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. MELOXICAM tablets, for oral use initial U.S. Approval: 2000

See full prescribing information for complete board warning See full prescribing information for complete board warning cardiovascular thrombotic events, including myocardiol infarction and stroke, which can be tabl. This risk may occur early in treatment and may increase with duration Melosicam is controlled.

of use (5.1)

Misolocium E. Contraindicated in the setting of coronary artery bypass graft (CABG)

Misolocium E. Contraindicated in the setting of coronary artery bypass graft (CABG)

Misoloc cause an increased risk of serious gastrointestinal (II) adverse events

textualing biselenily, ulceration, and perioration of the science for interface, which

symptoms. Biserry patients and patients with a prior history of peptic ulcer disease
and/or of biselenily are at greater fact for serious (II events (E.2)).

DOSAGE AND ADMINISTRATION
the lowest effective dose for the shortest duration consistent with individual patient treatment goals

nce daily in children \geq 60 kg in tablets are not interchangeable with approved formulations of oral meloxicam even if the gram strength is the same (2.6)

DOSAGE FORMS AND STRENGTHS
 Meloxicam Tablets: 7.5 mg, 15 mg (3)

CONTRAINDICATIONS
 Nove hypersentiality to melanization or any components of the drug product (4)
 Hotory or admiss, uniticatic, or other allergic-type reactions after taking applies or other NSAIDs (4)
 In the setting of CABG surgery (4)

History of statima, utilizati, or other alleryli-ripe inactions that the late paper of other READs (4)

In the setting of Cold surpey (4)

**MARINGS AND PRECADINGS

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ADVERSE REACTIONS
 MOSt common (a5% and greater than piacebo) adverse events in adults are dammen, upper respratory tract infections, dysepapea, and infectionable symptoms (6.1).

Adverse events closered in products reade surface in nature to the adult clinical trial experience.

Advance events closered in products reade surface in nature to the adult clinical trial experience.

with relation instantions they are consistent of the consistent of

Who nave office our exercing toward.

See 17 for PATIENT COUNSELING INFORMATION and Medication Guide.

Revised: 8/2024

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 B.7 Recall Impair 13.1 Carchogenesis, Mutagenesis, Impartment u re may 14 CLINICAS TUDIO SER DEMONSTRATE A TUDIO SER DEMONSTRATE A TUDIO SER DEMONSTRATE A TUDIO SER DEMONSTRATE AND SER DEMONSTRATE AND SER DEMONSTRATE AND SER DEMONSTRATE AND SER DESCRIPTION SER DESCRIPTION

WARNING RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS

- Cardiovascular Thrombotic Events

 *Nonstrovidal unti-inflammatory drugs (15AUDs) cause an increased

 *Nonstrovidal unti-inflammatory drugs (15AUDs) cause an increased

 *Nonstrovidal unti-inflammatory drugs (15AUDs) cause an increased

 infarction and stroke, which can be fatal. This risk way occur early in

 treatment and may increase with duration of use [see Warnings and

 *Mebokican is contraindicated in the setting of cornorsy artery bypass

 graft (CABG) surgery [see Contraindications (4) and Warnings and

 *Prevaotives (5.4)

Precautions (5.1) 1. Southern and Perforation (4) and Warnings and Bastrointestinal Bleedina. Ukeration, and Perforation (5.1) 1. SIADIO scause an increased risk of serious gastrointestinal (GI) adverse events including bleeding, ukeration, and perforation of the stomach or intestines, which can be fatal. These events can occur at any time during use and without warning symptoms. Elderly patients bleeding are at greater risk for serious GI events [see Warnings and Precautions (5.2)].

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

Meloxicam is indicated for relief of the signs and symptoms of rheumatoid arthritis [see Clinical Studies (14.1)].

1.3 Juvenile Rheumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course

Meloxicam is indicated for relief of the signs and symptoms of pauciarticular or polyarticular course juvenile Rheumatoid Arthritis in patients who weighs \succeq 60 kg [see Dosage and Administration (2.4) and Clinical Studies (14.2)].

Carefully consider the potential benefits and risks of meloxicam and other treatment options before deciding to use meloxicam. Use the lowest effective dosage for the shortest duration consistent with individual patient treatment goals [see Warnings and Precautions (3)].

After observing the response to initial therapy with meloxicam, adjust the dose to suit an individual patient's needs.

In adults, the maximum recommended daily oral dose of meloxicam is 15 mg regardles of formulation. In patients with hemodalysis, a maximum daily dosage of 7.5 mg is recommended (see Use is Pspecific Populations (8.7) and Clinical Pharmacology (12.3). Meloxicam may be taken without regard to timing of meals.

For the relief of the signs and symptoms of osteoarthrills 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.

2.3 Rheumatoid Arthritis

For the relief of the signs and symptoms of rheumatoid arthritis, the recommended starting and maintenance oral dose of mebxicam is 7.5 mg once daily. Some patients may receive additional benefit by increasing the dose to 15 mg once daily.

2.4 Juvenile Rheumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course

For the treatment of juvenile rheumatoid arthritis, the recommended oral dose of mebxicam is 7.5 mg once daily in children who weigh ≈ 60 kg. There was no additional benefit demonstrated by increasing the dose above 7.5 mg in clinical trials.

Meloxicam tablets should not be used in children who weigh <60 kg.

2.5 Renal Impairment
The use of meloxicam in subjects with severe renal impairment is not recommended. In patients on hemodialysis, the maximum dosage of meloxicam is 7.5 mg per day [see Clinical Pharmacology (12.3)].

2.6 Non-Interchangeability with Other Formulations of Meloxicam

Who documents that graves were depulsed in systemic exposure to other approved formulations of oral mebixiam. Therefore, meloxicam tables are not interchanged with other formulations of oral mebixicam product even if the total miligram strength is the same. Do not substitute similar does strengths of meloxicam tables with other formulations of oral mebixicam product.

3 DOSAGE FORMS AND STRENGTHS

- Mebxicam tablets, USP:

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

 15 mg; yellow cobured, round, flist bevelled tablets, debossed with "CIPLA" on one side and "159" on the other.

- CUNINAINDICATIONS
 Mebixiam is contraindicated in the following patients:
 Known hypersexibityly (e.g., anaphytictic reactions and serious skin reactions) to
 Known hypersexibityly (e.g., anaphytictic reactions and serious skin reactions)
 (5.7.5.2)
 Hattory of asthma, uttaria, or other alergic-type reactions after taking apairs, uttaria, or other lessage of the state of coronary artery bypass graft (CABG) surgery (see Warnings and Precautions (5.7.5.8))
 In the setting of coronary artery bypass graft (CABG) surgery (see Warnings and Precautions (5.7.5.8))

5 WARNINGS AND PRECAUTIONS

5.1 Cardiovascular Thrombotic Events

Cinical trisk of several COX-2 selective and nonselective NSAIDs of up to three years duration have shown an increased risk of serious cardiovascular (CV) thrombotic authority of the control of

Post-MI Patients

Post-ML Distincts.

