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

Meloxicam Tablets USP Initial U.S. Approval: 2000

- stitul U.S. Approvab. 2000

 MARNING CARDIOVASCULAR and GASTROMTESTMAL RISKS

 SARADING CARDIOVASCULAR and GASTROMTESTMAL RISKS

 PACKAGE Company of the Company o

Melouicam tablets are a non-steroidal anti-inflammatory drug indicated for
• Oznourthritis (OA) (1.1)
• Resumated Arthritis (RA) (1.2)

Use the bisest effective dose for the stortest duration consistent with individual treatment goals for the
0.60 (2.7) and 8.6. (2.7) and 9.6. (2.7) and 9.6

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

CONTRAINDICATIONS
 Known hypersensibility (e.g., analphylactioir reactions; and serious side reactions) to melosicam (4.1)
 Histopy of action, utercain, or other lisable allegic byte reactions after taking sayline or other NSABil (1.1)
 Use during the peri-operative period in the setting of coronary artery bypass graft (CABG) surgery (4.2)

Use during the perison perison because the setting of contrary artery bysass grift (LEGB) ungrey (4.2)
 WARMINGS AND PRECAUTIONS
 Particular and the perison of the

Red eletteron and celema. Should be sever was creative in preduction. The electry, those fined appellary records and other wren's light with long term such us the shirt custion in the electry, those or any celeman in articipants. The use of melasticant in patients with sever recall impairment is not consommedide (3.6). The electric control of the electric contr

Most common (a5% and grant the pictories) abdress events in adults are districts, upper receptionly rest infections, disposing, and otherwise place to the pictories of the

www.Ma.gorimedwatch.

DNUG INTERACTIONS

- Coccombast use of resistance and appire is not generally recommended because of the potential of

- Coccombast use of resistance and appire is not generally recommended because of the potential of

- Coccombast use with molecular necessaries interest patients about (2).

- Coccombast use with National Processaries patients about (2).

- Coccombast use with National Processaries patients about (2).

- Coccombast use with National Processaries patients about (2).

- Coccombast use with National Processaries patients about (2).

- Coccombast use with National Processaries patients and National Processaries (2).

- Coccombast use with National Proce

FULL PRESCRIBING INFORMATION: CONTENTS* WARNING: RISK OF CARDIOVASCULAR AND GASTROINTESTINAL EVENTS 1. INDICATIONS AND USAGE

- 1.1 Osteoarthritis (OA)
 1.2 Rheumatoid Arthritis (RA)
 2. DOSAGE AND ADMINISTRATION
 2.1 General Instructions
- 2.2 Osteoarthritis
 2.3 Rheumatoid Arthritis
 3. DOSAGE FORMS AND STRENGTHS
 4. CONTRAINDICATIONS

- 4.1 Alloys Reactions
 4.2 Caronay Supple Reactions
 5.2 Cardiovascular Thrombotic Events
 5. Cardiovascular Thrombotic Events
 5. Cardiovascular Thrombotic Events
 5.3 Hepain: Effects
 5.3 Hepain: Effects
 5.4 Hypertension
 5.5 Congestive Heart Failure and Edema
 5.5 Congestive Heart Failure and Edema
 5.5 Congestive Heart Failure and Edema
 5.5 Analysis Reactions
 5.3 Autoria Supplementaria Supplementari

- 5.13 Use in Patients with Pre-ex 5.14 Montoring 6. ADVERSE REACTIONS 6.1 Clinical Trials Experience 6.2 Post Marketing Experience 7.1 ACE-inhibitors 7.1 ACE-inhibitors 7.2 Asprim 7.3 Durretics 7.4 Montories and Patients 7.5 (Cyclosporine 7.5 Vectories)

- Cyclosporine Warfarin SE IN SPECIFIC POPULATIONS

- 13.1 Pregnant Delivery
 13.2 Labor and Delivery
 13.3 Nursing Mothers
 14.7 Pediatric Use
 15.4 Pediatric Use
 15.6 Hepatic Impairment
 15.7 Renal Impairment
 16.0 VOENDOSAGE
 16.0 DESCRIPTION
 16.1 CLINICAL PHARMACOLOGY
 12.1 Mechanism of Action
 12.2 Pharmacodynamics

- CLINICAL PHANA--2.1 Mechanism of Action
 2.2 Pharmacolynimins
 2.2 Pharmacolynimins
 3.1 Carriogenesis, Mutagenesis, Impairment of Fertilly
 13.1 Carriogenesis, Mutagenesis, Impairment of Fertilly
 13.1 Carriogenesis and Re

- 17. PATIENT COUNSELING INFORMATION
 17. Medication Guidentes
 17.3 Gastrontestral Effects
 17.3 Gastrontestral Effects
 17.4 Rejactions(Line)
 17.5 Allevers-Skin Reactions
 17.5 Allevers-Skin Reactions
 17.6 Effects
 17.7 Anaphylication effect of the state of

WARNING: RISK OF CARDIOVASCULAR AND GASTROINTESTINAL EVENTS

- Cardiovascular Risk
 Nonsteroidal anti-inflammatory drugs (NSAIDs) may cause an increased risk of serious cardiovascular (CV) thrombotic events, myocardial infarction, and stroke, which can be fatal. This risk may or risk factors for cardiovascular disease may be at greater risk [see Warnings and Precautions (5.11)].

 Meloskiam tablets are contraindicated for the treatment of right peripherative pain in the setting of coronary artery bypass graft (CMO) surgery [see Contraindications (4.2) and Warnings and Precaudions (5.11)].

(S-1)).

MSAIDs cause an increased risk of serious gestrointestinal (GI)

MSAIDs cause an increased risk of serious gestrointestinal (GI)

the stomach or intestines, which can be fatal. These events can occur at any time during use and without warning symptoms. Ellerly patients are at greater risk for serious gestrointestinal events [see

1.1 Osteoarthritis (OA) Meloxicam tablets are indical [see Clinical Studies (14.1)]. ted for relief of the signs and symptoms of osteoarthritis

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

2. DOSAGE AND ADMINISTRATION

2.1 General Instructions

Carefully consider the potential benefits and risks of meloxicam tablets USP and other treatment options before deciding to use meloxicam tablets USP. Use the lowest effective dose for the shortest duration consistent with individual patient treatment goals [see Warnings and Precautions (5.4)].

