Approval Package for:

Application Number: 019777/S032

Trade Name: ZESTRIL TABLETS

Generic Name: Lisinopril Tablets

Sponsor: Zeneca Pharmaceuticals

Approval Date: January 8, 1997

\mathbf{A}	D	plication I	Number	019777/S032	

APPROVAL LETTER

Public Health Service '

Rockville MD 20857

Food and Drug Administration



NDA 19-777/S-032

JAN 8 1997

Zeneca Pharmaceuticals
Attention: William J. Kennedy, Ph.D.
1800 Concord Pike
P.O. Box 15437
Wilmington, DE 19850-5437

Dear Dr. Kennedy:

Please refer to your December 3, 1996 supplemental new drug application (NDA) submitted under section 505(b) of the Federal Food, Drug, and Cosmetic Act for Zestril (lisinopril) 2.5, 5, 10, 20, and 40 mg Tablets.

The supplemental application provides for final printed labeling revised as follows:

CLINICAL PHARMACOLOGY, Pharmacodynamics and Clinical Effects, Heart Failure: The last sentence has been revised to add the following to the end of this subsection:

The once daily dosing for the treatment of congestive heart failure was the only dosage regimen used during clinical trial development and was determined by the measurement of hemodynamic response.

CLINICAL PHARMACOLOGY, Pharmacodynamics and Clinical Effects, Acute Myocardial Infarction: As we requested in our letter dated November 24, 1995, in the first sentence of the fourth paragraph, the word "a" has been revised to "the" in the phrase "consisting of the number of patients who had"

WARNINGS, Anaphylactoid and Possibly Related Reactions, Anaphylactoid reactions during membrane exposure: The phrase " (a procedure dependent upon devices not approved in the United States)" has been deleted, as we requested in our facsimile dated March 8, 1996.

WARNINGS, Hypotension: As we requested in our letter dated November 24, 1995, "e.g.," has been added before "systolic blood pressure of 100 mm Hg or lower" in the second sentence of the fourth paragraph.

PRECAUTIONS, Pediatric Úse: The word "children" has been replaced with "pediatric patients" as required by the regulations on the revision of the Pediatric Use subsection of the labeling [21 CFR 201.57 (f)(9)].

HOW SUPPLIED: To comply with the CDER Stability Committee Uniform Storage Statement, the first sentence of the fourth paragraph has been revised to "Store at controlled room temperature, 20-25° C (68-77° F) [see USP].

We have completed the review of this supplemental application and have concluded that adequate information has been presented to demonstrate that the drug is safe and effective for use as recommended in the final printed labeling included with your December 3, 1996 submission. Accordingly, the supplemental application is approved effective on the date of this letter.

We remind you that you must comply with the requirements for an approved NDA set forth under 21 CFR 314.80 and 314.81.

If you have any questions, please contact:

Ms. Kathleen Bongiovanni Regulatory Health Project Manager (301) 594-5334

Sincerely yours,

Raymond J. Lipicky, M.D.
Director
Division of Cardio-Renal Drug Products
Office of Drug Evaluation I
Center for Drug Evaluation and Research

\mathbf{A}	D	plication	Number	019777/S032

FINAL PRINTED LABEL

F.P.O. Pharmacoo supplied by IPR

PROFESSIONAL INFORMATION BROCHURE

ESTRII®

USE IN PREGNANCY

When used in pregnancy during the second and third frimesters, ACE inhibitors can cause injury and even death to the developing felus. When pregnancy is detected, ZESTRIL should be discontin-ued as soon as possible. See WARNINGS, Fetal/Neonatal Morbidity and Mogtality.

DESCRIPTION
Lisinoprii is an oral long-acting angiotensin converting enzyme inhibitor. Lisinoprii, a synthetic peptide derivative, is chemically described as (S)-1-f/N^{-C}-(T-Carboxy-3-phenylpropry)-1-ypyll-propried difydrate. Its empirical formula is C2₁H₃,N₃O₅-2H₂O and its structural formula is:

Lisinopril is a white to off-white, crystalline powder, with a molecular weight of 441.53. It is soluble in water and sparingly soluble in methanol and practically insoluble in ethanol.

ZESTRIL is supplied as 2.5 mg, 5 mg, 10 mg, 20 mg and 40 mg tablets for oral administration.

2.5 mg tablets - calcium phosphate, magnesium stearate, mannitol,

staton. 5,10 and 20 mg tablets - calcium phosphate, magnesium stearate, mannitol, red ferric oxide, starch. 40 mg tablets - calcium phosphate, magnesium stearate, mannitol, starch, yellow terric oxide.

starch, yellow ferric oxide.

CLINICAL PHARMACOLOGY

Mechanism of Action: Lisinopril inhibits angiotensin converting enzyme (ACE) in human subjects and animals. ACE is a pepticyl dipeptidase that catalyzes the conversion of angiotensin it to the vascoonstrictor substance, angiotensin 11. Angiotensin II also stimulates aldosterone secretion by the adrenal cortex. The beneficial effects of lisinopril in hypertension and heart failure appear to result primarily from suppression of the renin-angiotensin-aldosterone system. Inhibition of ACE results in decreased plasma angiotensin II which leads to decreased vasooressor activity and to decreased of soft of the renin-arity decrease in a small increase of serum potassium. In hypertensive patients with normal renal function treated with ZEFRIL alone for up to 24 weeks, the mean increase in serum potassium was approximately 0.1 mEQL; however, approximately 9% had a decrease greater than 0.5 mEQL in the study, patients treated with ZEFRIL and bydrochlorothizative for up to 24 weeks and a mean decrease in serum potassium of 0.1 mEQL; approximately 9% had a decrease greater than 0.5 mEQL in the same study, patients treated with ZEFRIL and bydrochlorothizative for up to 24 weeks had a mean decrease in serum potassium of 0.1 mEQL; approxiweeks had a mean decrease in serum potassion of 0.1 mEq.C. approximately 4% of patients had increases greater than 0.5 mEq.C. approximately 2% had a decrease greater than 0.5 mEq.C. See PRE-CAUTIONIS). Removal of angiotensin II negative feedback on renin secretion leads to increased plasma renin activity.

secretion leads to increased plasma renin activity.

ACE is identical to kininase, an enzyme that degrades bradykinin.
Whether increased levels of bradykinin, a potent vasodepressor peptide, play a role in the therapeutic effects of ZESTRIL remains to be elucidated. While the mechanism through which ZESTRIL lowers blood pressure is believed to be primarily suppression of the renin-angiotensin-aldosterone system, ZESTRIL is anilhypertensive even in patients with low-renin hypertension. Although ZESTRIL was anithypertensive in all races studied, black hypertensive patients (usually a low-renin hypertensive population) had a smaller average response to monotherapy than nonblack patients. Concomitant administration of ZESTRIL and Although advisorition that the reduced blood pressure in black and nonblack patients and any racial differences in blood pressure response were no longer evident.

differences in blood pressure in block and blooming patients and any racial differences in blood pressure response were no longer vide differences in blood pressure response were no longer vide differences in blood pressure response were no longer vide patients. Patramacokinetics and Metabolism: Following oral administration of ZESTRIL, peak serum concentrations in acute myocardial infarction patients. Declining serum concentrations are acute myocardial infarction patients. Declining serum concentrations exhibit a prolonged terminal phase which does not contribute to drug accumulation. This terminal phase probably represents saturable binding to ACE and is not proportional to dose. Lisinopril does not appear to be bound to other serum proteins. Lisinopril does not appear to be bound to other serum proteins. Lisinopril does not undergo metabolism and is excreted unchanged entirely in the urine. Based on urinary recovery, the mean extent of absorption of lisinopril is approximately 25%, with large intersubject variability (6%-60%) at all doses tested (5-80 mg). Lisinopril absorption is not influenced by the presence of food in the gastrointestinal tract. The absolute biographability of lisinopril is reduced to 16% in patients with stable NYHA Class II-IV congestive heaf failure, and the volume of distribution appears to be slightly smaller than that in normal subjects. The oral bioavailability of lisinopril in patients with acute myocardial infarction is similar to that in healthy volunteers.

