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

See rus prescribing information for complete boxed warning.

Nonsteroidal anti-inflammatory drugs (MSAIDs) 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 increase with duration of use. (5.1).

can be fail. This risk may occur early in treatment and may increase with dustion Medicacian is contradictated in the stating of comeany enterpt lypans graft (CABC). NSASOR cause an increased risk of serious gastrointestinal (CI) adverse events lockeding bleeding, decreation, and gendrection of the satisface for instantians, which symptoms. Elderly positions and patients with a prior history of popic user of symptoms. Elderly positions and patients with a prior history of spelic user disease add/or to bleeding as at a general risk for scales of contradicts. (3-1)

INDICATIONS AND USAGE
Im Tablets, USP is a non-sheroidal arti-inflammatory drug indicated for artificial (AU(1.1) ameliod Arthritis (BA) (1.2) sile Pheumatoid Arthritis (BA) in patients who seligh a 60 kg (1.3)

Disage and administration

Dosage and administration

Use the lowest effective dose for the shortest duration consistent with individual patient treatment goals

(2-1).

CASIN (22) and 84 CAS (23) Casting that 27 cap oper daily control of the control

Notion hypersensitivity to melascian or any components of the drug product (4)

Notions hypersensitivity to melascian or any components of the drug product (4)

Nisiony of altitime, unicular, or other altergic-type reactions after tailong aspin or other NSADs (4)

In the sating of CADS coursey (4)

The transport of the control of the day of the day

Most common (a) Se and greate that placefully district sends in adults and denhes, upper respiratory fact affection, dispersis, and influences descriptions (c).

 Adverse words closered in pediatric studies were similar in nature to the adult circuit trial experience (c).

To report SUSPECTED ADVERSE REACTIONS, contact Strides Pharma Inc. at 1-877-244-9825 or go to www.strides.com or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

- Section 2 - Sect

Infertility: NSAIDs are associated with revenible infertility. Consider withdrawal of meloxicam in women who have difficulties conceiving (8.3)

Manage :
 With his difficulties conceiving (8.3)

See 17 for PATIENT COUNSELING INFORMATION and Medication Guide.

Revised: 6/2021

FULL PRESCRIBING INFORMATION: CONTENTS* WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL 1.1 Disconstricts (DA) 1.2 Reburnation stricts (DA) 1.2 Reburnation stricts (DA)

1.1 Ostooarthritis (OA)
1.2 Rheumatoid Arthritis (RA)
1.3 Juvenile Rheumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course
2 DOSAGE AND ADMINISTRATION
2.1 General Dosing Instructions
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2.1 General Dosin (Instructions
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Pediatric Use
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11. DESCRIPTION
12.1 SECRIPTION
12.1 Nechanism of Action
12.3 Non-CLINICAL TOXICOLOGY
13.1 CACKIOGENER, Malagenesis, Impairment of Fertility
CLINICAL TOXICOLOGY
13.1 Cackinogenesis, Malagenesis, Impairment of Fertility
CLINICAL STUDIES

WARNING: RISK OF SERIOUS CARDIOVASCULAR AND GASTROINTESTINAL EVENTS

GASTROMESTHAL EVENTS

**Nonstrovidal anti-inflammatory drugs/IMEADS) cause an increased risk

**Nonstrovidal anti-inflammatory drugs/IMEADS) cause an increased risk
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inflammators and may licrease with duration of use [see Warnings and
intrastance and may licrease with duration of use [see Warnings and
IMEADS (audits) carterial decided in the setting of comonary artery lypiosis
graft (CABG) surgery [see Contraindications (4) and Warnings and
Pricoacionics (3.7)

Presentions (5.3)**

Continues the Management of the Management of the Management of M

1.1 Osteoarthrikis (OA)
Meloxicam Tablets, USP is indicated for relief of the signs and symptoms of osteoarthrikis [see Clinical Studies (14.1)].

Rheumatoid Arthritis (RA)
 Meloxicam Tablets, USP is indicated for relief of the signs and symptoms of rhe arthritis [see Clinical Studies (14.1)].

1.3 Juvenile Rheumatoid Arthritis (JRA) Pauciarticular and Polyarticular Course Meloxicam Tablets, USP is indicated for relief of the signs and symptoms of pauciarticular or polyarticular course juvenile Rheumatoid Arthritis in patients who weigh >60 kg (see Dosage and Administration (2.4) and Clinical Studies (12.4) and

A CUASMA AND ADMINION IN ATTOM

2. General Design Instructions

Carefully consider the potential booths and risks of Meloxicam Tablets, USP and other
thermaters options before designing to use Meloxicam Tablets. USP. Use the lowest
efficient losse for the shortest duration consistent with individual patient treatment
each (see Winnings and Percactions (6)).

After observing the response to their thermay with Meloxicam Tablets, USP, adjust the
loads in the management of the control of the control

In adults, the maximum recommended daily oral dose of Meloxikam Tablets, USP is 15 gregories for formulation. In patients with hermodalysis, a maximum daily dosage of 7.5 mg is recommended (see Use in Specific Populations (8.7), and Clinical Pharmacology (12-3)). #Eloxicam Tablets, USP may be taken without regard to timing of meals.

2.2. Osteoarthrikis

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

2.3 Rhoumatod Arthritis
For the reider of the signs and symptoms of rhoumatoid arthritis, the recommended
starting and multinease or and some of Newtochar Tables, USP is 7.5 mg once daily.
Some patients may receive additional benefit by increasing the dose to 15 mg once daily.

2.4 Juvenile Rheumatold Arthritis (JRA) Psuciarticular and Polyarticular Course
For the treatment of juvenile rheumatold arthritis, the recommended oral dose of
Molosciam Tablets, USP is 7.5 mg once day in children who weigh a60 kg. There was no additional benefit demonstrated by increasing the dose above 7.5 mg in clinical trists.

