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

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FULL PRESCRIBING INFORMATION: CONTENTS*
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1.2 Deacounter (SIGN)
1.2 Deacounter (Martin (Binne) (Binne)
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12 CLINEAL PREMINENCE.

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WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS

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recautions (5.1). I Carterion, and Perforation and Americanian (Satisfaction testal Balacian), Useration, and Perforation | * SSADIO Cause an increase of risk of serious gestrobesthal (GI) adverse events including blaeflay, Micration, and perforation of the stomach or intestitions, which can be little Times events can occur at a stomach or intestitions, which can be little Times events can occur at and patients with a prior history of peptit user disease moder GI bleeding are at greater risk for serious GI events [see Warnings and Precautions (5.4).

1 INDICATIONS AND USAGE

1. Institute Institute See Use (Institute Institute Ins

arthritis (see Linical Studies (14-1)).

1.3 Jovenile Rheumatold Arthritis (IRA) Pauciarticular and Polyarticular Cours
Meloxician tables are indicated for rolef of the signs and symptoms of pauciarticular or
polyarticular course (overein Rhaumatold Arthritis in patients who weigh selfo kg [see
Dosage and Arthritishatton (24) and Chicki Studies (14-2)].

2. Observal body partnersions.

Carofully, consider the potential benefits and risks of melouscam tablets and other treatment option. Both of the control to the potential benefits and risks of melouscam tablets. Use the lowest effective decapy to use melouscam tablets. Use the lowest effective decapy for the shortest duration consistent with individual patient treatment qualifies warmings and Processions (5). After observing their response to initial threapy with melouscam tablets, adjust the dose to said a mindful application heads.

In adults, the maximum recommended daily oral dose of meltoxicam tablets are 15 mg regardless of formulation. In platients with hemodalysis, a maximum daily dosage of 1.5 mg is recommended (see Use in Specific Populations (2.7 and Ginkal Pharmacology (2.2.9) [2.2.9) [

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3 DOSAGE FORMS AND STRENGTHS

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A CONTRANDICATIONS

Moleculam is contraindicated in the following patients:

• Known hypersensibility (a.g., anaphylicitic reactions and serious skin reactions) to malocition or any components of the during product [see Warnings and Precautions (

• History of asthma, uniticaria, or other allergic-type reactions after taking aspirin or

other NSAIDs. Severe, sometimes fatal, anaphylactic reactions to NSAIDs have beseroprated in such patients [see Warnings and Precautions (5.7, 5.8)]

In the setting of coronary artery bypass graft (CABG) surgery [see Warnings and Precautions (5.1)]

3 MANIBORS AND PRECIATIONS

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Post-M Patients

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5.3 Analysicals: Reactions

Miniscen has been some classified with pully-licit reactions in patients with and offlood Miniscen has been some classified with many lateral content and content of the discretization of an analysical creation content of the content

5.9 Serious Skin Reactions

NSAIDs. Reclaim protections

NSAIDs. Reclaim protections causes serious skin adverse reactions such as exclusive dermatiks, Stevens-Indexon Syndrome ISSI, and toxic spidermal necropies (IRIN), which can be falls These serious cents may occur without warring, inform patients about the sign said symptome of serious skin reactions, and to describtion the patients about the sign said symptome of serious skin reactions, and to describtion the serious skin reaction and the serious skin reaction and the serious skin reactions in SAIDs (IRI accordinated cated in patients with previous serious skin reactions in SAIDs (IRI accordinated cated).

A production of the contraction of the contraction

hemoglobin or hematocrit. — oy sayan or symptoms of ainenis, monitor in KSADs, including matoxicam, may increase the risk of biseding wents. Co-morbid conditions such a coagulation disorders or concomitant used wherein, other airticoagulation, arripitative agents (e.g., aspirin), serotomin reupstain einhibers (SSRIs) and assistant in nergolespirine reupstain einhibers (SSRIs) and assistant in nergolespirine reupstain einholds (SSRIs) may remease the risk. Monitor these palatiest for signs of biseding [see Drug Interactions (7)]. S. 2. Masking of Hollans-estimates.

5.12 Masking of Inflammation and Fever
The pharmacological activity of molecicam in reducing inflammation, and possibly fever may diminish the utility of diagnostic signs in detecting infections.

6 ADVISES REACTIONS
The following abrears reactions are discussed in greater detail in other sections of the discussion of a dis

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

Adults Osteoarthritis and Rheumatoid Arthritis

Observations and Philamental Annies.

