

www.pdr.net



BIOCON PHARMA LTD (IPR2020-01263) Ex. 1012, p. 001





Senior Vice President, Directory Services: Paul Walsh

Director of Product Management: Mark A. Friedman Associate Product Manager: Bill Shaughnessy Senior Business Manager: Mark S. Ritchin Financial Analyst: Wayne M. Soltis Director of Sales: Dikran N. Barsamian National Sales Manager, Pharmaceutical Sales: Anthony Sorce National Account Manager: Don Bruccoleri Senior Account Manager: Frank Karkowsky **Account Managers:** Marion Gray, RPh Lawrence C. Keary Jeffrey F. Pfohl Suzanne E. Yarrow, RN Electronic Sales Account Manager: Stephen M. Silverberg National Sales Manager, Medical Economics Trade Sales: Bill Gaffney Director of Direct Marketing: Michael Bennett List and Production Manager: Lorraine M. Loening Senior Marketing Analyst: Dina A. Maeder

Director, New Business Development and Professional Support Services: Mukesh Mehta, RPh Manager, Drug Information Services: Thomas Fleming, RPh Drug Information Specialist: Maria Deutsch, MS, RPh, CDE Editor, Directory Services: David W. Sifton Senior Associate Editor: Lori Murray Director of Production: Carrie Williams Manager of Production: Kimberly H. Vivas Senior Production Coordinator: Amy B. Brooks Production Coordinators: Gianna Caradonna, Maria Volpati Data Manager: Jeffrey D. Schaefer Senior Format Editor: Gregory J. Westley Key to FDA Use-in-Pr Index Editors: Johanna M. Mazur, Robert N. Woerner Art Associate: Joan K. Akerlind Senior Digital Imaging Coordinator: Shawn W. Cahill Digital Imaging Coordinator: Frank J. McElroy, III Electronic Publishing Designer: Livio Udina Fulfillment Manager: Stephanie DeNardi

Copyright © 2000 and published by Medical Economics Company, Inc. at Montvale, NJ 07645-1742. All rights reserved. None of the content of this publication may be reproduced, stored in a retrieval system, resold, redistributed, or transmitted in any form or by any means (electronic, mechanical, photocopying, recording, or be reported use in the prior written permission of the publisher. PHYSICIANS' DESK REFERENCE®, PDR®, PDR For Ophthalmodgy®, Pocket PDR®, and The PDR® Family Guide to Prescription Drugs® are registered trademarks used herein under license. PDR For Nonprescription Drugs and Dietary Supplements™, PDR Companion Guide™, PDR® for Herbal Medicines™, PDR® Medical Dictionary™, PDR® Nurse's Drug Handbook™, PDR® Nurse's Dictionary™, The PDR® Family Guide to Natural Medicines and Healing Therapies™, The PDR® Family Guide to Common Ailments™, The PDR® Family Guide to Over-the-Counter Drugs™, and PDR® Electronic Library™ are trademarks used herein under license.

Officers of Medical Economics Company: President and Chief Executive Officer: Curtis B. Allen; Vice President, New Media: L. Suzanne BeDell; Vice President, Corporate Human Resources: Pamela M. Bilash; Vice President and Chief Information Officer: Steven M. Bressler; Chief Financial Officer: Christopher Caridi; Vice President and Controller: Barry Gray; Vice President, New Business Planning: Linda G. Hope; Vice President, Business Integration: David A. Pitler; Vice President, Finance: Donna Santarpia; Senior Vice President, Directory Services: Paul Walsh; Senior Vice President, Operations: John R. Ware; Senior Vice President, Internet Strategies: Raymond Zoeller



ISBN: 1-56363-330-2

BIOCON PHARMA LTD (IPR2020-01263) Ex. 1012, p. 002

PRODUCT INFORMATION

it with D.H.E. 45® (dihydroergotamine mesylate) Injection, USP unless at least 6 hours have elapsed since your last injection. No more than 6 mL of D.H.E. 45® (dihydroergotamine mesylate) Injection, USP should be injected during a one-week period.

Learn what to do in case of an Overdose

If you have used more medication than you have been instructed, contact your doctor, hospital emergency department, or nearest poison control center immediately. How to use the D.H.E. 45[®] (dihydroergotamine mesylate) Injection, USP

- Use available training materials.
 Read and follow the instructions in the patient instruction booklet which is provided with the D.H.E. 45® (dihydroergotamine mesylate) Injection, USP package before attempting to use the product.
- If there are any questions concerning the use of your D.H.E. 45® (dihydroergotamine mesylate) Injection, USP, ask your Doctor or pharmacist.

2. Preparing for the Injection

• Carefully examine the ampul (glass vial) of D.H.E. 45® (dihydroergotamine mesylate) Injection, USP for any cracks or breaks, and the liquid for discoloration, cloud-iness, or particles. If any of these defects are present, use a new ampul, make certain it is intact, and return the defective ampul to your doctor or pharmacy. Once you open an ampul, if it is not used within an hour, it should be thrown away.

3. Locating an Injection Site

- Administer your subcutaneous Injection in the middle of your thigh, well above the knee.
- 4. Drawing the Medication into the Syringe
 Wash your hands thoroughly with soap and water.
 Check the dose of your medication.

 - Check the dose of your medication.
 Look to see if there is any liquid at the top of the ampul. If there is, gently flick the ampul with your finger to get all the liquid into the bottom portion of the ampul.
 Hold the bottom of the ampul in one hand. Clean the ampul neck with an alcohol wipe using your other hand. Then place the alcohol wipe around the neck of the ampul and heapt it acome her merging your them the ampul and break it open by pressing your thumb
 - against the neck of the ampul. Tilt the ampul down at a 45° angle. Insert the needle into the solution in the ampul.

