MELOXICAM- meloxicam tablet Denton Pharma, Inc. DBA Northwind Pharmaceuticals

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.

Tablets USP, for oral use Localize names tar, you use and
 Localized and the second can be full. This chain may accurately in traditional and a long long as well described of an (4.1). The start back and the s

RECENT MAJOR CHANGES Beard Warning 5/2026 Indicators and Usage, Juwelle Pheumatoli Arthrite (IRA) Pauciaticula (2020 Dosage and Administration, General Dosaing Instructions (21) 6/2026 Dosage and Administration, Journile Theumatoli Arthrite (IRA) Pauciaticul Dosage and Administration, Journile Theumatoli Arthrite (IRA) Pauciaticul Cosage and Administration, Journile Theumatoli Arthrite (IRA) Pauciaticul Dousge and Administration, Jovenie Rheumatolin (? 21) (2016 2.4) 62016 Wannings and Precautions, Gardessance Thomholic Genetic (S.1) 52016 Wannings and Precautions, Heart Fallure and Eliema (5.3) 52016 Wannings and Precautions, Heart Fallure and Eliema (5.3) 52016

INDICATIONS AND USAGE

INDICATIONS AND USAGE INDICATIONS AND USAGE Obtoorthis (OA) (12) Phoumatoid Arthritis (OA) (12) Phoumatoid Arthritis (OA) (12)

 Losses effective dosage for the shortest who weigh adding (1.2)
 Loss the lowest effective dosage for the shortest duration consistent with individual patient treatment goals
 (2.1)
 (2.2) and RA (2.2): Starting does: 7.5 mg once daily Dose may be increased to 15 mg once daily • JPA (2.4):

Known hypenenstölvity to meliautam or any components of the drug product (4)
 Hibby yil anthras, urtitani, or other allengi-type neutrinos after taking auplinis or other NSADs (4)
 In the setting of CAIGs surger (4)

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benefits an expected to actually in relat of monitoring main function (1.6) *interchantical Biographics* (2.6) extension reprove type in an analysisticit, rescandon score (1.5) productions and/orally activation of the state state of the production and/orally. Its for the state of the production of the state of the production of the state of the production of the state of the production of the state of the production of the state of

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b) Biomania
b) Bio

8.1 Pregnancy 8.2 Lactation 8.3 Females and Nales of Reproducti 8.4 Pediatric Use 8.5 Genatric Use 8.6 Hepatic Impairment 8.7 Renal Impairment

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FULL PRESCRIBING INFORMATION

WARNING: RISK OF SERI OUS CAI SCULAR AND GASTROINTESTINA

Cardiovascular Thrombotic Events
 Nonsteroidal anti-inflammatory dru

Nonstervidal anti-inflammatory drugs (NSADD) (auss an increased risk of serious cardiovarcubic thromototic evolute, including myocardial treatment and may increase with duration of use [see Warnings and Precasations (2 - 3 1).
 Contractional (2 - 3 1).
 Warnings and Precasations (2 - 3 1).
 Warnings and Precasations (2 - 3 1).

1 INDICATIONS AND USAGE

I INDICATIONS AND USAGE
I.1 Osteoarthritis (OA)
Molexian tablest are indicated for relief of the signs and symptoms of osteoarthritis [
see Clinical Studies (14.1)].

see Check Studied (141)] 1.2 Resumption further (ReA) Maiorizant Labeits are indicated for real of the signs and symptoms of rheumatoid mitris (Lae (Check) Statistics (14.1)) 1.3 Jownik Resumption (24.1)) 1.3 Jownik Resumption (24.1) Statisticant Labeits are indicated for first of the signs and programs characterization polysticature course journal Resumption (24.1) Statistics (12.1) Statistics (14.1)) Statistics (14.1) Statistics (14.1

2 DOSAGE AND ADMINISTRATION

2.1 General Dosing Instructions

c.1 General Dosing Instructions Carefully consider the potential benefits and risks of Meloscam tablets and other trainment option balls descripting tablets and tablets. Use the levest affective Warnings and Processorties (3) is a second with incidenal patient trainment goal (see Warnings and Processorties (3) is a second ball therapy with Meloscam tablets, adjust the dose to so at an individual patient's media.

suz an induktual pazient's needs. In aduts, the maximum recommended daily oral dose of Mekoscam tablets is 15 mg regardles of formulation, in patients with hemodulysis, a maximum daily dosage of 7, 12 23). Meloscam tablets may be taken without regard to timing of meak.

