## MELOXICAM- meloxicam tablet REMEDYREPACK INC.

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

v V	ARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS
	See full prescribing information for complete boxed warning.
car car	steroidal anti-inflammatory drugs (NSADs) cause an increased risk of serious diovascular thrombotic events, including myocardala infarction and stroke, whi be fatal. This risk may occur early in treatment and may increase with duratio se (5.1)
	oxicam is contraindicated in the setting of coronary artery bypass graft (CABG perv ( 4.5.1)
incl car svn	Ds cause an increased risk of serious gastrointestinal (GI) adverse events uding bleeding, ukceration, and perforation of the stomach or intestines, which be fatal. These events can occur at any time during use and without warning sptoms. Elderly patientsand patients with a prior history of peptic ukcer disease (or G i bleeding are at creater risk for serious Gi events (5.2)

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1) OA (2.2)and RA (2.3): Starting dose: 7.5 mg once daily Dose may be increased to 15 mg once daily

 Dose may be increased to 15 mg once daily
 PA (2.4):
 7.5 mg once daily in children ±60 kg
 Meloxican tables are not interfacepable with approved formulations of oral meloxicam even if the total milligram strength is the same (2.6) 

Known hypersentlikely to metalscame of any components of the drug product ( 4)
 History of antimum, uricania, or other allergic-type reactions after taking aspirin or other NSAIDs ( 4)
 In the setting of CABG surgery ( 4)

ADVERSE REACTONS
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with red important sectors (1) <u>Runting</u> (SARS) can reduce starburst effect of funcientials and thisaide diuretics. Monitor patients is accure diuretic efficacy including antihypertensive effects. (1) <u>Horizon (SARS) are associated with version (SARS)</u> <u>Horizon (SARS) are associated with version (SARS)</u> <u>Horizon (SARS)</u> <u>Horizon</u> who have difficultus concerning over, See 17 for PATIENT COUNSELING INFORMATION and Medication Guide. Revised: 3/2025

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## FULL PRESCRIBING INFORMATION

## WARNING RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS

Cardiovascular. Thrombolik. Events Nonstrovidal anti-inflammatory drugg (IkBLD) consea an increased Nonstrovidal anti-inflammatory drugg (IkBLD) consea an increased infrarction and stroke, which can be fast. This risk mayoccur early in treatment and maynresses with duration of use [see Warnings and Metholican is contraindicated in the setting of coronary artery bypass graft (CABG) surgery [see Contraindications (4) and Warnings and Precautions (5.1).

Precoutions (5.1). Gastionistamiadescing. Uteration. and Perforation • NSAIDS cause an increased risk of serious gastrointestinal (GI) adverse events licituding balending. Uteration. and perforation of the adverse events licituding balending. Uteration. and perforation of the any time during use and without warning symptoms. Elseriy patientstand patients with a point for point performance of the any time during use and without warning symptoms. Elseriy patientstand patients with a point of the point of the series of the warning and Precedition (2.1).

1 INDICATIONS AND USAGE

## 1.1 Osteoarthritis (OA)

Meloxicam is indicated for relief of the signs and symptoms of osteoarthritis [ see Clinical Studies (14.1)]. 1.2 Rheumatoid Arthritis (RA)

Mebxicam 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 $\geq$ 60 kg [ see Dosage and Administration ( 2.4) and Clinical Studies (14.2)].

## 2 DOSAGE AND ADMINISTRATION 2.1 General Dosing Instructions

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 ( 5) ].

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 regardless of formulation. In patients with hemodialysis, a maximum daily dosage of 7.5 mg is recommended [see Use is pSecific Populations (8 27) and Cinical Pharmacology (12.3)]. Meloxicam may be taken without regard to timing of meals.

2.2 Osteoarthritis

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

## 2.3 Rheumatoid Arthritis

2.3 KRBUINESON PALIFIES For the relief of the signs and symptoms of rheumatoid arthritis, 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.

2.4 Juvenile Rheumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course For the treatment of juvenile rheumatoid arthritis, the recommended oral dose of metoxicam is 7.5 mg once daily in children who weigh  $\approx$  60 kg. There was no additional benefit demonstrated by increasing the dose above 7.5 mg in cinical 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

2.0 non-interchangeaway with other Pointations of neokcam devices tables have not show equivalent systemic exposure to other formulations of or al mebokam. Therefore, meloxicam tables are not interchangeable with other formulations of or al meloxicam product, even if the total miligram strength is the same. Do not substitute similar does strengths of meloxicam tables with other formulations of or al meloxicam product.

## 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 other.

## 4 CONTRAINDICATIONS

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 Mexican is investigated in the following patients:
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 Fistory of asthma, urticaria, or other alergic-type reactions after taking apply or other KSADs. Severe, sometimes fatal, anaphylatic reactions at the taking sprin or other KSADs. Severe, sometimes fatal, anaphylatic reactions to KSADs have been reported in such patients (are Warmings and Precaudors (5.7,5.8))
 Precautions (5.1)

## 5 WARNINGS AND PRECAUTIONS

5.1 Cardiovascular Thrombotic Events 5.1 Cardiovascular Thrombotic Events
Chickinar this of event and nonselective HSAIDs of up to three years duration have shown an increased risk of serious cardiovascular (C) from those to the series of the series of the series cardiovascular (C) from those wellable data, its increase that the series (For C) thrombotic events is similar for all HSAIDs the relative increase in serious C) thrombotic events over baseline conferred by HSAID baseline transmission of the series absolute neclease of excess series CV thrombotic events, due to the increase in the absolute neclease of excess series CV thrombotic events, due to the increase of the absolute neclease of excess series CV thrombotic events, due to the increase of the thrombotic metrics they and a why as the first weeks of the series of the se

thromotor rask has been observed most consistently at higher does. To minimize the potential risk for an adverse CV event in KSAID-traded patients, use the lowest effective does for the shortest duration possible. Physicians and patients should remain alter for the development of such versit, throughout the entire treatment and the development of such versit, should be the entire treatment about the symptoms of seriors. CV events applied to the store to take if they courrent there is no consistent events associated with KSAID use. The concurrent use of apply and the KSAID such as mediocation, increases the they of serious gastrointestinal (CI) events [ see Warnings and Pre-autions 5.2 ).

Status Post Coronary Artery Bypass Graft (CABG) Surgery

Two large, controlled clinical trails of a COX-2 selective NSAID for the treatment of pain in the first 10-14 days following CABG surgery found an increased incidence of myocardial infarction and stroke. NSAIDs are contraindicated in the setting of CABG [see Contraindications ( 4)].