 - into the solution in the ampul.
 Draw up the medication by pulling back the plunger slowly and steadily until you reach your dose.
 Check the syringe for air bubbles. Hold it with the needle pointing upward. If there are air bubbles, tap your finger against the barrel of the syringe to get the bubbles to the top. Slowly and carefully push the plunger up so that the bubbles are pushed out through the needle and you see a drop of medication.
 When there are no air bubbles check the dose of the
 - When there are no air bubbles, check the dose of the medication. If the dose is incorrect, repeat steps 6 through 8 until you draw up the right dose.

5. Preparing the Injection Site

With a new alcohol wipe, clean the selected injection site thoroughly with a firm, circular motion from inside to outside. Wait for the injection site to dry before injecting.

6. Administering the Injection

- Hold the syringe/needle in your right hand. • With your left hand, firmly grasp about a 1-inch fold of
- skin at the injection site. Push the needle shaft, bevel side up, all the way into the fold of skin at a 45° to 90° angle, then release the
- fold of skin. While holding the syringe with your left hand, use your right hand to draw back slightly on the plunger.If you do not see any blood coming back into the sy-
- If you do not see any block coming back more sp-ringe, inject the medication by pushing down on the plunger. If you do see blood in the syringe, that means the needle has penetrated a vein. If this happens, pull the needle/syringe out of the skin slightly and draw back on the plunger again. If no blood is seen this time, inject the medication.
- Use your right hand to pull the needle out of your skin quickly at the same angle you injected it. Immediately press the alcohol wipe on the injection site and rub.

Check the expiration date printed on the ampul containing medication. If the expiration date has passed, do not use it. Answers to patients' questions about D.H.E. 45® (dihydro-ergotamine mesylate) Injection, USP

What if I need help in using my D.H.E. 45® (dihydroergotamine mesylate) Injection, USP?

If you have any questions or if you need help in opening, putting together, or using D.H.E. 45® (dihydroergotamine mesylate) Injection, USP, speak to your doctor or pharmacist.

How much medication should I use and how often?

Your doctor will have told you what dose to use for each normalization of the second state of the secon (dihydroergotamine mesylate) Injection, USP should be injected during a one-week period. Do not use more than this amount unless instructed to do so by your doctor.

If you have any other unanswered question about D.H.E 45® (dihydroergotamine mesylate) Injection, USP, consult your doctor or pharmacist.

*Trademark of Medical Economics Company, Inc. **REV: MAY 1998**

DIOVAN® [dī-o-van] valsartan Capsules

Rx only The following prescribing information is based on official labeling in effect July 1999.

Prescribing Information

USE IN PREGNANCY

When used in pregnancy during the second and third trimesters, drugs that act directly on the reninangiotensin system can cause injury and even death to the developing fetus. When pregnancy is detected, Diovan should be discontinued as soon as possible. See WARNINGS: Fetal/Neonatal Morbidity and Mortality.

DESCRIPTION

Diovan (valsartan) is a nonpeptide, orally active, and specific angiotensin II antagonist acting on the AT_1 receptor subtype.

Valsartan is chemically described as N-(1-oxopentyl)-N-[[2'-(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl]methyl]-L-valine. Its empirical formula is $C_{24}H_{23}N_5O_3$, its molecular weight is 435.5, and its structural formula is



Valsartan is a white to practically white fine powder. It is soluble in ethanol and methanol and slightly soluble in water.

Diovan is available as capsules for oral administration, con-taining either 80 mg or 160 mg of valsartan. The inactive ingredients of the capsules are cellulose compounds, crospovidone, gelatin, iron oxides, magnesium stearate, povidone, sodium lauryl sulfate, and titanium dioxide.

CLINICAL PHARMACOLOGY

Mechanism of Action

Angiotensin II is formed from angiotensin I in a reaction catalyzed by angiotensin-converting enzyme (ACE, kininase II). Angiotensin II is the principal pressor agent of the renin-angiotensin system, with effects that include vasocon-striction, stimulation of synthesis and release of aldosterone, cardiac stimulation, and renal reabsorption of sodium. Valsartan blocks the vasoconstrictor and aldosteronesecreting effects of angiotensin II by selectively blocking the binding of angiotensin II to the AT_1 receptor in many tis-sues, such as vascular smooth muscle and the adrenal gland. Its action is therefore independent of the pathways

for angiotensin II synthesis. There is also an AT_2 receptor found in many tissues, but AT_2 is not known to be associated with cardiovascular h_{0}^{-1} meostasis. Valsartan has much greater affinity (about 20,000-fold) for the AT₁ receptor than for the AT₂ receptor. The primary metabolite of valsartan is essentially inactive with an affinity for the AT₁ receptor about one 200th that of valsartan itself.

Blockade of the renin-angiotensin system with ACE inhibitors, which inhibit the biosynthesis of angiotensin II from angiotensin I, is widely used in the treatment of hypertension ACE inhibitors also inhibit the degradation of brady-kinin, a reaction also catalyzed by ACE. Because valsartan does not inhibit ACE (kininase II), it does not affect the response to bradykinin. Whether this difference has clinical relevance is not yet known. Valsartan does not bind to or block other hormone receptors or ion channels known to be important in cardiovascular regulation.