2.2.Osteaarthritis For the relief of the signs and symptoms of osteaarthritis the recommended starting and maintenance ceal dose of Makoucam tablets is 7.5 mg one day. Some patients may receive additional benefit by increasing the dose to 15 mg once addy.

may recease additional benefit by increasing the dose to 1s mg once day. 2.3 Rheumatoid Arthritis For the relief of the signs and symptoms of rheumatoid arthritis, the recommended starting and maintenance and dose of Melosicam tables is 7.5 mg once day. Some plaints may recease additional benefit by increasing the dose to 15 mg once day.

Jackies in the second second attribute (Jackies) and second and second s

2 5 Perce --- remainingairment The use of Meloxicam tablets in subjects with severe renal impairment is not recommended.

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

ung tawe Cincar Manmacology (12.3) }.
2.6 Non-Interchangeability with Other Formulations of Meloxicam Molecum tables have not them equivalent systemic exposure to other approved formations of or al meloxicam. Therefore, Meloxicam tables are not interchangeable with other formations of or all meloxicam products wort if the tables are not interchangeable to end of the system tables of the system of the system of the system of the system to end of the system of the system of the system of the system tables of the system tables of the system of the s

3 DOSAGE FORMS AND STRENGTHS Natorican Tablets USP. 7.5 m; Light yolwn round find the bevilet edged, tablet with U.S.L. debossed on one side and 7.5 sebossed correlation on the other side 1.5 m; Light yeak, raquitus sitespace, raquitus with U.S.L. debossed on one side and 15 debossed centrally on the other side

4 CONTRAINDICATORS Mexicum atlabits are contraindicated in the following patients: In Source hypersensitively (e.g., analyhipetic reactions and serious dim nanctions) to machine and any components of the drong product (g and Wrannige and Princations) (e.g. and any components) and the drong product (g and Wrannige and Princations) (e.g. and any components) (e.g. and any components) (e.g. and exact any effect of any components) (e.g. and exact any components) (e.g. and exact any proteint is includy product. [g and Wrannige and Princations (g 15, Salar) (e.g. and exact any components) (e.g. and exact any components) (e.g. and exact any proteint is includy product. [g and wrannige any product of any components) (e.g. and product of any effect of any components) (e.g. any effect of any component of any component of any effect of any components) (e.g. any effect of any

INGS AND PRECAUTIONS

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The accuracy to the transmission of the constrained calls of the stating of CMBC (see Terminal Termin

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

5.2 Gastrointestinal Bleeding, Ulceration, and Perforation

3.1 Gathaltistical blacking, Ukraslin, and Parlondim. McMa, hushing municipation, can case straing superstrational (1) giving strain strain black hushing multiple strates, which is a first strain strain strain black hushing multiple strates, which is a first strain strain strain black hushing multiple strain strain strain strain strain strain grants in high parlies that strain strain strain strain strain black hushing multiple strain strain strain strain black hushing multiple strain s

Additionally regardless and advanced for decisional and advanced for decisional and advanced for decisional advanced for the second advanced for the s

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

Inform patients of the warring signs and proptoms of hepatotock?) (e.g., nanosa, target, kthorpy, darhna, prottos, pandos, tytt copper quadret for interfaciencies, and "the prottos, pandos, tytt copper quadret for interfaciencies, and "the prottos, pandos, tytt, copper quadret for interfaciencies, and "the prottos, pandos, tytt, copper quadret for interface of \$\$ systemic multitations occur (e.g., essimplik, rat, etc.), discontinue Meixscan immediate, and particular electrical pandos of the patient [see Use in Specific Pepadation (d & and clinical Pharmacology (2.23).

Projections (s. b) and Linke arrhenizations(21,22-3). 5 3.4 Appentationals NSANDs, Including Melosciam, can lead to new onset or worsaning of preventions programming, and provide the standard sector of the standard detaction. Placets tables applications: converting earry and IACE (Intelbates, Baladid detaction), or a Damp Interactions (7, 7). Nonitor tobod prossure (BP) during the initiation of NSAD treatment and throughout the course of therapy.

5.5 Heart Failure and Edema

Additionally, fluid retention and edema have been observed in some patients treated with NSAIDs. Use of meloxicam may blant the CV effects of several therapeutic agents used to treat these medical conditions (e.g., diuratics, ACE inhibitors, or angiotensin receptor bickners [ABb3]] [see Drug Interactions (7)].

