MELOXICAM- meloxicam tablet NuCare Pharmaceuticals, Inc. HIGHLIGHTS OF PRESCRIBING INFORMATION
These highlights do not include all the information needed to use MELOXICAM TABLETS safely and effectively. See full prescribing information for MELOXICAM TABLETS. J.S. Approval: 2000 WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EV Ear hij practicing information for compiles based sarring.

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BOXED WARNING
WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL
EVENTS

EVENTS

- Monstroullar Thrombotic Events

- Nonstroullar and Influence of the Control of the Con

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

Studies (14-1):

1.2 Rheumatoid Arthritis (RA)

Meloxicam is indicated for relief of the signs and symptoms of rheumatoid arthritis [see

Clinical Studies (14-1)].

As Juvenile Reumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course Meloxiciam is indicated for relief of the signs and symptoms of pauciarticular or polyarticular course Juvenile Rhaumatoid Arthritis in patients 2 years of age and older [see Cincal Stotias (4-2)].

2 DOMES AND ADMINISTRATION
2.1 General Design Instruction
2.1 General Design Instruction
2.2 General Design Instruction
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options before activity in user mislancium, to as the leasest effective dosage for the
Proceedings (4) Limited and Proceedings (4) Limited Instructions (4) Limited Ins

remensal patents needs.

In addits, the marinum recommended daily oral dose of melecicam is 15 mg regardless of formation. In patients with hemodalysis, a maximum daily dosaps of 7.5 mg is recommended see but less rispectife populations (8.7) and odd clinical Pharmacology (12.3) J. Meloxicam may be taken without regard to timing of meals.

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

The State of the S

2.5 Renal Impairment
The use of melorican in subjects with severe renal impairment is not recommende
In patients on hemodalysis, the maximum dosage of meloxicam is 7.5 mg per day
Chical Pharmacology (12.3);

Clinical Pharmacology (12.3) 2.

S. Non-InterAmpability with Other Formulations of Meloxik am Meloxik mutablets have not shown equivalent systemic expourse to other approved formulations of an interdoctant. Therefore, resilicace an tables are not interchangeable to the contraction of an interdoctant motion, resilicace mutables are not affect-integrable to this same. Do not substitute similar does strengthe of meloxicam tablets with other formulations of oral meloxicam products.

3 DOSAGE FORMS AND STRENGTHS

Melloxicam bablets, USP
7.5 mg; yellow cobured, round, biconvex, tablets, debossed with "158" on one side and "C on the other.

1.5 mg; yellow cobured, round, flat beveiled tablets, debossed with "CIPLA" on one side and "150" on the other.

4 CONTRAMBICATIONS

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5 WARNINGS AND PRECAUTIONS

5 WARNINGS AND PRECAUTIONS

3.L Cardiovascular Thrombotic Events

Clinical risks of several COX.2 selective and nonselective NSAIDs of up to three years duration have shown an increased risk of serious cardiovascular (CV) thrombotic duration have shown an increased risk of the district serious cardiovascular (CV) thrombotic duration was considered to the contract of the contract

use appears to be similar in those with and without known CV disease or risk factor CV disease. However, patients with known CV disease or risk factors had a higher use against 1, to be sinker in those with and without hower Of bissues or in its factors for deadless inclinated with a section of the factor of the factor

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5.5 Heart Failure and Edema

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may demins the durity of diagnostic signs in detecting miscrems.

5.13 Laboratory Monitoring

Because serious G blooding, hepatotoxicity, and renal injury can occur without warning
symptoms or signs, consider monitoring patients on long-term MSAID treatment with a

CSC and a chemistry profile periodically is see Warnings and Procusionis (52, 53, 56).

Cité and a chamistry pride protectée, pla au Warning and Precasions (\$2, \$3, \$5, \$6).

A MONESE BLACTOR STATE DE L'ANDRE STAT Adults Osteoarthritis and Rheumatoid Arthritis

<u>Optional Printing and Philometrolis of Architects</u>

The melotiscers Plass of 2 Critical Fair disclarabase includes 10,122 OA patients and 1012 RA patients treated with melotical To 3 Englishy, 3505 OA patients and 1351 RA patients treated with melotical To 3 Englishy Melotical on those doctors was deministered to 65 Fair content of the Control of t

The content of the co

	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
Fall	0.6	2.6	0.0	1.3
influenza-like symptoms Central and Peripheral	5.1	4.5	5.8	2.6

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
Skin				

Raish ² 2.5 2.6 0.6 2.0

¹WHD preferred terms edema, edema dependent, edema peripheral, and edema legs combined
²WHD preferred terms rash, rash erytherabous, and rash maculo-papular combined

