## MELOXICAM- meloxicam tablet St. Mary's Medical Park Pharmacy

HIGHLIGHTS OF PRESCRIBING INFORMATION These highlights do not include all the information needed to use Meloxicam Tablets USP. usfely and effective), see full prescribing information for Meloxicam Tablets USP.

## Tablets USP, for oral use

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 Mark 12, Append 201
 Mark 12, Append

RECENT MAJOR CHANGES Beard Warning 5/2026 Indicators and Usage, Juwelle Pheumatoli Arthrite (IRA) Pauciaticula (2020 Dosage and Administration, General Dosaing Instructions ( 21) 6/2026 Dosage and Administration, Journile Theumatoli Arthrite (IRA) Pauciaticul Dosage and Administration, Journile Theumatoli Arthrite (IRA) Pauciaticul Cosage and Administration, Journile Theumatoli Arthrite (IRA) Pauciaticul Dousge and Administration, Jovenie Rheumatolin (? 21) (2016 24) 60206 Wainings and Precaritions, Gardessance Thomholic Genetic (S1) 50206 Wainings and Precaritions, Heart Fallure and Elema (53) 50206 Wainings and Precaritions, Heart Fallure and Elema (53) 50206

INDICATIONS AND USAGE INDICATIONS AND USAGE INDICATIONS AND USAGE Obtoosthist (OA) (12) Rhournatold Arthritis (OA) (12) Rhournatold Arthritis (OA) (12)

Losses effective dosage for the shortest who weigh adding (1.2)
 DoCAGE AND ADMINISTRATION
 Use the lowest effective dosage for the shortest duration consistent with individual patient treatment goals
 (2.1)
 OA(2.2) and RA(2.3):

Starting dose: 7.5 mg once daily Dose may be increased to 15 mg once daily • JPA (2.4):

Known hypenenstölvity to meliautam or any components of the drug product ( 4)
 Hibby yil anthras, urtitani, or other allengi-type neutrinos after taking auplinis or other NSADs ( 4)
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• <u>District</u> (KKC) and related an antityte Manual Man

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 1.2 Resemands Arthritis (PA)
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8.1 Pregnancy 8.2 Lactation 8.3 Females and Nales of Reproducti 8.4 Pediatric Use 8.5 Genatric Use 8.6 Hepatic Impairment 8.7 Renal Impairment

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FULL PRESCRIBING INFORMATION

WARNING: RISK OF SERI EVENTS OUS CAI SCULAR AND GASTROINTESTINA

Cardiovascular Thrombotic Events
 Nonsteroidal anti-inflammatory dru

Nonstervidal anti-inflammatory drugs (NSADD) ( suus an increased nich of enroux cardiovarcubur tomontotic evonts, including myocardial treatment and may increase with duration of use [ see Warnings and Precasations ( 3.1.).
 Constraint ( 1.1.).
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Warnings and revisions (3.1). The second sec

1 INDICATIONS AND USAGE

I INDICATIONS AND USAGE
I.1 Osteoarthritis (OA)
Molexian tablest are indicated for relief of the signs and symptoms of osteoarthritis [
see Clinical Studies (14.1)].

see Check Studied (141)] 1.2 Resumption further (Ref) Maiorizant Itabies: are indicated for reflet of the signs and symptoms of rheumatoid mitrifs (are chicked Station (14.1)) 1.3 Jownah Rheumatoid Arthrifts (RBA Placuktricitude and Polyariticular Conser-point Robust Constant Station (14.1)) Datages and Annowalism (12.1) and Check Station (14.2).

2 DOSAGE AND ADMINISTRATION 2.1 General Dosing Instructions

c.1 General Dosing Instructions Carefully consider the potential benefits and risks of Meloscam tablets and other trainance option balls descripting target meloscam tablets. Use the levelst effective Warnings and Processorties (1) and the second second patient trainance galaxies Warnings and Processorties (1) and the range with Meloscam tablets, adjust the dose to so at an individual patient's media.

suz an induktual pazient's needs. In aduts, the maximum recommended daily oral dose of Mekoscam tablets is 15 mg regardles of formulation, in patients with hemodulysis, a maximum daily dosage of 7, 12 23). Meloscam tablets may be taken without regard to timing of meak.

2.2.Osteaarthritis
For the relief of the signs and symptoms of osteaarthritis the recommended starting and maintenance ceal dose of Makoucam tablets is 7.5 mg one day. Some patients may receive additional benefit by increasing the dose to 15 mg once addy.

They revenue and a structure of the terms of the outer of 2 in the outer.
2.3 Rheumatoid Arthritis
For the relief of the signs and symptoms of rheumatoid arthritis, the recommended starting and maintenance and does of Melosizam tables is 7.5 mg once daily. Some additional breaks additional barrier by or receivery the does to 15 mg once daily.

Jackies and Jackies and Anthritis (BA). Parciacitational and Polyaritodas. 2.4 Jovenin Rohematold Anthritis (BA). Parciacitational and Polyaritodas. Ter the treatment of jovenin Hommatold attricits, the recommended and add so and Rohecamatolatics (2.5.7 mg once daily), in rohemator model and big. There was no additional boniff demonstrated by increasing the data above 7.5 mg in clinical trails. Network and the trained by the second and the second and the second and the second and the Rohecamatolatics in bolic and the second and the second and the second and the second and the Rohecamatolatics in bolic and the second and the second

In patients on hemodialysis, the maximum dosage of Meloxicam tablets is 7.5 mg per day [ see Clinical Pharmacology ( 12.3) ].

ung tawe Cincar Manmacology (12.3) }.
2.6 Non-Interchangeability with Other Formulations of Meloxicam Molecum tables have not them equivalent systemic exposure to other approved formations of or al meloxicam. Therefore, Meloxicam tables are not interchangeable with other formations of or all meloxicam products wort if the tables are not interchangeable to end other and the system of the system of the system of the system to the system of the system of the system of the system of the system to the system of the system of the system of the system of the system to the system of the system of the system of the system of the system to the system of the system o

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2.5 Renal impairment The use of Neloxicam tablets in subjects with severe renal impairment is not recommended.

