Response of Plasma Aldosterone to Postural Change, Diuretica, Angiotensin II, Sodium Restriction or Loading and Prostaglandin

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SUMMARY

This paper shows the result of plasma aldosterone which was measured by radioimmunoassay method using aldosterone-3-carboxymethoxamine-BSA or aldosterone-hemisuccinate-BSA. Normal control value at recumbency or upright, primary aldosteronism, uremia, essential hypertension and miscellaneous diseases were examined.

By diuretica, plasma aldosterone of normal volunteer was increased but that of patients with edema was decreased temporarily. The response of plasma aldosterone in edematous patient to the constant infusion of Angiotensin II, was stronger than that of normal volunteer. But primary aldosteronism with adenoma showed the sequentially decreased plasma aldosterone by the constant infusion of Angiotension II. Some of the primary aldosteronism showed the normal response to the sodium loading and restriction. By the pressure dose Prostaglandin A$_2$ or E$_1$ infusion, the plasma aldosterone showed a tendency to increase.

Additional Indexing Words:
Radioimmunoassay Edema Primary aldosteronism

The measurement of plasma aldosterone is widespread rapidly. Up to now, it was measured by the double isotope method but this contained much problem about reliability or reproducibility.

Recently the development of radioimmunoassay method, however, conquered this problem. Renin-angiotensin-aldosterone system has a feed-back system. But in some disorders this system does not always act as normal. The measurement of aldosterone is necessary to elucidate this mechanism. In this paper, not only renin-angiotensin system but also other factors which influence to plasma aldosterone were studied.

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Received for publication September 14, 1973.
MATERIALS AND METHODS

Several normal volunteers, 16 hospitalized primary aldosteronism, 8 uremic patients, 7 essential hypertensive patients, 11 congestive heart failure, 3 Cushing's disease, 2 Bartter's syndrome, 2 liver cirrhosis with ascites and 1 congenital adrenal hyperplasia were studied.

a) Influence of postural change
Plasma aldosterone of normal volunteers and patients suffering from miscellaneous diseases was measured in the morning at recumbency. The former was measured again after upright and walking during 2 hours and compared with the value of recumbency. All of these volunteers were not put on diet control during this experiment.

b) Influence of furosemide
Furosemide 100 mg was intramuscularly injected in normal volunteers and patients with edema. Before and after injection, at the interval of 15 min, the blood was drawn 6 times from cubital vein and aldosterone was measured. During these procedures, blood pressure, hematocrit, urine volume and urine electrolytes were measured.

c) Influence of Angiotensin II
Angiotensin II 10 ng per Kg per minute in 250 ml saline solution was intravenously drop infused during 1 hour in normal volunteers, primary aldosteronism and edematous patients. Plasma aldosterone was measured before and after procedure at the interval of 15 min 6 times. At the same time, blood pressure, hematocrit and pulse rate were observed.

d) Sodium restriction and loading
Plasma aldosterone of primary aldosteronism was measured in the following conditions:
1) at the recumbency in early morning,
2) at the same condition after 3 days of sodium restricted diet (almost 3 Gm of sodium chloride per day),
3) at the same condition after 3 days of sodium loaded diet (almost 20 Gm of sodium chloride per day).

e) Influence of Prostaglandin
Prostaglandin A2 50 to 140 ng per Kg per minute or Prostaglandin E1 100 ng per Kg per minute was constantly infused to a hypertensive patient and normal dogs.

RESULTS

a) Plasma aldosterone of normal volunteer and the patients suffering from miscellaneous diseases:

The result of Fig. 1 (left) shows plasma aldosterone of normal volunteer (recumbency or upright), primary aldosteronism, uremia (nephritis, malignant hypertension), essential hypertension, congestive heart failure, congenital adrenal hyperplasia and Bartter's syndrome. Normal range was distributed 0 to 28 ng/100 ml and the mean was 9.24 ± 6.28 (S.D.) ng/100 ml. In 8 of 16 cases of primary aldosteronism it was 50 to 244 ng/100 ml, and in other 8
cases it was 28 to 50 ng/100 ml. The mean was 70.3 ± 50.24 (S.D.) ng/100 ml. The patients of renal disease with uremia showed the upper limit of normal range or slightly elevated value and its mean was 27.15 ± 13.89 (S.D.) ng/100 ml. Seven cases of essential hypertension were within normal range and its mean was 10.55 ± 8.51 (S.D.) ng/100 ml.