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

In adults, the maximum recommended daily oral dose of meloxicam tablets USP is 15 mg regardless of formulation. In patients with hemodialysis, a maximum daily dosage of 7.5 mg is recommended [see Warnings and Precautions (5.6), Use in Specific

Populations (8.7) and Clinical Pharmacology (12.3)].

Meloxicam tablets USP 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 tablets USP 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

3. DOSAGE FORMS AND STRENGTHS

- Tablets:

 7.5 mg: Light yellow, round flat beveled edged tablet with UL debossed on one side and 7.5 debossed centrally on the other side.

 15 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet with UL debossed on one side and 5.5 mg: Light yellow, capsule shaped, bconvex, tablet yellow, ca

4. CONTRAINDICATIONS

4.1 Allergic Reactions

Meloxicam tablets are contraindicated in patients with known hypersensitivity (e.g. anaphylactoid reactions and serious skin reactions) to meloxicam.

Meb.xicam tablets should not be given to patients who have experienced asthma, urticaria, or allergic-type reactions after taking aspirin or other NSAIDs. Severe, rarely fatal, anaphylacit-like reactions to NSAIDs have been reported in such patients [see Warnings and Precautions (5.7, 5.13)].

4.2 Coronary Surgery

Mebxikam tablets are contraindicated for the treatment of peri-operative pain in the setting of coronary artery bypass graft (CABG) surgery [see Warnings and Precautions (5.1)].

5. WARNINGS AND PRECAUTIONS

5.1 Cardiovascular Thrombotic Events

5.1 Cardiovascular Thrombotic Events
Cinical rules of several COX-2 selective and nonselective NSAIDs of up to three years' duration have shown an increased risk of serious cardiovascular (CV) thrombotic vents, myocardial fraction, and stroke, which can be fast, AlhSAIDs, both COX-2 selective and nonselective, may have a samilar risk. Patients with income CV disease or advantage of the control of t

the steps to take if they occur.
Two large, controlled, clinical trials 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 [see Contraindications (4.2)].

There is no consistent evidence that concurrent use of aspirin mitigates the increased risk of serious CV thrombotic events associated with NSAID use. The concurrent use of aspirin and an NSAID does increase the risk of serious GI events [see Warnings and Precautions (5.2)].

5.2 Gastrointestinal (GI) Effects - Risk of GI Ulceration, Bleeding, and Perforation

5.2 Gastrointestral (GI) Effects - Risk of GI Ukeration, Bleeding, and Perforation

NAMD, indicated the movement of the control of the contr

patents and treetoric, special are should be taken in treasing the proposation. First, To maintain the profession factor for the profession of the professi

5.3 Hepatic Effects

5.3 Hepatic Effects
Solderine development of the control of the control

A palent with syndroms and/or signs suggesting liver dysfunction, or in whom an alternative test has occurred, should be evaluated for evidence of the development of the syndroms and the syndroms and the syndroms and the syndroms are syndroms consistent with level disease develop, or if system canadisciplina social (e.g., essingoliha, rish, etc.), discontinue meloxicam (see Use in Specific Populations (6.6) and Critical Pharmacology (12.3)).

5.4 Hypertension

5.4 Hypertension
NSAIDs, including mebxicam tablets, can lead to onset of new hypertension or worsening of price-existing hypertension, either of which may contribute to the increased incidence of CV events. NSAIDs, including mebxicam tables, should be used with caution in patients with hypertension. Blood pressure (BPI) should be monitored closely during the hitation of HSAID treatment and throughout the course of therapy. Patients saking ACE inhibitors, thiszides or loop duretics may have impared response to these therapies when taking NSAIDs.

5.5 Congestive Heart Failure and Edema

Fluid retention and edema have been observed in some patients taking NSAIDs. Use meloxicam with caution in patients with fluid retention, hypertension, or heart failure.

5.6 Renal Effects

5.5 Feat Effects
Some find effects
General invariance of the Sulbs, including meloxicam tablets, can result in renal copilarly necrosis, event invarificancy, acute renal falance, and other renal rings, Renal toxicxly has also been seen in patients in whom renal prostagations have a compensatory role in the maintenance of renal perfusion. In these patients of content of the submittenance of renal perfusion. In these patients of the patients are renal decompensation. Patients at greatest risk of this reaction are the renal decompensation. Patients at greatest risk of this reaction are those with impatient renal function, heart failure, beer dystantion, those taking duretes, ACE-inhibotrs, and angiotens in Il receptor antiagrants, and the safety. Discontinuation Apharmacolonical toxically in patients with and moderate renal impatient reveals data no dosage adjustments in these patient populations are required. Patients with severe renal impartment have not been studied. The use of meloxicam in patients whits sever renal impartment have not been studied. The use of meloxicam in patients with severe renal impartment have not been studied. The use of meloxicam in patients with severe renal impartment have not been studied. The use of meloxicam in patients with severe renal impartment have not been studied. The use of meloxicam in patients with severe renal impartment have not been studied. The use of meloxicam patients with severe renal impartment have not been studied. The use of problems was increased. Therefore 8 is recommended that meloxicam opage in this population not received. The patients with respect to the patients with patients with a patient with required renal received on patients with repaired renal received in the patient with required renal received in the patients with respect to the patient with repaired renal received in the patient with repaired renal received in the patient with respect to the pat

Use caution when initiating treatment with meloxicam in patients with considerable dehydration. It is advisable to rehydrate patients first and then start therapy with meloxicam. Caution is also recommended in patients with pre-existing kidney disea

The extent to which metabolites may accumulate in patients with renal impairment has not been studied with meloxicam. Because some meloxicam metabolites are excreted by the kidney, monitor patients with significant renal impairment closely.

