Low-protein diet supplemented with ketoacids ameliorates proteinuria in 3/4 nephrectomised rats by directly inhibiting the intrarenal renin–angiotensin system

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Abstract

Low-protein diet plus ketoacids (LPD + KA) has been reported to decrease proteinuria in patients with chronic kidney diseases (CKD). However, the mechanisms have not been clarified. As over-activation of intrarenal renin–angiotensin system (RAS) has been shown to play a key role in the progression of CKD, the current study was performed to investigate the direct effects of LPD + KA on intrarenal RAS, independently of renal haemodynamics. In this study, 3/4 subtotal renal ablated rats were fed 18% normal-protein diet (Nx-NPD), 6% low-protein diet (Nx-LPD) or 5% low-protein diet plus 1% ketoacids (Nx-LPD + KA) for 12 weeks. Sham-operated rats fed NPD served as controls. The level of proteinuria and expression of renin, angiotensin II (AngII) and its type 1 receptors (AT1R) in the renal cortex were markedly higher in Nx-NPD group than in the sham group. LPD + KA significantly decreased the proteinuria and inhibited intrarenal RAS activation. To exclude renal haemodynamic impact on intrarenal RAS, the serum samples derived from the different groups were added to the culture medium of mesangial cells. It showed that the serum from Nx-NPD directly induced higher expression of AngII, AT1R, fibronectin and transforming growth factor-β1 in the mesangial cells than in the control group. Nx-LPD + KA serum significantly inhibited these abnormalities. Then, proteomics and biochemical detection suggested that the mechanisms underlying these beneficial effects of LPD + KA might be amelioration of the nutritional metabolic disorders and oxidative stress. In conclusion, LPD + KA could directly inhibit the intrarenal RAS activation, independently of renal haemodynamics, thus attenuating the proteinuria in CKD rats.

Key words: Chronic kidney diseases; 3/4 Nephrectomised rats; Low-protein diet; Ketoacids; Renin–angiotensin system

Restriction of dietary protein intake has been reported to decrease urinary protein excretion and protect renal function in uraemic patients, and therefore it is one of the major treatment strategies for chronic kidney diseases (CKD)1,2. It was once believed that such protective effects were produced by haemodynamic changes3–5. However, further investigations showed that a low-protein diet (LPD) also has several non-haemodynamic protective effects, such as ameliorating oxidative stress, balancing protein metabolism, decreasing the expression of growth factors, preventing metabolic acidosis and reducing inflammation6–8. Furthermore, accumulating evidence has shown that reno-protective effects of LPD are also associated with the inhibition of intrarenal renin–angiotensin system (RAS).

Inappropriate activation of the intrarenal RAS is becoming increasingly important in the pathogenesis of renal injury than systemic RAS9. All of the RAS components are found in the kidney. Angiotensin II (AngII) concentrations within the kidney are 1000-fold higher than those in circulating blood, providing evidence for the functional importance of locally generated AngII10,11. In this way, the activation of intrarenal RAS is a key factor in CKD progression.

Several studies have investigated the relationship between different amounts of dietary protein and the activation of intrarenal RAS. Rosenberg et al.12 found renal renin mRNA levels to be higher in male rats fed a 50% protein diet than in those fed a 20% protein diet and lower in male rats fed a 6% protein diet. Arimura et al.13 reported mRNA levels of

Abbreviations: AngII, angiotensin II; AT1R, Ang type 1 receptor; CKD, chronic kidney diseases; FN, fibronectin; LPD, low-protein diet; KA, ketoacids; LPD + KA, low-protein diet plus ketoacids; Nx-LPD, 6% low-protein diet; Nx-LPD + KA, 5% low-protein diet plus 1% ketoacids; Nx-NPD, 18% normal-protein diet; RAS, renin–angiotensin system; TGF-β1, transforming growth factor-β1.

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angiotensinogen and renin to be lower in mice fed an 18 % protein diet than in those fed a 24 % protein diet. Correa-Rotter et al.(14) reported renal renin mRNA levels to be higher in CKD rats fed a 40 % protein diet than in those fed a 6 % protein diet. These studies indicated that high protein intake was linked to increased renin expression in normal rats, as well as in CKD rats. However, most of these investigations have focused only on local renin and angiotensinogen expression and may have overlooked the results of local AngII and AngII receptor expression. Furthermore, these studies failed to explore the possible mechanisms underlying the interaction between different protein intakes and intrarenal RAS.

