

INTENTIALIBETS OF PRESCRIBING INFORMATION
These highlights do not include all the information needed to use DUXAZOSIN TABLETS safely and effectively. See ultip rescribing information for DUXAZOSIN TABLETS.

OUXAZOSIN TABLETS.

(S.3) DOXAZOSIN tablets, for oral use Initial U.S. Approval: 1990

Signs and symptom.
 (BPH)
 Treatment of Hypertension

---DOSAGE AND ADMINISTRATION

OSAGE AND ADMINISTRATION.
 For the treatment of BPH. Initiate therapy at 1 mg once daily. Dose may be titrated at 1 to 2 week intervals. up to 8 mg once daily. (2.2)
 For the treatment of hypertension: initiate therapy at 1 mg once daily. Dose may be titrated as needed, up to 16 mg once daily. (2.3)

------DOSAGE FORMS AND STRENGTHS• Tablets: 1 mg, 2 mg, 4 mg, 8 mg. See 17 for PATIENT CONTRAINDICATIONS
 See 17 for PATIENT COUNSELING INFORMATION and other ingredient in douazosin other quinazolines, or any FDA-approved patient labeling.

-WARNINGS AND PRECAUTIONS-----

(5.3)

The most commonly reported adverse reactions from clinical trials are fatigue, malaise, hypotension, and dizziness. (6.1)

INDICATIONS AND USAGE
Dozazoin tablets are an alpha, adrenergic antagonist indicated for (1)

\* Signs and symptoms of Benign Prostatic Hyperplasia (Pib at 1-800-919A-1086 or www.fda.gov/medivatch.

- PRUG INTERACTIONS

- Strong cytochrome P450 (CVP) 3A inhibitors may increase exposure to foraccosin and increased risk of hypotension. (7.1)

- Oncomittant administration of doxazosin tablets with a phosphodies terase-5 (PDE-5) inhibitor can result in additive blood pressure lowering effects and symptomatic hypotension. (7.2) USE IN SPECIFIC POPULATIONS
 Hepatic Impairment: Monitor for hypotension. (8.6, 12.3)

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FULL PRESCRIBING INFORMATION

1 NBICATIONS AND USAGE
1 Benigh Prostable (hyperplasta (BPH)
1.1 Benigh Prostable (hyperplasta (BPH)
1.2 Hypertension

Dozazola hables are indicated for the treatment of hypertension, to lower blood pressure. Lowering blood pressure reduces the risk of fatal and nordisal cardiovascular events, primarily strokes and myocardial infarctions. These beniffels have been seen in controlled trials of antihypertensive drugs from a vive exercise of the second controlled controlled in the controlled controlled in the controlled controlled in the controlled c

benefits have been seen in controlled trials of antihyperference frugs from a wide variety of pharmacologic classes, including this drug.

Including as a proposed pressure should be part of comprehensive cardionascular risk management, including, as apropriate, light control, diabetes management, and temporate the temporate proposed pressure posts. For specific advice sodium intake. Many patients will require more than one drug to achieve blood pressure goals. For specific advice on goals and management, see published guidelines, such as those of the National High Blood Pressure Education Programs. Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure Characterism frugs.

Program's Joint National Committee on Prevention, Detection, Febulation, and Treatment of High Blood Pressure LNUC.

NUCL SHIPPS returned from the Committee of The Pressure LNUCL SHIPPS and The Committee of The Pressure LNUCL SHIPPS and The Committee of The Com

2 DOSAGE AND OMINISTRATION
2 1 Dosling Information
Following the initial dose and with each dose increase of doxazosin tablets, monitor blood pressure for at least 6 hours following administration. If doxazosin tablets administration is discontinued for several days, therapy should be restated using the initial dosing regimen.
2 2 Benign Prostatic Hyperplasia
The economised initial dosing off doxazosin tablets is 1 mg given once daily either in the morning or severing. The economised initial dosago off towazosin tablets is 1 mg given once daily either in the morning or severing. The economised by the dozen any bet traited at 1 mg of the dozen and the traited at 1 mg or the days the dozen any bet traited at 1 mg or the days of the traited at 1 mg or the days of the traited at 1 mg or the days of the traited at 1 mg or the days of the days of the traited at 1 mg or the days of the days of the days of the traited at 1 mg or the days of the days of the traited at 1 mg or the days of the days of the days of the traited at 1 mg or the days of the days

week intervals to 2 mg, and unterconverting the many of the many o

Dozazosin Tablets, USP are available containing dozazosin mesylate, USP equivalent to 1 mg, 2 mg, 4 mg or 8 mg of dozazosin.

