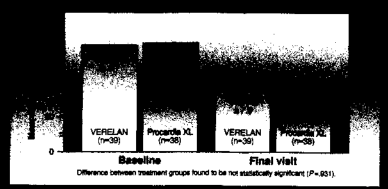


VERELAN

AS EFFECTIVE AS PROCARDIA XL IN REDUCING BP AT THE 24TH HOUR

Reduction in mean DBP measured 24 ± 2 hours after dosing



Results of a 12-week, randomized, double-blind, parallel, comparative study of patients with mild-to-moderate by perfension in 10-study sites nationwide. Patients not controlled on VERLLAN 240 mg/day svere titrated to 360 mg/day and, if needed, 480 mg/day, patients not controlled on Procardia XL30 mg/day were titrated to 60 mg/day and, if needed, 30 mg/day.

□ No significant difference between groups in the number of titrations to goal DBP (<90 mm Hg)

Procardia M. is a registered trademark of Phyor Inc.

Constipation, which can easily be managed in most patients, is the most frequently reported side effect of verapamil.

Please see brief summary of Prescribing Information including CONTRAINDICATIONS, WARNINGS, and PRECAUTIONS on last page.

VERELAN

EXCELLENT TOLERABILITY SIMILAR TO PLACEBO IN A DOUBLE-BLIND STUDY

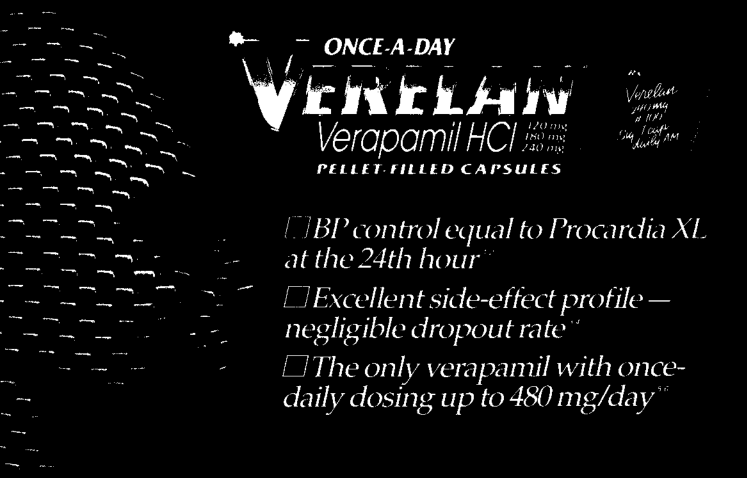
Incidence of side effects commonly associated with calcium channel blockers



Results of a 4-week, double bland-placebo-controlled study of patients with essential hypertension ATRITAN 120 ang/day n =28-240 mg/day n =20-placebo-n =26-20.

No patients discontinued VERFLAN therapy due to constipation, headache, dizziness, or edema





References: 1. Levy B. Rosenberg LN, Colasants DA. A comparison of VEREL AN® and Procardia® XL in the treatment of patients with midd to moderate hypertension. American College of Clinical Pharmacology. 21st Annual Meeting, 1992. Abstract. 2. Further analysis of Levy B. et al. (See reference 1.) Data on tile. Lederia Laboratories, Psarl River, NY. 3. Carr. AA, Bottini PB, Prisant LM, et al. Once-daily varapamil in the treatment of mid-to-moderate hypertension: a double-blind placebo-controlled dose-ranging study. J Clin Pharmacol' 1991;31:144-150, 490. 4. Further analysis of Carr AA, et al. (See reference 3.) Data on tile. Lederia Laboratories, Psarl River, NY. 5. VERELAN Prescripting Information. 3. Physicians' Desk Reference®, 48th ed. Montvale, NJ: Medical Economics Data; 1992;1181-1183 (Isoptin® SR) 2157-2159 (Islan® SR).

Brief Sommers

YERELAN*

Hease Pallet-Filled Capsules

For complete Prescribing Information, consult package insert

CLINICAL PRANMACOLOGY

Food does not affect the extent or rate of the absorption of verapamil from the controlled release VERELAN capsule.

Antioventricular block can occur in patients without prevaising condition defects (see WARRINGS).

Acceleration of ventricular rate and/or ventricular fibrilation has been reported in patients with atrial flutter or atrial fibrilation and a coexisting accessory AV pathway following administration of verapamil (see WARRINGS).

