Lixisenatide protects doxorubicininduced renal fibrosis by activating wNF- κ B/TNF- α and TGF- β /Smad pathways

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Abstract. – OBJECTIVE: The aim of this study was to investigate whether Lixisenatide, NF- κ B/ TNF- α , and TGF- β /Smad pathways exert clear regulatory roles in doxorubicin-induced renal fibrosis in rats, and to explore the possible underlying mechanism.

MATERIALS AND METHODS: 30 rats were randomly assigned into the sham group, the Doxorubicin (DOX) group and the Lixisenatide group, 10 in each group. Eight weeks after the specific animal procedure, serum and kidney samples of rats were collected. The serum levels of Cr and BUN were detected usin tive commercial kits. The activities of PIN ialdehyde (MDA), total antioxidant ca ity (T-AOC), catalase (CAT), reduced glutat (GSH), and superoxide dismutase (SOD) in ney homogenate were accessed using co mercial kits. Meanwhile, path esions kidney tissues were evaluation taining by ing and immune-histochemical st NEL assay, respectively. Also, of relotein J ative genes in NF-KPTN ys we ermined by pathways in rat ki al stainin immune-histoche Western blot, respective **RESULTS:** he Lixisena e group showed signin, antly h evels of Cr and BUN. Activities T-AOC, CAL and SOD in the Lixisen e group were icantly higher, where MDA activity was sonificantly lower group. Lixisenatide treatment the DQ thar viated **DOX-induced** pathologiren W. d cell a tosis in kidneys. Furcal le evels of relative genes rmore prote кВ/Т **TGF-**β/Smad pathways in gnificantly downregulated in ra neys w isenatide roup when compared with the the DC NS: Lixisenatide protects doxoruin-induced renal fibrosis in rats by inhibiting **B/TNF**- α and TGF- β /Smad pathways. Lixsenatide, Nrf2, NF- κ B/TNF- α , TGF- β /Smad, Doxorubicin, Renal fibrosis.

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kidney diseases (CKDs) Recent, chro lly^{1,2}. Due to the high have been concerned exi renal replace therapy, CKD brinheavy economic burden on affected patients their families Therefore, it is of great signimechanism of renal fibrosis ice to verify ssion, to r ent chronic renal failure and p al fib s progression as soon as posdel renal failure is the final stage of sible³. e development of various CKDs. The most balogical change of chronic renal failure

real fibrosis, including glomerular sclerosis and renal interstitial fibrosis. This may eventually lead to the loss of renal function⁴. Renal fibrosis is also an important process to determine chronic renal insufficiency. Meanwhile, the pathological changes can result in loss of normal nephron, proliferation of fibroblasts and myofibroblasts, as well as accumulation of excess extracellular matrix^{5,6}. Previous studies^{7,8} have also demonstrated that renal fibrosis has the characteristics of difficult treatment and progressive lesion, which is the key point in the treatment of chronic renal failure.

Doxorubicin (DOX) is the most widely used broad-spectrum and highly effective anti-tumor antibiotic. However, DOX causes severe kidney injury, which seriously limits its clinical application^{9,10}. Reactive oxygen species (ROS) produced by excessive DOX metabolism in the body may be the main cause of oxidative damage in kidney^{11,12}. Renal cells are highly energy-consuming, which require abundant ATP to maintain their vigorous metabolism and life activities¹³. Therefore, renal cells are very sensitive to oxidative stress and dysfunction caused by exogenous chemicals¹³. Furthermore, renal cells are major targets for DOX toxicity, whose damage is an important feature of DOX-induced nephrotoxicity¹⁴.

