Role of Aldosterone in the Remnant Kidney Model in the Rat

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Abstract

The renin-angiotensin-aldosterone system (RAAS) participates in the injury sustained by the remnant kidney. Our studies assessed the importance of aldosterone in that model and the response of aldosterone to drugs interfering with the RAAS. Initially, four groups of rats were studied: SHAM-operated rats, untreated remnant rats (REM), REM rats treated with losartan and enalapril (REM AIIA), and REM AIIA rats infused with exogenous aldosterone (REM AIIA + ALDO). The last group was maintained with aldosterone levels comparable to those in untreated REM rats by constant infusion of exogenous aldosterone. REM rats had larger adrenal glands and a 10-fold elevation in plasma aldosterone compared to SHAM. REM AIIA rats demonstrated significant suppression of the hyperaldosteronism as well as marked attenuation of proteinuria, hypertension, and glomerulosclerosis compared to REM. REM AIIA + ALDO rats manifested greater proteinuria, hypertension, and glomerulosclerosis than REM AIIA rats. Indeed, by 4 wk of observation all of these features of the experimental disease were similar in magnitude in REM AIIA + ALDO and untreated REM. In separate REM rats spironolactone administration did not reduce glomerular sclerosis but did transiently reduce proteinuria, lowered arterial pressure, and lessened cardiac hypertrophy. In summary, aldosterone contributes to hypertension and renal injury in the remnant kidney model. (J. Clin. Invest. 1996; 98:1063–1068.) Key words: aldosterone • renal failure • glomerulus • hypertension • renin

Introduction

 Interruption of the renin angiotensin-aldosterone system (RAAS) by converting enzyme inhibition or angiotensin II (Ang II) receptor antagonism dramatically alters the course of renal disease in the remnant kidney model (1–3). Furthermore, converting enzyme inhibition has proven clinically effective in slowing the decline in renal function of diabetic nephropathy (4). Reductions of systemic arterial pressure and glomerular pressure have generally been associated with these effects (1–3). Actions of Ang II in the kidney have been posited as the major mechanisms for maintenance of elevated glomerular pressure and reductions in glomerular pressure have been attributed, at least in part, to removal of the intrarenal effects of Ang II (1–3). Other nonhemodynamic but intrarenal actions of Ang II such as growth promoting effects and enhancement of ammoniagenesis may also contribute to progressive renal injury and relief from such actions may thereby reduce injury (1). However, the participation of circulating aldosterone in chronic glomerular damage and the possibility that the beneficial effects of RAAS blockade derive from aldosterone suppression have not been systematically examined. We report experiments designed to test the role of aldosterone in the remnant kidney model of progressive renal disease.

Methods

Study 1. Adult male Sprague-Dawley rats weighing 200–300 grams were subjected to sham (SHAM) renal ablation consisting of laparotomy and manipulation of the renal pedicles or subtotal renal ablation to produce the remnant kidney model (REM). Ablation consisted of right nephrectomy and infarction of a portion of the left kidney by ligation of one segmental renal artery. 3 d after reduction in renal mass, the rats with remnant kidneys had creatinines measured on tail vein blood. They were stratified on the basis of that measurement and assigned to one of three groups such that initial serum creatinines were equivalent in all REM groups. The first group with partial ablation consisted of rats with remnant kidneys and no other manipulation (REM). The second partially ablated group consisted of rats that were placed on a combination of enalapril and losartan in their drinking water (50 and 180 mg/liter, respectively) (REM AIIA). Both agents were used simultaneously with the plan of maximally inhibiting both production and action of Ang II. The third group of partially ablated animals (REM AIIA + ALDO) were treated similarly with losartan and enalapril but were subcutaneously infused with aldosterone by osmotic mini-pump at either 30 or 40 μg/kg per d. The rates of aldosterone infusion were determined by initially measuring plasma aldosterone in subgroups of SHAM, REM, and REM AIIA rats. Then, based on the data of Martin et al. and several preliminary trials with differing rates of infusion, these two rates were chosen as likely to bracket the level of aldosterone in untreated REM rats (5). These two rates yielded plasma aldosterone levels that were not significantly different from one another. For the 30 μg/kg per d infusion, plasma aldosterone averaged 504±139 pg/ml (n = 7). For the 40 μg/kg per d infusion, plasma aldosterone averaged 468±87 pg/ml (n = 6). Also systolic pressure and proteinuria were similar at these two rates of aldosterone infusion. The data for the two levels of aldosterone administration were, therefore, pooled into a single group (REM AIIA + ALDO) for the 2-wk study. Small numbers of deaths occurred in various groups; animals that died before the completion of the study were not included in any of the analyses. All rats were allowed free access to water and standard rat chow (protein 24% and sodium 0.44%); Teklad Premier Laboratory Diets, Madison, WI). 12–14 d after assignment to groups, the rats were placed in metabolic cages for collection of urine for 24 h. Systolic blood pressures were mea...

