IRGRIGHTS OF PRESCRIBING INFORMATION
These highlights do not include all the information needed to use MELOXICAMTABLETS valely and
effectively. See full prescribing information for MELOXICAMTABLETS.

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Melucicam is a non-servoidal neti-inflammatory drug indicated for:

Oureaumhrite (OA)(1.1)

Else umanoid Arthrite (RA)(1.2)

Jave elle Flee umanoid Arthrite (RA)

DOSAGE FORMS AND STRENGTIES
 Melasican Tables: 75 mg, 15 mg (3)

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Revised: 7/2019

FULL PRESCRIBING INFORMATION: CONTENTS*
WARNING RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS
11NDICATIONS AND USAGE

1.1 Obsocratifis (OA)
1.2 Rheumands Arthrife (RA)
1.3 Investle Rheumands Arthrife (IRA) Fauciarticular and Polyarticular Course
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FULL PRESCRIBING INFORMATION

WARNING RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS

WARNING REN OF SERIOUS CALIDOVASCILLAR AND GAST ROUNTESTRUA.

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*Non-restellar die fellemanners (reng, (NSMID)) came en increased rick of restructure candisvascular demodale cerveni, including speceror field intercision and new low, which can be faind. This rick may enser such in terceiore and may increase with disordine of use of the Marketian of candinal cerval for the special intercision and new low restruction of the origin of committee of the serious (careas) array hyposa garden (3.1). Engenimental interficial care fine to string circumstantial interficial care fine to string circumstantial interficial (3.1) and Womings and Frontantian (3.1). Engenimental interficial (3.1) and Womings and Frontantial (3.1) and the serious including before, described, and preferenties of the stume for interficial (3.1) and the serious including before, described, and preferenties of the stume for investigation, which can be fast, These events can exercise and printer of the stume for investigation of the stume for the serious (3.2). Meeting are at greater rick for serious CI events [see Viernings and Procusions (5.2)].

L1 Osteoarheids (OA)
Molessicamis indicated for relief of the signs and symptoms of osteoarheids; for Chical Studies
(441):
L2 Blommaniel Arthrids (UA)
Molessicamis indicated for relief of the signs and symptoms of rhomanoid arthrids [1 or Clinical Studies
(441):

1.3. Javenile Rheumanid Arthriki (JRA) Psuclaricular and Polyaricular Course
Metoricum is indicated for relief of the signs and symptoms of psuclaricular course plaveaile Rheumanid Arthritis in patients who weights 260 kg [see Dosage and Administration (2-4) and Claimol Studies (Jack)

2 DOSAGE AND ADMINISTRATION

2. General Daving Unstructions
Carefully consider the potential benefit and risks of nebotic an and other treatment options before
the consideration of the potential benefit and risks of nebotic and and other treatment options before
the consideration of the potential benefit of the consideration of th patients needs.

In adults, the maximum recommended daily oral dose of meloxicam is 15 mg regardless of formulation in patients with hermodulysis, a maximum daily dosage of 7.5 mg is recommended [see Use in Specific Populations (8.7 mg collision)]. The production (8.7 mg collision) from the collision of Primarcology (12.3)].

Meloxicam may be taken without regard to timing of meals.

Moltocamen for team visitors are posses to an approximate property of the strict of the signs and systems of some definit the recommended storting and minimizance and note to 15 mg earlier shelly. Some patient key receive additional basefully investigate doubter to 15 mg core earlier, the close to 15 mg core earlier, the close to 15 mg core and the core

All Javealta Rhomantaid Arthritis (IAS) Peaciariscular and Polyariscular Ceurse
For the recument of prevale rhemanoid arthritis, the recommended onal dose of melosticiam is 7.5 mg
see daily included with weigh 26 OE, priere was an additional benefit demonstrated by increasing
the dose down 7.5 mg included rishs.
Melosticiam tailors should not be used in children who weigh 5-60 kg.

