UII/GPR14 is involved in NF-κB-mediated colonic inflammation in vivo and in vitro

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Abstract. The present study was conducted to investigate the molecular mechanism of urotensin II (UII) and its receptor, G protein-coupled receptor 14 (GPR14), in colonic inflammation. Urantide, a special antagonist of GPR14, and GPR14-siRNA were used to inhibit GPR14 signaling in dextran sulfate sodium (DSS)-induced inflammation in mice and Caco-2 cells. The results showed that urantide alleviated rectal bleeding, histological injury and production of interleukin (IL)-17 and tumor necrosis factor-α (TNF-α) caused by DSS in mice. GPR14-siRNA transfection subsequent with GPR14 inhibition reduced DSS-induced interferon-γ (IFN-γ) production in Caco-2 cells. Meanwhile, both in vivo and in vitro data demonstrated that inhibition of UII/GPR14 alleviated nuclear factor-κB (NF-κB) activation caused by DSS. In conclusion, UII/GPR14 signaling was involved in DSS-induced colonic inflammation and its inhibition may serve as a potential therapeutic target, which may be associated with the NF-κB signaling pathway.

Introduction

Urotensin II (UII) and its receptor, G protein-coupled receptor 14 (GPR14), are widely expressed throughout the cardiovascular, pulmonary, central nervous, renal and metabolic systems (1,2). UII/GPR14 signaling has been reported to be involved in various biological functions under both physiological and pathological conditions, such as enhancing foam cell formation, modulating insulin and catecholamine release, and regulating food intake (3). Meanwhile, elevated plasma levels of UII and expression of UII and UT have been noted in various diseased conditions, including hypertension, atherosclerosis, heart failure, pulmonary hypertension, diabetes, renal failure and metabolic syndrome (3), suggesting a potential marker of disease activity. Meanwhile, pharmacological and genetic inhibition of the UII/GPR14 signaling pathway has been reported to serve as a protective mechanism in various pathological conditions (4), such as liver inflammation, ischemia-reperfusion injury and pulmonary arterial hypertension (5-7).

Recent studies suggest a potential immune inflammatory function of the UII/GPR14 system (8). In lipopolysaccharide challenged mice, inhibition of UII/GPR14 signaling by urantide treatment markedly alleviated the production of pro-inflammatory cytokines such as tumor necrosis factor-α (TNF-α), interleukin (IL)-1β and interferon-γ (IFN-γ) via inactivation of the nuclear factor-κB (NF-κB) pathway (9), which is known to be a major pro-inflammatory transcription factor controlling a wide number of genes, including cytokines (10). Meanwhile, urantide, a special antagonist of GPR14, has also been demonstrated to attenuate the inflammatory response via inhibition of p38 mitogen-activated protein kinase phosphorylation in lipopolysaccharide (LPS)-stimulated Kupffer cells (9). However, UII/GPR14 signaling in intestinal inflammation and the potential therapeutic function in intestinal inflammatory disease are still obscure. Thus, in the present study, we investigated UII/GPR14 expression and the effects of inhibition of UII/GPR14 via urantide treatment on dextran sulfate sodium (DSS)-induced inflammation in mice and Caco-2 cells.

Materials and methods

Animal model and groups. Thirty female Kunming mice (weighing, 23.14±1.37 g) were randomly assigned into three groups: a control group (control, n=10), a DSS-treated group (DSS, n=10) in which mice received 5% DSS (Kayon Biotechnology Co., Ltd., Shanghai, China) instead of tap water for 7 days to establish an inflammatory bowel disease (IBD) model, and a urantide group (DSS + UR, n=10) in which mice received 5% DSS and an intravenous injection of 0.6 mg/kg urantide (Peptides, Louisville, KY, USA) dissolved in saline. The control and untreated challenged animals received the same volume of saline alone. The dosage of urantide used in the present study was according to a previous study (4). All mice were housed in polycarbonate cages in a room with controlled temperature (25±3˚C), humidity (50±5%) and a
12 h cycle of light and dark. They were allowed free access to laboratory strip chows throughout the experimental period. After the experimental period, each animal was weighted to calculate average weight gain, and then each mouse was sacrificed and colon length and weight were measured. In addition, colon tissues from each mice were harvested and immediately frozen in liquid nitrogen and stored at -70°C for subsequent gene expression and western blot analyses. The present study was conducted according to the guidelines of the Declaration of Helsinki and all procedures involving animal subjects were approved by the Animal Welfare Committee of the Department of Gastroenterology and Yantai Municipal Laiyang Central Hospital.

