Responses of plasma concentrations of A type natriuretic peptide and B type natriuretic peptide to alacepril, an angiotensin-converting enzyme inhibitor, in patients with congestive heart failure

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Abstract

Background—Plasma concentrations of A type or atrial natriuretic peptide (ANP) and B type or brain natriuretic peptide (BNP) are increased in patients with congestive heart failure (CHF).

Objective—To examine the haemodynamic and hormonal responses, especially of ANP and BNP, to oral administration of an angiotensin-converting enzyme (ACE) inhibitor in patients with CHF and in controls.

Patients—12 patients with CHF and 11 controls.

Methods—Haemodynamic variables and plasma concentrations of ANP, BNP, and other hormones were serially measured for 24 hours after alacepril (37.5 mg) was given by mouth.

Results—Pulmonary capillary wedge pressure and systemic vascular resistance decreased significantly in both groups. The cardiac index increased only in the CHF group. In patients with CHF pulmonary capillary wedge pressure, systemic vascular resistance, and cardiac index were significantly changed from 1 to 12 hours after alacepril administration. Plasma ANP and BNP decreased significantly after alacepril was given to the CHF group: neither concentration changed in the control group. In the CHF group plasma ANP was significantly lower between 1 and 6 hours and was highly significantly correlated with pulmonary capillary wedge pressure. Plasma BNP, however, was significantly lower between 6 and 24 hours after alacepril and was not correlated with pulmonary capillary wedge pressure.

Conclusions—The response of plasma BNP after alacepril administration occurred later and lasted longer than the plasma ANP response. This may indicate that the mechanisms of synthesis, secretion, or degradation of the two peptides are different.

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A Type or atrial natriuretic peptide (ANP) has a wide range of potent biological effects including natriuresis, vasodilatation, and inhibition of renin and aldosterone secretion and it has an important role in the homeoeostasis of body fluids and blood pressure. The secretion of ANP is regulated mainly by atrial stretching and ANP is secreted not only by the coronary sinus but also directly into the atrial cavities. Plasma concentrations of ANP are higher in patients with congestive heart failure (CHF). B Type or brain natriuretic peptide (BNP) was first isolated from a porcine brain, and subsequently from the hearts of pigs and rats. BNP belongs to the same family of peptides as ANP and like ANP has strong vasodilating and natriuretic actions. We isolated human BNP from human atrium, determined its amino acid sequence, and established a specific radioimmunoassay for human BNP by developing a monoclonal antibody against it. We showed that BNP is a novel cardiac hormone secreted mainly from the ventricle and that plasma BNP is much increased in patients with CHF and those with acute myocardial infarction.

In patients with CHF the renin-angiotensin-aldosterone, arginine vasopressin, and sympathetic nervous systems are activated. Though the heightened activities of these vasoconstrictive agents help to maintain the perfusion pressure to vital organs, they can, by increasing both preload and afterload, ultimately contribute to the deterioration of cardiac function and high mortality. Though both ANP and BNP, vasodilating peptides, are secreted in large amounts in patients with CHF, the concentrations attained in these patients may not be sufficient to improve left ventricular function: intravenous infusion of both ANP and BNP improves left ventricular function by reducing both preload and afterload. There are many reports that treatment with angiotensin-converting enzyme inhibitors is beneficial in patients with CHF. Because atrial stretching is an important regulator of ANP release, it might be anticipated that angiotensin-converting enzyme inhibitors, because of the reduced preload, can induce a decrease in plasma ANP. However, there is no consensus on the effect of angiotensin-converting enzyme inhibitors on ANP release. Furthermore, there are few reports on the response of plasma BNP to the administration of ACE inhibitors in patients with CHF. In this study we examined the short term haemodynamic and hormonal
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responses, especially of ANP and BNP to alacepril (an ACE inhibitor) given by mouth to patients with CHF.

Patients and methods

Patients

We studied 12 patients with CHF (7 men and 5 women; aged 43–72 (mean 54)) and 11 controls (6 men and 5 women; aged 41–68 (mean 51)). The cause of CHF was dilated cardiomyopathy in eight patients, old myocardial infarction in three patients, and mitral regurgitation in one patient. Nine patients were in New York Heart Association class III and three were in class IV. The controls had complained of chest pain, but no cardiac disease was apparent on a coronary arteriogram and left ventriculogram.

Written informed consent was obtained from each patient and his or her family. This study protocol accords with the guidelines of the ethics committees of our hospitals.