# 3 DOSAGE FORMS AND STRENGTHS Natorican Tablets USP. 7.5 m; Light yolwn round find the bevilet edged, tablet with U.S.L. debossed on one side and 7.5 setbossed correlation on the other side 1.5 m; Light yeak, raquitus sitespace, raquitus with U.S.L. debossed on one side and 15 debossed centrally on the other side

INGS AND PRECAUTIONS

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The accuracy to the transmission of the constrained calls of the stating of CMBC (see Terminal Termin

Avoid the use of Meloxicam in patients with a recent ML unless the banefits are expected to outweigh the risk of recurrent CV thrombotic events. If Meloxicam is used in patients with a recent NI, monitor patients for signs of cardiac ischemia.

5.2 Gastrointestinal Bleeding, Ulceration, and Perforation

3.1 Gathaltistical blacking, Ukraslin, and Parlondim. McMa, hushing municipation, can case straing superstrational (1) sympathic strains and the strain of the strain strains and the stra

Additionally regardless and advanced for decisional part of a characteristic Constraints. The Construct Advanced advance

Elevations of ALT or AST (less than three times ULN) may occur in up to 15% of patients treated with NSAIDs including meloxicam.

Inform patients of the warring signs and proptoms of hepatotock?) (e.g., nanosa, target, kthorpy, darhna, prottos, pandos, tytt copper quadret for interfaciencies, and "the prottos, pandos, tytt copper quadret for interfaciencies, and "the original control of the proton (s.e. as in proton (s.e. as in proton (s.e. as in proton (s.e. as in proton of the proton (s.e. as in proton (s.e

5.5 Heart Failure and Edema

Additionally, fluid retention and edema have been observed in some patients treated with NSAIDs. Use of meloxicam may blant the CV effects of several therapeutic agents used to treat these medical conditions (e.g., diuratics, ACE inhibitors, or angiotensin receptor bickners [ABBs]] [see Drug Interactions (7)].

Avoid the use of Meloscam in patients with severe hant failure unless the bondfits are expected to solvewigh the risk of extremely have failure. If Meloscam is used in patients which serves hand failure, monthe patients for large of ensuring have failure. 5.6 Read Tookty and HyperKatamini <u>Baral Tookty</u> and HyperKatamini Baral Tookty, and Meloscam, cased anni failure, and other rend inputs?

Rend taxicity lact also been seen in patients in whom rend prostaglanders have a compensationary role in the maintenance of renal perfusion. In those patients, administration of a short DAM may cause a date sequence fractions for prostagland formation and, accounting, in renal also flow, which may proceeded event rend. matching and accounting, in the all about form, which may proceeded event rend. matching and the second second

The renal effects of Meloxicam may hasten the progression of renal dysfunction in patients with preexisting renal disease. Because some Meloxicam metabolites are excreted by the kidney, monitor patients for signs of worsening renal function.

Correct volume status in dehydrated or hypovolemic patients prior to initiating Meloxicam. Monitor renal function in patients with renal or hepatic impairment, heart failure, dehydration, or hypovolemia during use of Meloxicam ( see Drug Interactions ( 7)

J. No information is available from controlled chical studies regarding the use of Ministerions (J. No information is patients with advanced related as Avaid Teau and Mikescam in patients with advanced related as a studies. Avaid The use of Mikescam Endowed Networks and the studies of th

<u>Huperkalama</u> Increases in serum potassium concentration, including hyperkalamia, have been reported with use of NSAIDs, even in some patients without renal impairment. In palaints with normal renal function, these effects have been attributed to a hyporenisment', hypadiosteronism state.

reportentiame-opposition stream status. 5.7 Anaphylactic Reactions Midioxiam has been associated with anaphylactic reactions in patients with and without known hypersensibility to midioxic an and in patients with apprivilentiative actimatic game contraindications (4) and Warmings and Privacations (5.8).

Seek emergency help if an anaphylactic reaction occurs.

Sake energiency hulp 1 an anaphyticit reaction accurs. 32 Eascretistics of Atoma Naticet of Carpio SeashNot An obspotiation of patients with anitiation may how applies acative actions attem may based private, information comparison by anaphysics have a presention of Zan-matching have a private and with PSAMDs has been reported in such applies and according transmission and anitiation of the same acative activity and according transmission and anitiation of the same acativity and according transmission activity and anitiation of the same acativity and according transmission activity and anitiation of the same acativity and according transmission activity and anitiation of the same activity and according transmission of the same activity and activity and according transmission of the same activity acti

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may deminish the utility of diagnostic signs in detecting intections. 5.13 Laboratory Monitoring Bacusios services GI bilading, hepatotoxic/by, and renal hijiry can occur without warning symptome or signs, consider monitoring patients on long-term MSAID treatment with a CEC and a chemistry profile periodically lise Warnings and Procautions (5.2, 5.3, 5.6).

The contrast protection (2.5.2, 5.16)
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 Detable and the contrast protection of the contrast protection of the contrast protection (2.5.1)
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6.1 Clinka Thulk Experiment Because circle structure are conducted under webly varying conditions, adverse reaction realised and a this clinka of a dirig cannot be directly compared to rates in the Analysis of the clinka structure of reflect the rate does not an experiment and advection of the structure of the structure of the structure of Alabit The Mathieum O 2, doing of a structure benefative. Bit 70, 00 a structure on The Mathieum O 2, doing of a structure benefative. Bit 70, 00 a structure on The Mathieum O 2, doing of a structure benefative. Bit 70, 00 a structure on The Mathieum O 2, doing of a structure benefative. Bit 70, 00 a structure on The Mathieum O 2, doing of a structure benefative. Bit 70, 00 a structure on the Mathieum O 2, doing of a structure benefative. Bit 70, 00 a structure on the Mathieum O 2, doing of a structure benefative. Bit 70, 00 a structure on the Mathieum O 2, doing of a structure benefative. Bit 70, 00 a structure on the Mathieum O 2, doing of a structure of a structure. Bit 70, 00 a structure on the Mathieum O 2, doing of a structure of a structure. Bit 70, 00 a structure on the Mathieum O 2, doing of a structure of a structure. Bit 70, 00 a structure on the Mathieum O 2, doing of a structure of a structure. Bit 70, 00 a structure on the Mathieum O 2, doing of a structure of a structure. Bit 70, 00 a structure on the Mathieum O 2, doing of a structure of a