Avoid the use of Meloscam in patients with severe hant failure unless the bondfs are expected to solvewigh the risk of extremely have failure. If Meloscam is used in patients which serves hand failure, monthe patients for large of ensuring have failure. 5.6 Read Tookty and HyperKatamini <u>Baral Tookty</u> and HyperKatamini Baral Tookty, and Meloscam, cased anni dana, and other rend inputs?

Rend taxicity lact also been seen in patients in whom rend prostaglanders have a compensationary role in the maintenance of rend perfusion. In those patients, administration of a short DAM may cause a date sequence fractions for prostagland formation and, accounting, in rend labol flow, which may proceeded event rend. market and accounting, in rend labol flow, which may proceeded event rend. market and accounting in rend labol flow, which may proceeded event rend. market and accounting in rend labol flow, which may proceeded event rend. market and accounting in the proceeding have the event of the state of duratics and AcC Inholders or AMB, and the elderly. Discontinuation of NSAID therapy to caused followed by exercising the provincement durate.

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.

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

J. No information is available from controlled chical studies regarding the use of Ministerions (J. No information is patients with advanced related as Avaid Teau and Mikescam in patients with advanced related as a studies. Avaid The use of Mikescam Endowed Avaid Studies and Stu

<u>Huperkalama</u> Increases in serum potassium concentration, including hyperkalamia, have been reported with use of NSAIDs, even in some patients without renal impairment. In palaints with normal renal function, these effects have been attributed to a hyporenisment', hypadiosteronism state.

reportentiame-opposition stream status. 5.7 Anaphylactic Reactions Midioxiam has been associated with anaphylactic reactions in patients with and without known hypersensibility to midioxic an and in patients with apprivilentiative actimatic game contraindications (4) and Warmings and Privacations (5.8).

Seek emergency help if an anaphylactic reaction occurs.

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may deminish the utility of diagnostic signs in detecting intections. 5.13 Laboratory Monitoring Bacusios services GI bilading, hepatotoxic/by, and renal hijiry can occur without warning symptome or signs, consider monitoring patients on long-term MSAID treatment with a CEC and a chemistry profile periodically lise Warnings and Procautions (5.2, 5.3, 5.6).

The contrast protection (2.5.2, 5.16)
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 Detable and the contrast protection of the contrast protection of the contrast protection (2.5.1)
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6.1 Clinka Thulk Experiment Because circle structure are conducted under webly varying conditions, adverse reaction realised and a this clinka of a dirig cannot be directly compared to rates in the Analysis of the clinka structure of reflect the rate does not an experiment and a structure of the structure of reflect the rate does not an experiment Analysis of the analysis of the structure of the structure of the structure Databatement 2014 and and any not a bender to be structure of the structure Databatement and Databatement at athemes bender (1017) OA activities on 1013 BA

Characterizati and Rhammatiki Amritiki The Methods million 2016 of clinical trial database includes 10.122 OA patients and 1012 R patients translated with NetroCare 7.1 prophysy, 1955 OA patients and 1313 RA patients patients for at kase of clinical patients of the state one-yak Approximatily 2015 OF of these patients were translers for at kase one-yak Approximatily 2015 OF of these patients were translers for a kase one-yak Approximatily 2015 OF of these patients were translers for a kase one-yak Approximatily 2015 OF of these patients were translers of the kase one-yak Approximatily 2015 OF of these patients and the state of the clinical patients and the state of the sta tras. A 12-week multicenter, double-blind, randomized trial was conducted in patients with osteoarthritis of the knee or hip to compare the efficacy and safety of Neitoxicam with placebo and with an active control. Two 12-week multicenter, double-blind, randomized triak were conducted in patients with rheumatoid arthritis to compare the efficacy and

Sake energiency hulp 1 an anaphyticit reaction accurs. 32 Eascretistics of Atoman Related Carboph SeashNar An obspotiation of patients with anitiation may how supplies accellate actions attem may based private, information comparison by anaphysis, based patients of 22 metabolic private, information comparison of patients with private and the standard private and the SAMDA has been reported in such separis- samples and control actications of 10 Minor Machine and and a standard patients and control actication of 10 Minor Machine and action and and and the standard patients and an anti-patient for changes in the standard and and antipatient of a standard patients for changes in the standard patient and and an antipatient of a standard patients for changes in the standard patient and and an antipatient of a standard patient of the standard patient of the standard patient and and an antipatient of a standard patient of the standard patient of th

safety of Meloxicam with placebo.