Upon multiple dosing, lisinopril exhibits an effective half-life of accumulation of 12 hours.

Impaired renal function decreases elimination of listinopril, which is excreted principally through the kidneys, but this decrease becomes clinically important only when the glomentar filtration rate is below 30 mLmin. Above this glomerular filtration rate, the elimination hati-title is little changed. With greater impairment, however, peak and trough lisinopril levels increase, time to peak concentration increases and time to attain steady state is prolonged. Older patients, on average, have (approximately doubled) higher blood levels and the area under the plasma concentration time curve (AUC) than younger patients. (See DOSAGE AND ADMINISTRATION.) Listinoptil can be removed by hemodalysis. Studies in rats indicate that lisinopril crosses the blood-brain barrier poorty. Multiple doses of listinoptil in rats do not result in accumulation in

Stotes in rats involvate that instruction it coacts are to encourage and proofy. Multiple doses of lisinoprii in rats do not result in accumulation in any tissues. Milk of lactating rats contains radioactivity tollowing administration of VE (Isinoprii B, whole body autoradiography, radioactivity was found in the placenta following administration of labeled drug to pregnant rats, but now was found in the lettues.

Pharmacodynamics and Clinical Effects
Hypertension: Administration of ZESTRIL to patients with hypertension

Hyperfension: Administration of ZESTRL to patients with hypertension results in a reduction of both supine and standing blood pressure to about the same extent with no compensatory tachycardia. Symptomatic postural hypotension is usually not observed although it can occur and should be anticipated in volume and/or salt-depleted patients. (See WARNINGS.) When given logether with thiazide-type diuretics, the blood pressure lowering effects of the two drugs are approximately additive. In most patients studied, onsel of antihypertensive activity was seen at one hour after oral administration of an individual dose of ZESTRL, with peak reduction of blood pressure schieved by 6 hours. Although an antihypertensive effect was observed 24 hours after dosing with recommended single daily doses, the effect was more consistent and the mean effect was considerably larger in some studies with doses of 20 mg or more than with lower doses. However, at all doses studied, the mean antihypertensive effect was substantially smaller 24 hours after dosing than it was 6 hours after dosing than it

after dosing.
In some patients achievement of optimal blood pressure reduction may require two to four weeks of therapy.
The antihypertensive effects of ZESTRIL are maintained during long-term therapy. Abrupt withdrawal of ZESTRIL has not been associated with a rapid increase in blood pressure, or a significant increase in blood pres-sure compared to pretreatment levels.

e-response studies utilizing a once daily regimen were con-Two dose-response studies utilizing a once daily regimen were conducted in 438 mild to moderate hypertensive patients not on a diuretic. Blood pressure was measured 24 hours after dosing. An antihypertensive effect of ZESTRIL was seen with 5 mig in some patients. However, in both studies blood pressure reduction occurred sooner and was greater in patients treated with 10, 20 or 80 mg of ZESTRIL. In controlled clinical studies, ZESTRIL 20-80 mg has been compared in patients with mild to moderate hypertension to hydrochlorothizaide 12.5-50 mg and with atenol 65-20 mg, and in patients with moderate to severe hypertension to metoprotol 100-200 mg. It was superior to hydrochlorothizaide in effects on systolic and diastolic pressure in a population that was 3/4 caucasian. ZESTRIL was approximately equivalent to atenold and metoprotol in effects on diastolic blood pressure, and had somewhat greater effects on systolic blood pressure. Leasant, ESTINIT, was approximately equivalent to apenior and interpritor in effects on diastoric blood pressure, and had somewhat greater effects on systolic blood pressure.

ZESTRIL had similar effectiveness and adverse effects in younger and older (> 85 years) patients. It was less effective in blacks than in

and older (> 65 years) patients. It was less effective in blacks than in caucacians.

In hemodynamic studies in patients with essential hypertension, blood pressure reduction was accompanied by a reduction in peripheral arterial resistance with little or no change in cardiac output and in heart rate. In a study in nine hypertensive patients, following administration of ZESTRIL, there was an increase in mean renal blood flow that was not significant. Data from several small studies are inconsistent with respect to the effect of isinopril on glomerular filtration rate in hypertensive patients with normal renal function, but suggest that changes, if any, are not large. In patients with renovascular hypertension ZESTRIL has been shown to be well tolerated and effective in controlling blood pressure. (See PRECAUTIONS.)

Heart Fallure: During baseline-controlled clinical trials, in patients receiving digitalis and diuretics, single doses of ZESTRIL resulted in decreases in pulmonary capillary wedge pressure, systemic vascular resistance and blood pressure accompanied by an increase in cardiac out-

put and no change in heart rate.
In two placebo controlled, 12-week clinical studies, ZESTRIL as In two placebo controlled, 12-week clinical studies, ZESTRIL as adjunctive therapy to digitals and diverties improved the following signs and symptoms due to congestive heart failure: edema, rales, paroxysmal notemal dyspeae and jugular venous distention. In one of the studies, beneficial response was also noted for: orthopnea, presence of third heart sound and the number of patients classified as NYHA Class III and V. Exercise tolerance was also improved in this study. The effect of isinopni on mortality in patients with heart failure has not been evaluated. The once daily dosing for the treatment of congestive heart failure was the only dosage regimen used during clinical trial development and was determined by the measurement of hemodynamic response.

Asute Mycardial Infarction: The Grupon Islaino per lo Studio della

Acute Myocardial Infarction: The Gruppo Italiano per lo Studio della Sopravvienza nell'Infarch Miocardico (GISSI-3) study was a multicenter, controlled, randomized, unblinded clinical trial conducted in 19,394 Acute Myocardial Infarction: The Gruppo Italiano per to Studio della Sopravietras nell'Indrav Miccardico (ISISS-13) study was a multicenter, controlled, randomized, unbinded clinical trial conducted in 19,394 patients with acute myocardial infarction admitted to a coronary care unit It was designed to examine the effects of short-term (6 week) treatment with Isinoprii, Intrates, their combination, or no therapy on short-term (6 week) mortality and on longer-term death and markedly impaired cardiac function. Patients presenting within 24 hours of the orise of symptoms who were hemodynamically stable were randomized, in a 2 x 2 factorial design, to six weeks of either 1) ZESTRIL alone (n=4841), 21 intrates alone (n=4869), 3) ZESTRIL plus nitrates (n=4841), or 4) open control (n=4943). All patients received routine therapies, including thrombolytics (72%), aspiring (48%), and a beta-blocker (31%), as appropriate, normally utilized in acute myocardial infarction (MI) patients.

The protocol excluded patients with hypotension (systolic blood pressure s 100 mmHq), severe heart failure, cardiogenic shock, and renal dysfunction (serum creatinine 2 mg/dl. and/or protocol (see DDSAGE AND ADMINISTRATION).

Study treatment was withdrawn at six weeks except where clinical conditions indicated continuation of treatment.

The primary outcomes of the trial were the overal mortality at 6 weeks and a combined endpoint at 6 months after the myocardial infarction, consisting of the number of patients who died, had late (day 4) clinical congestive heart failure, or had extensive left ventricular damage defined as ejection fraction <35% or an akinelic-dyskinetic [A-D] score <45%.