Table 1b: Adverse Events (%) Occurring in ≥ 2% of Meloxicam Patients in two 12-

	Placebo	Meloxicam 7.5 mg daily	Meloxicam 15 mg dail
No. of Patients	469	481	477
Gastrointestinal Disorders	14.1	18.9	16.8
Abdominal pain NOS 2	0.6	2.9	2.3
Dyspeptic signs and symptoms 1	3.8	5.8	4.0
Nausea 2	2.6	3.3	3.8
General Disorders and Administration Site Co	enditions		
Influenza-like ilness 2	2.1	2.9	2.3
Infection and Infestations			
Upper respiratory tract infections- pathogen class unspecified ²	4.1	7.0	6.5
Musculoskeletal and Connective Tissue Disor	rders		
loint related signs and symptoms 1	1.9	1.5	2.3
Nervous System Disorders			
Headaches NOS 2	6.4	6.4	5.5
Skin and Subcutaneous Tissue Disorders			
Rash NOS 2	1.7	1.0	2.1
*MedDRA high level term (preferred terms): dyspeptic sig- aggrawated, enuctation, gastrointentinal instation), upper (jaryngits NOS, pharyngits NOS, sinvastis NOS), pich retu- aggrawated, joint crepitation, joint effusion, joint swelling *MedDRA preferred term: nausea, abdominal pain NOS, in NOS	respiratory tract ded signs and sy)	infections-patho mptoms (arthral)	gen unspecifier pis, arthralgia

The adverse events that occurred with meloxicam in $\approx 2\%$ of patients treated short-term (4 to 6 weeks) and long-term (6 months) in active-controlled osteoarthrikis trials are presented in Table 2.

	4 to 6 Weeks	6 Month	Controlled Trial	
	Meloxicam 7.5 mg daily	Meloxicam 15 mg daily	Meloxicam 7.5 mg daily	15 mg daily
No. of Patients	8955	256		306
Gastrointestinal	11.8	18.0	26.6	24.2
Abdominal pain	2.7	2.3	4.7	2.9
Constipation	0.8	1.2	1.8	2.6
Diarrhea	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 1	0.6	2.0	2.4	1.6
Pain	0.9	2.0	3.6	5.2
Central and Peripheral Nervous Sy		•		•
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				
Arthraigia	0.5	0.0	5.3	1.3
Back pain	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 ²	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

The Control term unit, value in primarium, see a second to the control term unit and the control term and population control term and the control term and t

Body as a Whole	allergic reaction, face edema, fatigue, fever, hot flushes, malaise, syncope, weight decrease, weight increase
Cardiovascular	angina pectoris, cardiac failure, hypertension, hypotension, myocardial infarction, vasculitis
Central and Peripheral Nervous:	System convulsions, paresthesia, tremor, vertigo
Gastrointestinal	colitis, dry mouth, duodenal ulcer, enuctation, esophagitis, gastric ulcer, gastritis, gastroesophageal reflux, gastrointestinal hemorrhagic duodenal ulcer, hemorrhagic gastric ulcer, intestinal perforation, melena, pancreatitis, perforated duodenal ulcer, perforated gastric ulcer, stomatitis ulcerative
Heart Rate and Rhythm	arrhythmia, palpitation, tachycardia
Hematologic	kukopenia, purpura, thrombocytopenia
Liver and Biliary System	ALT increased, AST increased, bilirubinemia, GGT increased, hepatitis
Metabolic and Nutritional	dehydration
Psychiatric	abnormal dreaming, anxiety, appetite increased, confusion, depression, nervousness, somnolence
Respiratory	asthma, bronchospasm, dyspnea
Skin and Appendages	alopecia, angioedema, bullous eruption, photosensitivity reaction, pruritus, sweating increased, urticaria
Special Senses	abnormal vision, conjunctivitis, taste perversion, tinnitus
Urinary System	abuminuria, BUN increased, creatinine increased, hematuria, renal failure

