MELOXICAM- meloxicam tablet Asciemed USA, Inc.

HIGHLIGHTS OF PRESCRIBING INFORMATION These highlights do not include all the information needed to use MELOXICAM TABLETS USP, safely and effectively. See full prescribing information for MELOXICAM TABLETS USP.

ELOXICAM Tablets USP, for oral use

Initial U.S. Approval: 2000

See full prescribing information for complete boxed warning. Nonstancial anti-influencemberry drugs (NSADD) course an increased risk of serious cardiovascular theorehotic events, including myscardial infarction and strake, which can be fatal. This risk may occur any in it treatment and may increase with duration microsoftware services and serv

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(2.1) GA (2.2) and RA (2.3): Starting dose: 7.5 mg once daily Dose may be increased to 15 mg once daily (PA (2.4):

\$A1 (24):
 \$A1 (24):
 \$A2 (24):

anemia (511,7)
 ADVERSE REACTIONS
 Most common (a5% and greater than placebo) advense events in addets are damhas, upper
 respinatory tract hetcione, dyspeptia, and influenza-like symptoms (61)
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5.2 Gastroineastual Bleeding, Ulcerulion, and Perforation 5.3 Hapatohistorik 5.3 Haut Tahura and Edema 5.3 Haut Tahura and Hoperkalamin 5.8 Baut Tahura and Hoperkalamin 5.10 Permit Annual Charlon and Hoper 5.10 Permit Annual Charlon and Hoper 5.11 January Moderning 5.12 Hauting of Internation and Fore 5.12 Hauting of Internation and Fore 5.13 Linearity Moderning 6. Chineck Time Experience

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FULL PRESCRIBING INFORMATION
WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL
EVENTS ______ EVENTS Cardiovascular: Thrombotk: Events • Rootstradial anti-Inflammadry drugs (IKSADD) cause an increased influction and travos, which can be fatal. This risk may occur early in treatment and may increase with duration of use (see Warning and Protections) (5.1). Constraindicate the satting of correary antery bypass graft (CABD) surgery (see Contraindications (4) and Warnings and Proceedings (5.1).

1 INDICATIONS AND USAGE

1.1 Osteoarthritis (OA) Meloxicam tablets are indicated for relief of the signs and symptoms of osteoarthritis [see Clinical Studies (14.1)].

see Carical Subate (24-27); 1.2 Rheumatoid Arthrikis (RA) Meloxicam tablets are indicated for rolled of the signs and symptoms of rheumatoid arthriki (see Chical Studies (24-27); artitrits (see Circuid Studies (14.1) [1.3] Jovennik Rheumatold Arthritis (JRA) Pauciarticular and Polyarticular Course Molociam tabatis are indicated for relief of the signs and symptoms of pauciarticular or polyarticular course jovenin Breumatoli Arthritis in patients who weigh e60 kg [see Dosage and Ambinstaton (2.4) and Clinical Studies (14.2).

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suit an individual patient's needs. In adults, the maximum recommended daily oral dose of Meloxicam tablets is 15 mg regardless of formulation. In patients with hemodialysis, a maximum daily dosage of 7.5 mg is necommended [see Use in Specific Populations (8.7) and Clinical Pharmacology (regardless or round-mig is recommended [see Use in Specific Populations , 12.3)]. Meloxicam tablets may be taken without regard to timing of meals.

For the relief of the signs and symptoms of osteoarthritis the recommended starting and maintenance erail dose of Meloxicam tablets is 7.5 mg once daily. Some patients may receive additional benefit by increasing the dose to 15 mg once daily.

