MELOXICAM- meloxicam tablet NCS HealthCare of KY, LLC dba Vangard Labs

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

WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTRO Nonstraidal anti-finamantary drag (NLRAD) cause a local sector of serious cardiovascular thrombolic works, including mycardial infarction and tarks, which use (5.1).
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RECENT Major Warnings and Precautions, Drug Reaction with Eosinophila and Systemic Symptoms (5.10) 04/2021 Warnings and Precautions, Fetal Toxicity (5.11) 04/2021

ngs and Prečautions, Ferial Toxicity (5.11) 04/2021 Micam is a non-steroidal anti-inflammatory drug indicated for: teoarthriss (0.4) (1.1) eumatoid Arthritis (RA) (1.2) endle Riveumatod Arthritis (RA) in patients who weigh ≥60 kg (1.3) Meloxi • Ost • Rho

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 Use to solve a solve the shortest duration constant with individual patient reasoners
 water annies (von solve a solve a

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

Known hypersensibility to melosicam or any components of the drug product (4)
 Hotsey of authors, unclarity, or other allergic-type reactions after taking apprin or other NSADs (4)
 In the setting of 2468 surgery (4)

Head of statistical website control and an experimentation of the statistical product (14)
 WARRINGS AND PRE-CALINDIS
 WARRINGS AND PRE-CALINDIS

ADVERSE REACTIONS
 Most common (25% and greater than placebo) adverse events in adults are diarrhea, upper
respratory trackindscrinds, oppergrama, and initianzes. Make symptoms (6.1)
 Adverse events observed in pediatric studies were similar in nature to the adult clinical trial experises
 (6.1)

(6.1) To report SUSPECTED ADVERSE REACTONS, contact Cipia Limited, India at 1-866-664-3268 or FDA at 1-800-FDA-1088 or www.fda.acovimedwatch. Data at a method and the sector of the sec

of metoscicam and analysis: Doess of apprin in not generally recommended (?) ACE tabilities, designed in Becerar Buckers (JBBI) of the Buckers Concentrate uses with ACE tabilities, advanced and the second secon

(7)
 USE IN SPECIFC POPULATIONS
 (minimum, NSADs are associated with reveable infertity. Consider withdrawal of melosicam in women
with have difficult convoluting (13)
See 17 for PATIENT COUNSELING INFORMATION and Medication Guide.
 Revised: 32023

FULL PRESCRIBING INFORMATION: CONTENTS* WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL

WARNING: RISK OF SERTUOS units... PENTS RECENT PARAGE SHAPU BACKE 1.1 Ostroarthins (GA) 1.2 Rhournatoid Arthriks (GA) 1.3 Juvenile Rheumatoid Arthriks (GA) Pauciarticular and Polyarticular Course Articula

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Pregnancy Lactation

Lactation Females and Males of Reproductive Potential Pediatric Use Geriatric Use Hepatic Impairment Renai Impairment /ERDOSAGE

8.7 Reinal Impairment
0 overktoosAct
1 DeScription
1 DeSc

Meloxicam is contraindicated in the setting of coronary artery bypass graft (CABG) surgery [see Contraindications (4) and Warnings and Precautions (5.1)].

1.2 Rheumatoid Arthritis (RA)

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

Learner of the second seco

Course 16 HOW SUPPLIED/STORAGE AND HANDLING 17 PATIENT COUNSELING INFORMATION * Sections or subsections omitted from the full prescribing information are not listed

FULL PRESCRIBING INFORMATION

WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS

AND GASTROINTESTINAL EVENTS Cardiovascular Thrombolic Eventa • Nonsteroidal anti-inflammatory drugs (INSAIDs) cause an increased ri of serious cardiovascular thrombotic events, including myocardial infarction and stroke, which can be fattal. Thin risk may occur early in treatment and may increase with duration of use [see Warnings and Precardions (J.D.).

Sastrointestinal Bleeding, Ulceration, and Perforation

MANDOWINGSING BEENDING, MICHAEL, MICHAE

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)].

2.1 General Dosing Instructions

Carefully consider the potential benefits and risks of meloxicam and other treatme options before deciding to use meloxicam. Use the lowest effective dosage for the shortest duration consistent with individual patient treatment goals [see Warnings 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 regardles of formulation. In patients with hemodialysis, a maximum daily dosage of 7.5 mg is recommended [see Use in Specific Populations (8.7), and Clinical Pharmacology (12.3)].

Meloxicam may be taken without regard to timing of meak

2.2 Osteoarthritis

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

2.3 Rheumatoid Arthritis

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.

may receive abdomatebeties by screasing the bose of a J may unce vary. 2.2 Apventile Returned J Arthrift B(JAR) Pauckarticular and Polyarticular Course for the treatment of juvelle rhoumatod arthrifts, he recommended and alose of mediciant 3.7 approx. The provided arthrifts, but recommended and alose of mediciant 3.7 approx. The provided arthrifts, but recommended and baset demonstrated by increasing the dise abover 3.7 mg in cleak trials. Mediciant tables abduint of but used in children who regly a 50 kg.

2.5 Renal Impairment

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

geability with Other Formulations of Meloxica 2.6 Non-Intercha

2.0 non-interChangeabacky with Utner formulations of relexican Mexican tables have not show equivalent systemic exposure to other approved formulations of oral mebokam. Therefore, meloxicam tables are not interchangeable with other formulations of oral mebokam product. even if the total milligram strength is the same. Do not substitute similar does strengths of meloxicam tables with other formulations of oral meloxicam product.

3 DOSAGE FORMS AND STRENGTHS

Moxickan tablets, USP: • 7.5 mg; yellow coloured, round, biconvex, tablets, debossed with "ISB" on one side and "C" on the other. • 15 mg; yellow coloured, round, fila beveled tablets, debossed with "CIPLA" on one side and "ISP" on the other.

4 CONTRAINDICATIONS

ContinuentIntentionation in the following patients:
 Known hypersensibility (e.g., anaphyticit reactions and serious skin reactions) to meloxicam or any components of the drug product Lie de Warnings and Precautions (5.7, 5.9).
 History of asthma, urtcaria, or other allergic-type reactions after taking asprin or other KSAIDs. Severe, sometimes fatal, anaphyticit reactions to KSAIDs have been reported in such patients (see Warnings and Precautions (5.7, 5.8)).
 Incasting and precautions (5.7, 5.8).
 Incasting and precautions (5.7, 5.8).
 Incasting and precautions (5.7, 5.8).