## Post-MI Patients

Paca-LH Dataterist. Observational studies conducted in the Danish National Registry have demonstrated the patients treated with NSAIDs in the post-MI period were at increased risk of reinfarctions. (C-related death, and a cause montable upporning in the First week of treatment. In this years in NSAID-treated patients compared to 12 per 100 percon years in non-NSAID respond patients. Although the absolute rade of death decides domewhat after the first year patients 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.

with a recent M, monitor patients for signs of cardia: achemia. In order in patients 5.2 Gastrointestabal Bleeding, Ukcration, and Perforation NeXDo, including molecular, can cause service gastrointestini (G) adverse events. In a service servic

## Risk Factors for GI Bleeding, Ulceration, and Perforation

Bak Exacts for GI Beedin, Uleration, and Perforation Patters with a prior history of peptic under disease and/or GI beeding who used NSADs had a greater than 10-foid increased risk for developing a GI beed compared to patients treated with NSADs include lange drandom of NSAD therapy, concomitant use of oral confronteness, aspira, anticogaulants, or selective sentonin resplate hinblors controls and the senton of NSAD therapy concomitant use of and controls the senton of national events of control and the sentonin resplate hinblors application of the senton of the senton of the senton of the senton postmarketing reports of fata GI events occurred in delay or debilated patients. Additionally, patients with advanced liver disease and/or cospulpatibility are at increased risk for GI beeling.

sk for G1 bedenig. Tradejis to Minnitze the G1 Risks in NSAID-treated patients: Use the lowest effective dosage for the shortest possible duration. A void ativitätistand of more than one NSAID at a time. A void axie in patients at higher risk unless benefits are expected to outweigh the increased risk of bieldenig, for so such patients, as weld as times with active G1 bedefing. Remain aist for signs and symptoms of G1 ukcration and bedefing during NSAID therapy.

therapy. If a serious GI adverse event is suspected, promptly initiate evaluation and treatment, and discontinue metoxicam unit a serious GI adverse event is niked out. In the setting of concomitant use of low-dose aspirin for cardiac prophylaxis, monitor patients more closely for evidence of GI bleeding [ see Drug Interactions (7) ].

## 5.3 Hepatotoxicity

5.3 Hepatotoxxcxy Elevations of 1.47 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 trial, cases of severe hepatic highry, including fulminant hepatits, liver necrosis, and hepatic falure have been reported.

necrosa, and hepatic faluer have been reported. Bendiors of 11 vr AST (lies its hin threat times ULN) any occur in up to 15% of patients treated with MSAIDs including metoxican. Inform patients of the warring signs and symptoms of hepatotoxicity (e.g., nausa), the symptomy distribution patients, plunder, right upper quadrant restricts and the symptomy distribution patients, plunder, right upper quadrant threads and the symptomy distribution patients, plunder, right upper quadrant threads and the symptomy distribution patients, plunder, right upper quadrant threads and the symptomy distribution of the patient (is set for a for specific Populations (8.8) and Clinical Pharmacology (12.3) [.

## 5.4 Hypertension

3.4 Rypertension NSAIDs, hicklaing méoxicam, can lead to new onset or worsening of preexisting hypertension, either of which may contribute to the increased incidence of CV events. Patients taking anglotensin converting enzyme (ACE) inhibitors, thiaded durietis, or loop duretics 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

and death. Additionally, fluid relation and edena have been observed is some patients treated with ISADIS. Use of melosciam may blant the CV effects of several therapeutic agents used to treat these medical contitions (cs. q. durites, ACE thebits, or angiotens's interestor blockers (RABS) (see Originary in patients with context, and the historic and the Addot the use of melosicam in patients with severt failer services the beenfits are Addot the use of melosicam in patients with severt failers in exection is a patients with severe heart failure, monitor patients for signs of worsening heart failers.

## 5.6 Renal Toxicity and Hyperkalemia Renal Toxicity

Renal Toxickit Long-term administration of NSAIDs, including metoxicam, has resulted in renal papilary necrosis, renal insufficiency, acute renal faiture, and other renal higary. Renal toxick) has also been seen in patients in whom renal postgalandins have a compensatory role in the maintenance of renal perfusion. In these patients, information and, seaso been seen in patients in whom renal postgalandins have a compensation role and the season of the season of the season of the formation and, secondary, in renal bolic flow, which may precipitae over renal decompensation. Patients at greatest risk of this reaction are those with impared renal decompensation. Patients are greatest risk of the reaction are those with impared renal decompensation in the toxics of the season of the deckry. Discontinuation of NSAID breagy is usually followed by recovery to the pretreatment state. The renal effects of moticizem and desease. Exclusic series moticizem and adjoints, maintenand patients and ACE in moticizem and desease. Exclusice series moticizem and patients with precessiting renal desease. Exclusice series moticizem metadores are excircted by the billowed, multicity abation to signs of varienting renal function.

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

# <sup>b</sup> Information is available from controlled directal studies regarding the use of Neohoxim in patients with advanced renal lisease. A nonline the use of neohoxican in patients with advanced renal directae unless the benefits are expected to outwey risk of worsening renal function. If medications used in patients with advanced re disease, monitor patients for signs of worsening renal function ( see Clinical Pharmacology ( 123.)].

## Hyperkalemia

<u>https://adminat</u> Increases in serum potassium 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 attributed to a hyporeninemic-hypoaldosteronism state.

5.7 Anaphyteic Reactions Netwicem has been associated with anaphyteicit reactions in patients with and without known hypersensibly to melokice and in patients with aspirin-sensible asthma [ see Contrandications (4) and Warnings and Precautions (5.8) ].

## Seek emergency help if an anaphylactic reaction occ

5.8 Exacerbation of Asthma Related to Aspirin Sensitivity 5.8 Exacerbation of Asthma Related to Aspirin Sensitivity A subopolistion of patients with astimum may have sepirin-ensitive asthma which may include chronic rhinosinustis complicated by nasal polys; severe, potentially fatal bronchrospara, majori roberiance to aspirin and other KSADS, Because cross-tempolity of the severe to a spiring and severe the SADS, Because cross-bronchrospara, majori resolution, and and the SADS, Because cross-bronchrospara, and and the severe severe severe the severe severe patients, mexican is contrandicated in patients with this form of aspiris sensibility ( without known aspiris sensibility), montor patients for changes in the signs and symptoms of asthma.

## 5.9 Serious Skin Reactions

3.3 serious sean neaccome (SAGDs, including mescamic, can cause serious size) and inter-section, serious mescamic, can cause serious size index index mescamic (SAGDs, including mescamic, can cause serious size) and serious readermain tecrolysis (TEN), which can be faal.HSADDs can also cause fixed drug eruption (TBD). FDE may serious series and series and series and series and series fixed drug eruption mescamic, inform patients about the signs and symptoms of serious site in reactors, and obscientification and the series and symptoms of serious site in reactors, and patient series and series and series and symptoms of series site in reactors, and patient series series and series and symptoms of series site in reactors, and patient series series and series and series and series and pervious series site mescines in bASID (see ContransCause in galants with pervious series).

previous service service services to NSADIS (see Contraindications (g)). 5.0 Drug Reaction with Eosinophila and Systemic Symptoms (DRESS) Drug Reaction with Eosinophila and Systemic Symptoms (DRESS) burg Reaction with Eosinophila and Systemic Symptoms (DRESS) therease the service of the service service of the servic

## 5.11 Fetal Toxicity

ature Closure of Fetal Ductus Arteriosus Avoid use of NSAIDs, including meloxicam, in pregnant women at about 30 weeks gestation and later. NSAIDs, including meloxicam, increase the risk of premature closure of the fetal ductus arteriosus at approximately this postational pae.