Table 1a depicts adverse events that occurred in x2% of the Meloxicam treatment groups in a 12-week placebo- and active-controlled osteoarthrifts trial. Table 1b depicts adverse events that occurred in x2% of the Meloxicam treatment groups in two 12-week placebo-controlled meumatoid arthrits trials.

	Placebo	Meloxicam 7.5 mg daily	Meloxicam 15 mg daily	Diclofenac 100 mg daily
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
Fal	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				
Pharyngkis	1.3	0.6	3.2	1.3
Upper respiratory tract infection	1.9	3.2	1.9	3.3
Skin				
Rash ²	2.5	2.6	0.6	2.0

	Placebo Meloxicam 7.5 mg daily Meloxicam 15 r			
No. of Patients	469	481	477	
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 0	Conditions			
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-				
oint 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 NOS *	1.7	1.0	2.1	

The adverse events that occurred with Meloxicam in a2% of patients treated short-term (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-Cr Trials

	4.6 Weeks Co	atralled Triple	6 Month Con	trolod Triple
	4-6 Weeks Co Meloxicam 7.5 mg daih			
No. of Patients	8955	256	169	306
Sastrointestinal	11.8	18.0	26.6	24.2
Abdominal pain	2.7	2.3	4.7	2.9
Constipation	0.8	12	1.8	2.6
Tiamba	1.9	27	5.9	2.6
Dyspeosia	3.8	7.4	8.9	9.5
Fahilance	0.5	0.4	3.0	2.6
Maricola Varicola	2.4	4.7	4.7	7.2
/omling	06	0.8	1.8	2.6
Body as a Whole	6.6	0.0	1.0	2.0
Accident household	0.0	0.0	0.6	2.9
dema "	0.6	2.0	2.4	1.6
Pain	0.9	2.0	3.6	5.2
Central and Peripheral Nervous S		2.0	3.0	3.2
Toriness	1.1	16	2.4	2.6
Hearlar he	2.4	27	3.6	2.6
lematologic				
Inemia	0.1	0.0	4.1	2.9
Musculoskeletal				
Arthraipia	0.5	0.0	5.3	1.3
lack pain	0.5	0.4	3.0	0.7
Paychiatric				
nsomoja	0.4	0.0	3.6	16
Respiratory				
Coughing	0.2	08	2.4	1.0
Joper respiratory tract infection	0.2	0.0	83	7.5
škin				
Pruritus	0.4	1.2	2.4	0.0
Rash 1	0.3	1.2	3.0	1.3
Urinary		-		
Micturition frequency	0.1	0.4	2.4	1.3
Jrinary tract infection	0.3	0.4	4.7	6.9

* WHO preferred terms edema, edema dependent, edema peripheral, and edema legs combined † WHO preferred terms rash, rash erythematous, and rash maculo-papular combined	
Higher doses of Meloxicam (22.5 mg and greater) have been acsociated with an increased risk of serious GI events; therefore, the daily dose of Meloxicam should not excised 15 mg.	
Pediatrics	
Pauciarticular and Polyarticular Course Juvenile Rheumatoid Arthritis (IRA)	
Three handbard and eighty-saves platest with pacatificitud and polyaritude cores (PA more expended to Maccan the Boass simply that that 100 ms 110 mg by the table simply that that 100 ms 110 mg by the table simply that that 100 ms 110 mg by that 100 ms 110 mg by that 100 ms 110 mg by the base simple s	
The following is a list of adverse drug reactions occurring in <2% of patients receiving	

ing the course of the trials. The ecific subgroup effect.	adverse events did not demonstrate an age
is a list of adverse drug reacting	ns occurrine in <2% of patients receiving
clinical trials involving approxim	
Vhole	aleroic reaction, face edima, fatious, fever, hot flishes, making, switches, weight increase
	angina pactoris, cardiac failure, hypertension, hypotension, hypotension, hypotension, hypotension, hypotension,
	nconvulsions, paresthesia, tremor, vertigo
tinal	colits, dry mouth, duodenal ulcer, eructation, esophagits, gastric ulcer, gastribis, gastrointestinal hemorrhage, hematemesis, hemorrhagic duodenal ulcer, hemorrhagic gastric ulcer, intestinal perforation, melena, pancreatitis, perforated duodenal ulcer, stomatitis ulcerative
and Rhythm	arrhythmia, palpitation, tachycardia
6	kukopenia, purpura, thrombocytopenia
liary System	ALT increased, AST increased, bilrubinemia, GGT increased, hepatitis
nd Nutritional	dehydration
	abnormal dreaming, anxisty, appetite increased, confusion, depression, nervousness, somnolence
	asthma, bronchospasm, dyspnea
	alopecia, angloedema, bulbus eruption, photosensitivity reaction, pruritus, sweating increased, urticaria
	abnormal vision, conjunctivitis, taste pervension, tinnitus
tem	abuminuria, BUN increased, creatinine increased, hematuria, renal failure

ritis

6.2 Post Marketing Experience

Body as a Who Cardiovascular Central and Pe Gastrointestin Heart Rate and Homstelook

