.2 Postmarketing Experience
In non-U.S. postmarketing surveilance, the following adverse reactions have been
reported during sort-approval use of dicidenac sodium. Because these reactions are
reported voluntarly from a population of uncertain size, it is not always possible to
reliably estimate their frequency or establish a cause releationship to drug exposure.

Abdominal pain, accidental injury, allergic reaction, asthenia, back pain, body odor, chest pain, edema, face edema, haltosis, headache, lack of drug effect, neck rigidity, pain

Leg cramps, myalgia

Merveus
Depression, dizines, drovishes, lethargy, paresthesia, paresthesia at application ste
Respiratory
Asthma, dypanea, laryngismus, laryngiss, pharyngiss
Shin and Appendages

>kun and Appendages

At the Application Site: Contact dermatitis, contact dermatitis with vesicles, dry skin, pruritus, reskin.

Other Skin and Appendages Adverse Reactions: Eczema, rash, pruritus, skin discoloration, urticaria

discoloration, urticana

Special Senses
Abnormal vision, blurred vision, cataract, ear pain, eye disorder, eye pain, taste perversion

Fortrioled dried studies showed that the concentrate use of NSAIDs and ensigners does of against does not produce any greater therapports, effect than the use of NSAIDs and ensigners does not produce any greater therapports, effect than the use of NSAIDs and expire was associated with a significant produce of the superior does not not a shading line NSAIDs and a NSAID and expire was associated with a significant produce of the superior does not not a shading line NSAIDs and NSAID a control of the contro oxin

coxin

Cox nect. ISANDS have produced elevations in plasma libium levels and reductions in renal libium clearance. The mean minimum lithium concentration increased 15%, and the renal clearance decreased by approximately 20%. This effect has been attributed to NSAID inhibition of renal prostaglandin synthesis.

During concentrated use of disclines: sodium and libium, monitor options for signs of libium buckly. hotresaste in direct. Concomitant use of ISAIDs and methotrevate may increase the risk of methotrevate toxicky (e.g., meutropenia, thrombocytopenia, renal dysfunction) buring concomitant use of diclofenac sodium and metho Concomitant use of diclofenac sodium and cyclosporine may increase cyclosporine's nephrotoxicky.

During concomitant use of diclofenac sodium and cyclosporine, monitor patients for signs or worsening renal function. Sump concommant use of discherac sodium and cyclesporine, monther patients for signs or environment use of months of the control of the contr

Circlemac and anticoappaints such as warfarin have a synegotic effect on bleeding. The concommant use of discliners, and anticoappaints from the steep regulated and such as a suffering have been required to the use of either drug above.

— Sention increase by plaidest, plays, a important rise in hermatiss. Case control and control epidemiologistic dates knowed that concerndant use of infering with prestron requirement and management of the use of either drug plays to the prestream of the prestr

Concomitant use of dictofenac sodium and pemetrexed may increase the risk of pemetrexed-associated myelosuppression, renal, and GI toxicity (see the pemetrexed prescribing information).

During concomitant use of dictofenac sodium and pemetrexed, in patients with renal impairment whose creating clearance ranges from 45 to 79 mL/min, monitor for myelosuppression, renal and GI toxicity. VSAIDs with short elimination half-lives (e.g., diciofenac, indomethacin) should be avoided for a period of two days before, the day of, and two days following administration pemetrexed

8 USE IN SPECIFIC POPULATIONS

emetrexed

8 USE IN SPECIFIC POPULATIONS

3.1. Pregnancy

Bib. Summar:

Bib. Summar:

Live of INSUIDs, Including dicinfers sodium, can cause premature closure of the fetal ductus arterious and fetal renal dyfunction leading to object/arminos and, in some cases, incendal renal respirate filescare of these ratios, limited oral and duction of dicinfers could make the second of the second oral second oral

nature Closure of Fetal Ductus Arteriosus

Use of NSAIDs, including diclofenac sodium, 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 olgohydramnios, and in some cases, neonatal renal impairment.

with case of feed read adjutances leading to algorithmic and in some cases, montal feral in great bandler. Tageding other particular in regarding other particular in regarding other particular in regarding other particular in regarding control of the particular in regarding in the particular in regarding control of the particula

etal/Neonatal 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 diclofenac sodium, can cause premature closure of the fetal ductus arteriosus (see Data).