Meloxicam Tablets, USP should not be used in children who weigh <60 kg

Renal Impairment
 The use of Moloxicam Tablets, USP in subjects with severe renal impairment is not recommended.

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

2.6 Non-Interchangeability with Other Formulations of Meloxicam

Meloxicam Tablets, USP have not shown equivalent systemic exposure to other approved formulations of oral meloxicam. Therefore, Meloxicam Tablets, USP are no interchangeable with other formulations of oral meloxicam product even if the total miligram strength is the same. Do not substitute similar dose strengths of Meloxicam Tablets, USP with other formulations of oral meloxicam product.

- Meloxicam Tablets, USP:

 7.5 mg: light yellow colored, oval shaped uncoated tablets engraved with 5 160 on one side and plain on other side.

 15 mg: yellow colored, oval shaped uncoated tablets engraved 5 161 on one side and plain on other side.

- -. Am summictATIODS
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5 WARNINGS AND PRECAUTIONS

S VARAMINGS AND PRECAUTIONS

3.1. Cardivascular Thremshorks: Family
Clinical trian of soweral CDU2 2 detective and monesteristics ISABD; of up to three years
durated have been an increased risk of a foreize cardiovascular (CDU2) thremshork
events, isolating impactabilit inference half year durated, which can be train Based on
events, isolating impactabilit inference half year durated, which can be train Based on
the residual concerns an ensemble of the property of the prope

about the symptoms of serious CV events and the steps to take if they occur.

There is no consistent soldness that concurrence and experimentation be increased assign and an MSARD, such as melosticam, increases the risk of serious gestrointested (old events [see thready and Procastion 6.7 of Section 1.2 o

Post-MI Patients

Pack ME Platinist

Observational studies conducted in the Danish National Registry have demonstrated that patients treated with NSAIDs in the post-Me period were at increased risk of reinfarction, CV-related edam, and alexane mersially period were at increased risk of reinfarction in the CV-related edam, and alexane mersially period period representation of treatment. In this year, in NSAID restance and alexane mersial period restaurance and restaurance and

the next rour years of rotow-up.

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

with a record M, monitor polaters for signs of cradics inchemia.

3.5 Castroinetsstate Milesdeuig Mucration, and Perforation
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MAMAD, including mitoscam, can cause sention gastroinetsiand (GII adversa events
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cocur at any time, with or without average symptomer. In palents treated with MAMADmitoscap symptomics (Upper GII Advers, possibility) profit profits or page
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Without risk. Risk Factors for Gi Bleeding, Ulceration, and Perforation

Bible Extent son for Ill Bibleofing, Mexicity, and Perforation
Platfers with a prior having of project used regions and/or of bleeding who used MSAIDs had a greater from 10-10-fed increased in 18 for developing and to bleed compared to patient had been provided by MSAIDs and the property of the proper

- Additionary, picteries with advanced her disease and/or couplingship are at increased Scientific to Missens, the Clinic Miss. NIASI (MISS of the Missens and Missens). If the Missens are already of the Missens and Missens are already to a time, precise of the Advanced Missens and Missens are already of the Missens and Missens and Missens are already of the Missens and Missens and Advanced Missens and Missens and

5.3 Hepatotoxicity

5.3 Hepatotoxicky Blevolators of AI, Tor AST (three or more times the upper limit of normal (ULNI) have been reported in approximately 1% of MSAID-treated patients in circular rate. In addition, memorises, and hepatic plants are the proported, in Activity plants the hepatiles, been encrose, and hepatic faither have been incoported, in Activity plants the hepatiles, been except and the AID of AID (Institute of AID) and the AID (

Treated with NaHUE recursing misosceam. Inform patients of the warring signs and symptoms of hepatotoxicity (e.g., naissa, inform patients of the warring signs and symptoms or the signs of the signs

5.4 Hypertension

5.4 Hypertension
NSAMD, including melasizam, can lead to new onset or worsening of preexisting hypertension, either of which may contribute to the in-creased incidence of CV events. Pleatents taking application inconverting regime (ACE) inhibitors, that abide duretics, or bog duretics may have impaided response to these therapies when taking NSAMD. [see Drug inferentiations].
Nontribute bods pressure (BP) during the initiation of NSAMD treatment and throughout the course of threating.

course of therapy.

5.5 Near Failure and Edema

The Cox band traditional RIAOD Trialists: Colaboration meta-analysis of randomized controlled randomized from the cox band traditional RIAOD Trialists: Colaboration meta-analysis of randomized controlled randomized and programmary two folial increase in hospitalizations for controlled randomized from the controlled randomized positions for the controlled randomized from the controlle

and death. Additionally, fluid retention and edema have been observed in some patients treated with NSAIDs. Use of meloxicam may blant the CV effects of several therapoutic agents used to treat these medical conditions (e.g., duringsets, AZE Pribliotrs, or applicents necessfor blockers [ARBS]) [see Drug Interactions (7)]. Available of the conditions of

seasows to covering mer as of winovaring beart failur. If induction is such is patients, who were here in Maria member and the patients with sover hear that member patients of significant in Maria Touckty and Hyperkalamin.

Amand Touckty and Hyperkalamin in Maria Maria

(7)1. Wo information is available from controlled clinical studies regarding the use of motivacion in patients with advanced renal disease. Aveid the use of motivacion in the first of the control of the tild of working in certain function. If methods can is used a planter with advanced renal disease, monitor patients for signs of worsening renal function [see Clinical Ammancology (22) and control of the control of the

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 hypereninemic-hypoaidosteronism state.

hypomenisms—bypositionisms state.

3.7 Anaphysick Execution

Missiscent has been associated with maphysick reaction in patients with and without hypomenisms, for modecare and a planters with apply-associative atthread (see the patients) of the patients with apply-associative atthread (see Seek emergency legs of an apply-plack reaction occurs.

