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BUDD LARNER, P.C. 150 JOHN F. KENNEDY PARKWAY



Moverse Reactions Excluding Non-Drug Related	Treatment Group					
	DynaCirc CR [®] (Isradipine) (N=422)	Ptecebo (N=188)				
Edema	15.2%	2.2%				
Headache	13 0%	12.4%				
Dizziness	4 7%	2 7%				
Fa tigue	4 3%	2.2%				
Abdominal Discomfort	2 8%	0 5%				
Mushing	19%	0.5%				
Constipation	1.7%	0.0%				
Palpitations	1 2%	0 0%				
Kausea	1.2%	1.6%				
Abdominal Distention	1 2%	0 0%				

The following adverse experiences were reported in 0.5%—1.0% or less of DynaCirc CR® (isradipine) or immediate-release DynaCirc® (isradipine) treated patients in hyportensive studies, or were noted in postmarketing experience with immediate-release DynaCirc® (isradipine) Capsules More serious events are shown in italics. The relationship of these adverse experiences to isradipine administration is

BKIN Pruritus, urticaria, angioedema
MUSCULOSKELETAL, backache/pain, joint pain, neck

muscoliosaeletale. Backachepain, joint pain, neck pain/sorestif, legs achepain, cramps of legs/feet RESPIRATORY: Dyspnea, nasal congestion, cough. CARDIOVASCULAR: Epistaxis, tachycardia, cheat pain, shortness of breath, hypotension, syncope, atrial or ventric-ular fibrillation, myocardial infarction, heart failure GASTROINTESTINAL: Diarrhea, vomiting, appetite in-

ed or decreased UROGENITAL Pollakiuria, impotence, dysuria, nocturia

CENTRAL NERVOUS Drowsiness, insomnia, lethargy, servousness, libido decrease/frigidity, impotence, depression, paresthesia (which includes numbress and tingling),

AUTONOMIC Dry mouth, hyperhidrosis, visual distur-

MISCELLANEOUS Weight gain, throat discomfort, drug

hver, leukopenia, elevated liver function tests. No gastrointestinal bleeding has been reported in clinical trais with DynaCirc CR® (isradipine) Controlled Release Tablets

In a long-term (one-year) DynaCirc CR[®] (isradipine) open-label, hypertension trial, the adverse events reported were generally the same as those seen in the short-term placebo-controlled studies. About 6% of DynaCirc CR^{Φ} (isradipine) treated patients discontinued the long-term trial due to ad-

With immediate-release DynaCirc® (isradipine) Capsules, most of the adverse experiences were transient, mild, and related to vasodilatory effects. The following table shows the most common adverse events reported in U.S. clinical trials for immediate-release DynaCirc (isradipine) Capsules, volunteered or elicited, and considered by the investigator to be at least possibly drug related.

|See second table at top of previous page| |In open-label, long-term studies of up to two years in dura-tion with immediate-release DynaCirc® (isradipine) Capsules, the adverse experiences reported were generally the same as those reported in the short-term controlled trials. The overall frequencies of these adverse events were slightly higher in the long-term than in the controlled stud-ies, but in the controlled studies most adverse reactions were mild and transient.

OVERDOSAGE

Although there is no well documented experience with DynaCirc® (isradipine) overdosage, available data suggest that, as with other dihydropyridines, gross overdosage would result in excessive peripheral vasodilation with subsequent marked and probably prolonged systemic hypotension. Clinically significant hypotension overdosage calls for active cardiovascular support including monitoring of cardiac and respiratory function, elevation of lower extremities and attention to circulating fluid volume and urine out A vasoconstrictor (such as epinephrine, norepinephrine or levarterenol) may be helpful in restoring vascular tone and blood pressure, provided that there is no contraindica-

and noted pressince provided matchere is no commanda-tion to its use Since uradipine is highly protein bound, di-signsis is not likely to be of benefit.

Significant lethality was observed in mice given oral doses of over 200 mg/kg and rabbits given about 50 mg/kg of teradipine. Rats tolerated doses of over 2000 mg/kg without effects on survival.