It is well known that AngII contributes to proteinuria through Ang type 1 receptor (AT1R) activation, and AT1R blockade can ameliorate proteinuria in human CKD patients and CKD animals(15,16). It has also been reported that LPD combined with losartan, an AT1R blocker, has additive effects on retarding renal deterioration in CKD rats by further reducing the concentration of transforming growth factor-β (TGF-β) (17). In this way, the association between anti-proteinuria and anti-fibrotic effects of LPD and AT1R blockade and whether they had any synergetic effects merit further exploration.

Ketoacids (KA), keto-analogues of essential amino acids, have been prescribed alongside LPD to patients with CKD(18). Low-protein diet plus ketoacids (LPD + KA) has more pronounced beneficial effects than LPD alone with respect to impeding the progression of CKD and retarding the initiation of kidney replacement therapy(19,20). However, the extra renoprotective effects of KA remain unclear. The association between KA and intrarenal RAS is also unclear.

Here it is hypothesised that the renoprotective effects of ameliorating proteinuria associated with LPD + KA treatment might be related to the direct inhibition of intrarenal RAS, in addition to the changes in renal haemodynamics. An in vitro experiment was designed here to evaluate the effects of LPD + KA on intrarenal RAS in 3/4 nephrectomised rats and another in vitro experiment was designed to investigate the direct impact of LPD + KA on RAS in mesangial cells with and without AT1R blockade. Proteomics and biochemical detection were performed to evaluate the possible mechanisms underlying these beneficial effects of LPD + KA.

### Methods

#### Animals

The experimental protocol was approved by the Animal Study Committee at Fudan University. Here, 3/4 nephrectomy was performed on 200–250-g male Sprague–Dawley rats (Laboratory Animal Institute of Fudan University), as reported previously(21). Animals were placed on normal-protein diet (NPD) (18 %, Nx-NPD, n 10), LPD (6 %, Nx-LPD, n 10) and LPD (5 %)+KA (1 %) (Nx-LPD+KA, n 10) 1 week after 3/4 nephrectomy surgery. The ten male Sprague–Dawley sham-operated rats fed NPD served as control. All the three diets were modified from AIN-95 (American Institute of Nutrition Rodent Diets), and contained the same energy content (14.6 kj (3.5 kcal/g), vitamins and minerals (Table 1). KA (compound α-ketoacid) was provided by Fresenius-Kabi. The active ingredients of KA were L-lysine monoacetate 105 mg, ketoanalogue α-leucine 101 mg, ketoanalogue α-valine 86 mg, α-phenylalanine ketoanalogue 68 mg, α-isoleucine ketoanalogue 67 mg, α-hydroxy analogue of methionine 59 mg, α-threonine 53 mg, α-histidine 38 mg, α-tyrosine 30 mg and α-tryptophan 23 mg. Animals were permitted free access to food and water and were housed in metabolic cages for 24 h of urine collection every 4 weeks. After 12 weeks of feeding, the rats were killed and the blood samples were obtained for biochemical detection. The cortex of kidney was immediately removed and prepared for histological, protein and mRNA analysis. For RIA and ELISA analysis, a portion of the renal cortex tissues was homogenised (on ice) in the buffer containing 0·1 M-phosphate and 10 μM-indomethacin at pH 7·4. The resulting homogenate was centrifuged for 30 min at 4000 rpm at 4°C. The supernatant was aspirated into polypropylene test tubes and stored at −70°C.

#### Culture of renal mesangial cells

Conditionally immortalised mouse glomerular mesangial cells were provided by Vanderbilt University Medical Center(22). They were cultured in Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 10 % (v/v) fetal bovine serum (FBS) (GIBCO), penicillin (100 U/ml) and streptomycin (100 μg/ml). Cultures were incubated at 33°C in 95 %O2/5 %CO2 with 100 U/ml interferon-γ for proliferation and 37°C in 95 %O2/5 %CO2 for intervention. After serum starvation for 12 h, cells were treated with DMEM supplemented with FBS as control and various kinds of serum (100 μl/ml) obtained from animal experiments with or without losartan (0·02 μM) for 48 h. The supernatant and cell pellets were collected through centrifugation at 3000 rpm for 10 min and stored at −70°C.