Oligohydramnios/Neonatal Renal Impairment

Published iterature 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.

Olgohydramics/Recontal Renal Impairment: Nucleible studies op optimiration; propris describe maternal NSAD use at about 30 inchibited studies optimiration; propris describe maternal NSAD use at about 30 olgohydramics, and in some cases, necessaria renal impairment. These adverse in olgohydramics, and in some cases, necessaria renal impairment. These adverse in olicitation in an energia, after days to week of the relatives, although 400 initiation. In many; cases, but not all, the decrease in amminic fluid was transient and reversels with cases of the during. There was been a limited of case report of which were reviewable. Some cases of necontal renal optimization required treatment with transies proceedings, such as exchanged restrictions of oliginal with transies proceedings.

Methodological limitations of these postmirelating studies and reports historial tack of a control or group, infliend featurement of the production of the control or group, infliend featurement of the geographic and concomitant use of other medications. These limitations preclude establishing related sentimes of the risk of adverse fetal and neonated outcomes with of the control of the mostly preterm inflients, the generalizability of certain reported risks to the full-term inflient exposed to MRSIBs brough material use is uncertain use in pretent and the control of the control

Animal Data

Anima

Risk Summary
Based on available data, dicidence may be present in human milk. The developmental and health benefits of brasstefening should be considered along with the mother's clicitationed for Child-Risk and any political adverse effects on the breastefer infant.

Data the Child-Risk from the underlying melanest concludes.

Data the Child-Risk from the underlying melanest concludes the conclude of the concludes the conclude of the concludes the Linear terms of the concludes the concludes

postparking period.

3. I Familias and Males of Reproductive Potential
Infortility

Familiar
Based on the mechanism of action, the use of prestaginarian mediated ISSAIDS, including disclosure, sodium and provided register of control foliation, which has been decidence, sodium and provided register of control foliation, which has been shown that deministration prostaginaries synthesis inhibitors has the potential to don't be a second or so the provided of the control foliation. In the control foliation is the potential to don't be a second or so the control foliation. Consider without control foliation. In such as do ISSAIDs, including discherace, sodium, in women who have difficulties conceiving or whom or undergroup in severagional or feeting.

8.4 Pediatric Use
Safety and effectiveness in pediatric patients have not been established.

8.5 Gerharte: Use Ellerly galderns, compared to younger patients, are a greater risk for NSAID-associated scriuss cardiovascular, gastrointesinal, and/or renal adverse reactions. If the averticated bowerf to the ellerly patient valuesing have patient risks, start dosing at workcapted bowerf to the ellerly patient valuesing have patient risks, start dosing at WARMINGS AND PERCAUTIONS (5.1, 3.2, 5.3, 5.6, 5.13).
WARMINGS AND PERCAUTIONS (5.1, 3.2, 5.3, 5.6, 5.13).
Of the 911 patient readed with clicitients coolium is seven controlled, Plasse 3 clinical traits, 444 subjects were 65 years of age and over. There was no age-related difference on open-habeds after yint, 33 subjects, were 65 years of age and over inchire patients.
Or subjects 75 and over. There was no difference in the incidence of adverse events with horsy term prepared to discribes. South for the clicity population.

18 OVERDOSAGE
Symptoms following acute IISAID overdosages have been typically limited to lethorary, drowsiness, nausea, vomiting, and epigastric pain, which have been generally reversible with supportive cere. Gastroinetistan biolegaring has occurried, hypotheristanis, naute rend with supportive cere. Gastroinetistanis biolegaring has occurried, hypotheristanis, naute rend AND REGISTATION (5.1.5.5.5.4.5.6).