Clinical evaluation of DSS colitis. Rectal bleeding and diarrhea were monitored daily. The appearance of blood in the stool was measured by hemoccult tests (Beckman Coulter), and was given a score from 0-4, defined as follows: 0 for no blood; 2 for positive hemoccult; and 4 for gross bleeding. The severity of diarrhea was given a score from 0-4, defined as follows: 0 for well-formed pellets; 2 for pasty and semi-formed stools; and 4 for liquid stools (11). All clinical scorings were performed in a blinded manner.

Histomorphometry determination. The morphological evaluation after DSS treatment was carried out using hematoxylin and eosin (H&E) staining according to a previous study (12). Briefly, one piece of each colon sample (0.5 cm) was main and eosin (H&E) staining according to a previous study (12). Fixation after DSS treatment was carried out using hematoxylin and eosin (H&E). All specimens were examined under a light microscope (Nikon, Tokyo, Japan).

The histological examination was performed in a blinded manner using a scoring system, previously validated and described (13). Three independent parameters were measured: severity of inflammation (0-3: none, slight, moderate and severe), depth of injury (0-3: none, mucosal, mucosal and submucosal and transmural), crypt damage (0-4, none, basal 1/3 damaged, basal 2/3 damaged, only surface epithelium submucosal and transmural), crypt damage (0-4, none, basal 1/3 damaged, basal 2/3 damaged, only surface epithelium intact, entire crypt and epithelium lost) and percentage of the involved area (0-4: 0%, 1-10%, 11-25%, 26-50% and 51-100%). All scores on the individual parameters together could result in a total score ranging from 0 to 14.

Cytokines. Intestinal segments were fragmented using an ultrasonic disruptor (Bandelin, Berlin, Germany) and homogenized in RIPA buffer (Takara, Tokyo, Japan). In addition, the homogenates were centrifugated at 10,000 rpm for 20 min, and the supernatants were used for further analysis.

IL-1β, IL-10, IL-17, TNF-α and IFN-γ were quantified using an enzyme-linked immunosorbent assay (ELISA) kit (R&D Systems, Inc., Minneapolis, MN, USA) according to the manufacturer's protocol.

NF-κB activity. Colonic NF-κB activity was measured according to ELISA kits (Cell Biolabs, San Diego, CA, USA).

Cell culture and treatment. Human colorectal adenocarcinoma-derived intestinal epithelial cells (Caco-2) were obtained from the American Type Culture Collection (ATCC; Manassas, VA, USA) and grown in Dulbecco's modified Eagle's medium (DMEM)/F12 supplemented with 1 mM sodium pyruvate, 20% fetal bovine serum (FBS) (HyClone, Logan, UT, USA), and 50 µM penicillin-streptomycin. The cells were then treated with 2% DSS for 4 days to induce inflammation according to a previous study (14).

siRNA transfection. Human GPR14 siRNA was obtained from Guangzhou RiboBio and transfected into cells using Lipofectamine RNAiMAX reagent according to the manufacturer's instructions (Invitrogen, Carlsbad, CA, USA). After transfection for 48 h, the medium over the cells was changed before subsequent treatments.