Patients receiving ZESTRIL (n=9672) (6.4% vs. 7.2%, respectively) at six weeks afraed numerically better on the combined end-point at 6 morths, the open nature of the assessment of heart failure, substantial toss following procloud any conclusion about this endopoint.

Patients with acute myocardial infarction, treated with ZESTRIL, had a higher (9.0% vers

Myocardial Infarction.

INDICATIONS AND USAGE Hypertension: ZESTRIL is indicated for the treatment of hypertension. It may be used alone as initial therapy or concomitantly with other classes of arithypertensive agents.

Heart Falture: ZESTRIL is indicated as adjunctive therapy in the man-gement of heart failure in patients who are not responding adequately to diuretics and digitalis.

Acute Mycardial Infarction: ZESTRIL is indicated for the treatment of hemodynamically stable patients within 24 hours of acute myocardial infarction, to improve survival. Patients should receive, as appropriate, the standard recommended treatments such as tromobytics, aspirin and

In using ZESTRIL, consideration should be given to the fact that another angiotensin converting enzyme inhibitor, captopril, has caused agranulocytosis, particularly in patients with renal impairment or collagen vascular disease, and that available data are insufficient to show that ZESTRIL does not have a similar risk. (See WARNINGS.)

In considering the use of ZESTRIL, it should be noted that in controlled ACE inhibitors have an effect on blood pressure that is less in black its than in nonblacks. In addition, ACE inhibitors have been associated with a higher rate of angioedema in black than in nonblack patier (see WARNINGS, Angioedema).

CONTRAINDICATIONS

CONTRAINDICATIONS
ZESTRIL is contraindicated in patients who are hypersensitive to this product and in patients with a history of angioedema related to previous treatment with an angiotensin converting enzyme inhibitor.

WARNINGS
Anaphylactold and Possibly Related Reactions: 'Presumably because anglotensin-converting enzyme inhibitors affect the metabolism of elcosanoids and polypeptides, including endogenous bradykinin, patients receiving ACE inhibitors (including ZESTRIL) may be subject to a variety of adverse reactions, some of them serious.

receiving ACE inhibitors (including LESTIFIL) may be subject to a value of adverse reactions, some of them serious.

Angloedema: Angloedema of the face, extremities, lips, tongue, glottis and/or larynx has been reported in patients treated with angloetersin converting enzyme inhibitors, including ZESTIFIL. This may occur at any time during treatment. ACE inhibitors have been associated with a higher rate of angloedema in black than in nonblack patients. ZESTIFIL should be promptly discordinued and appropriate therapy and monitoring should be provided until complete and sustained resolution of signs and symptoms has occurred. In instances where swelling has been confined to the face and lips the condition has generally resolved without treatment, although antihistamines have been useful in relieving symptoms. Angloedema associated with laryngeal edema may be latal. Where there is involvement of the tongue, glottis or farynx, likely to cause airway obstruction, appropriate therapy, e.g., subculandous episephyline sedution 1:1000 (0.3 ml.) of 5.5 ml.) and/or measures necessary to ensure a patent airway about be promptly provided. (See ADVERSE REACTIONS.)

Patients with a history of angloedema unrelated to ACE inhibitor therapy may be at increased risk of angloedema while receiving an ACE inhibitor. See also NIDICATIONS AND USAGE and CONTRANIDICATIONS).

Anaphylactala Reactions During Desensitization: Two patients under going desensitizing treatment with hymenoptera venom while receiving ACE inhibitors sustained life-threatening anaphylactoid reactions. In the same patients, these reactions were avoided when ACE inhibitors were temporarily withheld, but they reappeared upon inadvertent rechallenge.

remporarily minimols, but they reappeared upon madvertent rechallenge. Anaphylactoid Reactions During Membrane Exposure: Sudden and potentially life-threatening anaphylactoid reactions have been reported in some patients dialyzed with high-flux membranes (e.g., AM697) and treated concomitantly with an ACE inhibitor. In such patients, dialysis must be stopped immediately, and aggressive therapy for anaphylactoid reactions be initiated. Symptoms have not been relieved by antihistamines in these situations. In these patients, consideration should be given to using a different type of dialysis membrane or a different class of antihypertensive agent. Anaphylactoid reactions have also been reported in patients undergoing low-density lipoprotein apheresis with dextran sulfate absorption.

emt type of delaysis membrane or a different class of antihypertensive agent. Anaphylactori pactions have also been reported in patients undergring low-density lipoprotein apheresis with dextran suitate absorption. Hypotensics: Excessive hypotension is rare in patients with uncomplicated hypertension treated with ZESTRIL alone. Patients with heart failure given ZESTRIL commonly have some reduction in blood pressure, with peak blood pressure reduction occurring 6 to 8 hours post does, but discontinuation of therapy because of continuing symptomatic hypotension usually is not necessary when dosing instructions are followed; caution should be observed when initiating herapy. (See DOSAGE AND ADMINISTRATION).

Patients at risk of excessive hypotension, sometimes associated with oliquina and/or progressive zolerinia, and rarely with acute renal failure and/or death, include those with the following conditions or characteristics: heart failure with systolic blood pressure below 100 mmtlg, hyponatremia, high dose diuretic therapy, recent intensive diuresis or increase in divertic dose, renal dialysts, or severe volume and/or salt depletion of any etiology. It may be advisable to eliminate the diuretic (except in patients with heart failure), reduce the diuretic dose or increase and intake cautiously before kinkling therapy with ZESTRIL in patients at risk for excessive hypotension who are able to tolerate such adjustments. (See PRECAUTIONS, Drug Interactions and ADVERSE RECTIONS.)

Patients with scheme myocardial infarction in the GISSI-3 trial had a higher (9.0% versus 3.7%) inclidence of persistent hypotension (systolic blood pressure < 90 mmtlig for more than 1 hour) when treated with ZESTRIL. Treatment with ZESTRIL must not be initiated in acute myocardial infarction patients at risk of successive hypotension, therapy should be followed closely for the first two weeks of treatment and whenever the dose of ZESTRIL and/or diuretic is increased. Similar considerations may apply to patients with schemic heart or envero

diuretic may be necessary.

duretic may be necessary.

Leutopenia/Neutropenia/Agranulocytosis: Another angiotensim converting ercyme knithitor, captopnii, has been shown to cause agranulocytosis and bone marrow depression, rarely in uncomplicated patients but more frequently in patients with renal impairment especially if they also have a collagen vascular disease. Available data from clinical trials of ZESTRIL are Insufficient to show that ZESTRIL does not cause agranulocytosis at similar rates. Marketing experience has revealed rare cases of teukopenia/heutropenia and bone marrow depression in which a causal relationship to lisinopril cannot be excluded. Periodic monitoring of white blood cell counts in patients with collagen vascular disease and renal disease should be considered. ease should be considered.

asse should be considered.

Hepatic Faiture: Rarely, ACE inhibitors have been associated with a syn-frome that starts with cholestatic jaundice and progresses to fulminant repatic necrosis and (sometimes) death. The mechanism of this syndrome is not understood. Patients receiving ACE inhibitors who develop jaundice or marked elevations of hepatic enzymes should discontinue the ACE hibitor and receive appropriate medical follow-up.

Fetal/Necestal Morbidity and Mortality: ACE inhibitors can cause fetal of neonatal morbidity and death when administered to pregnant women. (Mecaulta) increasing the information, No. 2 millionous can cause to onatal morbidity and death when administered to prognant women il dozen cases have been reported in the world literature. Whe noy is detected, ACE inhibitors should be discontinued as soon a

pregnancy is observed, ALC inhibitors during the second and third trimesters of pregnancy has been associated with fetal and neonatal injury, including hypotension, neonatal skull hypoplasia, anurta, reversible or irreversible renal failure, and death. Oligoinydramnios has also been reported, pre-sumably resulting from decreased fetal renal function, oligothydramnios in this setting has been associated with fetal limb contractures, craniofacial deformation, and hypoplastic lung development. Permaturity, intraducting growth retardation, and patent ductus arteriosus have also been reported, which will be a for clear whether these ancurrence, were due to the APSown retardation, and patent ductus arteriosus have also been reported, though it is not clear whether these occurrences were due to the ACEinhibitor exposure

inhibitor exposure.