1. Abbreviations used in this paper: Ang II, angiotensin II; RAAS, renin-angiotensin-aldosterone system; REM, remnant; REM AIIA, REM rats treated with losartan and enalapril; REM AIIA + ALDO, REM AIIA rats infused with aldosterone.

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ured in the awake state by the tail cuff method. Rats were then returned to their standard housing for 1 d and killed on the following day by decapitation. Trunk blood was obtained for biochemical analyses and organs were removed for biochemical analyses and weighing. Adrenal weights are expressed as the sum of both glands.

Study 2. A second cohort of rats comprising the same four groups (except that the infusion of aldosterone was at 35 μg/kg per d in the REM AIIA + ALDO group) was maintained for 4 wk. Measurement of systolic arterial pressure, plasma potassium, and urine collections were obtained on these rats after 2 and 4 wk of assignment to groups. At 4 wk under methohexitol anesthesia, their kidneys were perfused fixed with formaldehyde, embedded in paraffin, sectioned, and stained with hematoxylin and eosin. The prevalence of glomerular sclerosis was determined on these sections in a blinded manner. For each rat, at least 100 glomerular profiles were assessed for the presence or absence of any sclerosis. The results are expressed as the percentage of glomeruli having any sclerotic component.

Study 3. A third cohort of rats made up of two groups (both REM) was maintained for 4 wk. These rats were fed the same chow as those of the other two cohorts but were fed powdered food. One group had spironolactone added to the chow at 5.66 grams/kg. These two groups of rats were studied in the same fashion as in study 2 for 4 wk with the degree of glomerulosclerosis determined at the conclusion of the experiment.

The drug was administered orally in the food because preliminary experiments demonstrated that spironolactone is insoluble in water and its subcutaneous administration as an aqueous suspension led to precipitate, apparently of the drug, within the subcutaneous tissue. Also spironolactone solubility in propylene glycol was excellent but when administered in this solvent, sterile inflammatory lesions developed. The dose of spironolactone was chosen by preliminary studies using the mineralocorticoid salt model of hypertension. Rats underwent unilateral nephrectomy alone or unilateral nephrectomy with normal saline as drinking water, and continuous infusion of aldosterone at 1.39 grams/kg per d by Alzet pump. Spironolactone, in the powdered food, at 5.66 grams/kg, reduced the increment in systolic pressure by 75%. However, spironolactone at 1.39 grams/kg did not influence weight or kidney weight SBP V, protein excretion rate; P, plasma aldosterone level; PRA, plasma renin activity; Adrenal RC, adrenal renin concentration.