- creat materials Experience
 The following adverse rescents have been diverted of adverse gaves argument is or diverse that the second of the se

Warning

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

	Table 3 Clinically Significant Drug Interactions with Meloxicam						
Drugs that In	erfere with Hemostasis						
Clinical Impact:	Matrician and addrosogulatis such as warrain have a symptize diffect on biadely. The concentrativit and embodynam and addrosogulation have as in protecting the system of the direct on a system of the						
intervention:	Mandor patients with concombant use of Makoicam with anticoopstants (e.g., wardarki), artipitative apents (e.g., applin), selective sendors in respitate inhibitors (SSRIs) for signs of bleacing [see Warnings and Procautions (5.11)].						
Aspirin							
	Controlled clinical studies showed that the concombant use of NSAIDs and analysis doses of aspirin dose not produce any greater therapeutic effect than the use of NSAID admeter 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 admet [see Warnings and Procuedoes (5.2)].						
intervention:	Concombant use of Milotinician and low dose aspirin or analysis: doses of aspirin is not generally recommended because of the increased risk of bleeding (see Warnings and Precautions (5.11)). Meloniciam is not a substitute for low dose aspirin for cardiovascular protection.						
	, Angiotensin Receptor Blockers, or Beta-Blockers						
canca impact:	KMD/s may deminish the authypertuniess effect of augioansis enverting enzyme (ACE) inhibers, augioansis incorport backers (AREA), or back-backers (Including propertunid). Indices that are added you can be applied (Carding these on Area), where any anti-anti-anti-anti-anti-anti-anti-anti-						
intervention:	During concomfant (see Of Mesicican and ACE inhibets, RM, et al., beta-factors, monther block pressure to artise data block pr						
Diuretics							
Clinical Impact:	Chick Landback, as welk is pont- matering deventions, howed that HSUD: reduced the nativenet effect of loop duretics (e.g., forosemide) and thiadel duretics in some patients. This effect has been attributed to be NSUD inhibition of revial prostagatand's synthesis. However, studies with forosemide agents and materixed are reduction in nativenet effect. Furscenides single and multiple dose planmacophramics and planmacophramics and planmacophramics and planmacophramics.						
intervention:	During concombant use of Makuciam with duratics, observe paraleters for signs of worsening remain function, in addition to assuring duratic efficacy including anth/sportansise effects (see Warnings and Precautions (5.6)).						
Lithium							
Cinical Impact:	NSAUs 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 productions in renal lithium clearance.						
intervention:	During concomitant use of Melocican and Bihum, monitory patients for signs of Bihum toxicity.						
Methotrexate							
	Concomitant use of NSAIDs and methotrexate may increase the risk for methotrexate toxickly (e.g., neutropenia, thrombocytopenia, renal dysfunction).						
	During concombant use of Meloxicam and methodresiate, monitor patients for methodresiate toxicity.						
Cyclosporine							
	Concomitant use of Meloxicam and cyclosporine may increase cyclosporine's nephrotoxicity.						
intervention:	During concomitant use of Melocican and cyclosporine, monitor patients for signs of worsening renal function.						
NSAIDs and S							
	Concombant use of metoxican with other NSAIDs or salicylates (e.g., offlurisal, salisalate) increases the risk of OI toxicity, with little or no increase in efficacy (see Warnings and Proceedings (5.2)).						
intervention:	The concentrate use of metoxicam with other NSADs or salcylates is not recommended.						
Pemetrexed							
	Concombant use of Meloxican and pemetrexed may increase the risk of pemetrexed associated myelosuppression, renal, and Gl toxicby (see the pemetrexed prescribing information).						
1	During concomitant use of Meloxicam and permitrixed, in patients with renal impairment whose creatinine clearance ranges from 45 to 29 mL/min, monitor for myelosuppression, renal and GI toxicity.						
intervention:	Nations's taking melosician should interrupt dosing for at least five days felore, the days fe						