#### Laboratory measurements

The levels of serum albumin, total protein, serum creatinine and blood urea nitrogen, TAG, total cholesterol, LDL, HDL, fasting blood glucose, insulin and homoeostasis model assessment of insulin resistance (HOMA-IR) were determined by automatic clinical chemistry analyzer (Modular Corporation). Coomassie Blue G-250 (Nanjing Jiancheng Bioengineering Institute) was used to detect 24-h proteinuria in the rats. The levels of malondialdehyde (MDA), protein carbonylation, GSH peroxidase

### Table 1. The composition of different protein diets

<table>
<thead>
<tr>
<th>Components (g/kg)</th>
<th>NPD</th>
<th>LPD</th>
<th>LPD + KA</th>
</tr>
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<tbody>
<tr>
<td>Maize starch</td>
<td>602</td>
<td>617</td>
<td>617</td>
</tr>
<tr>
<td>Casein</td>
<td>180</td>
<td>65</td>
<td>55</td>
</tr>
<tr>
<td>Soyabeen oil</td>
<td>30</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Fibre</td>
<td>50</td>
<td>50</td>
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<tr>
<td>Mineral salt</td>
<td>35</td>
<td>35</td>
<td>35</td>
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<tr>
<td>Vitamins</td>
<td>0·5</td>
<td>0·5</td>
<td>0·5</td>
</tr>
<tr>
<td>Chloride choline</td>
<td>2·5</td>
<td>2·5</td>
<td>2·5</td>
</tr>
<tr>
<td>Sucrose</td>
<td>200</td>
<td>200</td>
<td>200</td>
</tr>
<tr>
<td>KA</td>
<td>–</td>
<td>–</td>
<td>10</td>
</tr>
</tbody>
</table>

NPD, normal-protein diet; LPD, low-protein diet; LPD+KA, low-protein diet plus ketoacids.
Low-protein diet and renin–angiotensin system

Levels of AngII in the renal cortex were detected using RIA kits (provided by Shanghai Chinese Traditional Medical University). ELISA kits (R&D Systems) were used to measure the levels of AngII in serum and supernatant of cultured cells. All of the samples for measuring AngII were pre-treated by passing them through a Centricon-10 column with a cut-off of >10,000 Da (Amicon; EMD Millipore). The amount of fibronectin (FN) and TGF-β1 released in the cell culture medium was also detected using ELISA kits.

Real-time PCR

Total RNA was extracted from freshly removed cortex of rat kidney and cultured mesangial cells using Trizol® Reagent (Invitrogen). The reverse transcription system included the following: RNA template 5 μg, Oligo dt 1 μl, dNTP 2 μl, Moloney murine leukaemia virus reverse transcriptase (MMLV) 5× buffer 4 μl, RNA inhibitor 1 μl, MMLV 1 μl and diethyl pyrocarbonate H₂O.₆μl (First-strand cDNA Synthesis Kit, Fermentas). The fluorescence quantification PCR system included the following: SYBR Premix Ex Taq, 25 μl; Primer F, 1-0 μl (0-2 μM); Primer R, 1-0 μl (0-2 μM); ROX Reference Dye, 1-0 μl; DNA template, 40 μl; and dH₂O.180 μl with a sum volume of 50-0 μl (SYBR Premix Ex Taq Kit; TaKaRa). PCR reaction was carried out at 95°C for 3 s and at 60°C for 30 s for forty cycles (7500 Fast Real-Time RCR System; Applied Biosystems). The primers and renin probe used in the rat model were as follows – forward primer: 5’-ACC CTC CCC GAC ATC ATC GTA AAG TGC-3’, reverse primer: 5’-AGA ACT TGC GGA TGA AGG TG-3’, probe: 5’-TGG GAG GCA GGA CCT ACA TCA GCA-3’. Primers and probe for AT1R were as follows – forward primer: 5’-TGG ATT TCG AAT AGT GTC TGA GAC C-3’, reverse primer: 5’-AGA TGA TGC TGT AGA GGG TAG GTA T-3’, probe: 5’-TCA ACC CAG AAA AAC AAA ATG GCC C-3’. For cultured mouse mesangial cells, primers for FN were as follows – forward primer: 5’-CCA GGT TGA TGA TAC TTC CAT TGT T-3’, reverse primer: 5’-GCT CTG TGC TAC TGC CTT CTA CTG-3’. Primers for TGF-β1 were as follows – forward primer: 5’-TCC AAA CTA AGG CTC GCC AGT CC-3’, reverse primer: 5’-TGG TGC GGT CCA CCA TTA GTA C-3’. Primers for ATIR were as follows – forward primer: 5’-CAA AGG CCA AGT CGC ACT CA-3’, reverse primer: 5’-GAT GCT GTC ATG TCT GT-3’.