Characterizati and Rhammatiki Amritiki The Methods million 2016 of cline tri of clineatise includes 10.122 OA patients and 1012 R patients translated with NetroCare 7.1 prophysy, 1955 OA patients and 1313 RA patients patients for at kase of clineatism of the state of the state one-yake Approximatily 2015 OF of these patients were translers for at kase one-yake Approximatily 2015 OF of these patients were translers for at kase one-yake Approximatily 2015 OF of these patients were translers for a kase one-yake Approximatily 2015 OF of these patients were translers that the approximation of the state one-bodies characterization of the state of the state one-yake of the state tras. 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 Neitoxicam with placebo and with an active control. Two 12-week multicenter, double-blind, randomized trials were conducted in patients with rheumatoid arthritis to compare the efficacy and

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## safety of Meloxicam with placebo.

# Table 1a depicts adverse events that occurred in x2% of the Meloxicam treatment groups in a 12-week placebo- and active-controlled osteoarthrifts trial. Table 1b depicts adverse events that occurred in x2% of the Meloxicam treatment groups in two 12-week placebo-controlled meumatoid arthrits trials.

	Placebo	Meloxicam 7.5 mg daily	Meloxicam 15 mg daily	Diclofenac 100 mg daily
No. of Patients	157	154	156	153
Gastrointestinal	17.2	20.1	17.3	28.1
Abdominal pain	2.5	1.9	2.6	1.3
Diarrhea	3.8	7.8	3.2	9.2
Dyspepsia	4.5	4.5	4.5	6.5
Flatulence	4.5	3.2	3.2	3.9
Nausea	3.2	3.9	3.8	7.2
Body as a Whole				
Accident household	1.9	4.5	3.2	2.6
Edema 1	2.5	1.9	4.5	3.3
Fal	0.6	2.6	0.0	1.3
Influenza-like symptoms	5.1	4.5	5.8	2.6
Central a n d Peripheral Nervous System				
Dizziness	3.2	2.6	3.8	2.0
Headache	10.2	7.8	8.3	5.9
Respiratory				
Pharyngkis	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

	Placebo Me	loxicam 7.5 mg dail	Meloxicam 15 m
No. of Patients	469	481	477
Gastrointestinal Disorders	14.1	18.9	16.8
Abdominal pain NOS *	0.6	2.9	2.3
Dyspeptic signs and symptoms <sup>1</sup>	3.8	5.8	4.0
Nausea	2.6	3.3	3.8
General Disorders and Administration Site	Conditions		
Influenza-like illness	2.1	2.9	2.3
Infection and Infestations			
Upper Respiratory tract infections-	4.1	7.0	6.5
pathogen class unspecified <sup>†</sup>			
Musculoskeletal and Connective Tissue Dis			
oint related signs and symptoms <sup>†</sup>	1.9	1.5	2.3
Nervous System Disorders			
Headaches NOS *	6.4	6.4	5.5
Skin and Subcutaneous Tissue Disorders			
Rash NOS *	1.7	1.0	2.1

# The adverse events that occurred with Meloxicam in a2% of patients treated short-term (4 to 6 weeks) and long-term (6 months) in active-controlled osteoarthritis trials are presented in Table 2.

Table 2 Adverse Events (%) Occurring in ≈2% of Meloxicam Patients in 4 to 6 Weeks and 6 Month Active-Cr Trials

	4.6 Weeks Co	atrolled Triple	6 Month Con	trolled Triple
	4-6 Weeks Co Meloxicam 7.5 mg daih			
No. of Patients	8955	256	169	306
Sastrointestinal	11.8	18.0	26.6	24.2
Abdominal pain	2.7	2.3	4.7	2.9
Constipation	0.8	12	1.8	2.6
Tiarrhea	1.9	2.7	5.9	2.6
Zyspepsia	3.8	7.4	8.9	9.5
bhilence	0.5	0.4	3.0	2.6
Naricea	2.4	4.7	4.7	7.2
/ombing	0.6	0.8	1.8	2.6
Body as a Whole				
Accident household	0.0	0.0	0.6	2.9
dema	0.6	2.0	2.4	1.6
Pain	0.9	2.0	3.6	5.2
Central and Peripheral Nervous Sy	stem			
Dizziness	1.1	1.6	2.4	2.6
Headache	2.4	2.7	3.6	2.6
Hematologic				
Anemia	0.1	0.0	4.1	2.9
Musculoskeletal				
Arthralgia	0.5	0.0	5.3	1.3
Back pain	0.5	0.4	3.0	0.7
Psychiatric				
ainmoan	0.4	0.0	3.6	1.6
Respiratory				
Coughing	0.2	0.8	2.4	1.0
Jpper 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 • WHO preferred terms edema, edema dep	0.3	0.4	4.7	6.9

WHO preferred terms edema, edema dependent, edema peripheral, and edema legs combined WHO preferred terms rash, rash erythematous, and rash maculo-papular combined	
igher doses of Neloxicam (22.5 mg and greater) have been associated with an creased risk of serious GI events; therefore, the daily dose of Neloxicam should not xceed 15 mg.	
ediatrics	
auciarticular and Polyarticular Course Juvenile Rheumatoid Arthritis (IRA)	
The handback and eighty-seven patients with paracterization and polyaritation course ARA are expected to Miscourse Markan and the discent analysis of LSIS to 15.37 mod 15.7	
he following is a list of adverse drug reactions occurring in <2% of patients receiving leloxicam in clinical trials involving approximately 16,200 patients.	