Blockade of the angiotensin II receptor inhibits the negative regulatory feedback of angiotensin II on renin secretion, but the resulting increased plasma renin activity and angio tensin II circulating levels do not overcome the effect of valsartan on blood pressure. Pharmacokinetics

Valsartan peak plasma concentration is reached 2 to 4 hours after dosing. Valsartan shows bi-exponential decay kinetics following intravenous administration, with an average elimination half-life of about 6 hours. Absolute bioavailability for the capsule formulation is about 25% (range availability for the capsule formulation is about 25% (range 10%-35%). Food decreases the exposure (as measured by AUC) to valsartan by about 40% and peak plasma concentration (C_{max}) by about 50%. AUC and C_{max} values of valsartan increase approximately linearly with increasing dose over the clinical dosing range. Valsartan does not accumulate appreciably in plasma following repeated administration tion.

Metabolism and Elimination

Valsartan, when administered as an oral solution, is pri-30220906 marily recovered in feces (about 83% of dose) and urine BIOCON PHARMA LTD (IPR2020-01263) Exclerite0.12 tupe 003 for revisions

NOVARTIS PHARMACEUTICALS/2015

(about 13% of dose). The recovery is mainly as unchanged drug, with only about 20% of dose recovered as metabolites. The primary metabolite, accounting for about 9% of dose, is valeryl 4-hydroxy valsartan. The enzyme(s) responsible for valsartan metabolism have not been identified but do not seem to be CYP 450 isozymes.

Following intravenous administration, plasma clearance of valsartan is about 2 L/h and its renal clearance is 0.62 L/h (about 30% of total clearance).

Distribution

Ŗ

The steady state volume of distribution of valsartan after intravenous administration is small (17 L), indicating that valsartan does not distribute into tissues extensively. Valsartan is highly bound to serum proteins (95%), mainly serum albumin.

Special Populations Pediatric: The pharmacokinetics of valsartan have not been investigated in patients <18 years of age.

Geriatric: Exposure (measured by AUC) to valsartan is higher by 70% and the half-life is longer by 35% in the elderly than in the young. No dosage adjustment is neces-sary (see DOSAGE AND ADMINISTRATION).

Gender: Pharmacokinetics of valsartan does not differ significantly between males and females.

Renal Insufficiency: There is no apparent correlation between renal function (measured by creatinine clearance) and exposure (measured by AUC) to valsartan in patients with different degrees of renal impairment. Consequently, dose adjustment is not required in patients with mild-to-moderate renal dysfunction. No studies have been performed in patients with severe impairment of renal function (creatinine clearance <10 mL/min). Valsartan is not removed from the plasma by hemodialysis. In the case of severe renal disease, exercise care with dosing of valsartan (see DOSAGE AND ADMINISTRATION).

Hepatic Insufficiency: On average, patients with mild-to-moderate chronic liver disease have twice the exposure (measured by AUC values) to valsartan of healthy volun-teers (matched by age, sex and weight). In general, no dos-age adjustment is needed in patients with mild-to-moderate liver disease. Care should be exercised in patients with liver disease (see DOSAGE AND ADMINISTRATION).

Pharmacodynamics and Clinical Effects

Valsartan inhibits the pressor effect of angiotensin II infu-sions. An oral dose of 80 mg inhibits the pressor effect by about 80% at peak with approximately 30% inhibition per-sisting for 24 hours. No information on the effect of larger doses is available.

Removal of the negative feedback of angiotensin II causes a 2- to 3-fold rise in plasma renin and consequent rise in angiotensin II plasma concentration in hypertensive patients. Minimal decreases in plasma aldosterone were observed after administration of valsartan; very little effect on serum potassium was observed.

In multiple-dose studies in hypertensive patients with sta-ble renal insufficiency and patients with renovascular hypertension, valsartan had no clinically significant effects on glomerular filtration rate, filtration fraction, creatinine clearance, or renal plasma flow.

In multiple-dose studies in hypertensive patients, valsartan had no notable effects on total cholesterol, fasting triglycerides, fasting serum glucose, or uric acid.

The antihypertensive effects of Diovan were demonstrated principally in 7 placebo-controlled, 4- to 12-week trials (one in patients over 65) of dosages from 10 to 320 mg/day in patients with baseline diastolic blood pressures of 95-115. The studies allowed comparison of once-daily and twicedaily regimens of 160 mg/day; comparison of peak and trough effects; comparison (in pooled data) of response by gender, age, and race; and evaluation of incremental effects of hydrochlorothiazide.

Administration of valsartan to patients with essential hypertension results in a significant reduction of sitting, supine, and standing systolic and diastolic blood pressure, usually with little or no orthostatic change.

In most patients, after administration of a single oral dose, onset of antihypertensive activity occurs at approximately 2 hours, and maximum reduction of blood pressure is achieved within 6 hours. The antihypertensive effect per-sists for 24 hours after dosing, but there is a decrease from peak effect at lower doses (40 mg) presumably reflecting loss of inhibition of angiotensin II. At higher doses, however (160 mg), there is little difference in peak and trough effect. buring repeated dosing, the reduction in blood pressure with any dose is substantially present within 2 weeks, and maximal reduction is generally attained after 4 weeks. In long-term follow-up studies (without placebo control), the effect of valsartan appeared to be maintained for up to two years. The antihypertensive effect is independent of age, gender or race. The latter finding regarding race is based on pooled data and should be viewed with caution, because antihypertensive drugs that affect the renin-angiotensin system (that is, ACE inhibitors and angiotensin-II blockers) have generally been found to be less effective in low-renin hypertensives (frequently blacks) than in highrenin hypertensives (frequently whites). In pooled, random-ized, controlled trials of Diovan that included a total of 140 blacks and 830 whites, valsartan and an ACE-inhibitor control were generally at least as effective in blacks as whites. The explanation for this difference from previous findings is unclear.