The medicaces Than 20 of cited updid statutes includes 15.127 OA patients and 1021 Ma patients tracked with medicace 7.5 mg/sty. 250 OA patients and 1511 Ma patients patient for the state of the medicaces 7.5 mg/sty. 250 OA patients and 1511 Ma patients patients for a state of them 15.12 Ma patients of the state of them 15.12 Ma patients patients for a state of them 15.12 Ma patients of the state of the st

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

Table 1b depicts adverse events that occurred in x2% of the meloxicam treatment groups in two 12-week placebo-controlled resumatoid arthritis trials.

Table 1a Adverse Events (%) Occurring in ≥ 2% of Meloxicam Patients in a 12-Week Osteoarthritis Placebo- and Active-Controlled Trial

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 *	2.5	1.9	4.5	3.3
Fall	0.6	2.6	0.0	1.3
Influenza- like symptoms	5.1	4.5	5.8	2.6
Central and Peripheral Nervous System				
Dizziness				
	3.2	2.6	3.8	2.0
Headache	10.2	7.8	8.3	5.9
Respiratory				
Pharyngitis	1.3	0.6	3.2	1.3
Upper Respiratory Tract Infection	1.9	3.2	1.9	3.3
Skin				
Rash ²	2.5	2.6	0.6	2.0

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

	Placebo	Meloxican	
		7.5 mg daily	15 mg daily
No. of Patients	469	481	477
Gastrointestinal Disorders	14.1	18.9	16.8
Abdominal pain NOS 2	0.6	2.9	2.3
Dyspeptic signs and symptoms 1	3.8	5.8	4.0
Nausea 2	2.6	3.3	3.8
General Disorders and Administration Site C	onditions	•	•
Influenza-like illness ²	2.1	2.9	2.3
Infection and Infestations	•	•	•
Upper respiratory tract infections-	4.1	7.0	6.5
pathogen class unspecified 1 Musculoskeletal and Connective Tissue			1
	Disorders		
joint related signs and symptoms 2	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
*MedDRA high level term (preferred terms): dyspeptic si	ons and symptoms (d		pepsia

*Jacofd Ahrby I woul term (previewed terms): dyspeptic signs and symptoms (dyspepsia, dyspepsia aggressedes, dructation, quaterinshished inflatation, lugar resistants/rate inflations-aptivages unraper-filed (bayogita MOS, phayogita MOS, insulis MOS), joint related signs and symptoms (activation activation) and activation of the control point relations, injust resistants (past regulations), joint relations, injust resistants (past regulations) point relations, injust relations (past resistants) and most of the control term neurosa, abdominal pain MOS, influenza-like Biness, headaches MOS, and math MOS.

	4 to 6 Weeks Controlled Trials		Month Controlled Tria	
	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 *	0.6	2.0	2.4	1.6
Pain	0.9	2.0	3.6	5.2
Central and Peripheral Nervous System Dizziness Headache Hematologic	1.1 2.4	1.6 2.7	2.4 3.6	2.6 2.6
Anemia	0.1	0.0	4.1	2.9
Musculoskeletal Arthraigia Back pain	0.5 0.5	0.0	5.3	1.3
Psychiatric		0.4	3.0	0.7
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 [†]	0.3	1.2	3.0	1.3
Urinary Mcturition frequency	0.1	0.4	2.4	1.3
Urinary tract infection	0.3	0.4	4.7	6.9

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

measure care and moving approxim	
	Margic reaction, face edema, fatque, fever, hot flushas, malaise, syncope, weight docrease, weight increase
	ingna pectoris, cardiac falure, hyportension, hypotension, myocardial infarction, vascutilis
Central and Peripheral Nervous System	
	Colos, by mouth, doctoral user, excitation, ecophagia, guiter user, peortes, gastroecophagia refux, gastroecophagi
Heart Rate and Rhythm	arrhythmia, palpitation, tachycardia
Hematologic	kukopenia, purpura, thrombocytopenia
Liver and Bilary System	ALT increased, AST increased, bilirubinemia, GGT increased, hepatitis
	phydration Services S
	abnormal dreaming , anxistry, appetite increased, confusion, depression, nervous ress; somnolence
	asthma, bronchospasm, dyspnea
Skin and Appendages	lisiopicia, angisedema, bullous eruption, photosiensibivity reaction, pruntus, sweating increased, urticaria
	Abnormal vision, conjunctivitis, taste perversion, tinnitus
Urinary System	albuminuria, BUN increased, creatinine increased, hematuria, renal failure

7 DRUG INTERACTIONS
See Table 3 for clinically significant drug interactions with meloxicam. See also Warnings and Precautions (\$.2, \$.6, \$.11) and Clinical Pharmacology (12.3) .