ing the course of the trials. The ecific subgroup effect.	adverse events did not demonstrate an age
is a list of adverse drug reacting	ns occurrine in <2% of patients receiving
clinical trials involving approxim	
Vhole	aleroic reaction, face edima, fatious, fever, hot flishes, making, switches, weight increase
	angina pactoris, cardiac failure, hypertension, hypotension, hypotension, hypotension, hypotension, hypotension,
	nconvulsions, paresthesia, tremor, vertigo
tinal	colits, dry mouth, duodenal ulcer, eructation, esophagits, gastric ulcer, gastribis, gastrointestinal hemorrhage, hematemesis, hemorrhagic duodenal ulcer, hemorrhagic gastric ulcer, intestinal perforation, melena, pancreatitis, perforated duodenal ulcer, stomatitis ulcerative
and Rhythm	arrhythmia, palpitation, tachycardia
6	kukopenia, purpura, thrombocytopenia
liary System	ALT increased, AST increased, bilrubinemia, GGT increased, hepatitis
nd Nutritional	dehydration
	abnormal dreaming, anxisty, appetite increased, confusion, depression, nervousness, somnolence
	asthma, bronchospasm, dyspnea
	alopecia, angloedema, bulbus eruption, photosensitivity reaction, pruritus, sweating increased, urticaria
	abnormal vision, conjunctivitis, taste pervension, tinnitus
tem	abuminuria, BUN increased, creatinine increased, hematuria, renal failure

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## 6.2 Post Marketing Experience

Body as a Who Cardiovascular Central and Pe Gastrointestin Heart Rate and Homstelook

- creat materials Experience
 The following adverse rescents have been diverted of adverse gaves are proved in or diverse rescents and the beam provided on the following adverse rescents the beam provided on the following adverse rescents are provided on the second on th

Warning

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

	Table 3 Clinically Significant Drug Interactions with Melosicam
<b>Drugs that In</b>	terfere with Hemostasis
Clinical Impact:	Nanckar and autocapaters such as warfarn have a synergistic effect on bleeding. The concentrater use of interkican and autocapaters behave in concentrative and and autocapaters behave in concentrative
intervention:	Mandeor patients with concomitant use of Matorican with anticoaquistats (e.g., warfurh), antipatiakit agents (e.g., apprint), selective service in respirable inhibitors (SBRIs) for signs of bianding (see Warrings and Precautions ( S 11)).
Aspirin	
Clinical Impact:	Eontrolled chical studies showed that the concomitant use of NSAIDs and analysis closues of aspirin does not produce any greater therapeutic effect than the use of NSAIDs alone. In a clinical study, the concomitant use of an NSAID and aspirin was associated with a significantly increased incidence of GI adverse reactions as compared to use of the NSAID alone [see Warnings and Precautions (5.2)].
intervention:	Cancombant use of Miloxiciam and low done aspirin or analysis dones of aspirin is not generally necemmended because of the increased risk of biadeding (see Warnings and Proceedions (511)). Melonicam is not a substitute for the done aspirin for cardiovascular protection.
ACE Inhibitor	, Angiotensin Receptor Blockers, or Beta-Blockers
Clinical Impact:	KMD: may denied the authyperturences effect of anglements of entry of anglement converting enzyme (Call) inhibits; a babbediner (including organical). materials who are addening view (call and call an
	During concentration are Melocina, monther for signs of worsning concentration are ACE inhibers, and ACE inhibers, and ACE inhibers, and and ACE inhibers, and and ACE inhibers or additional and ACE inhibers are administrated concentrativity, patients should be adequately
Diuretics	
Clinical Impact:	Chinal catalogies, as wells a post- matering descriptions, howed has KSADs reduced the nativents effect of loop durates (a.g., tensemble and material durates (a.g., tensemble and material durates), however, studies with foresemble aperts and method and the state durates in a nativent of effect. Foresemble single and material durates (a.g., tensemble and that durates are not affect for a state of the state
intervention:	During concombat use of Melosicam with duratics, observe patients for signs of worsening renal function, in addition to assuring duratic efficacy including anthyperturnsive effects [see Warnings and Precautions [ 5.6]].
Lithium	
Clinical Impact:	NSAIDs have produced elevations in plasma ithum levels and reductions in renal ithium clearance. The mean minimum ithium concentration increased 15%, and the renal clearance decreased by approximately 20%. This effect has been attributed to NSAID inhibition of renal prostaglandin synthesis (see Clinical Pharmacology (12-3)).
intervention:	buring concomitant use of Miduxicam and Ithium, montor patients for signs of Rhium toxicity.
Methotrexate	
Clinical Impact:	Concomitant use of NSAIDs and methotrexate may increase the risk for methotrexate toxicity (e.g., neutropenia, thrombocytopenia, renal dysfunction).
Intervention:	buring concomitant use of Melaxicam and methodrostata, monitor patients for methodrostata toxicity.
Cyclosporine	
Clinical Impact:	Concomitant use of Meloxicam and cyclosporine may increase cyclosporine's nephrotoxichy.
Intervention:	buring concentral use of Melosican and cyclosporine, monitor patients for signs of worsering renal function.
NSAIDs and S	
Clinical Impact:	Concomitant use of meloxicam with other NSAIDs or salicylatus (e.g., dflunical, salicalate) increases the risk of GI toxicity, with little or no increase in efficacy [see Warnings and Precautions ( 5.2)].
Intervention:	The concentrate use of metazican with other NSADs or saleytates is not recommanded.
Pemetrexed	
Clinical Impact:	Concomitant use of Melosicam and permetrexed may increase the risk of permetrexed-associated myelosuppression, renal, and Gl toxicity (see the permetrexed prescribing information).
	During concomitant use of Meloxican and penetroxed, In patients with renal impairment whose creatinine clearance ranges from 45 to 79 mL/min, monitor for myelosuppression, renal and GI toxicity.
Intervention:	Materials taking melosiciam should Interrupt dosing for at least five days before, the days of, and two days following permetereed administration.
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USE IN SPECIFIC POPULATIONS
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trimester) [ see warnings and Precadoons ( 5.20) ].	
There are no adequate and well-controlled studies of Meloxicam in pregnant women. Data from observational studies regarding potential entryrolitat irisks of NSAID use in women in the first or second throatisms of pregnancy are inconclusion. In the general U.S. population, al chirally necognized pregnancies, regardless of drug exposure, have a background read or 2-4% for margin malformations, and 15-20% for pregnancy back	
In animal reproduction studies, embryofetal death was observed in rats and rabbits trated during the proof of organogeness with metricicat and rabbits of doses equivalent 0.8.5- and 6.5-tmost the maximum recommanded human dose (HRHD) of Metrician embryopeness with metalschan at an oral dose equivalent to 72-times the Metrici. In pro- desing dose and a dose and a star and a dose equivalent to 72-times the Metrici. In pro- desing dose and a dose and a dose and a star and a dose	e- 1.
Based on animal data, prostaglandins have been shown to have an important role in endomstrial vascular permeability, blastocyst implantation, and decidualization. In anima studies, administration of proctaglandin synthesis inhibitors, such as meloxicam, resulted in increased pre- and post-implantation loss.	a