## ohydramnios/Neonatal Renal Impairment

upunyatamanankenohala Benai Impairment Use of NSLB)s, chulding relexicum, at about 20 weeks gestation or later in pregnancy may cause feat irenal drysfunction leading to oligohydramios and, in some cause, neonatal renai majament. These adverse outcomes are seen, on average, after drays to weeks of treatment, athough oligohydramios is able en infrequently reported as soon al 4 hours after ASLB olisation. Oligohydramios is other uput of adverse with treatment discontinuation. Complications of polonged objohydramins may. for cause of impaired neonatal renail function, invasive procedures such as exchange transfusion or dialysis were required.

If NSAID treatment is necessary between about 20 weeks and 30 weeks gestation, mebxicam use to the lowest effective dose and shortest duration possible. Conside ultrasound monotring of anniabic fluid if mebxicam treatment estandes beyond 48 hours. Discontinue mebxicam if oligohydramniss occurs and follow up according to clinical practice *le sel Ds en Specific Populations Q*[]).

## 5.12 H

Anemia has occurred in NSAD-irrested patients. This may be due to occuit or gross blood loss, fluid retention, or an incompletely described effect on enythropolesis. If a patient treated with medioxicam has any signs or symptoms of anemia, monitor hemoglobin or hematocrit.

NRJDs, hculding meckkam, may increase the risk of bleeding events. Co-morbid conditions such as coaguitation disorders or concombant use of warfarin, other anticoaguitaria, majatelike agents (ac, a pin/s), serootin republic inhibitors (SSRI), anticoaguitaria, majatelike agents (ac, a pin/s), serootin republic inhibitors (SSRI), these patients for signs of bleeding (see Drug interactions (77).

## 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.

S.14 Laboratory Monikoring Because serious GI bleeding, hepatotoxicky, and renal hjury can occur without warning symptoms or signs, consider monitoring platents on bing-term NSAID treatment with a CEC and a chemistry profile periodicily just Warnings and Precaution (5.2, 5.3, 5.9).

## 6 ADVERSE REACTIONS

EAVERSE FEACTONE
The long adverse reactions are discussed in greater detail in other sections of the
 Cardivascutar Trombotic Events [see Boxed Warning and Warnings and
 Precautions (5.1)]
 Gi Bleeding, Uleration, and Perforation [see Boxed Warning and Warnings and
 Precautions (5.2)]
 Heart Falure and Edema Precautions (5.3)]
 Heart Falure and Edema (see Warnings and Precautions (5.5)]
 Heart Falure and Edema (see Warnings and Precautions (5.5)]
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 Heat Toxicky (see Warnings and Precautions (5.1)]
 Heat Toxicky Eve Warnings and Precautions (5.1)]
 Heat Toxicky Eve Warnings and Precautions (5.1)]

## 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

Laussianciffs and Information Artificia The metoicam Phase 2.7 chical trial database includes 10.122 OA patients and 10.12 RA patients treated with metoic Lan 7.5 mg/aloy, 305 OA patients and 13.11 RA patients patients for a tease 2.7 chical trial patients for a tease on year. Approximately 10.500 of these patients were treated in tem pacebo and/or archite-controlled to these patients were treated in tem pacebo and/or archite-controlled documentation of the set of the

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 meloxicam with placebo and with an active corrict) randomized trials were conducted in placebo trials were conducted in patients with theumatoid arthritis to compare the efficacy and safety of moloxicam 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 15 mg daily 157
 Meloxicam 100 mg daily 153
 Diclofenac 100 mg daily 137

 157
 154
 156
 153

 17.2
 20.1
 17.3
 28.1
 No. of Patients 157 Gastrointestinal 17.2 Gastrointesti Abdominal pain Diarrhea Dyspepsia Flatulence 2.5 3.8 4.5 4.5 1.9 2.6 3.2 4.5 3.2 3.2 3.8 4.5 3.2 3.9 6.5 3.9 7.2 -rpsepsisi 4.5 Fabalence 4.5 Hodyn a Whole 4.5 Fal 0.6 Influersz-kie 5.7 Heidsche 10.5 Respiratory Pharmydis 1.5 Pharmydis 1.5 Respiratory 1.9 Pharmydis 2.5 Kash 2.5 Kas 3.2 4.5 0.0 5.8 2.6 3.3 1.3 4.5 1.9 2.6

4.5 2.6 2.6 7.8 3.8 8.3 2.0 5.9 0.6 3.2 1.3 3.3

1.9 Rash<sup>2</sup> 2.5 2.6 0.6 2.0 "WHO preferred terms edema, edema dependent, edema peripheral, and edema legs combined us, and rash ma

## Table 1b Adverse Events (%) Occurring in ≥2% of Meloxicam Patients in two 12-Week Rheumatoid Arthritis Placebo-Controlled Trials

3.2

d terms rash, rash ervt

Placebo Meloxicam Meloxicam

		7.5 mg daily	15 mg daily
No. of Patients	469	481	477
Gastrointestinal Disorders	14.1	18.9	16.8
Abdominal pain NOS <sup>2</sup>	0.6	2.9	2.3
Dyspeptic signs and symptoms <sup>1</sup>	3.8	5.8	4.0
Nausea <sup>2</sup>	2.6	3.3	3.8
General Disorders and Administrat	ion Site Co	nditions	
Influenza-like illness <sup>2</sup>	2.1	2.9	2.3
Infection and Infestations			
Upper respiratory tract infections-patho class unspecified <sup>1</sup>	igen 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 <sup>2</sup>	6.4	6.4	5.5
Skin and Subcutaneous Tissue			
Disorders			
Rash NOS 2	1.7	1.0	2.1
<sup>1</sup> MedDRA high level term (preferred terms): o			
dyspepsia aggravated, eructation, gastroin infections-pathogen unspecified (laryngitis related signs and symptoms (arthralgia, art effusion, joint swelling)	NOS, pharyne	atis NOS, sinus	itis NOS), joint
<sup>2</sup> MedDRA preferred term: nausea, abdomina NOS and rash NOS	l pain NOS, in	fluenza-like ilin	ess, headache