These adverse effects do not appear to have resulted from intrauterine ACE-inhibitor exposure that has been limited to the first trimester. Mothers whose embryos and fetuses are exposed to ACE inhibitors only during the first trimester should be so informed. Nonethelass, when palients become pregnant, physicians should make every effort to discontinue the use of ZESTRIL as soon as possible. Rarely (probably less often than once in every thousand pregnancies), no alternative to ACE inhibitors will be found. In these rare cases, the mother schould be apprised of the potential hazards to their letuses, and serial ultrasound examinations should be performed to assess the intraamniotic environment.

mothers should be apprised of the potential hazards to their fetuses, and serial ultrasound examinations should be performed to assess the intraamniotic environment. If ofigohydramnios is observed, ZESTRIL should be discontinued unless it is considered lifesaving for the mother. Contraction stress testing (CST), a nonstress test (KST), or biophysical profiling (BPP) may be appropriate, depending upon the week of pregnancy. Patients and physicians should be aware, however, that eligohydramnios may not appear until after the letus has sustained irreversible injury. Infants with histories of in utero exposure to AEE inhibitors should be closely observed for hypotension, ofiguria, and hyperhalemia. It ofiguria occurs, attention should be directed toward support of blood pressure and renal perhission. Exchange transfusion or dialysis may be required as means of reversing hypotension and/or substituting for disordered renal function. Lisinopril, which crosses the placeta, has been removed from neonatal circulation by peritioneal dialysis with some clinical benefit, and theoretically may be removed by exchange transhusion, although there is no experience with the latter procedure.

No teratogenic effects of lisinopril were seen in studies of pregnant rats, mice, and rabbits. On a mg/kg basis, the doses used were up to 625 times (in mice), 188 times (in rabb in a).

PRECAUTIONS

Impaired Renal Function: As a consequence of inhibiting the reninangiotenstin-aldosterone system, changes in renal function may be anticipated in succeptible individuals. In patients with severe congestive heart failure whose renal function may depend on the activity of the renin-angiotensin-aldosterone system, treatment with angiotensin converting enzyme inhibitors, including ZESTRIL, may be associated with oliguita and/or progressive azotemia and rarely with acute renal failure and/or death.

and/or death.

In hypertensive patients with unilateral or bilateral renal artery stenosis, increases in blood urea nitrogen and serum creatinine may occur. Experience with another angiotensin converting enzyme inhibitor suggests that these increases are usually reversible upon discontinuation of ZESTRIL and/or diuretic therapy. In such patients, renal function should be monitored during the first few weeks of therapy.

Some patients with hypertension or heart failure with no apparent pre-existing renal vascular disease have developed increases in blood urea nitrogen and serum creatinine, usually minor and transient, especially when ZESTRIL has been given concomitantly with a diuretic. This is more likely to occur in patients with pre-existing renal inspariment. Dosage reduction and/or discontinuation of the diuretic and/or ZESTRIL may be required.

required.

Patients with acute myocardial infarction in the GISSI-3 trial, treated with ZESTRIL had a higher (2.4% versus 1.1%) incidence of renal dyslunction in-hospital and a six weeks (increasing creatinine concentration to over 3 mydL or a doubling or more of the baseline serum creatinine concentration). In acute myocardial infarction, treatment with ZESTRIL should be initiated with cauliton in patients with evidence of renal dyslunction. defined as serum creatinine concentration exceeding 2 mg/dL. If renal dysfunction develops during treatment with ZESTRIL (serum creatinine concentration exceeding 3 mg/dL or a doubling from the pre-treatment value) then the physician should consider withdrawal of ZESTRIL.

value) then the physician should consider withdrawal of Zesthile.

Evaluation of patients with hypertension, heart failure, or myocardial
infarction should always include assessment of renal function. (See
DOSAGE AND ADMINISTRATION.)

Hyperkalemia: In clinical trials hyperkalemia (serum potassium greater Hypertatemta: In clinical trials hypertalemia (serum potassium greater than 5.7 mEq/) occurred in approximately 2.2% of hypertensive patients and 4.8% of patients with heart failure. In most cases these were isolated values which resolved despite continued therapy: Hyperkalemia was a cause of discontinuation of therapy in approximately 0.1% of hypertensive patients; 0.6% of patients with heart failure and 0.1% of patients with myocardial infarction. Risk factors for the development of hyperkalemia include renal insufficiency, diabetes melitius, and the concomitant use of potassium-sparing diuretics, potassium supplements and/or potassium-containing sall substitutes, which should be used cautiously, if at all, with ZESTRIL. (See Drug Interactions.)

Cough: Presumably due to the inhibition of the degradation of endogenous bradykinin, persistent nonproductive cough has been reported with all ACE inhibitors, almost always resolving after discontin-uation of therapy. ACE inhibitor-induced cough should be considered in the differential diagnosis of cough.

Surgery/Anesthesta: In patients undergoing major surgery or during anesthesia with agents that produce hypotension, ZESTRIL may block angiotensin II formation secondary to compensatory renin release. If hypotension occurs and is considered to be due to this mechanism, it can he corrected by volume expansion.

Information for Patients

Angloedema: Angioedema, including laryngeal edema, may occur at any time during treatment with angiotensin converting enzyme inhibitors, including ZESTRIL. Patients should be so advised and told to report immediately any signs or symptoms suggesting angioedema (swelling of face, extremities, eyes, lips, tongue, difficulty in swallowing or breathing) and to take no more drug until they have consulted with the prescribing

Symptomatic Hypotension: Patients should be cautioned to report lightheadedness especially during the first few days of therapy. If actual syncope occurs, the patient should be told to discontinue the drug until

syncope occurs, me patient should be tool to discontinue the ording they have consulted with the prescribing physician.

All patients should be cautioned that excessive perspiration and dehydration may lead to an excessive fall in blood pressure because of reduction in fluid volume. Other causes of volume depletion such as vomiting or diarrhea may also lead to a fall in blood pressure; patients should be advised to consult with their physician.

Hyperkalemia: Patients should be told not to use salt substitutes containing potassium without consulting their physician.

Leukopenta/Neutropenta: Patients should be told to report promp ny indication of infection (e.g., sore throat, fever) which may be a sign

ZESTRIL® (lisinopril)

leukopenia/neutropenia.

Praguancy: Female patients of childbearing age should be told about the consequences of second- and third-trimester exposure to ACE inhibitors, and they should also be told that these consequences do not appear to have resulted from intrauterine ACE inhibitor exposure that has been limited to the first timester. These patients should be asked to report pregnancies to their physicians as soon as possible.

NOTE: As with many other drugs, certain advice to patients being treated ith ZESTRIL is warranted. This information is intended to aid in the safe and effective use of this medication. It is not a disclosure of all possible adverse or intended effects.