5.7 Anaphylactoid Reactions

As with order ESAIDS, anaphylectoid reactions have accurred in patients without known prior exposures to motivation. Most accurate making the group prior proposures to motivation. Motivations and build not be gloven to platents with new prior triad. This symptom complex typically occurs in asthmatic patients with one experience withinks with or without masal polypic, or more one pitch specific proportionally fatally branchespain after taking against or other MSAIDS, piec Contradictations (4.1) and analytication reaction occurs.

5.8 Adverse Skin Reactions

NSAIDs, including meboxicam tablets, can cause serious skin adverse events such as sexfolative dermatitis, Stevens-Johnson Syndrome (SJS), and toxic epidermain enrolysis (TRI), which can be fatal. These serious events may occur without warning. Inform patients about the signs and symptoms of serious six manifestations and only see of the drug at the first appearance of six in said or any other sign of hypersensibity.

5.9 .Pregnancy

5.9 :Pregnancy Starting at 30 weeks gestation, avoid the use of meloxicam, because it may cause premature closure of the ductus arteriosus [see Use in Specific Populations (8.1) and Patient Counseling Information (17.8)].

5.10 Corticosteroid Treatment

Mebxicam cannot be expected to substitute for corticosteroids or to treat corticosteroid insufficiency. Abrupt discontinuation of corticosteroids may lead to disease exacerbation. Slowly taper patients on prolonged corticosteroid therapy if a decision is made to discontinue corticosteroids.

5.11 Masking of Inflammation and Fever

The pharmacological activity of mebxicam in reducing fever and inflammation may diminish the utility of these diagnostic signs in detecting complications of presumed noninfectious, painful conditions.

5.12 Hematological Effects

Acenia may occur in patients receiving MSAIDs, including melouicam tables. This may be due to fluid retention, occul or gross oil bodo diss, or an incompletely described effect upon eythropoises. Retention on long-term treatment with NSAIDs, including meloxicam, should have their hemoglobin or hematocrit checked if they exhibit any signs or symptoms of anemia.

NSAIDs inhibit platelet aggregation and have been shown to prolong bleeding time in some patients. Unlike aspirin, their effect on platelet function is quantitatively less, of

shorter duration, and reversible. Carefully monitor patients treated with meloxicam who may be adversely affected by afterations in platelet function, such as those with coagulation disorders or patients receiving anticoagulants.

Coaguiton oscoress or patients receiving anticoaguinnts.

5.13 Use in Patients with Pre-existing Asthma
Patients with asthma may have apprin-sensible estima. The use of apprin in patients
with apprin-sensible estimal has been associated with severe bronchospasm, which
can be fastal. Since cross reactivity, including bronchospasm, between apprin and other
SASIDs has been reported in such apprin-sensible patients, medicinary and not to be
administered to patients with this form of apprin sensiblely and should be used with
cauchon in patients with pre-existing selection.

Because serious GI tract user altons and bleeding can occur without warning symptoms physicians should monitor for signs or symptoms of GI bleeding. Patients on long-term breatment with NSAIDs should have their CBC and a chemistry profile checked periodically. If clinical signs and symptoms consistent with we nor renal disease develop, systemic manifestations occur (e.g., esshophila, rsh, etc.) or if abnormal liver tests presist or wissen, mebicisen should be deloctrificated.

6. ADVERSE REACTIONS

6. ADVERSE REACTIONS

Because chical trible are conducted under widely varying conditions, adverse reaction rates observed in the clinical trisis of a drug cannot be directly compared to rates in the cinical trisis of another drug and may not reflect the rates observed in practice. The following serious adverse reactions are discussed elsewhere in the bleding:

Cardioviscustin information cerest light Beowed Warming and Warmings and Precautions (5.2)]

Hepatic effects [see Warmings and Precautions (5.3)]

Hepatic effects [see Warmings and Precautions (5.3)]

**Competitive Heart Statuse and celema [see Warmings and Precautions (5.5)]

**Competitive Heart Statuse and celema [see Warmings and Precautions (5.7)]

**Adverse skin reactions [see Warmings and Precautions (5.7)]

**Adverse skin reactions [see Warmings and Precautions (5.7)]

**Adverse skin reactions [see Warmings and Precautions (5.7)]

6.1 Clinical Trials Experience

6.1 Clinical Trials Experience
Adults
Osteoarthritis and Rheumatoid Arthritis
The metoscam Phase 2/2 clinical trial database includes 10,122 OA patients and 1012 RA
patients treated with metoscam 2.5 mg/day, 3,300 OA patients and 1351 RA patients
treated with metoscam 2.5 mg/day, 8400 Cam at these obsess was administered to 631
treated with metoscam 2.5 mg/day, 8400 Cam at these obsess was administered to 641
to 50,000 of these patients were treated in the placebo- and/or active-controlled costeoarthrists trials and 2363 of these patients were treated in ten placebo- and/or active-controlled remaintal arthrist Paris. Gastronietestical (G) adverse events were the most controlled on the placebo- and/or active-controlled remaintal arthrist Paris. Gastronietestical (G) adverse events were the most controlled on the placebo- and/or active controlled on the placebo- active and the placebo- and/or active controlled on the placebo- and/or active controlled on the placebo- and/or active controlled on the placebo- active active placebo- active active active placebo- active active placebo- active place

tribls. A 12-week multicenter, double-bind, randomired tribl was conducted in polisients with osteoathribls of the innee or hip to compare the efficacy and safety of medician who placebo and with an active control. The 21-week multicenter, double-bind, randomized tribls were conducted in patients with inheumatoid attribls to compare the efficacy and safety of medicians with placebo.