DOSAGE AND ADMINISTRATION

The dosage of DynaCirc CR® (isradipine) Controlled Release Tablets should be individualized. The recommended initial dose of DynaCirc CR® (isradipine) is 5 mg once-daily as monotherapy or in combination with a thiazide diuretic An anthypertensive response usually occurs within 2 hours, with the peak antihypertensive response occurring 8-10 hours post-dose; blood pressure reduction is maintained for at least 24 hours following drug administration. If necessary, the dose may be adjusted in increments of 5 mg at 2-4 week intervals up to a maximum dose of 20 mg/day Adverse experiences are increased in frequency above 10 mg/

DynaCirc CR® (isradipine) Controlled Release Tablets should be swallowed whole and should not be bitten or di-

The bioavailability (increased AUC) of immediate-release The bioavailability (increased AUC) of immediate-release DynaCirc® (isradipine) is increased in elderly patients (above 65 years of age), patients with hepatic functional impairment, and patients with mild renal impairment. Ordinarily, a starting dose of DynaCirc CR® (isradipine) 5 mg once-daily should be used in these patients

HOW SUPPLIED

(isradipine) Controlled Release Tablets:

5 mg: A light pink, round, standard biconvex and film coated tablet Printing is in red with "DynaCirc CR" in a semicircle with "5" centered below the semicircle.

Bottles of 30 controlled release tablets (NDC 65726-235-10) A beige, round, standard biconvex and film coated 10 mg: A beige, round, standard biconvex and unit contestablet Printing is in red with "DynaCirc CR" in a semicircle with "10" centered below the semi-circle.
Bottles of 30 controlled release tablets (NDC 65726-236-10)

Store and Dispense:

Below 86°F (30°C) in a tight container, protected from moisture and humidity

Rx only Revised: June, 2007 Distributed by. Reliant Pharmaceuticals, Inc Liberty Corner, NJ 07938

Address Medical inquiries to: Reliant Medical Inquiries

2655 Meridian Parkway Durham, NC 27713-2203 or Call: 877-311-7515 © 2007 Reliant Pharmaceuticals, Inc. 22352702

2002700 Shown in Product Identification Guide, page 329

LOVAZA™ lŏ-vā-zā]

(omega-3-acid ethyl esters)

Lovaza, a lipid-regulating agent, is supplied as a liquid-filled gel capsule for oral administration. Each one gram capsule of Lovaza (omega-3-acid ethyl esters) contains least 900 mg of the ethyl esters of omega-3 fatty acids These are predominantly a combination of ethyl esters of eicosapentaenoic acid (EPA - approximately 465 mg) and docosahexaenoic acid (DHA - approximately 375 mg). The structural formula of EPA ethyl ester is:

The empirical formula of EPA ethyl ester is $\rm C_{22}H_{34}O_2$, and the molecular weight of EPA ethyl ester is 330 51. The structural formula of DRA ethyl ester is

The empirical formula of DHA ethyl ester is $C_{24}H_{36}O_2$, and the molecular weight of DHA ethyl ester is 356.56. Lovesa capsules also contain the following mactive ingredients: 4 mg a-tocopherol (in a carrier of partially hydroge-nated vegetable oils including soybean oil), and gelatin, glyserol, and purified water (components of the capsule

CLINICAL PHARMACOLOGY

Mechanism of Action: The mechanism of action of Lovaza is not completely understood Potential mechanisms of action include inhibition of acyl CoA.1,2-diacylglycerol acyltransferase, increased mitochondrial and peroxisomal 8-oxidation in the liver, decreased lipogenisis in the liver, and increased plasma lipo-protein lipase activity. Lovaza may reduce the synthesis of triglycerdes (TGs) in the liver because EPA and DHA are poor substrates for the enzymes responsible for TG synthe sis, and EPA and DHA inhibit esterification of other fatty

acius Pharmacokinetic and Bioavailability Studies

In healthy volunteers and in patients with hypertriglyceri-demia (HTG), EPA and DHA were absorbed when administered as ethyl esters orally. Omega-3-acids administered as ethyl esters (Lovaza) induced significant, dose-dependent increases in serum phospholipid EPA content, though in-creases in DHA content were less marked and not dose-dependent when administered as ethyl esters. Uptake of terpatient when administered as cityl eachs of perfect of EPA and DHA into serum phospholipids in subjects treated with Lovaza was independent of age (<49 years vs ≥49 years) Females tended to have more uptake of EPA into serum phospholipids than males Pharmacokinetic data on Lovaza in children are not available.

Drug Interactions

Cytochrome P450-Dependent Monooxygenese Activities: The effect of a mixture of free fatty acids (FFA), EPA/DHA and their FFA-albumin conjugate on cytochrome P450dependent monooxygenase activities was assessed in human her microsomes. At the 23 µM concentration, FFA resulted in a less than 32% inhibition of CYP1A2, 2A6, 2C9, 2C19, 2D6, 2E1, and 3A. At the 23 µM concentration, the FFA-albumin conjugate resulted in a less than 20% inhibition of CYP2A6, 2C19, 2D6, and 3A, with a 68% inhibition being seen for CYP2E1. Since the free forms of the EPA and DHA are undetectable in the circulation (<1 µM), clinically significant drug-drug interactions due to inhibition of P450 mediated metabolism EPA/DHA combinations are not ex-