	Table 3 Clinically Significant Drug Interactions with Meloxicam
	iterfere with Hamostasis
Clinical Impact:	
	Section in relative by glidates pay, an important rich in himmottasis. Case-central and orbit or glidates pays an important rich in himmottasis. Case-central and orbit or glidates pays an important rich in himmottasis. Case-central and orbit or glidates pays an important rich in himmottasis. Case-central and orbit or page and an ISAUD allows. Market or pages and an important rich in himmottasis. Case-central and orbit or pages and an ISAUD allows. Market or pages and an important rich in himmottasis. Case-central and orbit orb
Intervention: Aspirin	Monitor palaints with concomitant use of melavicam with articoagulants (e.g., warfarin), antiplatekit agents (e.g., asprin), selective serotonin resuptake inhibitors (SSRIs), and serotonin nonspinephrine resuptake inhibitors (SNRIs) for signs of bleeding [see Warnings and Procautions (S.11)].
Clinical Impact: Intervention:	Sometivation formical studies showed that the concommand used of HSADDs and analysis of Security of Se
intervention:	Autocommunic was or menuticem and us to one adjust or an expectation activation to adjust or an extraordistrial to the contraction of the contract
ACE Inhibitor	PRINCE AND THE ADMINISTRATION OF THE ADMINIS
Cloical Impact	S. regionaries messages in the contents of the
	In patients who are elderly, volume-deplated (including those on disretic therapy), or have renal impairment, co-administration of an NSAID with ACE inhibitors or ARBs may result in deterioration of renal function, including possible acute renal failure. These effects are usually reversible.
Intervention:	During concomitant use of meloxicam and ACE inhibitors, Affilias, or betta-blockers, monitor blood pressure to ensure that the desired blood pressure is obtained.
	During concentrate use of medicious man ACE inhibitors or Afficis in patients who are adeletely, volume-depletel, or have impaired medicine, member for signs of encounting read Procustions (S. 60). Where three drivers are administrated concentrately. Journals volume depletely, or have impaired and medicine and function, member for signs of encounting read Procustions (S. 60). Where three drivers are desirable for the administrated concentrately. Journals volume depletely, outsines depletely and depletely and procustions (S. 60).
Diuretics	when these drugs are administered controllationly, patients stroom or abequately information at the degrining or the controllation at the degrining or the controllation at the degrining or the controllation at the degrining of
Clinical Impact:	Clinical studies, as well as post-
	marketing observations, showed that NSAIDs reduced the naturative defect of loop duretics (e.g., furosemide) agents and multiple dose pharmacodynamics and pharmacodynamics and pharmacodynamics and pharmacodynamics are not affected by multiple doses of meloxicam.
	During concomitant use of melanicism with distratics, observe patients for signs of worsening renal function, in addition to assuring distrated fracting metallypartnessive effects [see Warnings and Procautions (5.6)]
Lithium	
	ISSAIDs have produced elevations in plasma lithium levels and reductions in renal lithium clauracce. The mean minimum lithium concentration increased 51%, and the renal claurance decreased by approximately 20%. This effect has been attributed to RSAID inhibition of renal prostagalardin synthesis [see Circial Pharmacology (12.3)].
	During concernitant use of melonickam and Bilbum, member publishts for signs of Bilbum toxicity.
Methotrexat	
Clinical Impact:	Concentrant use of NSAIDs and multi-batevaste may increase the risk for multi-batevaste toxicity (e.g., neutropenia, thrombocytopenia, cand dysfunction).
Intervention:	During concendrant use of melexicam and methothexate, montor patients, method reside toxicky.
Cyclosporine	
Clinical Impact:	Concemitant use of melucicam and cyclosporine may increase cyclosporine may increase cyclosporine in a process of the cyclosporine may increase cyclosporine in a process of the cyclosporine may increase cyclosporine may increa
Intervention:	During concomitant use of meleoxicam and cyclosporine, member patients for signs of worsening renal function.
NSAIDs and 5	alicylates
Clinical Impact:	Concomitant use of medicicism with other MSAIDs or sale/plates (e.g., offlurins), sale/plates
Intervention:	The concomitant use of melanicam with other INSAIDs or salecylates is not recommended.
Pemetrexed	
Clinical Impact:	Concentrate use of meterican and pernetrosced may increase the risk of pernetrosced-associated myeloscopproscion, renal, and Gi toxicity (see the pernetrosced procuration).
Intervention:	During concentlations of melecizian and permeteread, in patients with renal impairment without creativine clearance point 95 to 79 millimin, the concentration and Contextby, Publishers taking melecizian should interrupt doing for at least the days following permeteread administration. In patients with residence indicates the content of the concentration and content of the production of the content of the con

8 USE IN SPECIFIC POPULATIONS

B USE IN SPECIAL POPULATIONS

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E.3. Prepaired.