Observational studies conducted in the Danish National Registry have demonstrated that patients treated with Nations in the post-file proof were all nereased risk of reinflatction. Cerv-distated death, and all-case mortality beginning in the File week of treatment. In this years in NSAID-treated patients compared to 12 per 100 person years in non-MSAID years in NSAID-treated patients compared to 12 per 100 person years in non-MSAID with the increased residue rate of death exclude somewhat after the first year post-fill, the increased residue risk of death in NSAID users persisted over at least the next turn years of follow-up.

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

5.2 Gestrointestinal Bleeding, Ukeration, and Perforation
NSAIDs, richally melosicam, can cause serious gastrointestinal (i) subverse events
NSAIDs, richally melosicam, can cause serious gastrointestinal (ii) subverse events
small intestine or large intestine, which can be falsal. These serious adverse events can all intestine, or large intestine, which can be falsal. These serious adverse events on corruit any nitro, with or without warring symptoms, in patients treated with NSAIDs.
Only one in the patients who develop a serious upper Gil adverse event on NSAID
NSAIDs occurred in approximately 1% of patients treated of 7-3 6-months, and in about 2-4% or patients treated for one year. However, even short-term NSAID therapy is not without risk.

Risk Factors for GI Bleeding, Ulceration, and Perforation

Idea: Exicitys for La Islanding. Liveration. And Interformation of Disbeding who used NSAIDs Partiests with a prior history of peptic used releases and/or CI bleeding who used NSAIDs without these risk factors. Other factors that increase the risk of CII bleeding in palients without these risk factors. Other factors that increase the risk of CII bleeding in palients restead with NSAIDs include bringer duration of ISAID therapy concomitate use of oral conficustorials, aspirit, anticalgulants, or selective sendonin reuptake inhibitors or conficustorials, aspirit, anticalgulants, or selective sendonin reuptake inhibitors postmarketing reports and CII and CII

- risk for of bleeding.

 Strategies to Minimize the GI Risks in NSAID-treated patients:

 Use the bwest effective dosage for the shortest possible duration.

 Avoid administration of more than one NSAID at a time.

 Avoid use in patients at higher risk unless benefits are expected to outweigh the noneased risk of bleeding, for solve platferts, as well as those with active GI bleed consider alternate the applies other than NSAID consider alternate the propose of the number of the decreasion and bleeding during NSAID therapy.

- therapy.

 If a serious GI adverse event is suspected, promptly initiate evaluation and treatment, and discontinue meloxicam unit a serious GI adverse event is ruled out.

 In the setting of concomitant use of low-dose aspiring for cardiac prophytaxis, monitor patients more closely for evidence of GI bleeding [see Drug Interactions (7)].

Elevations of ALT or AST (three or more times the upper limit of normal [ULN]) have been reported in approximately 1% of NSAID-treated patients in clinical trials. In addition, rare, sometimes fatal, cases of severe hepatic injury, including fulminant hepatitis, liver necrosis, and hepatic failure have been reported.

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

Treated with IN-ALUS INCLUDINg metoccam. Inform patients of the warning signs and symptoms of hepatotoxicky (e.g., naused. Tudgue, lethnay, diurrhea, purvitus, jaundice, right upper quadrant frederieness, and the systemic manifestations occur (e.g., essionophila, rash, etc.). discontinue melboicam inmediately, and perform a cinical evaluation of the patient [see Use in Specific Populations (8.6) and Cinical Privamacology (22.3)].

5.4 Hypertension

NSAIDs, including meloxicam, can lead to new onset or worsening of preexisting hypertension, either of which may contribute to the increased incidence of CV events. Patients taking anjotensin converting earryme (AEC) inhibitors, thisacked duretics, or loop duretcs may have impaired response to these therapies when taking NSAIDs [see Drug Interactions (7)].

Monitor blood pressure (BP) during the initiation of NSAID treatment and throughout the course of therapy.

5.5 Heart Failure and Edema

The Coxils and traditional NSAID Trialsts' Collaboration meta-analysis of randomized controlled trials demonstrated an approximately two-fold increase in hospitalizations for compared to pick-observed read increases and the compared to pick-observed read increases. In a Danish Hatsonal Registry study of patients with heart failure, NSAID use increased the risk of MI, hospitalization for heart failure, and death.

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

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

5.6 Renal Toxicity and Hyperkalemia

Renal Toxicky
Long-term administration of NSAIDs, including meloxicam, has resulted in renal papillary
necrosis, renal insufficiency, acute renal failure, and other renal injury,
necrosis, renal insufficiency, acute renal failure, and other renal injury,
Renal toxickly has abo been seen in patients in whom renal postaglandins have a
compensatory role in the maintenance of renal perfusion. In these patients,
of the renal control of the renal renal renal renal renal renal renal renal decompensation. Patients at greatest risk of this reaction are those with imparted renal
decompensation. Patients at greatest risk of this reaction are those with imparted renal
function, depleytation, hypootenish, heart failure, bed relyturation, those tables,
is usually followed by recovery to the pretreatments associations of rend defourcing in
the renal effects of medius remains when the renarression of rend defourcing in

is usually followed by fectorely to the prescentific state: The renal effects of mebxicam 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)).

No information is available from controlled clinical studies regarding the use of mebuicam in patients with advanced renal disease. Avoid the use of mebuicam in patients with advanced renal disease, and the use of mebuicam in patients with advanced renal disease, must be benefits are expected to according to the patients of the pati

Increases in serum potassium concentration, including hyperkalemia, have reported with use of NSAIDs, even in some patients without renal impairme patients with normal renal function, these effects have been attributed to a hyporeninemic-hypoaldosteronism state.

New York Reactions
Meboxin has been associated with anaphylicitic reactions is patients with and without bloom hypersensitivity to metox can and in patients with aspirin-sensitive astimal [see Contrainations (of and Warnings and Precautions 6.09]. Sees indirections for an aphylicitic reaction occurs.

5.8 Exacerbation of Asthma Related to Aspirin Sensitivity

5.8 Exacerbation of Asthma Related to Aspirin Sensithity
A subopolation of patters with asthma may have appin-resible asthma which may include chronic rhinosinustic complicated by nasal polyps; severe, potentially fetal bronchospasm; and/or intellenance to apprin and other MSAIDs. Because cross-reactively between aspirin and other MSAIDs has been reported in such appin-records reaching the property of the pr

5.9 Serious Skin Reactions

3-3 serious Skin Relactions

ASIGN, including relevation, can clause serious skin alverse reactions such as

NESID, which can be fatal. NSAIDs can also cause fixed drug engition (FIGE). The may

present as a more severe variant known as generalized bullous fixed drug engition (FIGE). The may

constitute a more severe variant known as generalized bullous fixed drug engition

(GSTGE), which can be fit chreatering. These serious events may occur without and

constitute the constitution of the constitutio

5.10 Drug Reaction with Eosinophilia and Systemic Symptoms (DRESS)

Drug Reaction with Eosinophilia and Systemic Symptoms (DRESS)
Drug Reaction with Eosinophilia and Systemic Symptoms (DRESS) has been reported in
patients taking INSAIDS such as meloxicam. Some of these events have been fatal or Intreateming, DRESS pixely, although one of exclusively, presents with Feer, rail, we
repeated the control of the control of the control of the control of the control
repaids, nephrits, hematological abnormalities, myocarditis, or myocitis. Sometimes
symptoms of DRESS may resemble an active vali refection, sciencipilia is often present.
Because this disorder is variable in its presentation, other organ systems not noted here
such as fever of hymnidaeropathy, may be present each hough rails in face deviant. If
such signs or symptoms are present, discontinue meloxicam and evaluate the patient
immediately.