Diovan-Cont.

Abrupt withdrawal of valsartan has not been associated with a rapid increase in blood pressure. The blood pressure lowering effect of valsartan and

thiazide-type diuretics are approximately additive The 7 studies of valsartan monotherapy included over 2000 patients randomized to various doses of valsartan and about 800 patients randomized to placebo. Doses below 80 mg were not consistently distinguished from those of pla-cebo at trough, but doses of 80, 160 and 320 mg produced dose-related decreases in systolic and diastolic blood pres-sure, with the difference from placebo of approximately 6-9/3-5 mmHg at 80-160 mg and 9/6 mmHg at 320 mg. In a controlled trial the addition of HCTZ to valsartan 80 mg resulted in additional lowering of systolic and dia-stolic blood pressure by approximately 6/3 and 12/5 mmHg for 12.5 and 25 mg of HCTZ, respectively, compared to valent no. valsartan 80 mg alone.

Patients with an inadequate response to 80 mg once daily were titrated to either 160 mg once daily or 80 mg twice daily, which resulted in a comparable response in both groups.

In controlled trials, the antihypertensive effect of once-daily valsartan 80 mg was similar to that of once-daily enalapril 20 mg or once-daily lisinopril 10 mg.

There was essentially no change in heart rate in valsartantreated patients in controlled trials.

INDICATIONS AND USAGE

Diovan is indicated for the treatment of hypertension. It may be used alone or in combination with other antihypertensive agents.

CONTRAINDICATIONS

Diovan is contraindicated in patients who are hypersensitive to any component of this product. WARNINGS

Fetal/Neonatal Morbidity and Mortality

Drugs that act directly on the renin-angiotensin system can cause fetal and neonatal morbidity and death when administered to pregnant women. Several dozen cases have been reported in the world literature in patients who were taking angiotensin-converting enzyme inhibitors. When pregnancy is detected, Diovan should be discontinued as soon as possible.

The use of drugs that act directly on the renin-angiotensin system during the second and third trimesters of pregnancy has been associated with fetal and neonatal injury, including hypotension, neonatal skull hypoplasia, anuria, reversible or irreversible renal failure, and death. Oligohydramnios has also been reported, presumably resulting from decreased fetal renal function; oligohydramnios in this setting has been associated with fetal limb contractures, craniofacial deformation, and hypoplastic lung develop-ment. Prematurity, intrauterine growth retardation, and patent ductus arteriosus have also been reported, although it is not clear whether these occurrences were due to expo-

These adverse effects do not appear to have resulted from intrauterine drug exposure that has been limited to the first trimester. Mothers whose embryos and fetuses are exposed to an angiotensin II receptor antagonist only during the first trimester should be so informed. Nonetheless, when patients become pregnant, physicians should advise the patient to discontinue the use of valsartan as soon as ossible

Rarely (probably less often than once in every thousand pregnancies), no alternative to a drug acting on the remin-angiotensin system will be found. In these rare cases, the mothers should be apprised of the potential hazards to their fetuses, and serial ultrasound examinations should be performed to assess the intra-amniotic environment.

If oligohydramnios is observed, valsartan should be discontinued unless it is considered life-saving for the mother. Contraction stress testing (CST), a nonstress test (NST), or biophysical profiling (BPP) may be appropriate, depending

biophysical profiling (BPP) may be appropriate, depending upon the week of pregnancy. Patients and physicians should be aware, however, that oligohydramnios may not appear until after the fetus has sustained irreversible injury. Infants with histories of in utero exposure to an angiotensin II receptor antagonist should be closely observed for hypo-tension, oliguria, and hyperkalemia. If oliguria occurs, attention should be directed toward support of blood pres-sure and perfusion. Exchange transfusion or dialysis may be required as means of reversing hypotension and/or may be required as means of reversing hypotension and/or substituting for disordered renal function. No teratogenic effects were observed when valsartan was

administered to pregnant mice and rats at oral doses up to 600 mg/kg/day and to pregnant rabbits at oral doses up to 10 mg/kg/day. However, significant decreases in fetal weight, pup birth weight, pup survival rate, and slight delays in developmental milestones were observed in studies in which parental rats were treated with valsartan at

tes in which parental rats were treated with valsartan at oral, maternally toxic (reduction in body weight gain and food consumption) doses of 600 mg/kg/day during organo-genesis or late gestation and lactation. In rabbits, fetotoxic-ity (i.e., resorptions, litter loss, abortions, and low body weight) associated with maternal toxicity (mortality) was observed at doses of 5 and 10 mg/kg/day. The no observed adverse effect doses of 600, 200 and 2 mg/kg/day in mice, rats and rabbits persent 9 6 and 0.1 timer rats and rabbits represent 9, 6, and 0.1 times, respectively the maximum recommended human dose on a mg/m² basis.

Hypotension in Volume- and /or Salt-Depleted Patients

Excessive reduction of blood pressure was rarely seen (0.1%) in patients with uncomplicated hypertension. In patients with an activated renin-angiotensin system, such as volume- and/or salt-depleted patients receiving high doses of diuretics, symptomatic hypotension may occur. This condition should be corrected prior to administration of Diovan, or the treatment should start under close medical supervision.

If hypotension occurs, the patient should be placed in the supine position and, if necessary, given an intravenous infusion of normal saline. A transient hypotensive response is not a contraindication to further treatment, which usually can be continued without difficulty once the blood pressure has stabilized.