Real-time PCR. Total RNA was isolated from liquid nitrogen pulverized tissues with TRIzol reagent, and then treated with DNase I (both from Invitrogen) according to the manufacturer's instructions. Synthesis of the first strand (cDNA) was performed with oligo(dT)18 and SuperScript II reverse transcriptase (Invitrogen). Primers were designed with Primer 5.0 according to the gene sequence of mouse to produce an amplification product. The primer sets used were as follows: β-actin, F, GTCCACCCTTCCACGAGATG and R, GAAAGGGTTGTAAAAAGCGACG; IL-1β, F, CTGGTACCTCGGATGGATG and R, GGAGATTGTGCCTGACTGT; IL-6, F, TGCAAGAAGCTTCCATCCAGT and R, GTGAAGTGGGAAGCAGCC; IL-10, F, ACAGCGCGGAGAACAGAATAC and R, CAGCTGCTCCTTGTTGAAAG; IL-17 F, TACCCTCAACGTTCAGTTCAG and R, TTTCCCTCGCAGATGCAC; IFN-γ F, ATGAAACGCTACAGCTACTGCATTGGCTT and R, CCTCAAACCTGGCAATCTCATGAATGC; TNF-α F, AGGCACTTCCCCAAAGAT and R, TGGAGGCTCTGGGCCCAGAA. Real-time PCR was performed according to previous studies (15,16). Relative expression was normalized and expressed as a ratio to the expression in the control group.

Western blotting. Proteins were extracted with protein extraction reagents in accordance with the manufacturer's instructions (Thermo Fisher Scientific, Inc., Waltham, MA, USA). Proteins from each sample (30 µg) were separated by SDS-polyacrylamide gel electrophoresis and electrophoretically transferred to a polyvinylidene difluoride (PVDF) membrane (Bio-Rad, Hercules, CA, USA). The membranes were blocked in 7% evaporated milk, diluted in Tris-buffered saline containing 0.1% Tween-20 (TBS-226T) at room temperature for at least 2 h, and then incubated overnight at 4°C with the following primary antibodies: U11 (ab194676), GPR14 (ab78449), IκBα (ab32518), IκBβ (ab7547), p-p65 (ab86299) and NF-κBp65 (ab7970) (Abcam, Inc., Cambridge, MA, USA).

Mouse β-actin antibody (Sigma) was used for protein loading control. After primary antibody incubation, the membranes were washed with TBS-T and incubated with alkaline phosphatase-conjugated anti-mouse or anti-rabbit IgG antibodies (Promega, Madison, WI, USA) for 2 h at room temperature. The membranes were washed with TBS-T followed by three washes with TBS; signals were detected by the addition of 5-bromo-4-chloro-3-238 indolylphosphate/nitroblue tetrazolium (BCIP/NBT) solution (Sigma), then quantified and digitally analyzed using ImageJ program (NIH, Bethesda,
The intensity of each band was measured and subtracted from the background. The expression ratio of target proteins was normalized against β‑actin (17).

Statistical analysis. All statistical analyses were performed using SPSS 17.0 software. Group comparisons were performed using one‑way analysis of variance (ANOVA) to test homogeneity of variances via Levene's test followed by Tukey's multiple comparison test. Values in the same row with different superscripts are significant (P<0.05), while values with the same superscripts are not significantly different (P>0.05).

Results

Clinical indices. As shown at Fig. 1, altered body weight, colon length and weight, rectal bleeding and histological scores indicated a colitis model, which is consistent with previous studies (18,19). Compared with the IBD group, urantide injection markedly reduced rectal bleeding and histological scores after DSS treatment, suggesting a protective role in the colitis model.

Inflammatory response in mice. In the present study, ileum and colon samples were collected to test cytokine concentrations. The results exhibited that DSS treatment increased IL-17, TNF-α and IFN-γ levels in the ileum, and IL-1β, IL-17, TNF-α and IFN-γ levels in the colon (P<0.05) (Table I). Compared with the IBD group, urantide significantly reduced ileal IFN-γ and colonic IL-17 and IFN-γ concentrations.

Table I. Effects of urantide injection on pro-inflammatory cytokines (pg/ml) in DSS-challenged mice.