3 DOSAGE FORMS AND STRENGTHS Meloxicam tablets, USP:

• 7.5 mg; yellow coloured, round, biconvex, tablets, debossed with "158" on one side and "C" on the

15 mg: yellow coloured, round, flat bevelled tablets, debossed with "CIPLA" on one side and "159" on the other.

4 CONTRAINDICATIONS

- A Constitution of the Constitution of the Constitution pricess:

 Network States and Constitution of the Constitution of Consti

5 WARNINGS AND PRECAUTIONS

S WARNINGS AND PRICALTIONS

Like dissist of the Promotive Event

Like dissist of the

pany occur. There is no consistent evidence that concurrent use of aspirin mitigans the increased risk of serious CV thrombotic evens associated with NSAID use. The concurrent use of aspirin and an NSAID, such as meloxicam, increases the risk of serious gastrointestinal (GI) events [see 'Hornings and Procuntion(S.2)].

Preconton(2, 1);
Same Post Coronary Armey Papass Graft (CARG) Surgery
Two large, comolled clinical trials of a COX-2 selective NSAID for the treatment of pain in the first
10-14 days following CARG surgery found a interested incidence of myocardial infarction and stroke.
NSAIDs are commiscionated in the seeing of CARG [see Constantialization (4)].

Based M (Endo)

Description of the Conference of the Parish National Regions have demonstrated that guidens is used before related to the post of the process of the proces

Avoid the use of molocicum inpatients with a recent MI unless the benefits are expected to outseigh the risk of recurrent CV thrombotic events. If meloxicam is used in patients with a recent MI, monitor patients for signs of cardiac ischemia.

patients for signs of conflict inclusion. An other control of the conflict inclusion of the conflict inclusion. An other control of the conflict including an other control of the conflict including confliction of the confl

- Administrating, posters for an administration to the Gaussian Configuration of an attenuated reaches (at Stangles as Ministration the Ga Riski in SNAD reveal patients).

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closely for relatives of Chiborologi (see Prog. Instructions; 77).

33 Hapainstockie;
Bavation of ALT or AST (three or more times the upper limit of normal [ULN]) have been reported in approximately 19 to ALMD-normal patients in clinical risks in Andidation; nor, consenters fact, cases of severe hapaics (steps; lackfulling infinisate hapatits, liver necrosis, and hapatic failure have been reground.

Bevations of ALT or AST (these than three times ULN) may occur in up to 15% of patients researd with NARIDs (actining and solutions).

NSAIDs including meloxicam Informapatient of the warring signs and symposus of hepatonxicity (e.g., nusses, Infigue, Jethangy, diarrhas, puritins, juantice, right upper quadratt underroses, and "In-like" symptoms. Jf clinical signs and symposus consistent with liver disease develops, or if systems mediseastions occur, contingulalis, ands, etc., bifscontine meloxicam immediately, and performs clinical evaluation of the provent (see Use 8-perce). For purpose of the year of the technique (22-23).

journal ner to te styrege reputation (as) unto Limitar transmission (2.2.2).

El Hypermania

NALDa, lechaling selecticum, can lead in new onto or woversing or perexisting hypermains, relieferance or within may combine to his interacted infections (CV events. Patient stating angionisms convention ensures (ACSL) inhibitors, shainted mitretic, or though disorders may have impaired response to these therepies wherein the (NALD) in terrol parameterize (7.1).

Maniate Bondo pressure (BP) during the initiation of NSAID resourcer and throughout the course of therapy.

therapy.

So Heart Railers and Edema
The Constitute and additional NSADT Trialers. Collaboration are analysis of probability of excendent data.
The Constitute and additional NSADT Trialers. Collaboration are analysis of probability of excent data and the constitute of the constitut

prospiralization for heart failure, and death.

Additionally, fluid resertion and edems have been observed in some patients treated with NSAIDs. Use
of melonicar may but the CV effects of several therapeutic agents used to treat these medical
conditions (e.g., distretics, ACE inhibitors, or angiourusin receptor blockers [ARBd-]) [see Drug
lateractions (7)].