NF- κ B/TNF- α exerts a multi-directional effect and unique regulatory mechanism that activates multiple pathways and genes¹⁵. The expression of NF- κ B/TNF- α exists in almost all mammalian cells, which is activated by a number of stimulating conditions¹⁶. NF- κ B/TNF- α pathway plays an important role in the normal physiological regulatory network of immune and inflammatory responses. Meanwhile, it also poses a crucial role in cancer and other diseases¹⁷. In addition, multiple growth factors have been confirmed to participate in the development and progression of renal fibrosis, including TGF-β1, TNF-α, PDGF, IGF-1, IL-4, ET-1, CTGF, and others¹⁷. Among them, TGF- β is a crucial inducing factor for ECM synthesis and accumulation. It has also been recognized as a basic fibrosis regulator. As is known to all, TGF- β 1 is a fibroblast chemotactic compound, which serves as the strongest stimulator of fibroblast collagen production. Currently, it is considered the most critical and important factor in fibrosis¹⁸.

Lixisenatide is a human glucagon-like peptide-1 (GLP-1) receptor agonist. It not only has a hypoglycemic effect of glucose concentration-dependent insulin release, but also exerts the of body mass reduction¹⁹. Pharmacolog dies have shown that Lixisenatide can st ate the proliferation and differentiation of isle cells, increase insulin synthesis, delay ga emptying and reduce food intal ixisenat has a high selectivity for GL s, whic is four times than that of **b** an GL receptor 20 ug/d agonists. Authors²¹ have cated t Lixisenatide is the best the Lixisenatidose. In recent yea the eff de on the occurr of organ and develo damage caused and ischem ve been well investiga ²¹. h tudy, we explored the role of Lix enatide in a ting DOX-induced renal fib is and its under mechanism. We aimed provide an experimental basis for cliof Lixisenatide in patients with nic plicatic rena

Man s and Methods

and Reagents

de was purchased from Sinopharm smical Reagent (Shanghai, China); Doxoruinjection was obtained from Qilu Pharmaceb al (Jinan, China); Relative commercial kits were purchased from Jiancheng Bioengineering Institute (Nanjing, China), including malondialdehyde (MDA), total antioxidant capacity (T-A-OC), catalase (CAT), reduced glutathione (GSH), superoxide dismutase (SOD), Cr (creatinine) and blood urine nitrogen (BUN) determinat

edures

Animals and Experimental P

30 male Sprague Dawley (S weighing 180-220 g (Vital River Laboratory I Techdard nology, Beijing, China) wer Jused in environment. Rats were domly assign three groups, including e sham roup, the group and the Lixise gro 10 rats in each eceived group. Rats in the han hrectomy and intra Itoneal n w normal e, rats in th saline. Mear group underwent n v and receive postoperative admin. ration X twice for constructing the renal fibrosis me After successful conthe model, ere intraperitonealstr fected with saline. The animal procedure in Lixisenatide youp was the same as that of the t intraperitoneal injection of group, wi odel construction. This study natide afte L ved b Animal Ethics Committee of wa. Nantona sity Animal Center.

Renal fibrosis model was established by right stirpation and repeated administration in rats. After one week of habituation, fats were anesthetized with 10% chloral hydrate (5 mg/kg) and placed in the prone position on the surgical table. Skin preparation (2.5-3.0 cm) in the right waist of the rat was performed. A 2.0-3.0 cm incision was cut to 0.5-1.0 cm on the right side of the spine. Subsequently, the right kidney was exposed and the perirenal capsule was bluntly dissociated using curved forceps. Renal artery and vein were clamped and fixed with hemostats to avoid the damage of adrenal tissue. The right renal artery and vein, as well as the ureter, were ligated with 3-0 suture, followed by removal of the right kidney. Finally, the incision was sutured layer by layer.

After surgery, 0.8 million unit/kg penicillin was intramuscularly injected into the hind limb of rats for three consecutive days to prevent infection. On the 7th day after surgery, the surgical incision suture was removed after the verification of healing. On the 7th day after the extirpation of the right kidney, rats were injected with 5 mg/kg DOX through the tail vein. Subsequently, 3 mg/ kg of doxorubicin was repeatedly injected on the 30th day after surgery. At the 8th week postoperatively, renal tissues were harvested for pathological examination. Extensive segmental or even glo-

merular sclerosis, balloon adhesion, renal tubular epithelial vacuolar degeneration, renal interstitial fibrosis and inflammatory cell infiltration indicated the successful construction of the renal fibrosis model in rats.