5.11 Fetal Toxicity

Premature Closure of Fetal Ductus Arteriosus

Avoid use of MSAIDs, Including metoxicam, in pregnant women at about 30 weeks
gestation and later NSAIDs, Including metoxicam, increase the risk of premature closure
of the fetal ductus arteriosus at approximately this gestational age.

Supernyaramios Menenatal Renal Impairment

Lie on NSAIDs, including metakeruan a about 20 weeks gestation or later in pregnancy may cause feat renal dysfunction leading to oligohydramnos and, in some cases, menenalat renal majement. These adverse outcomes are seen, on werage, after days to weeks of treatment, athough olgohydramnos base hen infrequently reported as soon as 48 hours after NSAID nistation. Olgohydramnos is often, but not always, remains own with treatment discontinuation. Complications of protonged olgohydramnos may, for access of majoral mental renal function, invalve procedures such as enchange transfusion or dialysis were required.

If NSAID treatment is necessary between about 20 weeks and 30 weeks gestation, limit mebxican use to the lowest effective dose and shortest duration possible. Consider utrasound monitoring of annibotic huild if mebxican treatment extends beyond 48 hours. Discontinue mebxican if olgohydramnios occurs and follow up according to clinical practice [see Use in Specific Populations [23].)

Anemia has occurred in NSAID-treated patients. This may be due to occult or gross blood biss, flut retention, or an incompletely described effect on erythropoles. If a hemoglobin or hematocirk.

NEMODIATION OF THE MADELER. MAY INCREASE THE FISK OF Bleeding events. Co-morbid conditions such as coaguitation disorders or concomitant use of warfarn, other anticoaguiatist, national such as capacity as continuous participation and propriet and participation and propriet and participation and propriet and participation and participa

5.13 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.

5.14 Laboratory Monitoring

Because serious GI bleeding, hepatotoxichy, and renal injury can occur without warnir symptoms or signs, consider monitoring patients on bng-term NSAID treatment with CBC and a chemistry profile periodically [see Warnings and Precautions (5.2, 5.3, 5.6)]

- 6 ADVERSE REACTIONS
 The following adverse reactions are discussed in greater detail in other sections of the biology.

 2017
 The control of th

6.1 Clinical Trials Experience

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to rates in the clinical trials of another drug and may not reflect the rates observed in practice. Adults

Osteoarthritis and Rheumatoid Arthritis

Osteoarthrits and Rheumatoid Arthritis

The meloxicam Phase 2/3 clinical trial disabase includes 10,122 OA patients and 1012 RA patients treated with meloxicam 7-3 mgdey, 3505 OA patients and 1351 RA patients graded with meloxicam 7-3 mgdey, 3505 OA patients and 1351 RA patients of the second patients for at least 6 months and to 312 patients for at least one year. Approximately 10,500 of these patients were treated in ten placebo- and/or active-controlled neumatoid arthritis trials. Gastrontestical (IGI) adverse events were the most frequently reported adverse events in all treatment groups across meloxicam trials.

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 mebxicam with placebo and with an active control. Two 12-week multicenter, double-blind, randomized trials were conducted in patients with rheumatoid arthritis to compare the efficacy and safety of mebxicam with placebo.

Table 1a depicts adverse events that occurred in ≥2% of the meloxicam treatment groups in a 12-week placebo- and active-controlled osteoarthritis trial.

Table 1b depicts adverse events that occurred in ≥2% of the meloxicam treatment groups in two 12-week placebo-controlled rheumatoid arthritis trials.

Table 1a Adverse Events (%) Occurring in ≥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 Perip	heral			
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				
Rash ²	2.5	2.6	0.6	2.0

rred terms rash, rash erythematous, and rash maculo-papular combined

	Placebo	Meloxicam	Meloxicam
		7.5 mg	15 mg
		daily	daily
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 Cond	litions	
Influenza-like illness ²	2.1	2.9	2.3
Infection and Infestations			
Upper respiratory tract infections-pathogen class unspecified ¹	4.1	7.0	6.5
Musculoskeletal and Connective			
Tissue Disorders			
Joint 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): dysp dyspepsia aggravated, eructation, gastrointest infections-pathogen unspecified (laryngkis NOS related signs and symptoms (arthralgia, arthralg	nal irritation , pharyngiti	, upper respir NOS, sinusiti	atory tract s NOS), joint

""", upper teapfattory tract
related signs and synthetions (attribute), anthralgia aggravated, joint crepitation, joint
elated signs and synthetions (attribute), arthralgia aggravated, joint crepitation, joint
elated, joint swelling)

Most Ag preserved term nausea, abdominal pain NOS, influenza-like illness, headaches
NOS, and rash NOS

The adverse events that occurred with meloxicam in $\ge 2\%$ of patients treated short-term (a to 6 weeks) and long-term (6 months) in active-controlled osteoarthrifs trials are presented in Table 2 Adverse Events (%) Occurring in $\ge 2\%$ of Meloxicam Patients in 4 to 6 Weeks and 6 Month Active-Controlled Osteoarthrifs Trials

	4 to 6 Weeks	Controlled Trials	6 Month	Controlled Trials
	Meloxicam 7.5 mg daily	Meloxicam 15 mg daily	Meloxicam 7.5 mg daily	Meloxicam 15 mg daily
No. of Patients	8955	256	169	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
Peripheral Nervous System Dizziness	1.1	1.6	2.4	2.6
Headache	2.4	2.7	3.6	2.6
Hematologic				
Anemia	0.1	0.0	4.1	2.9
Musculoskeletal	1	1	1	
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		1		
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	13
Urinary				
	0.1	0.4	2.4	1.3
Micturition frequency				

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

Pediatrizs

Pauciarticular and Polyarticular Course Juvenile Rheumatoid Arthritis (JRA)

Pauciatricular and Polyarticular Course Iuvenile Rheumatoid Arthrisk (IRBA)
Three hundred and egilty-seven paletest with pauciatricular and polyarticular course JRA
were exposed to meloxicam with doses ranging from 0.12 to 0.373 mg/kg per day in
tree clinical trials. These studies consisted of two 12-week multzeturer, double blind,
randomized trials (one with a 12-week open-label extension and one with a 40-week
pediatric studies with meloxicam were similar in nature to the adult chical tributes
pediatric studies with meloxicam were similar in nature to the adult chical tributes
experience, although there were differences in frequency. In particular, the following
most common adverse events, adomaing Jain, vonthing, darthes, headdent, and
in seveni c (2%) paletest receiving meloxicam. No unexpected adverse events directly
in sevenic consistency of the course of the tribute. The adverse events did not demonstrate an age
or gender-specific subgroup effect.