Avoid the use of meloxicam in padents with severe heart failure unless the benefits are expected to outseigh the risk of worsening heart failure. If meloxicam is used in padents with severe heart failure, monitor padents for signs of worsening heart failure.

5.6 Renal Toxicity and Hyperkalemia Renal Toxicity

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Long serro distinct (NSLD), ice (India) period can, has resulted in send populary accross,
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periodic never and compensation. Federa are generated to deline according and the server
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servery in the presentations.

construction of a color, and the charly. Decentification of NSAD heavy analysis for disciplinaries can be exceeded to the color of the

Increases in serum potassism concentration, including hyperkalenia, have been reported with use of NSAIDs, even in some patients without renal impairment. In patients with normal renal function, these effects have been antibuted to a hyporenismic-hypoaldossreoism state.

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and Henrings and Presentation (5.6);

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35 Seriem Sida Reactions

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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.

S.13 Laboratory Monitoring

Because serious GI bleeding, hepatotościty, and renal injury can occur without warning symptoms or signs, consider monitoring patients on long-turn NSAID resument with a CBC and a chemistry profile-profiledingly are Warnings and Presentations (S.2.5.3.5.4).

portion-leafly tow branching and Proceedinate (2.5, 2.5, 5.6).

AND/REM RELATIONS

The following adverse reactions are discussed in greater detail in other sections of the labeling:
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troquency reported adverse events in all resources goals across metocacum trans.

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the remaind arthrist is compare the efficacy and steps of melociacum treatment groups in a 12-even
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Table 1b depicts adverse events that occurred in 22% of the moloxicam treatment groups in two 12-week placebo-controlled rheumanid arthrifts trials.

Table 1a: Adverse Events (%) Occurring in 2.2% of Meloxicam Patients in a 12-Week Osteoarthritis Placebo- and Active-Controlled Trial

	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	5.1	4.5	5.8	2.6
Central and Peripheral				
Nervous System				
Dizziness	3.2	2.6	3.8	2.0
Headache	10.2	7.8	8.3	5.9
Respiratory				
Pharyogitis	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
WHO preferred terms edems, e-	dema depend	ent, edema periphera	al, and edema legs	combined
2WHO preferred terms rash, rash	h erythemator	as, and rash meculo-	papular combined	

	Placebo	Meloxicam 7.5 mg daily	Meloxicam 15 mg daily
No. of Patients	469	481	477
Gas trointestinal 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 Condi			
nfluenza-like illness ²	2.1	2.9	2.3
infection and Infestations			
Jpper respiratory tract infections- suftogen class unspecified ¹	4.1	7.0	6.5
Musculoskeletal and Connective Tissue Disorder	rs		
oint related signs and symptoms 1	1.9	1.5	2.3
Vervous System Disorders			
leadaches NOS 2	6.4	6.4	5.5
skin and Subcutaneous Tissue Disorders			
lach NOS 2		1.0	2.1

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 osteo arthritis trials are presented in Table 2.

	4 to 6 Weeks	Controlled Tris		Controlled Trials
	Meloxicam	Meloxicam		Meloxicam
		15 mg daily	7.5 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	8.0	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	8.0	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:	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
Mus culos keletal		•		•
Arthralgia	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
WHO preferred terms edems, ede				

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

serious CI evens, therefore, the displayate of melaciam should not receed to sag.

Polithric's

Benciaridate and Dibustical Course, hermite libraminal Architis LIBA.)