PRECAUTIONS

General

Impaired Hepatic Function: As the majority of valsartan is eliminated in the bile, patients with mild-to-moderate hepatic impairment, including patients with biliary obstruc-tive disorders, showed lower valsartan clearance (higher AUCs). Care should be exercised in administering Diovan to these patients.

Impaired Renal Function: As a consequence of inhibiting the renin-angiotensin-aldosterone system, changes in renal function may be anticipated in susceptible individuals. In patients whose renal function may depend on the activity of the renin-angiotensin-aldosterone system (e.g., patients with severe congestive heart failure), treatment with angio-tensin-converting enzyme inhibitors and angiotensin receptor antagonists has been associated with oliguria and/or progressive azotemia and (rarely) with acute renal failure and/or death. Diovan would be expected to behave similarly. In studies of ACE inhibitors in patients with unilateral or bilateral renal artery stenosis, increases in serum creati-nine or blood urea nitrogen have been reported. In a 4-day trial of valsartan in 12 patients with unilateral renal artery stenosis, no significant increases in serum creatinine or blood urea nitrogen were observed. There has been no longterm use of Diovan in patients with unilateral or bilateral renal artery stenosis, but an effect similar to that seen with ACE inhibitors should be anticipated. Information for Patients

Pregnancy: Female patients of childbearing age should be told about the consequences of second- and third-trimester exposure to drugs that act on the renin-angiotensin system, and they should also be told that these consequences do not appear to have resulted from intrauterine drug exposure that has been limited to the first trimester. These patients should be asked to report pregnancies to their physicians as soon as possible.

Drug Interactions

No clinically significant pharmacokinetic interactions were observed when valsartan was coadministered with amlo-dipine, atenolol, cimetidine, digoxin, furosemide, glyburide, hydrochlorothiazide, or indomethacin. The valsartanatenolol combination was more antihypertensive than either component, but it did not lower the heart rate more than atenolol alone.

Coadministration of valsartan and warfarin did not change the pharmacokinetics of valsartan or the time-course of the anticoagulant properties of warfarin.

CYP 450 Interactions: The enzyme(s) responsible for valsartan metabolism have not been identified but do not seem to be CYP 450 isozymes. The inhibitory or induction potential of valsartan on CYP 450 is also unknown.

Carcinogenesis, Mutagenesis, Impairment of Fertility

There was no evidence of carcinogenicity when valsartan was administered in the diet to mice and rats for up to 2 years at doses up to 160 and 200 mg/kg/day, respectively. These doses in mice and rats are about 2.6 and 6 times, respectively, the maximum recommended human dose on a mg/m² basis. (Calculations assume an oral dose of 320 mg/day and a 60-kg patient.)

Mutagenicity assays did not reveal any valsartan-related effects at either the gene or chromosome level. These assays included bacterial mutagenicity tests with Salmonella (Ames) and E coli; a gene mutation test with Chinese ham-ster V79 cells; a cytogenetic test with Chinese hamster ovary cells; and a rat micronucleus test.

Valsartan had no adverse effects on the reproductive performance of male or female rats at oral doses up to 200 mg/kg/ mance of male or temate rats at oral doses up to 200 hg/hg day. This dose is 6 times the maximum recommended human dose on a mg/m² basis. (Calculations assume an oral dose of 320 mg/day and a 60-kg patient.)

Pregnancy Categories C (first trimester) and D (second and third trimesters)

See WARNINGS, Fetal/Neonatal Morbidity and Mortality. **Nursing Mothers**

It is not known whether valsartan is excreted in human milk, but valsartan was excreted in the milk of lactating rats. Because of the potential for adverse effects on the nursing infant, a decision should be made whether to discontinue nursing or discontinue the drug, taking into account the importance of the drug to the mother. **Pediatric Use**

Safety and effectiveness in pediatric patients have not been established.

Geriatric Use

In the controlled clinical trials of valsartan, 1214 (36.2%) of

PHYSICIANS' DESK REFERENCE®

or safety of valsartan was observed in this patient population, but greater sensitivity of some older individuals cannot be ruled out.

ADVERSE REACTIONS

Diovan has been evaluated for safety in more than 4000 patients, including over 400 treated for over 6 months, and more than 160 for over 1 year. Adverse expe-riences have generally been mild and transient in nature and how are being for the safety of t and have only infrequently required discontinuation of ther-apy. The overall incidence of adverse experiences with

apy. The overall incidence of adverse experiences with Diovan was similar to placebo. The overall frequency of adverse experiences was neither dose-related nor related to gender, age, race, or regimen. Discontinuation of therapy due to side effects was required in 2.3% of valsartan patients and 2.0% of placebo patients. The most common reasons for discontinuation of therapy with Diwne ware bacdeobe and displaced with Diovan were headache and dizziness.

The adverse experiences that occurred in placebo-controlled clinical trials in at least 1% of patients treated with Diovan and at a higher incidence in valsartan (n=2316) than pla-cebo (n=888) patients included viral infection (3% vs. 2%), fatigue (2% vs. 1%), and abdominal pain (2% vs. 1%).

Headache, dizziness, upper respiratory infection, cough, diarrhea, rhinitis, sinusitis, nausea, pharyngitis, edema, and arthralgia occurred at a more than 1% rate but at about

the same incidence in placebo and valsartan patients. In trials in which valsartan was compared to an ACE inhib-itor with or without placebo, the incidence of dry cough was significantly greater in the ACE-inhibitor group (7.9%) than in the groups who received valsartan (2.6%) or placebo (1.5%). In a 129-patient trial limited to patients who had had dry cough when they had previously received ACE inhibitors, the incidences of cough in patients who received valsartan, HCTZ, or lisinopril were 20%, 19%, and 69% respectively (p<0.001). Dose-related orthostatic effects were seen in less than 1% of

patients. An increase in the incidence of dizziness was observed in patients treated with Diovan 320 mg (8%) compared to 10 to 160 mg (2% to 4%).