The following is a last of adverse adrug reactions occurring in <2% of patients receiving
meloxicam in clinical trials involving approximately 16,200 patients.

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, vascultis
Central and Peripheral Nervous	System convulsions, paresthesia, tremor, vertigo
Gastrointestinal	colitis, dry mouth, duodenal ulcer, eructation, esophagitis, gastric ulcer, gastritis, gastroesophageal reflux, gastrointestinal hemorrhage, hematemesis, 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	leukopenia, 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, prunitus, sweating increased, urticaria
Special Senses	abnormal vision, conjunctivitis, taste perversion, tinnitus
Urinary System	altuminuria RIIN increased creatinine increased hematuria renal failure

6.2 Postmarketing Experience

6.2 Postmarketing Experience
The following abuses reactions have been identified during post approval use of melanciam. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to relably estimate their frequency or establish a causal relationship to drug exposure. Decisions about whether to include an adverse event from spontaneous reports in labeling are typically based on one or more of the following factors: (1) sertusiness of the event. (2) number of reports, or (3) streight of following factors: (1) sertusiness of the event. (2) number of reports, or (3) streight of following factors: (1) sertusiness of the event. (2) number of reports, or (3) streight of following factors: (1) sertusiness of the event. (2) number of reports, or (3) streight of following factors: (1) sertusiness of the event. (2) number of reports, or (3) streight of following factors: (1) sertusiness of the event. (2) number of reports, or (3) streight of following factors: (1) sertusiness of the event. (2) number of reports, or (3) streight of following factors: (1) sertusiness of the event. (2) number of reports, or (3) streight of following factors: (1) sertusiness of the event. (2) number of reports, or (3) streight of following factors: (3) streight of factors: (3) streight of factors: (3) streight of factors: (3) streight of factors: (4) number of reports or (4) streight of factors: (4) number of reports or (4) number or (4) number of reports or (4) number of reports or (4) number of reports or (4) num

7 DRUG INTERACTIONS

7 DRUG IN TERACLIONA See Table 3 for cinically significant drug interactions with meloxicam. See also Warnings and Precautions (5.2, 5.6, 5.12) and Clinical Pharmacology (12.3).

Table 3 Clinically Significant Drug Interactions with Meloxicam

Orugs that Ir	nterfere with Hemostasis
Elinical Impact:	Mebxicam and anticoaguiants such as warfarin have a synergistic effect on bleeding. The concomitant use of mebxicam and anticoagulants have an increased risk of serious bleeding compared to the use of either drug abne. Serotonin release by platelets plays an important role in hemostasis. Case-control and cohort exidemiological studies
	showed that concomitant use of drugs that interfere with serotonin reuptake and an NSAID may potentiate the risk of bleeding more than an NSAID alone.
ntervention:	Monitor patients with concomitant use of meloxicam with anticoagulants (e.g., warfarin), antiplatelet agents (e.g., aspirin), selective serotonin reuptake inhibitors (SSRIs), and serotonin horepinephrine reuptake inhibitors (SNRIs) for signs of bleeding see Warnings and Precautions (S.12).
Spirin	
	Controlled cinical studies showed that the concomitant use of NSAIDs and analgesic doses of aspirin does not produce any greater therapeutic effect than the use of NSAIDs alone. In a clinical study, the concomitant use of an NSAID and aspirin was associated with a significantly increased incidence of GI adverse reactions as compared to use of the NSAID alone [see Warnings and Precautions (5.2)].
ntervention:	Concomitant use of meboxicam and low dose aspirin or analyseis doses of aspirin is not generally recommended because of the increased risk of bleeding (see Warnings and Precautions (5.12) Mebxicam is not a substitute for low dose aspirin for kardiovascular protection.
	s, Angiotensin Receptor Blockers, or Beta-Blockers
ilinical Impact:	NSAIDs may diminish the anthypertensive effect of anjotensin converting enzyme (ACE) inhibtors, anjotensin propranolo). In patients were dederly, volume-depleted (including those on diurest the trappy, or have renal impariment, coadministration of an NSAID with ACE inhibitors or ARBs may result in deterioration of renal function. Including oossible acute
	renal failure. These effects are usually reversible.
ntervention:	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 concomitant use of meloxicam and ACE inhibitors
	or AR8s in patients who are elderly, volume-depleted, or have mpaired renal function, monitor for signs of worsening renal function [see Warnings and Precautions (5-6)]. When these drugs are administered concomitantly, patients should be adequately hydrated. Assess renal function at the beginning of the concomitant treatment and periodically thereafter.
Diuretics	processes.
linical Impact:	Clinical studies, as well as post-marketing observations, showed that NSAIDs reduced the natriuretic effect of loop diuretics (e.g., furosemide) and thiazide diuretics in some patients. This effect

	has been attributed to the NSAID inhibition of renal prostaglandin synthesis. However, studies with furosemide agents and metoxicam have not demonstrated a reduction in natriured single and multiple does planmacolynamics and pharmacolynamics and manacolynamics and pharmacolynamics are not affected by multiple doses of metoxicam.
Intervention:	During concomitant use of meloxicam with diuretics, observe patients for signs of worsening renal function, in addition to assuring diuretic efficacy including anthypertensive effects [see Warnings and Precautions (5.6)].
Lithium	
Clinical Impact:	NSAIDs have produced elevations in plasma Ithium levels and reductions in renal lithium clearance. The mean minimum Ithium concentration increased 15%, and the renal clearance decreased by approximately 20%. This effect has been attributed to NSAID nihblion of renal prostaglandin synthesis [see Clinical Pharmacology (12.3)].
Intervention:	During concomitant use of meloxicam and lithium, monitor patients for signs of lithium toxicity.
Methotrexat	ė
Clinical Impact:	Concomitant use of NSAIDs and methotrexate may increase the risk for methotrexate toxicity (e.g., neutropenia, thrombocytopenia, renal dysfunction).
Intervention:	During concomitant use of meloxicam and methotrexate, monitor patients for methotrexate toxicity.
Cyclosporine	
	Concomitant use of meloxicam and cyclosporine may increase cyclosporine's nephrotoxicity.
Intervention:	During concomitant use of meloxicam and cyclosporine, monitor patients for signs of worsening renal function.
NSAIDs and !	Salicylates
Clinical Impact:	Concomitant use of meloxicam with other NSAIDs or salicylates (e.g., diffundal, salsalate) increases the risk of GI toxicity, with lttle or no increase in efficacy [see Warnings and Precautions (5.2)].
Intervention:	The concomitant use of meloxicam with other NSAIDs or salicylates is not recommended.
Pemetrexed	
Clinical Impact:	Concomitant use of meloxicam and pemetrexed may increase the risk of pemetrexed-associated myelosuppression, renal, and GI toxicity (see the pemetrexed prescribing information).
Intervention:	During concomitant use of meloxicam and pemetrexed, in patients with real impairment whose creatinine clearance ranges from 45 to 79 mL/min, mention for myelosuppression, real and Patients taking meloxicam should interrupt doshing for at least five days before, the day of, and two days following pemetrexal darministration. In patients with real many continuous in patients with creatinine clearance below 45 mL/min, the in patients with creating continuous many continuous procommended.