There handred and injudices are proposed to the control of the course of

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 Sys	stem convulsions, paresthesia, tremor, vertigo
Gas trointes tinal	colitis, dry mouth, duodenal ulcer, eructation, esophagitis, gastric ulcer, gastricis, gastrocucer, perforated duodenal ulcer, bemorrhagic duodenal ulcer, hemorrhagic gastric ulcer, intestinal perforation, melena, pancreatitis, perforated duodenal ulcer, perforated duodenal ulcer, stomatitis ulcerative
Heart Rate and Rhythm	arrhythmia, palpitation, tachycardia
Hematologic	leukopenia, purpura, thrombocyoppenia
Liver and Biliary System	ALT increased, AST increased, billimbinemia, GGT increased, hepatitis
Metabolic and Nutritional	dehydration
Psychiatric	abnormal dreaming, auxiety, appetite increased, confusion, depression, nervousness, sommolence
Respiratory	asthma, bronchospasm, dyspnea
Skin and Appendages	alopecia, angioedema, bullous eruption, phomsensitivity reaction, pruritus, sweating increased, urticaria
Special Senses	abnormal vision, conjunctivitis, name perversion, timitus
Urinary System	albuminuria, BUN increased, creatinine increased, hematuria, ronal failure

7 DRUG INTERACTIONS

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

Table 3 Clinically Significant Drug Interactions with Meloxicam	
Drugs that Interfere with Hemostasis	
Clinical Impacts Meloxic can and a microagalases such as warfarin have a synergistic effect on bleeding. The concomitant use of meloxic am and anicroagalases have an increased risk of serious bleeding compared to the use of either drug alone.	
Seronnin release by planeless plays an important role in hemistasis. Case-control and cohort epidemiological studies showed that concomitant use of drugs that interfere with seronnin relyation and an NSAID alone.	
Intervention: Monitor patients with concomitant use of meloxicam with anticoagulums (e.g., warfarin), antiplanter agents (e.g., aspirin), selective serotonin receptable inhibitors (SSRIs), and serotonin neceptable inhibitors (SSRIs) for signs of bleeding [see Wornings and Precontions (5.11)].	
As pirita	
Clinical Impacts Controlled clinical studies showed that the concomitant use of NSAIDs and analysesic doses of aspirin does not produce any greater therapeutic effect than the use of NSAID alone [see Wirmings and Precontions as compared to use of the NSAID alone [see Wirmings and Precontions (S.7)].	
Intervention: Concentrate use of melvicicam and low does aspirin or analysesic doses of aspirin is not generally recommended because of the increased risk of bleeding [see Warnings and Precontons (5.11)].	
Meloxicamis not a substitute for low dose aspirin for cardiovascular protection.	
ACE Inhibitors, Angiotensin Receptor Blockers, or Beta-Blockers	
Clinical Impact NSAIDs may diminish the antihypertensive effect of angionesin converting enzyme (ACE) inhibitors, angionesin receptor blockers (actuding proprantols).	
In parients who are elderly, volume-depleted (including those on discretic therapy), or have recal impairment, coadministration of an NSAID with ACE inhibitors or ARBs may result in desertionation of renal function, including possible acute renal failure. These effects are usually reversible.	
Sterwinds: During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince bridge steasure to seem that the desired blond pressure to seems that the desired blond pressure to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to bridge and the desired blond pressure to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to be a desired blond pressure to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to be a desired blond pressure to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to obtained. During concensiate use of milosiciaman ACE inhibitors, ARBs, no prince to obtained. During concensiate use of milosiciaman active to obtained. During concensiate use of milosiciaman act	
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Direction Statement of the Control o	-
Chical Imped Clinical studies, as well as cost-	
marketing observations, showed that NSAIDs reduced the nativeretic effect of loop disvetics (e.g., furosemide) and thiazide disvetics in some patients. This effect has been nativitied to the NSAID inhibition of renal prostaglandin synthesis. However, studies with furosemide agents and meloxicam have not demonstrated a reduction in nativeretic effect. Furosemide single and multiple dose pharmacolynamics and pharmacolynamics and pharmacolynamics are not affected by multiple dose	s of meloxicam
Intervention: During concomitant use of meloxicom with distractics, observe parients for singus of worsening renal function, in addition to assuring distracts of efficacy including anthypertensive effects [see Warnings and Procusions [5,6]].	
Lidium	
Clinical Impact, NSAIDs 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 prostaglandin synthesis [see Clinical Pharmacodogy (12-3)].	
Intervention: During concomitant use of meloxiscan and lithium, monitor patients for signs of lithium taxicity.	
Methotresate	
Clinical Impact; Concomitant use of NSAIDs and method exate may increase the risk for methodrexate toxicity (e.g., neutropenia, thrombocyspoenia, renal dysfunction).	
Intervention: During concomitant use of meloxicam and methorevane, monitor patients for methorevane training.	
Cyclosparine	
Clinical Impact Concomitant use of meloxicam and cyclosporine may increase cyclosporine's nephrotoxicity.	
Innevention: During conconstant use of meloxicam and cyclosporine, monitor patients for signs of worsening renal function.	
NSAIDs and Salicylates	
Clinical Impact; Concomitant use of moloxicam with other NSAIDs or salicylates (e.g., diffunisal, salsalate) increases the risk of GI toxicity, with little or no increase in efficacy [see Warnings and Precoutions (5.29).	
Intervention: The concomitant use of involvations with other NSAIDs or salicylans is not recommended.	
Pemetersed	
Clinical Impact Concomitant use of multivation and pennetreand may increase the risk of pennetreand-associated myelosuppression, renal, and GI taxicity (see the pennetreand pensetribing information).	
Intervention: During concomitant use of meloxicam and penetresed, in patients with renal impairment whose creatinine clearance ranges from 45 to 79 ml.lmin, monitor for myelosuppression, renal and GI toxicity.	
Patients taking meloxican should interrupt dosing for at least five days before, the day of, and two days following permet exed administration.	
In patients with creatinine clearance below 45 mL/min, the concomitant administration of meloxicam with permetward is not recommended.	