Experimental cycle threshold (Ct) values were normalised to glyceraldehyde-3-phosphate dehydrogenase (Fermentas) measured on the same plate, and the relative levels of gene expression were determined using the 2-ΔΔCt method.

Immunoblotting

Freshly removed cortex from rat kidneys were homogenised and extracted in lysis buffer containing 150 μM-NaCl, 100 μM TRIS buffered saline (pH 8.0), 1% Tween 20, 50 μM-dithio bisulfite, 1 μM-EDTA, 1 μM-phenylmethylsulfonyl fluoride, 10 μg/ml leupeptin, 10 μg/ml aprotinin, 10 μg/ml trypsin inhibitor and 10 μg/ml pepstatin. Protein concentration was determined using bichinchoninic acid protein assay (Sigma).

A measure of 20 μg of protein extract was separated on a 12% SDS-PAGE gel. Protein was transferred to a polyvinylidene difluoride membrane at 22 V overnight at 4°C. The membrane was washed three times with Tris-buffered saline with Tween (50 μM TRIS (pH 7.5), 150 μM-NaCl and 0.05% Tween 20) and then incubated in blocking buffer (150 μM-NaCl, 50 μM-TRIS, 0.05% Tween20 and 5% Carnation nonfat dry milk (pH 7.5)) for 1 h at room temperature. The membrane then was incubated with antibodies raised against TGF-β1 (1:100; Santa Cruz Biotechnology Inc.), renin (1:600; Professor Inagaki, Vanderbilt University), AT1R (1:200; Santa Cruz Biotechnology Inc.) and β-actin (1:2000; Sigma) overnight at 4°C. After three washes, the membrane was incubated with a horseradish-peroxidase-conjugated secondary antibody (1:10,000; Jackson Immuno Research Laboratories) for 1 h at room temperature followed by three 15-min washings. Antibody labelling was visualised using ECL (Amersham Biosciences).

Proteomics

Quantitative serum proteomics using isobaric tags for relative and absolute quantification (iTRAQ) was used to assess the up- and down-regulation induced by different protein intakes in nephrectomised rats, with detailed protocol described in previous works(25).

Statistics

All values were described as means with their standard errors. One-way ANOVA was used for analysis of difference between groups by SPSS statistic 11. Statistical significance was considered at P<0.05.

Results

Animal studies

The body weights of Nx-NPD, Nx-LPD and Nx-LPD + KA rats were lower than those of sham-operated rats but without statistically significance. Furthermore, there was no significant difference between body weights of Nx-NPD, Nx-LPD and Nx-LPD + KA rats. Other nutritional indices such as serum albumin and total protein were comparable in each group (Table 2).

The levels of serum creatinine and blood urea nitrogen in 3/4 nephrectomised groups were significantly higher than those in the sham group. Restriction of protein intake prominently reduced blood urea nitrogen levels in nephrectomised rats but had little impact on serum creatinine. The effect was retained in animals fed LPD + KA (Table 2).

There was significantly more urinary protein excretion in the Nx-NPD group than in the sham group. LPD decreased the urinary protein excretion and LPD + KA had more pronounced effects than LPD (Table 2).

The renal cortex expression of TGF-β1 was significantly higher in nephrectomised rats than in the sham group. LPD and LPD + KA significantly reduced protein levels of TGF-β1, and the level of TGF-β1 in the Nx-LPD + KA group was markedly lower than in the Nx-LPD group (Fig. 1A and B).
Immunoblot showed renin protein levels in the renal cortex to be markedly higher in the Nx-NPD group than in the sham group. Those in the Nx-LPD and Nx-LPD + KA groups were markedly lower than in the Nx-NPD group (Fig. 1(A) and (C)). Real-time PCR showed little difference of renin mRNA in the renal cortex among the sham, Nx-NPD and Nx-LPD groups. However, the Nx-LPD + KA group had significantly less renin mRNA (Fig. 1(E)).

