MELOXICAM- meloxicam tablet Aphena Pharma Solutions - Tennessee, LLC

INCORLIGHTS OF PRESCRIBING INFORMATION Three highlights do not include all the information meetind to use MELOXICAM TABLETS USP, cafely and effectively. See full prescribing information for MELOXICAM TABLETS USP.

MELOXICAM Tablets USP, for oral use

	WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS
	See full prescribing information for complete based warning.
•	Nonsteroldal anti-inflammatory drugs (NSAIDs) cause an increased risk of serious cardiovascular thrombotic evens, including myocardial infarction and streke, which can be fatal. This risk may occur early in transment and may increase with duration of use (5.1)
•	Melasicam tablets are contraindicated in the setting of coronary artery hypass graft (CABG) surgery (4, 5.1)
•	NSAUNC cause an increased risk of serious gastrointestinal (G1) adverse sevents including bleeding, underation, and performation of the stomach car intestinas, which can be faith. These sevents can occur at any time during use and without worning symptoms. Inderly patients and patients with a prior history of pentic altered floases and/or call bleeding are at restart risk (for serious G1 events 15.2).

Based Younning 2020 Microkowa nd Youg, Javedie Rheumaniki Arhivitis (IBA) Paucieticale and Polyanicular Course (L1) 6/2016 Dacaga and Administration, General Diving International (J.) 16/2016 Warning and Procanasa, Caefformacular Thomsteel Course (L1) 12/2016 Warning and Procanasa, Uner Rheura and Kenthin (RLA) Paucieticale and Polyanicular Course (L4)6/2016 Warning and Procanasa, Uner Rheura and Kenthin (RLA) (L2) 2016

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BOSAGE AND ADMENESTRATION Use the lowest effective douge for the shortest duration consistent with individual patient treatment goals (2.1) = 0.43 (2.2) used 8.43 (2)

.----Starting dose: 7.5 mg once daily Dose may be increased to 15 mg once daily • JRA (2.4):

To so go core daily in children 100 kg.
 Melosizan Tables are not interchangeable with approved formulations of oral melosicam even if the total milligram strength is the same (2.6).

Meknican Tables USP: 7.5 mg and 15 mg (3)

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FULL PRESCRIBING INFORMATION: CONTENTS* 1 INDICATIONS AND USAGE 1.1 Osneardtrifis (OA)
 1.2 Rhoumaold Arthrifis (RA)
 1.2 Rhoumaold Arthrifis (RA)
 1.3 Inventile Rhoumaold Arthrifis (JRA) Pauciarticular and Polyarticular Course
 DOSAGE AND ADMINISTRATION
 2.1 General Dosing Instructions

2.1 Contential Doising insertations 2.2 Oktoorarbitriis 2.3 Rhoumanoid Arthritis 2.4 Juwnite Rhoumanoid Arthritis (IRA) Pauciarticular and Polyarticular Course 2.5 Renal Impairment

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5.13 Laboratory Monitoring ADVERSE REACTIONS

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FULL PRESCRIBING INFORMATION

WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS

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1 INDICATIONS AND USAGE

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L3 Discontation are indicated for reliat of the signs and symptoms of theoremical arteristic (see Clinical Missicarabits are indicated for reliat of the signs and symptoms of theoremical arteristic (see Clinical Station (44.1)).

Statise (14.1), L3 Javonile Rheumanisid Arthrifis (JRA) Panciarticular and Polyarticular Course Meloscian tables are inficated for relief of the signs and symptoms of panciarticular or polyarticular course Javorile Rheumanid Arthrifis inpatients who weigh 560 kg [see Donage and Administration (24.1) and Chica's datase(14.2)).

2 DOSAGE AND ADMINISTRATION

2 DOSACE AND ADMINISTRATION 22 Octaval bosons in the section. Carefully consider the potential hearths and risks of Melosicantubles and other treatment options consistent with individual patient reasoning and pices Wanning, and Procentions (7). Alter observing the response to india therapy with Melosicantublers, adjust the dose to suit an individual patient week.

Identified particle werk in the second secon

Jecurating the dates to ing user any. 22 **Alexanatio Advectivit** For the reliaf of the signs and symptoms of throatmanial articlist, the recommended starting and minimums *even* along the Maxion and along in 17.2 mg, user a daily. Some patterns may receive additional both fity usersating the dates in 17.3 mg partice daily. 24 **Jecurity Elements of Description Constraints**, the recommended of additional **Descriptions** For the resonance of parents' horizontal arbitrist, the recommended of additional **Descriptions** For the resonance of parents' horizontal arbitrist, the recommended of additional **Descriptions** For the resonance of parents' horizontal arbitrist, the recommendance of the data data data data. Maximum **Descriptions Descriptions Descript**

2.5 Renal Impairment The use of Meloxicam tablets in subjects with severe renal impairment is not recommended.

In patients on hemodialysis, the maximum dosage of Melosican tablets is 7.5 mg per day [see Clinical Pharmacology (12.3)].