Assessment of Renal Function

The body weight of each rat was daily recorded before drug administration, and the changes in body mass in each group was calculated at the end of the animal procedure. After sacrifice, rat kidney was immediately harvested. The envelope and fat tissue were peeled off, followed by weighing the kidney mass. The renal index was calculated as kidney mass/body mass × 100%. Subsequently, 2 mL blood sample was collected. After 30 min of blood coagulation, blood samples were centrifuged at 3500 g/min for 10 min. Then the supernatant was harvested and the levels of Cr and BUN were detected using relative commercial kits.

Histological Examination

Kidney tissues were cut into coronal section, fixed with 10% paraformaldehyde and s with hematoxylin and eosin. Histological in kidney tissues were assessed by semi-qu tative detection of renal tubule injury and ne (0 grade = no injury, 1 grade = $\leq 10\%$, 2 gr = 46-7 = 11-25%, 3 grades = 26-45%, 4 and 5 grades = >76%). 5 field omly se lected for each sample, an he path gical le-5 sions were evaluated (ma ration 2

Terminal Deoxyr Leotia, Transferase dU Nick-End L Ving (TUNEL) Ass

TUNEL a y wa ducted to detect renal cell apopt is according the instructions of us Peroxidase h ApopTa Apoptosis Detectio t (Chemicon, Millipo, e, Billerica, MA). ction was counterstained with 5-µ araffin hem nd TUMEL-positive cells were random selected fields (magnificounted 200 ic each kidney sample were ing. for cel

al Measurements

the exposure of the proximal age of the second seco

the color of kidney turned from red to pale, the left kidney was harvested and preserved in liquid nitrogen for subsequent use. Kidney homogenate was prepared and the levels of MD/ CAT, GSH, and SOD were determined using refutive commercial kits.

Immuno-Histochemical Stain

Kidney slices were depa inized, d in ethyl alcohol and blocke 1th blocking on with rimary an for 30 min. After incul dies (TGF-β1, Smad 1α -A) at 4°C overnight and second t room perav an ture for 1 h, in nohistoe ts were kon Eclipse captured usi scope.

Western Blot

ed using the radio-im-Total protein was pitation assay PA) protein lysate m otime, Shanghai, China). The concentration extracted proton was quantified by the bicinninic acid ()) method (Pierce, Rockford, Π A). The p ein sample was separated by nhor and transferred to polyvinyligel (PVDF) membranes (Merck Mildene di ore, Billerica, MA, USA). Then the membranes bated with primary and secondary antiased on the standard protocols of Western olot. Chemiluminescence was used to expose the protein bands on the membrane.

Statistical Analysis

Statistical Product and Service Solutions 22.0 (SPSS) Software package (IBM, Armonk, NY, USA) was used for all statistical analysis. Data were expressed as $\bar{x\pm}s$. The *t*-test was used to compare the difference between the two groups. Categorical data were analyzed using χ^2 -test or Fisher's exact test. p<0.05 was considered statistically significant.

Results

Lixisenatide Improved Renal Function in DOX-Induced Renal Fibrosis

Compared with the sham group, body weight and renal index of rats in the DOX group were remarkably reduced after animal procedures (p<0.05). This indicated that DOX-induced renal damage and successful construction of the rat model. Higher body weight and renal index were observed in the Lixisenatide group (p<0.05, Figure 1A, 1B).