CLINICAL STUDIES

R

High Triglycerides: Add-on to HMG-CoA reductase inhibitor

The effects of Lovaza 4 g per day as add-on therapy to treatment with simvastatin were evaluated in a randomized, placebo-controlled, double-blind, parallel-group study of 254 adult patients (122 on Lovaza and 132 on placebo) with persistent high triglycerides (200-499 mg/dL) despite simva-statin therapy (Table 1). Patients were treated with openlabel sinvastatin 40 mg per day for 8 weeks prior to randomization to control their LDL-C to no greater than 10% above NCEP ATP III goal and remained on this dose The above NUEP AIP III goal and remained on this dose throughout the study. Following the 8 weeks of open-label treatment with sunvastatin, patients were randomized to either Lovaza 4 g, per day or placebo for an additional 8 weeks with sinvastatin co-therapy. The median baseline triglyceride and LDL-C levels in these patients were triglyceride and LDL-C levels in these patients were 268 mg/dL and 89 mg/dL, respectively. Median baseline non-HDL-C and HDL-C levels were 138 mg/dL and 45 mg/

The changes in the major inpoprotein lipid parameters for the Lovaza plus simvastatin and the placebo plus simva-statin groups are shown in Table 1.

(See table I below)

Lovaza 4 g per day significantly reduced non-HDL-C, TG, TC, VLDL-C, and Apo-B levels and increased HDL-C and LDL-C from baseline relative to placebo

Very High Triglycerides: Monotherapy

The effects of Lovaza 4 g per day were as domtzed, placebo-controlled, double-blind, parallel-group studies of 84 adult patients (42 on Lovaza, 42 on placebo) with very high triglyceride levels (Table 2). Patients whose baseline triglyceride levels were between 500 and 2000 mg/dL were enrolled in these two studies of 6 and 16

Continued on next page

Table 1: Response to the Addition of LOVAZA 4 g per day to On-going Sinvastatin 40 mg per day Therapy in Patients with High Triglycerides (200 to 499 mg/dL)

Parameter	LOV	LOVAZA + Simvastatin		Placebo + Simvestatin N=132				
	BL	EOT	Median % Change	B1.	EOT	Median % Change	Difference	P-Value
Non-HDL-C	137	123	-9.0	141	134	-2 2	-68	<0 0001
TG	268	182	-29.5	271	260	-63	-23.2	<0 0001
TC	184	172	-48	184	178	-1 7	-3 1	<0.05
VLDL-C	52	37	-27 5	52	49	-7.2	-20.3	<0.05
Apo-B	86	80	-4 2	87	85	-1.9	-2.3	<0.05
HDL-C	46	48	+3 4	43	44	-1 2	+4 6	<0.05
LDL-C	91	88	+0.7	88	85	-28	+3.5	=0 05

BL = Baseline (mg/dL); EOT = End of Treatment (mg/dL), Median % Change = Median Percent Change from Baseline, Difference = LOVAZA Median % Change - Placebo Median % Change



Lovaza-Cont.

weeks duration. The median triglyceride and LDL-C levels in these patients were 792 mg/dL and 100 mg/dL, respectively Median HDL-C level was 23 0 mg/dL The changes in the major lipoprotein lipid parameters for the Lovaza and placebo groups are shown in Table 2

Table 2: Median Baseline and Percent Change Fron Baseline in Lipid Parameters in Patients with Very High TG Levels (≥500 mg/dL)

, a						
Parameter		LOVAZA N=42		lacebo N=42		
	BL	% Change	BL	% Change	Difference	
TG	816	-44.9	788	+6 7	-51 6	
Non-HDL-C	271	-138	292	-36	-10.2	
TC	296	-97	314	-17	-8.0	
VLDL-C	175	-417	175	-09	-40 8	
HDL-C	22	+91	24	0.0	+91	
LDL-C	89	+44 5	108	-48	+49.3	

BL = Baseline (mg/dL); % Chg = Median Percent Change from Baseline, Difference = Lovaza Median % change Placebo Median % Change

Lovaza 4 g per day reduced median TG, VLDL-C, and non-HDL-C levels and increased median HDL-C from baseline relative to placebo Lovaza treatment to reduce very high TG levels may result in elevations in LDL-C and non-HDL-C in some individuals Patients should be monitored to ensure that the LDL-C level does not increase exces-

sively

The effect of Lovaza on the risk of pancreatitis in patients with very high TG levels has not been evaluated

The effect of Lovaza on cardiovascular mortality and morbidity in patients with elevated TG levels has not been

INDICATIONS AND USAGE

Very High Triglycerides
Lovaza is indicated as an adjunct to diet to reduce triglyceride (TG) levels in adult patients with very high (≥500 mg/ dL) triglyceride levels.