5.8 Executations of Anthera Related to Aughin Searchibery Anaphysics and Anaphys

symptoms of arbitms.

35. Serious Sikh Reactions
(NADA)s, including melavicam, can case serious skin adverse reactions such as
endicible dermaits. Sowers-Johnson Symborne (SS)s, and took optimum encreyiss.
(TRIU, with six tab in fair. Their serious overst irray score without it wantly, lifetim
used in entire case in the six table of the six table of the six table of melavicam is the rist appearance of six and or any other sign of decident the
hypersensity, Melavicam's continuidated in patients with previous serious skin
reactions in Sixther, I of Contributations of the

reacces on SMAIDs [see Contraindactors (4)].

5.3.10 Fung Reaction with Ecoloophila and Systemic Symptoms (DRESS) for particular of the State of the State of Systemic Symptoms (DRESS) has been reported in pointer its baling SMAIDs such a medicación. Sonder of these event has been final or life horsestering (DRESS typically, although and exclusively, presents with fiver, rash, hypothese proposition of the state of the seed of the seed for the seed of the seed

5.11 Fetal Toxicity Premature Closure of Fetal Ductus Arteriosus

Parameter. Cheese of Establishma Adertosia.

And Oliver of MASA, Anchology melancaria, my programs somen at about 30 weeks and out and MASA. Anchology melancaria, my programs some and a programs of the first ductus arteriors at approximately the spectational per present of the first ductus arteriors at a special per programs of the first ductus arteriors at a special per programs of the spec

example, include limb contractures and delayed lung maturation. In some postmarketing cases of impaired neonatal renal function, invasive procedures such as exchange transfusion or dialysis were required.

translusion or dialysis were required. If HSADD returned in encourage the IRSADD returned in encourage the encourage that the e

patient Tissues with miscurcam has any sign of symptomic abundant, montain MAGIAS, including miscurcam has any sign of symptomic abundant conditions such as coopylation disorders or concomitant used of martins, other such as the coopylation disorders or concomitant used of martins, other such as structure or comparison requirate inholisms (SMRISI may known such as the sixth Montant these patients for sign of blooking lose Drug Interactions (TS). SIMISI has been such as the This planmancological author) of molicitian in reducting Inflammation and possibly fover, may dimension but sign of dispositists signs in districting Inflammation and possibly fover, may dimension but sign of dispositists signs in districting Inflammation and possibly fover, may dimension but signs of dispositists signs in districting Inflammation and possibly fover, may dimension but signs of passible signs in districting Inflammation and possibly fover, may dimension but signs of passible signs in districting Inflammation and possibly fover, may dimension but signs of passible signs in the signs in the signs of the signs of the Tissue signs of the participant of the signs of the s

6 ADVERSE REACTIONS

- 6 ADVISED EACTIONS

 The Tollwaye advisors exactions are discussed in greater datal in other sections of the studies; the production of the studies; and the studies of the studies; and the studi

R. 1. Clinical Trials: Experience

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 or another drug and may not reflect the rates observed in practice.

The product of the conduction of th Adults
Osteoarthritis and Rheumatoid Arthritis

Observatives and Renomined Articles

The meloscam Place of Trickel I and Renomined Articles

The meloscam Place of Trickel I and Renomined I and I and

trial. A 12-week multicenter, double blind, randomized trial was conducted in patients with consourch for this time or by to consourch as efficiency and select production with the time or by the consourch as efficiency and select production with the ware conducted any patients with homestand arthrists to compare the efficiency and salely of rendoction with pacieties. A select production will be patient to the patients of the patients are the patients and the control in 2.7% of the residuction treatment groups in 2.1% well described and achieve certified obstance that selection and selection and selection of the control of selection selection selections. The selection of the control of the patients are the selection of the control of the patients are the selection of the control of the patients are the selection of the control of the selection treatment groups in 10.2 % on the patients are the selection of the s

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

| | Placebo | Meloxicam 7.5 mg daily | Meloxicam 15 mg daily | Diclofenac 100 mg dail |
|---|---------|---------------------------|--------------------------|---------------------------|
| 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* | 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† | 2.5 | 2.6 | 0.6 | 2.0 |

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

| | Placebo | Meloxicam 7.5 mg daily | Meloxicam 15 mg daily |
|---|---------|------------------------|-----------------------|
| No. of Patients | 469 | 481 | 477 |
| Gastrointestinal Disorders | 14.1 | 18.9 | 16.8 |
| Abdominal pain NOS [®] | 0.6 | 2.9 | 2.3 |
| Dyspeptic sign and symptoms [†] | 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- | | | |
| pathogen class unspecified† | 4.1 | 7.0 | 6.5 |
| Musculoskeletal and Connective Tissue | | | |
| Disorders | | | |
| oint related signs and symptoms† | 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 |
| MedDRA preferred term: nausea, abdominal pain NOS, influenza-li | | | |

MedDRA high level term (preferred terms): dyspeptic signs and symptoms (dyspepsia, dyspepsia aggravated, enuctation, gastrointstimal intation), upper esspratory tract infections-pathogen unspecified (laryngists NOS, pharyngists NOS, sinusists NOS), joint nistated signs and symptoms (anthraiga, authraigla aggravated, joint creptation, joint effision, joint swelling)

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

| | 4 to 6 Weeks 0 | | | |
|------------------------|---------------------------|--------------------------|---------------------------|--------------------------|
| | 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 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 |
| Back Pain | 0.5 | 0.4 | 3.0 | 0.7 |
| Psychiatric | | | | |
| nsomnia | 0.4 | 0.0 | 3.6 | 1.6 |
| Respiratory | | | | |
| Coughing | 0.2 | 0.8 | 2.4 | 1.0 |
| Joper 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 | | | | |
| Micturition | | | | |
| frequency | 0.1 | 0.4 | 2.4 | 1.3 |
| Urinary tract | | T | 1 | |
| infection | 0.3 | 0.4 | 4.7 | 6.9 |

Higher doses of meloxicam (22.5 mg and greater) have been associated with an increased risk of serbuss Gallewest, therefore, the daily dose of meloxic an should not exceed 1.6 mg.
Pediatrics
Paciatrics and Polyanticular Course Inventor Resourced Arthritis (IBA)

Pacietticals and Edularitical Course Journal Ribourantial Entitle (III).