Phoreacology (22.3). 22 Non-Inter-Changeability with Other Fermulations of Melnxicaan Molosican nables have on choose equivalent systemic exposure to obser approved formulations of oral melosican. Therefore, Melosican nables are on inter-changeabile with other formulations of oral melosican approach provide the systemic and an antipart of the same. Do not subfatture similar does aregular to Melosican nables with other termination of our almosican product.

DOSAGE FORMS AND STRENCTHS
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4 CONTRAINDICATIONS

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5 WARNINGS AND PRECAUTIONS

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they occur. There is no consider vertices that concurrence use of applicating stages the increased cities of orders CV therefore events associated with NSAD mark. The concurrence as of application and a NAAD, such as of the concurrence of the concurrence of the concurrence of the concurrence of the CV there are an an and the concurrence of the concurrence of the concurrence of the CV there are an an and the concurrence of the concurrence of the concurrence of the SV the large, constructed for a large start of LAC and CV and CV and the concurrence of paths the first SVAD are accommodated and is setting of LAC and the CONcurrence of the concurrence of the start of the start of the CONCurrence of the start of the CONCurrence of the concurrence of the start of the CONCurrence of the start of the CONCurrence of the concurrence of the start of the CONCurrence of the start of the CONCurrence of the concurrence of the concurrence of the start of the CONCurrence of the start of the CONCurrence of the concurrence of the concurrence of the start of the CONCurrence of the start of the CONCurrence of the concurrence of the concurrence of the start of the CONCurrence of the start of the CONCurrence of the concurrence of the concurrence of the start of the CONCurrence of the concurrence of the start of the CONCurrence of the concurrence of the concurrence of the start of the concurrence of the concurr

Nothin are communicated in the sening of LARL just calendarchoson (e)[]. <u>Nath Distant</u> Observational analises conducted in the Danish National Polytory have demonstrated in patients traverate with NSDBs in the polymery diverse and traverante. In this same cohere, the incidence or class in the cases mentioning in the first work of a variance, in this same cohere, the incidence or class in the processory and intervent the processory of the polymery of the polymery of the polymery processory and intervent the processory of the polymery of the polymery of the polymery of the polymery are not first year polymery. The polymery of polymery is a polymery processory and the polymery polym

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

12.1 Curvations that Blocking, Ukrasian, and Perlovation NNARN, keileding subscience, encourse serious paraintensiteding [2], sherves events technique interastical, biological contention, and periodical of the explanation, structure, small intensities, to traject interastical, biological, Thue serious adverse events catassets are applied by the encourse of the encourse periodical encourses of the encourse of the encourse of the encourse of the encourse of the periodical encourse of the encourse of the

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Learning on evaluate of converging the long manufactor (7): 23 Happanostic). Elevation of ALT or AST (here or more sizes the upper limit of normal (ULNI) have been reported proprisminely 1% of NSLB-rosen adjustics in Clickel traits. In Addition, rate, conversions fault, cases of severe hepatic isjury, including fulminant hepatitis, liver necrosis, and hepatic failure have been reported.

Elevations of ALT or AST (less than three times ULN) may occur in up to 15% of patients treated with NSAIDs including meloxicam.

Inform patterns of the varieties spins and symptoms of hopentrus/city (e.g., statuss, farigue, behavior, duration, particular, spin topologic and and there was and the facility of symptoms). If clicity is pre-toring the status of the pattern level (in a hope) the functional the level and status (12.5). A status of the sta

drangy. ES Jenet Fallers and Edenas The Cotth and radioised NSAD Trialistic Collaborationness analysis of randomized cosmolied trials memorande an agroundly non-fold increase ishospitalizations for heart failers (seCOS-2 memorander) and agroundly on fold increase ishospitalization for heart failers (seCOS-2 har Danish National Registry study of patients with heart failers, NSAD are increased her risk of ML hospitalization for heart failers, add doub.

Addisionally, fluid returnion and edema have bren observed in some pasterns treated with NSAIDs. Use of molocicammy bluer the CV effects of several therapeutic agress used to treat these medical conditions (e.g., districts, ACE inhibitors, or angiotensin receptor blockers [ARBs]) [see Drug himterictions (7)].

Avoid the use of Meloxicam in patients with severe heart failure unless the benefits are expected to outweigh the risk of worsening heart failure. If Meloxicam is used in patients with severe heart failure monitor patients for signs of worsening heart failure. 5.6 Renal Toxicity and Hyperkalemia

al Toxicity

Renal Toxicity Long-term administration of NSAIDs, including Meloxicam, has resulted in renal papillary necrosis, renal insufficiency, acute renal failure, and other renal injury.