Figure 1. Lixisenatide alleviated renalized tion in DC (n=10), the DOX group (n=10) and the provided group treatment groups; **C**, Serum level p creating in the dia different treatment groups. Data are present as mean \pm difference vs, DOX group (p

Meanwhile titative anal of renal ripheral blood level of function show that X group an Cr in the Lixisenatide group antly higher th was sig at of the sham group <0.05, Figure 1C). A. o, the peripheral vel of N in the DOX group and the blo arkedly higher than the Lixi up was 2<0.05. are 1D). Although Lixisham g ide ti narkably decreased BUN t be reversed to baseline. still c

in Archite Preserved Renal Architecture and Alleviated Architration

significant pathological lesions were observed kidneys harvested from the sham group. Pathological evaluation of kidney tissues in the DOX group showed significantly widening difaced renal fibrosis. **A**, Body weight of rats in the sham group 0); **B**, The ratio of renal weight/body weight in the different treatment groups; **D**, Blood levels of urea nitrogen in the *significant difference vs, sham group (p<0.05); #significant

fuse glomerular mesangial area, capillary lumen stenosis and focal segmental sclerosis of the glomerulus. Meanwhile, immune complex deposition was observed in the glomerular basement membrane, which was irregularly disordered. Most of the glomeruli were swollen and some glomeruli had a crescent formation. Mild to moderate hyperplasia of the mesangial matrix was found, and the number of sclerotic cells increased significantly. The renal sac was moderately dilated, with thickness of the cyst wall. Renal tubular epithelial cells showed granule degeneration, manifesting as cell disintegration, shedding and regeneration. Tubular lumen remarkably expanded with tubular shape in the lumen. Lymphatic-monocyte infiltration around the renal tubule was found, with immune complex deposition. Renal interstitium showed mild to moderate fibrosis and infiltration



Figure 2. Lixisenatide prevented DOX-induced renal pathological mions. Renal sectors were state osin, and examined under a light microscope (magnification 200³) in H&E staining mission statikidney tissues in the sham group, the DOX group, and the Lixisenation roup. **B**, Quantification of outer medulla of the three groups. **C**, Quantification of TUNEL-position of bata were expressed as mean \pm SD.*significant difference vs, sham group statistical significant difference vs, sham group is significant difference vs.

s were stained with hematoxylin and sson staining and TUNEL assay of fication of ATN score in cortex and number of 5 HPF in different groups. ignificant difference vs, DOX group

of inflammatory cells. The Lixisenatide gap presented slighter pathological beings than a DOX group (Figure 2A, 2B) and tubula necrosis (ATN) score in the DOX group was remarkably higher than the disenatide roup and the sham group (p<0.05, Figure 3B)

Lixisenatide Drewased the sectors of Renal Tuber Sectors Sector S

As shown in Figure 1 and 2C, the number of TUN1 positive cells in a ray tissues of the DOX and p was remarkably higher than the sham group The Linenatide group showed significant, T is EL-positive cells than that of the DOX g. (0.05)

natide peased ROS Iction and Tissue Impairment Incing Antioxidant Capacity

ficantly impair the antioxidant capacity of kisissue and increase the production of ROS. In a study, Lixisenatide administration remarkably restored the antioxidant capacity in renal fibrotic tissues. Compared with the DOX group, the level of MDA was significantly decreased after Lixisenatide administration (Figure 3A). Additionally, we also found a higher level of ROS in kidney tissues of the DOX group than that of the Lixisenatide group (Figure 3B). Moreover, results also indicated that the activities of T-AOC, CAT, GSH and SOD were remarkably higher in the Lixisenatide group compared with those of the DOX group (Figure 3C-3F).

Lixisenatide Activated NF-& B/TNF-a Signal Pathway in DOX-Induced Renal Fibrosis

To investigate whether Lixisenatide protected DOX-induced renal damage through activating NF- κ B/TNF- α pathway, serum samples of rats were collected. Subsequently, the levels of cytokines were detected. Results showed that the levels of TNF- α , IL-1 β , and IL-6 in the Lixisenatide group were markedly lower than the DOX group (Figure 4A-4C).