Three hundred and eight your explaints with prosecutional and optiviticular course JRA were exposed to miduscam with douce ranging from 0.12 to 0.15 mg/leg or day in the water exposed to miduscam with douce ranging from 0.12 to 0.15 mg/leg or day in the process of the process

genoer-specins supgroup erroct.

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

| | Table 3 Clinically Significant Drug Interactions with Melosicam |
|--|--|
| Body as a Whole | silergic reaction, face edema, fatigue, fever, hot flushes, |
| 1 - | mables, syncope, weight decrease, weight increase |
| Cardiovascular | angina pectoris, cardiac failure, hypertension, |
| | hypotension, myocardial infarction, vasculitis |
| Central and Peripheral Nervous System | Convulsions, parasthesis, tremor, vertigo |
| Gastrointestinal | collisis, dry mouth, duodenal ulicer, eructation, esophagitis, gastric ulicer, gastric ulicer, gastroescophageal reflux, gastrointestinal hemorrhagic deuodenal ulicer, hemorrhagic qastric ulicer, intestinal perforated duodenal ulicer, perforated duodenal ulicer, perforated duodenal ulicer, perforated duodenal ulicer, perforated dastric ulicer, stomatitis ulicerative |
| Heart Rate and Rhythm | arrhythmis, palpitation, tachycardia |
| Hematologic | eukopenia, purpura, thrombocytopenia |
| Liver and Biliary System | ALT increased, AST increased, bilirubinemia, GGT |
| | ncreaced, hepatitis |
| Metabolic and Nutritional | dehydration |
| Psychiatric | abnormal dreaming, anxiety, appetite Increased, |
| - | confusion, depression, nervousness, somnolence |
| Respiratory | asthma, bronchospasm, dyspnea |
| Skin and Appendages | alopecia, angioedema, bulious eruption, photosenstivity |
| | reaction, pruritus, sweating increased, urticaria |
| Special Senses | abnormal vision, conjunctivitis, taste perversion, tinnitus |
| Urinary System | albuminuria, BUN increased, creatinine increased, |
| | hometuris recal failure |

The following elevens: nearliers have been destribed during post approval or and monitorisms. Records the enterpolar are required within \$1.000 post approval or and uncertainties, it is not always possible to reliably estimate their Properties; or establish a country of the enterpolar and their post of their

7 DRUG INTERACTIONS
See Table 3 for clinically significant drug interactions with meloxicam. See also Warnings and Precautions (2.5, 5.6, 5.12) and Clinical Pharmacology (12.3).
Table 3 Clinically Significant Drug Interactions with Meloxicam