Study protocol

Figure 1 shows the study protocol. A Swan-Ganz catheter (Goodtec, USA) was inserted a day before the study and the patients were kept in bed overnight. The study patients did not receive any drugs for at least 24 hours before the procedure. Two serial baseline measurements were obtained in each patient 15 minutes apart and when haemodynamic variables were stable the patients were given alacepril (37.5 mg by mouth).

Haemodynamic measurements

Pulmonary capillary wedge pressure, pulmonary arterial pressure, right atrial pressure, and systemic blood pressure were measured 15 minutes before and immediately before alacepril was given by mouth; and 1, 2, 3, 4, 6, 8, 12, and 24 hours after alacepril administration. Cardiac output was measured in triplicate by the thermodilution technique at the same time as the pressure measurements. The heart rate was monitored continuously by the electrocardiogram (lead II). Cardiac index and systemic vascular resistance were calculated by standard formulas.

Blood sampling

Blood samples were taken from the pulmonary artery through a Swan-Ganz catheter at the same time as the pressure was measured. Blood samples for the measurement of plasma renin activity and plasma concentrations of aldosterone, catecholamine, ANP, and BNP were transferred to chilled disposable tubes containing aprotonin (1000 kallikrein inactivator units/ml) and ethylene-diaminetetraacetic acid (1 mg/ml). All the blood samples were immediately placed on ice and promptly centrifuged at 4°C. Aliquots of plasma were immediately stored at –80°C until the assay.

Measurement of plasma hormones

The plasma concentration of ANP was measured by a specific radioimmunoassay for a human ANP as reported elsewhere. This radioimmunoassay recognises a carboxyterminal sequence of ANP and the minimum limit of detection is 1 pg/tube. The intra- and inter assay coefficients of variation were 7.2% and 7.8% respectively. The cross reactivity with human BNP and C type natriuretic peptide was less than 0.01% on a molar basis. Plasma BNP was measured by a specific radioimmunoassay with a monoclonal antibody that recognised the ring structure of human BNP. The limit of detection was 1 pg/tube. The intra assay and inter assay coefficients of variation were 8.4% and 6.4% respectively. The cross reactivity with a human ANP was less than 0.01% and with C type natriuretic peptide it was less than 1% on a molar basis. Plasma renin activity and plasma aldosterone concentration were measured with commercial kits—renin radioimmunoassay beads (Dainabot, Tokyo, Japan) and an aldosterone radioimmunoassay kit II (Dainabot, Tokyo, Japan) respectively. Plasma noradrenaline was measured by high performance liquid chromatography combined with the trihydroxyindole fluorometric procedure (HLC8030 Tosoh, Tokyo, Japan).

Statistical analysis

All values were expressed as mean (1SE). We used one way analysis of variance for repeated measures with subsequent Dunnett's test for within group comparisons. Haemodynamic variables and hormone concentrations in the control and CHF groups were compared by unpaired t testing. A P value < 0.05 was regarded as statistically significant.

Results

Haemodynamic study

Figure 2 shows the results of the haemodynamic measurements. Pulmonary capillary wedge pressure decreased significantly from 1 hour after the administration of alacepril in both the control and CHF groups, and the lowest level was reached 3 hours after administration of alacepril in both the control and CHF groups. In both groups pulmonary capillary wedge pressure remained significantly reduced up to 12 hours after alacepril. At each time during the study there was a significant difference in pulmonary capillary wedge pressure between the control and CHF groups.

Cardiac index increased significantly from 1 hour after the administration in the CHF group and a maximum was reached 2 hours

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<td>Time course</td>
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Figure 1 Study protocol. Haemodynamic measurements included pulmonary arterial pressure, pulmonary capillary wedge pressure, right atrial pressure, and systemic arterial pressure.
after the administration of alacepril. A significant increase persisted up to 12 hours after the administration of alacepril to the CHF group. In the controls, however, there was no significant increase in cardiac index except at 6 and 12 hours. Cardiac index was significantly lower in the CHF group than in the control group before alacepril was given and 6, 8, 12, and 24 hours after it was given.

In both groups systemic vascular resistance was significantly lower 1 hour after alacepril. It was lowest in the CHF group 2 hours after alacepril and in the control group at 6 hours. A significant decrease persisted at 12 hours in both groups. There was a significant difference in systemic vascular resistance between the control and CHF groups at each time except 2 hours after alacepril was given.

Heart rate did not change during the first 4 hours in either group and increased significantly at 6 and 12 hours in the CHF group and at 6, 8, and 12 hours in the control group. Heart rate tended to be higher in the CHF group than in the control group, but none of the differences in heart rate between the two groups during the study period was significant.