Read tocidy has also been seen is agained in holes are all protoglateline have a comparatory tool is the minimum of experiments in the second second second second second second dependent enderstone in processing tank for each second ACE in histories or AREs, and the shelfy: Discontinuation of NADD therapy is smally followed by recovery is the preventioned second s

The renal effects of Meloxicam may hasten the progression of renal dysfunction in patients with preexisting renal disease. Because some Meloxicam metabolites are excreted by the kidney, monitor patients for signs of worsening renal function.

Increases in serum potassium concentration, including hyperkalentia, have been reported with use of NSAIDs, even in some patients without renal impairment. In patients with normal renal function, these effects have been antibuted to a hyporentiemetric hypolablestreorism status.

5.7 Anaphylactic Reactions Melosizem has been associated with anaphylactic reactions in patients with and without known hypersensitivity to meloscicam and in patients with aspirin-sensitive asshma [see Contraindications (4) and Worrings and Precoutions (5.8)].

Seek emergency help if an anaphylactic reacti

5.8 Exacerbation of Asthma Related to Aspirin Sensitivity

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5.10 Premature Closure of Fetal Ductus Arteriosu Meloxicam may cause permaner closure of the feal datus anteriosus. Avoid use of NSAIDs, including Meloxicam, in pregnant women starting at 30 weeks of gestation (third trimester) [see Use in Specific Populations (8.1)].

5.11 Hematologic Toxicity

5.11 Hemsthejk Textby Amerika has eccured in SK3D-zonard patients. This may be due to occult or grows blood loss, fluid remains, has an incomplicity due club difference or sphenopartics. If a plant transfer with Model Skinstandhuo Markan and Markan Skinstandhuo and skinstandhuo and skinstandhuo and skinstandhuo complicational distance of the skinstandhuo and skinstandhuo and skinstandhuo and plantandhuo and skinstandhuo and

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

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6 ADVERSE REACTIONS

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the laze or hip to compare the efficacy and safety of Meloxicam with placebo and with an active control. Two 12-week multicenter, double-blind, randomized trials were conduced in patients with rheumatoid arthrifts to compare the efficacy and safety of Meloxicam with placebo.

Table 1a depicts adverse events that occurred in 25% of the Melosican treatment groups in a 12-week placebo- and active-controlled outcoarthrifts trial. Table 1b depicts adverse events that occurred in 25% of the Melosican treatment groups in two 12-week placebo-controlled rehumand a strifts triats.

	Placebo	Meloxicam 7.5 mg daily	Meloxicam 15 mg daily	Diclofenac 100 mg daily
No. of Patients	157	154	156	153
Gastro intestinal	17.2	20.1	17.3	28.1
ubdominal pain	2.5	1.9	2.6	1.3
liarrhea	3.8	7.8	3.2	9.2
lyspepsia	4.5	4.5	4.5	6.5
Flatulence	4.5	3.2	3.2	3.9
lausea	3.2	3.9	3.8	7.2
lody as a Whole				
accident household	1.9	4.5	3.2	2.6
dema ¹	2.5	1.9	4.5	3.3
all	0.6	2.6	0.0	1.3
fluenza-like symptoms	5.1	4.5	5.8	2.6
entral and Peripheral Nervous vstem				
lizziness	3.2	2.6	3.8	2.0
eadache	10.2	7.8	8.3	5.9
espiratory				
harymitis	1.3	0.6	3.2	1.3
opper respiratory tract infection	1.9	3.2	1.9	3.3
kin				
ash ²	2.5	2.6	0.6	2.0

Abdominal pain NOS*	0.6	2.9	2.3
Dyspeptic signs and symptoms [†]	3.8	5.8	4.0
Nausea*	2.6	3.3	3.8
General Disorders and Administration Site O	onditions		
Influenza-like illness*	2.1	2.9	2.3
Infection and Infestations			
Upper Respiratory tract infections-	4.1	7.0	6.5
pathogen class unspecified [†]			
Musculoskeletal and Connective Tissue Dis-	arders		
Joint related signs and symptoms [†]	1.9	1.5	2.3
Nervous System Disorders			
Headaches NOS"	6,4	6.4	5.5
Skin and Subcutaneous Tissue Disorders			

Rash NOS⁵ 1.7 2.1 MoDRA performed serme mousee, abdominal pain NOS, influenza-like likews, headacties NOS, and rash NOS MoDRA high bend irom (preformed serme), dyspeptic high and symptomic (dyspeptic) aggravated, erectation, gaseizminual retarious, apper empiritory tract infection-pathogen sumpecided (lowingthe NOS, handlash NOS), junit related signal on all symptomic (methoday, antholog aggravated, juni certainto, the restanding)

The adverse events that occurred with Meloxicamin 22% of patients treated short-term (4 to 6 weeks) and long-term (6 months) in active-controlled osteoarthrifts trials are presented in Table 2.