# These are no sources on the effects of Meloscam during abor or delivery. In animal tables, PEADO: A block meloscam, initiate prostagandin synthesis, classe delayed patrumtico, and increase the incidence of stillarth.

Meloxica organog mg of M rabbits t the hear compari based or meloxica

greater, respectively, than the MRHD based on BSA compa throughout organogenesis.

Branghour depropersions. On administration of exploses, beginning and the spectration through latetions included the schedule of spectra, beginning and the spectration through latetions becamparation. **E.1 Lattone E.2 Lattone E.3 Lattone** Through any horton data available on whitefur metasicam is present in human mitik, gr Machiner of a spectration of the schedule of the schedule of the schedule of the Machiner of the schedule of the schedule of the schedule of the schedule of the Machiner of the schedule on schedule on the schedule of the schedule of the schedule of the Machiner of the schedule on schedule on the schedule of the schedule of the schedule of the Machiner of the schedule on schedule on the schedule of the sche

<u>Data</u> Animal Data

Meloxican was present in the milk of lactating rats at concentrations higher than those in plasma.

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8.7 Renail Impairment No dose adjustment is nocessary in patients with mild to moderate renail impairment. Patients with severe renail impairment have not been studied. The use of Meloxican in subjects with severe renail impairment is not recommended. In patients on hemologikas, mildiocam should not exceed 7.5 mg per day, Meloxicam is not diayable [ see Dosage and Administration ( 2.1) and Cline if hemorecology ( 2.12.3).

10 OVERDOSAGE Symptoms Robwing acute NSAID overrisosages have been typically limited to lethorgy, drewinses, nauce, wombing, and explastic pair, which have been generally reversible with supportive care. Gastrotietestral blanding has a cocurred, hyperinnesh, acute read and Prescedance (1, 5, 5, 5, 4, 6, 5).

Manage patients with symptomatic and supportive care following an KSAID overforsage. There are no specific antidotes. Consider emissis and/or activated characal (60 to 100 grams in adult). It of grams per kig of being weight in patient gradients and/or controls cathoritis in symptomatic patients seen within from hours of negations or in patients, skalamiser porticinate of a set of the second model dough, Forcata patients, skalamiser of units, hereoclapist, or hemoperfusion may not be useful doub to high protein binding.

There is limited experience with metoxican overdosage. Choistytramine is known to accelerate the charance of metoxican. Accelerated fermional of metoxican by 4 g oral Administration of choistynmine may be used fichioning and workdoage. For additional information about overdosage treatment, cal a policin control center (1-800-322122):

11 DESCRIPTION More can Tables: UP are a noncorrected and informationy drug (MSAD). Each tables contains 7.5 mp ang makerkam for oral administration. More can is charactering designates at 4-hydroxy-2-anety- $H_{\rm C}$ -methy-2-bitaph/2-H-1,2-bitrobilizme-3-carboranish-1,1-bitrok. The melecular walks is 351.4.1 is empirical formula is C \_2H 12H 2O \_{2} \_{2} = and E has the following structural formula:

# and a

Molectam is a pastel yellow sold, practically insoluble in water, with higher solubility observed in strong actits and basics. It is very slightly soluble in methand Makinzkam Makinzkam has plan values of 11 and 4.2. We have a 1.1 in or 4 consolution per 1.7. A Makinzkam Makinzkam Makinzkam kawa have based and a sublet as a tablet for oral administration containing 7.5 mg or 15 mg makinzkam.

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

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12.3 Pharmacokinetics

Table 4 Sing	le Dose and S	teady-State	Pharmacokine	tic Parameters for	Oral 7.5 mg and 15 m	g Meloxicam (Mean ani	1% CV) .
				Steady State			gle Dose
Pharmacokinetic Parameters (*	(CV) F					Renal failure (Fasted) H	lepatic insufficiency (Fasted)
		7.5 mg *	tablets	15 mg capsules	15 mg capsules	15 mg capsules	15 mg capsules
N		14	8	5	8	12	12
C max	[µg/mL]	1.05	(20)	2.3 (59)	3.2 (24)	0.59 (36)	0.84 (29)

t max	[h]	4.9 (8)	5 (12)	6 (27)	4 (65)	10 (87)
t 1/2	[h]	20.1 (29)	21 (34)	24 (34)	18 (46)	16 (29)
CL/f	[mL/min]	8.8 (29)	9.9 (76)	5.1 (22)	19 (43)	11 (44)
V 2/15	[L]	14.7 (32)	15 (42)	10 (30)	26 (44)	14 (29)
* The parameter values in the t † not under high fat conditions 2 Meloxicam tablets 5 V Z/f =Dose/(AUC+Kel)	table are from various studies					

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The process of a classication strategies. We the Realisticity distribution of the guards Declaration concentrational to important Buff, affer and dates, race profession of the important buffer and the properties of the strategies of the date and the properties of the strategies of the strategies of the strategies of the strategies of the dates and the strategies of the strategies of the strategies of the strategies of the classical constraints of the strategies of the strategies of the strategies of the classical constraints of the strategies of the strategies of the classical strategies of the strategies of the strategies of the classical strategies of the strategies of the strategies of the properties of the strategies of the strategies of the properties of the strategies of the strategies of the properties of the strategies of th

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Specific Populations -Pediatric

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Etisrly make (seS years of age) exhibited meloxicam plasma concentrations and staady-state pharmacolenatics initiar to young make. Etisrly females (seS years of age) had a 47% high ethod Crass as at compared to younger concentrations in the stellary female, the adverse every grofts was compared for both ethory patient population. A smaller fire for faction was found in stillary female patients in comparison to siderly make patients.