Deng Interactions

Dreg Interactions
Hypotension - Patients on Diuretic Therapy: Patients on diuretics and
especially those in whom diuretic therapy was recently instituted, may
occasionally experience an excessive reduction of blood pressure after
initiation of therapy with ZESTRIL. The possibility of hypotensive effects
with ZESTRIL can be minimized by either discontinuing the diuretic or
increasing the saft intake prior to initiation of treatment with ZESTRIL. If it
is necessary to continue the diuretic, initiate therapy with ZESTRIL at a
dose of 5 mg daily, and provide close medical supervision after the initial
dose until blood pressure has stabilized. (See WARNINGS, and DOSAGE
AND ADMINISTRATION.) When a diuretic is added to the therapy of a
patient receiving ZESTRIL, an additional antihypertensive effect is usually
observed. Studies with ACE inhibitors in combination with diuretics indicate that the dose of the ACE inhibitors are reduced when it is given
with a diuretic. (See DOSAGE AND ADMINISTRATION.)

with a durence. (See DUSANE AND ADMINISTRATION.)
Indomethaclae: In a study in 36 patients with mid to moderate hypertension where the antihypertensive effects of ZESTRIL alone were compared to ZESTRIL given concomitantly with indomethacin, the use of indomethacin was associated with a reduced effect, although the difference between the two regimens was not significant.

oftion Services in the two regimens was not significant.

Other Agents: ZESTRIL has been used concomitantly with nitrates and/or digoxin without evidence of clinically significant adverse interactions. This included post myocardial infarction patients who were received intravenous or transdermal introlycerin. No clinically important pharmacokinetic interactions occurred when ZESTRIL was used concomitantly with propranolo or hydrochrothiza/de. The presence of food in the stomach does not after the bioavailability of ZESTRIL.

ach does not after the bioavailability of 2ES HILL.

Agents Increasing Serum Potassium: ZESTRIL attenuates potassium
loss caused by thiazide-type diuretics. Use of ZESTRIL with potassiumspaning diuretics (e.g., spironolactione, triamerene or amilioride), potassium supplements, or potassium-containing salt substitutes may lead to
significant increases in serum potassium. Therefore, if concomitant use
of these agents is indicated because of demonstrated thypokalemia, they of these agents is multi-ded because of terministated hypotalisms, should be used with caution and with frequent monitoring of serum potassium. Potassium sparing agents should generally not be used in patients with heart failure who are receiving ZESTRIL.

Lithium: Lithium toxicity has been reported in patients receiving lithium concomitantly with drugs which cause elimination of sodium, including ACE inhibitors. Lithium toxicity was usually reversible upon discontinuation of lithium and the ACE inhibitor. It is recommended that serum lithium levels be monitored frequently if ZESTRIL is administered concomitantly with lithium.

with lithium.

Carcinggenesis, Mutagenesis, Impairment of Fertility: There was no evidence of a tumorigenic effect when issingorit was administered for 105 weeks to make and lemale rats at doses up to 90 mg/kg/day (about 56 or 9 times" the maximum recommended daily human dose, based on body weight and body surface area, respectively). There was no evidence of carcinogenicity when lisinopril was administered for 92 weeks to (make and female) mice at doses up to 135 mg/kg/day (about 84 times" the maximum recommended daily human dose). This dose was 6.8 times the maximum human dose based on body surface area in mice.

"Calculations assume a tuman weight of 50 kg and human body surface area of 156 mg."

Lisinopril was not mutagenic in the Ames microbial mutagen test with or without metabolic activation. It was also negative in a forward mutation assay using Chinese hamster lung cells. Lisinopril did not produce single strand DNA breaks in an in vitro alkaline elution rat hepatocyte assay. In addition, isinopril did not produce increases in chromosomal aberrations in an in vitro test in Chinese hamster ovary cells or in an in vivo study in mouse bone marrow.

mouse bone marrow.

There were no adverse effects on reproductive performance in male and female rats realed with up to 300 mg/kg/day of lisinopril. This dose is 188 times and 30 times the maximum human dose when based on mg/kg and mg/m2, respectively.

Pregnancy Categories C (first trimester) and D (second and third trimesters). See WARNINGS, Fetal/Neonatal Morbidity and Mortality.

trimesters). See WARNINGS, Petanvelonatal Morbinity and Morbany. Mursing Mothers: Mike to factating rats contains radioactivity following administration of ¹⁴C lisinopril. It is not known whether this drug is excreted in human milk. Because many drugs are excreted in human milk and because of the potential for serious adverse reactions in nursing infants from ACE inhibitors, a decision should be made whether to dis-continue nursing and/or discontinue ZESTRIL, taking into account the importance of the drug to the mother.

Pediatric Use: Safety and effectiveness in pediatric patients have not been established

ANYERSE REACTIONS
ZESTRIL has been found to be generally well tolerated in controlled clinical trials involving 1969 patients with hypertension or heart tailure. For the most part, adverse experiences were mild and transient.

Phyperfension:
In clinical trials in patients with hyperfension treated with ZESTRIL, discontinuation of therapy due to clinical adverse experiences occurred in 5.7% of patients. The overall frequency of adverse experiences bould not be related to total daily dosage within the recommended therapeutic

dosage range.

7 adverse experiences occurring in greater than 1% of patients with hypertension treated with ZESTRIL or ZESTRIL plus hydrochlorothiazide in controlled clinical trials, and more frequently with ZESTRIL and XESTRIL plus hydrochlorothiazide than placebo, comparative incidence data are listed in the table below:

PERCENT	OF PATIENTS	IN CONTROLLED ST	UDIES
{di	(n=1349) Incidence	ZESTRIL/ Hydrochlorothiazide (n=629) Incidence (discontinuation)	PLACEBO (n=207) Incidence (discontinuation)
Body as a Whole			4 5 45 65
Fatigue	2.5 (0.3)	4.0 (0.5)	1.0 (0.0)
Asthenia	1.3 (0.5)	2.1 (0.2)	1.0 (0.0)
Orthostatic Effects	1.2 (0.0)	3.5 (0.2)	1.0 (0.0)

Cardiovascular	_	_				
Hypotension	1.2	(0.5)	16	(0.5)	0.5	(0.5)
Digestive		,,		(0.0)	0.5	(0.5)
Diarrhea	2.7	(0.2)	2.7	(0.3)	2.4	(0.0)
Nausea	2.0		2.5		2.4	(0.0)
Vomiting	1.1	(0.2)	1.4		0.5	(0.0)
Dyspepsia	0.9	(0.0)	1.9		0.0	(0.0)
Muscoloskeletal		. ,		()	0.0	(0.0)
Muscle Cramps	0.5	(0.0)	2.9	(8.0)	0.5	(0.0)
Nervous/Psychiatric						(/
Headache	5.7	(0.2)	4.5	(0.5)	1.9	(0.0)
Dizzlness	5.4	(0.4)	9.2	(1.0)	1.9	(0.0)
Paresthesia	0.8	(0.1)	2.1	(0.2)	0.0	(0.0)
Decreased Libido	0.4	(0.1)	1.3		0.0	(0.0)
Vertigo		(0.1)	1.1	(0.2)	0.0	(0.0)
Respiratory				(<u>-</u>)		(/
Cough	3.5	(0.7)	4.6	(0.8)	1.0	$\{0.0\}$
Upper Respiratory				,,		(4.0)
Infection	2.1	(0.1)	2.7	(0.1)	0.0	(0.0)
Common Cold	1.1	(0.1)	1.3	(0.1)	0.0	(0.0)
Nasal Congestion	0.4	(0.1)	1.3	(0.1)	0.0	(0.0)
Influenza	0.3	(0.1)	1.1	(0.1)	0.0	(0.0)
Skin				,,		(,
Rash	1.3	(0.4)	1.6	(0.2)	0.5	(0.5)
Urogenital				,		,,
Impotence	1.0	(0.4)	1.6	(0.5)	0.0	(0.0)
Chart sale and bee						

Chest pain and back pain were also seen, but were more common on placebo than ZESTRIL.

Pacebo than ZESTRIL.

