WARNING: RISK OF CARDIOVASCULAR AND GASTROINTESTINAL EVENTS

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 Nons teroidal anti-tiflammatory drugs (NSAIDs) may cause an increased risk of serious
 cardions ordain (CV) thrombotic events, myocardia infarction, and struck, which can be
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 or risk factors for cardiovascular disease may be at greater risk [see Wornings and
 Procuntors (5.1)].
- Meloxican tablets are contraindicated for the treatment of perioperative pain in the setting of coronary artery bypass graft (CABG) surgery [see Contraindications (4.2) and Warnings and Precautions (5.1)].

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NSAIDs cause an increased risk of serious gastrointestinal (GI) adverse reactions including bleeding, udceration, and perforation of the stomach or intestines, which can be fatal. These events can occur at any time during use and without warming symptoms. Elderly patients are at greater risk for serious gastrointestinal events [see Wornings and Precunitors (G2)].

1. INDICATIONS AND USAGE

1.1 Os teo arthritis (OA)

Meloxicam tablets are i Studies (14.1)]. ndicated for relief of the signs and symptoms of osteoarthritis [see Clinical

1.2 Rheumatoid Arthritis (RA)

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 meloxican tablets USP and other treatment options before deciding to use meloxicam tablets USP. Use the lowest effective dose for the shorrest duration consistent with includeal patient reatment goals leg Warnings and Procarditors (S.A.). After observing the response to initial therapy with meloxicam tablets USP, adjust the dose to suit an individual patients needs.

instruction patients i necess.

In adults, the maximum recommended daily oral dose of meloxicam tablets USP is 15 mg regardless of formaliation. In patients with hemodialysis, a maximum daily dosage of 7.5 mg is recommended [see Warnings and Precutions (6.6). Use in Despetile Populations (6.7) and Clinical Pharmocology (12.3)].

Meloxicam tablets USP may be taken without regard to timing of meals.

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

For the relief of the signs and symptoms of rheumatoid arthritis, the recommended starting and maintenance or all 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.

3. DOSAGE FORMS AND STRENGTHS

- Tables:

 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, capasile shaped, biconvex, tablet with UL debossed on one side and 15 debossed centrally on the other side.

4.1 Allergic Reactions

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

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

4.2 Coronary Surgery

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

5. WARNINGS AND PRECAUTIONS

5.1 Cardiovas cular Thrombotic Events

5.1 Cardiovas cular Thrombotic Events
Clinical trisks of several COX-2 selective and nonselective NSAIDs of up to three years' duration have
shown an increased risk of serious cardiovascular (CV) thrombotic events, myocardial infaction, and
stools, which can be failed. All NSAIDs, both COX-2 selective and numelective, myo have a similar
stools, which can be failed. All NSAIDs, both COX-2 selective and numelective, myo have a similar
the potential risk for an adverse CV event in patients reased with an NSAID, the lowest effective does
should be used for the shortest duration possible. Physician and patients should remain alert for one
development of such events, even in the absence of previous CV symptoms. Patients should be informed
about the signs and/or symptom of services CV events and the steps to tale 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 Controllactions (42).

[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 Precoutions (5:2)].

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

5.2 Gastrointestinal (GI) Effects - Risk of GI Ukeration, Bleeding, and Perforation
NSAIDs, including meloxicam tables, can cause serious gastrointestinal (GI) adverse events including
inflammation, theeding, diceration, and perforation of the sunand, small insettine, or large insetine,
symptoms, in patients reason deep deproaction of the sunand, small insettine, or large insetine,
symptoms, in patients reason dwith NSAIDs. Only one in five patients who develop a serious upper GI
adverse event on NSAID therapy is symptomatic. Upper GI ulters, gross beleeding, or perforation
caused by NSAIDs, occur in approximately 1% of patients reason of 3 to 6 months, and in about 2 to
4% of patients reasted from one year. These enexts, containe with longer dataroin of itse, increasing the
short-term therapy is not without risk.
Prescribe NSAIDs, Including meloxicam tables, with examer causion in those with a prior history of
ulter disease or gastrointestinal bleeding. Patients with a prior history of peptic ulter-disease and/or
space of the patients of

5.3 Hepatic Effects

Dorderline elevations of one or more liver tests may occur in up to 15% of patients taking NSAIDs including meloxican tablets. These laboratory abnormalities may progress, may remain unchanged, may be transient with contaming therapy. Notable elevations of ALT or AST (approximately three or more times the upper limit of normal) have been reported in approximately 1% of patients in clinical trials with NSAIDs. In addition, tare case of severe bepatie reaction, including sandities and fatal rails with NSAIDs. In addition, tare case of severe bepatie reaction, including sandities and fatal response of them with fatal outcomes have been reported [see Adverse Reaction (6.11)].

A patient with symptoms and/or signs suggesting liver dysfunction, or in whom an abnormal liver test has occurred, should be evaluated for evidence of the development of a more severe hepatic reaction while on therapy with neloxicant. If clinical signs and symposon consistent with liver disease develop or If systemic manifestitions occur (e.g., es)mismological, etc.), discontinue meloxicam [see Use in Specific Populations (8.6) and Chincel Patronacology (12.3).

NSAIDs, including meloxicanulabless, can lead to onset of new hypertension or worsesting of pre-existing hypertension, either of which may committee to the increased incidence of CV events. NSAIDs, including meloxicanulables, should be used with caution in patients with hypertension. Blood pressure (BP) should be monitored closely during the initiation of NSAID reament and throughout the coarse of therapy.

Patients taking ACE inhibitors, thiazides or loop diuretics may have impaired 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.

Long-term definishment of NSAIDs, including meloxicam tables, can result in renal papillary necrosis, renal insufficiency, actue renal failure, and other renal injury. Renal moxicity has also been seen in padiers in whom renal prosagandins have a compensatory role in the maintenance of renal perfusion in these patients, administration of a nonsteroidal anti-inflammatory drug may cause a dose-dependent reduction in prostaglandin formation and, secondarily, in renal blood flow, which may precipitate overt renal decompensation. Patients at greatest risk of this reaction are twee with impaired renal furtion, the ref failure, liver dysfanction, dose taking diurties, ACE-inhibitors, and angionesis ill receptor amagonists, and the elderly. Discontinuation of NSAID therapy is usually followed by recovery to the precedenties state.

recovery to the pretearment state.