Table 1a depicts adverse events that occurred in 22% of the effocution of the production of t

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			ng Diclofenac 100
		mg daily	daily	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	n			
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

WHO preferred terms rash, rash erythematous, and rash maculo-papular combined

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

daily 481 18.9 2.9 5.8 3.3	15 mg daily 477 16.8 2.3 4.0 3.8
18.9 2.9 5.8 3.3	16.8 2.3 4.0
2.9 5.8 3.3	2.3 4.0
5.8 3.3	4.0
3.3	
	3.8
7.9	
2.9	
	2.3
7.0	6.5
1.5	2.3
6.4	5.5
1.0	2.1
	6.4

etuccasoni, gas uninessina irriadioni, opper iespratori succinculoris-painique i un'ispecia grandici invo-phanyojits NOS, sinustis. NOS), joint related signs and symptoms (arthralgia, arthralgia gravated, joint phanyojits NOS, sinustis. NOS), joint swelling)

2 medDRA preferred term: nouseea, abdominal pain NOS, influenza-like illness, headaches NOS, and rash NOS

The adverse events that occurred with meloxicam in $\ge 2\%$ of patients treated short-term (4-6 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							
		ntrolled Trials	6 Month Controlled Trials				
		Meloxicam 15 mg					
	daily	daily	daily	daily			
No. of Patients	8955	256	169	306			
Gastrointestinal	11.8	18.0	26.6	24.2			
Abdominal pain	2.7	2.3	4.7	2.9			
Constipation	0.8	1.2	1.8	2.6			
Diarrhea	1.9	2.7	5.9	2.6			
Dyspepsia	3.8	7.4	8.9	9.5			
Flatulence	0.5	0.4	3.0	2.6			
Nausea	2.4	4.7	4.7	7.2			
Vomiting	0.6	0.8	1.8	2.6			
Body as a Whole							
Accident household	0.0	0.0	0.6	2.9			
Edema ¹	0.6	2.0	2.4	1.6			
Pain	0.9	2.0	3.6	5.2			
Central and Peripheral Nervous Sys	tem						
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	0.1	0.4	4.7	6.9			
¹ WHO preferred terms edema, edema depen				3.3			

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

The following is a list of adverse druy reactions occurring in <2% of patients receiving mebricam or clinical trials involving approximately 16.200 patients.

Body as a Whole alergic reaction, face edema, fatigue, fever, hot flushes, malaise, syncope, weight decrease, weight increase Cardiovascular angina pectoris, cardioc failure, hypertension, hypotension, myocardial infarction, vascultis Body as a Whole

Cardiovascular

anging actors, cardisc failure, begreterison, hypothesis, makket, syncope, weight decrease, weight nere
anging actors, cardisc failure, begreterison, hypothesis, mycardisl affection, wasculis

Central and Peripheral Nervous System.com/ukbors, paresthesis, tereor, vertigo

Central and Peripheral Nervous System.com/ukbors, paresthesis, tereor, vertigo

Everal Refer and Bhythm

All Refer Refer and Bhythm

All Torcessed, Chaption, Tedryczeri, December System.

Metabolic and Nutritional

All Torcessed, Arthromema, Coff Torcessed, Department, Order System

Metabolic and Nutritional

dehydration

Metabolic and Nutritional

dehydration

All Torcessed, Chaptionema, Gotto Increased, hepatitis

dehydration

altomatic Arthromema, Gotto Increased, confusion, depression, nervousness, somnolence

authorized and Company and Compa

causal relationship to drug exposure. Decisions about whether to include an adverse event from spontaneous reports in labeling are typically based on one or more of the following factors; [1] send-curies of the event; [2] number of (reports, or [6]) Strength of following factors; [1] send-curies or event; [2] number of (reports, or [6]) Strength of experience or the Renature include acute urinary retention; agranulos (posts; alterations in mood (such as mood elevation), anaphylicatiof reactions including shock; experience multiforms; exfoliative dermatitis; interstitial reportis; juuniciae, liver failure. Stevens-johnson syndrome, and took explainmal necrolysis.

See also Clinical Pharmacology (12.3).

7.1 ACE-inhibitors

NSAIDs may diminish the anthypertensive effect of ACE-inhibitors. This interaction should be given consideration in patients taking meloxicam concomitantly with ACE-inhibitors.

7.2 Aspirin

When meloxicam is administered with aspirin (1000 mg three times daily) to healthy volunteers, an increase the AUC (10%) and C_{max} (24%) of meloxicam was noted. This clinical significance of this interaction is not known; however, as with other HSAIDS concomitant administration of meloxicam and aspirin is not generally recommended because of the potential for increased adverse effects.

Concomitant administration of low-dose aspirin with meloxicam may result in an increased rate of GI ulceration or other complications, compared to use of meloxican alone. Meloxicam is not a substitute for aspirin for cardiovascular prophylaxis.

7.3 Diuretics

7.3 Disrects

Clinical studies, as well as post marketing observations, have shown that NSAIDs can reduce the natriuretic effect of furosemide and thisuizes in some patients. This response has been attributed in hibition of real prostaginaries synthesis. However, studies with furosemide signitis and mebbox can have into demonstrated a reduction in natriuretic arrangement of the control of the

7.4 Lithium

In a study conducted in healthy subjects, mean pre-dose Rhium concentration and AUC were not reseed by 21 his a subjects receiving Billium doses ranging from B64 to 1072 mg when the subjects receiving Billium doses ranging from B64 to 1072 mg of the subjects of the subject of

7.5 Methotrexate

NSAIDs have been reported to competitively inhibit methotrexate accumulation in rabb kidney sixes. Therefore, NSAIDs may reduce the elimination of methotrexate, thereby enhancing the toxicity of methotrexate. Use caution when meloxicam is administered concomitantly with methotrexate [see Clinical Pharmacology (12-3)].

7.6 Cyclosporine

Mebickam, like other NSAIDs, may affect renal prostaglandins, thereby altering the renal toxicity of certain drugs. Therefore, concomitant therapy with mebickam may increase cyclosporine's nephrotoxicity. Use caution when meloxicam is administered concomitantly with cyclosporine.