5 WARNINGS AND PRECAUTIONS

5.1 Cardiovascular Thrombotic Events 5.1 Cardiovascular Thoromotic Events Clinical traits of several COA2 societies and nonselective MSAIDs of up to three years duration have shown an increased risk of serious cardiovascular (CV) thoromotic events, including negracial infraction (MB) and stroke, which can be fatal based on more tables, it is used to a serious CV increased to the series is conferred by MSAID use appears to be similar in those with and whoult known. Videase or risk factors had a higher for the series of the series CV increased or risk factors had a higher factor with the series CV increased or the series first of series CV thoromotic risk has been observed most consistently at higher doses. To main after the development of Jusci events, through and barlies found main after that development of Jusci events, through and barlies found abute the symptoms of series CV events in StalD-readed patients, use the barest affective dose for the shortest duration possible. Physicians and patients about the symptoms of series CV events in StalD-readed patients for a dispute the symptoms of series CV events in StalD-readed patients for the rest one about the symptoms of series CV events and the steps to take if the creased

There is no consistent evidence that concurrent use of aspirin mitgates the increased risk of serious CV thrombotic events associated with NSAID use. The concurrent use of aspirin and an NSAID, such as meloxicam, increases the risk of serious gastrointestinal G events [see Warnings and Precautions (5:2)].

val events (see warling) and precautors (5.27). Statu. Post Concours, Afters Papasa Caff (CABG) Surgery Two large, controlled chical trails of a COX-3 selective NSAD for the treatment of pain in the first 10-14 days following CABG surgery found an increased incidence of myocardial infrarction and ströke. NSAIDs are contraindicated in the setting of CABG [see Contraindications (4)].

Post-MI Patients

Pail: M Patients Discretational studies conducted in the Danish National Registry have demonstrated that public to tradeal with the study. The patient have been applied to the strength of the strength of the strength of the strength of the study. The strength of the strength of the strength of parts in RSAD strength patients compared to 12 per 00 person years in the NSAD strength of the strength patients compared to 12 per 00 person years in the NSAD strength of the strength patients (the strength patient) was a per 100 person years in the NSAD strength of the strength patients compared to 12 per 00 person years in the NSAD strength patient. The strength patients (the strength patient) was a per 100 person years in the NSAD strength patient. The strength patients (the strength patient) was a strength with the strength patients who the used modelman in patients with a recent M unless the benefits are expected with a recent Min monitor patients for signs of cardiac is chema.

5.2 Gastrointestinal, Bleeding, Ulceration, and Perforation

3.2 Gastrointestinal, Bleeding, Ukceration, and Perforation NSMDs, including meloxicam, can cause serious gastrointestinal (d) adverse events including influenzation, Ibeeding, uschraften, and performation of the esophagis, stomach, occur at any time, with or without warning symptoms, in patients without these risk factors. treated with NSADS. Only one in five patients with out these risk factors. treated with NSADS. Only one in five patients with out develop a serious upper GI adverse event on NSAD therapy is symptomatic. Upper GI uters, gross bleading, or adverse event on USAD therapy is symptomatic. Upper GI uters, gross bleading, or adverse event on USAD therapy is symptomatic. Upper GI uters, gross bleading, or months, and had ubd 2-4% of patients treated for one year. However, even short-term NSAD therapy is not without risk. Relification of the outer blead of the orderologing at GI bleeding how used NSADs has a greater than 10-40 increased risk for developing at GI bleeding monothering is of origo torticotest exist, aspin, anticoaguitants, or selective seriorin requirate inhibitors USAD bio incluse the for duration of program hash status. Next USADS is include bleed and on or general health status. Next USANDs include the origo arrange of RNAD therapy concombing in a of oral controcter exist, aspin, naticoaguitants, or selective seriorin requirate inhibitors USADB biotest with advolve disease and poor general health status. Next USADB is included with used NADS biotest with advolve disease and for couplupatity are at increased risk of Gi beeding.

reik of G biedenig. Strategies to Momine the GI Bicks in NSAIP1-treated patients: • Use the lowest effective dosage for the shortest possible duration. • Joed administration of more than on NSAID at a time. • Joed administration of more than on NSAID at a time. • Joed administration of more than on NSAID at a time. • Joed administration of the NSAIP. • Remain alert for signs and symptoms of GI ulcration and bleeding during NSAID • Breath alert for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration and bleeding during NSAID • Breath active for signs and symptoms of GI ulcration active for signs and symptoms of GI

- therapy, 1 is a serious GI adverse event is suspected, promptly hilate evaluation and treatment, and discontinue melaxicam until a serious GI adverse event is ruled out.

 In the setting of concorntant use of low-dose aspirit for cardiac prophylaxis, montor patients more closely for evidence of GI bleeding [see Drug Interactions (7)].

5.3 Hepatotoxicity

Elevations of ALT or AST (three or more times the upper limit of normal (ULN) have been reported in approximately 1% of NSAU-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.

Unserver while ranked services of the warming signs and symptoms of hepatotoxicky (e.g. nauses, hepatotic services) and the service services and the service services and the like symptoms). If clinical signs and symptoms consistent with live disease develop, or systemic manifestations occur (e.g. existipation), existing and the patient (see Use in Specific Population (6.8) and Chical Amanda (Constant) (2.5).

5.4 Hypertension

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

course of therapy. 5.5 Heart Failure and Edema The Coxb and traditional NSAID Traditists' Collaboration meta-analysis of randomized controlled traits demonstrated an approximately two-fold increase in hospitalization compared to pickeb-tradeet patients. In a Drainh National Registry Study of patients with heart failure, ISAID use increased the risk of Mi, hospitalization for heart failure, and each.

and death. Additionally, full retention and edema have been observed in some patients treated with NSAIDS. Use of meloxican may blunt the CV effects of several therapeatic agents used to rest these medical conditors (e.g., our direct, ACE hishlows, or angiotension receptor blockers (AdBis) [Lee anging inferencies (7)]. Awold the use of meloxican in padents with severe heart falure unless the benefits are expected to outwegh the risk of worksmip heart falure. If meloxicam is used in patients with severe heart falure, monitor padents for signs of worksmip heart falure.