Les UN SECTOR POPULATIONS 2.3 Property Balances Balan

There are no adequate and well-controlled studies of Meloxicam in pregnant women. Data from observational studies regarding potential embryofetal risks of NSAD use in women in the first or second threates of pregnancy are inconclusive. In the general U.S. population, all chically recognized pregnancies, regardises of drug exposure, have a background rate of 2-4% for major mellormations, and 15-20% for gregnancy loss.
In animit reproduction studies, emittyofield adeit was observed in rats and rabbes tratatad during the particle of organopanesis their themotexian at an at disces equivalent to 3.5- and 5.5-tmest the maximum recommande human dasa (MHHD) of Naturckian intravada richardon a dispath hant a discles equivalent to 7.5-tmest the MHD. If par- ticles and recommendation and an and at a set of the set of the set of the set of the display planticity. An adversation of the set of the set of the set of the set of the Nature of the set of the display planticity. An adversation of the set of the set of the set of the set of the Nature of the set of the propagenesis that and of the set of t
Based on animal data, prostaglandins have been shown to have an important role in endometrial vaccular permeability, blastocyst implantation, and decidualization. In animal studies, administration of prostaglandin synthesis inhibitors, such as meloxicam, resulted in increased pre- and post-implantation loss.

These are no sources on the effects of Meloscam during abor or delivery. In animal tables, PEADO: A block meloscam, initiate prostagandin synthesis, classe delayed patrumtico, and increase the incidence of stillarth.

Meloxica organog mg of M rabbits t the hear compari based or meloxica

greater, respectively, than the MRHD based on BSA compa throughout organogenesis.

Broughout oppropriate. Our administration of mediacion the preparent rate during the petition through lattices increased in technic of dynamics, adjustment and the second of the pre-tices comparison. **B.1 Catter B.2 Lesten B.2 Lesten** Throw are no hourns data available on whether metacicam is prepared in hours may dynamics the dynamics of the pre-second of the second of the pre-tices of the p

<u>Data</u> Animal Data

Number Later Meloxicam was present in the milk of lactating rats at concentrations higher than those in plasma.

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of MSADs, including Namer Larger, many management of the main sector of the main sector

8.7 Renail Impairment No dose adjustment is nocessary in patients with mild to moderate renail impairment. Patients with severe renail impairment have not been studied. The use of Meloxican in subjects with severe renail impairment is not recommended. In patients on hemologikas, mildiocam should not exceed 7.5 mg per day, Meloxicam is not diayable [see Dosage and Administration (2.1) and Cline if hemorecology (2.12.3).

10 OVERDOSAGE Symptom Robwing acute NSAID overdiscages have been typically limited to lethargy, drewiness, nauxel, combing, and expander pais, which have been generally reversible with supportive card. Gastrofisterial baseding has a cocurrent, Hyperinerion, acute read and Pracedance (13, 53, 54, 68, 50). These focusing bases read (ac bit harings)

Manage patients with symptomatic and supportive care following an ISAID overforage. There is no solver, the symptomatic care supportive care following an ISAID overforage. The symptometry of the symptomatic care of the symptometry is a symptometry of the symptomatic care of the symptomatic patients serve with in four hours of negation or in patients with a large overdage (3 to 10 them the incompression) and dataget for a durate, labeling of units, hereodays, or hemperfusion may not be useful due to they prove helper.

There is limited experience with metoxican overdisage. Choixityramine is internet accelerate the classrace of metoxican. Accelerator formould in metocican by 4 g oral does of choixityramine gluon Dive thread and you as dominational a a chical Table for additional information about overdisage treatment, cal a policion control center (1-800-222-1221).

11 DESCRIPTION More can Tables: UP are a noncorrected and informationy drug (MSAD). Each tables contains 7.5 mp ang makerkam for oral administration. More can is charactering designates at 4-hydroxy-2-anety- $H_{\rm C}$ -methy-2-bitaph/2-H-1,2-bitrobilizme-3-carboranish-1,1-bitrok. The melecular walks is 351.4.1 is empirical formula is C _2H 12H 2O _{2} _{2} = and E has the following structural formula:

and a

Molectam is a pastel yellow sold, practically insoluble in water, with higher solubility observed in strong actits and basics. It is very slightly soluble in methand Makinzkam Makenzam has plan values of 11 and 4.2. We have a 1.1 in or 4 consolution per 1.7. A Makenzam Makenzam water and a sublet as a tablet for oral administration containing 7.5 mg or 15 mg makezam.