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

## . Table 2 Adverse Events (%) Occurring in ≿2% of Meloxicam Patients in 4 to 6 Weeks and 6 Month Active-Controlled Osteoarthritis 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 dailv	
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	
Vomitina	0.6	0.8	1.8	2.6	
Body as a Whole					
Accident household	0.0	0.0	0.6	2.9	
Edema <sup>1</sup>	0.6	2.0	24	16	
Pain	0.9	2.0	3.6	5.2	
Central and Peripheral Nervous System					
Dizziness	1.1	1.6	2.4	2.6	
Headache	2.4	2.7	3.6	2.6	
Hematologic					
Anemia	0.1	0.0	4.1	2.9	
Musculoskeletal					
Arthralgia	0.5	0.0	5.3	1.3	
Back pain	0.5	0.4	3.0	0.7	
Psychiatric					
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 <sup>2</sup>	0.3	1.2	3.0	1.3	
Urinary					
Micturition frequency	0.1	0.4	2.4	1.3	
	0.3	0.4	4.7	6.9	
Urinary tract infection <sup>1</sup> WHO preferred terms of combined <sup>2</sup> WHO preferred terms r	0.3 dema, edema de	0.4 ependent, edem	4.7 a peripheral, and	6.9 edema leg	

24460 preterred terms rate, has herghtematus, and make heaplagute combined Higher doods of Viseous Gievantis therefore; the day doos of methoticam should not exceed 3 Tay. Residence of the discussion Gievantis therefore; the day doos of methoticam should not exceed 3 Tay. Residence of the discussion Gievantis therefore; the day doos of methoticam should not exceed 3 Tay. Residence of the discussion of the discussion of the discussion of the discussion Residence of the discussion of the discussion of the discussion of the discussion where exposed to mexican with discussion range from Casto 0.375 mg/s per day in three chical trials. These studies consisted of the 12-week multicenter, double-blind, vertexing on and one Ly-var open-tables strange from Casto 0.375 mg/s per day in three chical trials. These studies consisted of the 12-week multicenter, double-blind, septience, affording three were difference in integratery, in particular, the following provide, were more common in the pediatric than in the adult trials. Rash was reported and or specific subgroup effect. The following is a lot of adverse durg reactions out dimension damonstrate an age mexican in clinical trials involving approximately 16,200 patients.

allergic reaction, face edema, fatigue, fever, hot flushes, malaise, syncope, weight decrease, weight increase
angina pectoris, cardiac failure, hypertension, hypotension, myocardial infarction, vasculitis
ystem convulsions, paresthesia, tremor, vertigo
colitis, dry mouth, duodenal ulcer, eructation, esophagitis, gastric ulcer, gastritis, gastroesophageal reflux, gastrointestinal hemorrhage, hematemesis, hemorrhagic duodenal ulcer, hemorrhagic gastric ulcer, intestinal perforated, perforated duodenal ulcer, perforated gastric ulcer, stomatitis ulcerative
arrhythmia, palpitation, tachycardia
leukopenia, purpura, thrombocytopenia
ALT increased, AST increased, bilirubinemia, GGT increased, hepatitis
dehydration
abnormal dreaming, anxiety, appetite increased, confusion, depression, nervousness, somnolence
asthma, bronchospasm, dyspnea
abpecia, angioedema, bullous eruption, photosensitivity reaction, pruntus, sweating increased, urticaria
abnormal vision, conjunctivitis, taste perversion, tinnitus
albuminuria, BUN increased, creatinine increased, hematuria, renal failure

Building optimizes and the second sec

7 DRUG INTERACTIONS

SecTable 3 To Clinically significant drug interactions with metoxicam. See also Warnings and Precautions ( 5.2, 5.6, 5.12) and Clinical Pharmacology ( 12.3). Table 3 Clinically Significant Drug Interactions with Meloxicam

Dr	ugs that Ir	terfere with H	iemostasis		

Clinical Impact:	Meloxicam and anticoagulants such as warfarin have a
	synergistic effect on bleeding. The concomitant use of
	meloxicam and anticoagulants have an increased risk of serious
	bleeding compared to the use of either drug alone.
	Serotonin release by platelets plays an important role in
	hemostasis. Case-control and cohort epidemiological 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.
intervention:	Monitor patients with concomitant use of meloxicam with
	anticoagulants (e.g., warfarin), antiplatelet agents (e.g., aspirin),
	selective serotonin reuptake inhibitors (SSRIs), and serotonin
	norepinephrine reuptake inhibitors (SNRIs) for signs of bleeding (
	see Warnings and Precautions (5.12) 1.
Aspirin	see warnings and Precaudons ( 5.12) ].
Clinical Impact:	Controlled clinical studies showed that the concomitant use of
	NSAIDs and analgesic doses of aspirin does not produce any
	preater 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 I see Warnings
	and Precautions ( 5.2) 1.
ntervention:	Concomitant use of meloxicam and low dose aspirin or analgesic
	doses of aspirin is not generally recommended because of the
	increased risk of bleeding [ see Warnings and Precautions ( 5.12)
	1.
	Meloxicam is not a substitute for low dose aspirin for
	cardiovascular protection.
ACE Inhibitor	s, Angiotensin Receptor Blockers, or Beta-Blockers
linical Impact	
. innearmpace.	
	angiotensin converting enzyme (ACE) inhibitors, angiotensin
	receptor blockers (ARBs), or beta-blockers (including
	propranolol).
	In patients who are elderly, volume-depleted (including
	those on diuretic therapy), or have renal impairment.
	coadministration of an NSAID with ACE inhibitors or ARBs may
	result in deterioration of renal function, including possible acute
	renal failure. These effects are usually reversible.
ntervention:	During concomitant use of meloxicam and ACE inhibitors.
intervention:	During concomitant use of meloxicam and ACE inhibitors, ARBs, or beta-blockers, monitor blood pressure to ensure that
Intervention:	During concomitant use of meloxicam and ACE inhibitors, ARBs, or beta-blockers, monitor blood pressure to ensure that the desired blood pressure is obtained.
Intervention:	During concomitant use of meloxicam and ACE inhibitors, ARBs, or beta-blockers, monitor blood pressure to ensure that the desired blood pressure is obtained. During concomitant use of meloxicam and ACE inhibitors
Intervention:	During concomitant use of mebxicam and ACE inhibors, ARBs, or beta-blockers, monitor blood pressure to ensure that the desired blood pressure is obtained. During concomitant use of mebxicam and ACE inhibitors or ARBs in adients who are deferty, obume-deoleted, or have
Intervention:	During concomitant use of mebxicam and ACE inhibors, ARBs, or beta-blockers, monitor blood pressure to ensure that the desired blood pressure is obtained. During concomitant use of mebxicam and ACE inhibitors or ARBs in adients who are deferty, obume-deoleted, or have
Intervention:	During concomtant use of mebxicam and ACE inhibitors, ARBs, or beta-bockers, monitor blood pressure to ensure that the desired blood pressure is obtained. During concomtant use of mebxicam and ACE inhibitors or ARBs in patients who are elderly, volume-depleted, or have impaired renal function, monitor for signs of worsening renal
Intervention:	During concomitant use of meioxicam and ACE inhibitors, ABBs, on teah-blockers, monitor blood pressure to ensure that the desired blood pressure is obtained. During concomitant use of meioxicam and ACE inhibitors or ABBs in patients who are delerly, volume-depleted, or have impaired renal function, monitor for signs of worsening renal function (see Warnings and Precautions (5.50).
Intervention:	During concomtant use of meloxicam and ACE inhibitors, ABS, or beta-blockers, monto blood pressure to ensure that the desired blood pressure is obtained. During concomtant use of meloxicam and ACE inhibitors or ABS in patients who are edderly, volume-depeted, or have impared renal function, monitor for signs of worsening renal function [ see Warnings and Precautions ( 5.6)]. When these drugs are administered concomitantly.
Intervention:	During concomitant use of metoxicam and ACE inhibitors, ABS, or bica-bickers, monitor biod pressure to ensure that the desired biod pressure is obtained. During concomitant use of metoxicam and ACE inhibitors or ARBs in platients who are detery, volume depicted, or have mighted renal function, monitor for signs of worsening renal monitor and the determinant of the anti- metor and the determinant of the anti- determinant of the determinant of the determinant When these drugs are administered concomitantly, batterns should be adequately hydrated. Assess renal function at
Intervention:	During concomtant use of mebsicam and ACE Inhibors, BBs, or beb-bickers, minichr blod pressure to ensure that the denired blod pressure to obtained. Or ABS in platether with the advectory of the advectory or ABS in platether who are defery, vulnim and ACE inhibors intuction (see Warnings and Precaudions (5.6)). Tuctoring see the administered concontantifycom the beginning of the concontant reterment and periodically the beginning of the concontant reterment and periodically
Intervention:	During concomitant use of metoxicam and ACE inhibitors, ABS, or bica-bickers, monitor biod pressure to ensure that the desired biod pressure is obtained. During concomitant use of metoxicam and ACE inhibitors or ARBs in platients who are detery, volume depicted, or have mighted renal function, monitor for signs of worsening renal monitor and the determinant of the anti- metor and the determinant of the anti- determinant of the determinant of the determinant When these drugs are administered concomitantly, batterns should be adequately hydrated. Assess renal function at