• The 1 mg tablets are available as white to off-white caplet-shaped tablets, debossed with "AC 356" on one side and scored on the other side.

• The 2 mg tablets are available as white to off-white round tablets, debossed with "AC" and "357" on the scored side and plain on the other side.

• The 4 mg tablets are available as white to off-white round tablets, debossed with "AC 356" on the scored side and plain on the other side.

• The 8 mg tablets are available as white to off-white caplet-shaped tablets, debossed with "AC 359" on one side and scored on other side.

• The 8 mg tablets are available as white to off-white caplet-shaped tablets, debossed with "AC 359" on one side and scored on other side.

The Use of dozazonin tablets is contraindicated in patients with a hypersensitivity to dozazosin, other quinazolines (e.g. parsystic levership) or any off is components.

4 CONTINUATION.

The use of doxazosini tablets is contraindicated in patients with a hypersensitivity to doxazosin, other quin. (e.g., prazosin, terazosin), or any of its components.

5 WARNINGS AND PRECAUTIONS

5 WARNINGS AND PRECAU UND
5 1 Postural Hypothesian without symptoms (e.g., dizziesa) may develop within a few hours following
Postural Hypothesian with white symptoms (e.g., dizziesa) may develop within a few hours following
the postural Hypothesian with the postural Hypothesian was to been reported
that the man few hours after dorsing. As with other alpha-blookers, there is a potiential for syncope, especially after the
initial does or after an increase in dosage strength. Advise patient how to avoid symptoms resulting from postural
hypothesian and what measures to base should they develop.
Concomitant administration of docazzasin tablets with a PDE-5 inhibitor can result in additive blood pressure lowering
affective and commontate throughtendors.

Concomitant administration or usual content of the content of the

of file blocks, iris dilator rings, or viscoelastic substances. There does not appear to be a benefit of stopping alpha, blocker therapy prior to catariast surgery. Observe the property of the product of the produc

3.4 Priagnosm
Alpha, antagonists, including doxazosin, have been associated with priapism (painful penile erection, sustained for hours and unrelieved by sexual intercourse or masturbation). This condition can lead to permanent impotence if not

hours and unrensembly promptly featible promptly

in practice.

rostatic Hyperplasia (BPH)

ence of adverse events has been ascertained from worldwide clinical trials in 965 BPH patients. The
rates presented below (Table 2) are based on combined data from seven placebo-controlled trials involving

once-daily administration of doxazosin tablets in doses of 1 mg to 16 mg in hypertensives and 0.5 mg to 8 mg in normotensives. Adverse reactions occurring more than 1% more frequently in BPH patients treated with doxazos

Table 1. Adverse Reactions Occurring more than 1% More Frequently in BPH Patients Treated with

BODY SYSTEM	Doxazosin tablets N = 665	Placebo N = 300
NERVOUS SYSTEM DISORDERS		
Dizziness*	15.6%	9.0%
Somnolence	3.0%	1.0%
CARDIAC DISORDERS		
Hypotension	1.7%	0%
RESPIRATORY, THORACIC AND	MEDIASTINAL DISORDERS	
Dyspnoea	2,6%	0.3%
GASTROINTESTINAL DISORDER	S	
Dry Mouth	1.4%	0.3%
GENERAL DISORDERS AND ADI	MINISTRATION SITE CONDITIONS	
Fatigue	8.0%	1.7%
Oedema	2.7%	0.7%
*Includes vertigo		

'Includes vertigo
Other adverse reactions occurring less than 1% more frequently in BPH patients treated with doxazosin tablets versus placebo but plausibly related to doxazosin tablets include: palpitations. 
\*\*Phypertension\*\*
Divazosin tablets have been administered to approximately 4000 hypertensive patients in clinical trials, or whom 1679 were included in the hypertension clinical development program. In placebo-controlled studies, adverse events occurred in 49% and 40% of patients in the doxazosin and placebo groups, respectively, and led to discontinuation in 2% of patients in each group.