A Deepandy

RBS Summary

We of MSADIs, including metaskism, can cause premature closure of the fetal ductus
arterious and refail remail dysfunction leading to objoying armics and, in some cases,
menonatal remail implement. Because of these resis, limit does and duration of metaskism
use between about 20 and 30 weeks of gestation, and awoit metoxicam use at about 30
weeks of gestation and later in pregnancy (see Circlar Constructions, Data).

Premature Closure of Fetal Ductus Arteriosus

Oligohydramnios/Neonatal Renal Impairment

Use of NSAIDs at about 20 weeks gestation or later in pregnancy has been associated with cases of fetal renal dysfunction leading to oligohydramnios, and in some cases, neonatal renal impairment.

Data from observational studies regarding potential embryofetal risks of NSAID use in women in the first or second trimesters of pregnancy are inconclusive.

total roll disservational studies regarding ploteful entrolysteric for the store of such as women in the first of second invasions of pregnancy precised roles on the store of the second studies of t

The estimated background risk of major birth defects and miscarriage for the indicated population(s) is unknown. All pregnancies have a background risk of birth defect, loss, or other adverse outcomes. In the U.S. general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnancies 8.2% to 4% and 15% to 20%, respectively.

Avoid use of NSAIDs in women at about 30 weeks gestation and later in pregnancy, because NSAIDs, including meloxicam, can cause premature closure of the fetal ductus arteriosus (see Data).

Oligohydramnios/Neonatal Renal Impairment:

If an NSAID is necessary at about 20 weeks gestation or later in pregnancy, limit the use to the lowest effective dose and shortest duration possible. If meloxican treatment extended beyond 46 bours, consider montrion with utraxound for oligohydramios. If oligohydramios occurs, discontinue meloxicam and follow up according to clinical Labor or Delwey.

<u>Data</u> Human Data

Premature Closure of Fetal Ductus Arteriosus

Premature Closure of Fedial Ductus Arteriosus:

Published Renture reports that the use of HSAIDs at about 30 weeks of gestation and later in pregnancy may cause premature closure of the fedial ductus arteriosus.

Oliphoydraminos/Honotatal Renal Impramment:

Published studies and postmarketing reports describe maternal NSAID use at about 20 weeks operation or hater in pregnancy ascorded with Fedial renal alpharuction leading to outcomes are seen, on average, after days to weeks of treatment, although outports are seen, on average, after days to weeks of treatment, although outports are seen, on average, after days to weeks of treatment, although outports are seen and although of the properties are seen and although of the contract of the properties are seen and although of the contract of the properties are seen and although of the contract of the properties are seen after the properties are considered and although of the properties are seen after the properties are considered as the properties are seen after the properties are considered as the properties are seen after the properties are considered as the properties are seen after the properties are considered as the properties are seen after the properties are considered as the properties are seen as the properties ar

of which were irreversible. Some cases of neonatal renal dysfunction required treatment with the control of the

urrougnous organogeness.

Oral administration of meloxicam to pregnant rats during late gestation through lactation increased the incidence of dystocia, delayed parturition, and decreased offspring survival at meloxicam doses of 0.125 mg/kg/day or greater (0.08-times MRHD based on BSA comparison).

8.2 Lactation

8.2 Lactaton

Risk Summay

There are no human data available on whether meloxicam is present in human mills, 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 cincial need for
meloxicam and my optential adverse effects on the breastfed infant from the
meloxicam and my optential adverse effects on the breastfed infant from the

Data

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

8.3 Females and Males of Reproductive Potential

8.3 Females and Males of Reproductive Potential infettility.

Forales

Based on the mechanism of action, the use of prostagalanth-mediated NSAIDs, including mebxicam, may delay or prevent rupture of ovarian folicies, which has been associated with reversible infettility is nome women. Published animal studies have shown that administration of prostagalantial synthesis inhibitors have those prostagalantial administration of prostagalantial synthesis inhibitors have the prosterior of subject the prostagalantial synthesis inhibitors have the prostagalantial administration of prostagalantial synthesis inhibitors have the prostagalantial administration of prostagalantial prostagalantial synthesis and of NSAIDs, including mediators, in women who have difficulties conceiving or who are undergoing investigation of Interlity.

8.4 Pediatric Use

The safety and effectiveness of meioxicam 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

Elderly patients, compared to younger patients, are at greater risk for NSAID-associated serious cardiovascular, gastrointestinal, and/or renal adverse reactions. If the

anticipated benefit for the elderly patient outweighs these potential risks, start dosing at the low end of the dosing range, and monitor patients for adverse effects [see Warnings and Precautions $(5.1,\,5.2,\,5.3,\,5.6,\,5.14)$].

8.6 Hepatic Impairment
No dose adjustment is necessary labelers with mild to moderate hepatic impairment
Patients with sent hepatic impairment have not been adequately studied. Since melosition with caution is patients with sent from the patient in the patient is made to the patient in the patient is made to the patient with caution in patients with hepatic impairment [see Warnings and procedures (1.3) and Childel Pharmacology (1.2.3)].

8.7 Renal Impairment

No dose adjustment is necessary in patients with mild to moderate renal impairment. Patients with severe renal impairment have not been studied. The use of most office subjects with severe renal impairment is not recommended. In patients on hemodalysis meloxicam should not exceed 7.5 mg per day. Meloxicam is not dialyzable [see Dosage and Administration (2.1) and Clinical Pharmacology (2.3)].

10 OVERDOSAGE
Symptoms flolwing acute NSAID overdosages have been typically limited to lethargy, drowstess, naueses, contribing, and egigastric pair, which have been generally reversible refairs respiratory depression, and come have occurred, but were rare [see Warnings and Precaudios 15, 13, 25, 45, 45].

and Preculations (5.1, 5.2, 5.4, 5.6).

Manage patients with symptomats and supportive care following an ISADI overdosage. There are no specific antisother. Consider emests and/or activated charcoal (60 to 100 grams in adults, 11 to 2 grams per lay of body weight in pediating taletties) and/or cosmotic calhartic in symptomatic patients seen within four hours of ingestion or in patients with a large everdosage (5 to 10 times the recommended dosage); Forced high protein bridge of control in the commended dosage) forced high protein bridge.

There is limited experience with meloxicam overdosage. Cholestyramine is known to accelerate the clearance of meloxicam. Accelerated removal of meloxicam by 4 g oral doses of cholestyramine given three times a day was demonstrated in a clinical trial. Administration of cholestyramine may be useful following an overdosage.

For additional information about overdosage treatment, call a poison control center (1-800-222-1222).

Mebickam is a nonsteroidal and-inflammatory drug (MSAID). Each tablet contains 7.5 mg or 15 mg mebickam, USP for oral administration, Mebickam is chemically designated as 4-hydroxy-2-mebly-4-fr-ethips-4-taboxyl-2-fr-12-brevolbiase-in-3-carbonalide-1,3-doxxde. The molecular weight is 331.4. Its empirical formula is C1₂+1₃N₃O45₂ and it has the following surfured formula.