| | Meloxicam and anticoagulants such as warfarin have a |
|------------------|---|
| | synergistic effect on bleeding. The concomitant use of meloxicam and anticoagulants have an increased risk of serious bleeding compared to the use of either drug alone. |
| | meloxicam and anticoagulants have an increased risk of serious bleeding compared to the use of either drug alone |
| Clinical Impact: | |
| | hemostasis. Case-control and cohort epidemiological studies showed that concomitant use of drugs that interfere with |
| | snowed that concomitant use of drugs that interfere with serotonin reuptake and an NSAID may potentiate the risk of |
| | bleeding more than an NSAID alone. |
| | Monitor patients with concomitant use of meloxicam with |
| Intervention: | anticoagulants (e.g., warfarin), antiplatelet agents (e.g., aspirin), selective serotonin reuptake inhibitors (SSRIs), and serotonin |
| inter Peritori. | norepinephrine reuptake inhibitors (SNRIs) for signs of bleeding |
| Aspirin | [see Warnings and Precautions (5.12)]. |
| Aspirin | Controlled clinical studies showed that the concomitant use of |
| | NSAIDs and analgesic doses of aspirin does not produce any |
| Clinical Impact: | greater therapeutic effect than the use of NSAIDs alone. In a |
| unical impact: | 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)]. Concomitant use of meloxicam and low dose aspirin or analgesis |
| | concomitant use of meloxicam and low dose aspirin or analgesis doses of aspirin is not generally recommended because of the |
| Intervention: | doses of aspirin is not generally recommended because of the increased risk of bleeding (see Warnings and Precautions (5.12) |
| incervencon. |). Meloxicam is not a substitute for low dose aspirin for |
| | meioxicam is not a substitute for low dose aspirin for cardiovascular protection. |
| ACE Inhibitors | , Angiotensin Receptor Blockers, or Beta-Blockers |
| | NSAIDs may diminish the anthypertensive effect of angiotensin converting enzyme (ACE) inhibitors, angiotensin |
| | angiotensin converting enzyme (ACE) inhibitors, angiotensin receptor blockers (ARBs), or beta-blockers (including |
| | |
| Clinical Impact: | |
| | on diuretic 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. |
| | During concomitant use of meloxicam and ACE inhibitors, ARBs, or beta-blockers, monitor blood pressure to ensure that |
| | ARBs, or beta-blockers, monitor blood pressure to ensure that the desired blood pressure is obtained. |
| | |
| Intervention: | ARBs in patients who are elderly, volume-depleted, or have impaired renal function, monitor for signs of worsening renal |
| Intervention: | impaired renal function, monitor for signs of worsening renal |
| | function [see Warnings and Precautions (5.6)]. • When these drugs are administered concomitantly, patients |
| | should be adequately hydrated. Assess renal function at the |
| | beginning of the concomitant treatment and periodically thereafter. |
| Diuretics | |
| | Clinical studies, as well as post-marketing observations, showed |
| | that NSAIDs reduced the natriuretic effect of loop diuretics (e.g. |
| | furosemide) and thiazide diuretics in some patients. This effect has been attributed to the NSAID inhibition of renal prostaglandi |
| Clinical Impact: | synthesis. However, studies with furosemide agents and |
| | meloxicam have not demonstrated a reduction in natriuretic |
| | effect. Furosemide single and multiple dose pharmacodynamics and pharmacokinetics are not affected by multiple doses of |
| | meloxicam. |
| | During concomitant use of meloxicam with diuretics, observe |
| Intervention: | During concomitant use of meloxicam with diuretics, observe patients for signs of worsening renal function, in addition to assuring diuretic efficacy including antihypertensive effects [see |
| | Warnings and Precautions (5.6)]. |
| Lithium | |
| | NSAIDs have produced elevations in plasma lithium levels and reductions in renal lithium clearance. The mean minimum lithium |
| Clinical Impact: | concentration increased 15%, and the renal clearance decrease |
| Lanical Impact: | concentration increased 15%, and the renal clearance decrease by approximately 20%. This effect has been attributed to NSAID |
| | inhibition of renal prostaglandin synthesis [see Clinical Pharmacology (12.3)]. |
| | During concomitant use of meloxicam and lithium monitor |
| intervention: | During concomitant use of meloxicam and lithium, monitor patients for signs of lithium toxicity. |
| Methotrexate | |
| Clinical Impact: | Concomitant use of NSAIDs and methotrexate may increase the risk for methotrexate toxicity (e.g., neutropenia, |
| Ciriicar impact: | thrombocytopenia, renal dysfunction). |
| intervention: | During concomitant use of meloxicam and methotrexate. |
| | monitor patients for methotrexate toxicity. |
| Cyclosporine | Concomitant use of meloxicam and cyclosporine may increase |
| Clinical Impact: | cyclosporine's nephrotoxicity. |
| intervention: | During concomitant use of meloxicam and cyclosporine, monito |
| | patients for signs of worsening renal function. |
| NSAIDs and S | alicylates Concomitant use of meloxicam with other NSAIDs or salicylates |
| Clinical Impact: | (e.g., diflunisal, salsalate) increases the risk of GI toxicity, with |
| Ciriicar impact: | Ittle or no increase in efficacy [see Warnings and Precautions |
| | (5.2)]. |
| intervention: | The concomitant use of meloxicam with other NSAIDs or salicylates is not recommended. |
| Pemetrexed | |
| | Concomitant use of meloxicam and pemetrexed may increase |
| Clinical Impact: | the risk of pemetrexed-associated myelosuppression, renal, and GI toxicity (see the pemetrexed prescribing information). |
| | |
| | During concomitant use of meloxicam and pemetrexed, in patients with renal impairment whose creatinine clearance |
| | |
| | |
| | Patients taking meloxicam should interrupt dosing for at least five days before, the day of, and two days following pemetrexec |
| Intervention: | |
| Intervention: | |
| Intervention: | administration. In patients with creatinine clearance below 45 mL/min, the |
| intervention: | |

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy Risk Summary

Black Similary

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Only operative mixed beautiful and implainment of the mixed of the

Use of NSAIDs at about 20 weeks gestation or later in pregnancy has been associated with cases of fetal renal dysfunction leading to oligohydramnios, and in some cases, neonatal renal impairment.

neonatar renal impairment.

Data from observational studies regarding potential embryofetal risks of NSAID use in women in the first or second trimesters of pregnancy are inconclusive.

Data from observational studies regarding potential enhydred risks of NSAID use in women in this first is second trimeters for organizary are forcularly second regarding. In artisting reproduction studies, emboydread death was observed in rate and rate of 10.65 and 6.5-times from the maintain incommond human does Belloyil of individual to 10.65 and 6.5-times from the maintain incommond human does Belloyil of individual compresses with medicac and as or rail does equal-sector to 73-times the MRRIL in producing producing the common section of the common section of the common section of section produces and an experimental section of the common section of section produces and a section of the common section of the common section of section produces and decreased dispressing serviced as 0.05 times of section produces and section of the common section of section produces and section of the common section of section of the section producing section of section section of the section of section of section of the section of sec

eth and 15% to 20%, respectively.

[Plaid Considerations]

Febalishound Adverse Reactions

Anoid use of HAADEs in women able to 30 weeks gestation and later in pregnancy,

more influence, not make generative closure of the febal ductual

entresce, (see Table)

Oliphydramicos. Honoratid Renal Imparament:

In his Mich Dis necessary at ideas 270 weeks gestation or later in pregnancy, limit the use

to the bower effective does and shortest duration possible. If melociam treatment

to the bower effective does and shortest duration possible. If melociam treatment

places are the seed of the seed o

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.

studies, KSALDs, Including meloskoum while prostagilated synthesis, cause delayed parturation, and norse he hockered or laberth.

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Ries Summary.

There are no human data sociable on whether melonicam is present in human mile, or on the effects on breastfed infants, or on mile production. The developmental and health benefits of horselated pshoulb be considered along with the methor's clinical need for melosicam and any potential adverse effects on the breastfed infant from the memokscam or from the underlying material condition.

Table

Animal data

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

Females Based on the mechanism of action, the use of prostaglandin-mediated MSAIDs, including melascam, may delay or prevent rupture of ovarian folicities, which has been associated with reversible infertly is some women. Published animal studies have shown that with reversible infertly is some women. Published animal studies have shown that prostaglandin-mediated folicities regulared for ovalution. Small studies in women tracted with MSAID have also shown a reversible delay in ovalution. Consider withdrawd of MSAIDs, including melascizem in women who have difficulties conceiving or who are undeepong investigation of infertity.