Diovan has been used concomitantly with hydrochlorothiazide without evidence of clinically important adverse interactions.

Other adverse experiences that occurred in controlled clinical trials of patients treated with Diovan (>0.2% of valsartan patients) are listed below. It cannot be determined whether these events were causally related to Diovan.

Diovan. Body as a Whole: Allergic reaction and asthenia Cardiovascular: Palpitations Dermatologic: Pruritus and rash Digestive: Constipation, dry mouth, dyspepsia, and flatulence

Musculoskeletal: Back pain, muscle cramps, and myalgia Neurologic and Psychiatric: Anxiety, insomnia, paresthesia, and somnolence

Respiratory: Dyspnea Special Senses: Vertigo

Urogenital: Impotence Other reported events seen less frequently in clinical trials included chest pain, syncope, anorexia, vomiting, and angioedema

Post-Marketing Experience The following additional adverse reactions have been reported in post-marketing experience:

Hypersensitivity: There are rare reports of angioedema; Digestive: Elevated liver enzymes and very rare reports of hepatitis.

Clinical Laboratory Test Findings In controlled clinical trials, clinically important changes in standard laboratory parameters were rarely associated with administration of Diovan. Creatinine: Minor elevations in creatinine occurred in

0.8% of patients taking Diovan and 0.6% given placebo in controlled clinical trials. Hemoglobin and Hematocrit: Greater than 20% decreases

in hemoglobin and hematocrit were observed in 0.4% and 0.8%, respectively, of Diovan patients, compared with 0.1% and 0.1% in placebo-treated patients. One valsartan patient discontinued treatment for microcytic anemia.

Liver function tests: Occasional elevations (greater than 150%) of liver chemistries occurred in Diovan-treated patients. Three patients (<0.1%) treated with valsartan dis-

continued treatment for elevated liver chemistries. Neutropenia: Neutropenia was observed in 1.9% of patients treated with Diovan and 0.8% of patients treated with placebo.

Serum Potassium: Greater than 20% increases in serum potassium vero observed in 4.4% of Diovan-treated patients compared to 2.9% of placebo-treated patients. No patient treated with valsartan discontinued therapy for hyperkalemia.

OVERDOSAGE

Limited data are available related to overdosage in humans. Limited data are avaliable related to overdosage in induction The most likely manifestations of overdosage would be hypotension and tachycardia; bradycardia could occur from parasympathetic (vagal) stimulation. If symptomatic hypotension should occur, supportive treatment should be instituted.

Valsartan is not removed from the plasma by hemodialysis. Valsartan was without grossly observable adverse effects at single oral doses up to 2000 mg/kg in rats and up to

PRODUCT INFORMATION

mended human dose on a mg/m² basis). (Calculations assume an oral dose of 320 mg/day and a 60-kg patient.) DOSAGE AND ADMINISTRATION

The recommended starting dose of Diovan is 80 mg once daily when used as monotherapy in patients who are not

volume-depleted. Diovan may be used over a dose range of 80 mg to 320 mg daily, administered once-a-day.

80 mg to 320 mg daily, administered once-a-day. The antihypertensive effect is substantially present within 2 weeks and maximal reduction is generally attained after 4 weeks. If additional antihypertensive effect is required, the dosage may be increased to 160 mg or 320 mg or a diuretic may be added. Addition of a diuretic has a greater effect than dose increases beyond 80 mg. No initial dosage adjustment is required for elderly patients, for patients with mild or moderate renal impair-ment, or for patients with mild or moderate liver insuffi-ciency. Care should be exercised with dosing of Diovan in patients with hepatic or severe renal impairment. Diovan may be administered with other antihypertensive Diovan may be administered with other antihypertensive agents.

Diovan may be administered with or without food.

HOW SUPPLIED

Diovan is available as capsules containing valsartan 80 mg or 160 mg. Both strengths are packaged in bottles of 100 capsules and 4000 capsules and unit dose blister packages. Capsules are imprinted as follows

80 mg Capsule - Light grey/light pink opaque, imprinted CG FZF

Bottles of 100	NDC 0083-4000-0
Bottles of 4000	NDC 0083-4000-4
Unit Dose (blister pack)	NDC 0083-4000-6
Box of 100 (strips of 10)	
160 mg Capsule - Dark grey	light pink opaque, imprinted
CG GOG	Constant and and and an an an and an
Bottles of 100	NDC 0083-4001-01
Bottles of 4000	NDC 0082 4001 41

Unit Dose (blister pack) NDC 0083-4001-61 Box of 100 (strips of 10)

Store below 30°C (86°F). Protect from moisture.

Dispense in tight container (USP). C98-49 (Rev. 12/98)

Distributed by

Novartis Pharmaceuticals Corporation

East Hanover, New Jersey 07936 Shown in Product Identification Guide, page 325

DIOVAN HCT® [dī-o-van]

valsartan and hydrochlorothiazide **Combination Tablets** 80 mg/12.5 mg 160 mg/12.5 mg

Rx only The following prescribing information is based on official labeling in effect on July 1999.