	4-6 Weeks Co	Trials	6 Month Cor	
	Meloxicam 7.5 mg daily			
No. of Patients	8955	256	169	306
Gastro intestinal	11.8	18.0	26.6	24.2
Abdominal pain	2.7	2.3	4.7	2.9
Constitution	0.8	1.2	1.8	2.6
Diarrhea	1.9	2.7	5.9	2.6
Dyspensia	3.8	7.4	8.9	9.5
Flatulence	0.5	0.4	3.0	2.6
Nausea	2.4	4.7	4.7	7.2
Vomiting	0.6	0.8	1.8	2.6
Body as a Whole				
Accident household	0.0	0.0	0.6	2.9
Edema*	0.6	2.0	2.4	1.6
Pain	0.9	2.0	3.6	5.2
Central and Peripheral Nervous Sy	stem			
Dizziness	1.1	1.6	2.4	2.6
Headache	2.4	2.7	3.6	2.6
Hematologic				
Anemia	0.1	0.0	4.1	2.9
Mus culos keletal				
Arthralgia	0.5	0.0	5.3	1.3
Backpain	0.5	0.4	3.0	0.7
Psychiatric				
Insomnia	0.4	0.0	3.6	1.6
Respiratory				
Coughing	0.2	0.8	2.4	1.0
Upper respiratory tract infection	0.2	0.0	8.3	7.5
Skin				
Pruritus	0.4	1.2	2.4	0.0
Rash [†]	0.3	1.2	3.0	1.3

Default Distance Viscancian foregroups 0.1 0.4 2.4 1.3 Viscancian foregroups 0.3 0.4 4.7 6.9 * Vitio preferred names, neuk nodes deserve particului, al deserve profession, al neuk serve preferred and neuk serve p

Higher doses of Meloxicam (22.5 mg and greater) have been associated with an increased risk of serious GI events; therefore, the daily dose of Meloxicam should not exceed 15 mg.

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7 DRUG INTERACTIONS

See Table 3 for clinically significant drug interactions with meloxicam. See also Warnings and Precautions (5.2, 5.6, 5.11) and Clinical Pharmacology (12.3).

Table 3 Clinically Significant Drug Interactions with Melusican
Drugs dat Interfere vilh Hemostasis
Environment autore environment env
Marvemention: Munices parlents with conconstant use of Melonican with anticoagulants (e.g., varlaris), antiplatelet agents (e.g., aspiris), where/we servotation responde inhibitors (SSBb); and terenomics more pipeline responde inhibitors (SSBb); for signs of Meeding (see Warsing and Procandom (5.11)).
Napite Existed meet Channelle ditated suders showed for the concentrative of NSADD and analysis, does of anisotropy as conducted and the concentrative of an NSADD and assistives as assisted with a similarant increased indexes of Claberter tractions as ommered to use of the NSADD along two Warrings and Proceedings (2)
Encomitant use of Meloxican and low dose applin or analyses'c doses of aspirin is not generally recommended because of the increased risk of bleeding (see Warnings and Procardious (5.11)). Meloxicam is not a substante for low dose aspirin for cardiovascular protection.
ACE Inhibitors, Augiotensis Receptor Blackers
SUDs may dening the authypensitive effect of anginesis converging energies (AC2) inhibits, adjoined in Converging (angine (AC2)) inhibits or Add State (RAB), to the shele (AC2) inhibits or Add State (RAB) in the shele (AC2) inhibits or Add State (RAB) in the shele (RAB) in the s
Intervention: During concentrate are of Molecican and ACL inhibitors, ARBs, to bus, ARBs, to present to ensure that the desired blood pressure is establed. During concentrate are of Molecican and ACL inhibitors or ARBs in patients who are elderly, volume-depleted, or lave impaired read function, monitor for signs of vorsenting read functions. ARBs, to bus, ARBs, t
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Baind addes, as well a post-
tervention: buring concentratures of Melotiscian with disordes, observe patients for signs of versioning real fraction, in addition to assuring disorder efficacy including and Psecanions [56]).
Lihim
Cinical Impact, DSAIDs have produced elevations in plasma lithium levels and reductions in renal lithium 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 postagiandia synthesis [see Clinical Pharmacology (12,3)].
Intervention: During concentrat use of Melonican and lithium, monitor patients for signs of lithium suscity.
Methorexate
Clinical Impact: Concomitant use of NSAIDs and methotrexase may increase the risk for methorexase may increase the risk for methorexase toxicity (e.g., neuroperia, thrombocytopenia, trend dysfunction).
terrorentica: buring concentiarase of Molosican and nedosexue, motior patients for mediorescue molecy.
Cycles parine
Clinical Impace: Concominant use of Moloxicam and cyclosporine may increase cyclosporine's nephrotoxicity.
Intervention: During concentrat use of Melonican and cyclosportine, meniner patients for signs of worsening renal function.
NSAIDs and Salicylaus
Clinical Impace: Euroconstant use of melosicam with other NSAIDs or salicylans (e.g., diffunisal, salsalam) increases the risk of GI toxicity, with little or no increases in efficacy [see Warnings and Precautions (5.2)].
Intervention: The concentration use of neuloscican with other NSAIDs or salicylans is not recommoded.
Principal
Clinical Impact: Concomitant use of Moloxican and pometexed may increase the risk of penaterexed-associand myelosuppression, renal, and GI taxicity (see the penaterexed pescribing information).
During conconstant use of Melostican and permetexed, in patients with renal impairment whose creatinine clearance ranges from 45 to 79 mL/min, monitor for myelosuppression, renal and GI toxicity.
termention: patients taking melonican should immerphologing for at least five days before, the day of, and two days holitowing permervand administration.
In particus with cruation clearance below 45 al. Juin, the concentrat administration of mole science with permetered is not recommended.
LUSE IN SPECIFIC POPULATIONS