Kidney tissues of rats in the Lixisenatide group and the DOX group were collected as well. Western blot results showed that the protein levels of NF- κ B, TNF- α , VCAM-1, ICAM-1, and MCP-1 in the Lixisenatide group were significantly lower than the DOX group (Figure 4D).



dialdehyde Figure 3. Lixisenatide attenuated oxidative stress. A. Content of A) in kidney tissues. **B**, Content of ROS was expressed as arbitrary units per millimetre squarefield. C, of total oxidant capacity (T-AOC) in kidney

glutathione (GSH) in kidney tissues. F, tissues. D, Content of catalase (CAT) activity in kidne tissues. E, Conte were expressed as mean \pm SD. *significant difference vs, sham Content of superoxide dismutase (SOD) in kidney t group (p < 0.05); #significant difference vs, DOX §

Lixisenatide Activated TGF ad Pathway in DOX-Induce brosis Furthermore, immun stocher staining showed that positive TGE- β 1. ression Smad2/3 and α -SMA in DOX group were Aficant fer than the sham group. By tast, the Lix de group presented lower f these genes are 5A). ults also found that the Moreover, We 2rn br TGF- β 1, Smad2/3, ression level protein ex Smad4. agen I, collagen nd α-SMA were y downregulated in the Lixisenatide remar then co ared with the DOX group (Figro gure

scussion

ances have been made in the study al fibrosis, such as theories of intact hron, trade-off hypothesis, glomerular hypern, dyslipidemia, uremic toxins and nutrideficiencies. Renal fibrosis is a common tion pathway for various kidney diseases to develop into chronic renal failure^{1,3,4}. Therefore, renal fi-

brosis and other organ fibrosis are hotspots in medical researches⁵. Organ fibrosis is a pathological process in which abnormal growth and excessive deposition of extracellular matrix can be found in tissues due to inflammation-induced necrosis of organ parenchymal cells²¹. Organ fibrosis and sclerosis are different stages of development in the same pathological process. Meanwhile, organ fibrosis is an important intermediate link and a necessary pathway for further development of organ sclerosis^{21,22}. Scholars⁶⁻⁸ have indicated that renal fibrosis includes tubulointerstitial fibrosis and glomerular sclerosis. Meanwhile, the two lesions are closely related. It is generally believed that renal fibrosis is an inevitable step in the progression of chronic renal failure.

Injury stimuli first alter the local immune microenvironment, leading to renal intrinsic cell activation, pro-inflammatory cytokine production and release, mononuclear-macrophages, recruitment of T lymphocytes to lesioned renal tissue. Subsequently, mesangial cells, myofibroblasts and tubular epithelial cells can be activated or transdifferentiated by relevant cytokines. A large amount of ECM is produced and deposited,



Figure 4. Lixisenatide decreasion system of the protein system of the expression of NF- κ B/TNF- α relevant proteins. **A**, Content of TC (α in h, α and β an

which f er causes fibrou. formation, microva ar network reduction, enal tissue remodeli nchymal damage and even the renal p ction³⁻⁵, <u>Inflammation</u> is the basic loss patholog hange o KD and is also the trigbrosis, mainly characterig faci en infiltration and inflammatory immuh or secretio.^{21,22}. During renal fibrosis, the me itial injury is caused by the activatu Je cytokines and chemokines. It is erved that many chemokine receptors, such as 13, CX3CR1, CCR1, CCR2, monocyte cheic factor 1 and osteopontin are upregulated mo in lesioned kidneys^{23,24}. Current researches have demonstrated that the pharmacological effects of Lixisenatide on treating fibrotic lesions are closely related to the inhibition of the inflammation and collagen proliferation, as well as the enhancement of collagenase activity and collagenolysis²⁵.