5.6 Renal Toxicity and Hyperkalemia Renal Toxicity

Long-term administration of NSAIDs, including meloxicam, has resulted in renal papilary necrosis, renal insufficiency, acute renal failure, and other renal injury. necross, renai insurincency, acute renal naure, and ourer renal injury. Renal toxicity has also been seen in patients in whom renal prostaglandins have a compensatory role in the maintenance of renal perfusion. In these patients, administration of an ISAID may cause a dose-dependent reduction in prostaglandin formation and, secondarity, in renal blood flow, which may precipitate overt renal

decompensation. Patients at greatest risk of this reaction are those with impaired renal function, dehydration, hypovolemia, heart falure, liver dysfunction, those taking diuretics and Acti hinbitors or ARBs, and the deviry. Discontinuation of NSAID therapy is usually followed by recovery to the pretreatment state.

The renal effects of meloxicam may hasten the progression of renal dysfunction in patients with preexisting renal disease. Because some meloxicam metabolites are excreted by the kidney, monitor patients for signs of worsening renal function. Excrete up the known monten particle to agits of worsening renormation. Correct volume known early and end particle of hypoxylemic patients prior to initiating meloxicam. Monitor renal function in patients with renal or hepatic impairment, heart falure, dehydration, or hypovolemia during use of meloxicam (see Drug Interactions (7)).

(7). On information is available from controlled clinical studies regarding the use of mediax can in patients with advanced real disease. Avoid the use of networking in the form of the studies of th

Hyperkalemia

<u>Hypersonna</u> Increases in serum potassium concentration, including hyperkalemia, have been reported with use of NSAIDS, even in some patients without renal inpairment. In patients with normal renal function, these diffects have been attributed to a hyporeninemic-hypoaldosteronism state.

5.7 Anaphylactic Reactions Meloxicam has been associated with anaphylactic reactions in patients with and without known hypersensitivity to meloxicam and in patients with apprint-sensitive asthma [see Contraindications (a) and Warnings and Precautions (5.8)]. Seek emergency help if an anaphylactic reaction occurs.

5.8 Exacerbation of Asthma Related to Aspirin Sensitivity

5.8 Exacerbation of Asthma Related to Aspirin Sensitivity A subpopulation of patterns with asthma may have aspirin-ensitive asthma which may include chronic minosinustic complicated by naial polypis; severe, potentially fatal toronchrospian; and point relations of a soft and don'the SSIDS. Because Cross-nochrospitations, may and an experimentary of the soft and and soft assist, networks and soft and soft assist, networks and soft assist patients, networks in contraindicated in patients with this form of aspiri sensitivity (see Contraindicators) (20). When mekokicam is used in patients with precessing asimption without home.

5.9 Serious Skin Reactions

3.2 Joint and the second se

5.10 Drug Reaction with Eosinophilia and Systemic Symptoms (DRESS)

3.24 Units Reschin March Stanophila and Systemic Symptomic Units.37 Drug Reschico with Estinophila and Systemic Symptomic Direction Sha be been reported in the symptomic Direction State State State State State State State State State Retrostanting. DRESS typical, although not erclassively presents with their rash-typinadenogative, and/or facial seeling. Direct ricital amarchitations may include hepatish, nephritis, hematological abnormalities, myocardits, or myosits. Sometimes symptomic of DRESS may resente an active vial infection. Sociophila is often present, may be howled. It is important to node that eavy manifestations of hypersensibility, such as fever or improdenogative, may be preserved event hough rash in not evident. If such as you symptoms are present, discontinue metoxicam and evaluate the patient mendiately

5.11 Fetal Toxicity

Permature Closure of Fetal Ductus Arteriosus Avoid use of NSAIDs, including metoxicam, in pregnant women at about 30 weeks gestation and attern NSAIDs, including metoxicam, increase the risk of premature closure of the fetal ductus arteriosus at approximately this gestational age.

or the relativistical set to basis a support integer to establishing the Opphylarizativistical and the support of the set of SADADs, including methods to objectly attransition or later in pregnancy my cause feat area of spharic contrainable to objectly attransition or later in pregnancy my cause feat area of spharic contrainable to objectly attransition or later in pregnancy methods of the set of the methods of the set of the which is of the set of the set of the set of the set of the which is of the set of the set of the set of the set of the which is of the set o

If NSAID treatment is necessary between about 20 weeks and 30 weeks gestation, il metoxicam use to the lowest effective dose and shortest duration possible. Consider ultrasound monitoring of annitoti fuld if metoxicam treatment extends beyond 48 hours. Discontinue metoxicam if oligohydrammiss occurs and follow up according to clinical practice fee Use is Specific Populations (a.1).

5.12 Hematologic Toxicity

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

NRINGAUNTO INTERNACIA: NSAIDS, including meloxicam, may increase the risk of bleeding events. Co-morbid conditions such as coagulation disorders or concomtant use of warfarin, other anticoagulants, antiplatekt agents (e.g., aspirin), section in resplate inhibitors (SSRIs) and serotonin noreginephrine reuptake inhibitors (SRRs) may increase this risk. Monitor these patients for signs of bleeding less *Dray* 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 these diagnostic signs in detecting infections.

5.14 Laboratory Monitoring

Because serious GI bleeding, hepatotoxicity, and renal injury can occur without warning symptoms or signs, consider monitoring patients on long-term NSAID treatment with a CBC and a chemistry profile periodically lise Warnings and Precautions (52, 53, 5.6).

Control of the end of the en

6.1 Clinical Trials Experience

Because clinical triak 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

Categorithis and Bhommado Affrids. Detending and Bhommado Affrids. In envolvem Phase 22 Cricks of vol distabase includes 10,122 OA patients and 1018 PA patients with mean 22 Cricks of vol 2008. Detending and the mean 2008 of the patients of the states of the patients of the states patients for at least for norths and to 321 patients for at least one year. Approximately 10,500 of these patients were treated in ten placebe and/or active-controlled discover of the patients were treated in ten placebe and/or active-controlled subsection of the patients were the most frequently of a were were then most frequently reported adverse events in all treatment groups across meaxiam true.

