## MELOXICAM- meloxicam table Carilion Materials Management

IBGBLIGHTS OF PRINCEIBING INFORMATION
Three highlights do not include all the information needed to use MELOXICAM TABLETS safely and
effectively. See full proceeding information for NELOXICAM TABLETS.

MELOXICAM tablets, for oral use

MANISCE, 2018 OF SERVICE C. GEORGY CACCULA NO ROLL AND CONTROL EXTENSION OF SERVICE C. S

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Solida Processins, Conference of Trombusic Curare (3.1)

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# WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS

WARNING, BING OF SERIOUS CARIODVASCULAR AND GASTRONTESTINAL CONTROLLED TO A CO

## 1 INDICATIONS AND USAGE

1.1 Osteoarthritis (OA)

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

1.2 Rheumateid Arthritis (RA)

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

1.3 Juvenile Rheumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course Meloxican tablets are indicated for relief of the signs and symptoms of pauciarticular or polyarticular course Juverile Rheumaniel Arthritis in patients who weigh x60 kg [see Dosage and Administration (2.4) and Chited Studies (14.2)].

## 2 DOSAGE AND ADMINISTRATION

2.1 General Design Instructions
Carefully consider the powerful healths and ricks of endorsteam takes and other systems registers
Carefully consider the powerful healths and ricks of endorsteam takes and other systems register
Consistent with influsional pointer measures goods be the Warnings and Proceedings (5).

After observing the response to initial therapy with melostican tablets, adjust the dose to sait an included pattern serving expense.

In adults, the maximum recommended daily oral dose of metoxicam tablests are 15 mg regardless of formulation. In patients with homofullysis, a maximum daily dosage of 7.5 mg is recommended (see Use in Specific Population (8.7) and Clinical Pharmacology (72.5).

Metoxicam tablets may be taken without regard to fining of meals.

A second attention may be searn without regard to turning or meas.

2.2 On teneraltritis

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

2.3 Rheumatoid Arthritis

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

2.4 Juvenile Rheumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course

For the treatment of jovenile rheumanoid arthritis, the recommended or al dose of meloxicam tablets is 7.5 mg once daily in children who swigh >60 kg. There was no additional benefit demonstrated by increasing the dose above 7.5 mg in clinical tablets.

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

2.5 Renal Impairment

The use of meloxicam in subjects with severe renal impairment is not recommended.

In patients on hermolialysis, the maximum dosage of meloxicam is 7.5 mg per day [see Clinical Phormocology (2014)].

rnamencology (12:3).

2.6 Non-interchangeability with Other Formulations of Meloskicam
Meloskican tables have not shown equivalent systemic exposure to other approved for
meloskican. Therefore, meloskican tables are not interchangeable with other formulation
meloskican produce verifie to testi militigam steeping is the same. Do not substitute
steepings of meloskican tables with other formulations of oral meloskican product.

3 DOSAGE FORMS AND STRENGTHS

### 4 CONTRAINDICATIONS

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## 5 WARNINGS AND PRECAUTIONS

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they occur.

There is no consistent evidence that concurrent use of applint miniguages the increased risk of serious CV thrombotic events associated with NSAID use. The concurrent use of aspirit midra and NSAID, such as melonicam increases the risk of serious gastrointestinal (Cri) events leve hibrarings and Procustors (5-2)).

an effective file of the control of

Avoid the use of meloxicam in patients with a recent MI unless the benefits are expected to outweight the risk of recurrency CV thrombodic revens. If meloxicam is used in patients with a recent MI, monitor patients or signs of cardiac ischemia.

no idea for correst CV thresholder covers. In the control of the c

nature than the state of the st

cancar available of the planting less the an Apopter propasation (to a) and Lancar transmissing (12.1). SALTIP, including melocician, caniford in two ones or worsening of proceding hyperension, either of which may combine to be interested inclusion of CV weres. Plantin lange againstant conversion engage (ACL) shibitors, fluided districts, or floor districts may be be impaired response to these throughout the control of th

## 5.5 Heart Failure and Edema

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increases in serum potassium concentration, including hyperkalemia, have been reported with use of NSAIDs, even in some patients without renal impairment. In patients with normal renal function, these effects have been attributed to a hyporerismic—hypoildostronism state.

offices have been attributed to a pipporenamic—hypoatosisteronicins state.