One congenital adrenal hyperplasia was 66.1 ng/100 ml, 2 cases of Bartter's syndrome were 268 and 99 ng/100 ml respectively. Two patients of liver cirrhosis with ascites were as high as 38 and 107 ng/100 ml. Three Cushing disease were 10.5, 24.2 and 112.5 ng/100 ml.

b) Influence of the postural change:

Fig. 1 (right) shows plasma aldosterone of normal volunteer at recumbency and after walking for 2 hours in the morning. The mean value of 10 cases at recumbency was 7.31 ± 3.28 (S.D.) ng/100 ml. The mean value after walking of 8 out of above 10 cases was 11.25 ± 8.96 (S.D.) ng/100 ml. This was statistically significant.

c) Influence of diuretica:

Fig. 2 shows the plasma aldosterone before and after intramuscular injection of furosemide 100 mg in one volunteer. Each sample was drawn at an interval of 30 min. Hematocrit, urine electrolytes and volume were measured simultaneously. After the injection, urine volume and Na excretion were slightly increased. As K excretion was almost unchanged, Na/K ratio was increased. Before procedure, plasma aldosterone was 5.8 ng/100 ml. After that it was 12.9, 11.5, 22.3, 19.3, and 6.9 ng/100 ml respectively. This might mean that the dehydration and sodium diuresis increased the plasma aldosterone slightly.
Fig. 2. The response of plasma aldosterone level to furosemide 100 mg intramuscular injection in normal volunteer.

Same method was tried to the patients suffering from liver cirrhosis with ascites and congestive heart failure with edema (Fig. 3 left and right). Both patients showed markedly increased sodium excretion and increased Na/K ratio and decreased urine osmotic pressure.

In patient with ascites, 2 samples drawn before procedure showed 101 and 103 ng/100 ml and after diuresis the values in 30 min, 1, 2, 3, 4, 5, and 6 hours were 71.1, 67.9, 73.0, 68.1, 69.5, 69.5, and 58.8 ng/100 ml respectively. In patient of congestive heart failure, the values before procedure were 28.0 and 30.5 ng/100 ml and after 1, 2, 3, 4, and 6 hours of procedure, the values of plasma aldosterone were 10.3, 11.5, 6.0, 8.1, and 9.5 ng/100 ml respectively.

These results mean that the trend of plasma aldosterone in edematous patient exhibits a different behavior from normal volunteers.

Nine and 24 hours after furosemide procedure, the aldosterone of an edematous patient markedly increased, such as 73 and 125 ng/100 ml, although it was not pointed out in this figure. Such a phenomenon seems to show
so-called "rebound," while the cause is not clear.

d) Influence of Angiotensin II constant infusion:

Fig. 4 (left and right) shows the change of blood pressure and plasma aldosterone during 1 hour constant infusion of Angiotensin II at the rate of

Fig. 3. The response of plasma aldosterone level to furosemide 100 mg intravenous injection in liver cirrhosis (left). The response of plasma aldosterone level to furosemide of same dose in congestive heart failure (right).

Fig. 4. The response of plasma aldosterone level to Angiotensin II 10 ng/Kg/min infusion in 2 normal volunteers.
10 ng per Kg per minute. Plasma aldosterone of one volunteer in Fig. 4 (left) showed 15.8 ng/100 ml in pre-infusion and thereafter the values determined at 15 min interval after the beginning of constant infusion were 14.0, 19.4, 18.8, 23.4, and 29.1 ng/100 ml respectively. Another volunteer (Fig. 4, right) showed 10.3 ng/100 ml in pre-infusion and thereafter 16.4, 29.6, 34.4, 33.9, and 29.9 ng/100 ml respectively.

The increase of systolic and diastolic blood pressure were 26 mmHg and 44 mmHg in the former and 28 mmHg and 20 mmHg in the latter respectively. The former showed the slightly increased hematocrit, and the latter showed the decreased pulse rate. The same procedure was tried to the primary aldosteronism, who was confirmed to have had the adenoma of 16 mm diameter by operation.