Angust: Impairment: Federing a single 5 mg dates of maloscam there uses to marked difference in plasma concentrations in plasmits with mild (CHS-Hop) Case II (or modurate (CHS-Hop) Case II) is plasmit (memory modulate). The single concentration of the single of the mild of the single concentration of the single concentration of the single concentration with mild to modulate insplace regarisment. Frances with even heads: the plasma of the single concentration of the Market concentration of the single concentration of the single concentration of the Market concentration of the single concentration of the single concentration of the Market concentration of the single concentration of the single concentration of the Market concentration of the single concentration of the single concentration of the single concentration of the Market concentration of the single concentration of the single concentration of the single concentration of the Market concentration of the single concentration

# enall impairment. Total drug above an encontration of maintainan devenand and table isotance of maintainan encoderation that the base part of maintainane maintain free part values were similar in all groups. The higher medication statutors in subjects with reading anyments may be able to increased retarch to indrused maintainan which is available for higher employees and sub-reased retarch of unboard maintainan which is available have not been adapted traditional statutors and the source relation impairments is not encommended in an Dual statutor and the source relation impairments is not encommended in an Dual statutor (2015). Warnings and Presentation (2016) and the signative Regulation (2017).

Hemodaliyal Folowing a single dose of milioxicam, the free Cmax plasma concentrations were higher in galantic with reveal take is on chronic hemodaliyal. (15) file infraction) to concentration in plasmic therefore, additional dose; are not encessary after hemodaliyals; Meksicam is not dialyzable [see Dosage and Administration (2.1) and Use in Specific Population (2.7).

hemodality, Allicocch in the divide land bases and Administration (2.1) and the Marchinest Constraints (2.1) and the second sec

Methodrowate: A study in 13 resonance of entries (MA) parkets vehanced the effects of marked AMM and a final state on the bagerout effects of the parket state of the state o

## 13 NONCLINICAL TOXICOLOGY

13 NORCLIMICAL TOXICOLOGY 13. Carcinogenesis, Mudagenesis, Impairment of Fertility <u>Cercinopensis</u> 104 weeks) and mice (99 weeks) administered metrican at cardioses to 0.8 mg/staysh rata and up to 8.0 mg/staysh mice (pt to 15.4mg/stay). Area respectively, The maximum recommended human dese (BMHO) of 15 mg/stay Mexican basics on boys strate and (ESA) commended.

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

## int of Fertility

Meloxicam did not impair male and female fertility in rats at oral doses up to 9 mg/kg/day in makes and 5 mg/kg/day in females (up to 5.8- and 3.2-times greater, respectively, than the MRHD based on BSA comparison).

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The case of Philotecan for the management of signs and symptoms of esteen-thresh es-sented to the normality of the signs of the signs

14.2 juvenile Rh Course umatoid Arthritis (JRA) Pauciarticular and Polyarticular

# The use of Meloxicam for the treatment of the signs and symptoms of pauciarticular polyarticular course juvenile Rheumatoid Arthritis in patients 2 years of age and older was evaluated in two 12-week, double-blind, paralel-arm, active-controlled trials.

Both choice included three arms: negregate and the data of mildicates. In both mapping the second s

17 PATIENT COUNSELING INFORMATION Advice the patient for read the FDA-approved patient labeling (Medication Guide) that accompanies active prescription disperside). Additional Medication Guides can be obtained by calling Unichem at 1-866-562-4616. Inform patients, families or their caregivers of the following information before initiating therapy with an NSAID and periodically during the course of ongoing therapy. Cardiovascular Threehotic Events Advise patients to be alert for the symptoms of cardiovascular threehotic events, including cheat pair, shortwise of breath, weakness, or shuring of speech, and to support the symptoms to their threath-new provide immediately (Lee Warnings and Processitions (5.1)). Cardiovascular (Basering Ubcration, and Perforation

Cancertaintial basing. Unrandow and Parliadian Davis parkets in provide providence of a version of a basing in scheding oppositive, of a schedule in providence of a version of a version of a version of a version of the abase appendix of a version of version of a version of a version of a version of version

Hear Falser and Falsers Advise patients to be aim for the symptoms of congestive heart falser including stortistics of teaching including and proceedings of the stortight from Matthicen Analthicent Reactions. Homom patients of the signs of an analyticatic reaction (e.g., officer) yourship, swelling of the face or thready, instruct, patients to seak immediate emerging (e.g., p. ).

Advise patients to stop Meloxicam tablets immediately if they develop any type of rash and to contact their healthcare provider as soon as possible [ see Warnings and Precautions ( 5.9) ]. Precautions (5.9)]. <u>Female Facility</u> Advise females of reproductive potential who desire pregnancy that NSAIDs, including Maloxican tablets, may be associated with a reversible delay in ovulation [see Use in Specific Populations (3.3)]. Inform pregnant women to avoid use of Meloxicam tablets 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)

Avoid Concentrate Lise of IXSAIDs. Inform patients that the concentrate use of Meloxicam tablets with other NSAIDs or salkylates (e.g., diffurisis, salaatab) is not recommended due to the increased risk of gaterintestratin dexity, and Bitto on increase in efficiency (a well and and Precaution (5.2) and Drug Interactions (7.1). Nat't patients that RSAIDs may be present in voive the counter "medications for trainment of codes, fore; or insomnita.