5.7 Anaphylactic Reactions
Mediociam has been associated with anaphylactic reactions in patients with and without known hypersensitivity to melociciam and in patients with aspirits-tensitive aothma [see Controlledications (4) and Witwings and Preconditions (5.8)].

Seek emergency help if an anaphylactic reaction occurs

Sock energy by high II an anaphylactic reaction occurs.

3. Exercitation of Manian Related to Applies Sensitishly

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may leaked cheater informationation complication by anaphylyse, severe, promotally fault bronchesposes,
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the symptoms of authors.

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3.6 section of the storm of the sto

# the utility of diagnostic signs in detection; infections. SDI Laboratory Municipal Because serious Ci Bheeding, hopatonicity, and read injury can occur without warning symposus or the control of the

6 ADVERSE REACTIONS
The following adverse reactions are discussed in greater detail in other sections of the labeling:
Cardinavacach: Translation Extens [see those Warning and Warning and Precursions (5,1)]
High particular properties of the processing of the processing and Precursions (5,2)]
High particular processing and Precursions (5,3)]
High practical processing and Precursions (5,3)]
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High Tallian and Extension for Marriagon (4,3)
High Tallian and Extension for Marriagon (4,3)
Analyticatic Resistance for Versions (4,5)]
Serions (3,6)
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## 6.1 Clinical Trials Experience

and may are reflected as one-observed inpractice.

Andreads

General and the service of the serv

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

Table 1b depicts adverse events that 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	7.5 mg daily	15 mg daily	Diclofenac 100 mg daily
No. of Patients	157	154	156	153
Gas tro intes tinal	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 <sup>1</sup>	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				
Pharyogitis	1.3	0.6	3.2	1.3
Upper Respiratory Tract Infection	1.9	3.2	1.9	3.3
Skin				
Rash <sup>2</sup>	2.5	2.6	0.6	2.0

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

	Placebo	Meloxicam 7.5 mg daily	Meloxican 15 mg daily
No. of Patients	469	481	477
Gastro intestinal Disorders	14.1	18.9	16.8
Abdominal pain NOS <sup>2</sup>	0.6	2.9	2.3
Dyspeptic signs and symptoms 1	3.8	5.8	4.0
Nausea <sup>2</sup>	2.6	3.3	3.8
General Disorders and Administration Site Cond	itions	•	•
Influenza-like illness <sup>2</sup>	2.1	2.9	2.3
Infection and Infestations			
Upper respiratory tract infections- pathogen class unspecified <sup>1</sup>	4.1	7.0	6.5
Musculoskeletal and Connective Tissue Disor	ders		
Joint related signs and symptoms 1	1.9	1.5	2.3
Nervous System Disorders			
Headaches NOS <sup>2</sup>	6.4	6.4	5.5
Skin and Subcutaneous Tissue Disorders			
Rash NOS <sup>2</sup>	1.7	1.0	2.1
MedDRA high level term (preferred terms): dyspeptic sig	ns and symptoms (dys.	реркія, дукреркія а	ggravated,

The adverse events that occurred with meloxicam in 22% of patients treated short-term (4 to 6 weeks) and long-term (6 months) in active-como fled osteoarthritis trials are presented in Table 2.

Table 2Adverse Events (%) Occurring in ≥ 2% of Meloxicam Patients in 4 to 6 Weeks and 6 Month Active-Controlled
Octoor-thelist Trials

	4 to 6 Weeks Controlled Trials	ı	6 Month Controlled Trials	
	Meloxicam	Meloxicam	Meloxicam	Meloxicar
	7.5 mg daily	15 mg dails	7.5 mg daily	15 mg dail
No. of Patients	8955	256	169	306
Gastro intestinal	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
Ederma*	0.6	2.0	2.4	1.6
Pain	0.9	2.0	3.6	5.2
Central and Peripheral Nervous System Dizziness	1.1	1.6	2.4	2.6
Headache	2.4	2.7	3.6	2.6
Hematologic Anemia	0.1	0.0	4.1	2.9
Musculoskeletal				
Arthralgia	0.5	0.0	5.3	1.3
Backpain	0.5	0.4	3.0	0.7
Psychiatric				
Insomnia	0.4	0.0	3.6	1.6
Respiratory Coughing	0.2	0.8	2.4	1.0
Upper respiratory tract infection	0.2	0.0	8.3	7.5
Skin				
Pruritus	0.4	1.2	2.4	0.0
Rash <sup>†</sup>	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

I bill op referred som som, hand synthesismen, set from norder-project combined.