The inactive ingredients in Meloxicam tablets USP include colloidal silicon dioxide, crospovidone, lactose monohydrate, magnesium stearate, microcrystalline cellulose, povidone and sodium citrate dihydrate.

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12.3 Pharmacokinetics

12.3 Environmentational Environmentation of the second sector of the

Table 4 Single Dose and Steady-State Pharmacokinetic Parameters for Oral 7.5 mg and 15 mg Meloxicam (Mean and % CV) ".							
			Steady State			gle Dose	
Pharmacokinetic Parameters	i (%CV) H		ed) ¹ Elderly males (Fed) ²		Renal failure (Fasted) H	lepatic insufficiency (Fasted)	
		7.5 mg ⁺ tablets	15 mg capsules	15 mg capsules	15 mg capsules	15 mg capsules	
N		18	5	8	12	12	
Cmax	[ug/mL]	1.05 (20)	2.3 (59)	3.2 (24)	0.59 (36)	0.84 (29)	

t max	[h]	4.9 (8)	5 (12)	6 (27)	4 (65)	10 (87)
t 1/2	[h]	20.1 (29)	21 (34)	24 (34)	18 (46)	16 (29)
CL/f	[mL/min]	8.8 (29)	9.9 (76)	5.1 (22)	19 (43)	11 (44)
V 2/15	[L]	14.7 (32)	15 (42)	10 (30)	26 (44)	14 (29)
* The parameter values in the t † not under high fat conditions 2 Meloxicam tablets 5 V Z/f =Dose/(AUC+Kel)	table are from various studies					

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Specific Populations -Pediatric

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Etisrly make (seS years of age) exhibited meloxicam plasma concentrations and staady-state pharmacolenatics initiar to young make. Etisrly females (seS years of age) had a 47% high ethod Crass as at compared to younger concentrations in the stellary female, the adverse every grofts was compared for both ethory patient population. A smaller fire for faction was found in stillary female patients in comparison to siderly make patients.

Angust: Impairment: Federing a single 5 mg dates of maloscam there uses to marked difference in plasma concentrations in plasmits with mild (CHS-Hop) Case II (or modurate (CHS-Hop) Case II) is plasmit (memory modulated to the single concentration of the single concentration with mild to modulate insplace insplammer. Frances which are not head to insplant and the single concentration of the single concent

enall imperiment. Total ring above an encountage of making and the second of the standard of the standard with the second constraints with the the second constraints and the second constraints and

Hemodaliyal Folowing a single dose of milioxicam, the free Cmax plasma concentrations were higher in galantic with reveal take is on chronic hemodaliyal. (15) file infraction) to concentration in plasmic therefore, additional dose; are not encessary after hemodaliyals; Methodicam is not dialyzable [see Dosage and Administration (2.1) and Use in Specific Population (2.7).

hemodality, Allicocch in the divide land bases and Administration (2.1) and the Marchinest Constraints (2.1) and the second sec

Methodrowate: A study in 13 resonance of entries (MA) parkets vehanced the effects of marked AMM and a final state on the bagerout effects of the parket state of the state o

13 NONCLINICAL TOXICOLOGY

13 NORCLIMICAL TOXICOLOGY 13. Carcinogenesis, Mudagenesis, Impairment of Fertility Cercinogenesis Thare was not reveals in bunner incidence in leng-term carcinogenchy studies in rats (104 weeks) and mice (109 weeks) administered metricram at cardiodese to 0.08 mightsph in rata and up to 0.03 mightsph in mice (1pt 0.5 and 26 mise, respective), The maximum recommanded human dese (UHHO) of 15 mightsp Makekcam back on body strate and (155) commanden.

Mutagenesis

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.

nt of Fertility

Meloxicam did not impair male and female fertility in rats at oral doses up to 9 mg/kg/day in makes 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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The case of Philotecan for the management of signs and symptoms of esteen-thresh se-sent and the symptoms of the signs of

umatoid Arthritis (JRA) Pauciarticular and Polyarticular 14.2 juvenile Rh Course The use of Meloxicam for the treatment of the signs and symptoms of pauciarticular polyarticular course juvenile Rheumatoid Arthritis in patients 2 years of age and older was evaluated in two 12-week, double-blind, paralel-arm, active-controlled trials.