Heart Failure:

In patients with heart failure treated with ZESTRIL for up to four years, discontinuation of therapy due to clinical adverse experiences occurred in 1.0% of patients. In controlled studies in patients with heart failure, therapy was discontinued in 8.1% of patients treated with ZESTRIL for 12 weeks, compared to 7.7% of patients treated with placebo for 12 weeks. The following table lists those adverse experiences which occurred in greater than 1% of patients with heart failure treated with ZESTRIL or placebo for up to 12 weeks in controlled clinical trials, and more frequently on ZESTRIL than placebo.

Controlled Trials

	ZESTRIL (n=407) Incidence (discontinuation) 12 weeks		Placebo (n≠155) Incidence (discontinuation) 12 weeks		
Body as a Whole					
Chest Pain	3.4	(0.2)	1.3	(0.0)	
Abdominal Pain	2.2	(0.7)	1.9	(0.0)	
Cardiovascular				(/	
Hypotension	4.4	(1.7)	0.6	(0.6)	
Digestive		1,	0.0	(0.0)	
Diarrhea	3.7	(0.5)	1.9	(0.0)	
Mervous/Psychiatric	•	(0.0)	1.3	(0.0)	
Dizziness	11.8	(1.2)	4.5	(1.3)	
Headache	4.4	(0.2)	3.9	(0.0)	
Respiratory	***	(0.2)	3.3	(0.0)	
Upper Respiratory					
Infection	1.5	(0.0)	1.3	(0.0)	
Skin	1.0	(0.0)	1.3	10.0)	
Rash	1.7	(0.5)	0.6	(0.6)	

Also observed at > 1% with ZESTRIL but more frequent or as frequent on placebo than ZESTRIL in controlled trials were asthenia, angina pectoris, nausea, dyspnea, cough, and pruritus.

Worsening of heart failure, anorexia, increased salivation, muscle worksming of heart rather, anotexia, morease servation, measure cramps, back pain, myalgia, depression, chest sound abnormalities, and pulmonary edema were also seen in controlled clinical trials, but were more common on placebo than ZESTRIL.

Acute Myocardial Infarction: In the GISSI-3 trial, in patients treated with ZESTRIL for six weeks following acute myocardial infarction, discontinuation of therapy occurred in 17.6% of patients.
Patients treated with ZESTRIL had a significantly higher incidence of

hypotension and renal dysfunction compared with patients not taking ZESTRIL.

ZESTRIL.

In the GISSI-3 trial, hypotension (9.7%), renal dysfunction (2.0%), cough (0.5%), post infarction angina (0.3%), skin rash and generalized edema (0.01%), and angloedema (0.01%) resulted in withdrawal of treatment. In elderly patients treated with ZESTRIL, discontinuation due to renal dysfunction was 4 2%

renal dystunction was 4.2%.

Other clinical adverse experiences occurring in 0.3% to 1.0% of patients with hypertension or heart failure treated with ZESTRIL in controlled clinical trials and rarer, serious, possibly drug-related events reported in uncontrolled studies or marketing experience are listed below, and within each category are in order of decreasing severity.

and within each category are in order of decreasing severity.

Bedy 13: a Whole: Anaphylacloid reactions (see WARNINGS, Anaphylacloid Reactions During Membrane Exposure), syncope, orthosatic effects, chest discomilorit, pain, pelvic pain, analysis, edema, facial edema, virus infection, lever, chills, malase.

Cardiovascular: Cardiac arrest, myocardial infarction or cerebrovascular accident possibly secondary to excessive hypotension in high risk patients (see WARNINGS, Hypotension); pulmonary embolism and infarction, arrhythmias (including entricular tachycardia, atrial fabrications), palpitations, transient ischemic attacks, paroxysmal nocturnal dyspnea, orthostatic hypotension, decreased blood pressure, peripheral edema, vasculdis.

vascultis.

Digestive: Pancrealitis, hepatitis (hepatocellular or cholestatic jaundice) (see WARNINGS, Hepatic Fallure), womiting, gastritis, dyspepsia, heart-burn, gastrolintestinal cramps, constipation, flatutence, dry mouth, Hemadologic: Pare cases of bone marrow depression, hemolytic anemia, leukopenia/neutropenia and thrombocytopenia.

meinatorugu. Andre Cassa of tom meinator department meinatorugu.

mila, leukoperia/heutopenia and thrombocytopenia.

Endocrine: Diabetes mellitus.

Metabolic: Weight loss, dehyration, fluid overload, gout, weight gain, Musculoskeletal: Arthritis, arthraigia, neck pain, hip pain, low back pain, joint pain, lep pain, knee pain, shoulder pain, ammalni, lumbago, Nervous System/Psychiatric: Stroke, ataxia, memory impairment, tremor, peripheral neuropathy (e.g., dysesthesia), spasm, paresthesia, contusion, insomnia, somolence, hypersomnia, irritability and nervousness.

Respiratory System: Malignant lung neoplasms, hemophysis, pulmonary intiltrates, bronchospasm, asthma, pleural effusion, pneumonia, bronchilis, wheezing, orthopnea, painful respiration, epistaxis, taryoptis, smustis, phanryngea pain, pharyoptis, finitis, rhinorfrea.

Skin: Urticaria, alopecia, herpes zoster, photosenstivity, skin lesions, skin infections, pemphigus, erythema, flushing, diaphoresis, Other severe skin reactions have been reported rarely, including toxic epidermal necroysis and Stevens-Johnson syndrome; causal relationship has not been established.

Special Senses: Visual loss, diplopia, blurred vision, tinnitus, photo-

Problem assessmentation: Urogenilal System: Acute renal failure, oligurla, anuria, uremia, pro-gressive azotemia, renal dysfunction, (see PRECAUTIONS and DOSAGE AND ADMINISTRATION), pyelonephritis, dysurla, urinary tract infection, branch prin

Miscellasseus: A symptom complex has include a positive ANA, an elevated erythro dimentation cate, arthragical-mittis, myalagi, sever, vasculitis, esolidopitila and leukocytosis. Rash, photosenstitivity or other dermatological manifestations may occur alone or in combination with these symptoms.

ANGIGEDEMA: Anpipedema has been reported in patients receiving ZESTRIL (0.1%). Anpipedema associated with laryngeal edema may be Itala it angioedema of the face, extremities, kips, tongue, glottis and/or larynx occurs, treatment with ZESTRILs should be discontinued and appropriate therapy instituted immediately. (See WARNINKSS.) HYPDTENSION: In hypertensive patients, hypotension or synope was a cause of discontinuation of therapy in 0.5% of hypertensive patients. In patients with heart failure, hypotension occurred in 5.3% and synope occurred in 1.8% of patients. These adverse experiences were causes for discontinuation of therapy in 1.8% of hese patients. In patients with heart failure, hypotension occurred in 1.5% of hese patients in patients treated with ZESTRIL for six weeks after acute myocardial infarction, hypotension (systoic blood pressure st00 mmity) resulted in discontinuation of therapy in 9.7% of the patients. (See WARNINGS.) Fetal/Neconatal Mortality.

Fetal/Neconatal Mortality and Mortality.

Chegis: See PRECAUTIONS - Cough

Cough: See PRECAUTIONS - Cough

Clinical Laboratory Test Findings

Serum Electrolytes: Hyperkalemia (See PRECAUTIONS), hyponatremia.

Creatiniae, Blood Urea Nitrogen: Minor increases in blood urea nitrogen and serum creatinine, reversible upon discontinuation of therapy, were observed in about 2.0% of patients with essential hypertension treated with ZESTRIL alone. Increases were more common in patients receiving conomitant diuretics and in patients with renal aftery sterosis. (See PRECAUTIONS.) Reversible minor increases in blood urea nitrogen and serum creations with a consequence and serum creations with senal (Now of estimate and serum creations with a consequence and serum creations with a Now of estimate and serum creations with a Now of estimate and serum creations were observed in more procedurate.)