Adverse reactions occurring more than 1% more frequently in hypertensive patients returned with divazosin discontinuation in 2% of patients in each group.

Adverse reactions occurring more than 1% more frequently in Supretensive patients returned with divazosin discontinuation in 2% of patients in each group.

Adverse reactions occurring more than 1% more frequently in Supretensive patients returned with divazosin discontinuation and supported to the patients of the supported to the supported to the contribution of the supported to the

# Table 2. Adverse Reactions Occurring more than 1% More Frequently in Hypertensive Patients Treated with Doxazosin Tablets versus Placebo

BODY SYSTEM	Doxazosin tablets N = 339	Placebo N = 336
NERVOUS SYSTEM DISORDERS		
Dizziness	19%	9%
Somnolence	5%	1%
RESPIRATORY, THORACIC AND	MEDIASTINAL DISORDERS	
Rhinitis	3%	1%
RENAL AND URINARY DISORDE	RS	
Polyuria	2%	0%
REPRODUCTIVE SYSTEM AND E	BREAST DISORDERS	
GENERAL DISORDERS AND ADI	MINISTRATION SITE CONDITIONS	
Fatigue / Malaise	12%	6%

Other adverse reactions occurring less than 1% more frequently in hypertensive patients treated with docazonis habites versus placebo but plausibly related to docazonis habites use include verifigo, hypotension, of fullables, pletiska and oederma. Docazonis habites have been associated with decreases in white blood cell counts. *Laboratory Charge Observed in Clinical Studies* 

Lanksgenia Weufopenia
Decrasses in mean white blood cell (WBC) and mean neutrophil count were observed in controlled clinical
trials of hyportensive patients receiving doxazosin tablets. In cases where follow-up was available, WBC
and neutrophil counts returned to normal after discontinent of doxazosin tablets. No patients became
symptomatic as a result of the low WBC or neutrophil counts.

and neutrophil counts returned to normal after discontinuation of dovazasen tablets. No patients is symptomatic as a result of the low Wide Cor neutrophil county.

6.2 Postmarketing Experience.

6.2 Postmarketing Experience.

File following advances reactions have been identified during post-approval use of dovazaoin procession of the procession of t

Eye Disorders: Intraoperative Floopy Iris Syndrome [see Wamings and Precautions [5:2]]: Cardina Disorders: bandycardia: Respiratory, Thoracic and Mediastinal Disorders: bronchospasm aggravated; Gastrointestinal Boorders: vomiting: Hepatobiliary Disorders: cholestais, hepatitis oblisatic; Skin and Subuctaneous Tissue Disorders: utricaria; Musculoskeletal and Connective Tissue Disorders: muscle cramps, muscle weakness; Renal and Urinary Disorders: hematicin, inclurition disorder, inclurition frequency, nocturia; Reproductive System and Breast Disorders: gynecomastia, priapism.

# 7 DRUG INTERACTIONS

CPPA Inhibitors

"irrd studies suggest that doxazosin is a substrate of CPP34A. Strong CPP3A inhibitors may increase source to doxazosin. Monitor blood pressure and for symptoms of hypotension when doxazosin tablets used concomitantly with strong CPP34 inhibitors. See Clinical Pharmacology (12.3). Phosphodiesterase-5 (PDE-5) Inhibitors committed administration of doxazosin tablets with a phosphodiesterase-5 (PDE-5) Inhibitor can result additive blood pressure lowering effects and symptomatic hypotension. Monitor blood pressure and for rightons of hypotension few Warmigs and Procautions (5.1).