Lisane Considerations

In individuals with hypertriglyceridemia (HTG), excess body weight and excess alcohol intake may be important contributing factors and should be addressed before initiat-ing any drug therapy Physical exercise can be an important ancillary measure. Diseases contributory to hyperlipidemia, (such as hypothyroidism or diabetes mellitus) should be looked for and adequately treated. Estrogen therapy, thiszide diuretics, and beta blockers are sometimes asse with massive rises in plasma TG levels. In such cases, discontinuation of the specific etiologic agent, if medically in-dicated, may obviate the need for specific drug therapy for

The use of lipid-regulating agents should be considered only when reasonable attempts have been made to obtain satisfactory results with non-drug methods. If the decision is made to use lipid-regulating agents, the patient should be advised that use of lipid-regulating agents does not reduce the importance of adhering to diet (See PRECAUTIONS).

CONTRAINDICATIONS

Lovaza is contraindicated in patients who exhibit hypersensitivity to any component of this medication

PRECAUTIONS

Initial Therapy: Laboratory studies should be performed to ascertain that the patient's TG levels are consistently abnormal before instituting Lovaza therapy. Every attempt should be made to control serum TG levels with appropriate duet, exercise, weight loss in overweight patients, and control of any medical problems (such as diabetes mellitus and hypothyroidism) that may be contributing to the patient's TG abnormalities. Medications known to exacerbate HTG (such as beta blockers, thiazides, and estrogens) should discontinued or changed, if possible, before considering TGlowering drug therapy.

Continued Therapy: Laboratory studies should be per-formed periodically to measure the patient's TG levels during Lovaza therapy. Lovaza therapy should be withdrawn in patients who do not have an adequate response after 2 months of treatment.

Information for Patients:

Lovara should be used with caution in patients with known sensitivity or allergy to fish. Patients should be advised that use of lipid-regulating agents does not reduce the imporering to diet.

Laboratory Tests

In some patients, increases in alanine aminotransferase (ALT) levels without a concurrent increase in aspartate aminotransferase (AST) levels were observed. Alanine aminotransferase levels should be monitored periodically during Lovaza therapy

In some patients, Lovaza increased low-density hoporotein cholesterol (LDL-C) levels. As with any lipid-regulating product, LDL-C levels should be monitored periodically during Lovaza therapy

Drug Interactions.

Anticoagulants: Some studies with omega-3-acids demonstrated prolongation of bleeding time. The prolongation of bleeding time reported in these studies has not exceeded normal limits and did not produce clinically significant bleeding episodes Clinical studies have not been done to thoroughly examine the effect of Lovaza and concomitant anticoagulants Patients receiving treatment with both Lovaza and anticoagulants should be monitored periodi-

HMG-CoA reductase inhibitors: In a 14-day study of 24 healthy adult subjects, daily co-administration of simva-statin 80 mg with Lovaza 4 g did not affect the extent (AUC) or rate (Cmax) of exposure to simvastatin or the major active metabolite, beta-hydroxy simvastatin at steady state

Cytochrome P450-Dependent Monooxygenese Activities Omega-3-fatty acid containing products have been shown to increase hepatic concentrations of cytochrome P450 and activities of certain P450 enzymes in rats. The potential of Lovaza to induce P450 activities in humans has not been

Carcinogenesis, Mutagenesis, Impairment of Fertilit

In a rat carcinogenicity study with oral gavage doses of 100. 600, 2000 mg/kg/day by oral gavage, males were treated with omega-3-acid ethyl esters for 101 weeks and females for 89 weeks without an increased incidence of tumors (up to 5 times human systemic exposures following an oral dose of 4 g/day based on a body surface area comparison) Stan-dard lifetime carcinogenicity bioassays were not conducted

Omega-3-acid ethyl esters were not mutagenic or clastogenic with or without metabolic activation in the bacterial mutagenesis (Ames) test with Salmonella typhimurium and Escherichia coli or in the chromosomal aberration assay in Chinese hamster V79 lung cells or human lymphocytes Omega-3-acid ethyl esters were negative in the in vivo e micronucleus assav

In a rat fertility study with oral gavage doses of 100, 600, 2000 mg/kg/day, males were treated for 10 weeks prior to mating and females were treated for 2 weeks prior to and maning and remaies were treated for 2 weeks prior to and throughout mating, gestation and factation. No adverse effect on fertility was observed at 2000 mg/kg/day (5 times human systemic exposure following an oral dose of 4 g/day based on a body surface area comparison)

Pregnancy Category C:
There are no adequate and well-controlled studies in preg nant women It is unknown whether Lovaza can cause fetal harm when administered to a pregnant woman or can affect reproductive capacity. Loyaga should be used during cy only if the potential benefit justifies the potential risk