t:Clinical studies, as well as post-marketing observations, showed that NSAIDs reduced the natriuretic effect of loop diuretics (e.g., furosemide) and thaizid eduretics is some patients. This effect has been attributed to the NSAID inhibition of renal prostaglandir

	synthesis. However, studies with furosemide agents and metoxicam have not demonstrated a reduction in natriuretic effect. Furosemide single and multiple dose pharmacodynamics and pharmacokinetics 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 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 Pharmacology (12.3)].
Intervention:	During concomitant use of meloxicam and lithium, monitor patients for signs of lithium toxicity.
Methotrexat	
	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	
Clinical Impact:	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 S	Salicylates
	Concomitant use of meloxicam with other NSAIDs or salcylates (e.g., diflunisal, salsalate) increases the risk of GI toxicity, with ittle 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	
	Concomitant use of meloxicam and pemetrexed may increase the risk of pemetrexed-associated myelosuppression, renal, and GI toxicity (see the pemetrexed prescribing information).
Intervention:	Buring concomitant use of meloxican and penetrexed, in patients with real impairment vhose creating clearance range from 45 cp 3 m.Lmin, monitor for myelosuppression, real aid patients taking mechanism should be real to a strain patients taking mechanism should be and the strain the days before, the day of, and two days following penetrexed administration. In patients with creating be able with in patients with creatine clearance below 45 m.Lmin, the concomitant administration of meloxican with penetrexed is no concomitant, administration of meloxican with penetrexed is no concomitant.

## 8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy Risk Summary

If remains Exosure to virsue ductions in the basis of the setting setting to the set of the setting setting to the set of the set of

Mean-table vality in the second secon

organopresis at an oral dose equivalent to 2.6 and 26-times the MHID (see Data). Based on animal data, prostspändins have been shown to have an important role in endonetral vascular permeability, biatocyst implantation, and decisulatation. In animal studies, administration of prostspändern synthesis hibbors exu has metoxican, resulted in increased pre- and post implantation loss. Prostspändins also have been shown to prostagalind synthesis hibbors have been reported to impair kideny development when administered at clinically relevant doses. The estimated background risk of mage been reported to impair kideny development order advises automes. In the U.S. general population, risk of birth defect, toss, order advises automes. In the U.S. general population, rule estimated background risk and 33-to 20%, respectively in clinically recognized pregnances is 2% to friely and 33-to 20%, respectively.

Clinical Considerations Fetal/Neonatal Adverse Reactions

reumreunidal Adverse Reactions Premature Closure of Felal Ductus Arteriosus: Avoid use of NSAIDs in women at about 30 weeks gestation and later in pregnancy, because NSAIDs, including metboxicam, can cause premature closure of the felal ductus arteriosus ( see Jabla).

Labor or Delivery

There are no studies on the effects of meloxicam during labor or delivery. In animal studies, NSAIDs, including meloxicam, inhibit prostaglandin synthesis, cause delayed parturition, and increase the incidence of stillbirth.

<u>Data</u> Human Data

Premature Closure of Fetal Ductus Arteriosu

Published literature reports that the use of NSAIDs at about 30 weeks of gestation and later in pregnancy may cause premature closure of the fetal ductus arteriosus. Oligohydramnios/Neonatal Renal Impairment:

Olgohydramics/Neonatal Renal Impairment: Published studies and postmarkeling-proofs discribe maternal MSAID use at about 20 wess getation or later in pregnancy associated with fetal renal dysfurction leading to glophydramics. The discrimination of the second renal renal major maternal the second renal dysfurction have additional renal renal renal renal major maternal renal major maternal renal major maternal renal major maternal renal r

Animal Data Metockam was in chicadoperic, the alimitational operation of the data of the data metockam was in chicadoperic, the alimitation of the data of the data of the data of metockam based on BBA company. A mitotication of metockam to pregnant rabbits throughout embryogenesis produced an increased in cidence of septal defacts of the hermit ali an cardiacide of the programma of the MetoD based on BBA. Deface of the data based on BBA conversion). In rats and nabits, embryotethaity occurred at oral metockam dotes of 1 mitoglogian and 5 -5 fold throughout organogenesis.

## 8.2 Lactation

Risk Summary

There are no human data available on whether metoxicam is present in human milk, or on the effects on breakted infants, or on milk production. The developmental and health benefits of breakteding should be considered along with the mother's clinical need for meboxicam and any potential adverse affects on the breasted infant from the meboxicam of rom the underlying effects on the breasted infant from the meboxicam of rom the underlying effects.

## Data Animal 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 Infertility

Females

Females Based on the mechanism of action, the use of prostaglandh-mediated NSAIDs, including mebxicam, may delay or prevent rupture of ovarian foldiclas, which has been associated with reversible integrity is nome worms. Tubblind animal studie have shown that prostaglandh-mediated folkular rupture required for ovalation. Small studies in worms traded with NSAIDs have do shown are versible delay in ovalation. Small studies in worms of MSAIDs have do shown are versible delay in ovalation. Small studies in worms of MSAIDs have do shown are versible delay in ovalation. Small studies in worms of MSAIDs, including mediation, in women who have difficulties conceving or who are undergoing investigation of infertity.