BLP regulates
Blish Similars, in Intellige medication, during the third relations of programs; increases the right of
the resource cleanes of the fired dictions entries the NASIDs, including understann in
program women starting at 30 weeks of gestudion (third trimeway) I now Marriage and Procautions (5.10)

permanen coloure of the first distance sourceions. Avoid use of NNASID, strating restrictions, in a Deep and versus a security of all wheek of greated sourceived in the restriction of the security of the se

Meloxicam was not toratogonic when administered to pregnant rats during fetal organogenesis at oral doses up to 4 mplaying (2.6-fold greater than the MRINO of 15 mg of meloxicam based on BSA comparison). Administration of meloxicam being pregnant rabbits throughout embryogenesis produced an

increased incidence of sepal defects of the heart at an oral dose of 60 mg/kg/day (78-fold greater than the MRRID based on BSA comparison). The no effect level was 20 mg/kg/day (56-fold greater than the MRRID based on BSA conversion), it may and rabbit, enthroptobality occurred and card antendrican doses of 1 mg/kg/day and 5 mg/kg/day, respectively (0.55- and 6.5-fold greater, respectively), than the MRRID based on BSA comparison) when administrated motional orangements.

There are no human data available on whether motoxicam is present in human milk, or on the effects on breastfed infants, or on milk production. The developmental and health benefits of breastfeeding should be considered along with the mother's clinical need for motoxicam and any potential adverse effects on the breastfed infant from the melociacum or from the underlying maternal condition.

Libita Arrienal datas Meloxicam was present in the milk of lactating rats at concentrations higher than those in plasma.

8.3 Females and Males of Reproductive Potential

B. I Femiles and Males of Reproductive Potential biferilities.

Finals:

Finals:

Based on the mechanism of action, the use of proxigatudio-mediand NSAIDs, including melosicion, may dainy as present requires of sovial na folialists, which has been anoticined with reversible infertility intelligence of the control of th

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)].