Omega-3-acid ethyl esters have been shown to have an e bryocidal effect in pregnant rats when given in doses resulting in exposures 7 times the recommended human dose of 4 g/day based on a body surface area compariso

In female rats given oral gavage doses of 100, 600, 2000 mg/ kg/day beginning two weeks prior to mating and continuing through gestation and lactation, no adverse effects were observed in the high dose group (5 times human systemic ex-posure following an oral dose of 4 g/day based on body sur-face area comparison)

In pregnant rats given oral gavage doses of 1000, 3000, 6000 mg/kg/day from gestation day 6 through 15, no adverse effects were observed (14 times human systemic exposure following an oral dose of 4 g/day based on a body surface area comparison)

Increment rats given oral gavage doses of 100, 600, 2000 mg/kg/day from gestation day 14 through lactation day 21, no adverse effects were seen at 2000 mg/kg/day (5 s the human systemic exposure following an oral of 4 g/day based on a body surface area comparison). How or a given tesset on a body surrance areas comparison. How-ever, decreased live births (20% reduction) and decreased surrival to postnatal day 4 (40% reduction) were observed in a dose-ranging study using higher doses of 3000 mg/kg/ day (7 times the human systemic exposure following an oral lose of 4 g/day based on a body surface area compara

In pregnant rabbits given oral gavage doses of 375, 750, 1500 mg/kg/day from gestation day 7 through 19, no findings were observed in the fetuses in groups given 375 mg/ kg/day (2 times human systemic exposure following an oral lose of 4 g/day based on a body surface area comparison) However, at higher doses, evidence of maternal toxicity was observed (4 times human systemic exposure following an oral dose of 4 g/day based on a body surface area

Nursing Mothers

It is not known whether omega-3-acid ethyl esters are excreted in human milk Because many drugs are excreted in human milk, caution should be exercised when Lovaza is administered to a woman who is breastfeeding

Safety and effectiveness in pediatric patients under 18 years of age have not been established

Geriatric Use

A limited number of patients over 65 years of age were enrolled in the clinical studies. Safety and efficacy findings in subjects over 60 years of age did not appear to differ from those of subjects less than 60 years of age

ADVERSE REACTIONS

Treatment-emergent adverse events reported in at least 🧦 of patients treated with Lovaza 4 g per day or placebeing 8 randomized, placebe-controlled, double-blind, parallegroup studies for HTG are listed in Table 3 Adverse events. led to discontinuation of treatment in 3.5% of national treated with Lovaza and 2 6% of patients treated with

Table 3: Adverse Events in Randomized,
Placebo-Controlled, Double-Blind, Parallel-Group Studies
for Very High TG Levels (≥ 500 mg/dL) that Used LOVAZA 4 g per Day

BODY SYSTEM Adverse Event	LOVAZA (N = 226) n %		Placebe* (N = 236) n %	
Subjects with at least 1 adverse event	80	35 4	63	27.4
Body as a whole Back pain Flu syndrome Infection Pain	5 8 '10 4	2.2 3.5 4.4 1.8	3 3 5 3	1.0
Cardiovascular Angina pectoris	3	13	2	0.0
Digestive Dyspepsia Eructation	7 11	3 1 4 9	6	1.F 2.1
Skin Rash	4	18	1 .	94
Special senses Taste perversion	6	2.7	0	9.6

Adverse events were coded using COSTART, version \$5. Subjects were counted only once for each body system and for each preferred term

Additional adverse events reported by 1 or more patients from 22 clinical studies for HTG are listed below BODY AS A WHOLE Enlarged abdomen, asthenia, odor, chest pain, chills, suicide, fever, generalized fungal infection, malaise, neck pain, neoplasm, rh

arthritis, and sudden death
CARDIOVASCULAR SYSTEM. Arrhythmia, bypass gery, cardiac arrest, hyperlipemia, hypertension, migrown myocardial infarct, myocardial ischemia, occlusion, periodical infarct, myocardial ischemia, occlusion, migrown myocardial infarct, myocardial ischemia, occlusion, migrown myocardial infarct, myocardial ischemia, occlusion, migrown myocardial infarct, myocardial ischemia, occlusion, periodical ischemia, occlusion, periodic

eral vascular disorder, syncope, and tachycardia
DIGESTIVE SYSTEM Anorexia, constipation, dry nodysphagia, colitis, fecal incontinence, gastritis, gastriesis itis, gastrointestinal disorder, increased appetite, inter obstruction, melena, pancreatitis, tenesmus, and vossile HEMATOLOGIC-LYMPHATIC SYSTEM: adenopathy