Lipid metabolism often occurs in patients with CKD. Foam cells formed by macrophage phagocytic lipoproteins usually exist in areas of glomerular sclerosis and interstitial fibrosis^{4,8}. Some researches have shown that macrophages, mesangial cells and tubular cells can produce reactive oxygen free radicals and oxidized lipoproteins. Oxidize low-density lipoproteins stimulate the expressions of inflammatory and fibrotic cytokines, and induce cell apoptosis. Meanwhile, they also produce reactive oxygen free radicals, ulti-



Figure 5. Lixisenatide supplementation enhanced the expression to the tochemical staining of TGF- β 1, Smad2/3, and α -SMA. **B**, Protein I, collagen IV, and α -SMA in different groups. GAPDH was used a expression. Data were expressed as mean \pm SD. *significant difference DOX group (p<0.05).

els of TGF-β/A ssions of TGF oternal cor

relevant proteins. **A**, Immuno-his-Smad2/3, Smad4, Smad7, collagen to normalize the volume of protein (p<0.05); #significant difference vs.

mately leading to massive invasion of macr ges, apoptosis and accumulation of extracely matrix and tissue damage⁵⁻⁷.

Nuclear factor-kappa B () fast-re sponse transcription fact comm found in the form of heterodi of p50/ compo p65^{15,16}. Due to its binding in the cytoplasm in NF-KB can nactive be stimulated by tiple stimu which is subsequently d from IkB, located ve state, and bond to into the nuck in a The expres. evels of downstrethe κB sit am targ enes are regular NF- $\kappa B^{15,18}$. It is at the target genes of F-KB are widely know in pat ogical processes, such as hepainv 1, fibrosic and apoptosis¹⁶. A large tic n dies hav nown that the activity of number B is ar increased in patients with ating its role in the injury of epatitis coholic ste nohepatitis (NASH). Lipid penor ming NASH activates NF-κB. Subrc vated nuclear NF-κB promotes the duction of other inflammatory factors, aggraaflammatory response and in turn activates , eventually forming a positive feedback re-NF gulation. The gradual expansion of inflammatory response remarkably damages hepatocytes^{16,17}.

it studies have shown that certain factors and cytokines (such as IL-1) in DOX-induced renal fibrosis model are involved in the process of tubule-interstitial injury, including TGF-β, CTGF, IL-1, MCP-1, and osteopontin. They also promote the accumulation of extracellular matrix²⁶. Phenotypic transformation of mesangial cells, glomerular or tubular epithelial cells have been proved to be essential in renal tissue sclerosis or fibrosis. Meanwhile, it has been reported that under the induction of certain growth factors, renal interstitial fibroblasts can be transformed into myofibroblasts, renal tubular epithelial cells or glomerular epithelial cells (podocytes or Bowman's capsule epithelial cells). Previous works²⁷ have indicated that MCP-1 is capable of inducing trans-differentiation of renal tubular epithelial cells to myofibroblasts as well. The increased amount of renal interstitial fibroblasts is a crucial marker of interstitial fibrosis, serving as an evaluable marker of renal dysfunction and the prognosis of affected patients27. In addition, renal tubule-interstitial extracellular matrix, such as collagen I, II, IV, laminin, and fibronectin are significantly upregulated in DOX-induced renal fibrosis model.

In the present study, NF- κ B expression in the DOX group was significantly higher than that of

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the Lixisenatide group, indicating that ROS accumulation during the progression of renal fibrosis could activate NF- κ B. Moreover, Nrf2 was highly expressed in the Lixisenatide group than that of the DOX group, suggesting that Lixisenatide activated Nrf2 and alleviated inflammatory response induced by renal fibrosis.

Conclusions

We found that lixisenatide protects pathological kidney lesions, oxidative stress and inflammatory response after doxorubicin-induced renal fibrosis in rats by inhibiting NF- κ B/TNF- α and TGF- β /Smad pathways. Therefore, it might be a potential treatment target for renal fibrosis.

Conflict of Interests

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The authors declare that they have no conflict of interest.

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