<table>
<thead>
<tr>
<th>Cytokine</th>
<th>Control</th>
<th>DSS</th>
<th>DSS + UR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ileum</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IL-1β</td>
<td>363.16±26.14</td>
<td>392.28±13.75</td>
<td>389.38±28.17</td>
</tr>
<tr>
<td>IL-10</td>
<td>103.82±8.15</td>
<td>91.73±10.65</td>
<td>112.53±9.67</td>
</tr>
<tr>
<td>IL-17</td>
<td>175.39±20.33a</td>
<td>253.19±17.16e</td>
<td>207.27±11.56ab</td>
</tr>
<tr>
<td>TNF-α</td>
<td>253.38±26.47b</td>
<td>297.16±24.85a</td>
<td>313.47±25.14a</td>
</tr>
<tr>
<td>IFN-γ</td>
<td>293.29±27.92b</td>
<td>431.39±22.55a</td>
<td>361.54±31.54ab</td>
</tr>
<tr>
<td>Colon</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IL-1β</td>
<td>317.54±14.85b</td>
<td>475.65±62.42a</td>
<td>418.82±19.46a</td>
</tr>
<tr>
<td>IL-10</td>
<td>86.26±6.59</td>
<td>116.87±14.09</td>
<td>105.49±11.65</td>
</tr>
<tr>
<td>IL-17</td>
<td>117.69±19.74a</td>
<td>218.65±19.49a</td>
<td>151.11±26.49a</td>
</tr>
<tr>
<td>TNF-α</td>
<td>236.59±24.49b</td>
<td>378.65±34.57a</td>
<td>295.34±24.37ab</td>
</tr>
<tr>
<td>IFN-γ</td>
<td>251.68±15.94a</td>
<td>476.15±36.11a</td>
<td>387.12±25.43ab</td>
</tr>
</tbody>
</table>

DSS, dextran sulfate sodium; UR urantide; IL, interleukin; TNF-α, tumor necrosis factor-α; IFN-γ, interferon-γ. Data are presented as mean ± SEM. The values having different superscripted letters are significantly different (P<0.05; n=6 or 8).
Meanwhile, colonic expression of cytokines were further determined via RT-PCR (Fig. 2). The results showed that colonic IL-1β, IL-10, IL-17, TNF-α and IFN-γ were significantly upregulated in the IBD group (P<0.05). Urantide injection markedly downregulated IL-17 and TNF-α expression (P<0.05).

**UII levels and GPR14 expression in mice.** Urantide was used to inhibit UII/GPR14 signaling in the present study. We found that colonic UII levels were significantly higher in the IBD group when compared to that in the control group (P<0.05), while urantide treatment failed to influence the colonic UII concentration (P>0.05) (Fig. 3).

Meanwhile, our data exhibited that UII and GPR14 were significantly upregulated after DSS treatment (P<0.05) (Fig. 3), suggesting that UII/GPR14 is involved in DSS-induced inflammation. Although we failed to observe any significant difference in colonic UII expression after urantide injection, urantide markedly inhibited GPR14 expression (P<0.05) (Fig. 3).
NF-κB signal in mice. Colonic NF-κB activity was significantly increased after DSS treatment, while urantide markedly reduced NF-κB activity in the mice (P<0.05) (Fig. 4). Furthermore, colonic NF-κB signal-related proteins (i.e. IκBα, IκBβ, p65 and p-p65) were assessed and the results showed that DSS significantly increased p65 phosphorylation and urantide treatment inhibited p65 activation (P<0.05) (Fig. 4). IκB is an inhibitory protein of the NF-κB signaling pathway, and we found that DSS markedly inhibited IκBα expression, while urantide alleviated IκBα upregulation caused by DSS exposure.

UII/GPR14 mediates NF-κB signaling in Caco-2 cells. The in vivo data indicated that UII/GPR14 is involved in DSS-induced inflammatory response in mice. Thus, we further validated the effect of UII/GPR14 in vitro via GPR14-siRNA transfection. The results exhibited that GPR14-siRNA transfection markedly reduced GPR14 expression, which further inhibited DSS-induced IFN-γ production (P<0.05) (Fig. 5).

The western blot results showed that DSS exposure also increased GPR14 expression in vitro and inhibition of GPR14 via GPR14-siRNA transfection markedly reduced GPR14 expression (P<0.05) (Fig. 5). Yet, GPR14-siRNA transfection failed to influence IκBα and IκBβ, and GPR14 downregulation significantly inhibited p65 phosphorylation (P<0.05) (Fig. 5).

Discussion

Previous studies have suggested that the UII/GPR14 signaling pathway is involved in pro-inflammatory responses via mediating pro-inflammatory cytokine expression (4,9,20,21). In the present study, we used urantide, a special antagonist of GPR14, to block GPR14 and the results indicated that GPR14 inhibition alleviated DSS-induced colitis in mice evidenced by reduced rectal bleeding and histological injury, indicating a protective role of urantide in the colitis model.