8.5 Geriatric Use

B. Geriante Use

Derly patient, compared by unsper patient, see a grear risk for NAID-associated serious continuational agranoism data. And we read observe in actions. The arth paste bows for the delaying patients for a shipped bows for the delaying patients for adverse delayers. The arth paste bows from the delaying patients for adverse strong for the form of the delaying patients for adverse strong for the delaying patients for adverse strong for the delaying patients for adverse strong for the delaying patients for the delaying for the delay

19 OVERBOSAGE
Symposes following actors NSAID overdescapes have been specially intended to be following, described as a constant of the control of the contr

Process URITHING
There is limited experience with meloxicam overdosage. Cholestyvamine is known to accelerate the clearance of meloxicam Accelerated removal of meloxicam by 4 g oral doses of cholestyvamine given three times a day was doministeard in a Clinical trial. Administration of cholestyvamine may be useful following an overbrokage.

11 DESCRIPTION

11 DESCRIPTION
Meloxicamis a nessero cidal arti-inflammanry drug (NSAID). Each tablet comains 7.5 mg or 15 mg meloxicam, USP for oral administration. Meloxicam is chemically designated as 4-hydroxy-2-melys-N-6-mehyl-2-diazohyly-2-fi-1/2-benoridazina-3-carbo-tamide-1/1-dioxide. The molecular weight is 331-l. in empirical formula is c. p. in 37-30 g. 2-3 and then the following succurant formula:

Meloxicamis a pale vellow solid, practically insolable in water, with higher solability observed is strong acids and bases. It is very slightly solable in methanol. Meloxicam has an apparent partition coefficient (log P) $_{\rm app} = 0.1$ inn-octamibutlet $_{\rm P}$ P. 4. Meloxicam has péa values of 1.1 and 4.2.

Meloxicamis 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 arthrefores, colloidal silicon dioxide, sodium citrae dibedates menes issum searcar.

12 CLINICAL PHARMACOLOGY

LEAD Mechanism of ALL M

Abstration
The advance below with filling of melecular computer was 87% following a single and done of 37 mg or compared with 75% following in the property of the compared with 75% mg or compared wi

Table 4 Single Dose and Steady-State Pharmacokinetic Parame

	Steady State			Single Dose		
Pharmacokinetic Paran (%CV)	neters Healthy male adults	(Fed) Elderly males (Fe	d) Elderly females (F	rd) Renal failure (Fast	ed) Hepatic insufficiency (Fast	rđ)
	7.5 mg ³ tablets	15 mg capsules	15 mg capsules	15 mg capsules	15 mg capsules	
N	18	5	8	12	12	
Cmax [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)	
12 [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 -/T 4 [L]	14.7 (32)	15 (42)	10 (30)	26 (44)	14 (29)	

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probably responsible for the order to establish which account for 16% and 4% of the databasismed factories.

Account of the control of the co

ne analysis, utilizing population pharmacokinetics body-weight, but not age, was the single covariate for differences in the meloxicam apparent oral plasma clearance. The body-weight apparent oral clearance values were adequate predictors of meloxicam exposure in pediatric

patients. The pharmacokinetics of meloxicam in pediatric patients under 2 years of age have not been investigated. Gerianic

Gerianic Electry mules (el.5 years of age) exhibited meloxicam plasma concentrations and steady-stans plasma colinects similar to young makes. Electry for mules (el.5) years of age) that 47% indigated AUC age and age of the collectry for mules (el.5) years of age) that 47% indigated AUC age are constructed from the plasma of the collectry for mules, the developer and the collectry formals, the adverse reverse profile was comparable for both elderly passer postulation. A smaller free fraction was found in electry freathy age in comparison to delively make patients.

Sex Voung femules exhibited slightly lower plasma concentrations relative to young males. After single doese of 75 mg melosicam, the mean elimination half-life was 19.5 hours to refer femule group as compared to 23.4 hours for the mile group, As usualy stang, the date were similar [175] hours vs 21.4 hours, This (pharmacolkarict difference does to general est likely to be of linter clinical suportance, predicts, and the contraction of the co

generation produced and the production of the pr

offulphis wing a single dose of meloxicam, the free C $_{\rm max}$ plasms concentrations were higher in patients renal failure on chroric hermicallysis (1% free fraction) in comparison to healthy volunteers (2.3) articularly lens of the control of the configuration of the patient period of the control of

Administration (2.1) and the an Specific Populations (8.2)).