7.7 Warfarin

The effects of warfarin and NSAIDs on GI bleeding are synergistic, such that users of both drugs together have a risk of serious GI bleeding higher than users of either drug abne.

wome. Monotor anticoagulant activity, particularly in the first few days after initiating or changing medication therapy in platients receiving warfarin or similar agents, since these platients when administering melockam with warfarin since patients on warfarin may experience changes in INR and an increased risk of bleeding complications when a new medication is introduced [see Circla/Pharmacology (2.3.)].

8. USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

8.1 Pregnancy Category C. Category D starting 30 weeks gestation. There are no adequate and well-controlled studies in pregnant women. Meboxicam crosses the pice-real barrier. Prior to 30 weeks gestation, use meloxicam during pregnancy only if the potential benefit justifies the potential risk to the fetus. Starting at pregnancy only if the potential benefit justifies the potential risk to the fetus. Starting at pregnancy only if the potential benefit justifies the potential barrier of the ductus at refusion in the fetus may occur. If this drug is used during this time period in pregnancy, inform the patient of the potential hazard to a fetus (see Warrings and Presculture 1.59) and Potentic Countering information (1.78). Teratogenic Effects

Teratogenic Effects

Meboxicam was not teratogenic when administered to prepnant rats during fetal
organogenesis at oral doses up to 4 mgkgday (2.6-fold greater than the maximum
recommended human daly dose [MRNID] based on body surface area [BSA]
comparison). Administration of meboxicam to pregnant rabbts throughout
ormary one of the production at presented incidence of sepatial decision of the heart at an oral
entire or beging day. The in effect level was 20 mg/leg/day (12-fold greater than the
MRID based on FSA conversion).

In rats and rabbits, embryolethality occurred at oral meloxicam doses of 1 mg/kg/day and 5 mg/kg/day, respectively (0.65-and 6.5-fold greater, respectively, than the MRHD based on BSA comparison) when administered throughout organogenesis.

8.2 Labor and Delivery

The effects of melosciam on abor and delivery of prognant women are unknown. Oral administration of melosciam to pregnant rats during site gestation through licitation increased the nicition of dystocia, delegies parturation, and decreased offspring survival at melosciam doses of 0.125 mg/sg/dg/sy of greater lat least 11.25 times lower comparison).

8.3 Nursing Mothers

It is not known whether this drug is excreted in human milk; however, meloxicam was excreted in the milk of lactating rats at concentrations higher than those in plasma. An exception of the milk of lactating rats at concentrations higher than those in plasma serious adverse reactions in nursing infants from meloxicam a decision should be made whether to discontinue nursing or to discontinue the drug, taking into account the importance of the drug to the mother.

8.4 Pediatric Use

Use of this drug for a pediatric indication is protected by marketing exclusivity

As with any NSAID, caution should be exercised in treating the elderly (65 years and older)

ouer).

Of the total number of subjects in clinical studies, 5157 were age 65 and over (4044 in OA studies and 1113 in RA studies). No overal differences in safety or effectiveness were observed between these subjects and younger subjects, and other reported clinical experience has not identified differences in responses between the editery and younger patients, but greater sensitivity of some older individuals cannot be ruled out.

8.6 Hepatic Impairment

No dose adjustment is necessary in patients with mild to moderate hepatic impairment. Patients with severe hepatic impairment have not been adequately studied. Since medocace in significantly metabolized in the leve, huse of meboxan in these patients should be done with could be Warnings and Precautions (5.3) and Chikal Pharmacology (12.3).

8.7 Renal Impairment

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 meloxicam is subjects with severe renal impairment as not recommended. Thoughing a single does of meloxicam, the free C_{main} plaims concentrations were higher in patients with renal Countries of the patients of

10. OVERDOSAGE

10. OVENUSABLE
There is limited experience with meloxicam overdose. Four cases have taken 6 to 11 times the highest recommended dose; all recovered. Cholest prainie is known to accelerate the clearance of meloxicam.

Symptoms following acute NSAID overdose include letharry, drowsiness, nausea, vorning, and episatic pash, with a regenerally reversible with supportive care. Gastrointestrial bleeding can occur. Severe posicioning may result in hypertension, acute establishment of the control of the

worn merapeutic ingestion of NSAIDs, and may occur following an overdose. Patients should be managed with symptomatic and supportive care following an NSAID overdose. Administration of activated charcoal is recommended for patients who present 1.2 hours after overdose. For obstantiatiol overdose or severely symptomatic present 1.2 hours after overdose. For obstantiation of severely symptomatic mebruicam by 4 gm or aid doses of choles tyraimne quent three times a day was demonstrated in a finish at list. Administration of cholestypamne may be useful following an overdose. Forced duresis, alkalinization of urine, hemodalysis, or hemopertusion may not be useful tools to high protein but of the present of the present of the properties of the present of p

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

11. DESCRIPTION

Mebiciam, an oxicam derivative, is a member of the enoils acid group of nonsteroidal anti-inflammatory drugs (MSAIDs). Each tablet contains 7.5 mg or 15 mg mebiciam in oral administration Mebiciams is chemical designated as 4-hydroxy-2-methyl-k15-weight is 351.4. its empirical formula is C_{1,2}H_{2,1}N₂O₂O₂ and it has the following structur formula.



Mebxicam is a pastel yellow solid, practically insoluble in water, with higher solubility observed in strong acids and bases. It is very slightly soluble in methanol. Meloxicam has an apparent partition coefficient (log $\hat{\mathbf{P}}_{lago} = 0.1$ in n-octanol/buffer pH 7.4. Mebxicam has pike values of 1.1 and 4.2.

Meloxicam is available as a tablet for oral administration containing 7.5 mg or 15 mg meloxicam.

The inactive ingredients in Meloxicam tablets USP include colloidal silicon dioxide, crospovidone, lactose monohydrate, magnesium stearate, microcrystalline celluiose, povidone and sodium ctrate dihydrate.

12. CLINICAL PHARMACOLOGY

12. LURAchamin of Action
The mechanism of action of mekicam, like that of other MSAIDs, may be related to prostaglands synthesize (cycloxygenase) inhibition which is movived in the initial step prostaging the control of the composition of the composition of the composition of the composition of results in the repair of these composition sensitis in the repair. If it is not completely understood how reduced synthesis of these composition sensitis in the repair.