The levels of protein and mRNA of AT1R in the renal cortex of Nx-NPD were 1.46- and 1.45-fold higher than in the sham group, respectively (P < 0.05). Treatment with LPD and LPD + KA for 3 months significantly attenuated these changes, bringing levels nearly down to the control levels. However, there was no difference between LPD and LPD + KA (Fig. 1(A), (D) and (F)).

Systemic levels of AngII were significantly higher in Nx-NPD rats than in sham-operated rats, but those in the Nx-LPD and Nx-LPD + KA groups showed no great difference compared with those in the Nx-NPD group (Fig. 2(A)). By contrast,
renal cortex levels of AngII levels were significantly higher in Nx-NPD rats than in sham-operated rats. LPD and LPD + KA greatly decreased the level of AngII in the renal cortex (Fig. 2(B)).

In vitro studies in mesangial cells

To determine whether the different levels of protein intake had any direct, non-haemodynamic impact on intrarenal RAS, the cultured mouse mesangial cells were treated with various kinds of serum (100 μl/ml) from different animal groups for 48 h.

As the serum concentrations of AngII in nephrectomised rats were much higher than those of sham-operated rats (Fig. 2(A)), the baseline supernatant levels of AngII were also higher in the cultured cells treated with serum from nephrectomised rats than in the sham-serum-treated group, but no significant difference was found among three groups treated with serum from Nx-NPD, Nx-LPD and Nx-LPD + KA rats (Fig. 3(A)). After 48 h of cultivation, AngII levels were still higher in the Nx-NPD and Nx-LPD groups than in the sham group, and there was no significant difference between these two groups. However, the level was considerably lower in the Nx-LPD + KA group than in the Nx-NPD and Nx-LPD groups (Fig. 3(B)).

In addition, the Nx-NPD group had higher levels of AT1R protein and mRNA than the sham group (P < 0.05). Nx-LPD serum significantly reduced the over-production of AT1R protein and mRNA levels and LPD + KA was associated with less expression of AT1R than LPD (Fig. 4).

ELISA and real-time PCR indicated higher supernatant levels and mRNA synthesis of FN and TGF-β1 in the cells stimulated with Nx-NPD serum than in the sham-serum-treated group (P < 0.05). Nx-LPD and Nx-LPD + KA serum significantly reduced the supernatant and mRNA levels of FN and TGF-β1. LPD + KA, compared with LPD, further reduced the mRNA level of TGF-β1 (Fig. 5). To investigate the mechanisms by which LPD and LPD + KA decreased FN and TGF-β1, losartan (an AT1R blocker) was added to the cell cultural medium to show the differences in FN and TGF-β1 production. The supernatant and mRNA levels of FN and TGF-β1 in the Nx-NPD, Nx-LPD and Nx-LPD + KA groups were much lower (almost to normal) after losartan supplementation, and little difference was observed in the sham group before and after losartan treatment (Fig. 5).
Proteomics and biochemical detection

As the serum from nephrectomised rats fed NPD could directly activate the RAS in mesangial cells, there might have been some effectors in the serum associated with the local RAS activation. Therefore, proteomics was used to screen the differentially expressed proteins in the serum from each group of rats.

Here, 328 proteins were analysed, and in sum thirty-seven proteins (11% of all proteins) changed quantity. Among the thirty-seven proteins, the levels of 15 (41%) showed a statistically significant difference between sham-operated rats and nephrectomised rats fed different protein diets. These fifteen proteins were mainly involved in the physiological functions such as nutritional metabolism, oxidative stress and extracellular matrix synthesis (Table 3).