A 2-week multicenter, double bind, randomized trial was conducted in patients with osteoarthrits of the knee or hig to compare the efficacy and safety of meloxicam with placebo and with an active corror 1 in 0.2 week multicenter, double bind, randomized trial effect of meloxicam 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 Peripheral				
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
¹ WHO preferred terms edema, edema dependent, edema peripheral, and edema legs combined				

²WHO preferred terms rash, rash erythematous, and rash

Table 1b Adverse Events (%) Occurring in ≿ 2% of Meloxicam Patients in two 12-Week Rheumatoid Arthritis Placebo-Controlled Trials 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 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 Conditions			
Influenza-like illness ²	2.1	2.9	2.3
Infection and Infestations			
Upper respiratory tract infections-	4.1	7.0	6.5
pathogen class unspecified1		7.0	0.0
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
¹ MedDRA high level term (preferred terms): dyspeptic signs			
and symptoms (dyspepsia, dyspepsia aggravated,			
eructation, gastrointestinal irritation), upper respiratory tract			
infections-pathogen unspecified (laryngitis NOS, pharyngitis			
NOS, sinusitis NOS), joint related signs and symptoms (arthraigia, arthraigia aggravated, joint crepitation, joint			
(arthraigia, arthraigia aggravated, joint crepitation, joint effusion, joint swelling)			
² MedDBA preferred term: pausea, abdominal pain NOS			

² MedDRA preferred term: nausea, abdominal pain NOS, influenza-like ilness, headaches NOS, and rash NOS

The adverse events that occurred with meloxicam in \geq 2% of patients treated short-term (4 to 6 weeks) and bng-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 Osteoarthrikis Trials

	4 to 6 Weeks Contro		6 Month Controlled	
	Meloxicam 7.5 mg daily		am Meloxicam ailv7.5 mg dailv	Meloxicam 15 mg dail
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
Dyspeosia	3.8	7.4	8.9	9.5
Flatulence	0.5	0.4	3.0	2.6
Nausea	2.4	4.7	4.7	7.2
Vomiting	0.6	0.8	1.8	2.6
Body as a Whole				
Accident household	0.0	0.0	0.6	2.9
Edema ¹	0.6	2.0	2.4	1.6
Pain	0.9	2.0	3.6	5.2
Central and Peripheral Nervous Sy	stem			
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 ²	0.3	1.2	3.0	1.3
Urinary				
Micturition frequency	0.1	0.4	2.4	1.3
Urinary tract infection	0.3	0.4	4.7	6.9

Urinary tract infection ¹WHO preferred terms edema, edema dependent, edema peripheral, and edema legs combined ²WHO preferred terms rash, rash erythematous, and rash maculo-papular combined

Combined Higher doses of meloxicam (22.5 mg and greater) have been associated with an increased risk of serous GI events; therefore, the daily dose of meloxicam should not exceed 13 mg. *Pediatrica Pediatrica Pediatr*

The following is a list of adverse drug 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, malake, syncope, weight decrease, weight increase
Cardiovascular	angina pectoris, cardiac falure, hypertension, hypotension, myocardial infarction, vascultis
Central and Peripheral Nervous Sys	tem convulsions, paresthesia, tremor, vertigo
Gastrointestinal	colits, dry mouth, duodenal ulcer, eructation, esophagits, gastric ulcer, gastriks, 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, bulous eruption, photosensitivity reaction, pruritus, sweating increased, urticaria
Special Senses	abnormal vision, conjunctivitis, taste perversion, tinnitus
Urinary System	albuminuria, BUN increased, creatinine increased, hematuria, renal faliure

Common particle in the second second

7 DRUG INTERACTIONS

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

Table 3 Clinically Significant Drug Interactions with Meloxicam

Drugs that	Interfere with Hemostasis
Clinical 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 ether drug abine. Serotonin release by platelets plays an inportant role in hemostasis. Case-control and cohort epidemiological studies showed that concomiant use of drugs that interfere with serotonin respatie and an HS4D may portunitate the risk of beeling more than an HSAD abine.
Intervention:	Montor patients with concomtant use of metoxicam with anticoagulants (e.g., warfarin), antiplated egats (e.g., aspirin), selective serotonin reuptake inhibitors (SSRIs), and serotonin norephephrine reuptake inhibitors (SNRIs) for signs of bleeding (see Warnings and Precautions (5.22).
Aspirin	F
Clinical Impact:	Controled clinical studies showed that the concomitant use of NSAIDs and analgesic does of aspirh 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 adversere actions as compared to use of the NSAID alone [see Warninos and Precautions (5.21).
	Concombant use of medoxicam and low dose asprin or analgesic doses of asprin is not generally recommended because of the increased risk of bileeding (see Warnings and Precautions (5.12)). Metoxicam is not a substitute for low dose asprin for cardiovascular protection.
ACE Inhibit	ors, Angiotensin Receptor Blockers, or Beta-Blockers
Clinical Impact:	 NSAIDs may diminish the anthypertensive effect of angotensin converting enzyme (AEE) inhibits, angiotensin receptor blockers (ARBs), or beta-blockers (including propranolo). In patients who are elderly, volunci degited (including those on diuretic therapy), or have renal inpairment, coadministration of an MSAID with ACE inhibitors and ABBs may result in deterioration of renal function, including possible acute renal failure. These effects are usually reversible.
Intervention:	 During concomitant use of metoxicam and ACE whittons, ARBs, or beta-backers, and another back presence to ensure that the desired bload because of the another backers and ace whittons and ace whittons proteins who are defery, volume desired, or have impaired real function, monitor for signs of worsteining real function [see Warnings When these drugs are administered concomitantly, patients should be adequately hydrafed. Assess renal function are be beginning of the concomitant research and periodically threader.
Diuretics	L
Clinical Impact:	Clinical studies, as well as post-marketing observations, showed that NSAIDs reduced the nativariset certed of bog directics (e.g., furosemide) and thisized auretics in some patients. This effect has been attributed to with furosemide agents and melosicam have not demonstrated a reduction in nativartext effect. Furosemide single and multiple dose pharmacodynamics and pharmacointextes are not affected by multiple

does of melocitarius and premise the sense of the sense o

	including antihypertensive effects [see Warnings and Precautions (5.6)].
Lithium	
Clinical Impact:	NSAIDs have produced elevations in plasma lithium levels and reductions is renal lithium clearance. The mean minimum lithium concentration increases 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.
Methotrex	ate
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.
Cyclosporir	ie .
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.
NSAIDsand	Salicvlates
Clinical Impact:	Concomitant use of meloxicam with other NSAIDs or salicylates (e.g., diflunisal, salsalate) increases the risk of GI toxicity, with little or no increas in efficacy (see Warnings and Precautions (5.2)].
Intervention:	The concomitant use of meloxicam with other NSAIDs or salicylates is not recommended.
Pemetrexe	d
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, inpatients with renal impairment whose creatinice clearance ranges from 45 to 79 mL/min monitor for myebosuppression, renal and G1 tox/cty. Patients taking meloxicam should interrupt dosing for at least five days before. The day of, and two days following pemetrexet admitistration. Jaministration on meloxicam without pemetrexet admitistration administration on meloxicam without pemetrexet admitistration.