4.3 Pentanvistrios Expression.
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Table 2 Clinically Significant Drug Interactions with Milenkam Drug May 1 Manufacture with Namestania Drug May 1 Manufacture
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intervention: Concomitant use of meloxicam and low dose asserin or analosis doses of assirin is not operandy recommended because of the increased risk of bleeding I see Warnings and Precautions (5.11)).
Intervention: Concentrant use of meloxistam and low dose appirin or analysis doses of aspirin is not generally recommended because of the increased risk of bleeding [see Warnings and Precautions (\$.11)].
ACE Inhibitors, Angiotensin Receptor Biockers, or Beta-Blockers
Clinical prace; 155/4656; many demands the arithmysteriousive effect of angeleteria conversing extyres (ACE) in addition, and contained the arithmysteriousive effect of angeleteria conversing extyres (ACE) in addition, and are also are also are also are and experiment experted in a few and are also are also are are also are also are also are also are are also are also are also are also are are also are are also are a
Intervention: During concomitant use of meloxicam and ACE inhibitors, ARBs, or beta-blockers, monitor blood pressure to ensure that the desired blood pressure is obtained.
During concendrate use of missistems and AEC inhibitors or ARBs in patients who are alsolarly column-depleted, or have impaired renal function, months for segret of worsening renal function. [see literature of the production (5.6)]. When the seed using a war administrated concendrater, jumpated concendrater, furnature for section date adequately information, and the production (5.6).
Diuretics
Elicid larges: Elicid studies, as well as post- marked power booms to be included as well as post- marked power booms to be included as well as post- marked power booms to be included as well as post- marked power booms to be included as well as post- marked power booms to be included as well as post- marked power booms to be included as well as post- power booms to be included as power booms to be includ
Intervention: During concomitant use of melbricken with districks, observe patients for signs of worsening renal function, in addition to assuring duretic effects (see Warnings and Precautions (5.6)).
Uthlum
Clinical Impact MSAIDs have produced elevations in plasma filtium levels and reductions in renal thium clearance. The mean minimum thium concentration increased 15%, and the renal clearance decreased by approximately 20%. This effect has been attributed to NSAID inhibition of renal prostaglandin synthesis [see Clinical Pharmacobby (12.3)].
Intervention: During concomitant use of moloxicam and lithium, monitor patients for signs of lithium toxicity.
Methotrexate
Clinical Impact Economitant: use of NSAIDs and methotrecate may increase the risk for methotrecate toxicity (e.g., neutropenia, thrombocytopania, renal dysfunction).
Intervention: During concomitant use of maloxicam and methodrevate, monitor patients for methodrevate toxicity.
Cyclos portine
Clinical Impacts Concomitant use of meloxicam and cyclosporine may increase cyclosporine's neptirotoxicity.
Intervention: During concomitant use of melbrickim and cyclospoprine, monitor patients for signs of worsening renal function.
NSAIDs and Salicylates
Clinical Impact Concomitant use of meloxicsam with other NSAIDs or salicytates (e.g., offlurisal, sassialate) increases the risk of GI toxicity, with little or no increase in efficacy (see Warnings and Procustions (S.21).
Intervention: The concomitant use of melaxicam with other NSAIDs or salicylates is not recommended.
Pemetrexed
Clinical Impact Concomitant use of meloxicam and pernetrexed may increase the risk of pernetrexed-associated myelosuppression, renal, and Cli toxicity (see the pernetrexed prescribing information).
Intervention: During concomitant use of meloxicam and penetriesed, in patients with renal impairment whose creatmine clearance ranges from 45 to 79 mt./min, monitor for myelosuspression, renal and GI toxicity.
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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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11 DESCRIPTION

Motorizam is a monitoreoidal anti-inflammatory drug (MSAID). Each tablet contains 7.5 mg
or 15 mg motorizam, USP for oral administration. Nelexicam is chemically designated as
4-hydroxy2-methy4-Mc-5-methy4-2-milasoly2-2-M-2-benrosinative-3-cartox semised 1-ex4-hydroxy2-methy4-methy4-methy4-methy4-2-milasoly2-2-M-2-benrosinative-3-cartox semised 1-ex4-hydroxy2-methy4-methy

Meloxicam is a pale yellow solid, practically insoluble in water, with higher solubility observed in strong acids and bases. It is very slightly soluble in methanol. Meloxicam has an apparent partition coefficient (bg P) $_{\rm app}=0.1$ in n-octanol/buffer pH 7.4. Meloxicam has pKu values of 1.1 and 4.2.

Meloxicam is available as a tablet for oral administration containing 7.5 mg or 15 mg meloxicam, USP. The inactive ingredients in meloxicam tablets, USP include starch, microcrystalline cellulose, lactose anhydrous, colloidal silicon dioxide, sodium citrate dihydrate, magnesium stearate.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Milosiciam has analysisc, anti-inflammatory, and antipyretic properties.

The mechanism of action of melaxicam, like that of other MSAIDs, is not completely understood but involves inhibition of cyclossyspenses (COX-1 and COX-2). unrear-titude de invitación de protection de l'explosinguisme (CAA- altre CAA-2).