INFECTIONS AND INFESTATIONS Viral infe METABOLIC AND NUTRITIONAL DISORDIE Edema, hyperglycemia, increased ALT, and increased A MUSCULOSKELETAL SYSTEM: Arthralgia, arthr DISORDEM myalgia, pathological fracture, and tendon disorder. NERVOUS SYSTEM Central nervous system ne lepression, dizziness, emotional lability, facial paralysis, somnia, vasodilatation, and vertigo

RESPIRATORY SYSTEM Asthma, bronchitis, income cough, dyspnea, epistaxis, laryngitis, pharyngitis, page nia, rhinitis, and sinusitis

SKIN. Alopecia, eczema, pruritus, and sweating. SPECIAL SENSES. Cataract. UROGENITAL SYSTEM. Cervix disorder, end carcinoma, epididymitia, and impotence.

DRUG ABUSE AND DEPENDENCE

Lovaza does not have any known drug abuse or withde effects.

OVERDOSAGE

In the event of an overdose, the patient should be symptomatically, and general supportive care i stituted, as required

DOSAGE AND ADMINISTRATION

Patients should be placed on an appropriate lipid landet before receiving Lovaza, and should continue the during treatment with Lovaza. In chinical studies, was administered with meals

The daily dose of Lovaza is 4 g per day. The daily dose be taken as a single 4-g dose (4 capsules) or as two 3 (2 capsules given twice daily)

HOW SUPPLIED

Lovaza (omega-3-acid ethyl esters) capsules are applications are applications to the control of the control of

Recommended Storage: Store at 25°C (77°F), excursions permitted to 15°(59°-86°F) [see USP Controlled Room Temperature] freeze Keep out of reach of children.

stion will be superseded by supple



Placebo was corn oil for all studies

Novised: June 2007 Distributed by: Reliant Pharmaceuticals, Inc. Liberty Corner, NJ 07938 Address Medical Inquiries to Reliant Medical Inquiries de PPD \$655 Meridian Parkway Durham, NC 27713-2203 r Call: 877-311-7515 14252713 PRINTED IN USA 2007 Reliant Pharmaceuticals, Inc. Shown in Product Identification Guide, page 329

RYTHMOL® SR

ryth-mul] ronafenone hydrochloride) extended release

CAPSULES

DESCRIPTION

RYTHMOL SR (propafenone hydrochloride) is an antiarrhythmic drug supplied in extended-release capsules of 225, 325 and 425 mg for oral administration. The structural formula of propagenone HCl is given below:

2'-{2-Hydroxy-3-(propylamino) -propoxy}-3-phenylpropiophenone hydrochlonde

Propagenone HCl has some structural similarities to betablocking agents. Propafenone HCl occurs as colorless crystals or white crystalline powder with a very bitter taste. It is slightly soluble in water (20°C), chloroform and ethanol. Rythmol SR are capsules filled with cylindrical-shaped 2 × 2 mm microtablets containing propaganone and the following mactive ingredients antifoam, gelatin, hypromellose, red iron oxide, magnesium stearate, shellac, sodium lauryl sulfate, sodium dodecyl sulfate, soy lecithin and titanium

CLINICAL PHARMACOLOGY

Mechanism of Action

Propafenone is a Class 1C antiarrhythmic drug with local anesthetic effects, and a direct stabilizing action on myocardial membranes The electrophysiological effect of propagenone manifests itself in a reduction of upstroke velocity (Phase 0) of the monophasic action potential. In Purkinje fibers, and to a lesser extent myocardial fibers, proparenone reduces the fast inward current carried by sodrum ions. Diastolic excitability threshold is increased and effective refractory period prolonged Propafenone reduces spontaneous automaticity and depresses triggered activity Studies in anesthetized dogs and isolated organ preparations show that propafenone has beta-sympatholytic activ ity at about 1/50 the potency of propranolol. Chinical studies employing isoproterenol challenge and exercise testing after single doses of propafenone indicate a beta-adrenergic blocking potency (per mg) about 1/40 that of propranolol in man In clinical trials with the immediate release formula-tion, resting heart rate decreases of about 8% were noted at the higher end of the therapeutic plasma concentration range. At very high concentrations in vitro, propagenone can inhibit the slow inward current carried by calcium, but this calcium antagonist effect probably does not contribute to an-tiarrhythmic efficacy. Moreover, propafenone inhibits a vanety of cardiac potassium currents in in vitro studies (i.e. the transient outward, the delayed rectifier, and the inward rectifier current) Propafenone has local anesthetic activity approximately equal to procaine. Compared to propafenone approximately equat to protein compare to proper include the main metabolite, 5-hydroxypropafenone, has similar so-dium and calcium channel activity, but about 10 times less beta-blocking activity (N-depropylpropafenone has weaker sodium channel activity but equivalent affinity for betareceptors).