## R LISE IN SPECIFIC POPULATIONS

A USE IN SPECIFIC POPULATIONS
8.1 Frequancy
Risk Summary
The limited available data with doxazosin tablets in pregnant women are not sufficient to inform a drugassociated risk for major brith identits and miscarriage. However, untreated hypertension during pregnancy
processes and the second of the control of the contr

Hyperte...
Data
\*\*rimal Data
\*\*tivity

Animal Data Rediacativity was found to cross the placenta following oral administration of labelled doxazosin to pregnant rats. Studies in pregnant rabbits and rats at daily and doses of up to 41 mg/kg and 20 mg/kg, respectively, lepisam drug occentrations of 10 and 4 times, respectively, the human AUC exposures with a 12 mg/day therapeutic dose), have revealed no evidence of adverse developmental effects. A dosage regimen of 82 mg/kg/day in the rabbit was associated with reduced fetal survival. In peri – and postnatal statics in ast, postnatal development at maternal doses of 40 mg/kg/day or 50 mg/kg/day of doxazosin jabout 8 times human AUC exposure with a 12 mg/dg/da therapeutic dosely was delayed, as evidenced by slower body weight gain and slightly later appearance of anatomical features and reflexes.

Risk Summary

There is limited information on the presence of doxazosin in human milk (see Data). There is no infor on the effects of doxazosin on the breastfed infant or the effects on milk production.

Data

tar ingle case study reports that doxazosin is present in human milk, which resulted in an infant dose of less in 1% of the maternal weight-adjusted dosage and a milk/plasma ratio of 0.1. However, these data are ufficient to confirm the presence of doxazosin in human milk.

than 1% of the missions and insufficient to ordinite the presence of doxazosin in human mins.

8.4 Pediatric Use
The safety and effectiveness of doxazosin tablets have not been established in children.

8.5 Geratric Use
Benign Prostatic Hyperplasia (BPH)
The safety and effectiveness profile of doxazosin tablets was similar in the elderly (age > 65 years) and younger (age > 65 years) patients.

\*\*Americancian\*\*

The Safety and effectiveness profile or douacoast towards and the safety and effectiveness profile or douacoast towards. Hypertransical Clanical studies of douacoast habets did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects. Other reported clinical experience has located in the safety of the safety

VILLIAMONIA
erince with Occazosin tablets overdosage is limited. Two adolescents, who each intentionally ingested
mg doxazosin tablets with diclofenac or acetaminophen, were treated with gastric lavage with activated
cost and made full recoveries. A two-year-old orbit who accidently ingested 4 mg doxazosin tablets
cost leaded with gastric lavage and remained normothersive during the five-hour ememerancy reven

observation period. A six-month-old child accidentally received a crushed 1 mg tablet of doxazoist tablets and was reported to have been drowsy. A 25-year old female with chronic relevance of the property o

In IESCRIPTION

Dozazoni tablets, USP are a quinazoline compound that is a selective inhibitor of the adpha, subtype of alpha-adrenergic receptors. The chemical name of dozazoni mesylate is 1-(4-Amino-6.7-dimethoxy-2-quinazoliny)-4-(1.4-benzodioxan-2-ylcazbonyl) piperazine methanesustronac. The molecular romale for dozazonis mesylate is C<sub>21</sub>H<sub>21</sub>N<sub>2</sub>0, • CH<sub>1</sub>0,S and the molecular weight is 547.6. It has the following structure:

Doxazosin mesylate, USP is freely soluble in dimethylsulfoxde, soluble in dimethylformamide, slightly soluble in methanol, ethanol, and water (50% at 25°C), and every slightly soluble in use and contains in mig. 2 ma. 4 may and 8 mg of doxazosin as the free base. The inactive ingredients for all bables are microcystalline cellulose, anhydrous lactose, sodium starch glycottle, magnesium stearest and sodium lauty cellulose.