E.4. Prepaired.

E.5. Prepa

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 stilbirth.

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800-222-1222).

DESCRIPTION

**DESCR



Melanciam, USP is a pale yellow promoter, practically inscalable in water, stightly solable in extense, solable in dismetry/formandes, very stightly solable in ethanol (65 %) and in melanuol. Molecular has an apparent partition colliferation (69 %) gen — 0.1 in no-citamytiseffler pit 7.4. Melanciam has plas values of 1.1 and 4.2. Each melanciam tables (USP intended for or administrations constain 7.5 mg or 15 mg Each melanciam tables (USP intended for or administrations constain 7.5 mg or 15 mg each melanciam tables, proteins and solables (included intended intend

12 CLINICAL PHARMACOLOOY

12.1 Mechanism of Artican

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Table 45ingle Dose and Steady-State Pharmacokinetic Parameters for Oral 7.5 mg and 15 mg I (Mean and % CV) ¹

		(Mean and	76 CV) -		
Pharmacokinetic Parameters (% CV)				Single Dose	
	(Fed) ²	stiderly males (Fed)	(Fed) ²	(Fasted)	Hepatic insufficiency (Fasted)
	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 16 [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 _/f 4[L]	14.7(32)	15 (42)	10 (30)	26 (44)	14 (29)
² The parameter value	is in the table are from vi	rious studies			

*The parameter values in to 2 not under high flat condition 2 Meloxic am tablets 4 V y/f mDose/(AUC+K e/)

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Size
Voting females exhibited slightly lower plasma concentrations relative to young males.
After single doses of 7.5 mg melanicam, the mean elemination half-file was 19.5 hours.
After single doses of 7.5 mg melanicam, the mean elemination half-file was 19.5 hours are the data wave similar LTP folium's 2.1, 5 hours.) This planned collected for the operation of the data wave similar LTP folium's 2.1,5 hours.) This planned collected for dispressions of the collected for the product of the collected for the collected for the product of the collected for the product of the collected for the collected fo and no appreciable ofference in the Cmax or Timax across genders. Hepatiz Impairment February a single 15 mg dose of midd (xikla-hugh Class I) om moderate (Child-hugh Class I) om renderate (Child-hugh Class I) om renderate (Child-hugh Class I) om renderate (Child-hugh Class I) om the control of the child-hugh Class II of the child-hugh Class II of the child I of

(Child-Pugh Class III) have not been adequately studied [see Warnings and Precautions (5.3) and Use in Specific Populations (8.6)].

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February 2 (1997)

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protestioning 595. This resoluted in a discression in 1₅₀ from 192 hours to 12.5 mours and a 35% reduction in AUC. This supposts the existence of a reducidation pathway for
missiscam in the gastroinestrial struct. The clinical relevance of this interaction has not
been established. Christians

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

Digionin

Maloricam 15 mg once dally for 7 days did not alter the plasma concentration profile of digions that P-acetylrigosin administration for 7 days at clinical doses. In white testing found no protein binding drug interaction between digions and melos ican.

Libitim

Libbum in a study conducted in healthy subjects, mean prie dose thinms concentration and ALC in a study conducted with healthy subjects, mean prie dose thinms concentration and ALC interest and the second control of the

binding size from Printeractions (7):

Warfarin
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13 NONCLINICAL TOXICOLOGY

13.1. Cartinogenesis, Mutagenesis, Impairment of Fertility

Cartinogenesis in International Programment of Fertility

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14.2 Juvenile Rheumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course

14.2 journals Benemental Arthritis (IAA) Pacciarticular and Polyerticular (IAA) procession of the Polyerticu

16 HOW SUPPLIED/TORACE AND HANDLING
Maisstarm Tables USF). 15 mg are yellow, round-shaped, fife breviald edge, uncotated
face declosed with 2 2-2 and 5 m on one side and plan on other side and are supplied as
NCC 6601-1303-2 80/THLS OF 3
NCC 6601-1303-2 80/THLS OF 15
NCC 66071-303-3 80/THLS OF 50
NCC 66071-3 80/THLS OF 50
NCC 66071-3 80/THLS OF 50
NCC 66071-3 80/T

Storage
Store at 20' to 25' C (66' to 77' F) [see USP Controlled Room Temperature]. Keep melaction tablets in a dry place.
Disperse tablets in a tigit container.
Keep this and all medications out of the reach of children.