Manage patients with symptomatic and supportive cere following in ARSIAD overdosages. There are no specific artistations. Breads in an office commended due to a possibility of automatic particular symptoms and supportive cere following in dischlerate scalable particular symptoms and supported cere following in dischlerate scalable particular symptoms and supportive cere following in dischlerate scalable particular symptoms and symptoms and support symptoms and symptoms and symptoms and symptoms and symptoms are supported symptoms. The symptoms are symptoms and symptoms are symptoms and symptoms and symptoms are symptoms and symptoms. The symptoms are symptoms and symptoms are symptoms and symptoms are symptoms. The symptoms are symptoms are symptoms and symptoms are symptoms and symptoms are symptoms. The symptoms are symptoms are symptoms and symptoms are symptoms and symptoms. The symptoms are symptoms are symptoms are symptoms and symptoms are symptoms. The symptoms are symptoms are symptoms are symptoms. The symptoms are symptoms are symptoms are symptoms are symptoms and symptoms are symptoms. The symptoms are symptoms are symptoms are symptoms are symptoms. The symptoms are symptoms are symptoms are symptoms are symptoms are symptoms. The symptoms are symptoms are symptoms are symptoms are symptoms are symptoms. The symptoms are symptoms are symptoms are symptoms are symptoms are symptoms. The symptoms are symptoms are symptoms are symptoms are symptoms are symptoms. The symptoms are symptoms are symptoms are symptoms are symptoms are symptoms. The symptoms are symptoms are symptoms are symptoms are s

AL DESCRIPTION

Cholimans Solution Topical Solution USP, 1.5% w/w is a nonsterroidal anti-inflammatory drug, available as a clear, contriess to fairthy pric crange solution for topical application. Discription Solution Solution



Each 1 mL of solution contains 16.05 mg of diclofenac

The inactive ingredients in diclofenac sodium topical solution USP include: alcohol, dimethyl sulfoxide (DMSO, 45.5% w/w), glycerin, propylene glycol and purified water.

12.1 Mechanism Of Action
Dicbfenac has analysis, anti-inflammatory, and antipyretic properties.

The mechanism of action of dicbfenac sodium, like that of other NSAIDs, is not completely understood but involves inhibition of cyclosoxygenase (COX-1 and COX-2) Decimae is a potential member of processing and pro

12.3 Pharmacokinetics

After topical administration to healthy human volunteers of single and multiple maximum doses of dicbfenac sodium topical solution, 40 drops (approximately 1.2 ml.) to each kene (80 drops total dose), the following dicbfenac pharmacokinetic parameters were obtained: (see Table 2).

Table 2: Single-Dose (80 drops) and Multiple Dose (80 drops four times daily for 7 days) Diclofenac Sodium Topical Solution, 1.5% w/w Pharmacokinetic

Parameters			
Pharmacokinetic Parameters			
	Normal Adults [N=18] (Age: 18 to 55 years)	Normal Adults [N=19] (Age: 18 to 55 years)	
	Single Dose	Multiple Dose Four times daily for 7 days	
AUC _{0-t}	177.5 ± 72.6 ng.h/mL	695.4 ± 348.9 ng.h/mL	
AUC _{0-inf}	196.3 ± 68.5 ng.h/mL	745.2 ± 374.7 ng.h/mL	
Plasma C _{max}	8.1 ± 5.9 ng/mL	19.4 ± 9.3 ng/mL	
Plasma T _{max} (h)	11.0 ± 6.4	4.0 ± 6.5	
Plasma t _{1/2} (h)	36.7 ± 20.8	79.0 ± 38.1	
K _{el} (h-")	0.024 ± 0.010	0.011 ± 0.004	
CL/F (L/h)	244.7 ± 84.7*	**	

Dicbfenac systemic exposure from dicbfenac sodium topical solution application (4 times daily for 1 week) was approximately 1/3 of the dicbfenac systemic exposure from the Solaraze (dicbfenac topical gel) application (twice daily for 4 weeks).

Distribution
Diciofenac is more than 99% bound to human serum proteins, primarily to albumin

Disblanc diffuse this and so of the sproved filed. Officials into the joint occurs when plants levels are higher than those in the sproved filed, after which the process reverses and syroxial fluid levels are higher than plasms levels. It is not known whether officials in this high pirt plays a role in the effectiveness of disblance. Elimination Metabolism:

Net abolism:

Free dictriems metabolises have been identified in human plasma and urine. The metabolises include if hydroxy, -3-hydroxy, -45-dhydroxy, -45-d

The pharmacokinetics of diclofenac sodium topical solution has not been investigated in pediatric patients.

pediatr. Asce:

Pharmackinetic differences due to race have not been studied.