Iz CLINICAL PHRAIMACOLORY

12.1 Michanism of Action

12.1 Michanism of Action

13.1 Michanism of Action

14.1 Michanism of Action

The symptoms associated with benign prostatic hyperplasia (BPH)

The symptoms associated with benign prostatic hyperplasia (BPH), such as urinary frequency, nocturia, weak stream, hesitancy, and incomplete emptying are related to two components, anatomical (static) and functional (dynamic). The static component is related to an increase in prostate size caused, in part. by a proliferation of smooth muscle cells in the prostatic strona. However, the severity of BPH symptoms and the degree of urethral obstruction of not correlate well with the size of the prostatic. The dynamic component of BPH is associated with a correlate with the size of the prostatic. The dynamic component of BPH is associated with a case is mediated by the alpha, adrenoceptor, which is present in high density in the prostatic strona, prostatic cassile and bladed encel. Blockade of the alpha, receptor decreases urethral resistance and may relieve the obstruction and BPH symptoms and improve urine flow.

Hypertansion

resistance and may relieve the obstruction and BPH symptoms and improve urine flow. Hypertension

The mechanism of action of doxozosin tables is selective blockade of the alpha, (postjunctional) subtype of adversept; exceptions. Studies in normal human subjects have shown that doxozosin subtype of adversept; exceptions. Studies in normal human subjects have shown that doxozosin systolic presor effect of nonepinephrine. Doxozosin and prozosin have similar abilities to adaptance phenylephrine. The antihypertensive effect of doxozosin tables results from a decrease in systemic vascular resistance. The parent compound doxozosin is primarily made to the parent compound doxozosin of doxozosin control to the compound compound compound compound to doxozosin (primary low for a fixed principles of the protection of even the most potent compound compound to the parent drug indicate that the contribution of even the most potent from the primary fixed protection of specific proposity small. The 6'- and 7'-hydroxy meta-olites have demonstrated antioxidant properties at concentrations of 5 pl.M. in vite.

Benign Prostatic Hyperplasia (BPH)
Administration of doxazosin tablets to

Administration of doizazoish tablets to patients with symptomatic BHH resulted in a statistically significant improvement in maximum timms flow rate (see Ginicina Studies (14.1). Effect on Normotensive Patients with Beeliga Prostatic Hyperplasia (BHH). Although blockade of alpha, ademoceptors also lowers blood pressure in hypertensive patients with increased peripheral vascular resistance, dozozon tablets treatment of normotensive men with BHH dot not result in a clinically significant blood pressure less than 95 mm/leg and/of distatic blood pressure less than 95 mm/leg and of distance that 10 mm/leg and 10 mm/leg

ministration of doxazosin tablets results in a reduction in systemic vascular resistance. In lemts with hypertension, there is little change in cardiac output. Maximum reductions in blood sisure usually occur 2 to 6 hours after dosing and are associated with a small increase in inding heart rate. Like other alpha, advernency blocking agents, doxazosin has a greater effect blood pressure and heart rate in the standing position. 3 Pharmacokinetics

# 12.3 Pha

12.3 Pharmacokinetics
Absorption
Alter oral administration of therapeutic doses, peak plasma levels of doxazosin tablets occur at douct 2 to 3 hours. Boavailability is approximately 65%, reflecting first-pass metabolism of doxazoain by the liver. The effect of food on the pharmacokinetics of doxazoain tablets of doxazoain tablets of doxazoain tablets of doxazoain tablets are maintained plasma concentration (Lg.) and 12% in the rare under the concentration-time curve (AILO) cocurred when doxazoain tablets were administered with food. Neither of these differences is clinically significant.
In a crossover study in 24 months of the pharmacokinetics and safety of in a crossover study in 24 months of the pharmacokinetics, and safety of in a crossover study in 24 months of the pharmacokinetics, and safety of in a crossover study in 24 months of the pharmacokinetics, and safety of in a crossover study in 24 months of the pharmacokinetics and safety of control of the pharmacokinetics. Nowever, 11% less than that after exempl dosing and time to peak concentration after evening dosing occurred significantly later than that after morning dosing the lime to peak CR versus 3.5 hours, lower than the safety of the crossover study of the

stabolism account in the process of the fiver, mainly by 0-demethylation of the nazonine nucleus or hydroxylation of the benzodioxan molety. In vitro studies suggest that primary pathway for elimitation is via CVP3441, however, CVP2D6 and CVP22G9 metabolic however as also involved to a lesser extent. Although several active metabolities of oxozosin to be ben defittified, the pharmacokinetics of these metabolities have not been characterized.