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 (log P) $_{\rm app}=0.1$ in n-octanol/buffer pH 7.4. Meloxicam has pKa 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, bictose anhydrous, colbidal silicon dioxide, sodium citrate dihydrate, magnesium stearate.

12.1 Mechanism of Action

Mebxicam has analysisk, anti-riflammatory, and antipyretic properties.

The mechanism of action of mebxicam, like that of other NSAIDs, is not completely understood but involves inhibition of cyclooxygenase (COX-1 and COX-2).

Medioxicam is a potent inhibitor of prostaglandin synthesis in vitro. Medioxicam is a potent prostaglandin synthesis in vitro. Medioxicam is a potent prostaglandin synthesis in vitro. Medioxicamisjandinis sensitize afferent nerves and potentiate the action of bradykinin in inducing pain in animal models. Prostaglandins are readitors of inflammation. Because medioxicam is an inhibitor of prostaglandin synthesis, its mode of action may be due to a decrease of prostaglandin synthesis. Its mode of action may be due to a decrease of prostaglandin synthesis.

12.3 Pha

Absorption

Absorption

The absolute blowalshifty of metoxicam capsules was 89% following a single oral dose
of 30 mg compared with 30 mg fV bobus injection. Following single intravenous doses,
obese proportional pharmacolinetics were shown in the range of 5 mg to 60 mg, or
over the range of 7.5 mg to 3.5 mg. Mean Cras, was achieved within four to five hours
after a 7.5 mg mediscent mablet was then under fasted conditions, ridicating a
prolonged drug absorption. With multiple dossing, steady-state concentrations were
reached by 10 ps. 3.4 econd meboxicam concentration peak occurs around 12 to 14
hours
of the consideration of the second conditions of the conditions

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

Pharmacokinetic	Steady Stat	:e		Single Dose		
Parameters (%CV)	Healthy male adults (Fed) ²	Elderly males (Fed) ²	Elderly females (Fed) ²	Renal failure (Fasted)	Hepatic insufficiency (Fasted)	
	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)	
Tmax [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-#4 [1]	14.7 (32)	15 (42)	10 (30)	26 (44)	14 (20)	

¹ The parameter values in the table are from various studies

2 not under high fat conditions 3 Medicar an advantage of the property of the

Distribution

The statement of distribution 1953 of melocican is approximately 3D. Lebeusize in 18 4% Should to human plasma proteins (promary albumin) with in the threspectal color range. The fraction of protein briding is independent of drug concentration, over the chickally relevant concentration range, but cercases to 19-95 in patients with resul and 10% following a radioableed dose, over 90% of the radioactivity detected in the plasma was present as unchanged melocican.

Medician concentrations in symbol fluid, after a single oral dose, range from 40% to 50% of those in plasma. The free fraction is symbol fluid is 2.5 times higher than in plasma, due to the bewer abumin content in symovial fluid as compared to plasma. The significance of this penetration is unknown.

Elimination Metabolism

Medicanian extractively metabolised in the New Seldvicem metabolises include 5: Medicanian skicem file of death in row 8-50 melidated metabolism formed by oxidation of an intermediate metabolise 5-hydroxymethyl medicaran which is also excreted to a lessor exercit (9% of door, in vivo studies indicate that CP2CO (cytochrome PASO metabolismy enzyme) plays an important role in this metabolic system of the properties of t

Excretion

Excretion

Meboxiam excretion is predominantly in the form of metabolites, and occurs to equal setters in the urine and feets. Only faxes of the unchanged parent compound are extensively as the second of the control of the control

The mean elimination half-life $(t_{1/2})$ ranges from 15 hours to 20 hours. The elimination half-life is constant across dose levels indicating linear metabolism within the therapeutic dose range. Plasma clearance ranges from 7 to 9 mLlms. Specific Populations

Pediatric

Pediatrix
After single (0.25 mg/kg) dose administration and after achieving steady state (0.375 mg/kg)dg/k), there was a general trend of approximately 30% lower exposure in younge patients (2 to 8 years old is compared to the older patients (7 to 16 years old). The state of the older patients of the state old is compared to the older patients of the state of those in the adult patients, when using All C values normalized to a dose of 1.05 mg/kg (see Dosage and Administration (2.4)). The medicate mener (50) elimination half-life was 13.2 (10.3) and 13.0 hours (3.0) for the 2 to 6 year old patients, and 7 to 16 year old patients, respectively.

In a covariate analysis, utilizing population pharmacokinetics body-weight, but not age, was the single predictive covariate for differences in the meloxicam apparent oral plasma

² not under high fat conditions

clearance. The body-weight normalized 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

The pharmacokine been investigated

cuerant.

Belly missel (165) years of page onlibbed resolvant plants concentrations and the page of th

Sex Young formales exhibited slightly lower glasma concentrations relative to young males. After single doses of 7.5 mg nebox can, the mean elimination half all was 19.5 hours for the fielding rouge as compared to 2.3 hours for the mide group. At steady state, the data were similar (1.79 hours vs 21.4 hours). This pharmacokinetic difference due to gender is likely to be of title chical importance. There was intenty of pharmacokinetics and no appreciable difference in the C_{max} or T_{max} across genders.

Hepatic Impairment

Following a single 1.5 mg dose of mebxicam there was no marked difference in plasm concentrations in patients with mild (Chid-Pught Class I) or moderate (Chid-Pught Chid-Pught Ch

Renal Impairment

Renal Impairment
Mebiciscan pharmacokinetics have been investigated in subjects with mild and moderalt renal impairment. Total drup plasma concentrations of mebiciscan decreased and total clearance of mebiciscan increased with the degree of renal impairment while free clearance of mebiciscan increased with the degree of renal impairment subjects with renal covariance of the contraction of t

Hemodialysis

Following a single dose of meloxicam, the free C_{max} plasma concentrations were higher in patients with resul failer on chronic hemodalysis (15) free fraction) in comparison to concentration in plasma; the officer additional doses are not necessary after hemodalysis. Neloxicam for not dislyzable [see Dosage and Administration (2,1) and Use in Specif Populations (8,27)].

Drug Interaction Studies

academ: When MSAIDs were administred with agains, the protein binding of MSAIDs were described about the care of free MSAID was not aftered. When makes a deministered with against (1000 mg three times daily) to healthy volunteers. It tended to increase the AUC (10%) and C_{MS}. (24%) of mexicx.n. The clinical significance of this interaction is not known. See Table 3 for clinically significant drug interactions of NSAIDs with apprin [see Organ [seed as a contract of the cont

was span see God grandsound (2).

Choistsyramine Pretreatment for four days with choistsyramine significantly increased the clearance of mebukam by 50%. This resulted in a decrease in 1/2, from 19.2 hours to 125 hours, and a 35% reduction in AUC. This suggests the existence of recirculation and a 55% reduction in AUC. This suggests the existence of the interactions halvesy for mebukam in the gistrontestinal tract. The chical relevance of the interactions have not been established.

Cimetidine: Concomitant administration of 200 mg cimetidine four times daily did not alter the single-dose pharmacokinetics of 30 mg meloxicam.