Mean blood pressure decreased significantly from 1 hour after administration in both the control and CHF groups, and was lowest at 2 hours in both the control and CHF groups. The decrease was still significant 12 hours after alacepril in both groups. Mean blood pressure tended to be higher in the CHF group than in the control group, but none of the differences between the two groups was significant.

HORMONAL RESPONSES
Figure 3 shows the result of the hormonal responses. Plasma ANP decreased significantly from 1 hour after the administration in the CHF group, and was lowest 2 hours after oral alacepril. In the CHF group plasma ANP remained significantly lower 6 hours after the administration. In the controls, however, plasma ANP did not change in the 24 hours of the study. Throughout the study plasma ANP was significantly higher in the CHF group than in the controls.

Plasma BNP did not change during the first 4 hours after alacepril. In the CHF group it began to decrease significantly from 6 hours...
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Figure 4 Percentage changes in plasma ANP and BNP in patients with congestive heart failure (CHF).

Pre, pre-administration (100%), * P < 0.05, ** P < 0.01 compared with values at time 0.
† P < 0.05, †† P < 0.01 control vs CHF.

Figure 5 Correlation of Δ PCWP with Δ ANP (upper) and Δ BNP (bottom) in the congestive heart failure (CHF) group. There is a significant linear correlation between Δ PCWP and Δ ANP, but not between Δ PCWP and Δ BNP.

Δ PCWP, change in pulmonary capillary wedge pressure compared with the pre-administration value. Δ ANP, change in plasma ANP compared with the pre-administration value. Δ BNP, change in plasma BNP compared with the pre-administration value.

Correlation between pulmonary capillary wedge pressure and plasma ANP and BNP

Figure 5 shows the correlation between the change in pulmonary capillary wedge pressure from the value before alacepril (Δ PCWP) and the change in plasma ANP or BNP from the value before alacepril (Δ ANP or Δ BNP). There was a significant correlation between Δ PCWP and Δ ANP (n = 108, r = 0.521, P < 0.001) but there was no correlation between Δ PCWP and Δ BNP (n = 108, r = -0.089, P = NS).

Discussion

HAEMODYNAMIC RESPONSES TO ALACEPRIL

It is widely accepted that vasoconstrictor systems such as the renin-angiotensin-aldosterone system, sympathetic nervous system, and vasopressin are excessively activated in patients with CHF. Many studies show the efficacy of angiotensin-converting enzyme inhibitors in the treatment of CHF. We examined the haemodynamic and hormonal responses to alacepril, an angiotensin-converting enzyme inhibitor, in patients with CHF.

Alacepril is a sulphhydryl-containing prodrug of captopril that is converted into

and reached the lowest value 12 hours after administration: a significant decrease in plasma BNP persisted at 24 hours. In the controls, however, there was no change in plasma BNP in the 24 hours after alacepril. Throughout the study period plasma BNP was significantly higher in the CHF group than in the control group.

Plasma noradrenalin did not change significantly after alacepril was given to the CHF group. In the controls, however, it increased significantly from 1 hour, when it reached a peak. Plasma noradrenalin was still significantly raised at 12 hours in the control group.

Plasma noradrenalin in the CHF group was significantly higher before administration and 4, 8, and 12 hours after administration of alacepril than in the control group.

In both groups plasma renin activity increased significantly after alacepril and reached a peak at 6 hours. It remained significantly increased at 12 hours in the CHF group and at 6 hours in the controls. Plasma renin activity was significantly higher at 8 and 12 hours in the CHF group than in the controls.

Plasma aldosterone was significantly lower at 2, 3, 4, 8, and 12 hours after alacepril in the CHF group but did not change significantly in the control group. It tended to be higher in the CHF group than in the control group.

Figure 4 shows the percentage change in plasma ANP and BNP after alacepril. Plasma ANP was significantly reduced from 1 hour to 6 hours after alacepril and plasma BNP was significantly reduced at 8, 12, and 24 hours. The percentage reduction in ANP at 1, 2, and 3 hours after alacepril was significantly greater than the reduction in BNP.

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Correlation between pulmonary capillary wedge pressure and plasma ANP and BNP

Figure 5 shows the correlation between the change in pulmonary capillary wedge pressure from the value before alacepril (Δ PCWP) and the change in plasma ANP or BNP from the value before alacepril (Δ ANP or Δ BNP). There was a significant correlation between Δ PCWP and Δ ANP (n = 108, r = 0.521, P < 0.001) but there was no correlation between Δ PCWP and Δ BNP (n = 108, r = -0.089, P = NS).