pathways are also involved to a lesser extern. Although several active metabolites of doxazosin have been identified, the pharmacokinets of these metabolities have not been characterized. Excretion

Pleama elimination of doxazosin is biphasic, with a terminal elimination half-life of about 22 hours. Steady-state studies in hypertensive patients given doxazosin doses of 2 mg to 16 mg once daily showed linear kinetics and dose proportionality. In two states, following the grant opinion of the proportion of the p

natric pharmacokinetics of doxazosin tablets in young (< 65 years) and elderly ( $\succeq$  65 years) givens were similar for plasma half-life values and oral clearance. all Inpairment is

nal impairment rimacokinetic studies in elderly patients and patients with renal impairment have shown no nifficant alterations compared to younger patients with normal renal function. patic impairment

Hepatic Impairment
Administration of a single 2 mg dose to patients with cirrhosis (Child-Pugh Class A) showed a
40% increase in exposure to doxazsisi. The impact of moderale (Child-Pugh Class S) or severe
(Indi-Pugh Class S), hepatic impairment on the pharmacokinetics of doxazosin is not known
(see Use in Specific Psyculations (8.6)). The properties of the properties o

doxazosin (e.g., cimetidine).

metidine
healtity volunteers, the administration of a single 1 mg dose of doxazosin on day 1 of a fourty regimen of oral cimetidine (400 mg twice daily) resulted in a 10% increase in mean AUC of

the company of th ייניים עוברים ע

of digoxin, varfarin, phenytoin, or indomethacin.

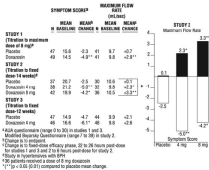
13 NONCLINICAT TOXICOLOGY
13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility
13.1 Carcinogenesis, Mutagenesis. Chronic dietary administration (up to 24 months) of doxazosin mesylate at maximally tolerated doses of 40 mg/kg/dgy in rats and 120 mg/kg/dgy in mice revealed no evidence of carcinogenesis engles of the service of the control o



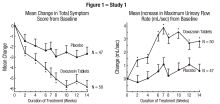
13.2 Animal Toxicology and Pharmacology
An increased incidence of myocardial necrosis of fibrosis was observed in long-term (6 to 12 months) studies in rats and mice reposure 8 times human ALIC exposure in rats and somewhat equivalent to human  $C_{\rm cons}$  exposure in mice). Findings were not seen at lower doses. In dogs no cardiobxicity was observed following 21 months of ord dosing at doses that resided in maximum plasma concentrations  $C_{\rm cons}$  14 mems the  $C_{\rm cons}$  exposure in humans receiving a 12 mg/day therepeaks close or in Wistar rats at  $C_{\rm cons}$  exposures 15 times human  $C_{\rm cons}$  opens. There is no evidence that similar desires occur in humans.

Internal C<sub>max</sub> proposes. There is no evidence that similar inclusions occur in humans can be produced to the control of the c

### Table 3. SUMMARY OF EFFECTIVENESS DATA IN PLACEBO-CONTROLLED TRIALS



"(")» c 0.68 (0.01) compared to plezebo mean change in one fixed-to-set subty (Stuby 2), obazzosin labellet therapy (4 mg to 8 mg, once daily) resulted in a significant and sustained improvement in maximum urinary flow rate of 2.3 mL/sec to 3.3 mL/sec (1861) 3 compared to blacebo (0 1 mL/sec), in this study, the only shad, in which verely evaluations were made, and the study of the study of the study of the study in the study of the study in the study of the study in the study of t



red to Placebo; + p < 0.05 Compared to Baseline; Doxazosin Titration to Maximum of 8 mg.