## 8.4 Pediatric Use

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

8.5 Genetric 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, 5.24) ].

## 8.6 Hepatic Impairment

No dose adjustment is necessary in patients with mid to moderate hepatic imparment Patients with severe hepatic imparment have not been adequately studied. Since mebokam is againgrandly metabolized in the lever and hepatotoxicity may courcr, use mebokam with caution in patients with hepatic impairment [ see Warnings and Precautions (5) and China Al hematoby (21.2).

## 8.7 Renal Impairment

No dose adjustment is necessary in patients with mid to moderate renal impairment. Patients with severe renal impairment have not been studied. The use of melokkam in subjects with severe renal impairment is not recommended. In patients on hemodajissi melokkam should not exceed 7.5 mg per day. Melokkam is not dalyzable [see Dosage and Administration (2.1) and Chiera Pharmacology (12.3)].

## 10 OVERDOSAGE

20 OPENLOSMC Symptons following acute NSAID overdosages have been typically imited to lethargy, drowsiness, nause, voniting, and epigastrc pain, which have been generally reversible with supportive care. Gostrometistant beefing has occurred. Hypertension, acute renal Warningsand Precautions ( 5.1, 5.2, 5.4, 5.6)

Warningsand Precautions (5.1, 52, 54, 56, 0). Manage patients with symptomatic and supportive care following an NSAID overdisage. There are no specific antiotics. Consider emissia and/or activated charcold (50 to 100 construct cathart in symptomatic apatients seem within four hours of highestion or in patients with a large overdisage (5 to 10 times the recommended disage). Forced option and a strategies of the symptometic apatient is seen within four hours of highestion or in patients with a large overdisage (5 to 10 times the recommended disage). Forced option area balance of metodicana. Accelerated removal of metodisand by 4 g oral doses of cholestyramine given three times a day was demonstrated in a clickal trial. For addimistration of cholestyramine given tures useful following on exercising and For addimistration of cholestyramine given submit following on web disages. For addimistration of cholestyramine given submit following on exercising 1. For addimistration of cholestyramine given submit following on exercising the for addimistration of cholestyramine given the submit following on exercising the for addimistration of cholestyramine given thread times and y was demonstrated in a chickal trial. For addimistration of cholestyramine given thread times and y was demonstrated in a chickal trial.

## 11 DESCRIPTION

Mexicam is a nonsteroidal anti-inflammatory drug (NSAID). Each tablet contains 7.5 mg or 15 mg meloxicam, USP for oral administration. Meloxicam is chemically designated as 4-hydroxy--methyh-IV-Smethyt-2-hitalox/12-41-2-bearcothitaina-8-achrosanide-1.1-dioxide. The molecular weight is 351.4. Its empirical formula is C <sub>14</sub>H <sub>13</sub>N <sub>3</sub>O 45 <sub>2</sub>and t has the following structural formula:



Mebxicam is a pale yelow solid, practically insoluble in water, with higher solubility observed in strong acids and bases. It is very slightly soluble in methanol. Mebxicam has an apparent partition coefficient (log P)  $_{\rm app}$ = 0.1 in *n*-octanol/buffer pH 7.4. Mebxicam 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, lactose anhydrous, colloidal silicon dioxide, sodium citrate dihydrate, magnesium stearate.

## 12 CLINICAL PHARMACOLOGY

## 12.1 Mechanism of Action

Meloxicam has analogsic, anti-inflammatory, and antipyretic properties

Metoxicam has analgetic, anti-hilmmnatory, and antityretic properties. The mechanism of action of metoxicam, like that of other NSADDs, is not completely understood but involves inhibitor of cyclooxygenate (COX-1 and COX-2) Metoxicam is a pacter inhibitor of prostagilantis synthesis is virtize. Metoxicam before the strength of the strength of the strength of the strength of the sensible afferent nerves and potentiate the action of brankytism in inducing pain in inhibitor of prostagilandin synthesis. Is mode of action may be due to a decrease of prostagilandin synthesis. Is mode of action may be due to a decrease of prostagilandin synthesis.

## 12.3 Pharmacokinetics

1.1.3 PrimmackinetCs Absorban The absorba bioavailability of meloxicam capsules was 89% following a single oral dose of 30 mg compared with 30 mg fV bolus injection. Following single intravenous doses, dose proportional pharmacchietics were shown in the range of 5 mg to 80 mg. After over the range of 1.5 mg to 3.5 mg. Medica mg to 80 mg. After over the range of 1.5 mg to 3.5 mg. Medica mg to 80 mg. After after a 7.5 mg meloxicam table was taken under fasted conditions, indicating a prolonged drug absorban. With multiple boding, staesly-state concentrations were reasoned to 40 mg. A mg to 40 mg to 10 mg to 40 mg to 10 mg to 40 mg to 10 mg to 40 mg to 10 mg to 40 mg to 10 mg to 40 mg to 10 mg to 40 mg to 10 mg to 40 mg to 10 mg to 40 mg to 10 mg to 40 mg to 10 mg to 40 mg to

Meloxicam capsules have been shown to be bioequivalent to meloxicam tablet Table 4 Single Dose and Steady-State Pharmacokinetic Parameters for Oral 7.5 mg and 15 mg Meloxicam (Mean and % CV)1

Pharmacokinetic Parameters	Steady Stat	e	Single Dos	Single Dose		
Parameters (%CV)	Healthy male adults (Fed) <sup>2</sup>	Elderly males (Fed) <sup>2</sup>	Elderly females (Fed) <sup>2</sup>	Renal failure (Fasted)	Hepatic insufficiency (Fasted)	
	7.5 mg <sup>3</sup> 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 2/f 4[L]	14.7 (32)	15 (42)	10 (30)	26 (44)	14 (29)	

ter values in the table are from various studies <sup>1</sup>The param

<sup>2</sup>not under high fat conditions

<sup>3</sup>Meloxicam tablets <sup>4</sup>V <sub>2</sub>/f =Dose/(AUC•K <sub>el</sub>)

Food and Antacid Effects

Food and Anacké Effects Administration of medicaria capacities following a high fat breakfast (75 g of fat) resulted in mean paix drug kevés (Lie, cm.) being increased by approximately 275% while the administration of medicaria capacities and administration of the second sec

concombant administration of anraces. <u>Distribution</u> <u>Distri</u>

## Elimination Metabolism

Metabolism Metabolism Metoxcam is extensively metabolized in the liver. Mebokcam metabolites include 5-carboxy mebokcam (60% of dose), from P-450 mediated metabolism formed by oxidation of an intermediate metabolism (5-yhdroxymethy) metabolism (6-yhdrox), (includes), and (includes), invitational (includes) indicate that CP2C9 (includes), and (includes), invitational (includes) indicate that CP2C9 (includes), and (includes), and (includes) indicate that CP2C9 (includes), and (includes), and (includes), and (includes), and (includes), is probable for the other two metabolites which account for 15% and 4% of the administered does, respectively, All the four metabolites are not known to have any in vicopharmacological activity.