Referencia de la potent inhibitor of protectiguisme symphosis (vivo. Melanciam concentrations marchand during therapy havap produced in rise of direction. Protectiguisment concentrations de l'explositation de l'expl

12.3 Pharmacokinetics Absorption

13.2 Pharmacolontest.

Ministration.

See a second second

Table 4 Single Dose and Steady-State Pharmacokinetic Parameters for Oral 7.5 mg and 15 mg Meloxicam (Mean and % CV) ¹

	Steady State			Single Dose	
Pharmacokinetic Param (%CV)	neters Healthy male adults (I	ed) ² Elderly males (Fed	i) ² Elderly females (Fed)	Renal failure (Fast	ed)Hepatic insufficiency (Faster
	7.5 mg ³ tablets	15 mg capsules	15 mg capsules	15 mg capsules	15 mg capsules
N	18	5	8	12	12
C _{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)
CL/f [mL/min]	8.8 (29)	9.9 (76)	5.1 (22)	19 (43)	11 (44)
V ₂ /Y ⁴ [L]	14.7 (32)	15 (42)	10 (30)	26 (44)	14 (29)
The parameter values in the t					

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Digoxin: Meloxicam 15 mg once daily for 7 days did not alter the plasma concentration profile of digoxin after \$-acstyldigoxin administration for 7 days at clinical doses. In who testing found no protein binding drug interaction bottween digoxin and melaxicam. profile of logical alley pour logical processing of the control of

13 MONCHINECA TOXICOLOGY

TOXI

Mutasenesis
Meloxicam was not mutagenic in an Ames assay, or clastogenic in a chromosome aberration assay with human lymphocytes and an in vivo micronuclaus 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 males and 5 mg/kg/day in females (up to 5.8- and 3.2-times greater, respectively, than the MBHD based on BSA comparison).

14 CHRCAL STUDIES
14.1 Otherwise the Benemated Arthritis
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SOSTIGATE.

saturage Store at 20 to 25 C (68 to 77 F) [See USP Controlled Room Temperature]. Keep meloxicam tablets in a dry place.

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

Face this and all medications and of the resist of Children.

37 PATRIC COMMISSION SHOPMATON

Ank this plainter to resist for \$1.0 Appears on planet lishing (Medication Guide) that accumpates each price originate (appears) and the state of the following features for the following features of the following features feat

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Advise patients to be alert for the symptoms of congestive heart failure including shortness of breast, unexplained weight gain, or oderna and to contact their healthcare provider if such symptoms occur [see Warnings and Precaucitors (5.5)].

antane, acid Helactions
Advise patients to stop meloxicam immediately if they develop any type of rash and to contact their healthcare provider as soon as possible [see Warnings and Precautions (5.0)].

contact that healthcare provider as soon as possible (see Warnings and Pricactions).

Advise financial configurations of regressional values of several providers of the PASAD, including advancam, may be associated with a reversible only in roundation; less than Space?

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Topics of (CASC):

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The risk of getting an uter or bleeding increases with:

Open Native of Etamuch Vietne, or stomach or intestinal bleeding or other age with use of EAGLOS.

Stating madicines called "Conficionationists", "andicoapulates", Open health Oncrossing dose of MAIDS.

o smoking drinking alcohol

NSAIDs should only be used:

• exactly as prescribed

• at the lowest dose possible for your treatment

• for the shortest time needed

at the level date possible for your treatment
 More are MARION.
 MORAL or are without a recommendation from the level of the

Tell your heathcare provider about all of the medicines you take, including prescription or over-the-counter medicines, vitaminis or herbal supplements. NSAIDs and some off medicines can interest 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 Nonsteroidal Anti-inflammatory Drugs (NSAIDs)?"

• new or worse high blood pressure

new or worse high blood pressure
 heart failure
 fiver problems including liver failure
 kidney problems including kidney failure
 low red blood cells (anemia)

Bit foll discourses—we like threatening skin reactions in the threatening skin reactions
 Chen raide effects of PSAIDs include : stomach pain, consignation, diarrhea, gas, hearthum, nuasea, vomiting, and dischess.

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

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

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	roduct Info	mation									
,	roduct Type		HUMAN PRE	SCRPTION	item (Sou			NDC 680	DC-68071-2186(NDC-69097- 99)		
	oute of Admin	istration	CINAL								
A	ctive Ingred	ient/Active	Moiety								
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м	BLOXICAM (UNII	VG2QFE3CGL	(MELCOCAM	LMILVS2QFE	(acar)		MILORC	MM.		15 mg	
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