Inverse even additional bandh by increasing that does to 15 mg once daily. 2.3 Housenedia dividentità for the relia of the signs and symptomic of theumatail antrhits, the recommendent tartitistica and methanican additional dividentità di 3.5 mg once daily. Some constraint and the signs and signature dividentità di 1.5 mg 2.4 journels Rhommataid Archittis (UA) Parcufarchicale and Polynchichat Con-text the transmitta di sectional di territori di sectional di se di Mascana tables in 5.5 mg once daily in childron who weight additis. The mession Mascana tables in 5.5 mg once daily in childron who weight additis, The mession Mascana tables in 5.5 mg once daily in childron who weight additis. The childra that has a section about dimensional dividentity of the section of the sign. There was no dimensional and the section of the section of the sign of the section of the section of the sign of the section of the sign of the section of the sign of the section of the se

additional benefit demonstrated by increasing the doke above 7.5 mg in clini Meloxicam tablets should not be used in children who weigh <60 kg. 2.5 Renal Impairment The use of Meloxicam tablets in subjects with severe renal impairment is not recommended.

In patients on hemotialysis, the maximum dosage of Meloxicam tablets is 7.5 mg per day [see Clinkal Pharmacology (12.3)].

varg i are cinital material (21.2) § 2.6 Non-Interchangeability with Other Formulations of Melosicam bioloxicam tables have not tabion equivalent systemic exposure to other approved formations of oral melosicam. Therefore, Melocicam tables are not interchangeable with other formations of oral melosicam product work The bioloxicam tables, with the tables the same. Do not catchetic entropy of the same strengths of Melosicam tables, with other formations of one interchange product.

3 DOSAGE FORMS AND STRENGTHS

Meloxicam Tablets USP: • 7.5 mg: Light yellow, round flat beveled edged, tablet with U & L debossed on one side and 7.5 debossed centrally on the other side

4 CONTRAINDICATIONS

4 CONTRANDICATIONS Web/cam tables are contraindicated in the following patients: • Known hypersensitivity (e.g., anaphytictic reactions and serious skin reactions) to malocican or any components of the drug product (2 see Warnings and Precautions (5.7, 5.0) • History of authma, unicaria, or other allergic-type reactions after taking asplrin or

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Populations (6.6) and Chield Pharmacology (12.3) [3.4 Appentations) NADAS, Including Mediculary, can load to new oncore or versening of prevakting hyperturison, abit or with many contributed the increased includence of CV events. Potents Barling angularismic converting surgrays (AES) Inhibitors, Barlado duretics, or poly Interactions (7.7), contrigorism the Microbiol State Monitor biologo pressure (BB) during the Initiation of MSAD treatment and throughout the course of thranky).

Contrast owneys: 5.5 Heart Fallware and Edema The Carb and traditional HSAID Trillists: Collaboration meta analysis of randomized controlled this demonstrated and approximately teoriodin forcusa in hospitalizations for compared to glocek-notated patients. Is a Datient National Registry study of patients with Natra Talare, NSAID use increased the risk of M₁ hospitalization for heart failure, and datah.

and dath. Additionaly, fluit retention and edema have been observed in some patients treated with ISADIG. Use of melanizam may bank the CV effects of several therapeutic appents used to treat these medical controls (e.g., during controls, and endowed to blockers (DABs)]) see Drug Interactions (7.1). Accord the use of Matchian in patients with severe heart failure unless the benefits are expected to outweigh the risk of enersieng heart failure. If Matocianis is used to patients with severe heart failure, monitor patients for edges of enersieng heart failure.

5.6 Renal Toxicity and Hyperkalemia <u>Renal Toxicky</u> Long-term administration of NSAIDs, including Meloxicam, has resulted in renal papilary necrosis, renal insufficiency, acute renal failure, and other renal injury.

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Correct volume status in dehydrated or hypovolemic patients prior to initiating Neloxicam. Monitor renal function in patients with renal or hepatic impairment, heart ailure, dehydration, or hypovolemia during use of Meloxicam [see Drug Interactions (7)

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5.9 Serios Stan Reactions SUBJb, locking moticam, can cause serious skin adverse reactions such as exclusion dermätik, Stewars-jonnen Syndrome (S)S, aut toxic späarmal necrolysis (TR), which can be tall: These serious cances through varing, inform patient about this signs and symptome of various skin reactions, and to discontinue the used Metalicatian is for its appearance of skin rank array other used Metalicatian is for stage paranetor about rank or any other used metalicatian is shown and the stage paranetor about the stage reactions to MSAIDE [see Contraindications (4)].