Highly donous of reductions (2.75 any of grouns) where associated out this nicessand disk of nestional Crievens, therefore, the stilly donous of multi-crievan client and nest exceed 15 mg.

Politicisc

The following is a list of adverse drug reactions occurring in <2% of patients receiving met clinical trials involving approximately 16, 200 nations.

cinical trials involving approximately 16,2	OU DAILYES.
Body as a Whole	allergic reaction, face edems, fatigue, fever, bot flushes, mulaise, syncope, weight decrease
	sagina percoris, cardiac falture, hypermenion, hyposemion, myocardial interction, vascutifis
Central and Peripheral Nervous System	
	Solids, dry routh, dundred silver, recutation, ecophagists, guartic starer, guarties, guaroscophaged serlus, guaroinential benormbage, homenewage, dundred sleer, homenewage, benormbagis guartic store, insential perforation, melma, purcusation, perforand dundred store, perforand guartic store, sounded stores, perforand store, perforand dundred store, perforand dundred store, perforand dundred store, perforand dundred store, perforand store, perforance
	arhythmia, palpitation, tachycardia
Hematologic	leukopenia, purpura, thrombor ytopenia
Liver and Biliary System	ALT increased, AST increased, bilirubinentia, GGT increased, hepatitis
	Sehydration
	abnormal dreaming, amény, appetite increased, confusion, depression, servousness, sommolence
Respiratory	jashima, bronchospasm, dyspmea
	dopecia, angiordemu, bullous eruption, phonosemińskiy roaction, praritus, sweading increased, urticaria
	abnormal vision, conjunctivitis, taste perversion, timitus
Urinary System	albuniumia, BUN increased, creatinin increased, hemateria, rend failure

6.2 Personar-kring Experience

The following adverse reactions have been identified during post-approval use of melocicion. Because these reactions are producted volumely from an populationed uncertaint size, it is one slowey possible to whether the interest of the control of

7 DRUG INTERACTIONS

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

	fere with Hemostasis
	defection and autocogalisms such as sufficial horal superficial feet on Selection, The concentrate use of reduction and autocogalism horal an interessed risk of serious behavior, output of the contract of t
	Monitor parkets with concominant use of melosticam with anticognilars (e.g., wartarini, antiplateler agreess, e.g., spairini, selective seronnin receptake inhibitors (SSRRs) and seronnin nonepimplerine response inhibitors (SSRRs) for signs of bleeding (see Worrings and Processions (S.11)).
Aspirin	
Clinical Impact:	Controlled clinical studies showed that the concomitant use of NSAIDs and analgesic doses of aspirindoses more produce any greater therapeautic effect than the use of NSAIDs alone. In a clinical study, the concomitant use of an NSAID and aspirin was associated with a significantly increased incidence of GI adverse reactions as compared to use of the NSAID alone [see Worning and Procurios (5,21)].
Intervention:	Concentinate use of melanicam and love does applien or analystic does not applien is not generally recommended because of the increased risk of bleeding (see Wornings and Precentions (5.17)).
	Meloricami no a substant for two done against for cardiovascular protection.  Analysis of Bernard Melorica and Bernard Melorica and Analysis of Bernard Melorica and Bernard Melo
Clinical Impact:	NSAID, was desirable the artificiation of the control of the contr
	During concentions are of mixtic constant CLC Statistices, APSBs, or the inclination, exceeds the design feature of the control found pressure is remove that the desired found pressure is remove that the desired found pressure is removed to the control found pressure is removed to t
Diuretics	
	Tillicial studies, a word as proc- understanding conservation, because of the number of MEXID reduced the number of referred long furners (e.g., number of an interpretation of the NEXID inhibition from proximal indicates when it formatted a general reduction in number of referred long furners (e.g., number of the proximal indicates as an effectably multiple doses of architectured in the contraction of the number of the contraction of the number of the number of the contraction of the number of the num
Intervention:	During concentiants use of mobissions with disserties, observe patients for signs of worsening renal function, in addition to assuring disserted efficies (see Hierorings and Procussions (5.6)).
Lithium	
	NSAIDs here produced elevations in glassus lithium/levels and reductions in presal lithium/levels and reductions in reseal lithium/levels and reductions in research lithium/l
Intervention:	buring concentrate use of nucleacions and fidhum, monitor patients for signs of lithium moticity.
Methotrexate	
Clinical Impact:	Concentinate use of NSAIDs and methorenesse may increase the risk for methorenesse may increase
Intervention:	buring concentrate use of nelvolucian and mediceneus, monitor patients for mediceneus molicity.
Cyclos porine	
Clinical Impact:	Concentinates use of medical constraints of cyclospories may increase cyclospories as suphressicity.
Intervention:	During concentrations used on the obsticutable to the concentration of the concentration used on the concentration used on the concentration used on the concentration used to
NSAIDs and Sa	Kryburs 1
	The conconstitute use of medionicians with other NSAIDs or salley lates is neterocommended.
Pemetrexed	
Clinical Impact:	Concentions on an of melanicam and prevenence of my increase the risk of premerceard-associated mynoleogoporosists, even, and GL tracking (see the premerceard previously), information).
Intervention:	preserved, in patients with read and preserved, in patients with read impairment whose creatables character ranges from \$5 to 30 at, this, monitor for myn-loogoperssion, read and CI toxicity. Patients taking melociates should interrupt dosing for at least five days before, the day of, and not only a formation of the preserved administration of melociates and preserved. In patients with read read administration of the preserved administration of melociates and preserved and interrupt dosing for at least five days before, the day of, and not only a formation of the preserved administration of melociates and preserved. In patients with the read administration of melociates and preserved administration of melociates and preserved.