As shown in Fig. 5 (left), the plasma aldosterone of pre-infusion was as high as 56.8 ng/100 ml. At an interval of 15 min after the start of constant infusion, the values of plasma aldosterone were 54.5, 43.2, 38.5, 30.9, and 30.3 ng/100 ml. Systolic and diastolic blood pressure were increased 27 mmHg and 15 mmHg respectively. Fig 5 (right) shows the result in this patient obtained by the same procedure after 18 days of adrenalectomy. The plasma aldosterone before infusion was 2.0 ng/100 ml and after infusion it was 5.8, 5.0, 10.7, 24.1, and 8.7 ng/100 ml respectively. This shows the same reaction with normal persons.

Fig. 6 shows the result in another case of primary aldosteronism who had an adenoma of 18 mm diameter. The plasma aldosterone before procedure.
was 66.7 ng/100 ml. At an interval of 15 min after that, the values were 38.5, 44.8, 36.9, and 33.4 ng/100 ml. Hematocrit was decreased slightly from 40% to 38%. Systolic and diastolic blood pressure were increased 37 mmHg and 20 mmHg respectively.

Fig. 7 (left) shows the result in nephrotic syndrome. Before procedure it was 32.2 ng/100 ml and after that the values were 54.1, 78.9, 78.8, 52.0, and 56.0 ng/100 ml. Hematocrit was decreased markedly from 44.5% to 39%. Systolic and diastolic blood pressure were increased 43 mmHg and 38 mmHg respectively.

Fig. 7 (right) shows the result of idiopathic edema. Before procedure it was 28.1 ng/100 ml, after the beginning of infusion 2 samples of 15 and 30 min showed as high as 74.5 and 70.3 ng/100 ml, but in spite of continuation of infusion, the values were gradually decreased to 27.3 and 21.7 ng/100 ml and after the stop it became 13.7 ng/100 ml.
Fig. 7. The response of plasma aldosterone level to Angiotensin II 10 ng/Kg/min infusion in nephrotic syndrome (left) and idiopathic edema (right).

Fig. 8. The change of plasma aldosterone to Angiotensin II 10 ng/Kg/min infusion in normal volunteer, patients with edema and primary aldosteronism. Each plot was subtracted from the pre-infusion value.
Fig. 8 shows the summary of these results which is displayed as the value subtracted from before procedure. This figure shows that the edematous patients are more sensitive to Angiotensin II as compared with normal control and primary aldosteronism is less sensitive.

e) Influence of sodium restriction and loading:
Six patients of primary aldosteronism were examined. Plasma at recumbency was drawn before procedure, after sodium chloride 3 Gm/day and 20 Gm/day during 3 days. The plasma aldosterone of these 3 points was measured. In spite of adenoma, 2 of 6 cases showed the marked response to the sodium restriction and loading (Fig. 9). The values of each case were 54.4, 133.6, 112.0 ng/100 ml and 105.5, 145.0, 78.5 ng/100 ml respectively. Other 4 cases of primary aldosteronism, however, made no response to sodium restriction and loading. Two cases of normal persons who were shown by broken line showed the normal fluctuation.

f) Influence of Prostaglandin A_2 or E_1:
Fig. 10 shows the influence of Prostaglandin A_2 to the patient with malignant hypertension. Prostaglandin A_2 50 ng per Kg per minute was...
Fig. 10. The response of plasma aldosterone level to Prostaglandin A₂ 50 ng/Kg/min infusion in malignant hypertension.

Fig. 11. The response of plasma aldosterone level in dog to Prostaglandin A₂ 120 ng/Kg/min in saline solution (left). The same experiment obtained from Prostaglandin A₂ 140 ng or E₁ 100 ng/Kg/min infusion using 4 dogs (right).
infused during 1 hour continuously. Blood pressure was gradually decreased and the excretion of urinary sodium was slightly increased. Despite no change of plasma cortisol, the plasma aldosterone was gradually increased. The values were 30.0, 26.1, 60.5, 89.1, 149.3, and 67.8 ng/100 ml respectively.

Fig. 11 (left) is an example which was tried to a dog. Before using of Prostaglandin A\textsubscript{2} 120 ng per Kg per minute, saline solution 500 ml was infused. The values of each spot were 8.4, 5.2, 3.4, 5.5, 5.7, 16.3, 14.9, 15.0, 17.8, 18.2, 13.9, and 12.6 ng/100 ml respectively. The increased urinary volume and sodium excretion will be due to the sodium diuresis. The decreased plasma aldosterone will be also due to volume expansion. Prostaglandin A\textsubscript{2} 120 ng per Kg per minute in saline solution 500 ml, however, increased the plasma aldosterone.