To confirm the results of proteomics, the animals were subjected to biochemical detection. Regarding carbohydrate metabolism, we found that with a comparable level of fasting...
Low-protein diet and renin–angiotensin system

blood glucose the secretion of insulin and HOMA-IR was greatly increased in 3/4 nephrectomised rats and LPD partially reduced the degree of insulin resistance. LPD + KA had further beneficial effects on reducing insulin and HOMA-IR compared with LPD (Table 4). Regarding lipid metabolism, the levels of TAG, total cholesterol and LDL were higher in 3/4 nephrectomised rats than in the sham group. LPD had no significant effects on reducing these levels, but LPD + KA markedly reduced the levels of TAG and total cholesterol. There was no difference in high-density lipoprotein levels among these four groups (Table 4). Regarding amino acid metabolism, serum levels of tryptophan, valine and isoleucine were measured, and no difference was found between these four groups (Table 4). Regarding oxidative stress, the levels of MDA in serum and renal cortex and the level of protein carbonylation in serum were higher in 3/4 nephrectomised rats than in the sham group. The levels of antioxidant enzymes such as GSH-PX and SOD were lower in the serum and renal cortex of 3/4 nephrectomised rats than in the sham group. LPD and LPD + KA greatly ameliorated the changes, compared with NPD. LPD + KA also had better effects than LPD alone on the MDA, SOD and GSH-PX levels (Fig. 6 and 7).

### Discussion

The present study has demonstrated that LPD + KA could significantly inhibit intrarenal RAS activation, attenuating the proteinuria in CKD rats. Mesangial cells were treated with serum derived from each of the experimental groups, and results showed that the serum from 3/4 nephrectomised rats fed NPD directly induced higher expression of AngII, AT1R, FN and TGF-β1 in the mesangial cells than that of sham-operated rats fed NPD. Serum from 3/4 nephrectomised rats fed LPD + KA significantly inhibited these abnormalities. Proteomics and biochemical detection suggested that direct inhibition of intrarenal RAS by LPD + KA, independently of renal haemodynamics, might be associated with improvement of nutritional metabolic disorders and oxidative stress in CKD rats.

In 1980s, LPD was found to slow the progression of experimental CKD(5). Since then, the reno-protective effects of LPD have been verified through clinical trials and animal studies(24–26). LPD has been found to reduce 24-h proteinuria, ameliorate glomerular sclerosis, suppress TGF-β1 expression and maintain nutritional status, which were consistent with the current findings. Our results also showed that LPD brought...
blood urea nitrogen levels down well below NPD but with slight changes in serum creatinine level, consistent with previous reports. To some degree, this indicates the retardation, but not reversal, of renal function in CKD by dietary intervention.

Haemodynamic changes are believed to play an important role in the reno-protective effects of protein restriction. However, the reno-protective effects extend far beyond such haemodynamic benefits. Further investigation has indicated the possible non-haemodynamic mechanisms of reno-protection.
through LPD. The present work reports the excessive oxidative stress and insulin resistance in renal ablated rats fed NPD, which were found to be ameliorated by LPD and LPD + KA. Interestingly, the additional non-haemodynamic action in improving lipid disorders was only detected in the Nx-LPD + KA group, not in the Nx-LPD group, suggesting that KA might have a direct impact on lipid profiles, as previously reported in clinical trials.

Furthermore, the association between dietary protein intake and intrarenal RAS in normal and CKD rats has been addressed in the recent years. In the present study, a comprehensive observation of systemic and intrarenal RAS was carried out in 3/4 nephrectomised rats fed different amounts of protein. The results showed that NPD not only increased the hormonal AngII levels in nephrectomised rats but also up-regulated the expression of renin protein, AngII and AT1R in the kidney, indicating local over-activation of RAS. Although restricted protein intake did not reduce the AngII levels in the circulation, both LPD and LPD + KA mitigated the changes of renin and its downstream AngII synthesis and AT1R expression in the kidney. Therefore, the reno-protective effects of LPD and LPD + KA were partly because of its down-regulation of intrarenal RAS.

The mechanisms by which LPD inhibits renin protein remain speculative. Some studies demonstrated that LPD resulted in a decrease in the production of PG in the kidney and inhibition of Na re-absorption in the loop of Henle, which then increases Na delivery to the macula densa, which may modulate the release of renin in nephrectomised rats. The decrease in renin protein contributed to the reduced intrarenal levels of AngII. Low AngII levels may up-regulate the expression of glomerular AT1R expression. However, in the present study the expression of AT1R in the renal cortex was inhibited in both Nx-LPD and Nx-LPD + KA rats, accompanied by reduced intrarenal AngII levels, suggesting that other mechanisms might be involved in the regulation of AT1R in the renal cortex.