reaction to Robibli (an Carbathandonia (d.). 3.03 Portantico Carbathandonia (d.). Biolican may cause premative closure of the field addrag and the second of an end of the second second of the second of the second of the second of the second second of the second of the second of the second of the second second of the second of the second of the second of the second baseline the second of the second of the second of the second baseline the second of the second of the second of the second baseline the second of the second of the second of the second baseline the second of the second of the second of the second baseline the second of the second of the second of the second baseline the second of the second of the second of the second baseline the second of the second second of the second of the second of the second of the second second of the second of the second of the second of the second second of the second of the second of the second of the second second of the second of the second of the second of the second second of the second of the second of the second of the second second of the sec

himogolen or hematocrtz. NSAIDs, including Metoxicam, may increase the risk of biaeding events. Co-morbid conditions such as caaguidan disorders or concomitant use of warfarin, other anticoaguiders, antipitatiet agents (ca., apprin), servorism regutate inhibitors (SSMs) and serdorin nonpinghrin erugataia inhibitors (SMtB) may increase this risk. Nonfor these patients for signs of baseding is adr long inhibitors (7).

5.12 Masking of Inflammation and Fever 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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CBC and a chemistry profile prioritaging (in we therapy and Procession (5.5, 3.5, 3.6) **CAROPER LACTOR** The following partner reactions are discussed in grater data in other sections of the langes, and the section of the section of the section of the langes, and the section of the section of the section of the langes, and the section of the section of the section of the langes, and the section of the section of the section of the langes, and the section of the section of the section of the langes, and the section of the section of the section of the langes, and the section of the section of the section of the langes of the section of the section of the section of the langes of the section of the section of the section of the langes of the section of the section of the section of the langes of the section of the section of the section of the langes of the section of the section of the section of the langes of the section of the section of the section of the langes of the section of the section of the section of the langes of the section of the section of the section of the langes of the section of the section of the section of the langes of the section of the section of the section of the langes of the section of the section of the section of the langes of the section of the section of the section of the section of the langes of the section of the section of the section of the section of the langes of the section of the section of the section of the section of the langes of the section of the section of the section of the section of the langes of the section of the section of the section of the section of the langes of the section of the langes of the section of the section of the section of the section of the langes of the section of

• Installing "tracking" loss Warning and Procedures (5.11) [
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truis. A 12-week multicenter, double-blind, randomized trial was conducted in patients with osteoarthritis of the inse or hip to compare the efficacy and safety of Meloxicam with placebo and with an active control in the 12-week multicenter, double-blind, randomized trials were conducted in patients with rheumateid arthritis to compare the efficacy and safety of Meloxicam with placebo. that were consistent of plaints with momento arona to compare the entacy staffor of Makrison with placksto. Table 1a dejects adverse events that occurred in z2% of the Meloxican treatment groups in a 12-week plackbo- and active-controlled osteoarthrifts trial. Table 1b dejects adverse events that occurred in z2% of the Meloxican treatment groups in the 12-week plackbo-controlled arthritis trial.

Table 1a Adverse Events (%) Occurring in ±2% of Meloxicam Patients in a 12-Week Osteoarthritis Placebo- and Active-Controlled Trial Placebo 7.5 mg daily Meloxicam JS Dictofenac 7.5 mg daily g daily daily

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In patients 8 USE IN SPECIFIC POPULATIONS 8.1 Pregnancy Risk Summary