124 Hypertension
In a poided analysis of latedob controlled hypertension studies with about 300 hypertensive polients per
In a poided analysis of latedob controlled hypertension studies with about 300 hypertensive polients for
In In It is many priven store daily, lovered blood pressure at 24
In In In It is many polient and blood pressure at 24
In In In It is many polient and blood pressure effects (1 to 6 hours) were larger by about 50% to 75% (6. through values were
about 55% to 76% opeak effect, with the larger peak-trough differences seen in systolic pressures. There
was no apparent difference in the blood pressure response of Caucasians and blacks or of patients above
and below age 65. In the same patient polypation, patients receiving dovazosin tablets gained a mean of 0.6
kg compared to a mean loss of 0.1 kg for placebo patients.

TABLE 4

TABLE 4

TABLE 4

TABLE 5

TO THE TABLE 5

TABLE 7

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## Mean Changes in Blood Pressure from Baseline to the Mean of the Final Efficacy Phase in sives (Diastolic BP <90 mmHg) in Two Double-blind, Placebo-controlled U.S. Studies with

PLACEBO (N=85)		Doxazosin tablets (N=183)						
Sitting BP (mmHg)	Baseline	Change	Baseline	Change				
Systolic	128.4	-1.4	128.8	-4.9*				
Diastolic	79.2	-1.2	79.6	-2.4*				
Standing BP (mmHg)	Baseline	Change	Baseline	Change				
Systolic	128.5	-0.6	128.5	-5.3*				
Diastolic	80.5	-0.7	80.4	-2.6*				
to <0.0E compared to a	Jacobo							

16 HOW SUPPLIED/STORAGE AND HANDLING
Divazosin Tablets, USP are available as tablets for oral administration. Each tablet contains dovazosin
meystee. USP quivalent to 1 mg. 2 mg. 4 mg or 8 mg of dovazosin.
The 1 mg tablets are available as white to off-white capiet-shaped tablets, debossed with "AC 356" on one side and scored on the other side. They are supplied as follows:
Bottles of 100 tablets
1000 0832-0856-1
Bottles of 500 tablets
1000 0

The 2 mg tables are available as white to off-white round tablets, debossed with "AC" and "357" on the sorred side and plain on the other side. They are supplied as follows:

NOC 0832-0857-10

Bottles of 500 tablets

NOC 0832-0857-15

Bottles of 500 ballets
NDC 0825-0857: The 4 mg tablets are available as white to off-white round tablets, debossed with "AC 359" on the scored side and plain on the other side. They are supplied as follows:
NDC 0822-0359: 10
NDC 0822-0359-11

nmended Storage: Store at 20° to 25°C (68° to 77°F) [See USP Controlled Room Temperature.]

17 PATENT COUNSELING INFORMATION

Advise the patient for read the FDA-approved patient labeling (Patient Information).

Advise the patient for read the FDA-approved patient labeling (Patient Information) at the initiation of therap

Advise patients of the possibility of synopopal and orthostatic symptoms, especially at the initiation of therap

and urged to avoid driving or hazardous tasks for 24 hours after the first dose, after a dosage increas

and after interruption of therapy when treatment is resumed. Advise patients to report symptoms to the
healthcare provider.

Distributed by
UPSHER-SMITH LABORATORIES, LLC
Maple Grove, MN 55369
200201 Revised 1217

Patient Information Doxazosin Tablets, USP (dox-AZE-oh-sin)

## What are doxazosin tablets?

Doxazosin tablets are a prescription medicine that contain doxazosin mesylate and are called an "alpha-blocker" Doxazosin tablets are used to treat:

the symptoms of benign prostatic hyperplasia (BPH)

high blood pressure (hypertension)

It is not known if doxazosin tablets are safe and effective

## Who should not take doxazosin tablets? Do not take doxazosin tablets if you:

 are allergic to doxazosin, other quinazolines, or any of the ingredients in doxazosin tablets. See the end of this Patient Information leaflet for a complete list of ingredients in doxazosin tablets

### What should I tell my healthcare provider before taking doxazosin tablets? Before taking doxazosin tablets, tell your healthcare provider about all of your medical conditions, including if you:

have had low blood pressure, especially after taking other medicine. Signs of the low blood pressure include fainting, dizziness, and lightheadedness.