A section of the sect

There are no suffices on the effects of Meloxican during labor or delivery. In animal studies, NSAIDs, including meloxicam, inlibit prostaglandin synthesis, cause delayed parturition, and increase the incidence of stillbirth.

Data

. Animal Data

A shard Mut Mutan Mutan

Data Asimi Data Molocican was presert in the milk of lactating rats at concentrations higher than those in plasma. B.3 Females and Males of Reproductive Potential

Infertility

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Fonds
Based on mechanism of action, the use of prestagilation-medianed SSADDs, including Medvatian, and during a prevent rupture of cardinal taillistics, which has how an acticuted with reventible infertibly inhibitors have been prevented to discontegorous galaxies. The second second

8.4 Pediatric Use

The safety and effectiveness of meloxicam in pediatric JRA patients from 2 to 17 years of age has been evaluated in three clinical trials [see Dosage and Administration (2-3), Adverse Reactions (6.1) and Clinical Studies (14-2)].

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8.6 Henatic Imnairment

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Ex Hogds Impairment, Nortes with while the understare length (impairment, Borten with while the observation of the second sec

10 OVERDOSAGE Symposs following acues NSAID overdesages have been specially limited to behave downsizes, mouse, vositing, and objestic juin, which have been generally reversible with supportive care. Garavinensiable Beenfing has occurred. Hypermission, acute renal failure, resplantery depression, and com have occurred, bewere mer give behavioring and Proteations (15, 25, 25, 45, 56).

Manage paierns with symptomatic and supportive care following an NSAID overdosage. There are no specific and/one. Consider movies: and/or actioned detected gibts 100 games in adults, 10 × 2 games for human of inspections. The specific adult and the specific adult and the specific adult of the specific adult and the specific adult and the specific adult adu

There is limited experience with melosican overdosage. Cholestynamine is known to accelerate the clearance of ankoncam. Accelerated removal of melosicam by 4 goal doses of cholestynamine given following an overdosageman and in a clinical and anticationation of cholestynamine given following an overdosage the state of the stat

11 DESCRIPTION Moloxican Tables USP are a nonstrenidal anti-inflammony drug (NSAID). Each tablet contain 7.5.7 or 15 ng moloxican for on al administration. Moloxicam is chemically designated as 4-hydroxy-2-methyl-K-G-anthyl-2-diaastyly-21/2-12-bromobilizato-2-achoxanide-1-1-dioxide. The moloxic wights 353.4.8 ks antipriced Journal is CapityNg-Oky and I that following screenard formula:



Meloxicam is a passel yellow solid, practically insoluble in water, with higher solubility observed strong acids and bases. It is very slightly soluble in mechanol. Meloxicam base an apparent particlon coefficient (og 1999 pp = 0.1 in no-cambifietp H7 3-4 Meloxicam base (3-1 and 4-2. Meloxicam is available as a tablet for oral administration containing 7.5 mg or 15 mg meloxicam.

The inactive ingredients in Moloxican tablets USP include colloidal silicon dioxide, crospovidone lactose monohydrato, magnesium stearate, microcrystalline cellulose, povidone and sodium cirtate dhydrate.

12 CLINICAL PHARMACOLOGY

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a Mortsmann and Mortsmann a

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Melasican excretion is productionly in the form of metadolaw, and occurs to equit examine in the arise and frees. Only passes of the anchored passes compound are excreted inducing (1875), and exception of the second s

Podiatric

After single (0.25 mplg) does administration and thre schering study case (0.35 mplg) days, there was a general torel of approximately 30% lever exposure is symapping pattern (21 to 5 years old 0.3 monoport of the short pattern bala molecular improvement influence of the symaphical schering study and the schering study of the s

Geriatric

Etherly males (+65 years of age) exhibited melosican plasma concentrations and stoady-state plasmacoilastics: similar to yong males. Etherly formics (+65 years of age) had a 2% higher ALCase memilization. Despite the increased trut concentration in the development, makes a store sever profile was comparable for both etherly patient populations. A smaller free fraction was found in etherly fremark galaxies in comparisons to leftry males galaxies.