A pharmacokinet is study in patients with mild and moderate renal impairment revealed that no dosage adjustments in these patient populations are required. Patients with severe renal impairment have not been studied. The use of melox cain impairment share not make the patient population is not recommended. A study performed in patients on hemodalysis revealed that although overall Cmm, was diminished in this population, the proportion of free drop not bound to plasma was increased. Therefore it is recommended that meloxicam dosage in this population not exceed 7.5 mg per day. Closely monitor the renal function of patients with impaired renal function who are taking meloxicam (see Dosage and Administration (2.1), Use in Specific Populations (8.7) and Clinical Pharmacology (12.2).

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 disease.

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

3.. Analopyaction treatmons.
As with other NSAIDS, analophactioid reactions have occurred in patients without known prior exposure to melasticam Mediactions should rule be given to patients with the aspiritariad. This symptom which is the property of the

5.8 Adverse Skin Reactions

NSAIDs, including melosic antiblets, can cause serious skin adverse events such as exfoliative dermatitis, Sevens-Johnson Syndrome (SJS), and notic epidermal necrolysis (TEN), which can be fatal. These serious events my occur without varning, liformpainters about the signs and symptoms of serious skin manifestations and discontine use of the drug at the first appearance of skin rash or any other sign of hypersensitivity.

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

Meloxicam 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 meloxicam in reducing fever and inflammation may diminish the utility of these diagnostic signs in detecting complications of presumed noninfectious, painful conditions.

Anemia may occur in patients receiving NSAIDs, including meloxicam tables. This may be due to fluid retention, occult or gross GI blood loss, or an incompletely described effect upon erythropolesis. Patients on long-term treatment with NSAIDs, including meloxicam, should have their hemoglobin or hematocrit checked if they exhibit any signs or symptoms of anemia.

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NSAIDs inhibit pateet aggregation and have been shown to prolong, bleeding time in some patient Unlike aspirin, their effect on plateler function is quartitatively less, of shorter duration, and revers Carefully montion patients retead with meloxicam who may be adversely affected by alterations in plateler function, such as those with coagulation disorders or patients receiving anticoagulants.

5.13 Use in Patients with Pre-existing Asthma

Patients with asthma may have aspirin-sensitive asthma. The use of aspirin in patients with aspirin-sensitive asthma has been associated with severe bronchospam, which can be fauld. Since crosss reactivity, including bronchospam, between aspirin and other PKADDs has been reported in such aspirin-sensitive patients, emboxic an should not be administered to patients with this form of aspirin-sensitivity and should be used with caudion in patients with pre-existing asthma-

5.14 Monitoring

Because serious GI tract ulcerations and bleeding can occur without warring symptoms, physicians should monitor for signs or symptoms of I bleeding, Patients on long-term reasment with NSAIDs should have their CER. and a chemistry portlic checked periodically, it clitical signs and symptoms should have their CER. and a chemistry portlic checked periodically, it clitical signs and symptoms consistent with liver or tend disease develop, systemic manifestations occur (e.g., essimphilia, rash, etc.) or if abarrant liver iests persist or worsen, melociacum should be discontinued.

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

- and may not reflect the rates observed in practice.

 The following retrous adverse reactions are discussed elsewhere in the labeling:

 Cardiovascular thrombotic evenue [see Boord Worning and Wornings and Precautions (5, 1)]

 Castroninestallar lefters risks of Cal Liceration, beeding, and perforation [see Boord Worning and Wornings and Precautions (5, 2)]

 Hepacier effects [see Warnings and Precautions (5, 4)]

 Congestive heart failure and cleening see Warnings and Precautions (5, 5)]

 Renal effects [see Warnings and Precautions (5, 6)]

 Anaphylaction faceations [see Warnings and Precautions (5, 7)]

 Adverse skin reactions [see Warnings and Precautions (5, 8)]

6.1 Clinical Trials ExperienceAdults

6.1 Clinical Trials Experience Adults

Chesonarthizia and Bheumatoid Arthritis

The metostcam Phase 2.0 clinical trial database includes 10,122 OA patients and 1012 RA patients treated with melostcam 75 mg/day, 3,050 CA patients and 1351 RA patients reated with melostcam 15 mg/day, 3,050 CA patients and 1351 RA patients reated with melostcam 15 patients for a least one year. Approximately 10,500 of these patients were reated in ten placebo- and/or active-controlled osteoarthirits trials and 258.3 of these patients were reated in ten placebo- and/or active-controlled remanoid arthritis visils. Castroliteration (Cl) adverse evenes were the most frequently reported adverse events in all treatment groups across melostcam trials.

A 12-week multicenter, double-blind, randomized trial was conducted in patients with osteoarthritis of the knee or hip to compare the efficacy and safety of meloxicam with placebo and with an active cortori. Two 21-week multicenter, double-blind, randomized trials were conducted in patients with rheumatoid arthritis to compare the efficacy and safety of meloxicam with placebo.

Table 1a depicts adverse event shat occurred in 22% of the meloxicam treatment groups in a 12-week placebo- and active-controlled osteoarthritis trial.

Table 1b depicts adverse event shat occurred in 22% 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 | 2.8 | 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 Rash² | 2.5 | 2.6 | 0.6 | 2.0 |

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

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

| | Placebo | Meloxicam 7.5 mg daily | Meloxicam 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 |
| Nause a ² | 2.6 | 3.3 | 3.8 |
| General Disorders and Administration Site Conditions Influenza-like illness ² | 2.1 | 2.9 | 2.3 |
| nfection and Infestations Upper Respiratory tract infections-pathogen class unspecified ¹ | 4.1 | 7.0 | 6.5 |
| Musculoskeletal and Connective Tissue Disorders Joint related signs and symptoms ¹ | 1.9 | 1.5 | 2.3 |
| Nervous System Disorders Headaches NOS ² | 6.4 | 6.4 | 5.5 |
| Skin and Subcutaneous Tissue Disorders Rash NOS ² | 1.7 | 1.0 | 2.1 |

¹ MedDRA high level term (preferred terms): dyspeptic signs and symptoms (dyspepsia, dyspepsia aggravated, eructation, gastrointestinal irritation), upper respiratory tract infections-pathogen unspecified.