Days International Conference and Conference and

Digitals. Molto Cam 15 ng one daily for 7 days did not due to a plante concentration profile of digitals and price of pages and the contraction for 7 days at clinical done. In vitro useing front on protein digitals and provide the contraction for 7 days at clinical done. In vitro useing front on protein the contraction of the contract

13 NONCLINICAL TOXICOLOGY

D. Carcinogenesis, Mitagenesis, Impairment of Fertility

Galizinogenesis

There was no increase intumer incidence in long-neuron conjugately motion in rare (104 weeks). There was no increase intumer incidence in long-neuron or 0.5 maylgody in not only up to 0.5 maylgody in not 0.5 maylgody in not only up to 0.5 maylgody in not only up to 0

Munapenesis Meloxicam was not munapenic in an Arnes assay, or classogenic in a chromosome aberration assay with human lymphocytes and an in vivo micronucleus uses in mouse bone marrow.

maximent 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 MRHD based on BSA comparison).

placeho. The use of microcard for the management of signs and symptoms of concentration was evaluated in six double-billed, active controlled trails a made the U.S. singing from 4 works' in 6 months' demands in his double-billed, actives controlled trails a made the U.S. singing from 4 works' in 6 months' demands in the procession. By mighty and officience SR 100 mighty and consistent with the efficiency were in the U.S. trail.

The use of microcard for the recement of the sign and symptoms of rheumanid admiriles was evaluated in a 12-week, double-billed, committee minimization trial. Microcard per S. print, part S. print, part

14.2 Juvenile Rheumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course

The use of meloxicam for the treatment of the signs and symptoms of pauciarticular or polyarticular course. Investile Rheumatoid Arthrifes in patients. 2 years of age and older was evaluated in two 12-week, double-billing, parallel-arm, active-corrolled trials.

work, double-linds, parallel-sum, artic-controlled trials.

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MON CORPORTISE-15 Besties of 2000

NOC (1997-138-15 15 Besties of 2000

Dispense tablets in a tight container.

Keep this and all medications out of the reach of children.

IP PATENT COUNSELING INFORMATION
About the patient are and the TDA approved patient labelling (Medication Guide) that accompanies each
prescription-labelline in read the TDA approved patient labelling (Medication Guide) that accompanies each
prescription and the patient of the control of the following information before initiating therapy with
an INSAID and periodically during the course of ongoing therapy.

Advise patients to be alert for the symptoms of cardiovascular thrombotic events, including chest pain, shortness of breath, weakness, or slurring of speech, and to report any of these symptoms to their healthcare provider immediately [see Wernings and Precardioses (5.1)].

noatticare provider immulately (see warming and Precautions (3.1)). Generalization Bladfani, Ulcaration, and Perforation Advise patients to report symptoms of afectations and bleeding, including epigastric palis, dyspepsia, media, and humanissis to their baddieres provider. In the setting of concentrations of for-shore supiral for centle prophylatis, inform patients on the increased risk for the signs and symptoms of GI bleeding [see Wormings and Precautions (2.5)].

Happinonicity

Information with a warring signs and symposes of happinonicity (e.g., names, fatigue, bethaupy, dardrea, parties, junatice, right upper quadrant nonderness, and "fin-like" symposes, if these occur, or produce to stop moto licitims and seek immediate moderal therapy (see Verenage and Percentions (5.3)).

[S.3]

5.39 J. Heart Falture and Edoma.
Advice patients to be alert for the symptoms of congestive heart failure including shortness of breath, unexplained weight gain, or edoma and to contact their healthcare provider if such symptoms occur [see Workings and Procustors (4.5)].