Drug Interaction Studies

Aspiria:

When MSAIDs were administered with aspiris, the protein binding of MSAIDs were reduced, although the clearance of the MSAID was no stated. The chical synfance makes of the MSAIDs with a spirit (see ABUG MTERACTIONS (7)).

Synfants design from the Committee of the MSAIDs with a spirit (see ABUG MTERACTIONS (7)).

13.3 Carcinogenesis, Mutagenesis, Impairment of Fertility Carcinogenesis

Technopmicity studies in mice and rats administered diciplena: sodium topical solution as a delary constituent for 2 years resulted in no significant increases in tumor incidence ad does up to 2 mightagles processponding to approximately 0.33 and 0.7- (MRHI) of diciplena: sodium topical solution (based on apparent bioavailability and body surface area compression).

surface area comparison). In a dermal carcinomicty study conducted in abino mice, daily topical applications of debefores column for their years at concentrations up to 0.03% disorderes column for debefores column for a column for their years and column for their years and column for their years and their column forgate. In a photococar mognitive, to surface of their years and years

Mutagenesis
Dischience, was not mutagenic or clastogenic in a battery of genotoxicity tests that included the bacterial reverse mutation assay, in with mouse it preploma point mutation assay, chromotomate of moute of preploma point mutation assay, clamotomate cells write, and an assay of tone matrices cells write, and an assay of tone matrices cells.

Impairment of Fertility
Fertility tradies have not been conducted with disclosines sociation topical solution. Dischience odium administrated to make and fermale rate at doses up to 4 regligation (1.4-dod of the Mitty) of dischience sociation packs allowed to a proper to bowokalatily and body surface area comparation) and not affect fertity. Studies have not been conducted to determine to salley on 160% on retility.

Ocube Effects

No adverse effects were observed using indirect ophthalmoscopy after multiple daily demail application to rate for 26 weeks and minings for 52 weeks of DMSO at twice the concentration found indicitiens south usepoles alouther. Nebstate studies of Demail or call admiration of DMSO in batch, storp and pay it described refractive change of or all admiration of DMSO in batch, storp and pay it described refractive change of a country of the contraction of the DMSO in batch, and the country of the DMSO in th

14 CLINICAL STUDIES

14 CLINICAL STUDIES

14. Studies in Orthoparchimis of the Knee

The use of disclience sodium trapical solution for the *restment of the signs and

The use of disclience sodium trapical solution for the *restment of the signs and

Clinical social social solution of the solution of the solution of the solution of the solution topical solution and solution of all ordinary for turnes as day of 12 weeks. Disclience excipients and one topical vericies solution (65.5% who PMSO with other excipients), applied directly but the solution (65.5% who PMSO with other excipients), applied directly but the solution (65.5% who PMSO with other excipients), applied directly but the solution (65.5% who PMSO with other excipients), applied directly but the solution (65.5% who PMSO with other excipients), and other solution of the solution

Table 3: Change in treatment outcomes after 12 weeks of treatment in one study of efficacy of Diclofenac Sodium Topical Solution, 1.5%

w/w				
	Study I Mean baseline score and mean change in efficacy variables after 12 weeks of treatment			
Efficacy Variable				
	Mean Baseline score	Diclofenac Sodium Topical Solution	N=154 Topical placebo1 N=155	Topical vehicle ² N=161
WOMAC pain score (Likert 3.1, 0- 20)	13	-6.0	-4.7	-4.7
WOMAC physical function (Likert 3.1, 0-68)	42	-15.7	-12.3	-12.1
POHA (0-4)	2.3	-1.0	-0.4	-0.6
² placebo formulation included 2.3 ² vehicle formulation included 45.5				