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy Risk Sum mary

teach sources provides the second second

Premature Closure of Fetal Ductus Arteriosus Use of NSAIDs, including meloxicam, at about 30 weeks gestation or later in pregnancy increases the risk of premature closure of the fetal ductus arteriosus.

Olgohydrannios/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.

With class to treat retai opularition example oblight professional real impartments. Botas from observational studies regarding potential embryofetal risks of NSAD use in women in the first or second trimsters of pregnancy are inconclusive. In animal reproduction studies, embryofetal death was observed in rats and rabbs totaled during the profield of organopare and a constraint of an isome explosion totaled during the profield of pregnancy are used to the profield of any totaled during the profield of pregnancy are used in constraints increased incidence of sepath heart defects were observed in rabbs to treat both. The the and post-natal reproduction studies, there was an increased incidence of dystocia, not be treatogener, effects were observed in rabbs to treated with motellow. The organized regarding and any postage in rats and rabbs to treated with motellow. The distance of the regression and any postage in rats and rabbs to treated with motellow. The distance regarding are also and any postage in the any state of the motellow and any regarding and any postage in the set was an increased incidence of dystocia, to the treatogener, effects were observed in rats and rabbs to treated with motellow and and organogenesis at an oral dose equivalent to 2.5 and 26-times the MRHD [are Data]. Based on animal data, postage indinario hystines is hinkings, such as metocia, man shown to have an important role in feda klinge development. In published animal studies, postagenesis role and incidence of dystocia, and to have an important role in feda klinge development. In published animal studies, postagenesis and any brites in thirdings the state of the indicated to have an important role in fred klinge development. In published animal studies, postagenesis and any brites and role based role to the role klinger development when administered at clincially relevant doses.

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 automes. In the U.S. general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnancies is 2% to 4% and 15% to 20%, respectively. Clinical Considerations

Fetal/Neonatal Adverse Reactions

Premature Closure of Fetal Ductus Arteriosus: 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 HSAID is necessary advoir 20 weeks gestation or lister in preparatory, hit has not to the based information and an advoired in the strength of the stre

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. Data

Human Data Premature Closure of Fetal Ductus Arteriosus: Published Rereature 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.

The the programs i may dependent of the contract of the contra

will investe proceedings, such as exchange drain usion of pages. Methodobigid impairments of these postantiseting studies and reports include lack of a control group; linked information regarding dose, duration, and timing of drug exposure; and concominant use of other medications. These imitations preclude establishing a relable estimate of the risk of adverse fetal and neonatal outcomes involved maternal KSAD use Decause the published safety data on investal outcomes involved mostly preterm inflants, the generalizability of certain reported risks to the full-term inflance exposed to ASUBs through material use is uncertain. Animal Data

Annu Data Mexicam was not teratogenic when administered to pregnant rats during fetal organogenesis at oral doses up to 4 mg/kgd/wj. (2-fold greater than the MRH0 d 15 mg/ mexicam based on BSA companies). Administration of metalscart to pregnant, the heart at an oral dose of 60 mg/kgd/wj. (78-fold greater than the MRH0 based on BSA comparison). The net effect well was 20 mg/kgd/wj. (78-fold greater than the MRH0 based on BSA conversion). In rats and rabbis, emphysichtafly occurred at oral mexicam dose of 10 mg/kgd/wj. (78-fold greater than the MRH0 based on BSA conversion). In rats and rabbis, emphysichtafly occurred at oral mexicam dose of 1 mg/kgd/wj. (78-fold greater than the MRH0 based on BSA conversion). In rats and rabbis, emphysichtafly occurred at oral mexicam dose of 1 mg/kgd/wj. (78-fold greater than the MRH0 based on GSA comparison) when administered throughout organogenesis.

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

8.2 Lactation

Risk Summary

There are no human data available on whether meloxicam is present in human mik, or on the effects on breastfel infants, or on mik production. The developmental and health benefits of breastfeeding should be considered along with he mother's chical need for meloxicam and any potential adverse effects on the breastfeel infant from the meloxicam or from the underlying maternal condition.

Interaction of the one of the one

8.3 Females and Males of Reproductive Potential

Infertility

Females

Females Based on the mechanism of action, the use of prostaglandin-mediated NSADs, including mdox.cam, may deby or prevent rupture of ovarian follows, which has been associated animistration of prostaglandin synthese inhibitors has the posterial to disrupt. prostaglandin-mediated followir rupture required for ovalution. Small studies in women rustad with NSADs have also shown are versible delay in ovalution. Conside withdrawal of NSADs, including mdox.cam, in women who have difficulties conceiving or who are undergraph mediation of identity.

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

6.3 vertant v use Eiterley 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 defly patient outweight these potential risks, start dosing at the low end of the dosing range, and monitor patients for adverse effects [see Warnings and Precoutions [5, 25, 25, 35, 45].

6.6 Repairle inpairment No dose adjustment is necessary in patients with mild to moderate hepatic impairment Patients with search explaid; majamarin heur not been adequately studied. Since metoxicam with cauton in patients with hepatic impairment [see Warnings and Precautions (5.3) and Criciael Paramacology (2.2.3).

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 metoxicam in subjects with severe renal impairment is not recommended. In patients on hemodalaysis metoxicam should not exceed 7.5 mg per day. Mebxicam is not dialyzable [see Dosage and Administration (2.2) and Clinkal Pharmacology (2.23)].

10 OVERDOSAGE

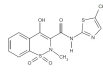
Symptons following acute NSAID overdosages have been typically limited to lethargy, drowsiness, nausea, vomking, and epigastrc pain, which have been generally reversible with supportive care Gastrontestand bleeding has occurred. Hypertension, acute renal failure, respiratory depression, and coma have occurred, but were rare [see Warnings and Precautions [5.1, 5.2, 5.4, 5.6].

and Precutions (5.1, 5.2, 5.4, 5.0). Manage patients with symptomizet and supporter care following an KSAD overall symptomizet There are no specific antiother. Consider emesis and/or activated charcoll (60 to 100 grams in adults). To grams per kg of oddy weight in pedaticer, patients) and/or somotic cathartic in symptomizic patients seen within four hours of ingestion or in address with a large overdisage (5.1 to 10 times the recommended disage), Forced high protein binding of urine, hemodalysis, or intemperfusion may not be useful due to high protein binding concerns the hemodalysis, or intemperfusion may not be useful due to high protein binding concerns the hemodalysis.