To explore whether LPD and LPD + KA had any direct effects, in addition to the changes of haemodynamics, on intrarenal RAS activation in CKD rats and to identify the underlying mechanisms, the mesangial cell, which plays a key role in the pathological changes that take place during glomerular sclerosis and responses to variable stimuli via local RAS, was selected for further experimentation. The serum samples derived from nephrectomised rats fed different dietary protein were added to the culture medium for the first time. The supernatant levels of AngII in 0-h culture medium of mesangial cells were measured, and no significant differences were observed between Nx-NPD, Nx-LPD and Nx-LPD + KA. In this way, the effects of original AngII from different sera of these groups on the expression of AT1R were excluded. Because, according to previous articles and reviews, there is no evidence that FN and TGF-β1 have any regulatory effects on AT1R expression in mesangial cells, the levels of FN and TGF-β1 in 0-h culture medium were not measured. As expected, Nx-NPD serum directly up-regulated AT1R expression accompanied by the enhanced productions of FN and TGF-β1, which are both well-recognised parameters associated with glomerular sclerosis. Both Nx-LPD and Nx-LPD + KA serum reduced the expression of AT1R significantly, but LPD had no effects on the levels of AngII in the supernatant of mesangial cells, suggesting that non-haemodynamic effects of LPD on intrarenal RAS may be mediated by decreasing expression of AT1R. LPD + KA could decrease both of the AngII levels in the supernatant and AT1R in the mesangial cells. As shown above, because AngII contributes to proteinuria through AT1R activation, and because AT1R blockade can ameliorate proteinuria in human CKD patients and CKD animals, the effects of LPD and LPD + KA in reducing proteinuria are partly mediated by inhibiting AT1R in CKD rats. Meanwhile, the high expression levels of TGF-β1 and FN induced by Nx-NPD serum were reduced to control levels by losartan treatment, as in previous findings in animal studies. Nx-LPD and Nx-LPD + KA serum treatment had the same effects as losartan on TGF-β1 and FN in mesangial cells, further indicating that the changes of some components in the serum induced by dietary protein intake may have direct regulatory effects on the expression of AT1R in the kidney, and then participate in the pathogenesis of glomerular sclerosis.

The present proteomics analysis further revealed that the main changes in the circulation of nephrectomised rats caused by different protein diets involved proteins related to nutritional metabolism, oxidative stress and extracellular matrix synthesis. Some studies have demonstrated that elevated levels of amino acids or glucose, insulin, LDL, TNF-α and IL-1 could stimulate the expression of AT1R in the mesangial cells and vascular smooth muscle cells. Thus, different protein-diet-induced changes of some nutritional metabolites, especially insulin and lipid components confirmed by biochemical detection in this study, might have direct effects on the up-regulation of AT1R in the kidney. Moreover, the reactive oxygen species were found to be substantially enhanced in the circulation and renal cortex of nephrectomised rats fed NPD, whereas LPD and LPD + KA decreased the severity, consistent with the previous study. The enhanced oxidative stress might directly activate intrarenal RAS and may exacerbate nutritional disorders or inflammatory status.

Our results demonstrated that the additional effects of KA supplementation of LPD were on the reduction of proteinuria, improvement of insulin and lipid metabolism, and amelioration of oxidative stress and glomerular sclerosis. This suggested that LPD + KA might have better effects on renal protection in nephrectomised rats than LPD alone. Notably, this is the first report to show that KA had the effect beyond LPD with respect to inhibiting renin, AngII and AT1R expression in the kidney. Although the expression of AT1R in the renal cortex showed no difference between Nx-LPD and Nx-LPD + KA groups, in culture of renal mesangial cells the expression of AT1R induced by serum from Nx-LPD + KA group was lower than that in the Nx-LPD group. This discrepancy might be attributable to the fact that AT1R in different regions of the kidney had different susceptibilities to KA supplementation of LPD, and it merits further exploration. These anti-RAS effects of KA might have been caused by the more pronounced improvement in glucose and lipid metabolism in nephrectomised rats fed LPD + KA than in those fed LPD alone, but the direct effect of KA cannot be completely excluded and is worth further exploration.
In conclusion, LPD + KA was found to directly inhibit intrarenal RAS, independently of renal haemodynamics, attenuating the proteinuria in CKD rats, and the underlying mechanisms might be associated with the improvement of nutritional metabolism and oxidative stress.

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