USE IN PREGNANCY

When used in pregnancy during the second and third trimesters, drugs that act directly on the renin-angiotensin system can cause injury and even death to the developing fetus. When pregnancy is detected, Diovan HCT should be discontinued as soon as possible. See WARNINGS: Fetal/Neonatal Morbidity and Mortality.

DESCRIPTION

Diovan HCT is a combination of valsartan, an orally active, specific angiotensin II antagonist acting on the AT1 receptor Subtype, and hydrochlorothiazide, a diuretic. Valsartan, a nonpeptide molecule, is chemically described

as N-(1-oxopentyl)-N-[[2'-(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl]methyl]-L-Valine. Its empirical formula is $C_{24}H_{29}N_5O_{32}$ its molecular weight is 435.5, and its structural formula is



Valsartan is a white to practically white fine powder. It is soluble in ethanol and methanol and slightly soluble in water.

Hydrochlorothiazide USP is a white, or practically white, practically odorless, crystalline powder. It is slightly soluble in water; freely soluble in sodium hydroxide solution, in nin wear, needy southe in southin hydroxide southout, in the butylamine, and in dimethylformamide; sparingly soluble in methanol; and insoluble in ether, in chloroform, and in dilute mineral acids. Hydrochlorothiazide is chemically de-scribed as 6-chloro-3,4-dihydro-2H-1,2,4-benzothiadiazine-7-sulfonamide 1,1-dioxide. Hydrochlorothiazide is a thiazide diuretic. Its empirical formula is $C_7H_8ClN_3O_4S_2$, its molecular weight is 297.73, and its structural formula is [See chemical structure at top of next column]



Diovan HCT tablets are formulated for oral administration with a combination of 80 mg or 160 mg of valsartan and 12.5 mg of hydrochlorothiazide USP. The inactive ingredients of the tablets are colloidal silicon dioxide, crospovidone, hydroxypropyl methylcellulose, iron oxides, magnesium stearate, microcrystalline cellulose, polyethylene glycol, talc, and titanium dioxide.

CLINICAL PHARMACOLOGY Mechanism of Action

Angiotensin II is formed from angiotensin I in a reaction catalyzed by angiotensin-converting enzyme (ACE, kininase II). Angiotensin II is the principal pressor agent of the renin-angiotensin system, with effects that include vasoconstriction, stimulation of synthesis and release of aldos-terone, cardiac stimulation, and renal reabsorption of sodium. Valsartan blocks the vasoconstrictor and aldosterone secreting effects of angiotensin II by selectively blocking the binding of angiotensin II to the AT_1 receivery blocking the sues, such as vascular smooth muscle and the adrenal gland. Its action is therefore independent of the pathways for angiotensin II synthesis.

There is also an AT_2 receptor found in many tissues, but AT_2 is not known to be associated with cardiovascular homeo-stasis. Valsartan has much greater affinity (about 20,000fold) for the AT_1 receptor than for the AT_2 receptor. The primary metabolite of valsartan is essentially inactive with an affinity for the AT1 receptor about one 200th that of valsartan itself.

Blockade of the renin-angiotensin system with ACE inhibi-tors, which inhibit the biosynthesis of angiotensin II from angiotensin I, is widely used in the treatment of hypertension. ACE inhibitors also inhibit the degradation of brady-kinin, a reaction also catalyzed by ACE. Because valsartan does not inhibit ACE (kininase II) it does not affect the response to bradykinin. Whether this difference has clinical relevance is not yet known. Valsartan does not bind to or block other hormone receptors or ion channels known to be important in cardiovascular regulation.

Blockade of the angiotensin II receptor inhibits the negative regulatory feedback of angiotensin II on renin secretion, but the resulting increased plasma renin activity and angiotensin II circulating levels do not overcome the effect of valsar-

sin in circulating revers to her or retent the relation of the second research of the relation proximately equivalent amounts. Indirectly, the diuretic action of hydrochlorothiazide reduces plasma volume, with consequent increases in plasma renin activity, increases in aldosterone secretion, increases in urinary potassium loss, and decreases in serum potassium. The renin-aldosterone link is mediated by angiotensin II, so coadministration of an angiotensin II receptor antagonist tends to reverse the potassium loss associated with these diuretics.

The mechanism of the antihypertensive effect of thiazides is unknown.

Pharmacokinetics

Valsartan

R

Valsartan peak plasma concentration is reached 2 to 4 hours after dosing. Valsartan shows bi-exponential decay kinetics following intravenous administration, with an average elimination half-life of about 6 hours. Absolute bioavailability for the capsule formulation is about 25% (range 10%-35%). Food decreases the exposure (as mea-(range 10%-35%). Food decreases the exposure (as measured by AUC) to valsartan by about 40% and peak plasma concentration (C_{max}) by about 50%. AUC and C_{max} values of valsartan increase approximately linearly with increasing dose over the clinical dosing range. Valsartan does not accumulate appreciably in plasma following repeated administration administration

Metabolism and Elimination

Valsartan

Valsartan, when administered as an oral solution, is primarily recovered in feces (about 83% of dose) and urine (about 13% of dose). The recovery is mainly as unchanged drug, with only about 20% of dose recovered as metabolites. The primary metabolite, accounting for about 9% of dose, is valeryl 4-hydroxy valsartan. The enzyme(s) responsible for valsartan metabolism have not been identified but do not seem to be CYP 450 isozymes.

Following intravenous administration, plasma clearance of valsartan is about 2 L/h and its renal clearance is 0.62 L/h (about 30% of total clearance).