Young females exhibited slightly lover plasma concentrations relative to young males. After single doses of 7.5 mg Monitorian, the man elimination half-life was 19.5 hours for the female group as compared to 23.4 hours for the male group. At standy stane, the data were similar (17.5 hours vs 21.4 hours are similar (17.5 hours vs 21.4 hours of the male group. At standy stane, the data were similar (17.5 hours vs 21.4 hours of the similar (19.5 hours vs 21.4 hours of the male group. At standy stane, the data were similar (17.5 hours vs 21.4 hours of the similar (19.5 hours vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar (19.5 hours of the similar vs 21.4 hours of the similar vs 21.4

hours). This pharmacokinetic difference due to gender is likely to be of little clinical importance. There was linearity of pharmacokinetics and no appreciable difference in the Cmax or Tmax across genders.

genders. Folgosia Inpairment Following a single 15 mg dose of molosicam there was no marked difference in plasma (concentrations inpairments with and Clade Hugh Class 1) or molerate (Clade Pugh Class 1) playade (inpairment) Molosoge adjustment is accessar justices with mith to molecule a single class 1) playade (inpairment) Norskey adjustment is accessar justices with mith to molecule subject inpairment. Forfane with survey hugher (inpairment) (Clade Hugh Class 1) Provanders (Clade Hugh Class 1) players with mith to molecule subject inpairment. Forfane with survey hugher (inpairment) (Clade Hugh Class 1) Provanders (Clade Hugh Class 1) Norskey and the molecule subject inpairment. Forfane with survey hugher (inpairment) (Clade Hugh Class 1) Norskey and the molecule subject inpairment. For a subject in the molecule subject in the molecule subject inpairment. For a subject in the molecule subject in the molecule subject inpairment. For a subject in the molecule subject in the molecule subject in the molecule subject inpairment. For a subject is normalised in the molecule subject in the molecule subject inpairment. For a subject is subject in the molecule subject in the molecule subject inpairment. For a subject is subject in the molecule subject inpairment inpairment in the molecule subject inpairment. For a subject is subject in the molecule subject in the molecule subject inpairment. For a subject inpairment is a subject in the molecule subject in the

Proceeding (2) and Use it specify Population (BD): Also Important Development Development

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Data Baracka Statisti Data Baracka Statisti Angene Webs SSADA: were addictioned with aspirit, the protein binding of NSADs were reduced, abrough the cleanare of free NSAD was and abreak. Webs Monitorianin administered with aspirits (100 mg how it may daily binding) without were administered with a spirit administered with a spirit administered and a spirit administered with a spirite law Daily binding without were administered with a discontrast of the spirite administered with a spirite law Daily binding without the consequence of the spirite administered with a spiritered with a spirite additional spiritered with a spiritered

store presente obtainers of a loss of an endotrem. Disposite Melocitican IS on gover a daily for 7 days did not alter the plasma concentration profile of digoximalter B-acetyldigoximadministration for 7 days at clinical doses. Is vitro testing found no protein binding drug interaction between digoxim and melocitican. umming drog uteraction between digoxin and melosican. Liftium: In a study confacted in healthy subjects, mana pro-dose liftium concentration and AUC were increased by 21% is subjects receiving liftium doses ranging from 804 to 1072 parked ally with melosican 15 mg QD every day as compared to subjects receiving liftium alone [see Drug Interactions (7).

Moneyour: A study in 12 thermanial arbitist (RA) pairies evaluated the effects of multiple does of multication of planneciliantics of productions and arbitistic and arbitrary a digitation effect on all manuscinations: of align does on studients and hypothese and the arbitrary of the arbitrary of the arbitrary of the arbitrary of hypothese the effect of align does of arbitrary and arbitrary of hypothese the arbitrary of the arbitrary of the arbitrary of hypothese the arbitrary of the arbitrary of the arbitrary of hypothese the arbitrary of the arbitrary of hypothese the arbitrary of hypothese the arbitrary of the arbitrary of hypothese the hypothese hyp

13 NONCLINICAL TOXICOLOGY

IN NORCLINICAL FORCEOLOGY IN LCarcingeness's Managenesis, Implement of Fertility <u>Carcinemusia</u> There was no increase in names incidence in long-averacarizangenicity studies in na (104 weeks) and implicitly in nise (104 to 5.3md 2.5.4ms, respectively, de maximum recommende Human dose [MBRIII] of 15 mg/Mg/Malanzam March domarky studies are MSA.

[MRtt1]/of 15 mg/say Mercana and a second se

moairment of Ferti Meloxicam did not impair male and female fertility in rats at oral doses up to 9 mg/kgiday in males and 5 mg/kgiday in females (up to 5.8- and 3.2-dimes greater, respectively, than the MRHD based on BSA commarison).