Inform patients not to use low-dose aspirin concomitantly with Meloxicam tablets until they talk to their healthcare provider [ see Drug Interactions ( 7) ]. For current prescribing information, call Unichem at 1-866-562-4616.

Real examples is, it grande 2006 2006 2006 2006 2006 2006 2007 2

Cardiovascular Thrombotic Events Nonstaroidal anti-inflammatory drugs (NSAIDs) cause an increased risk of serious cardiovascular thrombotic events, including myocardial infarction and stroke, which can be fatal. This risk may occur early in treatment and may

Serious Skin Reactions

Avoid Concomitant Use of NSAIDs

Use of NSAIDs and Low-Dose Aspirin

For current prescribing information, Manufactured by: UNICHEM LABORATORIES LTD. Pilerne Ind.: Estate, Pilerne, Bardez, Goa 403511, India Markoted by:

Hasbrouck Heights, NJ 07604 06-R-09/2016

Dispense tablets in a tight container. Keep this and all medications out of the reach of children.

makick am kine ginapit. 11 HOM SUPPLICATIONED AND MANCLINE Makickan tables (UP are available as supply solver, mod. flut, uncessed ab table constraining matchicks (TS) may be available as follow: Makickan Tables (UP 7) may be available as follow: Notickan Tables (UP 7) may



# Table La expeste adverse events that accounted as that of the Medica actuation and accounted as the second actual actual

## Abdominal pain NOSMedDRA preferred term: nausea, abdominal pain NOS, influenza-like liness, headaches NOS, and rash NOS

Advances and MCMERGE Age reference term, makes, advanced age and MCMERGE Age reference term makes, advanced age and mCMERGE ag

General Disorders and Administration Site Conditions

Influenza-like illness 2.1 2.9 2.3 Infection and Infestations

tory tract infections-pathogen class unsp ecified

Upper Respiratory tract infections-pathogen class 4.1 7.0 6.5 Musculoskeletal and Connective Tissue Disorders

Joint related signs and symptoms 1.9 1.5 2.3 Nervous System Disorders

Headaches NDS 6.4 6.4 5.5 Skin and Subcutaneous Tissue Disorders

Matorian 7.5 mg day
Matoran 1.5 mg day</

Cardi angina pe vasculitis -uncurra Central and Peripheral Nervous System convulsions, paresthesia, tremor, vertigo Gastrointestinal Colitis, dry mouth, duodenal ulcer, eructation, esophagtis, gastric ulcer, gastričis, gastroscophagai irdluz, gastrointestinal hemorrhage, hematamasis, hemorrhagi duodenal ulcer, hemorrhagic gastric ulcer, hitschal perforation, melena, parcreatits, perforated duodenal ulcer, perfortand gastric ulcer, stomatis ulcerative



Pruritus 0.4 1.2 2.4 0.0 RashWHH combinen 0.3 1.2 3.0 1.3 Urinary

tous, and rash maculo-papular

edema, edema dependent, edema peripheral, and edem

Body as a Whole

Dizziness 1.1 1.6 2.4 2.6 Headachs 2.4 2.7 3.6 2.6 Hernatolr

Anemia 0.1 0.0 4.1 2.9 Musculo

Arthraigia 0.5 0.0 5.3 1.3 Back pain 0.5 0.4 3.0 0.7 Psychiatric

Insomnia 0.4 0.0 3.6 1.6 Respirato

Coughing 0.2 0.8 2.4 1.0 Upper res 0.2 0.0 8.3 7.5 Skin

y tract infectio

Heart Rate and Rhythm arrhythmia, palpitation, tachycardia Hematologic

leukopenia, purpura, thrombocytopenia Liver and Bilary System

ALT increased, AST increased, bilirubinemia, GGT increased, hepatitis

## Metabolic and Nutritional dehydration

Psychiatric

abnormal dreaming, anxiety, appetite increased, confusion, depression, nervousness, somnolence

somnoence Respiratory asthma, bronchospasm, dyspnea Skin and Appendages alopecia, angioedema, bulbus eruption, photosensibility reaction, pruritus, sweating increased, urticaria

A lange of the second secon

Meloxicam and anticoagulants such as warfarin have a synergistic effect on bleeding. The concombant use of meloxicam and anticoagulants have an increased risk of serious bleeding compared to the use of either drug alone.

Serotonin release by platelets plays an important role in hemostasis. Case-control and cohort epidemiological studies showed that concomitant use of drugs that interfere with serotonin reuptake and an NSAID may potentiate the risk of bleading more than an NSAID alone.

## Intervention :

Monkor patients with concentrant use of Meloxiam with anticologulants (e.g., warfarin), antipatient agents (e.g., aspirin), selections exercision respirate influence (SSRs), and Warning and Procustoms (S-11)). Warning and Procustoms (S-11).

Clinical Impact :

# Curical impact : Controlled child situations showed that the concomitant use of NSAIDs and analgesic doese of apprint does not produce any greater threapoutic effect than the use of NSAIDs alone, is a final a study, the concomitant use of a nASD and apprint wait ASD and apprint and a study of the concomitant use of an ASD and apprint waits to use of the NSAID alone (see Warnings and Precautors (5-21)). Intervention :

Concombant use of Meloxican and buy dose acprin or analgesic doses of asprin is not generally recommended bacause of the increased risk of bleeding (see Warnings and Precactions (S11)) Meloxican in real acated for low dose acprin for and/ouscular protection. Actionations, Angotensin Receptor Blockers, or Beta - Blockers Christ Impact :

NSAIDs may diminish the antihypertensive effect of angiotensin converting enzyme (ACE) inhibitors, angiotensin receptor blockers (ARBs), or beta-blockers (including propranolo).

In patients who are elderly, volume-depleted (including those on diuretic therapy), or have renal impairment, coadministration of an HSAID with ACE inhibitors or ARBs may result in determation of renal function, including possible acute renal failure. Thise effects are usually reversible.