Efficacy Variable	Study II Mean baseline score and mean change in efficacy variables after 12 weeks of treatment			
	Mean Baseline score	Diclofenac Sodium Topical Solution N=164	Topical vehicle ¹ N=162	
WOMAC pain score (Likert 3.1, 0- 20)	13	-5.9	-4.4	
WOMAC physical function (Likert 3.1, 0-68)	42	-15.3	-10.3	
PGA (0-4)	3.1	-1.3	-1.0	

16 HOW SUPPLIED/STORAGE AND HANDLING

16 HOW SUPPLIED/STORAGE AND HANDLING

Dichlene Scolum Loyde Solubation (19.1 15) wile is supplied as a clear, colorless to fairly pirk carges esolution containing 18.05 mg of dicherace sodium per mid obtained, in a white hind from by polyethylene bodite with a white be density polyethylene doctor.

NOC Number and Size

5 T.O.2. (150 mil bottle in cartons of one

NOC 4 68180-5380.1

Storage

Storage 25°C (77°F); excursions permitted to 15 to 30°C (59 to 86°F) [See USP Controlled Room Temperature].

17 PATIENT COUNSELING INFORMATION

1) PATIENT COUNSELING INFORMATION
After the petient read the FGA approved patient bleing (Nedication Guiste) and Instructions for the Inflat accompanies and injercition dispersed. Inform patients, decidence, souther register and interest of the Inflat and Inflat a

Gastromotestinal sisecong, Useration, and verioration Advise patients to report symptoms of ulcerations and bleeding, including epigastric pain, dyspepsis, melena, and hematemesis to their health care provider. In the setting of concommant use of low-dose supplier for cardiacy prophysics, finding malents of the increased risk for and the signs and symptoms of Gi bleeding [see WARNINGS AND PRECAUTIONS 5.2].

PRECUTIONS 5.2].

Hypatotoxicity Mepatotoxicity warning signs and symptoms of hepatotoxicity (e.g., nausea, fissigus, lethrary, puratus, derrikes, junnouse, right upon quadrate trademess, and "thus for upon signs, the property program of the prog

See CUIT RAINDLATIONS (4) and WARRINGS ARU PRE-CAUTIONS (5.7)].

Serious Skin Reactions <u>including DRESS</u>

Advise patients to stop <u>taking</u> diciofenac sodium immediately if they develop any type o rash <u>ar fever</u> and to contact their healthcare provider as soon as possible (see WARRINGS ARU PRE-CAUTIONS (5.10, 5.11)).

Female Fertility

Framis Fortilly lotter programs women to avoid use of disoletime, softem topical solution and other NSAIDs starting at 30 weeks gestation because of the risk of the prevaluer cointing the feetal ductus attension. If treatment with discherines sodium topical solution is needed for a program woman between about 20 to 30 weeks spectation, soften her that the 4th hours [see Warrings and Precaudions (5.11) and the in Specific Populations (8.1)]. Fetal Toolsky

***Retail lookicky**
Inform pregnant women to avoid use of diciofenac sodium topical solution and other INSAIDs starting at 30 weeks gestation because of the risk of the premature closing of the feel ducture. **arterousis (see MARMINGS AND PRECAUTIONS (5.11) and USE W SPECIFIC POPULATIONS (8.1).

SPECIFIC POPULATIONS (8.1).
Avoid Concomitant Use of NSAIDs

Avoid Concommant Use or INAIUS Inform patients that the concommant use of diciofenas sodium topical solution with other INAIUS or solicylates (e.g., offlinios), solution to not recommended due to the increased risk of guaranteemant buskley, and title or no crease in efficacy, fores that NIAIUS may be present in "over the counter" medications for treatment of colds, fever, or insormis.

Use of NSAIDS and Low-Dose Aspirin

Inform patients not to use low-dose aspirin concomitantly with diclofenac sodium topical solution until they talk to their healthcare provider [see DRUG INTERACTIONS (7)].

Eye Exposure

Eye Exposure Instruct patients to avoid contact of dicbfenac sodium topical solution with the eyes and mucosa. Advise patients that if eye contact occurs, immediately wash out the eye with water or saline and consult a physicisin if irritation persists for more than an hour. Prevention of Secondary Exposure

Instruct patients to avoid skin-to-skin contact between other people and the knee(s) to which diclofenac sodium topical solution was applied until the knee(s) is completely dry.