17 PATIENT COUNSELING INFORMATION

17 PATION COURSEAND INFORMATION
Adults the planet for their off Designation granted bashing (Medicator Gasies) that
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Hepatedoxickly
Inform patients of the warring signs and symptoms of hepatedoxickly (e.g., naucea,
Indigos, Minago, durinks, printins, junidos, right upper quadrant tenderness, and
fully symptoms, If these outer, inviting patients to be to immission and seak inventibles

Heart Patient and Edmand
Annie patients to be and for the symptoms of competition have failure including
shortests of breath, unequilated easily gain, or edoms and to contact their healthcare
provider float symptoms count for sell symptoms and Princations (5.3).

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Interference section of the second providers of the contract of the c

Inform pregnant women to avoid use of meloxicam and other NSAIDs starting at 30 weeks gestation because of the risk of the premature closing of the fetal ductus arteriosus [see Warnings and Precautions (5.10) and Use in Specific Populations (8.1)

anteriors (as are training and Prevailables (5.3) and the in Figure Republicing (8.1). And Cancembatt Use of HSMDs. Under particular that the concentration and restriction with other MSADs or subsplies (see, divented, Madella on introducement and see to the concealed rise of the concealed rise of the concealed rise of prevailables (5.3) and the particular (5.3) and the particular that particular that MSADs may be prevailables (5.3) and the particular (5.3) and the particular of cells, New 4 restricts the FMSADs and PSADs of the MSADs and PSADs of the MSADs and the subsplit of the MSADs and PSADs of the MSADs and PSADs of the MSADs and the subsplit of the MSADs and Concealed the MSADs of the MSADs and the subsplit of the MSADs and the MSADs and provided and subsplit of the MSADs and the MSADs and provided and subsplit of the MSADs and the MSADs and the MSADs and subsplit of the MSADs and the MSADs and the MSADs and subsplit of the MSADs and the MSADs and subsplit of the MSADs and the MSADs and subsplit of the MSADs

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NSAIDs can cause serious side effects, including:

• Increased risk of a heart attack or stroke that can lead to death. This risk may be appear park in treatment and may increase.

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on with increasing deep of PSASDS.

on with togety used MSASDS.

Do not take MSASDS register to rafter a heart surgery called a "coronary
bryans upon't CASDS."

Avoid Laking MSASD offer a recent heart attack, unless your healthcare
provider less you to. You may have an increased risk of another heart attack,
you take MSASD with a winter a recent heart attack of another heart attack
you take MSASD with a recent heart attack of another heart attack. If you take NSAIDs after a recent heart attack.

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

a mytime during use

o without warning symptoms

that may expect the stomach of the stomach of

o that may cause owner.

The risk of getting an ulcer or bleeding increases with:

o past history of stomach ulcers, or stomach or intestinal bleeding with use of

a billing membrane utilité l'outremander, "introcagalistes", "SSRe", er "SRels," er "SRels," er "SRels," er "SRels," er "SRels," er sommer de l'outremander de l'outre de l'outr

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PACKAGE LABEL.PRINCIPAL DISPLAY PANEL



me	ELOXICAN iloxicam table							
P	roduct Info	mation						
P	oduct Type		HUMBS PRESCRIPTION DRUG	ten		MDC-68 051)	171-3130	NDC 6838
R	oute of Admin	istration	ONAL					
Aı	tive Ingred	ient/Activ	e Molety					
		loor	edient Name		D:	sis of Str	ength	Streng
м	ILOXICAM (UNI	VSZQPRICSI) (MILOSECAM - LINELVS 3 QF	R3CQF)	MILI	SKICAM		15 mg
In	active Ingn	edients						
			Ingredient Name				- 1	trength
	CTOSE MONOR							
	MONESHUM STE.							
	JCON DIOXIDE							
			78 (UNII: 822547895K)					
	VIDONE (LAS. I		INE UNI OPSKIZDIGUI					
CF	OSPOVIDONE	UNI 25.78306	(540)					
Pi	ospovoose roduct Char	acteristics	(ARTTOM)	Score			no so	úre.
Pi Co	ospovoose roduct Char dor sape	acteristics	(342)	Size			Brun	
Pi Co Sil	ospovoose roduct Char	acteristics	(ARTTOM)	Size	rt Code			
Pi Co	roduct Char eler sape swor setales	acteristics	(ARTTOM)	Size	rt Code		Brun	
Pi Co	roduct Char dor uspe wor estains	acteristic yelow NOUNE	(MILLOW)	Size			2C/20	
Pi Co	roduct Char eler sape swor setales	acteristic yelow NOUNE	(ARTTOM)	Size	rt Code Marketin Dat		ZC 20	
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