Both choice included three arris: negregate and the data of mildicates. In both mapping the second second

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NCC 10914-010-00; Bottiss of 90 Strongo Store at 20 ° to 25 ° (56 ° to 77 ° 97) [see USP Controlled Room Temperature]. Keep Malockam Tablets USP in a dry place Dispunse tablets in a dryt container. Keep this and all medications out of the reach of children.

17 PATIENT COUNSELING INFORMATION Advice the patient to read the FDA-approved patient tabeling (Medication Guide) that accompanies each prescription dispensed. Additional Medication Guides can be obtained by calling Unichem at 1-866-582-4616.

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TRADECOUNCE V Inform patients of the warning signs and symptoms of hepatotoxicity (e.g., nausea, fatigue, lethargy, darrhea, pruritus, jaundice, right upper quadrant tenderness, and "flu like" symptoms). If these occur, instruct patients to stop Meloxicam tables and seek immediate melocita therapy (see Warnings and Procedurios (5.3.3).

Hard Fahre and Fahres Advise patients to be also for the symptoms of compactive heart fahres including attorneds of the same section of the symptoms and the contact fair haltbcare bottoms and the same section of the symptoms and the contact fair haltbcare attarbard fahrest families and the same section (e.g., difficult sections), seeling of the lace or threads, bottom at patient to seak remote and the same of the lace or threads. The same section (e.g., difficult sections), seeling of the lace or threads, bottom at patients to seak remote and the same sector (e.g., difficult sections).

Surbout Stith Reactions Andrea patients to stop Meterican tablets immobiliarly if they develop any type of rach to construct the Mathematic provided as can possible (are Marrings and Press, 1997). The Mathematican and the Mathematican and the Mathematican Advised formalism of reproductions potential and self-to programmy that MARINs, including Meterican tablets, may be associated with a reversible delay in ovulation (are Usin in Service Provided on (3.3).

Spectre repositions (is 3) [<u>Fail Toxics</u>] Inform pregnant women to avoid use of Melovkam tablets and other NSAIDs starting 30 weeks gestartion because of the risk of the premature closing of the fetal ductus anteriosus [see Warnings and Precautions (5.10) and Use in Specific Populations (8.1).

Avoid Concember Use of NSAIDs Inform patients that the concember use of Meloxicam tablets with other NSAIDs or Regulate (e.g., dimits), statistable is not recommended due to the increased risk of Avoid (e.g., dimits), and avoid (e.g., dimits), and avoid (e.g., dimits), and Proceedings (5.2) and Drog Interactions (7.7). J. Avoit patients that NSAIDS may be present in "over the counter" medications for transmert of code, lever, or insomnia.

Medication Guide for Nonsteroldal Anti-Inflammatory Drugs (NSAIDs) What is the most important information 1 should know about medicines called Nonsteroidal Anti-Inflammatory Drugs (NSAIDs)? NSAIDs can cauge earlies called (March Industra) kid Nonstervidal Anti-Inflammatory Drugs (NSAIDs)? AIDs can cause serious side effects, including: ncreased risk of a heart attack or stroke that can lead to death. This risl rhappen early in treatment and may increase: with increasing doses of NSAIDs

i with increasing doses of MARIDS. In Damappen and Discose before a other a heart surgery caked a "converse year by pages graft (CAMC), and attack, unkes part healthcar provide the styre of the complex on a traceast is do attacher healthcar provider test years to complex on a traceast land. If years attack if year attacher that attacher attacher attacher attacher in years attacher attacher attacher. Ben attacher attacher attacher attacher healthcar attacher attacher attacher healthcar attacher attacher attacher healthcar attacher attacher healthcar healthca

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Lise of NSAIDs and Low-Dose Aspirin Inform patients not to use low-dose aspirin concomitantly with Nell they talk to their healthcare provider [see Drug Interactions (7)].

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

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

Cardiovascular Thrombotic Events

Hepatotoxicity

specific transmission of forwards a function of instantial forwards of the ADA sector transmission of the ADA sector and the ADA

a exactly as protoched to the based been possible for your treatment of for the identicate one packed or What we are stated with the state of the state of the state of the state of the What we are stated as an and the state of the state of the Mass of the state of the state of the state of the state of the madical conditions such as different types of arthritis, mentrual cramps, and other types of short-term pain.

The second second

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

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average of the artis, age, hands and read.
 If you take too much of your NSAID, call your healthcare provider or get medical help right away.
 These are not all the possible side effects of NSAIDs. For more information, ask your healthcare provider or pharmackt about NSAIDs.

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

Principal Display Panel NDC: 70934-010-30



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