In patients with hepatic insufficiency, metabolism is delayed and elixination half-life prolonged up to 14 to 16 hours (see PRECAUTIONS), the volume of distribution is increased, and plasma clearance reduced to about 30% of normal.

Severe LV dysfunction (see MARKINSS), hypotension (systolic pressure <90 mmHg) or cardiogenic shock, sick sinus synfrome (if no pacemaker is present), second- or third-degree AV block (if no pacemaker is present), alreal flutter/dibrikation with an accessory bypass tract (eg. WPW or LGL syndromes), (see MARKINSS), hypersensitivity to

MARKINGS.

Verapamil should be avoided in patients with severe LV dysfunction (eg., ejection fraction < 30%) or moderate-to-severe symptoms of cardiac failure and in patients with any degree of ventricular dysfunction if they are receiving a beta blocker. Control milder heart failure with optimum digitalization and/or distracts before VEREL AN is used. Verapamil may occasionally produce hypotension. Elevations of liver enzymes have been reported.

Several cases of hepaticellular injury have been demonstrated to be produced by verapamil. Periodic monitoring of liver function in patients on verapamils in grudent. Some patients with paroxysmal and/or chronic atrial further/librillation and an accessory pathway (eg., WPW or LGL syndromes) have developed an increased antegrade conduction across the accessory pathway bypassing the AV node, producing a very rapid ventricular response or ventricular fibrillation after receiving IV verapamil (or digitals). Because of this risk, oral verapamil is contraindicated in such patients. AV block may occur (second- or third-degree, D. 8%). Development of marked first-degree block or progression to second- or third-degree block requires reduction in dosage or, rarely, discontinuation and institution of appropriate therapy. Sinus bradycardis, second-degree AV block, sinus arrest, pulmonary edema and/or severe bypotension were seen in some critically cardia, second-degree AV block, sinus arrest, pulmonary edema and/or severe hypotension were seen in some critically Ill patients with hypertrophic cardiomyopathy who were treated with verapamir.

PRECAUTIONS

Verapamil should be given cautiously to patients with impaired hapatic function (in severe dysfunction use about 30% of the normal dose) or impaired renal function, and patients should be monitored for abnormal prolongation of the PR interval or other signs of overdosage, Verapamil may decrease neuromuscular transmission in patients with Duchamer muscular dystophy and may prolong recovery from the neuromuscular blocking agent veruronium. He may be necessary to decrease verapamil dosage in patients with attenuated neuromuscular transmission. Combined therapy with beta-adrenengic blockers and verapamil may result in additive negative reflects on hear fatte, afforement conduction and or cardiac contractifity, there have been reports of excessive bradypardia and AV block, including complete heart block. The risks of such combined therapy may outweigh the benefits. The combination should be used only with caution and

VERELANO Verapamii HCI

close monitoring. Decreased metoproloi clearance may occur with combined use. Chronic varapamil treatment can increase serum digozin levots by 50% to 75% during the first week of therapy, which can result in digitals toxicity. In patients with hepatic cirrihosis, verapamil may reduce total body clearance and extrarenal clearance of digitoxin. The digoxin does should be reduced when verapamil is given and the patient carefully monitored. Verapamil will usually have an additive effect in patients receiving blood pressure-lowering agents. Disopyramide should not be given within 48 hours before or 24 hours after verapamil administration. Concomitant use of flectaintie and verapamil and quintidine therapy in patients with hypertrophic cardiomyopathy should be avoided, since significant hypotension may result. Verapamil has been given concomitantly with short- and long-acting intrates without any undestable interactions. Interaction between cirreditine and chronically administered verapamil has not been studied. In healthy volunteers, clearance of verapamil was reduced or unchanged. Concomitant use of liftigum and verapamil may result in a lovering of sarum fithium levels or increased sensitivity to liftigum. Patients receiving both drugs must be monitored carefully. Verapamil may increase carbamazepine concentrations during combined use. Ritampin may reduce verapamil bioavailability. Pherohabstical may increase verapamil clearance, Verapamil may increase serum levels of cyclosporine. Concomitant use of inhalation anesthetics and calcium antagenists needs careful titration to avoid excessive cardiovascular depression. Verapamil may prometate the activity of neuromuscular folloxing opens to currace lives of explaints of the surface of the prometal control of the surface verapamic depolaritingly designed tooking agents (currace lives) of explaints of the surface of the prometal of explaints of the surface of the prometal of explaints of the prometal control tooking agents (currace lives) of explaints of the surface of