Electrophysiology:

Electrophysiology studies in patients with ventricular tachycardia (VT) have shown that propagenone prolongs atmoventricular (AV) conduction while having little or no effect on sinus node function. Both atmoventricular (AV) nodal conduction time (AH interval) and His-Purkinge conduction time (HV interval) are prolonged. Propafenone has little or no effect on the atrial functional refractory period, but AV nodal functional and effective refractory periods are pro-longed In patients with Wolff-Parkinson-White (WPW) syn-drome, RYTHMOL immediate release tablets reduce conduction and increase the effective refractory period of the accessory pathway in both directions (see ADVERSE RE-ACTIONS/Electrocardiograms).

Studies in humans have shown that propafenone exerts a negative inotropic effect on the myocardium Cardiac eatheterization studies in patients with moderately impaired ventricular function (mean C I = $2.61 \, \mathrm{L/min/m^2}$), utilizing

Table 1: Analysis of tachycardia-free period (days) from Day 1 of randomization

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Parameter	225 mg BID (N = 126) n (%)	325 mg BID (N = 135) n (%)	425 mg BID (N = 136) n (%)	Placebo (N = 126 n (%)	
Patients completing with terminating event+	66 (52)	56 (41)	41 (30)	87 (69)	
Comparison of tachycardia-free periods					
Kaplan-Meier Median	112	291	*	41	
Range	0 - 285	0 - 293	0 ~ 300	0 - 289	
p-Value (Log-rank test)	0 014	< 0 0001	< 0 0001		
Hazard Ratio compared to placebo	0 67	0.43	0 35		
95% CI for Hazard Ratio	(0.49, 0 93)	(0 31, 0.61)	(0.24, 0.51)		

- Fewer than 50% of the patients had events. The median time is not calculable
 Terminating events comprised 91% atrial fibrillation, 5% atrial flutter, and 4% PSVT.

intravenous propagenone infusions (loading dose of 2 mg/kg over 10 mm+ followed by 2 mg/min for 30 mm) that gave mean plasma concentrations of 3.0 µg/mL (a dose that produces plasma levels of propafenone greater than does rec-ommended oral dosing), showed significant increases in pulmonary capillary wedge pressure, systemic and pulmo vascular resistances and depression of cardiac output and cardiac index.

Pharmacokinetics and Metabolism:

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Absorption/Bioavsilability: Maximal plasma levels of propagenone are reached between three to eight hours foling the administration of RYTHMOL SR. Propafenone is known to undergo extensive and saturable presystemic biotransformation which results in a dose and dosage form dependent absolute bioavailability; e.g., a 150 mg immediate release tablet had an absolute bioavailability of 3 4%, while a 300 mg immediate release tablet had an absolute bioavailability of 10.6%. Absorption from a 300 mg dose was rapid, with an absolute bioavailability of 21.4%. At still larger doses, above those recommended, bioavailability of propafenone from immediate release tablets increased still further

Relative bioavailability assessments have been performed between RYTHMOL SR capsules and RYTHMOL immediate release tablets. In extensive metabolizers, the biograilability of propafenone from the SR formulation was less than that of the immediate release formulation as the more gradual release of propafenone from the prolonged-release preparations resulted in an increase in overall first pass metabolism (see Metabolism) As a result of the increased first pass effect, higher daily doses of propafenone were required from the SR formulation relative to the immediate release formulation, to obtain similar exposure to propafenone. The relative bioavailability of propafenone from the 325 twice daily regimens of RYTHMOL SR approximates that of RYTHMOL immediate release 150 mg three times daily regimen. Mean exposure to 5-hydroxypropafenone was about 20-25% higher after SR capsule administration than after immediate-release tablet administration

atter immediate-release tablet administration.
Food increased the exposure to propafenone 4-fold after single dose administration of 425 mg of RYTHMOL SR. However, in the multiple dose study (425 mg dose BID), the difference between the fed and fasted state was not significant. Distribution: Following intravenous administration of propafenone, plasma levels decline in a bi-phasic manner consistent with a two compartment pharmacokinetic model. The average distribution half-life corresponding to the first phase was about five minutes. The volume of the central compartment was about 88 liters (1 1 L/kg) and the total me of about 252 liters.

In serum, propafenone is greater than 95% bound to proterns within the concentration range of $0.5-2~\mu g/mL$. Protein binding decreases to about 88% in patients with severe hepatic dysfunction.