### Table I. Data from Study 1 at End of 2 Wk

<table>
<thead>
<tr>
<th>Group</th>
<th>Final body weight</th>
<th>Body weight gain</th>
<th>Left kidney weight</th>
<th>SBP mm Hg</th>
<th>U_{PROT} mg/24 h</th>
<th>P_{ALDO} pg/ml</th>
<th>Adrenal weight mg</th>
<th>PRA ng/ml per h</th>
<th>Adrenal RC ng/mg per h</th>
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<tr>
<td>SHAM</td>
<td>336</td>
<td>±16</td>
<td>11.1</td>
<td>118</td>
<td>10</td>
<td>50</td>
<td>51</td>
<td>3.1</td>
<td>2.2</td>
</tr>
<tr>
<td>(n = 6 or 7)</td>
<td></td>
<td>12</td>
<td>0.06</td>
<td>9</td>
<td>2</td>
<td>12</td>
<td>4</td>
<td>0.7</td>
<td>1.2</td>
</tr>
<tr>
<td>REM</td>
<td>277*</td>
<td>66</td>
<td>1.37</td>
<td>185*</td>
<td>121*</td>
<td>526*</td>
<td>63*</td>
<td>4.3</td>
<td>1.3</td>
</tr>
<tr>
<td>(n = 11–25)</td>
<td></td>
<td>26</td>
<td>0.23</td>
<td>26</td>
<td>54</td>
<td>250</td>
<td>4</td>
<td>2.08</td>
<td>0.8</td>
</tr>
<tr>
<td>REM AIIA</td>
<td>285*</td>
<td>74</td>
<td>1.16</td>
<td>125*</td>
<td>31*</td>
<td>181*</td>
<td>56*</td>
<td>13.3*</td>
<td>3.6</td>
</tr>
<tr>
<td>(n = 5–10)</td>
<td></td>
<td>17</td>
<td>0.25</td>
<td>14</td>
<td>12</td>
<td>124</td>
<td>4</td>
<td>12.3</td>
<td>2.4</td>
</tr>
<tr>
<td>REM AIIA + ALDO</td>
<td>302*†</td>
<td>66*</td>
<td>1.49*†</td>
<td>149*†</td>
<td>87*†</td>
<td>487*†</td>
<td>60</td>
<td>5.2</td>
<td>0.50*†</td>
</tr>
<tr>
<td>(n = 9 or 13)</td>
<td></td>
<td>14</td>
<td>0.23</td>
<td>21</td>
<td>28</td>
<td>114</td>
<td>7</td>
<td>5.0</td>
<td>0.20</td>
</tr>
</tbody>
</table>

Values are means±standard deviations. *P < 0.05 vs SHAM; †P < 0.05 vs REM; §P < 0.05 vs REM AIIA. SBP, conscious systolic blood pressure; U_{PROT}, protein excretion rate; P_{ALDO}, plasma aldosterone level; PRA, plasma renin activity; Adrenal RC, adrenal renin concentration.
exaggerated in REM AIIA. The REM AIIA + ALDO group demonstrated suppression of adrenal renin activity when compared with SHAM and REM AIIA.

Systolic blood pressure was highest in the REM group at 185±26 mmHg. The systemic hypertension was essentially prevented in the REM AIIA group with blood pressures reduced to 125±14 mmHg. This effect of losartan and enalapril was in part reversed with aldosterone infusion in the REM AIIA + ALDO group, which developed a systolic blood pressure of 149±21 mmHg. This pattern of effects on systemic blood pressure was mirrored in the variation of proteinuria among the groups. The untreated REM group had 10-fold elevation of proteinuria compared with the SHAM group (121 ±54 vs 11±2 mg/d). As with hypertension, proteinuria was nearly vitiated by enalapril and losartan in the REM AIIA group (33±13 mg/d). Aldosterone infusion caused proteinuria of 87±28 mg/d, a value significantly greater than both REM AIIA and SHAM. Thus, at 2 wk ~1/3 of the systemic hypertension and 2/3 of the proteinuria present in REM, but obviated in REM AIIA, was reproduced by aldosterone infusion in the REM AIIA + ALDO group.

Study 2. Body weights were similar among the three groups with the remnant kidneys while those in SHAM group had greater final body weights (Table II). However, by 4 wk the increment in body from the time of assignment to their groups was similar among all four groups. As in the 2-wk study, adrenal weight was least in the SHAM group. Also, blockade of Ang II production and action again blunted this adrenal hypertrophy in the REM AIIA group. Plasma potassiums were within the normal range and similar in the SHAM, untreated REM, and REM AIIA + ALDO groups. However, in the REM AIIA group potassium levels were distinctly elevated.