Reversible (upon discontinuation of verapamil) nonobstructive, paralytic ileus has been infrequently reported in asso-

Reversible (upon discontinuation of verapamil) nonobstructive, paralytic fleus has been infrequently reported in asso-ciation with the use of verapamil.
In clinical trials with 285 hyperfensive patients on VERELAN for more than 1 week, the following adverse reactions were reported: constipation (7.4%); headache (5.3%); dizziness (4.2%); lethargy (3.2%); dyspopsia (2.5%); rash (1.4%); ankie edema (1.4%); sleep disturbance (1.4%); mysigla (1.1%); in clinical trials of other formulations of vera-amil NCI (8 – 4.954); the following reactions have occurred at rates greater than 1.0%; constipation (7.3%); dizziness (3.3%); nausea (2.7%); hypotension (2.5%); adema (1.9%); headache (2.2%); rash (1.2%); CHFipulmonary adema (1.8%); talegue (1.7%); broydcardia (14.5%); N/W block-total 1*, 2*, 3* (1.2%); 2* and 3* (0.5%); flushing (0.5%); elevated here enzymes (see MARKHRGS).

The following reactions, reported in 1.0% or less of patients, occurred under conditions (open trials, marketing experience) where a causal relationship is uncertain. Cardievascutar: angina pectoris, atrioventricular dissociation, chest pain, claudication, repocardial infarction, applications, purpura (vascullis), syntopes. Bypastive System: darrhea, dry mouth, pastroinistriand distribus, ginglival hyperplass. Nervices at Lymphestic ecchymosis or bruising. Nervoess System: cerebrovascular accident, confusion, equilibrium disorders, insomaia, muscle cramps, paresthesis, psychotic symptoms, stakiness, somolence. Registratory; dyspense. Midia rathrigia and rash, exambema, his, exambema, blus, exambema, also, styperformations, maculae, sweating, urticaria, Stevens-Johnson syndrome, erythema multiforme. Speciel Senses: blurred vision. Uraparatial: gynecomassia, impotence, increased urination, spotty mentativation.



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						Side							
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1 Based on Cetaleno and data from Facts and Comparisons,* and used only to reflect relative side effects. The hydrocodone component in LORGET* 10/650 may cause sedation, nausea, vomiting, and constitution.

New Lorcet 10/650

Each tablet contains: 10 mg hydrocodoné bitartrate (Warning: May be habit-forming) and 650 mg acetaminophen.

The Phone-In Pain Relief with the Most Power

*In most states

For references and brief summary of prescribing information, see adjacent page.

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NewLorcet 10/650 (1)

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References:

1. Acute Pain Management Guideline Panel. Acute Pain Management:
Operative or Medical Procedures and Trauma. AHCPR Pub. No. 92-0032.
Rockville, MD. Agency for Health Care Policy and Research. Public Health
Service, US Department of Health and Human Services February 1992.
See especially pages iii, 5, 17, and 113. 2. Drug Information for the Health
Care Professional, Volume 18. Rockville. MD: US Pharmacopeial
Convention, 1992. 3. Catalano RB. The medical approach to management
of pain caused by cancer. Seminars in Oncology. 1975;2:379–392.

4. Narcotic agonist analgesics, in Kastrup EK (ed): Facts and
Companisons St. Louis, J.B. Lippincon Company, 1990, p. 242.

of pain caused by cancer. Seminars in Oncology 1975;2:379–382.

A Narcottic agonist analgesis. In Kastrup Et. (ed): Facts and Companisons St. Louis, J.B. Lippincon Company, 1990, p. 242.