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Alacepril is a sulphhydryl-containing prodrug of captopril that is converted into
captopril via an intermediate desacetyl-alacepril chiefly in the intestine, liver, and to a lesser extent the kidneys. Alacepril has a prolonged duration of action and reduces blood pressure in patients with hypertension.

Alacepril reduced pulmonary capillary wedge pressure and systemic vascular resistance and increased cardiac index, indicating that it relieved both preload and afterload, and improved left ventricular function in patients with CHF. This accords with the results of studies of other ACE inhibitors. Treatment with alacepril maintained a reduction of both preload and afterload for as long as a half day. This result clearly indicates that alacepril is useful for the treatment of CHF.

RESPONSES OF NORADRENALINE, RENIN, AND ALDOSTERONE TO ALACEPRIL

Some short-term trials of enalapril and captopril showed a significant reduction in plasma noradrenaline; others did not.

In the present study we found that plasma noradrenaline increased after alacepril was given by mouth to the controls, probably in response to the reduction of systemic arterial pressure. In the CHF group, however, noradrenaline concentrations did not change, even when systemic arterial pressure decreased. This is probably because the arterial baroreflex was reduced in patients with CHF.

Plasma renin activity was increased after alacepril administration in the controls and the CHF group, and plasma aldosterone concentration tended to decrease in both groups, which is regarded as indirect evidence of a decrease in angiotensin II activity. These results accord studies of other ACE inhibitors.

RESPONSES OF ANP AND BNP TO ALACEPRIL

Plasma ANP decreased from 1 hour after alacepril administration, and remained significantly lower up to 6 hours after alacepril was given by mouth. The secretion of ANP is mainly regulated by stretching of the atria and its plasma concentrations are increased in patients with CHF. ANP is also secreted by the ventricles in patients with CHF. Thus in the present study reduced atrial overload is thought to have contributed to the decreased plasma concentration of ANP. This suggestion accords with the significant correlation between the degree of change of the plasma ANP and that of pulmonary capillary wedge pressure.

A novel finding in the present study is the different responses in plasma ANP and BNP after alacepril administration. BNP was first isolated from a porcine brain, and subsequently from the hearts of pigs and rats. BNP belongs to the same peptide family as ANP, and like ANP may be involved in the regulation of blood pressure and fluid volume. We showed that BNP is a novel cardiac hormone secreted mainly by the ventricle, and that its plasma concentration is greatly increased in patients with CHF and those with acute myocardial infarction. None the less the secretion patterns of ANP and BNP vary with underlying cardiac disorders in CHF. Plasma BNP showed a much delayed and sustained response to alacepril compared with that of ANP, and plasma BNP was reduced for up to 24 hours, when pulmonary capillary wedge pressure had already returned to the pre-administration value. There was no significant correlation between the degree of change in plasma BNP and that of pulmonary capillary wedge pressure, suggesting that the secretion of BNP is not regulated by arterial pressure.

We have reported that clearance of BNP from circulation is slower than that of ANP. This may have contributed to the delayed reduction of plasma BNP. However, because BNP has a half life of a few minutes, the increase in plasma BNP that lasts several hours after alacepril can not be explained by this mechanism.

ANP is stored in granules in atrial myocytes (regulated pathway) and is secreted mainly in response to stretching of the atrial wall. This explains the rapid decrease in plasma ANP when pulmonary capillary wedge pressure falls. We found, however, that BNP was not reduced for up to 24 hours after synthesis in the ventricle, as evidenced by increased synthesis of messenger RNA in response to stimulus (constitutive pathway).

In the present study the synthesis of BNP in ventricles may have continued for several hours, even after the reduction of ventricular overload. The precise mechanism of the slow fall in plasma BNP remains uncertain. Other factors also may be involved in the mechanism.

In conclusion, plasma concentrations of both ANP and BNP decreased significantly after alacepril, an ACE inhibitor, was given by mouth. Plasma BNP showed a much delayed and prolonged response compared with that of ANP, which decreased rapidly in parallel with the reduction of pulmonary capillary wedge pressure. This difference may be accounted for by different mechanisms of synthesis, secretion, and degradation for ANP and BNP. The study also showed that the administration of alacepril, an ACE inhibitor, improved left ventricular function by reducing both preload and afterload and is useful for the treatment of CHF.

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Responses of plasma concentrations of A type natriuretic peptide and B type natriuretic peptide to acelopril


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