During concomitant use of Meloxican and ACE inhibitors, ABBs, or bata-blockers, monitor blood pressure to ensure that the desired blood pressure is obtained. During concomitant use of Meloxican and ACE Inhibitors or ABBs in patients who are electric values degletad, or have impaired read function, months for algois of ensuring new concomitant, passes thould be adequarely hydraut. Assess: read function at the bagroing of the concomitant treatment and particularly threadfare. Durings of the concomitant treatment and particularly threadfare.

Clinical Impact :

Choice causes, as we as poor numericing observations, showed that XRADP reduced the hardwards effect of a bop Survival erg. 5, increasingly introduced and processing of patients. This effect has been attributed to the NADD Inhibition of neural processing of applications. This effect has been attributed to the NADD Inhibition of maniper and demonstrated a velocition in number call effect. Proceeding single and multiple does of malabilities and patients were sense of single and multiple does effect and parameters and an effect. Proceeding single and multiple does effect and parameters and an effect and the single soft and the malabilities.

During concomitant use of Melsoican with divertices, observe patients for signs of worsaming remail function. In addition to assuring divertic efficacy including architypertransive effects (see Warnings and Precautions (5.6)): Librium Clinical Impact :

NSAIDs have produced elevations in plasma lithium levels and reductions in renal Rhium clearances. The mean minimum Rhium concentration increased 15%, and the renal clearance decreased by approximately 20%. This different has been attributed to NSAID imbition of renal prostaglandin synthesis (see Clerical Pharmacology (12.3)). Intervention :

During concomitant use of Neloxicam and Ithium, monitor patients for signs of Ithium toxicity.

# Methotrexate Clinical Impact :

Concombant use of NSAIDs and methotrexate may increase the risk for methotrexate toxicity (e.g., neutropenia, thrombocytopenia, renal dysfunction).

During concomitant use of Meloxicam and methotrexate, monitor patients for methotrexate toxicity.

# Cyclosporine Clinical Impact

icam and cyclosporine may increase cyclosp Concomitant use o nephrotoxicity.

## Intervention :

During concomitant use of Nelsociam and cyclosporine, monitor patients for signs of worsaning renal function. NSAIDs and Salociates Clerical Impact: :

Concording use of meloxicam with other NSAIDs or saleylates (e.g., dflunical, saleabab) increases the risk of GI toxicity, with little or no increase in efficacy (see Warnings and Precautions (5.2)). Intervention:

The concombant use of meloxicam with other NSAIDs or salicylates is not recommended.

Pemetrexed Clinical Impact use of Meloxicam and permetroxed may increase the risk of permetroxed yelosuppression, renal, and GI toxicity (see the permetroxed prescribing

associated m information).

During concomitant use of Neloxicam and pemetrexed, in patients with renal impairment whose creatinine clearance ranges from 45 to 79 mL/min, monitor for myelosuppression, renal and GI toxicity.

## Patients taking meloxicam should interrupt dosing for at least five days before, the day of, and two days following permetrexed administration.

In patients with creatinine clearance below 45 mL/min, the comeloxicam with pemetrexed is not recommended.

# milk, or on the effects on breastled infants, or on milk production. The mmmal and health benefits of breastleeding should be considered along with the sciffical mode (or Molockam and any potential alovers effects on the breastled arm was present in the milk of lactating rats at concentrations higher than those no. <text><text><text><text><text><text><text> developm mother's infant froi Meloxican in plasma Infertility I NSAIDs, i has been have show

Steady State Single Dose Pharmacokinetic Parameters (% CV ) Healthy male adults (Fed ) not u Elderly males (Fed ) Elderly females (Fed ) Renal failure (Fasted ) Hepatic insufficiency ( Fasted )

7 . 5 mgMeloxicam tablets tablets 15 mg capsules 15 mg capsules 15 mg capsules 15 mg capsules N





LOT# 77777	EXP 19-17	LOT#???	2000 LO	# 77-77	ECP 1	10.99
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MELOXICAM meloxicam tablet	naug side	CATCONIA	ALED HOOM 1	EMPERAT	URE 151-301	0.007-00171
Product Information						
Product Type	HEMAN PRE DRUG	SCRIPTION	item Code (Source)	1	DC-60760-608 24)	NEC 21300-
Route of Administration	a CRAL					
Active Ingredient/Ac	tive Moiety					
	ogredient Nam			Basis c	f Strength	Strengt
MELDINGAM (JAN: VOJOPR)	COL) (MELONCAN	- UNIT VG2 QFE	icer)	MELCOOCAN	4	7.5 mg
Inactive Ingredients						
	Ingredi	ont Name			:	Strength
CELLULOSE, MICROCRYST	ALLINE (UNIL OPT	R32061U)				
CROSPOVIDONE (UNII 2571						
LACTORE MONOHYDIATE		3				
MAGNESIUM STEAKATE (LI						
POVIDONE K30 (UNII U725)						
SEJCON DIOXIDE (UNIL ET)						
TRISODIUM CITRATE DINVI	SKATE (UNI: 8225	4789540				
Product Characteris	tics					
Color	yellow	Score			no score	
Shape	ROUND	Size			7040	
Flavor		imprint 1	ode		U(L)7,5	
Costains						

٠	item Code	Packa	ge Descript	ion	Marketing Start Date	Marketing End Date
1	NDC 60760- 600-10	10 in 1 BOTTLE, PLN Combination Product	TE; Type D P	65 a	03/03/2016	
1	404-30	30 in 1 BOTTLE, PLM Combination Product			02/03/2016	
	400-60	Combination Product		03/03/2016		
				03/03/2016		
-		Product		03/05/2018		
	NDC-60760- 600-14	14 in 1 BOTTLE, PLM Combination Product	TE Type D P	65 a	03/05/2018	13/31/2019
	Marketing Application Number or Monograph Category Citation		Citation		Date	Date
	Category	approxim	Citation	dane graph	Marketing Start Date	Marketing End Date
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