There is imited experience with meloxicam overdosage. Cholestyramine is known to accelerate the clearance of meloxicam. Accelerated removal of meloxicam by 4 g oral does 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).

11 DESCRIPTION

• co-store room Mookam is a nonsteroidal anti-inflammatory drug (NSAID). Each tablet contains 7.5 mg or 15 mg meloxicam, USP for oral administration. Neloxicam is chemically designated as Alyddroxy?- methydrix/Hostpvi/Jzrt12-brandblame3-achamide1.1-dioxide. The molecular weight is 351.4. Its empirical formula is Caj4hgMg052 and k has the following structural formula:



Meloxicam 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. Meloxicam has an apparent partition coefficient (log P)_{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 celulose, lactose anhydrous, colloidal silicon dioxide, sodium citrate dihydrate, mannei im strongen.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action Meloxicam has analgesic, anti-inflammatory, and antipyretic proper The mechanism of action of meloxicam, like that of other NSAIDs, is not completely understood but involves inhibition of cyclooxygenase (COX-1 and COX-2). Metokcam is a potent inhibitor of prostaglandin synthesis in vitro. Metokcam concentrations resched during threepy have produced in vivo effects. Prostaglandins sensitize afferent nerves and potentiate the acton of bradykin in inducing pain in animal modes. Prostaglandins are metidators of inflammation. Because metokcam is an inhibitor of prostaglandins are metidators of inflammation. Because metokcam is an inhibitor of prostaglandins are unditors of antimation. Because metokcam is an prostaglandins in peripheral issues.

12 3 Pharmacokinetics

Abacettain The absolute biowniability of metoxic am capsules was 89% following a single oral dose of 30 mg compared with 30 mg // bolas injecton. Following single intravenous dose, index of a single with a single of 1.5 mg // absolute single s s have been shown to be bioequiv ent to meloxicam tab

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

Steady State			Single Dose					
		Healthy male adults (Fed) ²	Elderly males (Fed)2	Elderly females (Fed) ²	Renal failure (Fasted)	Hepatic insufficiency (Fasted)		
(%CV								
		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)		19 (43)	11 (44)		
V2/f 4	[L]	14.7 (32)	15 (42)	10 (30)	26 (44)	14 (29)		

¹The parameter values in the table are from various studies

2not under high fat conditions

³Meloxicam tablets 4Vz/f =Dose/ (AUC•Kel)

Food and Antacid Effects

rook are what at circles. A definition of the second seco

Distribution

Distribution The mean volume of distribution (Vss) of meloxicam is approximately 10 L Meloxicam is -99.4% bound to human pistema proteins (primarly abumin) within the threspectic does range. The faction of protein binding is dependent of drog concentration, over the clinkally relevant concentration range, but decreases to -99% in patients with real decase. Meloxicam penetation in high nummar of bloot cells, the road boxing. S less than decase. Meloxicam penetation in high nummar of bloot cells, the road boxing. S less than ware present as unchanged meloxicam. Meloxicam concentrations in synowid hids, after a single or of does, range from 40% to 50% of those in plasma. The firef raction in synowid hidd as Compared to plasma. The significance of this penetation is unknown.

Elimination Metabolism

Metabolism Metabolism Metakcam is extensively metabolised in the liver. Metakcam metabolites include 5-sociation of an intermediate metabolite 3-hydronymethyl metakcam withich is also excreted to a laisex extent (%) of doars, in vitro studies distante that CPZC9 (cylocchrome PASD metabolities ensyme jalkys an important role in this metabolit. By notably responsible for the other two metabolities with court for 10 km and % of the administered doar, respectively. All the four metabolities are not known to have any in vice pharmacological actually. . Excretion

Excretion Moxicam excretion is predominantly in the form of metabolites, and occurs to equal extents in the urine and feccs. Only traces of the unchanged parent compound are excreted in the urine [0.2%] and recel; 16(%). The extern of the urinary excretion was secreted in the urine [0.2%] and recel; 16(%). The extern of the urinary excretion found in urine in the form of metaxicam, and the 3-hydroxymethyl and 3-Carboxy metabolites, respectively. There is significant bilary and/or entred is secretion of the drug. The was demonstrated when on al administration of holestyramine following a single V does of metaxicam decreased the AUC of metakcam p30.

The mean elimination of H⁻¹fe (t₁), 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 mL/min.

<u>Pediatric</u> After single (0.25 mg/kg) dose administration and after achieving steady state (0.375 mg/kg/skg), there was a general trend of approximately 30% lower exposure in younger patients (2 to 6 years old) as compared to the older patients (7 to 16 years old). The older patients had hockican exposure similar Single doso 1 sg/kg/t reduced (steady state) to Those in the adult patients, when using ALC values inormalized to a dose of hind few sizes (2 to 10) and 12.0 hours (3.0) for the 2 to 6 year old patients, energy to 16 year old patients, respectively.

To year our patients, respectively. In a covariate analysis, utilizing population pharmacokinetics body-weight, but not age, was the single predictive covariate for differences in the metexicam apparent oral plasma clearance. The body-weight normalized apparent oral clearance values were adequate predictors of metoxicam exposure in pediatric patients.

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

Genation: Electry males (ze 55, years of gel) simbled metoxical planta concentrations and gel) failed a 27% higher AUC, and 32% higher Cauca, as compared to younge females (5 5) years of gel) and a 27% higher AUC, and 32% higher Cauca, as compared to younge female (5 5) years of gel) after body wegls normalization. Begies the Increased total concentrations in the elderly females, the adverse event profile was compared to younge for the adverse of the adverse event profile was compared for body compareton to define ymate platers.

Young females exhibited slightly lower plasma concentrations relative to young males

After single doses of 7.5 mg meloxican, the mean elimination half-life was 19.5 hours for the female group as compared to 23.4 hours for the male group. At steady state, the data ware similar (13.7 hours 24.1 A hours). This pharmacoichnet difference due to gender is likely to be of little chical importance. There was linearly of pharmacoichnetics and no apprecisible difference in the Cmar of Tmax across genders.