Digorie: Meloxicam 15 mg once daily for 7 days; did not alter the plasma concentration profile of digoria nater §-acetyldigoxin administration for 7 days at clinical doses. In vitro stress figurant on protein binding drug in interaction between digoxin and meloxicam.

Lithium: In a study conducted in healthy subjects, mean pre-dose Bhamic oncentration and ALC were becreased by 21% in subjects receiving final modes ranging from 804 to receiving thium alone [see Drug Interactions 2D].

Methorozeada & shady in 3.1 Remainded airrish [Rin] plastics evaluated the effects of multiple doses of meloxicam on the pharmacokinetics of methodrozeada shad on the ave a significant effect on the pharmacokinetics of single doses of methodrozeada shad on the ave a significant effect on the pharmacokinetics of single doses of methodrozeada shad on the ave a significant effect on the pharmacokinetics of single doses of methodrozeada shad on the average and the significant effect on the pharmacokinetics of single doses of methodrozeada shad on the average and the significant effect on the pharmacokinetics of single doses of methodrozeada shad on the depth of the significant effect on the pharmacokinetic of single doses of methodrozeada shad on the depth of the significant effect on the pharmacokinetic of single doses of methodrozeada shad on the significant effect on the pharmacokinetic of single doses of methodrozeada shad on the significant effect on the pharmacokinetic of single doses of methodrozeada shad on the significant effect on the significant effect on the pharmacokinetic of single doses of methodrozeada shad on the significant effect on the pharmacokinetic of single doses of methodrozeada shad on the significant effect on the significant effect on the significant effect on the pharmacokinetic of single doses of methodrozeada shad on the significant effect on the signi

human serum binding sites [see *Drug Interactions (2*]].

Warfair: The effect of meloxicam on the anticoagulant effect of warfair was studied in a group of healthy subjects receiving daily doses of warfair in their produced an INR international Normalized Ratio between 12 and 13.8 in these subjects, meloxicam did international Normalized Ratio between 12 and 13.8 in these subjects, molecular did not the subject of the subject in the subj

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

<u>Carcinogenesis</u>

Net Explorations.

There was no increase in tumor incidence in long-term carcinogenichy studies in rats (104 weeks) and mice (99 weeks) administered meloxicam at oral doses up to 0.8 mg/gdgyl ar sand up to 8.0 mg/gdgyl ar mice (up to 0.5- and 2.6-times, respectively, the maximum recommended human dose (MRHD) of 15 mg/day meloxicam based on body surface area (1654) (cromparison).

based on body surface area [BSA] comparison).

Midiagnesis.

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 marrows.

Inneal ment of Testility.

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).

14 CLINICAL STUDIES

14.1 Osteoarthriks and Rheumatoid Arthriks
The use of indexican for the treatment of the signs and symptoms of osteoarthriks of
12.75 mg. 7.3 mg and 15 mg daily was compared to place. The four primary
endpoints were investigator's global assessment, patient policy
endpoints were investigatory and the global assessment, patient policy
endpoints and the global assessment patient patients
endpoints and the global assessment patients
endpoints compared with patients
endpoints compared with patients.

placebo.

The use of meloxicam for the management of signs and symptoms of osteoarthrists was evaluated in six double-bind, active-controlled trials outside the U.S. ranging from 4 weeks to 6 months of water for meloxicam, in doses of 7.5 months of the controlled trials outside the U.S. ranging from 4 weeks to 6 months of water for use from the meloxicam, in doses of 7.5 mogidity and consistent with the efficacy seen in the U.S. trial. This was remarked and consistent with the efficacy seen in the U.S. trial. The use of meloxicam for the treatment of the signs and symptoms of freemental arthrist was evaluated in a 12-week, double-blind, controlled multinational trial. Meloxicam (7.5 mg.) 15 mg. and 2.5 mg days laws compared to placedo. The primary with the properties of the properties of the primary with the properties of the properties of the primary with paticols. No incremental benefit was observed with the 2.5 mg dose compared to the 15 mg dose.

14.2 Juvenile Rheumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course

14.2 juvenile Rheumatold Arthrist (RIA) Auculetricular and Polyarticular Course
Course

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16 HOW SUPPLIED/STORAGE AND HANDLING

Mebxicam tablets, USP 7.5 mg are yellow coloured, round, biconvex tablets, debossed wth "158" on one side and "C" on the other.

Mebxicam tablets, USP 15 mg are yellow coloured, round, flat bevelled tablets, debossed wth "CIPLA" on one side and "159" on the other.

Meloxicam tablets, USP 7.5 mg are available as follows: NDC 69097-158-07

Bottles of 100 Bottles of 500 Bottles of 100

Store at 20°C to 25°C (68°F 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.

17 PATIENT COUNSELING INFORMATION

Advise the patient to read the FDA-approved patient labeling (Medication Guide) that accompanies each prescription dispensed.

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

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 Warnia and Precautions (5_1)].

Gastrointestinal Bleeding, Ulceration, and Perforation

Advise patients to report symptoms of ulcerations and bleeding, including epigastric pain, dyspepsia, mee'nd, and hematemes is to their healthcare provider. In the setting of concombant use of or-wides eapir for cardiac prophysixs, inform patients of the increased risk for the signs and symptoms of GI bleeding [see Warnings and Precautions (2,2)].

Hepatotoxicity

Inform patients of the warning signs and symptoms of hepatotoxicity (e.g., nausea, fatigue, lethargy, diarrhea, pruritus, jaundice, right upper quadrant tenderness, and "flu-like" symptoms. If these occur, instruct patients to stop meloxicam and seek immediate medical therapy [see Warnings and Precautions (5.3)].

Heart Failure and Edema

Advise patients to be alert for the symptoms of congestive heart failure including shortness of breath, unexplained weight gain, or edema and to contact their healthcare provider if such symptoms occur [see Warnings and Precautions (5,5)].

Anaphylactic Reactions

Inform patients of the signs of an anaphylactic reaction (e.g., difficulty breathing, swelling of the face or throat). Instruct patients to seek immediate emergency help if these occur [see Contraindications (4) and Warnings and Precautions (5.ZI)].

Serious Skin Reactions, including DRESS

Advise patients to stop taking meloxicam immediately if they develop any type of rash or fever and to contact their healthcare provider as soon as possible [see Warnings and Precautions (5.9), 5.10)].

Advise females of reproductive potential who desire pregnancy that NSAIDs, including metaxicam, may be associated with a reversible delay in ovulation [see Use in Specific Populations (8,3)].

Female Fertility Fetal Toxicity

The second was the second of the side of medical and other ISAID; starting at 30 miles (pagination because of the side of the premature chaign of the field discuss anterious). If treatment with melaxicam is needed for a pregnant woman between about 30 to 30 weeks gestation, advise her that she may need to be monitored for oligibility dimnios, if treatment continues for longer than 48 hours [see Warnings and Precautions(2,1)] and size in Specific Populations (23), and size in Specific Populations (23), and size in Specific Populations (23), and size in Specific Populations (24).

Avoid Concomitant Use of NSAIDs

ANNA LANGEMENT USE OF TRANSIS.