Anaphylactic Reactions

Anotheric Heartinia

Melton patients of the signs of an anotheric reaction (e.g., difficulty breading, swelling of the face or frond), history patients is set its medium energousy help if these occur | not Commissionium (d) Stiffens | patients | patient

resums are proviner as soons a prosision par warming our retreatment (3-17). Fremile Fertilia: Advise fermales of reproductive potential who desire preguncy that NSAIDs, including meloxicam, may be associated with a reversible delay in ovulation [see Use in Specific Populations (8.3)]. Field Toxicity.

Inform programs women to avoid use of moloxicam and other NSAIDs starting at 30 weeks gestation because of the risk of the premiture closing of the fread ductus arteriosus [see Wornings and Procustions (S.1)].

because of the risks of the presumes constained the fired duction services II. We Varnings and Proceedings (2.1) and the Superly Repulsion (2.1).

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Which is most important information I should have about medicines called Nessawaidal Anti-inflammatory Days (NSAIII)s)

Which is most important information I should have about medicines called Nessawaidal Anti-inflammatory Prompt (NSAIII)s)

NSAIIIs can come services side effects, including:

NSAIIIs can come services side of the called the called the data. This risk may happen early in which increasing doses of NSAIIIs

of the increasing doses of NSAIIIs.

Do not take NSAIDs right before or after a heart surgery called a "coronary artery bypass graft (CABG)."

Combut.

A beart surgery called a "coronary artery bypass god
A solid salage NSAIDs after a recent heart stack, miles; your healthcare prevides rafe, you to,
You may hear an interested fish of amende heart stack if you take NSAIDs after a recent heart
stack.

The second risk of the clings, afters, and many (note for the control of the control of the control of the control of the clings, afters, and many (note for the control of the contr

eased risk of bleeding, ukers, and tears (perforation) of the esophagus (tube leading 1 the mouth to the stomach), stomach and intestines:

anytime during use
 without warning symptoms
 that may cause death

The risk of getting as where or bleeding increases with:

| Spatitions of Standard Meeting, consends a instead bleeding with on at NSAIDs a caller age
| Spatial Confessional Confessional

NSAIDs should only be used:

• exactly as prescribed
• at the lowest dose possible for your treatment
• for the shortest time needed

What are NSAID:)

Note are used as weap aim and redores, wetting, and host different mode from medical conditions such as different types of about the medical conditions. What shaded not take NSAID: The note take NSAID: The note take NSAID as made, klows, or other allergic reaction with aspirition or any other NSAID.

*right before or after heart hyposis and particular and other types and other types and other types are described by the state of the particular and the state of the state hyposis and the state of the stat

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Product Informa	tion					
Product Type	HUM	AN PRESCRIPTION DRUG	hem Code (Source	NDC SETIE-1ISE(N	DC:69297-158	
Rusts of Administra	tion ORAL					
Active Ingredien	/Active Moiety					
	Ingredie	nt Nume		Basis of Strength	Strength	
MELOXICAM (UNIL V		CAM-UNEVG2QF83CG		LOXICAM	7.5 mg	
Inactive Ingredie						
		Ingredient Name			Strength	
MAGNESIUM STEAR		0				
SELICON DIO XIDE (U SODIEM CITRATE U						
Product Characte	ristics					
	yellow			no score		
Shape	gellow ROUND	Size		ton		
Shape Flavor			1+			
Shape Flavor Coutains	ROUND	Size Imprint Co-		Rmm Cytisk	ring End Da	
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Shaps Flavor Contains Packaging # Item Code I NDC 58 118-1158-9	Parks to a 1 BLISTER PACK	Size Imprint Co- inge Description	Marketing n Product 02/87/2018	Ross Cytis Cytis Start Date Marks	ting End Da	

 Extablishment
 Name
 Address
 IDSEI
 Business Operations

 Clinical for Infrare Withdraws, LLC
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