Hepatic Impa

majesis: impairment Teologing a single bining does of metodocate makes ways no marked afference in joinney footbarred participation and the single bining bining bining bining bining II) hepsite impairment compared to healthy volunteers. Protein binding of metoxem was not affected bin phasitic impairment. No dospase adjustment is necessary in palatests was not affected bining bini

Renal Imnairn

Renal Impairment Metockam pharmatokinetics have been investigated in subjects with mild and moderate renal impairment. Total drug pissma concentrations of metockam decreased and total clearance of metockam in increased with the darge of renal impairment while the ALC clearance of metockam increased with the darge of renal impairment they have mpairment may be due to increased fraction of unbound metockam which is available for hepatic metoblem and subsequence excretion. No dosea eguistment is mecsarry in patients with mild to moderate renal impairment. Patients with severe renal impairment (5.6) and Use in Specific Populations (8.7).

Hemodialysis

International and the second s

Drug Interaction Studies

Again: When HSAIDs are administered with aspin, the portexh binding of HSAIDs were related, although the charance of the HSAID was not ablend. When metadoxican is administered with aspin (1000 mg three times day) to healthy volunteers, t tended to increase the AUC (1054) and C_{SAI} (24%) of meta/carm. The clinical significance of this interaction is not known. See Table 3 for clinical significant drug interactions of NSAIDs with aspin (1800 mg interactions (7/s).

with aspire issee Drug Interactions (7)). Cobestyrainer Pertextament for four days with choisstyraine significantly increased the clearance of metoxicam by 50%. This resulted in a decrease in 12, 7 rom 13.2 hours to 12.5 hours, and 35% reduction ha AULC. This suggests that existence of a rediculation pathway for metoxicam in the gastroiredstimil tract. The clinical relevance of this interaction has to been stabilisher. Cimeditine: Concomitant administration of 200 mg cimetistine four times daily did not alter the single docs harmacokinets: Cal 30 mg metoxicam.

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 vitro testing found no protein binding drug interaction between digoxin and meloxicam.

testing town no protein binding drug interaction between digoxin and meoxicam. Libraim: In a study conducted in healthy subjects, mean pre-doces Ithum: concentration and ALU were increased by 21% in subjects receiving Ithum doses ranging from Bdds 1027 on ywice adjiw Min ebokam. It is no gO every day as compared to subject receiving Ithum alone [see *Drug Interactions (T)*]. Mechoreaset A study in 13 hematodia and arthis (RA) patients evaluated the effects of multiple doses of metoxicam on the pharmacokinetics of methodroxate taken once weekly. Metoxicam do In this was a significant effect on the pharmacokinetics of single human serum binding alete [see *Drug Interactions (T)*].

human serum binding sites [see Drug Interactions (7)]. Warfarit: The effect of mediciarian on the anticoagulant effect of warfarin was studied in a group of healthy subjects receiving daily closes of warfarin that produced an NR (international Kornaulice Ratio) between 12 and 1.8. In these subjects, meldioxican did determined by profitoribant time. However, one subject showed an increase in INR from 15 0.2.1. Catalon houdd be used them animitatering medication with warfars since paleties to warfarin may experience changes in NR and an increased risk of bleeting complications when an even medication is introduced [see Drug Interactions (7)].

13 NONCLINICAL TOXICOLOGY 13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

13.1 Carcinogeness, Mutageness, Impartment of Yertiky Carcinogenesis There was no increase in tumor incidence in long-term carchogenicity studies in rats (186) weeks) and mice (39 weeks) and individual and and a sets up to 0.8 (186) weeks) and mice (39 weeks) and animitate of mice and a sets up to 0.8 (186) weeks) and mice (39 weeks) animitate of mice and a set of the mice and the sets of the mice and the sets of the mice and the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the sets of the sets of the sets of the mice and the sets of the mice and the sets of the mice and the sets of the sets o

Mutagenesis

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

Impairment of Fertility

MedoxLam 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- greater, respectively, than the MRHD based on BSA comparison).

14 CLINICAL STUDIES

14.1 Osteoarthritis and Rheumatoid Arthritis The use of melokiam for the treatment of the signs and symptoms of osteaarthrits of the inner and hip was evaluated in a 12-week, double-bind, controlled trial. Melokicam (375 mg. 73 mg. and 15 mg daily was compared to placed. The four primary endpoints were investigators global assessment, patient global assessment, patient placed assessment, and the IMOMES core (as ela-diminister of quotischinaria addressing daily showed significant improvement in each of these endpoints compared with. 5 mg divertion.

The use of meloxicam for the management of signs and symptoms of osteoarthritis was evaluated in six double-bind, active-controlled triais outside the U.S. ranging from 4 week's to 6 montris' duration. In these triais, the diffectory of meloxicam, in doese of 7.5 mg/day and 15 mg/day, was comparable to proxicam 20 mg/day and dicb/fenac SR 100 mg/day and 15 mg/day.

migray and consistent with the effects years in the U.S. Two and the migray and dictifences 3.11 to the use of metackarshi to a 15 restantment of the signs and symptoms of metackarshi to a 15 migray of the migra

14.2 Juvenile Rheumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course

The use of meloxicam for the treatment of the signs and symptoms of pauciarticular o polyarticular 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.

We evaluated in two 12-week, sound-time, par element, par element, market element and the sound of the sound in the sound of the sound in the sound of the sound in the sound of the sound

The effract analysis used the ACR tealtrix 30 responder definition, a composite of parent and investigator assessments, counts of active joints and joints with limited range of motion, and enfrances and the activity of a strate points and points with limited range similar in all three groups. In both studies, and no difference was observed between the moticity and groups.

16 HOW SUPPLIED/STORAGE AND HANDLING Meloxicam tablets, USP 7.5 mg are yelow coloured, round, biconvex tablets, debos: with "156" on one side and "C" on the other. Meloxicam tablets, USP 15 mg are yellow coloured, round, flat bevelled tablets, debossed with "CIPLA" on one side and "159" on the other.

with "CIPLA" on one side and "159" on the other. Meloxicam tablets, USP 7.5 mg are available as follows: NDC 0615-8040-39 Bistercards of 30 NDC 0615-8040-30 Unit dose boxes of 30 NDC 0615-8040-30 UTIL GOSC GOALS 2. Meloxicam tablets, USP 15 mg are available as follows: NDC 0615-8124-39 Bilstercards of 30

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. dc00mpattles each prescription togeneter. 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 pair, shortness of breath, weakness, or slurring of speech, and to report any of these symptoms to their healthcare provider immediately [see Warnin and Precadultors [5,1].

and Precautons (5.1). Gastrahestnäh Beeling, Likeration, and Perforation Advise patients to report symptome of ukeration and bleeding, including epigastric for consolitation used for the object of the object of the object of the increased risk for the signs and symptoms of Gi bleeding Isee Warnings and Precautors (5.2).