8.4 Pediatric Use

The safety and effectiveness of meloxicam in pediatric JRA patients from 2 to 17 years of age has been evaluated in three clinical trisls [see Dosage and Administration (2.3), Adverse Reactions (6.1) and Clinical Studies (14.2)].

8.6 Hepatic Impairment
No dose adjustment is necessary in patients with mild to moderate hepatic impairment
No dose adjustment is necessary in patients with mild to moderate hepatic impairment
patients and the service part in reported in these not done adducted by a may occur use
meloscare with caution in patients with hepatic impairment [see Warnings and
meloscare with caution in patients with hepatic majorment [see Warnings and
meloscare of Circlar Patients (2007, 12.3)].

8.7 Renal impairment
No docage adjustment is necessary in patients with mild to moderate renal impairment.
No docage adjustment is necessary in patients with mild to moderate renal impairment members are not supported to the patients on hemodistysts, mildisciss with severe renal impairment is not recommended, in patients on hemodistysts, mildisciss mithod not exceed 7.5 mg per day. Metockam is not dislyrable (see Dosage and Antimistration (2.1) and official informations (12.3) in large Dosage and Antimistration (2.1) and official informations (12.3).

[see Bosage and Administration (2.1) and Christ Pharmacology (2.2)), IN OFFEDORAGE
Spragman Balance acutal SISSID overforsages have been byzickly initiated to lathage, program Balance acutal SISSID overforsages have been been promisely reversible with supportive care. Castroinstation bleeding has occurred, High preferration, scalar result with supportive care. Castroinstation bleeding has occurred, High preferration, scalar result failed, regardance programs, and come have excerned. Not were are late Marriages Manage patients with symptomized and supportive care following an HSAID overforsage, from a not support selform. For a supportive care following an HSAID overforsage gramm in stable, 10 2 years per lay of body weight is political patiently active gramman stable, 10 2 years per lay of body weight is political patiently active programs. The program of the programs of

11 DESCRIPTION

Motoclann is a nonsteroidal anti-inflammatory drug (MSAID). Each light yellow colored
motoclann table Loration 7.5 mg or 15 mg misolacam for oral administration.

Motoclann is chemically designated as 4-hydroxy2-archity44/5-methy42-2thacohy2-4t12-benostoklanni-2-chrostowanide-1,1-doise; The motoclary design is 351.4 its
empirical formula is C1,941,94(0.45) and it has the following structural formulas:

Nelockom II. a genetly effor useful precisely involvable in eatin, with higher solidary in decreased in remorp and solidary for the viriality's volvers in members. Mobic can have an apparent partition coefficient (log IP_{ings} = 0.1 in no-ctanoloculer part 7.4. Metoxicam has pika values of 1.3 and 4.2.

Molockom Tablets, USP is available as a tablet for or all administration containing 7.5 mg or 3 fing middlesses.

or 15 mg meloxicam.

The inactive ingredients in Meloxicam Tablets, USP include crospovidone, lactose mononlydrate, magnesium stearate, microcrystalline cellulose, povidone, sodium citrate dihydrate and yellow iron oxide (in 15 mg tablets only).

12 CLINICAL PHARMACOLOGY

Is a CLINICAL PHRAINACOLOGY

12.1 Mechanism of Action

Motorcam has avaignest, active ferrimentory, and aritypretic properties.

Motorcam has avaignest, active ferrimentory, and aritypretic properties.

The mechanism of a storo of Metorcams, like the der of hort (NSAIDs, is not completely understood but involves inhibition of cyclosovygenses (CDC-1 and CDC-2).

Motorcam is a potent whither of prostagations symbols in vitro. Neison device of the control of the control

12.3 Pharmacokinetics

13.3 Pharmacohients.

Miscontinia
The absolute bleavability of miscolutarn capsales was 89% following a single and dose of 30 mg company and m30 mg N tokas isjection. Following single intravelous dose, multiple and dose of 30 mg company and 10 mg company and 10 mg company and 10 mg company and 10 mg leavable company

Table 4 Single Dose and Steady-State Pharmacokinetic Parameters for Oral 7.5 mg and 15 mg Meloxicam (Mean and % CV)²

| | | Steady | | | | ie Dose |
|--|----------|--------------------------------|--|--|------------------------------|--------------------------------------|
| Pharmacokinetic Parameters (%CV) | | male | Elderly males (Fed) ² | Elderly females (Fed) ² | Renal failure (Fasted) | Hepatic insufficiency (Fasted) |
| | | 7.5 mg ³ tablets | 15mg capsules | 15mg capsules | 15mg capsules | 15mg 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 _{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) |
| | | 14.7 (32) | 15 (42) | 10 (30) | 26 (44) | 14 (29) |
| | | | | | | |

Find and Anticet Effects.

Administration of medication operates following a high fat broadest [75 g of fist) resulted in more past drug levels (i.e., Cross) being increased by agreement by 27% which the value of the control of the

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dose range, Pleanns Liberauce ranges from 7 to 9 mil. rans.

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Bleder mite (165 years of age) ein hibited metaksicam plasma concentrations and
status)-status pharma-cokinetics similar to young mails. Elderly fermites (165 years of
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concentrations in the ordinery femiliae (164 years) on the compared to younger femiliae
concentrations in the ordinery femiliae. See device over profess are compared to to both
cellerly potent populations. A strainfer fee faction was found in identify femile patients.

In comparation to deliver than plasmers.

Young females exhibited slightly lower plasma concentrations relative to young males. After single doses of 7.5 mg Melacus am Tablets, 195°, the mean elimination half-life was standard to the control of the control

Renal Imperioriest

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Drug InteractionStudies

Does InteractionStudies.