INDICATIONS AND USAGE: For the relief of moderate to moderately severe pain CONTRANDICATIONS: Hypersensitivity to acetaminophen or hydrocodone. WARNINGS: Respiratory Depressions: A high doss or in sensitive patients, hydrocodone may produce dose-related respiratory depression by acting directly on the train stem respiratory center. Hydrocodone also affects the center that controls respiratory inflythm, and may produce irregular and periodic breating. Need layer sent leneaged intracrinist Pressure: The respiratory depression relects of narcotics and their capacity to elevate cerebrospical fluid pressure may be markedly exaggerated in the pressure of head injury, other intracrinal lesons or a pressisting increase in intracrinal pressure. Further more, narcotics produce adverse reactions which may obscure the diagnosis or clinical course of patients with acute abdominal conditions. PRECAUTIONS: Special Rike Patients: As with any narcotic analgesic agent. Lorden 10550 should be used with caution in elderly or debilitated patients and those with severe impairment of hepatic or renal function. Nypothy-rodism. Addisons disease; prostatic hypertrophy or userial stricture. The usual pressultions should be observed and the possibility of respiratory depression should be kept in mind. Lorgen Reliable; Hydrocodone suppressively agents, or other CRS depressants (notioning apches) concernitarity with Lorgen 10650 may etihot; an additive CRS depressants including acception concernitarity with Lorgen 10650 may etihot; an additive CRS depressant floridocodone. The use of MAG inhibitors of tricyclic amideratesants with hydrocodone by requirations may increase the effect of either the amideratesant of hydrocodone has been shown to be fertadepenic. Pleace of high patient in the patient is decided the patient of the development of the patient in t produce dose-related respiratory depression by acting directly on the brain stem respiratory center. Hydrocotione as affects the center that controls respiratory rythm, and may produce regular and periodic breating it significant respiratory depression occurs, it may be antagonized by the use of nationane hydrochloride. Apply other support the measures when indicated. PMIP &BUBE AND DEPENDENCE: Loncat* 10/650 is subject to the Federal Controlled Substances Act (Schedule III). Psychic dependence, physical dependence, and tolerance may develop upon repeated administration of narcolics; therefore. Lorcet* 10/650 should be prescribed and administered with caution. However, psychic dependence is mightly to develop when Lorcet* 10/650 is used for a short time for the treatment of pain. BVERDOSABE: Acetaminophene: Signs and Symptoms: In acute acetaminophene overdosage, dose-dependent, optentially fallal hepatic necross is the most serious adverse effect. Renal lubular necross, propolycemic coma, and this mobile of the statement of pain. BVERDOSABE: Acetaminophene: Early symptoms following a portentially hepatotoxic overdose may include: na usea, vomiting, diaphoresis and general midalise. Climical and laboratory evidence of hepatic toxicity may not be apparent until 45 to 72 hours post-ingestion. Hydrocodone: Signs and Symptoms: Serious overdose with hydrocodone is characterized by respiratory rate and chamby skin, and sometimes brandpards and hypotension in severe overdosage, apina. Circulatory collapse, cardisc arrest and death may skin, and sometimes brandpards and hypotension in severe overdosage, apina. Circulatory collapse, cardisc arrest and death may skin, and sometimes brandpards and hypotension in severe overdosage, apina. Circulatory collapse, cardisc arrest and death may skin, and sometimes brandpards and hypotension in severe overdosage, apina. Circulatory collapse, cardisc arrest and death may skin, and sometimes brandpards and hypotension in severe overdosage, apina. Circulatory collapse, cardisc arrest

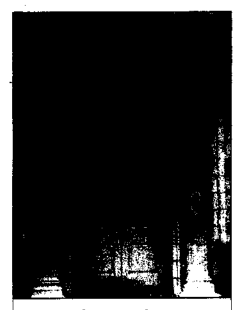


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Morrow JD, Margolies GR, Rowland J, Roberts LJ 2nd. Evidence that histamine is the causative toxin of scombroid-fish poisoning. N Engl J Med 1991; 324:716-20.

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Clinical Experience Network (CEN). A large-scale, office-based study evaluates the use of a new class of nonsedating antihistamines. A report from CEN. J Am Board Fam Pract 1990; 3:241-58.

Book

Rakel RE. Textbook of family practice. 4th ed. Philadelphia: WB Saunders, 1990

Chapter in Book

Haynes RC Jr. Agents affecting calcification: calcium, parathyroid hormone, calcitonin, vitamin D, and other compounds. In: Gilman AG, Rall TW, Nies AS, Taylor P, editors. Goodman and Gilman's the pharmacological basis of therapeutics. 8th ed. New York: Pergamon Press, 1990.

Government Agency

Schwartz JL. Review and evaluation of smoking cessation methods: the United States and Canada, 1978-1985. Bethesda, MD: Department of Health and Human Services, 1987. (NIH publication no. 87-2940.)

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