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

⁽laryngitis NOS, pharyngitis NOS, sinusitis NOS), joint related signs and symptoms (arthralgia, arthralgia aggravated, joint crepitation, joint effusion, joint swelling)

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

| | 4 - 6 Weeks Controlled | | 6 Month Controlled Trials | |
|--|------------------------|-----------------------|---------------------------|-----------------------|
| | Trials | | | |
| | Meloxicam 7.5 mg daily | Meloxicam 15 mg daily | Meloxicam 7.5 mg daily | Meloxicam 15 mg daily |
| No. of Patients | 8955 | 256 | 169 | 306 |
| Gastrointestinal | 11.8 | 18.0 | 26.6 | 24.2 |
| Abdominal pain | 2.7 | 2.3 | 4.7 | 2.9 |
| Constipation | 0.8 | 1.2 | 1.8 | 2.6 |
| Diarrhea | 1.9 | 2.7 | 5.9 | 2.6 |
| Dyspepsia | 3.8 | 7.4 | 8.9 | 9.5 |
| Flatulence | 0.5 | 0.4 | 3.0 | 2.6 |
| Nausea | 2.4 | 4.7 | 4.7 | 7.2 |
| Vomiting | 0.6 | 0.8 | 1.8 | 2.6 |
| Body as a Whole | | | | |
| Accident household | 0.0 | 0.0 | 0.6 | 2.9 |
| Edema ¹ | 0.6 | 2.0 | 2.4 | 1.6 |
| Pain | 0.9 | 2.0 | 3.6 | 5.2 |
| 0 - 1 - 10 - 11 - 12 0 - | | | | |
| Central and Peripheral Nervous System Dizziness | | | | 2.6 |
| Dizziness Headache | 1.1 | 1.6 | 2.4 | |
| 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 |
| · | 0.0 | 0.4 | 5.0 | 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 |
| · · · · · · · · · · · · · · · · · · · | | | | |

¹ WHO preferred terms edema, edema dependent, edema peripheral, and edema legs combined ² WHO preferred terms rash, rash exphematous, and rash muculo-papular combined Higher doose of meloxicam (22.5 mg and greater) have been associated with an increased risk of serious GI events; therefore, the daily dose of meloxicam should not exceed 15 mg.

The following is a list of adverse drug reactions occurring in less than 2% of patients receiving meloxicam in clinical trials involving approximately 16,200 patients.

Body as a Whole

Cardio social registraction, face edems, faigue, fever, hot flishes, malaise, syncope, weight decrease, weight increase
agina pectoris, cardiac faiture, hypernesion, hypomenion, myocardial infarction, vasculist

Cartio and Peripheral Nervous Systems—serving.

Gastrointestinal

Hernat Rate and Rkythm

Hernat Rate and Rkythm

Hernat Resistration

He

Hematologic Liver and Biliary System Metabolic and Nutritional

ALT increased, AST increased, unitarized, unitarized debythration abnormal dreaming, anciety, appetite increased, corfusion, depression, nervousness, somnolence asthma, brorchospasm, dysporea alopecia, angloedems, ballous eruption, photosensitivity reaction, prurints, sweating increased, urticaria abnormal vision, coinjunctivitis, usite preversion, diminus albuminaria, BUN increased, creatinine increased, hematuria, renal failure

Metabolic and Nutritio Psychiatric Respiratory Skin and Appendages Special Senses

Urinary System

6.2 Post Marketing Experience

6.2 Post Marketing Experience.
The following adverse reactions have been identified during post approval use of meloxicam. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to enterplaidy estimate their frequency or resultable actuals 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) extensionses of the event, (2) number of reports, or (3) strength of casals relationship to the drug. Adverse reactions reported in worldwide post marketing experience of the literature in Ludie cature turnary retendron agrantacy toxics; alterations in mod (such as mod elevation), analypiactoid reactions including short, erythern multiforms; evolution deventation; insensitial applicits, jundices, liter failure; Seeven solhonon syndome, and total exploration records.

7. DRUG INTERACTIONS

See also Clinical Pharmacology (12.3).

7.1 ACE-inhibitors

NSAIDs may diminish the antihypertensive 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_{DAR} (24%) of meloxicam was noted. The clinical significance of this interaction is not known; however, as with other NSAIDs concentual 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 meloxicam alone. Meloxicam is not a substitute for aspirin for cardiovascular prophylaxis.

7.3 Diuretics

7.3 Directics
Clinical studies, as well as post mariesting observations, have shown that NSAIDs can reduce the nartiruretic effect of furosenide and thiazides in some patients. This response has been antihuted to inhibition of real prostaglantins synthesis. However, studies with furosenide agents and reoloxican have not demonstrated a reduction in nartiruretic effect. Furosenide single and multiple dose pharmacodynamics and pharmacolivations can reduct a multiple doses of melosiciam. Nevertheless, during concordinant therapy with melosicam patients should be observed closely for signs of renal failant per lew furnings and Percunitors (Sol), as well as to essent educate directic efficacy.

7.4 Lithium

In a study conducted in healthy subjects, mean pre-dose lithium concentration and AUC were increased by 21% in subjects receiving lithium doses ranging from 804 to 1072 mg wice daily with reloxicant 15 mg every day a compared to subjects receiving lithium alone. These effects have been attributed to inhibition of renal prossiglantin synthesis by meloxicant Closely monitor patients on lithium resument for signs of lithium too occurrence and the contract of th

7.5 Methotrexate

NSAIDs have been reported to competitively inhibit methotrexate accumulation in rabbit kidney slices. Therefore, NSAIDs may reduce the elimination of methotrexate, thereby enhancing the toxicity of methotrexate list caution when meloxicam is administered conconitantly with methotrexate [see Clinical Pharmacology (12.3)].

7.6 Cyclosporine

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

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 alone.

Monitor articoagulant activity, particularly in the first few days after initiating or changing meloxicam therapy in patients receiving warfarin or similar agents, since these patients are at an increased risk of bleeding than with the use of either drug alone. Use caution when administering meloxicam with warfarin since patients on warfarin may experience changes in Rnd an increased risk of bleeding complications when a new medications is introduced goes (milked pharmacology (12.3)).

8. USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Pregnancy Category C; Category D starting 30 weeks gestation

reguancy casegory c, casegory L staring at weeks gestation.

There are no adoption and well-controlled studies in preguant women. Meloxicam crosses the placeral barrier. Prior to 30 weeks gestation, use meloxicam during preguancy only if he potential benefit in Meloxicam controlled to the properties of the properties of the potential benefit in Meloxicam controlled to the properties of the potential benefit in Meloxicam controlled to the properties of the potential benefit in Meloxicam controlled to the properties of the potential bazard to a feasi great Worlings and Percentation (2.9) and Patient Counseling Information (17.2). Teratogenic Effects

**ranogenic Effects

Meloxicam was not teratogenic when administered to pregnant rats during fetal organogenesis at oral
doses up to 4 mg/kg/day (2.6-fold greater than the maximum recommended human daily dose [MRHD]
based on body surface area [BSA] comparison). Administration of meloxicam to pregnant rabbits
throughout enthrogenesis producted in increased incidence of sepati defects of the heart at an oral
dose of 50 mg/kg/day. The me effect level was 20 mg/kg/day (26-fold greater than the MRHD based on
BSA conversion).