Excretion Medixican excretion is presonnianelly in the form of metabolities, and accurs to equal extents in the urine and feeces. Only fraces of the unchanged parent compound are excreted in the urine (20%) and feece (1.6%). The extent of the urinary accurstion was confirmed for unlabeled multiple 7.5 mg doses: 0.5%, 6%, and 13% of the dose were included the second second second second second second second second metabolities, respectively. There is significant bilary and/or entral second second base of molecular discression that U.C. of molecular by 50%. The mean elimination half-left t<sub>1.0</sub>) ranges from 15 hours to 20 hours. The elimination fail 4% is constant across dose levels indicating laser metabolism within the therapeutic dose of molecular across dose levels indicating laser metabolism within the therapeutic dose of any parent devance ranges from 10.9 mLmm. Specific Regulations **Beditize After** single (0.25 mg/kg) dose administration and after achieving steedy state (0.375 mg/kg/dos), there was a general trend of approximately 30% lowes reposure in younger department melocitican programs similar (single dose) or sightly reduced (steady state) to those in the adult patients, when using AUC values normalized to a dose of 2.5 mg/kg (steady bulk), tubies to polyage in the theolice on registry reduced (steady state) to those in the adult patients, when using AUC values normalized to a dose of 3.5 mg/kg (steady bulk), tubies to polyage in the to the steady bulk to not age. Excretion

The second particle analysis, utilizing population pharmacokinetics body-weight, but not age, was the single predictive covariate for differences in the meloxicam apparent oral plasma 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 been investigated.

## Geriatric

Eatery make (165) sures of app) exhibited metocicam planna concentrations and study-state planna accientics similar up young metos. Eatery females (165) years of age) had a 47% halper ALC\_gand 33% higher *C* max\_ase compared to younger females (165) years of age) after body vegits normalization. Explain the increased total or both eatery states and the study and the state of the state and total or both eatery plants: populations. A smaller free fraction was found in eidenty female platents in comparison to defay make platents.

Sex

Toung formulas exhibited signify lower plasma concentrations relative to young mulas. After single dates of 7.5 mg networks m. Its means information hard few with 15.5 hours for the female group as compared to 23.4 hours. For the male group, At steady states the data were similar (12.9 hours vs. 21.4 hours). This pharmacokitest difference du gender is likely to be of Rite clinical importance. There was linearly of pharmacokitest and no appreciable difference in the <code>cmax0T magnetross genders</code>. Hepatic Impairment

Hepatic Impairment Following a single 15 mg dose of mebxicam there was no marked difference in plasm concentrations in patients with mill (Dhili-Pugh Class I) or moderate (Chili-Pugh Class Was not affected by hepatic Impairment. No dosage adjustment is necessary in patient with mild to moderate hepatic Impairment Patients with seven hepatic Impairment (<u>L3</u>) and Use in Specific Populations (<u>R4</u>).

(3.2) and Use in Specific Populations (20): Real Impairment Mebuckam pharmacokinetics have been investigated in subjects with mid and moderate real impairment. Tayle bad ong bares accusentrations of metricular intervenes and total values were similar in all prougs. The higher mebuckam clearance in subjects with real paramet ray bad is to increased intraction of unknown molecular which is available in patients with mid to moderate real impairment. Patients with severe renal impairment ray be used in storesteer law of medication which were real-ing attemption of the severe real impairment of the severe renal impairment recursions (20) and Use in Specific Populations (20).

Prications (<u>56</u>) and use in space Hendrally/S Following a single does of metodamic, the fine C <sub>subp</sub>lana concentrations were higher backets with recent billive on charact, themodily/siz (1% free fraction) in comparison to heathy outneers (0.3% free fraction). Hemodily/siz dd not lower the total drug concentration in plasms: therefore, additional does are not necessary after hemodily/siz. MedixCam is not dilyzable [ see Dosage and Administration (<u>2.1</u>) and Use in Specific Populations (<u>2.1</u>).

In Specific Populations (E.D.). Drug Interaction Studiet Appirity/Nen KSADs were administered with asprin. Iter protein binding of NSADs were restuced, abrough the clearance of free NSAD was not altered. When makes an a administered with asprin 10000 mg threat time daily to heathy valuaters, it tended theraction are to the original clearance of the NSADs were administered. The second with appin is see Drug Interactions (2). Chestry-anime Pretentiment for four days with chestry-amine significantly increased the clearance of medvician by 50%. This resulted in a decrease in 12, from 12. Duots cricculation pathway for medvican in the gastronitational tract. The clinical relevance of this interaction has not been established.

ine:Concomitant administration of 200 mg cimetidine four times daily did not alter gle-dose pharmacokinetics of 30 mg meloxicam. *Cimetidii* the sing

Digoxin:Meloxicam 15 mg once daily for 7 days did not alter the plasma concentration profile of digoxin after β-acetyldigoxin administration for 7 days at clinical doses. In vitrotesting found no protein binding drug interaction between digoxin and meloxicam. Values and not no process of any of a second second

receiving Ithum also I [see Drug Interactions (2)]. The second set of subjects Mechoreaacta Astrophysical Tai heumatodia and anticity. Biol pairemis evaluated the effects of multiple doess of metoxican on the pharmacolitics of methoreaacta taken once weeky. Metoxican do not have a significant effect on the pharmacoliticatics of single and the second sec

13 NONCLINICAL TOXICOLOGY

# 2. Intercharter Understein, Impairment of Fertility <u>Carcinopensis</u>, Mutagenesis, Impairment of Fertility <u>Carcinopensis</u> There was no Evrose in tumor incidence in long-term carchogenicity studies in rats (104 wesk) and mice (99 wesk) administered metockam at oral doses up to 0.8 mg/qdg/ab in rats and up to 8.0 mg/qd/ab in mice (up to 5.0 and 2.6 times, respectively, the maximum recommended human dose (MRHD) of 15 mg/day melockam based on toby structure area (BSA) comparison).

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

Impairment of Fertility

Meloxicam did not impair male and female fertility in rats at oral doses up to 9 mg/kg/day in 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 Osteoarthritis and Rheumatoid Arthritis 

placebo. The use of meloxicam for the management of signs and symptoms of osteoarthritis was evaluated in six double-bihd, active-controlled trials outside the U.S. ranging from 4 week's to 6 montrol's duration. In these trials, the efficacy of meloxicam, in doese of 7.5 mg/day and 15 mg/day, was comparable to proxicam 20 mg/day and dicbfenac SR 100 mg/day mid consistent with the efficacy seen in the U.S. trial.

registry and considered with the care constrained or migrate and dicheries. SR 10 are the care of remainscalar for the service of the Bull. SN that care of remainscalar particles was evaluated in a 12 years, double blag controlled multitational trial. Medician (T-Sm) 15 mg, and 22 sm updates) and goally was compared to placebo. The primary endpoint in this study was the ARX20 response refer, a composite measure of clinka, and 35 mg days showed significant improvement in the primary endpoint compare with placebo. No incremental benefit was observed with the 225 mg dose compared to the 15 mg dose.