Application Site Reactions

Diciofenac sodium topical solution can cause a localized skin reaction at the application site. Advise patients to contact their physicians as soon as possible if they develop any type of localized application site rash.

- type of localed application set erab.

 Special Application instructions

 Instruct patients not to apply discherac sodium topical solution to open skin wounds

 Instruct patients not to apply discherac sodium topical solution to open skin wounds

 Instruct patients is on the control of the con

and the parameter to minimize or avoid exposure of treated kinetics to indust of artificial sunight.

The brands listed and trademarks of their respective owners and are not trademarks of role redorse Lipin Pharmaceuticals, inc. or its products.

Lipin Pharmaceuticals, inc.

Lipin Lipi

Dispense with Medication Guide available at: www.lapin.com/dicboodropicahol-mg.pdf
Medication Guide for Monateroidal Anti-inflammatory Drugs (MSAIDs)
What is the most important information is should know about medicines called
Nonsteroidal Anti-inflammatory Drugs (MSAIDs)
MSAIDs can cause serious side effects, including;

• Increased risk of a heast attack or stroke that can lead to death. This risk

• with increasing doses of MSAIDs

• with increasing doses of MSAIDs

• with brogge use of MSAIDs not take NSAIDs right before or after a heart surgery called a "coronary ery bypass graft (CABG)."

artery bypass graft (CABG).** A recent heart attack, unless your heabthcare provider tells you to. You may have an increased risk of another heart attack. If you train to You may have an increased risk of another heart attack if you train the your provider tells you to You may have an increased risk of the other tells of the your provider tells you have a provider to you have a supplied to the other your and tens (perforation) of the septiment of the other your anything the young the your proposes. I will not you have y

* most may cause occur
 The risk of settling an ulcer or bleeding increases with:
 * path history of stometh ulcer, or stometh or insetted bleeding with use of NSAIDs
 * path history of stometh ulcer, or stometh or insetted bleeding with use of NSAIDs
 * horearing dose of NSAIDs
 * horearing dose of NSAIDs
 * smoking
 * smoking
 * object on the stomethy of the stomet

NSAIDs should only be used:

• exactly as prescribed

• at the lowest dose possible for your treatment

• for the shortest time needed

What are NSAIDs?

What are Results:

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

Who should not take NSAIDs?

Do not take NSAIDs:

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

other NSAIDs. • right before or after heart bypass surgery.

- rgib before or after heart bypass surgery.

 Refore taking Kallofs, tell your healthcare provider about all of your medical conditions, including if your healthcare provider about 2 days of the provider and the provider and

your rouy. You should not take NSAIDs after 20 weeks of pregnancy.

**A or breatflending of pints to treast feet.

Tell your healthcare provider about all of the medicines you take, including the period of the pe

Get emergency help right away if you get any of the following symptoms: • shortness of breath or trouble breathing • shortness of breath or trouble breathing • chest pain • swelling of the face or throat • swelling of the face or broat • weakness in one part or side of your body

- Stop taking your NSAID and call your healthcare provider right away if you get any of the following symptoms:
- vomit blood
 more tired or weaker than usual
 there is blood in your bowel movement or it is black and sticky like tar

- there is bboom is p...
 diarrhea
 it ching
 unusual weight gain
 your sikn or eyes look yelow
 skin rash or blettes with fever
 skin rash or blettes with fever
 swelling of the arms, legs, hands and feet
 fluike symptoms

If you take too much of your NSAID, call your healthcare provider or get medical help right away.

medical nep 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. You may also report side effects to Lupin Pharmaceuticals, Inc. at 1-800-399-2561 or visit our website at www.lupinpharmaceuticals.com.

- Other information about NSADD

 Appin is an NSADD but 4 soles not increase the chance of a heart attack. Appin can

 Appin is an NSADD but 4 soles not increase the chance of a heart attack. Appin can

 be soled and of the chance of the chance of a heart attack. Appin can

 be some fine of resistance.