BUSE IN SPECIFIC COPILATIONS

8.1 Prepairs

8.1 Prepairs

Use of NSADs, including imbustions, during the first disease of prepairsy increases the risk of prepairs of the risk disease services. Avoid one of NSADs, including understanning in pregnant vinewastering at 25 weeks of greation (their disease) pice Vintening and Prevention (E.10). The research of the risk of prepairs and the research of the risk of the risk of prepairs and the research of the risk of

miller methods, and 15-20% for pregnancy loss.

In similar repredictions under extraction of their has onlivered in ran and radiolis travarid during the period of oppragments with methods came could have equivalent to 0.55- and 5.5-dens for mantams of the contraction of the cont

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

Animal Data 
Motoricam was not transported when administeneed to pregnant zon during fixed organogenesis at road 
dones up to a singlicities (2.6-fold goare than the MRIII of 1.8 get of and six cambood on BRA. 
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BLA Lemma 

There are no human data available on whether substicating power in human milk, or onthe effects on the Maximum 

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2. Floral Impairment

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No four a againment in secretary in pairms, with sald to medicate read impairment have reless mades. The time of industricant in studies; with severe read impairment have reless mades and impairment have read to impairment in the recommended. In pairms on hemotodaylor, in relocate inhelidad not exceed 7.5 mg per day.

Melroscom is not dissipatable (per Dossage and Administration (2.1) and Clinical Pharmacology (2.3)).

HOVERDOACE

Sympose filability age to NSAID everdinage have been spicially listed to be leftings, deswisses, mane, sweeting, and to greater join, which have been goverably reversible with supportive care. A second process of the pr

II DESCRIPTION
Meloxicanis a nonseroidal anti-inflammany drug (NSAID). Each yellow neloxican table contain
Sen que 15 mg or 15 mg meloxicanis (neloxicanis) dentecidades (Aspara de a 4-bydroxy
2-angly), 4-6-angly), 2-6-langly), 2-16-2-bennolatanis -2-anbountés 1-1-dentes (Aspara de a 4-bydroxy
angly), 4-6-angly), 2-6-langly), 2-16-2-bennolatanis -2-anbountés 1-1-dentes (Aspara de a 4-bydroxy
angly), 4-6-angly), 4-6-angly), 4-6-angly and 4-bydroxy
angly a 5-31-4 is a regional formation 4-cytophysiological that the following streams and formation
and the following streams and formation 4-bydroxy
and 4-bydroxy
angly a 5-16-4 is a regional formation 4-cytophysiology and the following streams and formation
and the following streams and formation 4-bydroxy
and 4-bydrox



Meloxicam, USP is a pale yellow powder, practically insoluble in water, slightly soluble in acetone, soluble in dimenylformamide, very slightly soluble in ethanol (96 %) and in methanol. Meloxicam has an apparent partition coefficient (log  $P_{log}p = 0.1$  in n-octanolbuffer pH 7.4. Meloxicam has pKa values of 1.1 and 4.2.

of L1 and 4.2. Each meloxicam tablet, USP intended for oral administration comains 7.5 mg or 15 mg of meloxicam, le addition, each tablet comains the following inactive ingredients: colloidal silicon dioxide, crospovidone, lacrose monthydraw, magnesium stearate, microcrystalline celluloise, povidone and sodium citaxe dibydrate.