Nonteratogenic Effects

roomerungens. Ejects in In rais and rabbits, embryolethality occurred at oral meloxicam doses of 1 mg/kg/day and 5 mg/kg/day, respectively (0.63-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 meloxicam on labor and delivery of pregnant women are unknown. Oral administration of meloxicam to pregnant rask during late gestation through lactation increased the incidence of dystoclar delayed parturition, and decreased offspring survival at melociam doses of 10-125 mg/kg/dup or graper (at least 12.5 times lower than the maximum recommended human daily dose based on body surface area commarison).

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. Because many drugs are excreted in

human milk and because of the potential for serious adverse reactions in nursing infants from meloxican 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 exclus

As with any NSAID, caution should be exercised in treating the elderly (65 years and older). 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 overall differences in safety or effectiveness were observed between these subjects and younger subjects, and younger subjects, and younger subjects and younger subjects and younger subjects, and younger patients, but greater sensitivity of some older individuals cannot be ruled out.

Robert Empairment

No dose adjustment is necessary in patients with mild to moderate bepatic impairment. Patients with severe bepatic impairment have not been adequately studied. Since nelsociam is significantly metabolized in the liver, the use of nelsociam in these patients should be done with caution [see Warnings and Precautions (5.3) and Clinical Pharmacology (12.3)]

8.7 Renal Impairment

6.7 Renal Impairment: Patients with severe renal impairment. Patients with severe renal impairment have not been studied. The use of meloxicamin subjects with severe renal impairment have not been studied. The use of meloxicamin subjects with severe renal impairment is not recommended. Following a single does of meloxicam, there fee Ca_{man} Bisman concentrations were higher in patients with neural lands not chronic benefit higher in patients with neural lands not chronic benefit higher in patients with neural lands not chronic benefit higher than the patients with the patients of the patients with the patients of the patients of the patients of the patients with the patien

10. OVERDOSAGE

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

recommended dose; all recovered. Cholestystamine is intown to accelerate the clearance of meloxicams. Symptoms following actue NSAID overdose include lengthey, dosvariens, nusses, vontring, and epigastric pain, which are generally reversible with supportive care. Gastroiterstand bleeding can occur. Severe pointoning my result in hypertension, actue result faulture, heaping depression, come, consulsions, cardiovascular collapse, and cardiac arrest. Anaphylicatiol reactions have been tyported with therepaetic ingestion of NSAIDs, and my occur following an NSAID revolution. Patients should be managed with symptomatic and supportive care, and my occur following an NSAID revolution. Patients should be managed with symptomatic patients of NSAIDs, and my occur following an NSAID revolution. For substantial overdose or severely symptomatic patients, activated charcoal may be administered repeatedly. Accelerated removal of meloxicamby 4 gm oral doses of cholestyranine given hree times a day was demonstrated in a clinical trial. Administration of to choestyranine may be useful following an overdose. Forced distress, skilalinization of urine, hemodulysis, or hemperfusion may not be useful.

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

11. DESCRIPTION

Meloxicam, an oxicam derivative, is a member of the emolic acid group of nonsteroidal anti-inflammancy drugs (NSAIDs). Each biblect contain 5.5 mg or 15 mg meloxicam for oral administration (Meloxicam) is chemically designated as 1-ph/qnvs2-rednely-16-f-melyty2-linalys). 24:1-12-bencontainer-8-carboxamide-11-dioxide. The molecular weight is 351.4. Its empirical formula is C₂(11)P₃(N)C₃C₃ and thus the following surecural formula.



Meloxicam is a pastel yellow solid, practically insoluble in water, with higher solubility observe strong acids and bases. It is very slightly soluble in methanol. Meloxicam has an apparent partition coefficient (log $P_{\text{lapp}} = 0.1$ 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

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

12. CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

The mechanism of action of meloxicam, like that of other NSAIDs, may be related to prostaglandin synthesase (cyclooxygeness) inhibition which is involved in the initial steps of the arachidonic acid cascade, resulting in the reduced formation of prostaglandins, thromboxanes and prostacylin. It is not completely understood how reduced synthesis of these compounds results in therapeutic efficacy.

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

12.3 Pharmacokinetics

Asonption.

The absolute bioavailability of meloxicam capsules was 89% following a single oral dose of 30 mg compared with 30 mg IV boits injection. Following single intravenues doses, dose-proportional compared with 30 mg IV boits injection. Following single intravenues doses, dose-proportional behavior in the proportional control of t

Table 3 Single Dose and Steady-State Pharmacokinetic Parameters for Oral 7.5 mg and 15 mg Meloxicam (Mean and % CV)¹

| | | Steady State | | | Single Dose | |
|---------------------------------|----------|---------------------|----------------|--------------------|------------------------|--------------------------------|
| Pharmacokinetic Paramete CV) | rs (% | Healthy male adults | Elderly males | Elderly females | Renal failure (Fasted) | Hepatic insufficiency (Fasted) |
| | | (Fed)2 | (Fed)2 | (Fed) ² | | |
| | | 7.5 mg 3 tablets | 15 mg capsules | 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) |
| Martin | III 1 | 147(22) | 1E (43) | 10 (20) | 26 (44) | 14 (20) |

 $^{^1}$ The parameter values in the table are from various studies 2 not under high fat conditions 3 Meloxicam tables 3 Meloxicam tables 3 Vz/f = Dose(AUC-Kel)

rood and Amacid Effects.

Administration of meloxic am capsules following a high fat breakfast (75 g of fat) resulted in mean perform levels (e.g., e.g., and period fat, e.g., and fat, e.g., and e

The mean volume of distribution (*15) of real-ord-camb a approximately 10 L. Meltodc and 1:-90.4% bound to human plane protein (primetry) abunday within the therapeut of so range. The fraction of protein binding is independent of drug concentration over the clinically relevant concertration range, and tecreases to "99% in patients with real disease. Melockcampeneration into human red blood cells, after oral dosing, is less than 10%. Following a radiolabeled dose, over 90% of the radioactivity detected in the planent was present as unchanged redoction.

Meloxicam concentrations in synovial fluid, after a single oral dose, range from 40% to 50% of those in plasms. The free fraction in synovial fluid is 2.5 times higher than in plasms, due to the lower abure content in synovial fluid as compared to plasms. The figrificance of this penetration is unknown.