The pattern of awake systolic blood pressure differences within this cohort of rats was similar at 2 wk to that observed in the cohort used for the 2-wk study (Fig. 1, A). Specifically, the highest pressures were observed in untreated REM AIIA with restitution of a large fraction of the increment in arterial

### Table II. Data from Study 2 at End of 4 wk

<table>
<thead>
<tr>
<th></th>
<th>Final body weight</th>
<th>Body weight gain</th>
<th>Left kidney weight</th>
<th>SBP</th>
<th>Heart weight</th>
<th>( U_{\text{Prot}V} )</th>
<th>( P_k )</th>
<th>Adrenal weight</th>
<th>Glomerular sclerosis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>grams</td>
<td>grams</td>
<td>grams</td>
<td>mmHg</td>
<td>grams</td>
<td>mg/24 h</td>
<td>mM</td>
<td>mg</td>
<td>%</td>
</tr>
<tr>
<td>SHAM (n = 6)</td>
<td>347</td>
<td>99</td>
<td>1.19</td>
<td>118</td>
<td>1.03</td>
<td>19</td>
<td>5.0</td>
<td>46</td>
<td>1.9</td>
</tr>
<tr>
<td>REM (n = 7)</td>
<td>±18</td>
<td>±16</td>
<td>0.08</td>
<td>10</td>
<td>0.05</td>
<td>6</td>
<td>0.6</td>
<td>2</td>
<td>1.2</td>
</tr>
<tr>
<td>REM AIIA (n = 7)</td>
<td>307</td>
<td>97</td>
<td>1.47*</td>
<td>109</td>
<td>0.88*</td>
<td>30</td>
<td>6.2*</td>
<td>57*</td>
<td>4.7*</td>
</tr>
<tr>
<td>REM AIIA + ALDO (n = 6)</td>
<td>312</td>
<td>106</td>
<td>2.03*</td>
<td>186*</td>
<td>1.28*</td>
<td>217*</td>
<td>4.5*</td>
<td>60*</td>
<td>25.3*</td>
</tr>
</tbody>
</table>
| Values are means±standard deviations. *P < 0.05 vs SHAM, †P < 0.05 vs REM, §P < 0.05 vs REM AIIA. \( P_k \), plasma potassium level; SBP, conscious systolic blood pressure; \( U_{\text{Prot}V} \), protein excretion rate.

**Figure 1.** (A) Conscious systolic blood pressure, measured at 2 and 4 wk in rats in the 4-wk study. (B) Urinary protein excretion at 2 and 4 wk in rats in the 4-wk study. Numerical values for the 4-wk data are provided in Table II. The error bars are standard errors of the means.
pressure by exogenous aldosterone infusion in the REM AIIA + ALDO group. This pattern persisted in the measurements made at 4 wk by which point an even larger fraction of the increment in blood pressure sustained in the untreated REM group was reproduced by aldosterone infusion in the REM + ALDO group. As with blood pressure, the patterns of proteinuria observed in the 2-wk study were essentially reproduced at 2 wk in this cohort (Fig. 1, B). This pattern of proteinuria progressed such that by 4 wk essentially identical levels of proteinuria were observed in the REM AIIA + ALDO group as in the untreated REM group. Proteinuria continued to be almost completely prevented in the REM AIIA group even by 4 wk.

Glomerular sclerosis was greatest in the untreated REM and the REM AIIA+ALDO groups (37.2±26.5 and 25.3±11.1%, respectively). Thus, aldosterone infusion despite ongoing enalapril and losartan administration reproduced most of the glomerulosclerosis seen in the untreated REM group. As expected, SHAM animals and those remnant animals in the REM AIIA group manifested similar prevalences of sclerotic glomeruli (1.9±1.2 and 4.7±5.2%, respectively), which were substantially lower than seen in the other two groups.

Study 3. In the rats studied for the effects of spironolactone (see Table III), neither body weight nor food intake was altered by spironolactone intake. The average dose of spironolactone was 420 mg/d per kg of rat weight as measured on the last day. Plasma potassium and adrenal weights were not affected by the drug. However, it reduced systolic arterial pressure at 4 wk by 18 mmHg. Heart weight was also lower with spironolactone therapy. The drug diminished proteinuria by ~40% at 2 wk but this was not statistically significant by 4 wk. Likewise, at 4 wk, there was no detectable difference in the degree of glomerulosclerosis with spironolactone treatment.