## 12 CLINICAL PHARMACOLOGY

IZ CLINICAL PHARMACOLOGY

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21. Pharmacocknets.

Absorption

22. Pharmacocknets.

Absorption

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26. Pharmacocknets.

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

Pharmacokinetic Parameters (% CV)	Steady State			Single Dose	
	Healthy male adult (Fed) <sup>2</sup>	Elderly males (Fed)	Elderly females (Fed) <sup>2</sup>	Renal failure (Fasted)	Hepatic insufficient (Fasted)
	7.5 mg <sup>3</sup> tablets	15 mg capsules	15 mg capsules		
N	18	5	8	12	12
Cmax [µg/mL]	1.05 (20)	2.3 (59)	3.2 (24)	0.59 (36)	0.84 (29)
t <sub>max</sub> [h]	4.9 (8)	5 (12)	6 (27)	4 (65)	10 (87)
tio [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-441L1	14.7(32)	15 (42)	10 (30)	26 (44)	14 (29)
CL/I (mL/min) V <sub>2</sub> /I <sup>4</sup> [L]	8.8 (29) 14.7(32) is in the table are from reditions	9.9 (76) 15 (42)	5.1 (22)	19 (43)	11 (44)

## Food and Antacid Effect

Medicinary and an experimental of the property of the property

In a covarian analysis, utilizing population pharmacokinetics body-weight, but not age, was the single predictive covariate for differences in the meloticism apparent oral plasma clearance. The body-weight normalized apparent oral clearance values were adequate predictors of meloxicism exposure in pediatric parterns.



Voting femilies exhibited (slight) tower plasms concurrenced most ratio to young radio. Are sight tower 0.7.5 mg molecules the most editorishment lift: lever 19.5 mours for the femilie group at compared to 2.4 shows for the make group. At usuady stars, the data were similar (17.5 hours vo. 21.4 There was linearly of pharmscolandics and no appreciable difference in the Crass or Trace across species.

togramient in inferiorimental part consequent Ammunitation (2.7), whereasy and reconstitution (5.3) as the Memoritalysis. Hemocralisation of the Memorital Part of the Memorita

Canscidine

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

Disjoini Digarin

Melosican 15 mg once daily for 7 days, did not alter the plasma concentration profile of digostin after β-acryddigostin aftrinistration for 7 days ar clinical doses.

In vitro resting found no proxim binding drug interaction between digostin and melosican.

Lithium in a study conducted in hashthy subjects, mean pre-dose lithium concentration and AUC were increased by 21% is subjects receiving lithium doses ranging from 804 to 1072 mg noice daily with meloxicam 15 mg QD every day as compared to subjects receiving lithium alone [see Drug harroctions (7)]. Methotocous

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The effect of indiscission of the autocopilates effect of workins was stoffed in a group of healthy

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13 NONCLINICAL TOXICOLOGY

II NONCLISECAL I UNICCULIED.

II Carcinegensis's Managements, Impairment of Fertility
Garcinegensis

There was no increase in tumor incidence in long-term carcinegenicity studies in cast (164 weeks) and
management in the case of the c

Mutagenesis

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

Impairment of Fertility

Meloxicand did not impair male and female ferfility 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-dimes greater, respectively, than the MRHD based on BSA commarison.

H. CLINICAL STORMS
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The use of individual methods and the sign and symptom of conventration of the law and hybrid and the sign of the law and hybrid methods and the sign of the law and hybrid methods and the sign of the law and hybrid methods and the sign of the law and hybrid methods and a

with plactics, No incremental hostifiction was observed with the 225 and thost comprised to the 15 mg date.

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HOW SUPPLIED
Product 68151-4319
NDC: 68151-4319-0 1 TABLET in a PACKAGE

17 PATIENT COUNSELING INFORMATION

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

Advise the patient to read the TDL-approved prient labeling (Medication Guide) that accompanies such prescription disponant.

Minimpatient, Institution of their caregivers of the following information before initiating therapy with interruptions, Institution of the opinion feetings of the contraction of the Contract

(5.3).

Mart Faltur and Edma

Advise patients to be alter for the symptoms of congestive heart falture including shortness of breach, unexplained voiging sign or orderns and to contact their healthcare provider if such symptoms occur [see Worrings and Precondition (5.3).