have any planned eye surgery
have prostate cancer or a history of prostate cancer. Your healthcare provider may have you checked for prostate cancer before you start taking and while you take doxazosin tablets.

have liver problems

are pregnant or plan to become pregnant. It is not known

if doxazosin will harm your unborn baby.

are breastfeeding or plan to breastfeed. It is not known if doxazosin passes into your breastmilk.
Talk to your healthcare provider about the best way to

feed your baby if you take doxazosin tablets

## Tell your healthcare provider about all the medicines you take, including prescription and over-the-counter medicines, vitamins, and herbal supplements. Doxazosin tablets may affect the way other medicines work, and other medicines may affect the way doxazosin tablets work causing side effects.

Especially tell your healthcare provider if you take:

other medicine for high blood pressure, medicine to treat erectile dysfunction (ED) called a phosphodiesterase type 5 (PDE-5) inhibitor. The use of doxazosin tablets with PDE-5 inhibitors can lead to a drop in blood pressure or to fainting.

Know the medicines you take. Keep a list of them to show your healthcare provider and pharmacist when you get a

## How should I take doxazosin tablets?

 Take doxazosin tablets exactly as your healthcare provider tells you to take it.

Your healthcare provider will tell you how many doxazosin tablets to take and when to take them.

Your healthcare provider may need to change your dose of doxazosin tablets until it is the right dose for you.

## What should I avoid while taking doxazosin tablets?

Do not drive or perform any hazardous task until at least 24 hours after you have taken doxazosin tablets if you are taking:

your first dose of doxazosin tablets

 Doxazosin tablets for the first time after your healthcare provider has increased your dose of doxazosin tablets

Doxazosin tablets for the first time after any breaks

(interruptions) in your treatment with doxazosin tablets

### What are the possible side effects of doxazosin tablets?

## Doxazosin tablets may cause serious side effects, including:

- A sudden drop in blood pressure, especially when you first start treatment or when there is an increase in your dose of doxazosin tablets, is common but can also be serious. This may cause you to faint, or to feel dizzy or lightheaded. Your risk of having this problem may be increased if you take doxazosin tablets with certain other medicines that lower blood pressure including PDE-5 inhibitors. Your healthcare provider may monitor your blood pressure while you take doxazosin tablets. See "What should I avoid while taking doxazosin tablets?
- Eye problems during cataract surgery. A condition called Intraoperative Floppy Iris Syndrome (IFIS) can happen during cataract surgery if you take or have taken alpha-blockers such as doxazosin tablets. If you need to have cataract surgery, be sure to tell your healthcare provider if you take or have taken doxazosin tablets.
- A painful erection that will not go away. Doxazosin tablets can cause a painful erection (priapism), which cannot be relieved by having sex. If this happens, get medical help right away. If priapism is not treated, you may not be able to get an erection in the future.

The most common side effects of doxazosin tablets are:

- weakness or lack of energy (asthenia)
- dizziness

Tell your healthcare provider if you have any side effect that bothers you or that does not go away. These are not all the possible side effects of doxazosin tablets. For more information, ask your healthcare provider or pharmacist.

Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

### General information about the safe and effective use of doxazosin tablets.

Medicines are sometimes prescribed for purposes other than those listed in a Patient Information leaflet. Do not used oxazosin tablets for a condition for which it was not prescribed. Do not give doxazosin tablets to other people, even if they have the same symptoms you have. It may harm them.

This Patient Information leaflet summarizes the most important information about doxazosin tablets. For more information, ask your healthcare provider. You can ask your healthcare provider or pharmacist for information that is written for healthcare professionals.

## What are the ingredients in doxazosin tablets?

Active ingredient: doxazosin mesylate

Inactive ingredients: microcrystalline cellulose, anhydrous lactose, sodium starch glycolate, magnesium stearate and sodium lauryl sulfate.

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For more information, go to www.upsher-smith.com or call 1-888-650-3789.

This Patient Information has been approved by the U.S. Food and Drug Administration.

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