Metabolism: There are two genetically determined patterns of propagenone metabolism. In over 90% of patients, the drug is rapidly and extensively metabolized with an elimination half-life from 2-10 hours. These patients metabolize propafenone into two active metabolites. 5-hydroxypropafenone which is formed by CYP2D6 and Ndepropylpropafenone (norpropafenone) which is formed by both CYP3A4 and CYP1A2. In less than 10% of patients, metabolism of propagenone is slower because the 5-hydroxy metabolite is not formed or is minimally formed. In the patients, the estimated propafenone elimination half-life ranges from 10-32 hours. Decreased ability to form the 5-hydroxy metabolite of propagenone is associated with a diminished ability to metabolize debrisoguine and a variety of other drugs such as encannde, metoprolol, and dextrometh-orphan whose metabolism is mediated by the CYP2D6 iso-zyme. In these patients, the N-depropylpropafenone metab-olite occurs in quantities comparable to the levels occurring in extensive metabolizers.

As a consequence of the observed differences in metabolism. administration of RYTHMOL SR to slow and extensive metabolizers results in significant differences in plasma concentrations of propafenone, with slow metabolizers achieving concentrations about twice those of the extensive metabolizers at daily doses of 850, mg/day At low doses the

differences are greater, with slow metabolizers attaining concentrations about three to four times higher than extensive metabolizers. In extensive metabolizers, saturation of the hydroxylation pathway (CYP2D6) results in greater-than-linear increases in plasma levels following administra-tion of RYTHMOL SR capsules. In slow metabolizers, propagenone pharmacokinetics are linear. Because the dif-ference decreases at high doses and is mitigated by the lack of the active 5-hydroxy metabolite in the slow metabolizers and because steady-state conditions are achieved after four to five days of dosing in all patients, the recommended dosng regimen of RYTHMOL SR is the same for all patients. The large inter-subject variability in blood levels require that the dose of the drug be titrated carefully in patients with close attention paid to chinical and ECG evid toxicity (see DOSAGE AND ADMINISTRATION).

The 5-hydroxypropafenone and norpropafenone metabolites have electrophysiologic properties similar to propafenone in vitro. In man after administration of RYTHMOL SR, the 5-hydroxypropafenone metabolite is usually present in con-centrations less than 40% of propafenone. The norpropatenone metabolite is usually present in concentrass than 10% of propafenone

Inter-Subject Variability:

With propafenone, there is a considerable degree of intersubject variability in pharmacokinetics which is due in large part to the first pass hepatic effect and non-linear pharmacokinetics in extensive metabolizers. A higher degree of inter-subject variability in pharmacokinetic parameters of propafenone was observed following both single and multiple dose administration of RYTHMOL SR capsules. Inter-subject variability appears to be substantially less in the poor metabolizer group than in the extensive metabo-lizer group, suggesting that a large portion of the variability s intrinsic to CYP2D6 polymorphism rather than to the for

The clearance of propafenone is reduced and the elimination half-life increased in patients with significant hepatic dys-function (see PRECAUTIONS). Decreased liver function also increases the bioavailability of propafenone. Absolute bioavailability assessments have not been determined for the RYTHMOL SR capsule formulation. Absolute bioavailability of RYTHMOL immediate release tablets has been demonstrated to be inversely related to indocyanine green clearance, reaching 60-70% at clearances of 7 mL/min and

Stereochemistry:

RYTHMOL is a racemic mixture. The R- and S-enantiomers of propafenone display stereoselective disposition characteristics in vitro and in vivo studies have shown that the R-isomer of propafenone is cleared faster than the S-isomer via the 5-hydroxylation pathway (CYP2D6). This results in a higher ratio of S-propagenone to R-propagenone at steady state. Both enantiomers have equivalent potency to block sodium channels; however, the S-enantiomer is a more po-tent \$\mathcal{G}\$-antagonist than the R-enantiomer. Following administration of RYTHMOL immediate release tablets or RYTHMOL SR capsules, the S/R ratio for the area under the plasma concentration-time curve was about 1.7 The S/R ratios of propafenone obtained after administration of 225, 325 and $425\,$ mg RYTHMOL SR are independent of dose. In addition, no difference in the average values of the S/R ratios is evident between genotypes or over time

Clinical Trials:

RYTHMOL SR has been evaluated in patients with a history of electrocardiographically documented recurrent episodes of symptomatic atrial fibrillation in two randomized, double-blind, placebo controlled trials

RAFT: In one US multicenter study (Rythmol SR Atrial Fibrillation Trial, RAFT), three doses of RYTHMOL SR (225 mg BID, 325 mg BID and 425 mg BID) and placebo were compared in 523 patients with symptomatic, episodic atrial fibrillation. The patient population in this trial was 59% male with a mean age of 63 years, 91% White and 6% Black. The patients had a median history of atrial fibrillation of 13 months, and documented symptomatic atrial fi-

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