Amplytactic Revictions

Anaphylacid Reactions

Helmapatien no for in signs of an anaphylacide reaction (e.g., difficulty breading, welling of the face or french, hauser patients in seek immediate energiency help if these occur [nor convolutioning (2)].

Servina Sila Bractions

Servina Sila Bractions

Acide patients in storparolocioni (2)/3.

Acide patients in storparolocioni immedianely if they develop any type of rash and in contact their beaffice are provider as soon as possible jue binnings and Percantion (5.5)).

Franké Fertilly

Frauls Freilly
Africe females of propodactive potential who desire preguncy that NSAIDs, its fading melosicam, may be assected with a reversible deep in outdaten just the 3-pospele populations (3/3).
Frail Teakby
Hefrim pregnaturement no words use of multivational deep NSAIDs, starting a 30 weeks genetion because of the risks of the presumer closing of the freal dectus arterious (see Niembygs and Precentions (3/4)) and the na jusqu'ell populations (3/4).

(6.10) and the in-people Populations (E.P.).

Associ Cancermanian of the SNADD.

Internations that the concentrate on the individual with sides (NSADD) or subsylvan (i.g., definitional, and the state of the concentration of the individual with sides (NSADD) or subsylvan (i.g., definitional, state of the individual sides (i.g., definitional, state of the individual sides), and tide or subsylvant (i.g., definitional, state of the individual sides), and tide of the individual sides (i.g., definitional sides), and the individual sides (i.g., definitional sides), and the individual sides (i.g., definitional sides). The individual sides (i.g., definitional sides) is the individual sides (i.g., definitional sides). The individual sides (i.g., definitional sides) in the individual sides (i.g., definitional sides). The individual sides (i.g., definitional sides) in the individual sides (i.g., definitional sides). T

o with increasing doses of NSAIDs o with longer use of NSAIDs Do not take NSAIDs right before or after a heart surgery called a "coronary artery

Do not user NSALDS rigin better or airer a treat surgery cause a - ceremany artery bypass graft (CABG)."

Avoid taking NSAIDs after a recent heart attack, unless your healthcare provider tells you to. You may have an increased risk of another heart attack if you take NSAIDs after a recent heart attack. antark.

Increased risk of bleeding, ukers, and tears (perforation) of the esophagus (tube leading from the mouth to the stomach), stomach and intestines:

NSAIDs

a stating and effective called "reinforcementals", "nationagelanes", "SSBIs", or "SNBIs"

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What are SNAIDs.

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What are SNAIDs.

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If you have been an authors seek, News, or other allergic reaction with aspirate are up other NAIDs, regished these are described powers and the power and an authorized power and a conditions.

Before taking NAIDs, the Byear he habitance provider above all of your media of conditions.

In which were take the power and the power and the power and a conditions are all the power and power and the power and power and the power and the

life-threatening allergic reactions
 Other side effects of NSAIDs include: stomach pain, constipation, diarrhea, gas,
 hearthurn, nausea, vomiting, and dizziness.

Bit de de comming this reactions
 Other side effects of SNAMs in subsets entend pain, conséguion, diarrhes, gas,
 Interfaces, mouses, veniting, and diarries.
 Cent energence hydre place way if you get any of the following symptoms:
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 Colon signifique de face or thread
 veniting of the face of the face

Meloxicam 7.5 mg tabs



Product Informa	tion					
Product Type		HUMAN PRESCRIPTION DRUG	hem Code (Sourc	e) NDCS815	-4319(NE	C68382-050
Ruste of Administra	ation	ORAL				
Active Ingredien	nt/Active M	loiety				
		neredient Name		Basis of Str	rorth	Strength
MELOXICAM (UNII V	VG2QFR3CGL	) (MELO XICAM - UNILVGJQFR ) CGI	) [6	ELOXICAM		7.5 mg
Inactive Ingredie	ents					
		Ingredient Name				itrength
CROSPOVIDONE (UP						
LACTO SE MONOSTE MAGNESIUM STEAR						
SILICON DIDXIDE (L					-	
		3614) (E/UNII 822547895K)			-	
POVIDONES (UNIL F)						
		INE (UNIL OPTRIZOS (U)				
	eristics					
Color	VELLOW		Score		80 KC	130
Color Shape			Size		6 mm	100
Color Shape Flavor Contains	VELLOW					139
Color Shape Flaver Coutsins Packaging	VELLOW	OUNDS	Size Imprint Code		6 mm 2C/25	
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