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cancar migrace.	Main/com ad anticoagularity such as warfurh have a springistic effect on blading. The concentrative use of meta/com and anticoagularity such as warfurh have a springistic effect on blading. The concentrative use of meta/com and anticoagularity have a springistic effect on blading. The concentrative use of anticoagularity have a springistic effect on blading compared to a such as a defined and approximate and anticoagularity have a springistic effect on blading. The concentrative use of anticoagularity have a springistic effect on blading more than an HSAID alone.
	Manhor patients with concomitant use of Makoulcam with anticoagulants (e.g., marfarin), antiplatikel agents (e.g., marfarin), antiplatikel
Aspirin	
Clinical Impact:	Controlled clinical studies showed that the concombant use of NSAIDs and analassic doses of aspirin does not produce any oreater therapeutic effect than the use of NSAIDs alone. In a clinical study, the concombant use of an NSAID and aspirin was associated with a significantly increased incidence of Gi adverse reactions as compared to use of the NSAID alone [see Warnings and Precautions [5,2]].
intervention:	Doromitant use of Meloscam and low dose appin or analysis doses of appin's in ot generally recommended because of the increased risk of bielding [see Warnings and Precautions (511]]. Meloscam is not a substitute for low dose appin for cardiovascular protection.
ACE Inhibitor	. Anajotensin Receptor Blockers
	NEX/Ds may dminish the antiwaritensive effect of anoiotensin converting enzyme (ACE) inhibitors, anoiotensin receptor blockers (including angranado).
Clinical Impact:	In statistics who are elderly, volume-depleted lincluding those on divertic therapy), or have renal insamment, coadministration of an ISAID with ACE inhibitors or ARBs may result in deterioration of renal fautre. These effects are usually reversible.
Intervention:	During concentration of Malocana and ACE Inhibitors, BMA in a second and ACE Inhibitors and ACE Inhibitors and ACE Inhibitors of Halo concentration (second and ACE Inhibitors, BMA in a second and ACE Inhibitors and ACE Inh
Diuretics	
Clinical Impact:	Cinical studies, as well as post-
canca impact:	marketing observations, showed that NSAIDs reduced the matriaretic effect of loop diaretics (e.g., forosemide agents and melosicam have not demonstrated a reduction in natriaretic effect. Furosemide single and multiple doses pharmacodynamics and pharmacokinetics are not affected by multiple doses of melosicam.
	During concombart use of Melosicam with dureks, observe patients for signs of wessening nema function, in addition to assuring dureks (efficacy including anthypertonsive effects [see Warnings and Precautions (5.6)].
Lithium	
	NSAIDs have produced elevations in plasma lithum levels and reductions in renai lithum clearance. The mean minimum lithum concentration increased 15%, and the renal clearance decreased by approximately 20%. This effect has been attributed to NSAID inhibition of renal prostaglandin synthesis (see Clenical Pharmacology (12-3)).
	During concombant use of Mikexican and Ithium, montor patients for signs of Ithium texcity.
Methotrexate	
Clinical Impact:	Economitant use of NSAIDs and methotrexite may increase the risk for methotrexite texicity (e.g., neutropenia, threnal dystunction).
intervention:	During concomitant use of Makoxican and methodrowska backty.
Cyclosporine	
Clinical Impact:	Enrommant use of Meloxicam and cyclosporine may increase cyclosporine's nephrotoxicity.
intervention:	During concomitant use of Melaxicam and cyclosporine, monitor patients for signs of worsening renal function.
NSAIDs and S	alcvistes
Clinical Impact:	Concomitant use of meloxicam with other NSAIDs or salicylates (e.g., diflurisal, salsalate) increases in effects (see Warnings and Procautions (5.2)].
Intervention:	The concombant use of metavican with other HSAIDs or sale/attas is not recommanded.
Pemetrexed	
	Economitant use of Meloxicam and permetrexed may increase the risk of permetrexed-associated myelosuppression, renal, and Gl toxicity (see the permetrexed prescribing information).
	During concomitant use of Melosicam and permitterend, in patients with remainingairment whose creativine clearance ranges from 45 to 79 mL/min, monitor for myelosuppression, renal and GI toxicity.
Intervention:	Patients taking meloxicam should interrupt dosing for at kaast five days before, the days of Johnson administration.