(See PRECAUTIONS.) Reversible minor increases in blood ura nitrogen and serum creatinine were observed in approximately 11.6% of patients with heart failure on concomitant diuretic therapy. Frequently, these abnormalities resolved when the dosage of the diuretic was decreased. Hemoglobit and tematerit: Small decreases in hemoglobin and hematocrit (mean decreases of approximately 0.4 g% and 1.3 vol%, respectively) occurred frequently in patients treated with ZESTRIL but were rarely of clinical importance in patients without some other cause of anemia. In clinical trials, less than 0.1% of patients discontinued therapy due to anemia.

anemia. In clinical usis, less than 0.1% of parients discontinuous tree-apy due to anemic occurred. (See WARNINGS, Hepatic Failure.)

In hypertensive patients, 2.0% discontinued therapy due to laboratory adverse experiences, principally elevations in blood urea nitrogen (0.5%), serum creatinine (0.5%), and serum potassium (0.4%).

In the heart failure trials, 3.4% of patients discontinued therapy due to laboratory adverse experiences; 1.8% due to elevations in blood urea nitrogen and/or creatinine and 0.6% due to elevations in serum potassium. In the myocardial infarction trial, 2.0% of patients receiving ZESTRIL discontinued therapy due to renal dysfunction (increasing creatinine concentration); less than 1.0% of patients discontinued therapy due to other laboratory adverse experiences: 0.1% with hyper-laiemia and less than 0.1% with hyper-laiemia and les

OVERDOSAGE

UVERLUSAGE

Following a single oral dose of 20 g/kg no lethality occurred in rats, and death occurred in one of 20 mice receiving the same dose. The most likely manifestation of overdosage would be typotension, for which the usual treatment would be intravenous infusion of normal saline solution. Earnem would be intravenous intradictly is.

Listnooril can be removed by hemodialysis.

DOSAGE AND ADMINISTRATION

Hypertension
Initial Therapy: In patients with uncomplicated essential hypertension
Initial Therapy: In patients with uncomplicated essential hypertension
of on directic therapy, the recommended initial dose is 10 mg once a
day. Dosage should be adjusted according to blood pressure response.
The usual dosage range is 20 to 40 mg per day administered in a single
daily dose. The antihypertensive effect may diminish toward the end of
the dosing interval regardless of the administered dose, but most commonly with a dose of 10 mg daily. This can be evaluated by measuring
blood pressure just prior to dosing to determine whether satisfactory
control is being maintained for 24 hours. If it is not, an Increase in dose
should be considered. Doses up to 80 mg have been used but do not
appear to give greater effect. If blood pressure is not controlled with
ZESTRIL. atone, a low dose of a diuretic may be added,
hydrochlorothiazide, 12.5 mg has been shown to provide an additive
effect. After the addition of a diuretic, it may be possible to reduce the
dose of ZESTRIL.

Diuretic Treated Patients: In hypertensive patients who are currently being treated with a diuretic, symptomatic hypotension may occur occasionally following the initial dose of ZESTRIL. The diuretic should be discontinued, if possible, for two to three days before beginning therapy with ZESTRIL to reduce the likelihood of hypotension. (See WARNINGS.) The dosage of ZESTRIL, should be adjusted according to blood pressure response. If the patient's blood pressure is not controlled with ZESTRIL.

response. If the patient's blood pressure is not controlled with ESSIDE, adone, duretic therapy may be resumed as described above. If the diuretic cannot be discontinued, an initial dose of 5 mg should be used under medical supervision for at least two hours and until blood pressure has stabilized for at least an additional hour. (See WARNINGS

pressure has statinged for at least air additional induit, loss transmission and PRECAUTIONS, Drug Interactions.)

Concomitant administration of ZESTRIL with potassium supplements, potassium sat substitutes, or potassium-saring diuretics may lead to increases of serum potassium. (See PRECAUTIONS.)

Dosage Adjustment in Renal Impairment: The usual dose of ZESTRIL (10 mg) is recommended for patients with creatinine clearance > 30 ml/min (serum creatinine of up to approximately 3 mg/dL). For patients with creatinine clearance > 10 ml/min (s mg/dL), the first dose is 5 mg once daily. For patients with creatinine clearance > 10 ml/min (susually on hemodialysis) the recommended initial dose is 2.5 mg. The dosage may be titrated upward until blood pressure is controlled or to a maximum of 40 mg daily.

Renal Status	Creatinine Clearance mL/min	Initial Dose mg/day	
Normal Renal Function to Mild Impairment	> 30	10	
Moderate to Severe Impairment	≥ 10 ≤ 30	5	
Dialveis Patients*	- 10	25**	

* See WARNINGS, Anaphylactoid Reactions During Membrane

Exposure.

Dosage interval should be adjusted depending on the blood pressure response.

response.

Heart Failure
ZESTRIL is indicated as adjunctive therapy with diuretics and digitals. The recommended starting dose is 5 mg once a day. When initiating treatment with Isingrif in patients with heart failure, the initial dose should be administered under medical observation, especially in those patients with low blood pressure (systolic blood pressure below 100 mm/lg). The mean peak blood pressure lovering occurs six to eight hours after dosing. Observation should continue until tolood pressure is stable. The concomitant diuretic dose should be reduced, if possible, to help minimize trypovolemia which may contribute to hypotension. (See WARNINGS and PRECAUTIONS, Drug Interactions) I the appearance of trypotension after the initial dose of ZESTRIL does not preclude subsequent careful dose litration with the drug, following effective management of the hypotension.

on. <u>sa) effective dosage range is 5 to 20 mg per day administered as</u>

Dosage Aljestment in Patients with Heart Fallure and Ranal Impairment or Hyponatramia: In patients with heart failure who have hyponatramia (serum sodium < 130 mEq.L.) or moderate to severe renal impairment (creatinine clearance 5 30 mL/min or serum creatinine > 3 mg/dL), therapy with ZESTRIL should be initiated at a dose of

Actual artists interction: in homodynamically stable patients withing of the onset of symptoms of acute myocardial infarction. The first SESTRIL is or ng given orally, followed by 5 mg after 24 hours, benedicted to the stable of the stable

pnare, the standard recommended treatments such as thrombolytics, aspirin, and beta-blockers. Patients with a low systolic blood pressure (< 120 mmHg) when treatment is started or during the first 3 days after the intarct should be given a lower 2.5 mg oral dose of ZESTBIL (see WARNINGS). It hypotension occurs (systolic blood pressure < 100 mmHg) a daily maintenance dose of 5 mg may be given with temporary reductions to 2.5 mg if needed. If profonged hypotension occurs (systolic blood pressure < 90 mmHg for more than 1 hour) ZESTRIL should be withdrawn. For patients who develop symptoms of heart failure, see DOSAGE AND ADMINISTRATION, leart Failure.

Heart Failure.

Dosage Adjustment is Patients With Myocardial Infarction with Renal Impairment: In acute myocardial infarction, treatment with ZESTRIL should be initiated with caution in patients with evidence of read dysfunction, defined as serum creatinine concentration exceeding 2 mg/dl. No evaluation of dosing adjustments in myocardial infarction patients with severe renal impairment has been performed.

severe renal impairment has been percorneu. Use in Etlery', in general, blood pressure response and adverse expe-riences were similar in younger and older patients given similar doses of ESETRIL Pharmacokinetic studies, however, indicate that maximum blood levels and area under the plasma concentration time curve (AUC) are doubled in older patients, so that dosage adjustments should be made with particular caution.