12.2 Pharmacodynamics

Meloxicam exhibits anti-inflammatory, analoesic, and antipyretic activities.

12.3 Pharmacokinetics

Absorption

Asserption. The abdotac boxelebility of medicion capacities was 97% following a single on ideas. The abdotac boxelebility of medicion capacities was 97% following a single of medicion capacities of the proportion of the proporti

			Steady Sta		Single Dose		
Pharmacokinetic Parameters (%CV)	Healthy male adults (Fed) ²		Elderly males (Fed) ²	Elderly females (Fed) ²	Renal failure (Fasted) Hepatic insufficiency (Fas		
		7.5 mg ³ tablets	15 mg capsul	es 15 mg capsules	15 mg capsules	15 mg capsules	
N		18	5	8	12	12	
C _{max}	[µg/mL]	1.05 (20)	2.3 (59)	3.2 (24)	0.59 (36)	0.84 (29)	
t _{max}	[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-/f4	[L]	14.7 (32)	15 (42)	10 (30)	26 (44)	14 (29)	

Food and Antacid Effects

From all with Exercises of medical resolution (see Superior Manufacture) on medical control and Administration of medical resolution (see Superior Manufacture) and the second proper position of the proper position of the proper position of the property of the second propert

Distribution

Distribution

The mean volume of distribution (Vss) of meloxicam is approximately 10 L. Mebxicam is -99.4% bound to human pisams proteins (primarly abumin) within the threspect (one range. The fraction of protein briding is dependent of drug concentration, over the clinically relevant concentration range, but decreases to -99% in patients with real disease. Meloxicam peer atola in the human are blood cets, after orid dosing, a less than 100 processing as the patient of the concentration of the response of the residentity detected in the plasma was present as in unchanged meloxicam.

Mebickam concentrations in synovial fluid, after a single oral dose, range from 40% to 50% of those in plasma. The free fraction in synovial fluid is 2.5 times higher than in plasma, due to the lower albumin content in synovial fluid as compared to plasma. The significance of this peretration is unknown.

Metabolism

Metabolism which is extensively metabolized in the liver, Meloxicam metabolites include 5'carboxy metabocam (60% of dose), from P-450 metabed metabolism formed by
secreted to a less-received (9% of dose), in the properties of the properties of

Excretion

Excretion Medician excretion is predominantly in the form of metabolites, and occurs to equal extents in the urine and feecs. Only fraces of the unchanged parent compound are excreted in the urine (0.2%) and feecs (1.6%). The extent of the urinary excretion was confirmed for uniabeled multiple 7.5 mg doses: 0.5%, 6%, and 1.3% of the dose were found in urine in the form of mebicacins, and the 5-hydroxymethyl and 5-catoboxymethyl and 5-dectory and the contract of the contract

Special Populations

Special Populations

Editoria

Edito

Gender

The time is exhibited sightly lower placens concentrations relative to young males. After single dose of 75 mg meloscum, the mean elementation had life was 155 hours for the female group as compared to 23.4 hours for the male group. At steady state, the data were similar (179 hours vs 214 hours). This pharmacoknetic difference due to gender is likely to be of little clinical importance. There was linearly of pharmacoknetics and no appreciable difference in the Cam, or Timax efforcs genders.

Hepatic Impairment

Headst (maximum). Following a stopp 1.5 mg dose of meloxicam there was no marked difference in plasma concernations in patients with mild (full-Puly). Class II or moderate (full-Puly) Class III) headst (maginater compared to headst volunteers, Profest binding of meloxicam was not affected by hepatic impairment. No dosage adjustment is necessary in patients with mild to moderate hepatic impairment. Roletox with seven hepatic impairment (Clab Puly) Class IIII) have not been adequately studied. [see Warnings and Precautions 5.3, and User in Specific Pepulations of See

(3-3) and use on species. Propagations (as (b)). Remail Impairment Medician pharmacolites have been investigated in subjects with mild and moderate meal impairment. Total drug plasma concentrations of medician decreased and total remail inpairment. Total drug plasma concentrations of medician decreased and total values were similar in all groups. The higher medician character is subjects with renal impairment may be due to increased fraction of unbound medician which is available in patients with the propagation of the

Populations (8.7); Hermodalskis Following a single dose of meloxicam, the free C_{max} plasma concentrations were higher in plasties that real failure on chronic hemodalysis (1% free fraction) is comparison to healthy volunteers (0.3% free fraction). Hemodalysis did not lower the total drug concentration in plasmit, therefore, additional doses are not necessary after the longitudes. Hebixicam's in off cultipaths, like bodings and Administration (2.1), warrings and Practions (5.6) and the in Specific Population (6.7).

Warnings and Precautions (5.6) and Use in Specific Populations (6.7)!.

Normal Interactions and International Specific Populations (6.7).

Aspire: When meloxicam is administered with aspire (1,000 mg three times daily) to healthy voluntees, stended to increase the AUC (10%) and C_{max} (24%) of meloxicam. The clinical significance of this interaction is not known [see Drug Interactions (7.2)]. Observations: Pretentiented for four days with cholestymenine significantly increased the clearance of meloxicam by 50%. This resulted in a decrease in typ, from 19.2 hours to 12.5 hours, and 35% reduction in AUC. This suggests the existence of a recirculation pathway for meloxicam in the gastroniestinal tract. The clinical reevance of the interaction has not been established.