Table 3 Clinically Significant Drug Interactions with Meloxi

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

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Abdominal pain	2.7	2.3	4.7	2.9
Constipation	0.8	1.2	1.8	2.6
Diamhea	1.9	2.7	5.9	2.6
Dyspepsia	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 System				
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
Musculoskeletal				
Arthralgia	0.5	0.0	5.3	1.3
Back pain	0.5	0.4	3.0	0.7
Psychiatric				
linsomnia	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
Urinary				
Micturition frequency	0.1	0.4	2.4	1.3
Urinary tract infection	0.3	0.4	4.7	6.9
* WHO preferred terms edema, edema depender	it, edema peripheral, and eder	na legs combined		

The adverse events that occurred with Meloxicam in a2% of patients treated short-te (4 to 6 weeks) and long-term (6 months) in active-controlled osteoarthritis trials are presented in Table 2. Table 2 Adverse Events (%) Occurring in ≥2% of Meloxicam Patients in 4 to 6 Weeks and 6 Month Active-Controlled Ost Trials

No. of Patients Gastrointestinal

Gastrointestinal Disorders	14.1	18.9	16.8
Abdominal pain NOS *	0.6	2.9	2.3
Dyspeptic signs and symptoms ¹	3.8	5.8	4.0
Nausea	2.6	3.3	3.8
General Disorders and Administration Site Co	inditions		
Influenza-like ilness *	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 Disor	ders		
oint related signs and symptoms ¹	1.9	1.5	2.3
Nervous System Disorders			
Headaches NOS	6.4	6.4	5.5
Skin and Subcutaneous Tissue Disorders			
Rash NOS	1.7	1.0	2.1
 MedDRA preferred term: nausea, abdominal pain NOS, it 	nfuenza-like ilness, head	aches NOS, and rash NO:	5
 MedDRA high level term (preferred terms): dyspeptic sig gastrointestinal initiation), upper respiratory tract infectio related signs and symptoms (arthraíois, arthraíois agons) 	rs-oathopen unspecified	Gannotis NOS, pharmol	ta NOS, sinusitis NOS), joint

2.5 2.6 0.6 2.0 Skin Rash ² nh2 2,5 2,6 un Table 2,5 2,6 Un Table 2,5 2,6 Un Table 19 Adverse Events (%) Occurring in 2,5% of Medison Hannis in two 12-Week Rheumatold Activities Placebo- Centroled Tuble 3,5 and aday Malaxican 15 ang aday Placebo Mediscan 7,5 ang aday Malaxican 15 ang aday 469 441 477 443 477 No. of Bations

No. of Patients	157	154	156	153
Gastrointestinal	17.2	20.1	17.3	28.1
Abdominal pain	2.5	1.9	2.6	1.3
Diarrhea	3.8	7.8	3.2	9.2
Dyspepsia	4.5	4.5	4.5	6.5
Flatulence	4.5	3.2	3.2	3.9
Nausea	3.2	3.9	3.8	7.2
Body as a Whole				
Accident household	1.9	4.5	3.2	2.6
Edema ¹	2.5	1.9	4.5	3.3
Fail	0.6	2.6	0.0	1.3
Influenza-like symptoms	5.1	4.5	5.8	2.6
Central a n d Peripheral Nervous System				
Dizziness	3.2	2.6	3.8	2.0
Headache	10.2	7.8	8.3	5.9
Respiratory				
Pharyngitis	1.3	0.6	3.2	1.3
Upper respiratory tract infection	1.9	3.2	1.9	3.3

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undergring investigation of infertility. **LA valancie:** Use The solarity of efficiency of a set of the solar set of the

Wrontype and Proceeding (\$1, 5, 5, 5, 5, 5, 5, 5, 1). **24. black transmission 24. black transmission 25. black transmission 26. black transmission 26. black transmission 27. black tr**

11 DESCRIPTION Nakorkam Tablets USP are a nonstarodial and inflummatory drug (NSAID). Each tablet constants / S my mitting. Are for construct a phase and a construct any decigning mitting. The motional area of the second and a construct any decigning 1.1. decisits. The motional area of the second area of the second area of the second area of the faith the following interturbal formula is C 14H 32H 30.6.5 and the number of interturbal formula is C 14H 32H 30.6.5 and