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intesting found no protein binding day interaction between disjoint and metoccam. Lettlam in a study contacted in healthy subject, mean per does little mocentration and AUI over the creations of the subjects receiving littlem does ranging from BUI or subjects receiving littlem does ranging from BUI or subjects receiving littlem and the subjects or receiving littlem and little little

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

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.

Impairment of Fertilty

Imparment of Fertiley

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

14 CLINICAL STUDIES

14 CLINICAL STUDIES

The use of relabelism for the treatment of the signs and symptoms of estocartrists of the tense and type and symptoms of estocartrists of the tense and type sendanting in 12-week, double bird, controlled trial. Midisciscan endpoints were investigated in 91 and secsionary, patter global assessment, patter global patterns, patt

pictories. The sure of management of signs and symptomic of extraorthrites are flower of microscount for the management of signs and symptomic of extraorthrites are seed to form the signs of the medical and 31 mights, which companies the princes are 30 miles and 41 miles of 12 miles and 12 miles and

14.2 yourself Resumed of Arthritis (JRM) Paradistriction and Polypriticular Course
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Meloxicam Tablets, USP 7.5 mg is available as light yellow colored, oval shaped uncoated tablet engraved 5 160 on one side and plain on other side.

Meloxicam Tablets, USP 15 mg is available as yelow colored, oval shaped uncoated tablet engraved 5 161 on one side and plain on other side.

Meloxicam Tablets, USP 7.5mg are available as follows Bottles of 100 NDC: 64380-715-06 Bottles of 500 NDC: 64380-715-07

Bottles of 500 NDC: 64380-715-07

Meloxicam Tablets, USP 15mg are available as follows:
Bottles of 100 NDC: 64380-716-06

Bottles of 500 NDC: 64380-716-07

Bottles of \$50 NDC: 64380-716-07

Stornage

Storn at 25°C (77°F); excursions permitted to 15° to 30° C (59°to 86°F) [see USP

Controlod Room Temperature! Keep Neboxicam Tablets, USP in a dry place.

Disperse tablets in sight container.

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

17 PATIENT COUNSELING INFORMATION Advise the patient to read the FDA-approved patient labeling (Medication Guide) that accompanies each prescription dispensed.

Inform patients, families or their caregivers of the following information before initiating therapy with an NSAID and periodically during the course of ongoing therapy.

Transcription as manutal and personally during the course of ongoing therapy. Cardiouscular Transcription (Cardiouscular Transcriptions) and cardiouscular thrombotic events, Advice patients to be alert for the symptoms of cardiouscular thrombotic events, including cheet spin, shortness of breath, weakness, or stairing of speech, and to report any of these symptoms to their healthcare provider immediately [see Warnings and Procaddors (6.73)]. and Precautions (5.11).

Gastrointestral Bleeding, Ulteration, and Perforation

Advice patients to proof virginame of diserations and bleeding, including epigastric
pair, dyseppsis, melans, and hematemesis to their healthcare provider. In the setting
concomitant use of the wides are pin'n for cardiac prophysiks, inform patients of the
increased risk for the signs and symptoms of GI bleeding [see Warnings and Precaudic
(5.21).

Hepatotoxicity

<u>Historicack</u>:
Intriducing the warring signs and symptoms of hepatotoxicky (e.g., nauces, flutgue, kithrapy, distribue, purious, jundice, right upper quadret involutions, see Intriducing the history, few Warrings and Percautions (5.73).

<u>Historical Research Continued to Percautions (5.73).</u>

Hast Tables and Edema
Africa patient to be airst for the symptomic of congestive heart faither including shortness of brastly, unexplained weight gain, or edoms and to contact their healthcare provided in 2014. The provided is 2014. The provided in 2014 the provided of 2014 they provided 2014 the

shortness of breath, unexplained weight gain, or ediems and to contact their healthcare provider it such preference occur (see Winning and Precadation (s. g., difficulty Installation).

Establishment: Beautism.

Establishment: Beautism of the provider of the second (s. g., difficulty breathing) seeking of the fine or throad). Instruct patients to seek immediate energency help if these occur (see Contrandications (d. g. and Precadation (S. 7).

Advise patients to stop taking Neilorian Tablets, USP immediately if they develop any strangs and Precadation (S. 7).

Estimate Establishment (S. 9.8, 10.0) **. The Neilorian providers is stoom in procision later Warrings and Precadation (S. 9.8, 10.0) **. The Neilorian Providers is stoom in procision later (S. 10.0) **. The Neilorian Providers is stoom in procision later (S. 10.0) **. The Neilorian Providers (S.

Use of NSAIDs and Low-Dose Aspirin
Inform patients not to use low-dose aspirin concomitantly with Meloxicam Tablets, USP
until they talk to their healthcare provider [see Drug Interactions (7)].

Strides Pharma Science Limited

Bengaluru-562106, India



SPL MEDGUIDE

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aarmea, gas. heartburn, nausea, vomting, and dizziness.

Set emergency help right away if you get any of the following symptoms:
shortness of breath or trouble breathing
c chest pain
weakness in one part or side of your body
staircraft speech
swelling is the face or throat

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

ou get any of the Tollowing symptoms:
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If you seculd it now reformate about MSAID, tak why your healthcare should MSAID that it written for health professional, for a propert SIMPERTE ADVISES REACTION, contact STAIRS Pharma for, at 1.377-244-9315 or go to symmuticides.com or contact FDA at 1-007-TB-118ER or work flagmentaries.

PACKAGE LABEL.PRINCIPAL DISPLAY PANEL 100 tablets Neloxicam Tablets, USP 7.5mg Rx Only Strides Pharma Inc.