 $\label{lem:concomitant} \textit{Cimetidine:} Concomitant administration of 200 mg cimetidine four times daily did not alter the single-dose pharmacokinetics of 30 mg meloxicam.$

carrescence Concomitant animostration of 200 mg cimetatine four times daily did not alter the single-doce pharmacoinetics of 300 mg meloxide the planta concentration. Digouin, Meloxican 15 mg norce daily for 7 days did not alter the planta concentration. Digouin, Meloxican 15 mg norce daily for 7 days did not alter the planta concentration to the string found no protein binding drugs interaction between digoris and meloxics. In who testing found no protein binding drugs interaction between digoris and meloxics. In which the concentration and AUC were increased by 22 Mis negligible, seeing protein protein binding drugs and all concentration and AUC were increased by 22 Mis negligible, seeing this modes rangering from 804 to receiving thism alone like the Drug Interaction (7-4), days as compared to subjects receiving thism alone like the Drug Interaction of the pharmacointeraction and the pharmacointeraction of the pharmacointe

13. NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment Of Fertility

Carchoppens: There was no increase in tumor incidence in long-term carchopenics, but not so in tumor incidence in long-term carchopenics, but note in rats (104 weeks) and mixe (99 weeks) administered meloxicam at oral dose up to 0.8 mg/kg/dly in rats and up to 3.0 mg/kg/dly in receipt up to 0.5 mg/kg/dly in receipt to 10.5 mg/kg/dly

Mutagenesis: Meloxicam was not mutagenic in an Ames assay, or clastogenic in a chromosome aberration assay with human lymphocytes and an in vivo micronucle test in mouse hope marchy.

² not under high fat cond 3 Meloxicam tablets 4 V₂/f= Dose/(AUC+Kel)

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-fold greater, respectively, than the maximum recommended human daily dose based on body surface area comparison).

14. CLINICAL STUDIES

14.1 Osteoarthritis and Rheumatoid Arthritis

1.4.1. Ostcoarthritis and Rheumatoli Arthritis
The use of mebuscan for the treatment of the signs and symptoms of octeoarthritis of
the knee and the was evaluated in a 1.2-week, double-bind, controlled trial. Moloricam
(3.75 mg. 7.5 mg. and 1.5 mg dally was compared to placebo. The four primary)
endpoints were investigator's global assessment, patient global assessment, patient pain
endpoints were investigator's global assessment, patient global assessment, patient pain
eassessment, and total WOMAC score is an exi-administered questionnaire adviressing
dally showed significant improvement in each of these endpoints compared with
placebo.

The use of mebuscan for the management of signs and symptoms of osteoarthritis was
evaluated in six double-bind, active-corrolled trials outside the U.S. ranging from 4
weeks' to 6 months' duration. In these trials, the effizacy of mebuscam, in does of 7.5
when the control of the six of

16. HOW SUPPLIED/STORAGE AND HANDLING

Meboixcam Tablets USP are available as light yellow, round, flat, uncoated tablets containing meboxcam 7.5 mg or as light yellow, obbong, biconvex, uncoated tablets containing meboxcam 15 mg. The 7.5 mg tablet is impressed with letter U and L on one side and tablet code 7.5 on the other side. The 15 mg tablet is impressed with letter U and L on one and L on one side and tablet code 1.5 on the other side.

Meloxicam Tablets USP 7.5 mg are available as follows:

35356-808-15 Bottles of 15

35356-808-30 Bottles of 30

35356-808-60 Bottles of 60 35356-808-90 Bottles of 90 Meloxicam Tablets USP 15 mg are available as follows:

Store at Controlled Room Temperature 20 $^{\circ}$ -25 $^{\circ}$ C (68 $^{\circ}$ -77 $^{\circ}$ F) [See USP]. Keep Mebxicam Tablets USP in a dry place

Dispense tablets in a tight container.

Keep this and all medications out of the reach of children

Patients should be informed of the following information before initiating therapy with an NSAID and periodically during the course of ongoing therapy.

17.1 Medication Guide

Inform patients of the availability of a Medication Guide for NSAIDs that accompanies each prescription dispensed, and instruct them to read the Medication Guide prior to using meloxicam tablets.

17.2 Cardiovascular Effects

1.1.4 Cardiovascular strects

NABLOS schulding molecular tablets, may cause serious CV side effects, such as MI or stroke, which may result in hopishalization and even death. Although serious CV events of the control of the control

17.3 Gastrointestinal Effects

17.3 Gastrointestinal Effects

NSAIGS relating mission can believe, can cause G decomfort and , ranky, serious G in NSAIGS relating mission can be leding, which may result in hospitablication and red edit and serious of the serious and bleeding, and so however the constraint and serious serious of the serious and bleeding and solvand serious discharge serious G tract ut develops and solvand and serious model and serious seriou

17.4 Hepatotoxicity

Inform patients of the warning signs and symptoms of hepatotoxicity (e.g., fatigue, lethargy, prurtus, jaundice, right upper quadrant tendemess, and 's symptoms). If these occur, instruct patients to stop therapy and seek imm medical therapy [see Warnings and Precautions (5.3)].

17.5 Adverse Skin Reactions

17.5 Advere Skin Recture (Sharper) and the Sharper (Sharper) and th

17.6 Weight Gain and Edema

Advise patients to promptly report signs or symptoms of unexplained weight gain edema to their physicians [see Warnings and Precautions (5.5)].

17.7 Anaphylactoid Reactions

Inform patients of the signs of an anaphylactoid reaction (e.g., difficulty breathing, swelling of the face or throat). Instruct patients to seek immediate emergency help [see Warnings and Precautions (5.71).

17.8 Effects During Pregnancy

Starting at 30 weeks gestation, meloxicam should be avoided as premature closure of the ductus arteriosus in the fetus may occur [see Warnings and Precautions (5.9) and Use in Specific Populations (8.1)].

Please address medical inquiries to 1-866-562-4616

Manufactured by: UNICHEM LABORATORIES LTD.

Pilerne Ind. Estate, Pilerne, Bardez, Goa 403511, India

Marketed by:



Rochelle Park, NJ 07662

R-03-03/2011

Red3-03/2011
Medication Guide for Non-Steroidal Anti-Inflammatory Drugs (NSAIDs.)
(See the end of this Medication Guide for a list of prescription NSAID medicines.)
What is the most important Information I know about medicines called NonSteroidal Anti-Inflammatory Drugs (NSAIDs)?