Inform patients that the concommant use of mebosicam with other NSAIDs or salkylates (e.g., diflunisal, salkslatel is not recommended due to the increased risk of quastrointestinal toxicity, and title or no horses in efficiency (see Warnings and Precautions (52) and Drug interactions (71). Alert patients that NSAIDs may be present in over the counter" medications for treatment of colds, feee, or insommit

Use of NSAIDs and Low-Dose Aspirin

Cipla USA, Inc.

10 Independence Boulevard, Suite 300,

Warren. NI 07059

Revised: 08/2024

Medication Guide for Nonsteroidal Anti-Inflammatory Drugs (NSAIDs) What is the most important information I should know about medicines called Nonsteroidal Anti-Inflammatory Drugs (NSAIDs)?

NSAIDs can cause serious side effects, including: Increased risk of a heart attack or stroke that can lead to death . This risk may happen early in treatment and may increase: • with increasing dose of NSAIDs • with oney use of NSAIDs

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

ortery bypass graft (CABG)." Avoid taking NSAIDs after a recent heart attack, unless your healthcare provider tells you to You may have an increased risk of another heart attack if you take MSAIDs after a recent heart attack. If you take MSAIDs after a recent heart attack. If you take MSAIDs after a recent heart attack. If you take MSAIDs after a recent heart attack. If you take MSAIDs after a recent heart attack. If you take MSAIDs after a recent heart attack. If you take MSAIDs after a recent heart attack. If you take MSAIDs after a recent heart attack. If you have MSAIDs after a recent heart attack. If yo

The risk of getting an ulcer or bleeding increases with:

- past history of stomach uicres, or stomach or intestinal bleeding with use of NSAIDs o older age
 lating medicines called 'conflosteroids', "anticoagulants", "SSRIs", or "SNRIs" opon health
 poor health
 longer use of NSAIDs
 smoking
 dinking also of NSAIDs

NSAIDs should only be used:
 exactly as prescribed
 at the lowest dose possible for your treatment
 for the shortest time needed What are NSAIDs?

What are NoAlDs:

NSAIDs are used to treat pain and redness, swelling, and heat (inflammation) from medical conditions such as different types of arthritis, menstrual cramps, and other types of short-term pain.

Who should not take NSAIDs?

- wino should not case insalius?

 If you have had an asthma attack, hives, or other allergic reaction with aspirin or any other NSAIDs.

 if you have had an esthma attack, hives, or other allergic reaction with aspirin or any other NSAIDs.

 right before or after heart bypass surgery.

- right before or after heart bypass surgery.

 Before taking NSAIDS, tell your healthcare provider about all of your medical conditions, including if you:
 have high blood pressure
 have high blood pressure
 have high blood pressure
 have high blood pressure
 have assume
 have assume
 many pains to become pregnant. Taking NSAIDs at about 20 weeks of
 pregnancy or later may harm your unborn baby. If you need to take ISAIDs for
 prangency or later may harm your unborn baby. If you need to take ISAIDs for
 practice that 20 years when you are between 20 and 30 weeks of pregnancy, your
 healthcare provider may need to monitor the amount of fluid in your womb around
 you have. You should not take NSAIDs after about 30 weeks of
 are breastfeeding or plan to breast feed.

Tell your healthcare provider about all of the medicines you take, including prescription or over-the-counter medicines, vitamins or herbal supplements. NSAIDs and some other medicines can interact with each other and cause serious side effects. Do not start taking any new medicine without talking to your healthcare provider first.

What are the possible side effects of NSAIDs? NSAIDs can cause serious side effects, including

See "What is the most important information I should know about medicines called Nonsteroidal Anti-inflammatory Drugs (NSAIDs)?"

- called Nonsteroidal Anti-friammatory Drugs (NSAIDs)?*

 New or worse high blood pressure

 heat if salze

 but red blood cells (ralemis)

 but red blood cell

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

- shortness of breath or trouble breathing
 chest pain
 weakness in one part or side of your body
 swelling of the face or throat

- Tausce
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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 pharmacist about NSAIDs.

Call your doctor for medical advice about side effects. You may report side effects to FDA at 1:800-FDA-1888.

- FDA at 1-800-FDA-1088.

 Other information about NSAIDs:

 Aspir's is an InSAID but it does not increase the chance of a heart attack. Aspir's in a third interference of a heart attack. Aspir's in the stomach and intestines.

 Some INSAIDs are sold in lower doese without a prescription lower-the-countert. Talk to your healthcare provider before using over-the-counter NSAIDs for more than 10 days.

General information about the safe and effective use of NSAIDs

Medicines are sometimes prescribed for purposes other than those listed in a Medication Guide. Do not use NSAIDs for a condition for which it was not prescribed. Do not give NSAIDs to other people, even if they have the same symptoms that you have. It may harm them.

If you would like more information about NSAIDs, talk with your healthcare provider. Yo can ask your pharmacist or healthcare provider for information about NSAIDs that is written for health professionals.

This Medication Guide has been approved by the U.S. Food and Drug Administration

Cipia USA, Inc. 10 Independence Boulevard, Suite 300, Warren, Nj 07059 Revised: 08/2024

PACKAGE LABEL.PRINCIPAL DISPLAY PANEL
NDC 69097-138-07 RX ONLY
Meloxicam
Tablets, USP
7.5 mg
PHARMACIST: PLEASE DISPENSE
WITH MEDICATION GUIDE
100 Tables
Cipla

S/N EXP LOT Area for Batch overprinting (Serial No., Expiry & Lot will be overprinted during commercial packing)

NDC 69097-159-07 Rx ONLY Meloxicam Tablets, USP 15 mg PHARMACIST: PLEASE DISPENSE WITH MEDICATION GUIDE 100 Tables Cipla



me	ELOXICAN eloxicam tablet							
Pi	roduct Infor	mation						
Pr	oduct Type		HUMAN PI	RESCRIPTION DRUG	Item C	ode (Source)	NDC	69097-158
Ro	oute of Admini	istration	ORAL					
Ac	tive Ingredi	ient/Activ	e Moiety					
		Ing	redient Na	me		Basis of St	rength	Strength
м	LOXICAM (UNI:	VG2QF83CG	L) (MELOXICA	M - UNI:VG2QF83CGL)		MELOXICAM		7.5 mg
In	active Ingre	dients						
			Ingredia	ent Name			Str	enath
м	AGNESIUM STEA	RATE (UNII:						
SIL	LICON DIOXIDE	(UNII: ETJ7Z 6	XBU4)					
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	lor		FLLOW	Score			nn sente	
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	eloxicam tablet							
P	roduct Infon	mation						
P	roduct Type		HUMAN I	PRESCRIPTION DRUG	Iten	n Code (Source)	NDC	69097-159
R	oute of Admini	stration	ORAL					
A	ctive Ingredi	ent/Act	tive Moiety					
		le le	gredient N	ame		Basis of Str	ength	Strengt
				AM - UNI: VG2QF83CGL)		MELOXICAM		15 mg
Ir	nactive Ingre	dients	loare	dient Name				trenath
c	ELLULOSE, MICR	OCBYST					_	Liengui
	AGNESIUM STEA							
	DOIUM STEARATI							
	olor		YELLOW	Score		no	score	
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