Chemical Structure Chemical Structure

Cancel Structure Mokicam is a pastel yoko dilu practica pricabili watar, with higher solubility observed in thorong acids and based. It is very tightly solubili in matured. Mokicamento high a value of 11 and 42. Mokicam is available as a table for oral administration containing 15 mg mokicame. In actical engineering the historical tables (10 becluica cidade acids disolito) crospositiona, latcisse monohyrizar, magnissian tasarate, microcrystalline calutose, provisiona and acidem table displays.

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Pharmacokinetic Parame	(4.4.4)	7.5 mg ¹ tablets	15 mg capsules	15 mg capsules	15 mg capsules	epatic insufficiency (Fas 15 mg capsules
		18	5	15 mg cupsules	12	12
max	(µg/mL)	1.05 (20)	2.3 (59)	3.2 (24)	0.59 (36)	0.84 (29)
max	[h]	4.9 (8)	5 (12)	6 (27)	4 (65)	10 (87)
1/2	(h)	20.1 (29)	21 (34)	24 (34)	18 (46)	16 (29)
Ú.	(mL/min)	8.8 (29)	9.9 (76)	5.1 (22)	19 (43)	11 (44)
2H3	[L]	14.7 (32)	15 (42)	10 (30)	26 (44)	14 (29)
The parameter values in the t not under high fat conditions	table are from various studies					

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been invergated. Geniroit: Eleviry music (e55 years of age) exhibited methiciane plasma concentrations and study-state plasmaceativetics similar to young make. Elserly formatic (e65 years of formatic (e55 years) of age) after to young make. Elserly formatic (e65 years) of concentrations in the source (for the source end of the source end to the elserly plasmaceativetics). A result with for function was found in elserly formate postance in the source plasma plasma.

Sex Set (nonsec exhibited is light) here decrease overstotene relative to young mixes, there single docs or 25 mg Meliocate. The mean elementary Meliot Mel well by 15 hours for the formal group as compared to 33.4 hours for the main group, At statusty state, the data were similer (17) hours V 31.4 hours for the main group, At statusty state, the grades is likely to be of BBC efficiency. They are used insafely of planmace/melions. Meant for the mean state of the state of the state of the planmace/melions. Meant for administration of the state of the state of the state of the planmace/melions.

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was aspiring see Drug Interactions (7) 1. Cholostyramine: Protreatment for four days with cholisstyramine significantly increased the charance of methodizam by 50%. This resulted is a discrease in try, from 19.2 hours; to 12.5 hours, and a 35% reduction in AUC. This suggests the existence of a rescruction pathway for methodizam in the garcronizational tract. The circuit arekennes of

this interaction has not been established. Cimetéline: Concomitant administration of 200 mg cimetéline four times daily did not aiter the single-dose pharmacokinetics of 30 mg meloxicam.

Digoxin: Maloxicam 15 mg once daily for 7 days did not alter the plasma concentration profile of digoxin after F-acetyldigoxin administration for 7 days at clinical doses. In etro testing found no protein binding drug interaction between digoxin and meloxicam.

metry there are another behavior below the second s

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

3.1.1 carcinogeness, protogeness, impairment of variety Carcinogenesis There was no horness in humor inclusion in long-term carcinogenicity studies in rats: indigating in rats and up to 8.0 migration material part of 0.5 - migrates. The respectively, the maximum recommended human does (MRHD) of 15 mg/day Matexian based on holy variate area (BSA) comparison).

Meloxicam was not mutagenic in an Ames assay, or clastogenic in a chromosome aberration assay with human lymphocytes and an in vivo micronucleus test in mouse bone marrow.

Impairment of Fertility Meloxicam did not impair male and female fertility in rats at oral doses up to 9 mg/kg/day in male: and 5 mg/kg/day in females (up to 5.8- and 3.2-times greater, respectively, than the MRHD based on BSA comparison).