500 tablets Meloxicam Tablets, USP 7.5mg Rx Only Strides Pharma Inc.



100 tablets Meloxicam Tablets, USP 15mg Rx Only Strides Pharma Inc.

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500 tablets Meloxicam Tablets, USP 15 mg Rx Only Strides Pharma Inc.

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| p | roduct Infor | mation | | | | | | |
| P | roduct Type | | HUMAN PRE | S CRIPTION DRUG | Item C | ode (Source) | NOC | 64280-716 |
| R | oute of Admin | istration | ORAL | | | | | |
| A | ctive Ingredi | ent/Active | Moiety | | | | | |
| | | Ingre | edient Nam | 10 | | Basis of St | rength | |
| H | ELOXICAM (LIVI) | VG2QF83CGL) | (MELOXICAM | - LINIE VG2QFR3CGL) | | MELOXICAM | | 15 mg |
| li | nactive Ingre | dients | | | | | | |
| | | | Ingredie | ent Name | | | | Strength |
| c | ELLULOSE, MICE ROSPOVIDONE (ERRIC OXIDE YEI | UNI: 25783065 | 561) X43902MRT) | | | | | |
| H | AGNESIUM STEA | RATE (UNIX 7) | 009796(20) | | | | | |
| H | | RATE (UNIX 7) NE U725QUEZ | 009796(20) 12X) | | | | | |
| P | AGNESIUM STEA | RATE (UNII: 7) NE: U725QW73 (UNII: 1Q73Q2) | 009796(30) (2X) (ULR) | | | | | |
| P | AGNESIUM STEA DVIDONE K30 (U DDIUM CITRATE | RATE (UNIT 7) NE U725QUEY (UNIT 1Q73Q2) Incterístics | 009796(30) (2X) (ULR) | Score | | | no score | |
| H P S | AGNESIUM STEA DVIDONE K20 (U DDIUM CITRATE FORGUET Chara plor hape | RATE (UNIT 7) NE U725QUEY (UNIT 1Q73Q2) Incterístics | 009796(30) (2X) (ULR) | Score Size | | | lmm | |
| M P S P C S F | AGNESIUM STEA DVIDONE K30 (U DDIUM CITRATE TODUCT Chara Dior | RATE (UNII: 7) NE: U725QWP2 (UNII: 1Q72Q2) scteristics YSL | 009796(30) (2X) (ULR) | Score | | | | |
| M P S P C S F C | AGNESIUM STEA DVIDONE K20 (U DDIUM CITRATE FORGUET Chara plor hape lavor | RATE (UNII: 7) NE: U725QWP2 (UNII: 1Q72Q2) scteristics YSL | 009796(30) (2X) (ULR) | Score Size | | | 3mm 5161 | |
| H P S P C S F C | AGNESIUM STEA DVIDONE K30 (U DDIUM CITRATE roduct Chari olor hape lavor pontains | RATE (UNII: 7) NII: U725Quera (UNII: 1073QU acterístics VIII OW | 009796(30) (2X) (ULR) | Score Stre Imprint Code | Mark | eting Start Date | S161 Mark | eting En |
| H P S P C S F C | AGNESSIM STEA DVNDONE K20 (III) DDNUM CITRATE roduct Chara Discr hape lavor ontains ackaging Rem Code NDC 66280-716-06 | RAYE (JUNE 7) NE U725QWP 2 JUNE 1073QWP 2 JUNE 1073QWP 3 ACTORISTICS VEL OVI Pa 100 in 1 80T Product | 009796(20) (220) (JUR) (LLOW AL. SICKAGE Des | Score Stre Imprint Code | Mark | Date | S161 Mark | eting End Date |
| H P S P C S F C | AGNESSIM STEA DVNDONE K20 (III) DDNUM CITRATE roduct Chara Discr hape lavor ontains ackaging Rem Code NDC 66280-716-06 | RAYE (JUNE 7) NE U725QWP 2 JUNE 1073QWP 2 JUNE 1073QWP 3 ACTORISTICS VEL OVI Pa 100 in 1 80T Product | 009796(20) (220) (JUR) (LLOW AL. SICKAGE Des | Score Stre Imprint Code | Mark | Date 09 | S161 Mark | eting En |
| MAS POSEC P # 1 2 | AGNESSUM STEATONDOME KID (UDDOWN CITRATE roduct Characolor polar avor potatins ackaging Rem Code NCC 66280-716- 07 | RAYE (UNII: 7) WE U725QWP 2 (UNII: 10720QWP 2 (UNII: 10720QWP 1072 | COSTRECION 1220 1220 1220 1220 1220 1220 1220 122 | Score Stre Imprint Code | Mark 05/14/20 | Date 09 | S161 Mark | eting En |
| MAS POSEC P # 1 2 | AGNESSIN STEE ON THE MODEL OF THE MODE | PPa loo in 1 sor Product | COSTRECION 1230 1240 1240 1240 1240 1240 1240 1240 124 | Score Size Imprint Code imprint Code icription ict a Combination ict a Combination | Mark 05/24/20 05/24/20 | Date 09 | Imm S161 Mark | Date |
| POSEC P # 1 2 | AGNESSUM STEATONDOME KID (UDDOWN CITRATE roduct Characolor polar avor potatins ackaging Rem Code NCC 66280-716- 07 | PPa loo in 1 sor Product | 009796(20) 222) juR9 iLLOW M. inckage Den TLE; Type 0: n TLE; Type 0: n | Score Stre Imprint Code Imprint Code Interprint Code Interpritation Interprint Code Interprint Code Interprint Code Interprint | Mark 05/24/20 05/24/20 | Date 09 09 rketing Start Date | Imm S161 Mark | eting Enc Date |

Labeler - Sintern Names Genera Limbel (\$0272814)

Registrant - Sintern Names Genera Limbel (\$0272814)

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