Metabolism

Meloxicam is extensively metabolized in the lizer. Meloxicam teadolists include 5'-carboxy meloxicam (66%) of dose), from P-45 mediated metabolists formed by oxidation of an intermediate metabolite 5'-hydroxymethyl meloxicam which is also excreted to a lesser extent (9% of dose), in vitro studies indicate that CVPE2G (systorborn P-50 metabolites) (engage navine) plays a minoritant role in this metabolic pathway with a minor comribation of the CVPAA4 isozyme. Patters' peroxidate extivity is probably responsible for the other to metabolites which excure for 16% and 46% of the admissrated dose, respectively. All the four metabolites are not lizewist have any in vito pharmacological activity. Exercision

Excretion Melosicam excretion is predominantly in the form of metabolites, and occurs to equal extents in the urine and feese. Only traces of the unchanged parent compound are excreted in the urine (0.2%) and feese (1.6%). The stear of the urinary excretion was confirmed for unabeled multiple 7.5 mg does 0.5%, 6%, and 1.3% of the dose were found in urine in the form of melosicam, and the 5-hydroxymethyl and 5'c-arboxy metabolites, respectively. There is significant ballary and/or enterplace accretion of the drug. This was demonstrated when oral administration of cholestyramine following a single IV does of melosicam decreased the AUC of melosicam by 50%.

inconcame creases use ACL on neconcamy 50%.

The mean elimination half-life (1), 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.

Special Populations

Geriatric

Gentanic.

Eiderly mules (265 years of age) exhibited meloxicam plasma concentrations and steady-state pharmacokimetics similar to young males. Eiderly formatic (265 years of age) had a 47% higher AUC₁₈ to make the contraction of the deletyle frames, the aboves event profile was comparable for both elderly patient populations. A smaller free fraction was found in elderly frames, the contraction of the contr

Young females exhibited slightly lower plasma concentrations relative to young males. After single doses of 75 mg meloxicam, the mean elimination half-life was 19.5 hours for the female group compared to 23.4 hours for the mile group. At steady stars, the data were similar (179 hours vs 21.4 hours). This pharmacolisteic difference due to gender is likely to be of little clinical importance. There was linearity of pharmacolisteic and no appreciable difference in the _mag. of T_max across.

Hepatic Impairment

English (mailtean). Following a single 15 mg dose of meloxicam there was no marked difference in plasma concentration in patients with mild (Child-Pugh Class I) or moderate (Child-Pugh Class II) bepaste impairment of the control of

Renal Impairment

Meloxicampharmacokinetics have been investigated in subjects with mild and moderate read impairment. Total drug glisarue concernations of meloxicam decreased and total clearance of meloxicam impairment. Total drug glisarue concernations of meloxicam decreased and total clearance of meloxicam impairment may be due to increased fraction of unbound meloxicam clearance in subjects with renal impairment may be due to increased fraction of unbound meloxicam which is available for bepatic metabloisam and subsequent excretion. No dosage adjustment is necessary in patients with surface for meloxicam in subjects with severe renal impairment and to be then adequately suddied. The use of meloxicam in subjects with severe renal impairment is not recommended [see Warnings and Precountors (5.6) and Use in Specific Populations (8.7)].

Hemodialysis

Intermonatives:

Following a single dose of meloxicam, the free C_{max} plasma concentrations were higher in patients with renal failure on chronic bemodalysis (1% free fraction) in comparison to healthy volunteers (3.% free fraction). Hemodalysis did not love the total drug concentration in plasms; therefore, additional doses are not necessary after hemodalysis. Meloxic and is not dislayable: [see Dosage and Additional doses are not necessary after hemodalysis. Meloxic and is not dislayable: [see Dosage and the Mallimatronic C.3]. Wormings and Precusations (5.5) on the in Specific Populations (6.7):

Drua Interactions

Drug Interactions
Applie: When meloxicam is administered with aspirin (1000 mg three times daily) to healthy volunteers, it tended to increase the AUC (10%) and contact (10%) of meloxicam. The clinical significance of this interaction is not incoming fear by interactions (7.2)).
Cholestyonnine: Perteamment for four days with cholestyrumine significantly increased the clearance of meloxicamby 50%. This resulted in a decrease in true, from 120 bours to 12.5 bours, and a 35% reduction in AUC. This suggests the existence of a recirculation pathway for meloxicam in the gastroitestial rate. The clinical relevance of this interactions have there exhall be a superior of the interaction has not been established.

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

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.

Lithium: In a study conducted in healthy subjects, mean pre-dose lithium concentration and AUC were increased by 21% in subjects receiving lithium doses ranging from 804 to 1072 mg twice daily with melouscan IS mg QD every day as compared to subjects receiving lithium alone [see Drug Interaction (7-64)].

(C-4)). Whethorexate: A study in 13 rheumatoid arthritis (RA) patients evaluated the effects of militiple doses of meloxicam on the pharmacokinetics of methorexate taken once weekly. Meloxicam did not have a significant effect on the pharmacokinetics of single doses of methorexate. In vitro, methorexate did not displace meloxicam from its human serum binding sites [see Drug Interactions (7:5)].

displace meloxican from its human serum bit anticogular effect of warfarin was studied in a group of healthy subjects received or meloxican mode and excogular effect of warfarin was studied in a group of healthy subjects received and in the produced an INR (international Normalized Radio) between 1 and 1.8. In these subjects, neloxic and for a other warfarin parmacolateries and average anticoagular effect of warfarins as effectives and the subject is shown an increase in INR from 1.5 in 2. Leations bould use used when antistratering meloxicat may warfarin sire patients on warfarin may experience changes in INR and an increased risk of bleeding complications when a new medicalization is involuced from Emphasization.

13. NONCLINICAL TOXICOLOGY

Last Customogeness, insuparatives in impairment of Ferning Correlangements. There was no increase in numor incidence in long-serm carcinogenicity studies in rats (104 weeks) and riscre (69 weeks) administrated inclusion and road does so up to 6.0 mig (agilty) in rats and death) of the control on stay with themsel hypothyc size and an in vive intermedies that in more been murrow, impairment of Fernings. Melosciacum data for impair male and from le retailing in rats as a coal doese up to 9 algorithm in the control of the contro

14.1 Osteoarthritis and Rheumatoid Arthritis

In the use of melocition for the resource of the signs and symptoms of osteoarthritis of the lose and hip was evaluated in a 12-week, double-blint, controlled trial. Meloxicam (3.75 mg. 7.5 mg., and 15 mg daily was compared to placebo. The flow primary endpoins were investigant's 50 plack assessment, patient global a

The use of meloxicam for the management of signs and symptoms of osteoarthritis was evaluated in double-blind, active-controlled trials outside the U.S. ranging from a week' to 6 months' doutsion. Here see tails, the efficacy of meloxicam, indoes not 1.5 miles from the second control prioration 20 migday and cloferac SR 100 mg/day and consistent with the efficacy seen in the U.S. ratio.

trial.