NSAID medicines may increase the chance of a heart attack or stroke that can lead to death. This chance increases:

• with longer use of NSAID medicines
• in people who have heard disease

NSAID melicines should rever be used right before or after a heart surgery called a correnancy artery lypass gant (CABLO).

NSAID melicines can cause obcers and bleeding in the stomach and intestines time during frestment. (Gers bleeding:

- can happen without warning symptoms
- my cause celebrate.

The chance of a person getting an ulcer or bleeding increases with:

• taking medicines called "corticosteroids" and "anticoagulants"

• longer use

• smoking

• driking atchol

older age
 having poor health

NSAID medicines should only be used:

• exactly as prescribed

• at the lowest dose possible for your treatment

• for the shortest time needed

for me shortest time needed
 What are Non-Steroidal Anti-Inflammatory Drugs (NSAIDs)?
 NSAID medicines 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 a Non-Steroidal Anti-Inflammatory Drug (NSAID)?

Do not take an NSAID medicine:

• If you had an asthma attack, hives, or other allergic reaction with aspirin or any other NSAID medicine

• for pain right before or after heart bypass surgery

for pan right before or after heart bypass surgery
 about all of your medical conditions
 about all of your medical conditions
 about all of the medicines you take. NSALD, and some other medicines can interact about all of the medicines you take. NSALD and some other medicines can interact show to your healthcare provider and pharmacist.
 for you are preparant.
 for you are preparant.
 for you are preparant.
 for you are preparant.
 for you are preparant.

What are the possible side effects of Non-Steroidal Anti-Inflammatory Drugs (NSAIDs)?

Serious side effects include:

• heart attack

- stroke
 high resure
 high resure

Get emergency help right away if you have any of the following symptoms: • consense of breath or trouble breathing • weakness in one part or side of your body • subrred speech • wedness in our the face or throat

- swelling of the face or throat Stop your NSAD medicine and call your healthcare provider right away if you have any of the following symptoms: - naussea - more the reaker than usual - your skin or eyes bok yellow - stomach pan - fluide symptoms - there is blood in your bowel movement or it is black and sticky like tar - unusual weeping - skin reak or discuss with feer - swelling of the same, and legs, hands and feet

These are not all the side effects with NSAID medicines. Talk to your healthcare provider or pharmacist for more information about NSAID medicines.

- or pharmacist for more information about MSAID medicines.

 Call your doctor for medical advice about side effects. You may report side effects to FDA at 1800-FDA-1088.

 Other information about Non-Steroidal Anti-Inflammatory Drugs (INSAIDs)

 Aspirin is an MSAID medicine but it does not increase the chance of a heart attack. Aspirin can asset bedeen in the Paris, stomach, and intertiens. Aspirin can also bedeen in the Paris, stomach, and intertiens. Aspirin can also so the side of the Paris of

NSAID medicines that need prescription

Generic Name	Tradename
Celecoxib	Celebrex
Diclofenac	Cataflam, Voltaren, Arthrotec (combined with misoprostol)
Diflunisal	Dolobid
Etodolac	Lodine, Lodine XL
Fenoprofen	Nalfon, Nalfon 200
Flurbiprofen	Ansaid
Ibuprofen	Motrin, Tab- Profen, Vicoprofen* (combined with hydrocodone), Combunox (combined with oxycodone)
Indomethacin	Indocin, Indocin SR, Indo-Lemmon, Indomethagan
Ketoprofen	Oruvail
Ketorolac	Toradol
Mefenamic Acid	Ponstel
Meloxicam	Mobic
Nabumetone	Relafen
Naproxen	Naprosyn, Anaprox, Anaprox DS, EC- Naprosyn, Naprelan, Naprapac (co- packaged with lansoprazole)
Oxaprozin	Daypro
Piroxicam	Feldene
Sulindac	Cinoril
Tolmetin	Tolectin, Tolectin DS, Tolectin 600

Vicoprofen contains the same dose of fluprofen as over-the-counter (OTC) NSAIDs, and is usually used for less than 1 days to treat pain. The OTC NSAID label warns that long-term continuous use may increase the risk of heart attack or stroke.

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

Manufactured by: UNICHEM LABORATORIES LTD.

Pilerne Ind. Estate, Pilerne, Bardez, Goa 403511, India Marketed by:



Rochelle Park, NJ 07662 13002550

R-02-09/2010



	ELOXICAI								
mi	eloxicam table	20							
P	roduct Info	rmation	1						
P	roduct Type		HUMAN PRI			em Code NDC-31		5356-808(NDC:29300-	
R	oute of Admir	nistration	ORAL						
Δ	ctive Ingred	lient/Δc	tive Moiety						
			aredient Nan	ne		Basis of St	renath	Strength	
					MELOXICAM		7.5 mg		
Ir	active Ingr	edients							
				ent Name			S	Strength	
	ROSPOVIDONE								
	ICTOSE MONO! AGNESIUM STE		UNI: EWQ57Q8I5:	0			_		
	LICON DIOXIDE								
			PRATE (UNI: B22	547895K)					
PC	VIDONE K29/3	32 (UNI: 39	ORMINZPEO)						
CI	LLULOSE, MIC	ROCRYST	ALLINE (UNI: OP2	R32D61U)					
	roduct Chai	racteris							
	olor		YELLOW	Score			no score 7mm		
	nape		ROUND	Size					
Flavor				Imprint 0	ode		U;L;7;5		
C	ontains								
D	ackaging								
۳					M	arketing Start	Mark	etina End	
2	Item Code		Package D	escription		Date		Date	
1	NDC:35356- 808-15	Combinat	15 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product			7/2007	10/11/2	019	
2	NDC:35356- 808-30 NDC:35356-	30 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product			03/0	7/2007			
3	808-60	60 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product				7/2007			
4	4 NDC:35356- 90 in 1 BOTTLE, PLASTIC; Type 0: Not a 03/07/2007 Combination Product								
N	larketing	Inforr	nation						
	Marketing Category			er or Monog tion		arketing Start Date		eting End Date	
Ab	IDA	ANDAG	77927		03/0	7/2007			