Instantional Control (1997) and Control (1997) a

Heat failure and Edema Advise patients to be alert for the symptoms of congestive heart failure including shortness of breasth, unexplained weight gain, or edema and to contact their heathcare provider if such symptoms occur [see Warnings and Precautions (5.5)].

proved is source and a section of the source of the source

Serious Skin Reactions. Including DRESS Advise patients to stop meloxicam immediately if they develop any type of rash and to contact their healthcare provider as soon as possible (see Warnings and Precautions

(5.9,5.10)].

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

Populations (8.3). Facil Toxicity Inform pregnant women to avoid use of metoxicam and other NSAIDs starting at 30 weeks gest inform pregnant women to avoid use of metoxication of the facil ducta. Second Secon

Additional and a second and a second second

Use of NSAIDs and Low-Dose Aspirin Inform patients not to use low-dose aspirin concomitantly with mebxicam until they talk to their healthcare provider (see Purg) Interactions (7)]. Manufactured by: Cipia, Ltd., Kurkumbh, India

Manufactured by: Cipa, Eds., Manufactured for: Cipa USA, Inc. 10 Independence Boulevard, Suite 300 Warren, NJ 07059 Revised: 5/2021 21089452

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 hannen early in treatment and may increase:

with increasing doses of NSAIDs
 with longer use of NSAIDs

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

artery bypass graft (CABG). Work taking MSLOB after a recent heart attack, unless your healthcare provider tells you to. You may have an increased risk of another heart attack in Increased risk of beleding, uncertained in the source of the source of the esophagus (Lube leading from the mouth to the stomach), stomach and interesting: the store of the source of the

that may cause death
 The risk of getting an ulcer or bleding increases with:
 back history of stomsch ulcres; or stomach or intestinal bleding with use of
 ISADs
 Labing medicines called "corticosteroids", "anticoagulants", "SSRIs", or "SNRIs"
 Increasing does of ISADs
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smoking drinking alcohol older age poor health advanced liver dise

advanced wer 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, sweling, 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?

If you have had an asthma attack, hives, or other allergic reaction with aspirin or any other NAIDs.
 right before or after heart bypass surgery.

Before taking RISDS, tell your healthcare provider about all of your medical conditions, including if you: have have high blood pressure. have estimate the pressure in the pressure of the state of the state of a represent or plan to become pregnant. Taking NSAIDs at about 20 weeks of pregnancy or later may have uniform baky. If you need to take ISAIDs for healthcare provider may need to motion the about 20 weeks of pregnancy or later may have uniform taken to the state of healthcare provider may need to motion the amount of taken to your baby. You should not take NSAIDs after 30 weeks of pregnancy.

Tell your healthcare provider about all of the medicines you take, includ prescription or over-the-counter medicines, vitamins or herbal supplem NSIDs and some other medicines can iteract with each other and cause serious effects. Do not start taking any new medicine without taking to your healthcare provider first.

Besthore norvider first:
 What are the possible effects of NSAIDs?
 What are the possible effects of NSAIDs?
 NSAIDs can cause serious side effects, including:
 See "What is the most important information I should know about medicines
 caled Monatcredid Anth-Informatory Drugs (NSADs)?"
 * new or works high blod pressure
 * lever problems including keffect and
 * life-treatening alerge reactions
 * life-treatening alerge reactions
 * Other side effects of MSAIDs include: stomach pain, constipation, darrhea, gas,
 heartfund inclusae, writing, and dizzness.

Restaulin, hasse, vontang, and aczess. Get energency heb pight away 'I you get any of the following sympton - shortness of breath or trouble breathing weakings in one part or side of your body - sizerd speech - swelling of the face or throat

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

nausea
 more tired or weaker than usual
 diarrhea
 kthing

liarrhea :ching :our skin or eyes look yellow ndigestion or stomach pain

indigetation or stomach pain
 Unive symptoms
 there is blood in your bowel movement or it is black
 vomit blood
 there is blood in your bowel movement or it is black and sticky like tar
 unsual weight or blacks with fever
 skin ratio or blacks with fever

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 pharmacsit about NSAIDs.

heathcare provider or pharmackit about NSADS. Call your dots for markical about call of effects. You may report side effects to FDA at 1.800-FDA-1088. Other Minimum Report ISADE Cases Beeling the beins, stomack, and institute. Aspin can also cause used in the stomach and intestines. Some NSADE are sold in lower does without a prescription (over-the-counter). Takk to your heathcare provider before using over-the-counter NSADE 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 Additional 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.

ham them. If you would like more information about NSAIDs, tak with your healthcare provider. You can ask your pharmactic or healthcare provider for information about NSAIDs that is written for health proteinsional. This Metication Guide has been approved by the U.S. Food and Drug Administration. Manufactured by C.J., Litt, Arkinnah, India

Manufactured for: Cipla USA, Inc. 10 Independence Boulevard, Suite 300 Warren, NJ 07059

. sed: 5/2021

PRINCIPAL DISPLAY PANEL - 7.5 MG

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MELOXICAM meloxicam tablet

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		Ingr	edient Name				Basis of	Strengt	h	Strengt
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Ir	nactive Ingr	edients								
		AIME. ETITIC	Ingredient	Name					S	trength
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c	ELLULOSE, MIC	ROCRYSTALLI	NE (UNI: OP1R320	061U)						
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CELLUGSE, MICROCRYSTALLINE (UNI: 0718320510) MAGNESIUM STEARATE (UNI: 70077M630) SODIUM STEARATE (UNI: 00725X07G) Product Characteristics Color yellow Score no score Shape ROUND Size 10mm

Flavor	Imprint Code	CIPLA:159
Contains		
Packaging		

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 Package Description
 Marketing Start
 Marketing End Date

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Establishment Name Address ID/FEI Business Operations NCS HealthCare of KY, LLC dba Vargard Labs (050032743) repark(0515-8040, 0515-8124) Revised: 3/2023 NCS HealthCare of KY, LLC dba Vangard Labs