The use of meloxicam for the treatment of the signs and symptoms of rheumatoid arthrids was evaluated in a 12-week, double-blind, controlled multitudional trial. Meloxicam (7.5 mg, 1.5 mg, and 2.5 mg daily) was compared to placebox. The primary endpoint in this study was the ACR20 response rate, a composite measure of clinical, laboratory, and functional measures of RA response. Patients receiving meloxicam 27.5 mg and 15 mg daily showed significant improvement in the primary endpoint compared with placebo. No incremental benefit was observed with the 22.5 mg dose compared to the 15 mg dose.

16. HOW SUPPLIED/STORAGE AND HANDLING

Molosicum Tables US9 are available as lighty rellow, round, flux uncound tables containing molosicum 15 mg. and tables containing molosicum 15 mg. The 7.5 mg tables its impressed with letter U and L nones side and tables containing molosicum 15 mg. The 7.5 mg tables its impressed with letter U and L nones side and tablet code 7.5 on the other side. The 15 mg tables its impressed with letter U and L nones side and tablet code 15 on the other side. The Molosicum Tables USP 7.5 mg are available as follows:

NDC 29300-124-13; Bottles of 30 NDC 29300-124-01; Bottles of 100

NDC 29300-124-10: Bottles of 1000

NDC 29300-124-50; Bottles of 5000

Meloxicam Tablets USP 15 mg are available as follows:
NDC 29300-125-13; Bottles of 30

NDC 29300-125-01; Bottles of 100

NDC 29300-125-10; Bottles of 1000 NDC 29300-125-50; Bottles of 5000

Store at Controlled Room Temperature 20^{0} - 25^{0} C (68^{0} - 77^{0} F) [See USP]. Keep Meloxicam Tablets USP in a dry place

Dispense tablets in a tight conta

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

17. PATIENT COUNSELING INFORMATION

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

1/24 Curantwockular Euercs
NSAIDS including moisscam tablets, may cause serious CV side effects, such as Mi or stroke, which
may result in hospitalization and even death. Although serious CV events can occur without worning
weakbers, sluring of speech, and should also for medical ability dece when observing any indicative sign or
symptoms. Patients should be apprised of the importance of this follow-up [see Warnings and
The Precentings 62,31].

17.3 Gas trointes tinal Effects

I/A Gustromersime Tructo.

NatIOS including protocae mabbes, can cause GI disconfort and, rarely, serious GI side effects, such as ulcers and breeding, which may result in hospitalization and even teath. Although serious GI signs and sympoms of ulcerations and bleefing, and should as for medical advice when observing any indicative signs or sympoms including epigantic pain, dyspepsia, melena, and hematemesis. Datients should be apprised of the importance of his follow-up for worming and Precumson (GZ)).

17.4 Hepatotoxicity

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

17.5 Adverse Skin Reactions

17.5 Adverse Skin Reactions
NSAIDS, including melosicam tables, can cause serious skin side effects such as exfoliative dermatitis, Stevens-Johnson Syndrome (SIS), and toxic epidermal necrolysis (TEN), which may result in hospitalization and even death. Although serious skin reactions may occur without warring, patients should be altert for the signs and symptoms of skin rash and blisters, fever, or other signs of hypersensitivity such as itsching, and should ask for medical advice when observing any indicative signs or symptoms. Advise patients to stop the drug immediately if they develop any type of rash and contact their physicians as soon as possible (see Warrings and Prezentions (S.B).

17.6 Weight Gain and Edema

Advise patients to promptly report signs or symptoms of unexplained weight gain or 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 (5.7)].

17.8 Effects During Pregnancy tation, meloxicam should be avoided as premature closure of the ductus

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

UNICHEM LABORATORIES LTD.

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



13003755 R-03-03/2011

Medication Guide for Non-Steroidal Anti-Inflammatory Drugs (NSAIDs.)

Settlements Gaine for Four-Settlement Ami-immunitary Drugs (ISSALIDS.)

(ISSet the end of his Medication Gaide for a list of prescription ISSAID medicines.)

What is the most important information I know about medicines called Non-Steroidal Anti-Inflammatory Drugs (ISSAIDS)

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

with longer use of NSAID medicines
 in people who have heart disease

NSAID medicines should never be used right before or after a heart surgery called a "coronary artery bypass graft (CABG)." NSAD medicines can cause uclers and bleeding in the stomach and intestines time during treatment. Ulcers bleeding:

• can happen without warring symptoms
• my cause death

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

taking medicines called "corticosteroids" and "anticoaculants"

- longer use
 smoking
 drinking alcohol
 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

What are Non-Steroidal Anti-Inflammatory Drugs (NSAIDs)?

NSAID medicines are used to treat pain and redness, swelling, and heat (inflammation) from medical

NSAID medicines are used to treat pain and reciness, swicconditions 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

- Tell your healthcare provider:

 about all of your medical conditions.

 about all of your medical conditions.

 NSAIDs and some other medicines can impract with each other and cause serious side effects. Keep all sit of your medicines to show to your healthcare provider and pharmacist.

 If you are pregament, NSAID medicals should not be used by preganat and women late in their pregameny.

 If you are breastfeeding, Talk to your doctor.

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

- What are the possible side effects of Non-Steroidal Sections kielder:

 heart attack:

 heart attack:

 strole is strole in the strole is strole in the s

Other side effects include:

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

 shormers of breath or trouble breathing
 chest pain
 exhest pain
 weakness in one part or side of your body
 slurred speech
 swellings of the face or throat

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

- nausea more tired or weaker than usual
- itching vour skin or eyes look yellow

- your skin or eyes look yellow stomach pain flu-like symptoms wmit blood there is blood in your bowel movement or it is black and sticky like tar unusual weight gain skin rash or blusters with fever swelling of the arms 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.

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 (NSAIDs)

 Aspirin is an NSAID medicine but it does not increase the chance of a heart attack. Aspirin can cause bleeding in the brain, stomach, and intestines. Aspirin can also cause ulcers in the stomach and
- the country in the train, shortern, and mesures. Aspirin can also cause diversing in stomach and intestines.

 Some of these NSAID medicines are sold in lower doses without a prescription (over-the-counter). Talk to your healthcare provider before using over-the-counter NSAIDs for more than 10 days.