**Discussion**

Our studies demonstrate that adrenal hypertrophy and hyperaldosteronism attend the hypertension, proteinuria, and glomerulosclerosis characteristic of the remnant kidney model. The adrenal enlargement has been previously reported, but the elevated plasma aldosterone has not been previously commented upon in this experimental disease (7). As expected, combined therapy with the Ang II receptor blocker, losartan, and the converting enzyme inhibitor, enalapril, lessened the hypertension, proteinuria, and glomerulosclerosis of the remnant kidney (1–3). This regimen also attenuated the hyperaldosteronism. Most importantly, the reproduction of the hyperaldosteronism by exogenous steroid infusion on the background of the pharmacologic blocking maneuvers restored most of the arterial hypertension, proteinuria, and glomerulosclerosis observed with the untreated, subtotally ablated kidney.

Hyperaldosteronism has been noted as a component of clinical chronic renal insufficiency. Hené and colleagues described elevation in plasma aldosterone in patients with stable chronic renal insufficiency of various etiologies (8). In their cross-sectional analysis of subjects with a range of renal function, the level of aldosterone appeared to increase as creatinine clearance fell below ~70 ml/min, rising ultimately three- to fourfold over the values measured above that clearance. Other clinical investigations have also identified heightened aldosterone levels in renal insufficiency. For example, Berl et al. studied eight subjects, whose average creatinine clearance was 14 ml/min, and five of them had plasma aldosterone levels above the normal range (9). Likewise nine subjects with even better renal function (average inulin clearance = 27 ml/min), who were studied by Reams and Bauer, had an average plasma aldosterone level more than fourfold greater than normal values (10, 11). The significance of this hyperaldosteronism to progression has not been much mooted but Walker did note a significant correlation between aldosterone level and rate of renal decay in a longitudinal study of patients with diabetes (12). While elevated plasma aldosterone levels have been remarked upon in clinical renal impairment, hyperaldosteronism has not been a recognized feature of the remnant kidney model. Adrenal hypertrophy had been previously linked to this experimental model. Specifically, Morrison documented increased adrenal weight after subtotal ablation and in qualitatively evaluating this growth he reported that the zona glomerulosa was more prominently widened than the reticulo (7). Thus, our studies extend this latter observation and demonstrate that, like the clinical condition, the remnant kidney model manifests an increased plasma aldosterone.

Pharmacologic interdiction of the RAAS strikingly mitigates the injury incurred by the untreated remnant kidney (1–3). Although the effect of this therapy on aldosterone had not been previously investigated, several other observations in this experimental model do suggest that aldosterone may contribute to its progressive injury. Quan and coworkers performed adrenalectomy in rats with subtotal nephrectomy (13). However, despite large doses of replacement glucocorticoid, sev-
eral cardinal features of the disease including proteinuria, histologic measures of renal injury, and hypertension were still less than in similarly ablated rats with intact adrenal glands. The authors did not replace aldosterone having provided a high salt ration instead. But since mineralocorticoids were not maintained, we speculate that their absence accounted for the mitigation of renal disease after adrenalectomy even with glucocorticoid therapy. Of additional interest, heparin administration provides remarkably complete protection from injury in the remnant kidney model (14–16). Although this effect may emanate from any of several actions of heparin, suppression of aldosterone production should be considered as one of these potentially beneficial actions. Finally, our present data supply yet more direct evidence that aldosterone provokes the key elements of this experimental disease, since the infusion of this steroid alone largely reproduced the syndrome.