Hydrochlorothiazide

Hydrochlorothiazide is not metabolized but is eliminated rapidly by the kidney. At least 61% of the oral dose is eliminated as unchanged drug within 24 hours. The elimination half-life is between 5.8 and 18.9 hours. Distribution

Valsartan

The steady state volume of distribution of valsartan after intravenous administration is small (17 L), indicating that valsartan does not distribute into tissues extensively. Valsartan is highly bound to serum proteins (95%), mainly serum albumin.

Hydrochlorothiazide

Hydrochlorothiazide crosses the placental but not the blood-

NOVARTIS PHARMACEUTICALS/2017

Special Populations

Pediatric: The pharmacokinetics of valsartan have not

Geriatric: The pharmatoxinetics of valsartan have nor been investigated in patients <18 years of age. Geriatric: Exposure (measured by AUC) to valsartan is higher by 70% and the half-life is longer by 35% in the el-derly than in the young. No dosage adjustment is necessary (see DOSAGE AND ADMINISTRATION).

Gender: Pharmacokinetics of valsartan does not differ sig-nificantly between males and females.

Race: Pharmacokinetic differences due to race have not been studied.

Renal Insufficiency: There is no apparent correlation be tween renal function (measured by creatinine clearance) and exposure (measured by AUC) to valsartan in patients with different degrees of renal impairment. Consequently, dose adjustment is not required in patients with mild-to-moderate renal dysfunction. No studies have been performed in patients with severe impairment of renal function (creatinine clearance <10 mL/min). Valsartan is not removed from the plasma by hemodialysis. In the case of severe renal disease, exercise care with dosing of valsartan (see DOSAGE AND ADMINISTRATION).

Thiazide diuretics are eliminated by the kidney, with a terminal half-life of 5-15 hours. In a study of patients with impaired renal function (mean creatinine clearance of 19 mL/min), the half-life of hydrochlorothiazide elimination

was lengthened to 21 hours. Hepatic Insufficiency: On average, patients with mild-to-moderate chronic liver disease have twice the exposure (measured by AUC values) to valsartan of healthy volun-teers (method by are sense of the sens teers (matched by age, sex and weight). In general, no dos-age adjustment is needed in patients with mild-to-moderate liver disease. Care should be exercised in patients with liver disease (see DOSAGE AND ADMINISTRATION). Pharmacodynamics and Clinical Effects

Valsartan – Hydrochlorothiazide

In controlled clinical trials including over 1500 patients, 730 patients were exposed to valsartan (80 and 160 mg) and concomitant hydrochlorothiazide (12.5 and 25 mg). A factorial trial compared the combinations of 80/12.5 mg, 80/25 mg, 160/12.5 mg and 160/25 mg with their respective com-ponents and placebo. The combination of valsartan and hydrochlorothiazide resulted in additive placebo-adjusted de creases in systolic and diastolic blood pressure at trough of 15-21/8-11 mmHg at 80/12.5 mg to 160/25 mg, compared to 7-10/4-6 mmHg for valsartan 80 mg to 160 mg and 6-10/3-5 mmHg for hydrochlorothiazide 12.5 mg to 25 mg alone.

In another controlled trial the addition of hydrochlorothiazide to valsartan 80 mg resulted in additional lowering of systolic and diastolic blood pressure by approximately 6/3 and 12/5 mmHg for 12.5 mg and 25 mg of hydrochlorothiazide, respectively, compared to valsartan 80 mg alone.

The maximal antihypertensive effect was attained 4 weeks after the initiation of therapy, the first time point at which blood pressure was measured in these trials.

blood pressure was measured in these thans. In long-term follow-up studies (without placebo control) the effect of the combination of valsartan and hydrochlorothia-zide appeared to be maintained for up to two years. The antihypertensive effect is independent of age or gender. The overall response to the combination was similar for black and non-black patients.

There was essentially no change in heart rate in patients treated with the combination of valsartan and hydrochlorothiazide in controlled trials.

Valsartan

Valsartan inhibits the pressor effect of angiotensin II infu-sions. An oral dose of 80 mg inhibits the pressor effect by about 80% at peak with approximately 30% inhibition persisting for 24 hours. No information on the effect of larger doses is available.

Removal of the negative feedback of angiotensin II causes a 2- to 3-fold rise in plasma renin and consequent rise in angiotensin II plasma concentration in hypertensive patients. Minimal décreases in plasma aldosterone were observed after administration of valsartan; very little effect on serum potassium was observed.

In multiple-dose studies in hypertensive patients with sta-ble renal insufficiency and patients with renovascular hypertension, valsartan had no clinically significant effects on glomerular filtration rate, filtration fraction, creatinine clearance, or renal plasma flow.

In multiple-dose studies in hypertensive patients, valsartan had no notable effects on total cholesterol, fasting triglycerides, fasting serum glucose, or uric acid.

The antihypertensive effects of valsartan were demonstrated principally in 7 placebo-controlled, 4- to 12-week trials (one in patients over 65) of dosages from 10 to 320 mg/day in patients with baseline diastolic blood pressures of 95-115. The studies allowed comparison of once-daily and twice-daily regimens of 160 mg/day; comparison of peak and trough effects; comparison (in pooled data) of response by gender, age, and race; and evaluation of incremental effects of hydrochlorothiazide.

Administration of valsartan to patients with essential hypertension results in a significant reduction of sitting, supine, and standing systolic and diastolic blood pressure, usually with little or no orthostatic change.

In most patients, after administration of a single oral dose, onset of antihypertensive activity occurs at approximately 2 hours, and maximum reduction of blood pressure is achieved within 6 hours. The antihypertensive effect per-

BIOCON PHARMA LTD (IPR2020-01263) Ex. 1012, p. 005