Pro-inflammatory cytokines and inflammation response in the gastrointestinal tract play a vital role in the progression of IBD (10,22-25). However, the effect of UII/GPR14 signaling on DSS-induced pro-inflammatory cytokine generation is not fully understood. In the present study, we found marked increases in pro-inflammatory cytokine (i.e. IL-1β, IL-10, IL-17, TNF-α and IFN-γ) levels and expression in the DSS challenged mice. Inhibition of the UII/GPR14 signal using urantide significantly reduced IL-17 and TNF-α generation, suggesting that the releases of these cytokines may be a consequence of UII/UTR system activation induced by DSS exposure. In lipopolysaccharide/D-galactosamine-induced liver inflammation, urantide has been demonstrated to prevent increases in pro-inflammatory cytokines such as IL-1β, TNF-α and IFN-γ (4). Meanwhile, the in vitro data revealed

![Figure 5. Effects of GPR14-siRNA on (A and B) GPR14 and pro-inflammatory cytokines, (C) IL-1β, (D) IL-10, (E) IL-17, (F) TNF-α and (G) IL-6 and (H and I) protein expression in DSS-challenged Caco-2 cells. Data are presented as mean ± SEM. The values having different superscripted letters are significantly different (P<0.05; n=3-6).](image-url)
that urantide treatment alleviated IFN-γ production caused by DSS in Caco-2 cells. Thus, we speculated that UII/GPR14 signaling may be involved in DSS-induced inflammation and inhibition of UII/GPR14 may exhibit an anti-inflammatory function in vivo and in vitro models.

UII/GPR14 signaling has been confirmed to mediate inflammatory response (9), while activity of the UII/GPR14 signal has never been evaluated in DSS-induced inflammation. In the present study, we found that the UII/GPR14 signal was significantly activated after DSS exposure evidenced by the upregulation of UII and GPR14. Similarly, Liang et al also reported that liver UII and GPR14 expression and serum UII levels were markedly enhanced in lipopolysaccharide/D-galactosamine-induced liver inflammation in mice (4). Meanwhile, both urantide and GPR14 siRNA transfection inhibited GPR14 expression, which further alleviated DSS-induced inflammation in mice and Caco-2 cells.

To investigate the mechanisms underlying the protective effect of UII/GPR14 signal inactivation on DSS-induced inflammation, we determined the effect of urantide and GPR14 siRNA transfection on signaling molecules of the NF-κB pathway. In the colon and Caco-2 cells. The NF-κB signaling pathway has been found to be associated with the expression of various cytokines and chemokines in response to cellular stimuli such as inflammation and oxidative stress (26-29). In the present study, NF-κB signaling was activated after DSS treatment in vivo and in vitro, while inhibition of the UII/GPR14 signal via urantide or GPR14 siRNA transfection significantly inactivated NF-κB via reducing p65 phosphorylation. NF-κB activation requires IkBs, an inhibitory protein of NF-κB (25,26,29). The present data revealed that urantide markedly alleviated the inhibitory effect of DSS on IkBα. Thus, we concluded that UII/GPR14 may mediate IkBα/NF-κB signaling in DSS-induced inflammation. Inhibition of the NF-κB signaling pathway has been indicated to be a potential therapeutic target in inflammatory diseases, including IBD. For example, inactivation of the NF-κB pathway was found to downregulate pro-inflammatory cytokine expression in an IBD murine model (30). Meanwhile, pyrrolidine dithiocarbamate, an antioxidant agent and NF-κB inhibitor, has been confirmed to reverse the activation of the NF-κB signaling pathway and alleviate colonic inflammation caused by DSS treatment (31).

In conclusion, UII/GPR14 signaling is involved in DSS-induced inflammation in mice and Caco-2 cells. Inhibition of UII/GPR14 via urantide or GPR14 siRNA transfection alleviated DSS-induced inflammation and NF-κB activity. Thus, the beneficial effect of UII/GPR14 inhibition on the inflammatory response may be associated with the NF-κB signaling pathway.

References