The effects of converting enzyme inhibition on the course of clinical renal insufficiency have been generally salutary (1, 4, 10, 18). Furthermore, aldosterone appears to respond to this therapy, although this effect has not been extensively examined. Bauer and Reams reported that the average baseline value of plasma aldosterone of 234 pg/ml in their subjects fell to 135 pg/ml after 1 mo of enalapril administration (10). Rui-lope and colleagues, also studying patients with renal insufficiency, observed a significant decline of plasma aldosterone from 266 to 105 pg/ml with 6 mo of captopril treatment (18). However, by 12 mo this level had risen to 234 pg/ml, a value insignificantly different from the baseline. That aldosterone secretion may “escape” from the suppressive actions of converting enzyme inhibitors has been noted in studies of these drugs in other clinical settings but the therapeutic consequences of such escape are uncertain (18). Moreover, the possibility that the beneficial actions of these agents in clinical renal insufficiency derive, at least in part, from a diminution in aldosterone levels has not been investigated.

Mineralocorticoids have been held responsible for scarring and injury in extrarenal components of the cardiovascular system. Weber and colleagues have adduced evidence that myocardial fibrosis can result from mineralocorticoid action, evidence recently confirmed by Young et al. (19, 20). These data also suggest that the fibrogenic and hypertrophic effects may not be solely a response to systemic hypertension engendered by these steroids. In the present study, the effects of spironolactone to lessen cardiac hypertrophy are also consistent with the findings of these investigators. Our data, however, suggest that aldosterone does contribute to the arterial hypertension of the remnant kidney model. But, the mechanism by which it does so is yet uncertain. Both sodium retention and vasoconstrictive effects of mineralocorticoids have been incriminated in the hypertension of the classic mineralocorticoid/salt model and both categories of action may be at play in partial renal ablation (21–23). We doubt, however, that salt retention is an entirely sufficient explanation for our findings in the subtotal ablation model. We hold this view based on previous studies both from our laboratory and those of others demonstrating virtually no influence on arterial hypertension of diets differing widely in sodium content (14, 24, 25). Vascular actions of aldosterone, for example, to enhance ion permeability in vascular smooth muscle, to reset baroreceptors, or to amplify local vasoconstrictor systems likely ally in the hypertensive action of aldosterone both in the remnant kidney and the mineralocorticoid/salt model (26–29). Glomerular capillary pressures were not measured but we suspect are elevated in the aldosterone supplementation since they are clearly increased in the mineralocorticoid salt model of hypertension (30). Such an effect would contribute to injury.

Nonhemodynamic actions of aldosterone may also participate in its renal as well as cardiac fibrotic consequences. In vitro studies of cultured mesangial cells revealed increased type IV collagen production after incubation with aldosterone and if such an effect obtained in vivo, the potential consequences for glomerular matrix and basement membrane expansion are easily envisioned (31). Although the distal tubule is usually considered the target for aldosterone action in the kidney, transcripts for the mineralocorticoid receptor have been detected in the glomerulus, albeit at lower levels than in the distal tubular epithelium, and they might serve to mediate steroid actions at this site (32). Also, nongenomic effects of aldosterone mediated through membrane rather than nuclear receptors, although not yet examined for glomerular cells, might transduce direct renal responses outside of the distal tubule (33). Thus, both hypertensive and more direct cellular actions of aldosterone, including fibrogenic ones, may account for its sclerosing influence on the remnant glomerulus.

Administration of the mineralocorticoid receptor blocker, spironolactone, had antihypertensive effects and a transient effect on proteinuria. It also mitigated cardiac hypertrophy. However, glomerular structural injury was not notably lessened. The reasons for these relatively modest effects, despite the demonstration of a sizable action of exogenous aldosterone in the rats with remnant kidneys receiving converting enzyme inhibition and angiotensin II receptor blockade, are not clear. Perhaps mineralocorticoid receptor blockade was incomplete or aldosterone may be acting through additional receptors such as the plasma membrane sites described by Wehling and colleagues (33). These latter membrane transducers’ effects are not inhibitable by spironolactone analogs (34).

In summary, our studies demonstrate that hyperaldosteronism occurs in the remnant kidney model. Angiotensin converting enzyme inhibition and angiotensin II receptor antagonism reduced the elevated aldosterone levels and appeared to function, in large part, through this suppressive effect on aldosterone. Aldosterone inflicts much of the injury in the subtotal ablation model and may do so generally in chronic progressive renal disease.

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References