14 CLINICAL STUDIES

14.1 Osteoarthritis and Rheumatoid Arthritis Int. Uncertainties and Automation Artificial suggest and synapses of consecutivity of the laws and high too see the final sector of the laws and high too set of the laws and high too set of the laws and high too sectors and the laws and high t

was previous two increments of architectus (RA) Paraclaritotic and an Mybarchard Course 12-20 new infle Neurannois M Architectus (RA) Paraclaritotic and Mybarchard Course The use of Meloxicam for the twomwer of the signs and symptoms of paraclaritotilar or polyarticular course J lowerile Rhumanial Architectus (RA) Paraclaritotic and Section 2019 and Section 2

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NDC 29300-124-30; Bomes of 5,000 Meloxican Tablets USP 15 mg are available as follows: NDC 29300-125-13; Bomles of 30 NDC 29300-125-01; Bomles of 100

NDC 29300-125-10; Bonies of 1,000 NDC 29300-125-50; Bonies of 5,000

Storage Store at 20⁶ to 25⁶C (68⁶ to 37⁶F) [see USP Controlled Room Temperature]. Keep Meloxican Tables: USP in a dry place

Dispense tablets in a tight comainer. Keep this and all medications out of the reach of children.

17 PATIENT COUNSELING INFORMATION Advise the patient to read the FDA-approved patient labeling (Medication Guide) that accompanies each prescription dispersed.

prescription dispensed. Additional Medication Guides can be obtained by calling Unichem at 1-866-562-4616.

Informpatients, families or their caregivers of the following information before initiating therapy with an NSAID and periodically during the course of ongoing therapy.

Endirected Threather's twen Arbite patients to address or during of speech, and a report any of these symptoms on their consects of broch, works or, subtracting of speech, and as report any of these symptoms in their discontent of the symptoms of the symptoms of the symptoms of the symptoms framework of the symptoms of the symptoms of the symptoms of the symptoms and the symptoms of the symptoms of the symptoms of the symptoms of the symptoms and the symptoms of the symptoms of the symptoms of the symptoms of the symptoms and the symptoms of the symptoms of the symptoms of the symptoms of the symptoms the symptoms of the s

Heranonicity: Informpatients of the variing tigm and symptoms of hepanitoxicity (e.g., nussea, Infigue, Induzy, darbas, purities, juantice, right upper quadrant underress, and "The-Jaie" symptoms). If these occur, instruct patients to stop Meloxic and tablets and seek immediate medical therapy [see Warnings and Proceedings].

HeartFathers and Edven Advise patients to be also: for the symptoms of congestive heart failure including shortness of breach, userpland weight and Printadion (53). Manchische Roestann, 653). Manchische Roestann

Amplituation for actions Inform patients of the signs of an anaphylactic reaction (e.g., difficulty breathing, swelling of the face or throad). Instruct patients to seek immediate emergency help if these occur [see Contraindications (4) and Warnings and Prevenzions (27.7)].

Serious Skin Reactions

Advise patients to stop Meloxican tablets immediately if they develop any type of rash and to contact their healthcare provider as soon as possible [see Warnings and Precusions (5.9)].

<u>Femile Femility</u> Advise females of reproductive potential who desire pregnancy that NSAIDs, including Meloxicam tublets, may be associated with a reversible delay in ovulation [see Use in Specific Populations (8.3)]. Fetal Toxicity

True Trooting Inform pregnant women to avoid use of Meloxican tablets and other NSAIDs starting at 30 weeks gestation because of the risk of the premater closing of the feal dactus arteriosus [see Warnings and Prevancions (S. 10) and Use in Specific Populations (8.1)].

Avoid Concomitant Lise of NSAIDs

Action Loncontinue Lise of NSAID3 Informations in the concentrate use of Melosican tablets with other NSAID5 or salicytants (e.g., diffusional, salication) is one recommended due to the increased risk of gaussimential sub-ting, and like diffusional salication is one recommended due to the increased risk of gaussimential sub-ting, and like that NSAID5 may be present in "over the counter" medications for treatment of colds, fever, or insomain.

Use of NSAIDs and Low-Dose Aspirin

omitantly with Meloxican tablets until they talk to their Inform patients not to use low-dose aspirin concomitantly with Meloxica healthcare provider [see Drug Interactions (7)]. For current prescribing information, call Unichem at 1-866-562-4616.

Manufactured by: UNICHEM LABORATORIES LTD. Pilerne Ind. Estate,m un. ristate, Pilerne, Bardez, Goa 403511, India Manufactured for:

Unichem Logo

Hashrouck Heights, NJ 07604 07-R-09/2017 13009858 SPL MEDGUIDE

Medication Guide for Nonsteroidal Anti-inflammatory Drugs (NSAIDs)
What is the most important information I should know about medicines called Nonsteroidal Ant inflammatory Drugs (NSAIDs)?
NSAIDs can cause serious side effects, including:
Increased risk of a heart attack or stroke that can lead to death . This risk may happen early in
reament and may increase: with increasing doses of NSAIDs
 with longer use of NSAIDs Do not take NSAIDs right before or after a heart surgery called a "coronary artery bypass
raft (CABG)."
Avoid taking NSAIDs after a recent heart attack, unless your healthcare provider tells you to. You may have an increased risk of another heart attack if you take NSAIDs after a recent heart
anack.
 Increased risk of bleeding, ulcers, and tears (perforation) of the esophagus (tube leading from the mouth to the stomach), stomach and intestines:
the mouth to the stomach), stomach and intestines:
without warning symptoms
that may cause death The risk of getting an ulcer or bleeding increases with:
 parts or getting an uncer or one-cong increases while part history of stomach ulcers, or stomach or intestinal bleeding with use of NSAIDs taking medicines called "corticosteroids", "articoagelants", "SSRIs", or "SNRIs"
 taking medicines called "corticosteroids", "anticoagulants", "SSRIs", or "SNRIs" increasing doses of NSAIDs
s smoking s drinking alcohol
a older age
a dvanced liver disease b bleeding problems
a exactly as prescribed a at the lowest dose possible for your treatment
a for the shortest time needed
What are NSAIDs?
NSAIDs are used to treat pain and redness, swelling, and heat (inflammation) from medical conditions such as different types of arthritis, menstrual cramps, and other types of short-term pain.
Who should not take NSAIDs?
Do not take NSAIDs: • if you have had an asthma attack, hives, or other allergic reaction with aspirin or any other NSAIDs.
right before or after heart bypass surgery.
Before taking NSAIDs, tell your healthcare provider about all of your medical conditions, including if you:
 have liver or kidney problems
 have high blood pressure
 have asthma are premant or plan to become pregnant. Talk to your healthcare provider if you are considering
are prognant or plan to become prognant. Talk to your healthcare provider if you are considering aking NSAIDs during prognancy. You should not take NSAIDs after 29 weeks of prognancy. are breastfeeding or plan to breast feed.
 are breastfeeding or plan to breast feed. Tall your boother was provider about all of the medicines you take, including prescription or over
are constructing point of the stretch. If your health care provider about all of the medicines you take, including prescription or over the -counter medicines, vitamins or herbal supplements. NSADS and some other medicines can increat with each other and cances sorious side effects. Do not start taking any new medicines without the effects of the start of the start start and the start and the start and the start start start and the start start and the start start and the start start start and the start start start and the start sta
interact with each other and cause serious side effects. Do not start taking any new medicine without
talking to your healthcare provider first. What are the possible side effects of NSAIDs?
NSAIDs can cause serious side effects, including:
See "What is the most important information I should know about medicines called Nonsteroid Anti-inflammatory Drugs (NSAIDs)?"
heart failure liver problems including liver failure
low red blood cells (anemia) life-threatening skin reactions
 Inte-Intreatening sam reactions Ilfe-threatening allergic reactions Other side effects of NSAIDs include: stomach pain, constipation, diarrhea, gas, heartburn, nausea,
Other side effects of NSAIDs include: stomach pain, constipation, diarrhea, gas, hearthurn, nausea,
romiting, and dizziness. Get emergency helo right away if you get any of the following symptoms:
Get emergency help right away if you get any of the following symptoms: • shormess of breath or trouble breathing
chest pain weakness in one part or side of your body
 swelling of the face or throat
Stop taking your NSAID and call your healthcare provider right away if you get any of the
loBoving symptoms : • Nausea • more fired or weaker than usual
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Limited Manufactured by: UNICHEM LABORATORIES LTD. Filerene Ind. Essaw, Filerene, Bardez, Goa 403511, India Manufactured for:

PHARMACEUTICALS (USA). INC.

Hasbrouck Heights, NJ 07604 06-R-09/2017 13009858

This Medication Guide has been approved by the U.S. Food and Drug Administration. Revised: September 2017

Revised Segurates 2017
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Cookeville, TN 38506 201811153H

PRINCIPAL DISPLAY PANEL - 7.5 mg NDC 71610-189 - Meloxicam USP 7.5 mg - Rx Only



PRINCIPAL DISPLAY PANEL - 15 mg NDC 71610-190 - Meloxicam, USP 15 mg - Rx Only

CCC FREEDOWN Medication, USP Torg. 30 Tables Medication (USP Street) Medication (USP Street)



Product Informat	tion						
Product Type		HUMAN PRI	SCRIPTION DRUG	Item Code (Sou	rce) NDC/1	6 10 - 19 0 (NI	C 29300-125
Reate of Administra	sion	ORAL					
Active Incredient							
Active ingredien	OLCENE	Incredient Na			Basis of S		Strength
MELOXICAM (UNE V	constance				MELOXICAM	araga	15 mg
Inactive Ingredie	nts						
			dient Name				Strength
CELLULOSE, MICRO			12136 1U)				
CROSPOVIDONE (UN LACTOSE MONORITI							
	TE (INE	10.05756(00)					
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Establishment

 Nom
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 Business Operations

 Adjurts Planta Schlimt - Francese, LLC
 (D213145
 (B27522/V111-815, Y101-810)

 Broked 11/2018
 Appena Planta Schlimtor - francese, LLC