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Le HOW SUPPLEIS-COMME AND HANCLING Management and the second sec

Storage Store at 20 ° to 25 °C (68 ° to 77 °F) [see USP Cont Temperature]. Keep Meloxicam Tablets USP in a dry place alled Boom Dispense tablets in a tight container. Keep this and all medications out of the reach of children.

17 PATIENT COUNSELING INFORMATION Arkies the patients for and the F/D-Approved patient takeling (Matcastin Galoi) that Academaness and procession departial. Additional Matcastin Galois can be obtained by calling Unkhem at 1.866-Homm patients, financia on that complexing on the following information before initiating therapy with an KSMD and periodically during the course of organity therapy. Certostance after Theoretic Fuence

theory into HSGD and particiting when the course of energy through.
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Anderbarder balantions inform patients of the signs of an anaphysicitic reaction (e.g., difficulty breathing, swelling of the face or throad), instruct patients to seek immediate amergency hap if these secces (see Controland, allow (4) and Warnings and Procession (5, 7).]. <u>Satisfue Sahn Instructors</u> Advise patients to stop Meloncian tablets immediately if they develop any type of rach and to contact their healthcare provider as soon as possible (see Warnings and Procession (4).

Frequencies (2-49); Emails Fettility: Advise fermals of reproductive potential who desire pregnancy that NSAIDs, including Meloxicam tablets, may be associated with a reversible delay in ovulation [see Use in Specific Populations (8-3)]. Fetal Toxicity

Inform pregnant women to avoid use of Meloxicam tablets and other NSAIDs starting 30 weeks gestation because of the risk of the premature closing of the fetal ductus arteriosus [see Warnings and Procautions (5.10) and Use in Specific Populations (8.1

anteria. So are Yanniga and Pracellanis (\$1) and Use in Specific Population (\$1) https://www.specific.execution.com/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/specific/s

SPL MEDGUIDE

Medication Guide for Honsterokial Anti-Inflammatory Druge (HSAIDs) What is the most important Information I should know about medicines NADADs can cause services side effects, hicklings: • Increased risk of a heart attack or stroke that can lead to death. This risk may happen early in technicat

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

Nausea	
 more tired or weaker than usual 	
darrhea	
Itching vour skin or eves look vellow	
jour sen or eyes look yeldw indicestion or stomach pain	
flu-like symptoms	
vomit hingd	
 there is blood in your bowel movement or it is black and sticky like tar 	
 unusual weight gain 	
 skin rash or bisters with fever 	
 swelling of the arms, legs, hands and feet 	
If you take too much of your NSAID, call your healthcare provider or medical helo right away.	get
These are not all the possible side effects of NSAIDs. For more information, asi	k unter a
healthcare provider or pharmacist about NSAIDs.	c you
Call your doctor for medical advice about side effects. You may report side effe	icts to
FDA at 1-800-FDA-1088.	
Other information about NSAIDs:	-
 Aspirin is an NSAID but it does not increase the chance of a heart attack. As 	pirin car
cause bleeding in the brain, stomach, and intestines. Aspirin can also cause ulc	ers in th
stomach and intestines.	
 Some NSAIDs are sold in lower doses without a prescription (over-the-count 	ter). Tab
to your healthcare provider before using over-the-counter NSAIDs for more th daws	an 10
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 prese	cobod D
not give NSAIDs to other people, even if they have the same symptoms that vo	
It may harm them.	
If you would like more information about NSAIDs, talk with your healthcare pro	vider. Yo
can ask your pharmacist or healthcare provider for information about NSAIDs	that is
written for health professionals.	
Additional Medication Guides can be obtained by calling Unichem at 1	-866-
562-4616.	
The other trademarks referenced are owned by third parties not affiliated with Laboratories Limited	Unichen
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Enovachem PHARMACEUTICALS	
Torrance, CA 90501	
This Medication Guide has been approved by the U.S. Food and Drup Administr	ation.
PACKAGE LABEL.PRINCIPAL DISPLAY PANEL	

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