HOW SUPPLIED

made with particular caulion.

#NOW SUPPLED

2.5 mg Tablets (MDC 8316-8135) white, oval, biconvex, uncoated tablets identified as "ZESTRIL 2 1/2" on one side and "135" on the other side are supplied in bottles of 100 tablets. ZESTRIL 2.5 mg tablets are nanufactured by Zenece Pharmaceuticals.

5 mg Tablets (MDC 8318-8139) pink, capsule-shaped, biconvex, bisected, uncoated tablets, identified "ZESTRIL" on one side and "130" on the other side are supplied in bottles of 100 tablets and 1000 tablets, and unit dose packages of 100 tablets of 100 tablets identified "ZESTRIL 10" debossed on one side, and "131" obbossed on the other side are supplied in bottles of 100 tablets, 1000 tablets, 3000 tablets, and unit dose packages of 100 tablets of 100 tablets. 20 mg Tablets (MDC 8310-8132) red, round, biconvex, uncoated tablets identified "ZESTRIL 20" debossed on one side, and "132" debossed on the other side are supplied in bottles of 100 tablets, 1000 tablets, 3000 tablets, and unit dose packages of 100 tablets. 3000 tablets, and unit dose packages of 100 tablets. 400 mg Tablets (MDC 8310-9132) red, bottles of 100 tablets, 3000 tablets, and unit dose packages of 100 tablets. 40 mg Tablets (MDC 8310-9134) yethow, round, biconvex, uncoated tablets identified "ZESTRIL 40" debossed on one side, and "134" debossed on the other side are supplied in bottles of 100 tablets. Store at controlled room temperature, 20-25°C (68-77°F) [see USP]. Protect from moisture, freezing and excessive heat. Dispense in a tight container.

¶Registered trademark of Hospal Ltd.

Manufactured by: IPR Pharmaceuticals Inc. Distributed by:



Rev I 08/96

Application Number	019777/S032	

LABELING REVIEW

JAN 8 ROCT

RHPM Review of Labeling

NDA:

19-777/S-032 Zestril (lisinopril) Tablets

Date of submission:

December 3, 1996

Date of receipt:

December 4, 1996

Applicant:

Zeneca Pharmaceuticals Group

Based on the labeling for NDA 19-915/S-009 Monopril (fosinopril sodium) Tablets for the treatment of CHF (approved May 2, 1995), we issued supplement request letters on May 4, 1995 for ACE inhibitors approved for the treatment of CHF with once-daily dosing (Prinivil and Zestril), asking for the following labeling change:

CLINICAL PHARMACOLOGY, Pharmacodynamics and Clinical Effects, Heart Failure: Please add the following as the last sentence in this subsection:

The once daily dosage for the treatment of congestive heart failure is a consequence of being the only dosage regimen used during clinical trial development and does not represent a known optimum dosage schedule.

Zeneca responded with a submission dated November 8, 1995. We issued a second supplement request letter dated December 1, 1995, asking for revision of the above sentence to:

The once daily dosage for the treatment of congestive heart failure was the only dosage regimen used during clinical trial development and was determined, perhaps erroneously, by the measurement of hemodynamic responses.

Zeneca responded with a submission dated February 1, 1996. We sent out a third supplement request letter dated May 1, 1996, asking for revision of the above sentence to:

The once daily dosage for the treatment of congestive heart failure was the only dosage regimen used during clinical trial development and was determined by the measurement of hemodynamic responses.

Zeneca has responded with this supplement. In addition to the above statement, they have included revisions requested in our November 24, 1995 approval letter for S-023, our facsimile transmission of March 8, 1996, the regulations on pediatric labeling, and the November 16, 1995 CDER Stability Committee Uniform Storage Statement Memorandum.

Review: The submitted final printed labeling has been revised as follows:

CLINICAL PHARMACOLOGY, Pharmacodynamics and Clinical Effects, Heart Failure:

The last sentence has been revised to add the following to the end of this subsection:

"The once daily dosing for the treatment of congestive heart failure was the only dosage regimen used during clinical trial development and was determined by the

measurement of hemodynamic response."

CLINICAL PHARMACOLOGY, Pharmacodynamics and Clinical Effects, Acute Myocardial Infarction:
As we requested in our letter dated November 24, 1995, in the first sentence of the fourth paragraph, the word "a" has been revised to "the" in the phrase "consisting of the number of patients who had ..."

WARNINGS, Anaphylactoid and Possibly Related Reactions, Anaphylactoid reactions during membrane exposure:

The phrase "(a procedure dependent upon devices not approved in the United States)" has been deleted, as we requested in our facsimile dated March 8, 1996.

WARNINGS, Hypotension:

As we requested in our letter dated November 24, 1995, "e.g.," has been added before "systolic blood pressure of 100 mm Hg or lower" in the second sentence of the fourth paragraph.

PRECAUTIONS, Pediatric Use:

The word "children" has been replaced with "pediatric patients" as required by the regulations on the revision of the Pediatric Use subsection of the labeling [21 CFR 201.57 (f)(9)].

HOW SUPPLIED:

To comply with the CDER Stability Committee Uniform Storage Statement, the first sentence of the fourth paragraph has been revised to "Store at controlled room temperature, 20-25° C (68-77° F) [see USP].

Recommendation: I will prepare an approval letter for Dr. Lipicky's signature. This supplement falls under 21 CFR 314.70(b), Supplements requiring FDA approval before the change is made.

Kathleen F. Bongiovanni 12:23-76

cc:

19-777/S-032 HFD-110 HFD-111/KBongiovanni HFD-111/SBenton

kb/12/23/96.

\mathbf{A}	D	plication	Number	019777/S032

CHEMISTRY REVIEW

DEC 11 1996

CHEMIST'S REVIEW	1. ORGANIZATION HFD-110	2. NDA Number 19-777				
Zeneca Pharmaceutical:	3. Name and Address of Applicant (City & State) Zeneca Pharmaceuticals Wilmington, DE 19850-5437					
5. Drug Name Zestril	6. Nonproprietary Name Lisinopril	7. Amendments & Other (reports, etc) - Dates				
8. Supplement Provides For Final Printed Labeling Package Insert (PI).						
9. Pharmacological Catego Antihypertensive	ory 10. How Dispensed [X] Rx OTC	11. Related IND(s)/ NDA(s)/DMF(s)				
12. Dosage Form(s) TCM	13. Potency(ies) 2.5, 5, 10, 20, 40 mg	NDA 19-558 Prinivil, Merck				
14. Chemical Name and Str	ructure	15. Records/Reports Current				
: 		Yes No				
		Yes No				
16. Comments:						
Changes have been maderequests from the Age	de in the following sectioncy:	ons in compliance with				
(1) CLINICAL PHARMA Heart Failure						
(2) CLINICAL PHARMA Acute Myocardia	ACOLOGY - Pharmacodynamic al Infarction	and Clinical Effects,				
(3) WARNINGS - Anap	phylactoid Reactions Duri	ng Membrane Exposure				
(4) WARNINGS - Hypo	otension					
(5) PRECAUTIONS - F	Pregnancy, Pediatric Use					
(6) HOW SUPPLIED - controlled room	The storage statement is temperature, 20-25°C (6	s changed to read "Store at 8-77°F) [See USP]."				
The revised PI is des	signated "Rev J 08/96."					
17. Conclusions and Recom	mendations:					
APPROVABLE in regard	APPROVABLE in regard to the technical aspects of the labeling.					
18.	REVIEWER					
Name James H.Short	stimula / Kroys	Date Completed 9 Dec 96				
Distribution: Original Jacket Reviewer Division File CSO						

jhs/12/9/96/N19-777.832

R/D Init: RWolters/