## 14.2 Juvenile Rheumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course

The use of meloxicam for the treatment of the signs and symptoms of pauciarticular oplyarticular course juvenile Rheumatoid Arthritis in patients 2 years of age and older was evaluated in two 12-week, double-blind, parallel-arm, active-controlled trials.

The standard set is the 4 standard state and the state of maximum; or meaxiam and 15 mg/ggag or haproxen. The effracy analysis used the ACR Pediatrix 30 responder definition, a composite of parent and investigator assessments, counts of active joints and joints with Imited rang of motion, and entythrocyte sedimentation rate. The proportion of responders were similar in all three groups in both studies, and no difference was observed between the mebokam does groups.

## 16 HOW SUPPLIED/STORAGE AND HANDLING

Meloxicam tablets, USP 7.5 mg are yellow coloured, round, biconvex tablets, debossed with "158" on one side and "C" on the other. NDC: 70518-2631-00 NDC: 70518-2631-00 NDC: 70518-2631-01 NDC: 70518-2631-02 PACKAGING: 60 in 1 BOTTLE, PLASTIC PACKAGING: 30 in 1 BLISTER PACK PACKAGING: 30 in 1 BLISTER PACK Store at 20°C to 25°C (68°F to 77°F) [see USP Controlled Room Temperature]. Keep meloxicam tablets in a drv place. metoxicam tablets in a dry place. Dispense tablets in a tight container. Keep this and all medications out of the reach of children. Repackaged and Distributed By: Remedy Repack, Inc 625 Kolter Dr. Suite #4 Indiana, PA 1-724-465-8762

## 17 PATIENT COUNSELING INFORMATION

Ar Primer Consistent in Constraints in Constraints and Advise the patient to read the FDA-approved patient labeling (Medication Guide) that accompanies each prescription dispensed. Inform patients, familise or the 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 and Precadions (52).

Advise patients to report symptoms of ulcerations and bleeding, including epigastric pain, dyspepsia, melena, and hematemesis to their healthcare provider. In the setting of

concomitant use of low-dose aspirin for cardiac prophylaxis, inform patients of the increased risk for the signs and symptoms of Gi bleeding [ see Warnings and Precautions (5.2)].

## Hepatotoxicity

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

Heart Failure and Edema

Heart Failure and coema Advise paients 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 (d) and Warnings and Precautions (5,2)).

Service Skin Reactions, including DBKS Advise patients to stop taking mebxicam immediately if they develop any type of rash or fever and to contact their healthcare provider as soon as possible [ see Warnings and Precautions (Sp. 540)].

## Female Fertility

Advise females of reproductive potential who desire pregnancy that NSAIDs, including metoxicam, may be associated with a reversible delay in ovulation [ see Use in Specific Populations (8.3)].

Populations (EU): Facil Toxicty: Inform preparat women to avoid use of meloxicam and other NSAIDS starting at 30 weeks gestation because of the risk of the premature closing of the fetal ductus arterious. If treatment with meloxicam is needed for a preparat woman between about objective and the treatment of the previous of the previous of the set of the disploytial minits. It treatment continues for longer than at hours I see Warnings and Precautions(5,1)) and Use in Specific Populations (6,2).

recasions\_\_\_\_\_ in a use in specific reputations (g\_\_\_) i. Assi Concombatility and NSAIDS. Inform patients that the concomband use of mebicians with other NSAIDS or sakylates (g\_s\_\_\_) distants, adjusted is not recommended by the the increased risk of and proceedings of the patient of the specific of the patients of the NSAIDS or sakylates Proceedings (g\_\_\_) and Drug Interactions (g\_\_\_) and the patients that MSAIDs may be present in Yower the counter medications for transment of costs, feec, or is soma.

Use of NSAIDs and Low-Dose Aspirin Unter un name and universe against a series of the series

(724) 465-8762

(1/24) 405-062 Medication Guide for Nonsteroidal Anti-Inflammatory Drugs (NSADS) What is the most important information I should know about medicines called Nonsteroidal Anti-Inflammatory Drugs (NSADS)? NSSDDs can cause serious side effects, including: NSADS can cause serious side effects, including: neuroscience of the heart attack or stroket that can lead to death. This risk may import early in treatment and may increase : with increased risk ones of NSADS.

with increasing doses of NSAIDs
 with longer use of NSAIDs

• with onger use of NAUGS Don't take NASIDS right before or after a heart surgery called a "coronary after yoppas graft (CABO.)" Axoid taking NASIDs after a recent heart attack, unless your heakhcare Axoid taking NASIDs after a recent heart attack. I'you take NASIDs after a recent heart attack.

The risk of getting an ulcer or bleeding increases with:

advanced liver disease
 bleeding problems

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

## What are NSAIDs?

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?

Who should not start NSADs: • I you have had an asthma attack, hives, or other allergic reaction with aspirin or any other NSADs: • right before or after heart bypass surgery.

Ingit before or after heart bypass surgery.
Before taking KSADS, fell your healthcare provider about all of your medical
controls that any subscription of the second second

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

Nearch are provider first. What are the possible side effects of NSAIDs? NSAIDs can cause serious side effects, includir

Insulus call cause serious site enterch, including: See "What is the most important information I should know about medicines called Nonsteroidal Anti inflammatory Drugs (NSAIDs)?" Houri failure I berg problems including kindly failure I berg problems kin

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

shortness of breath or trouble breathing @ slurred speech chest pain @ sweling of the face or throat weakness in one part or side of your bddy

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

nausea	<ul> <li>there is blood in your bowel movement or it is black</li> </ul>
<ul> <li>more tired or weaker than usual</li> </ul>	vomit blood
<ul> <li>diarrhea</li> </ul>	<ul> <li>there is blood in your bowel movement</li> </ul>

	or it is black and sticky like tar
<ul> <li>itching</li> </ul>	<ul> <li>unusual weight gain</li> </ul>
<ul> <li>your skin or eyes look yellow</li> </ul>	<ul> <li>skin rash or blisters with fever</li> </ul>
<ul> <li>indigestion or stomach pain</li> </ul>	<ul> <li>swelling of the arms, legs, hands and feet</li> </ul>
<ul> <li>flu-like symptoms</li> </ul>	

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 heathcare 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-1088.

FDA at 1-800-FDA-1088. **Other information about 15XIDS:** Apprint an information about 15XIDS: Apprint an information about 15XIDS: Apprint and information about 15XIDS: the stronger and information about the stronger about the stronger about the stronger the stronger about the stronger about the stronger about the stronger about the stronger Some SXADE are stold in low does without a prescription (over-the-counter). Takk 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 Medical 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. You 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. Repackaged By / Distributed By: RemedyRe ack Ind

625 Kolter Drive. Indiana. PA 15701

(724) 465-8762

(724) 465-8762 DRUG: MELOXICAM GENERIC: meloxicam DOSAGE: TABLET ADMINSTRATION: ORAL

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