NSAID medicines that need prescription

Generic Name Tradename
Celecoxid Celebrax
Celetria Catalian, Voltaren, Arthrotec (combined with misoprostol)
Diffunisal
Dolobid Celine(Lodine XL
Fenoproten Maifon, Nalion 200
Flurbipotoren Anada
Anada Carallam Voltarea, Arthroise (combined with misoprostol)
Dolobbid
Lodine, Lodine XL
Naffon, Nalfon 200
Ansaid
Motrin; Tab
Profece, Nicoprofera* (combined with hydrocodone), Combunox (combined with oxycodone)
Indoor, Indoories, SR, Indo-Lemmon, Indomethagan
Toradol
Honosei
Honos Duprofen Profen, Y....
Indomethacin Adoctin, Radicin SR, Itaa.
Convail
Ketorporfen
Ketorolac
Mefenauric Acid Ponstel
Meloxicam
Naprosen
Naprosen
Naprosen
Piroxicam
Pi lbupro fen

Manufactured by:
UNICHEM LABORATORIES LTD.

Pilerne Ind. Estate,

Pilerne, Bardez, Goa 403511, India



Rochelle Park, NJ 07662 13002550 R-02-09/2010

PACKAGE LABEL - MELOXICAM 7.5 MG TABLET



PACKAGE LABEL - MELOXICAM 15 MG TABLET



| melo | LOXICAM exicam tablet | | | | | | | | | |
|--|---|--|---|-----------------|-----------------|-----------|------|-----------|-------------|-------------|
| Pro | oduct Informatio | on | | | | | | | | |
| Pro | duct Type | HU | MAN PRESCRIPTIO | ON DRUG | Item Cod | le (Sourc | e) N | DC:1659 | 0-434(ND | C:29300-124 |
| Rou | te of Administrati | on OF | IAL | | | | | | | |
| Act | ive Ingredient/ | Active Moiety | | | | | | | | |
| | | Ingred | lient Name | | | | Basi | s of Str | ength | Strengtl |
| MEL | OXICAM (UNII: VG | 2QF83CGL) (MELC | XICAM - UNIEVG | QF83CGL |) | h | ŒLOX | ICAM | | 7.5 mg |
| Ina | ctive Ingredien | ts | | | | | | | | |
| CRO | SPOVIDONE (UNII: | 68401960MK) | Ingredient Na | me | | | | | | Strength |
| | TOSE MONOHYDE | | 7Q8I5X) | | | | | | | |
| MAC | NESIUM STEARAT | TE (UNIL 70097M6 | B0) | | | | | | | |
| SILI | CON DIO XIDE (UNI | II: ETJ7Z6 XBU4) | | | | | | | | |
| TRIS | SODIUM CITRATE | DIHYDRATE (UNI | EB22547B95K) | | | | | | | |
| | IDONE K29/32 (UN LULOSE MICROC | | | | | | | | | |
| | | | | | | | | | | |
| | duct Character | istics yellow | Sco | ore | | | | no | score | |
| Cole | or . | | Sco | | | | | no 7m | | |
| Colo | or pe | yellow | Siz | | | | | 7m | | |
| Colo Shap Flav Con | or pe or tains | yellow | Siz | e | | | | 7m | ım | |
| Cole Shap Flav Con | or pe or tains kaging | yellow ROUND | Siz | e print Code | | | | 7m U;i | im L;7;5 | |
| Cold Shap Flav Con | or pe or tains kaging Item Code | yellow ROUND | Siz Imj e Description | e print Code | e urketing S | Start Dat | ie. | 7m U;i | im L;7;5 | End Date |
| Cold Shap Flav Con Pac | or pe or tains kaging | yellow ROUND | Siz Imj e Description LE | e print Code | | Start Dat | ie. | 7m U;i | im L;7;5 | End Date |
| Cold Shap Flav Con Pac | pe pe or tains :kaging Item Code DC:16590-434-10 | yellow ROUND Packag 10 in 1 BOTT | Siz Imj e Description LE LE | e print Code | | Start Dat | e | 7m U;i | im L;7;5 | End Date |
| Cole Shap Flav Con Pac # 1 Ni 2 Ni 3 Ni | pe pe tains ckaging Item Code DC:16590-434-10 DC:16590-434-20 | yellow ROUND Packag 10 in 1 BOTT 20 in 1 BOTT | e Description LE LE | e print Code | | Start Dat | æ | 7m U;i | im L;7;5 | End Date |
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| Pac Flav Con Pac # Ni | r pe cor tains kaging Item Code DC:165300-434-10 DC:165300-434-20 DC:165300-434-30 DC:165300-434-30 DC:165300-434-30 | yellow ROUND Packag 30 in 1 BOTT 20 in 1 BOTT 30 in 1 BOTT 56 in 1 BOTT 60 in 1 BOTT | Siz Imp e Description LE LE LE LE | e print Code | | Start Dat | ee | 7m U;i | im L;7;5 | End Date |
| Cole Shap Flav Con Pac # 1 Ni 2 Ni 3 Ni 4 Ni 5 Ni 6 Ni 7 Ni | pe or ratins -kaging -kaging -tem Code -b0:16590-434-10 -b0:16590-434-20 -b0:16590-434-30 -b0:16590-434-30 -b0:16590-434-60 -b0:16590-344-60 -b0:16590-344-60 | yellow ROUND Packag 20 in 1 BOTT 20 in 1 BOTT 30 in 1 BOTT 30 in 1 BOTT 56 in 1 BOTT 56 in 1 BOTT 50 in 1 BOTT | Siz Imp | e print Code | | Start Dat | ·e | 7m U;i | im L;7;5 | End Date |
| Cole Shap Flav Con Pac # 1 Ni 2 Ni 3 Ni 4 Ni 5 Ni 6 Ni 7 Ni | r pe cor tains kaging Item Code DC:165300-434-10 DC:165300-434-20 DC:165300-434-30 DC:165300-434-30 DC:165300-434-30 | yellow ROUND Packag 30 in 1 BOTT 20 in 1 BOTT 30 in 1 BOTT 56 in 1 BOTT 60 in 1 BOTT | Siz Imp | e print Code | | Start Dat | ce | 7m U;i | im L;7;5 | End Date |
| Colo Shaji Flav Con Pac # | pe or ratins -kaging -kaging -tem Code -b0:16590-434-10 -b0:16590-434-20 -b0:16590-434-30 -b0:16590-434-30 -b0:16590-434-60 -b0:16590-344-60 -b0:16590-344-60 | yellow ROUND Packag 10 in 1 BOTT 20 in 1 BOTT 30 in 1 BOTT 56 in 1 BOTT 180 in 1 BOTT 280 in 1 BOTT | Siz Imp | e print Code | | Start Dat | ee | 7m U;i | im L;7;5 | End Date |
| Cole Shal Flav Con Pace # 1 Ni 2 Ni 3 Ni 4 Ni 5 Ni 6 Ni 7 Ni 8 Ni Ma | pe pe per tains -kaging Item Code -Citi530-434-10 -DC:16530-434-20 -DC:16530-434-30 -DC:16530-434-30 -DC:16530-434-82 -DC:16530-434-82 -DC:16530-434-82 | yullow BOUND Packag 10 in 1 BOTT 20 in 1 BOTT 56 in 1 BOTT 90 in 1 BOTT rmation | Siz Imp | e print Code | arketing S | Start Dat | | Ma | um L:7:5 | End Date |