 Some RSADDs are sold in lower doses without a precription forwer because I, risk

 Some RSADDs are sold in lower doses without a precription forwer because I, risk

 Some RSADDs are sold in lower doses without a precription forwer than 10 days.

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 have them them. It is more information about NSAIDs, this with your healthcare provider that is written for health professional.

The Medication Guide has been approved by the U.S. Food and Drug Administration. Neuroland Professional.

The Medication Guide has been approved by the U.S. Food and Drug Administration. Neuroland Professional. Professional Professiona

Steps for using diclofenac sodium topical solution:

Step 1. Wash your hands with soap and water before applying diclofenac sodium topical solution:

Step 2. Put 10 drops of diclofenac sodium topical solution **either** on your hand **or** directly on your knee (see **Figure A**).





Step 3. Spread dicibfenac sodium topical solution evenly on the front, back and sides of your knee (see **Figures 8** and C). Repeat steps 2 and 3, three times so that your knee is completely covered with a **total** of 40 drops of dicibfenac sodium topical solution.



Step 4. If your healthcare provider has prescribed diclofenac sodium topical solution for both kinese, repeat steps 2 and 3 for the other kines.

After you use dichlofenac sodium topical solution:

Wash your hands with so

topical solution.

Do not:

I created kine or allow another person to touch the kines treated with

dischlerate solden topical solution until your kines in completely dry.

cover your kines which challenge until your kines in completely dry.

cover your kines which challenge until your kines is completely dry.

medicines on your kines until a is completely dry.

medicines on your kines until a is completely dry.

until his anise where or a both for a less to 3 nimites after you put dischlerate sodium

use heating pads or cover the treated area with bandages where you have applied
dischlerates solution.

use heating pads or cover the treated area with bandages where you have applied
dischlerates solution.

Until the properties of the complete in the complete in the control of the complete in the complete

*store accentance southun bipocal solution 25°C (D*P*I); accurating permitted to 15 to 30°C (D*9 to 80°P). See this Centrole floor morrogenature.

Keep discherace southun topical solution and all medicines out of the reach of children. This instructions for the has been approved by the U.S. Food and Drug Administration.

The largest install control of their respectitio general seed are not trademosks of the largest large floor of the seed of

January 2021 266841

266841
Dictofenac Sodium Topical Solution
1.5% w/w
NDC: 68180-538-01
Container Label: 5 FL.OZ. (150 mL) bottle



Dictorenac Sodium ropical Solution
1.5% w/w
NDC: 68180-538-01
Carton Label: 5 FL.OZ. (150 mL) bottle in cartons of one



P	roduct Info	rmation				
Product Type			HUMAN PRESCRIPTION DRUG	Item	Code (Source)	NDC:68180-538
Ro	oute of Admi	nistration	TOPICAL			
A	ctive Ingre	dient/Activ	e Molety			
				Basis of Strength	Strength	
DI	DICLOFENAC SODIUM (UNI: QTG126297Q) (DICLOFENAC - UNI: 14408QL0L1)				DICLOFENAC SODIL	M 16.05 mg in 1 mL
In	active Ingr	edients				
Ingredient Name					Strength	
M	COHOL (UNI:)	IKS95@V9CMI)	•			
DI	METHYL SULF	OXIDE (UNI: Y	OWEV969BH)			
	YCERIN (UNI:					
	OPYLENE GLY		19Q167V3)			
Pi	ackaging				tarketing Start	Marketing Er
*	Item Code		Package Description		Date	Date Date
	NDC:68180- 538-01	1 in 1 CARTO	N	07	06/2016	
1		150 mL in 1 fi Combination	IOTTLE, DROPPER; Type 0: Not a Product			
М	larketing					
	Marketing Category	Appli	cation Number or Monograph Citation	Ma	rketing Start Date	Marketing En
	DA .	4ND4204			5/2016	

Labeler - Lugin Primmendicals, Inc. (89333071)

Registrant - Lunu (18730181)

Establishment

Suns Address Date

Unit (1870) Dasbess Operations

Unit (1870) Media Collegistic State

Unit (1870) Med

evised: 12/2023 Lupin Pharmaceuticals