Fig. 11 (right) shows the various types of effect to plasma aldosterone, when Prostaglandin A\textsubscript{2} or E\textsubscript{1} was infused, but it seems that the effect was not so strong as in Angiotensin II infusion.

**DISCUSSION**

Recently by the development of radioimmunoassay method, the measurement of plasma aldosterone became easy and rapid. Although the measurement of urine aldosterone excretion and secretion rate so far in use gave us many useful information, it was impossible to observe the plasma aldosterone level in short time interval.

This paper showed not only the plasma aldosterone in various diseases but also the sequential change of it in various stimulation. At recumbency in morning of normal volunteer without diet control, all values were less than 10 ng/100 ml, the values of plasma aldosterone in primary aldosteronism were scattered widely.

The maximum was 244 ng/100 ml and minimum was 28 ng/100 ml which was near the upper value of normal range. These results were almost same with the data of Fukuchi\textsuperscript{11} and Brown.\textsuperscript{8} From these results it will be said that the upper limit of normal range can not always deny the primary aldosteronism.

In fact, the plasma aldosterone in a primary aldosteronism showed the day-to-day variations. Although chronic nephritis with uremia showed the upper limit of normal control, some cases of malignant hypertension showed high value. All of essential hypertension were normal value, but congestive heart failure with edema was relatively high value. Congenital adrenal hyperplasia, Bartter's syndrome and liver cirrhosis with edema showed markedly high value.
Gowenlock (1959) reported the urinary aldosterone excretion due to postural change. The plasma aldosterone due to postural change is found in the paper of Wolff and Torbica (1963). Michalakis and Horton reported the relation between plasma renin and aldosterone in postural change.

Sympathicotonia caused by upright stimulates the afferent arteriole of glomeruli which induces the increased renin secretion. This increased renin accelerates the aldosterone secretion through the Angiotensin II. Upright from recumbency decreases the hepatic blood flow, consequently the plasma aldosterone increases. In all cases of control the plasma aldosterone after 2 hours of upright and walking more or less increased.

Many papers describing about the relation between edema and aldosterone are found. Luetscher (1954) found that nephrotic patient excreted urine containing much aldosterone, thereafter this was watched as the substance causing edema.

It is now recognized, however, that 1) the congestive heart failure in stable equilibrium of hemodynamic condition does not always show the high urinary aldosterone excretion, 2) the patient of primary aldosteronism who is excreting much aldosterone in urine does not always show edema, 3) by the administration of a large amount of aldosterone during 1 or 2 weeks, so-called escaped phenomenon appears. This escaped phenomenon, however, does not appear in congestive heart failure. Such a fact comes into problem when discussing about the role of aldosterone in edema.

In the above-mentioned results, the plasma aldosterone of normal control increased by diuresis. This fact will be explained by the feed-back mechanism due to sodium diuresis.

On the other hand, that of patients with edema was immediately decreased by diuresis. This fact may mean that the improvement of regulatory mechanism of water and electrolytes decreased the plasma aldosterone. Even in such patient, however, the plasma aldosterone was markedly increased after disappearing of diuretic effect as described before. Such a phenomenon cannot be explained only from the improvement of homeostasis. It looks like the body has set about to strive again for the restoration of lost sodium and water.

In normal control Angiotensin II stimulates the secretion of aldosterone as already known. Aldosterone secretion in nephrotic syndrome showed the higher sensitivity to Angiotensin II infusion than normal control.

In primary aldosteronism the plasma aldosterone was decreased by Angiotensin II infusion unexpectedly. Ganong (1966) said that the long time suppressed plasma renin activity decreased the reactivity of aldosterone stimulation to Angiotensin II, on the other hand, the increased plasma renin activity
increased it.

Response of plasma aldosterone to low or high sodium intake was seen in 2 cases of primary aldosteronism with adenoma. This suggests the participation of auto-regulatory system in some cases of primary aldosteronism.

Recently Fichman and Horton (1972) reported that Prostaglandin A increased the plasma aldosterone significantly. His result agrees with this paper. In over a half of his cases the increased plasma aldosterone was unaccompanied by the increased plasma cortisol or plasma renin activity. From these results he concluded that Prostaglandin may either have a direct physiological role in the regulation of aldosterone secretion or may influence aldosterone indirectly consequent to Prostaglandin induced natriuresis.

REFERENCES