| Produ | ct Informatio | n | | | | | | |
|---|--|---|----------------------|----------|---------------|-------------|----------------|--------------|
| Produc | t Type | HUMAN P | RESCRIPTION DRU | G Item C | Code (Source | NDC:165 | 90-469(NI | OC:29300-125 |
| Route o | f Administratio | m ORAL | | | | | | |
| Active | Ingredient/A | Active Moiety | | | | | | |
| | | Ingredient N | lame | | | Basis of St | trength | Strengtl |
| MELOX | ICAM (UNII: VG2 | QF83CGL) (MELOXICA) | 4 - UNIEVG2QF83C | GL) | М | ELOXICAM | | 15 mg |
| Inactiv | e Ingredient | s | | | | | | |
| | | | edient Name | | | | | Strength |
| | OVIDONE (UNIE | | | | | | | |
| | | ATE (UNII: EWQ57Q8153 E (UNII: 70097M6130) | 1) | | | | | |
| | N DIO XIDE (UNI | | | | | | | |
| | or Dio Albe (Com | | | | | | | |
| | DILIM CITRATE I | HINDRATE (UNII- B225) | (7895K) | | | | | |
| POVIDO | NE K29/32 (UN | HYDRATE (UNI: B225- II: 390 RMW2PEQ) RYSTALLINE (UNI: OP1 | | | | | | |
| PO VIDO CELLUI Produ | NE K29/32 (UN | II: 390 RMW2PEQ) RYSTALLINE (UNII: OPI is tics yellow | | | | | no score | |
| POVIDO CELLUI Produ Color Shape | ONE K29/32 (UN LOSE, MICROCE | II: 390RMW2PEQ) RYSTALLINE (UNII: OPI | R32D6 1U) | | | | 12mm | |
| POVIDO CELLUI Produ Color Shape Flavor | ONE K29/32 (UN LOSE, MICROCE ct Characteri | II: 390 RMW2PEQ) RYSTALLINE (UNII: OPI is tics yellow | R32D61U) Score | t Code | | | | |
| POVIDO CELLUI Produ Color Shape Flavor | ONE K29/32 (UN LOSE, MICROCE ct Characteri | II: 390 RMW2PEQ) RYSTALLINE (UNII: OPI is tics yellow | R32D6 IU) Score Size | t Code | | | 12mm | |
| Produ Produ Color Shape Flavor Contain | ONE K29/32 (UN LOSE, MICROCE ct Characteri is | B: 390 RMW2PEQ) FYSTALLINE (UNIR OPE SITICS yellow CAPSULE | Score Size | t Code | | | 12mm | |
| Produ Cell U Produ Color Shape Flavor Contain Packa | ONE K29/32 (UN LOSE, MICROCS Ct Characteri is | E-390 RMW2PEQ) YYSTALLINE (UNIR OPE SSICS yellow CAPSULE Package Desi | Score Size | | og Start Date | | 12mm U;L;15 | End Date |
| Produ Cellu Produ Color Shape Flavor Contain Packa | ONE K29/32 (UN LOSE, MICROCI Ct Characteri is ging Item Code 16590-469-15 | E 390 RAW2PEQ) VYSTALLINE (UNR OPE Stics yellow CAPSULE Package Dess 15 in 1 BOTTLE | Score Size | | ng Start Date | | 12mm U;L;15 | End Date |
| Produ Cellu Produ Color Shape Flavor Contain Packa | ONE K29/32 (UN LOSE, MICROCE Ct Characteri is ging Item Code 16:590-469-15 16:590-469-28 | E 390BAW2FEQ) YYSTALLINE (UNE OPE Stics yvllow CAPSULE Package Desi 15 in BOTTLE 28 in BOTTLE | Score Size | | og Start Date | | 12mm U;L;15 | End Date |
| Produ Cell U Produ Color Shape Ravor Contain Packa 1 NDC: 2 NDC: 3 NDC: | DNE K29/32 (UN LOSE, MICROCE Ct Characteri is ging Item Code 16590-469-15 16590-469-28 | Package Desi | Score Size | | og Start Date | | 12mm U;L;15 | End Date |
| Produ Cell.UI Cell.UI Color Shape Flavor Contain Packa # NDC: 2 NDC: 3 NDC: 4 NDC: | ONE K29/32 (UN LOSE, MICROCE Ct Characteri is ging Item Code 16:590-469-15 16:590-469-28 | E 390BAW2FEQ) YYSTALLINE (UNE OPE Stics yvllow CAPSULE Package Desi 15 in BOTTLE 28 in BOTTLE | Score Size | | sg Start Date | | 12mm U;L;15 | End Date |
| Produ Cellui Cellui Color Shape Flavor Contain Packa # 1 NDC: 2 NDC: 3 NDC: 4 NDC: 5 NDC: | DNE K29/32 (UN LOSE, MICRO CE CE Characteri is ging Item Code 16590-469-15 16590-469-30 16590-469-30 | Package Des Package Des Package Des B in BOTTLE So is BOTTLE So is BOTTLE | Score Size | | og Start Date | | 12mm U;L;15 | End Date |
| Produ | DNE K29/32 (UN OSE, MICROCI Ct Characteri Iss ging Item Code 15/90-469-15 16/590-469-28 16/590-469-30 16/590-469-36 | Package Designation of the Bottle Package Designation of the Bottle B in 180TILE 56 in 180TILE 56 in 180TILE 59 in 180TILE | Score Size | | ng Start Date | | 12mm U;L;15 | End Date |
| Produ Cellui Produ Color Raver Contain Packa I NDC: NDC: NDC: NDC: Marl | DNE K29/32 (UN. OSE, MICROCI Ct Characteri Iss ging Item Code 65/50-469-15 55/90-469-30 65/90-469-30 65/90-469-30 | Package Designation of the Bottle Package Designation of the Bottle B in 180TILE 56 in 180TILE 56 in 180TILE 59 in 180TILE | Score Size Imprin | Marketin | ng Start Date | e M | 12mm U;L;15 | End Date |