The Bradykinin B2 Receptor Gene Is a Target of Angiotensin II Type 1 Receptor Signaling

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Cross-talk between G protein–coupled receptors (GPCR) is known to occur at multiple levels, including receptor heterodimerization and intracellular signaling. This study tested the hypothesis that GPCR cross-talk occurs at the transcriptional level. It was demonstrated that the bradykinin B2 receptor gene (BdkrB2) is a direct transcriptional target of the angiotensin II (AngII) type 1 receptor (AT1R) in collecting duct cells. AngII induced BdkrB2 mRNA expression in mouse inner medullary collecting duct cells, and this effect was abrogated by AT1R blockade; in contrast, AT2R blockade was ineffective. Actinomycin D, an inhibitor of gene transcription, abrogated AngII-stimulated BdkrB2 expression. In addition, AngII produced dosage- and time-dependent increases in B2 receptor protein levels (2.9 ± 0.4 fold; P < 0.05). AngII stimulated phosphorylation of cAMP response element binding protein (CREB) on Ser-133 and assembly of p-CREB on the BdkrB2 promoter in vivo. Moreover, AngII induced hyperacetylation of BdkrB2 promoter–associated H4 histones, a chromatin modification that is associated with gene activation. Mutations of the CRE abrogated AngII-induced activation of the BdkrB2 promoter. AngII-treated inner medullary collecting duct cells exhibited augmented intracellular calcium signaling in response to bradykinin, confirming the functional relevance of AT1B2 receptor signaling. Finally, studies that were conducted in angiotensin type 1 receptor (Agtr1)-null mice revealed that BdkrB2 mRNA levels were significantly lower in the renal medulla of Agtr1+/− and Agtr1α/− than in Agtr1+/+ and Agtr1α/− mice. It is concluded that BdkrB2 is a downstream target of the AT1R-CREB signaling pathway. Transcriptional regulation represents a novel form of cross-talk between GPCR that link the renin-angiotensin and kallikrein-kinin systems.


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AngII signaling converges on the p53-CRE enhancer to stimulate BdkrB2 gene transcription. Our results indicate that AT1R signaling activates CREB phosphorylation and in vivo assembly of p-CREB on the BdkrB2 promoter in conjunction with histone hyperacetylation. AngII stimulates BdkrB2 gene transcription in IMCD3 cells via the AT1R. Studies in genetically targeted mice demonstrated that control of BdkrB2 gene by AngII occurs in the renal medulla and is mediated via the AT,R subtype A (AT1A) rather than AT1B. Thus, under conditions of augmented AngII and AT1R signaling, B2R expression will be enhanced, thereby maintaining a balance of these two powerful counter-regulatory systems.

Materials and Methods

Animals and Tissues

The animal study protocol was performed in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals and approved by the Institutional Animal Care and Use Committee of Tulane University and Louisiana State University. Tissues were harvested from adult male wild-type, Agtr1A/B−/− mice (3 mo of age; n = 3 each). Detailed procedures and genotyping were previously published (19). Total RNA was extracted from the heart, kidney medulla, and kidney cortex using the Trizol reagent (Invitrogen, Carlsbad, CA). Total RNA from mouse embryonic kidneys (E13.5 d) was extracted with RNeAqueous-4PCR Kit (Ambion, Austin, TX).

Cell Culture

IMCD3 cells (20) were obtained from American Type Culture Collection (Rockville, MD) and maintained in DMEM/F-12 that contained 10% FBS (Invitrogen) at 37°C in a humidified incubator with 5% CO2. The cells were plated in duplicate in six-well plates at 2 to 3 × 105 cells/well 2 d before AngII treatment. For avoidance of the confounding effects of growth factors that are present in the serum on B2R expression, IMCD3 cells were grown in serum-free medium for 24 h before treatment. Icatibant (a specific B2R antagonist, 10−6 M) and candesartan (a specific AT1 receptor antagonist, 10−6 M) were chosen on the basis of their ability to inhibit the hemodynamic and growth-related effects of exogenously infused intravenous BK or AngII (21–25).

Immunofluorescence Staining

IMCD3 cells were cultured on chamber slides to 40 to 50% confluence, washed once with ice-cold 1× PBS, and fixed in ice-cold 4% paraformaldehyde/0.1% glutaraldehyde solution at room temperature for 10 min. After removal of the fixatives, cells were permeabilized with ice-cold PBS that contained 1% Triton X-100 for 15 min at room temperature and blocked with ice-cold PBS that contained 1% BSA for 10 min. Cells were incubated with a polyclonal rabbit B2R antibody (AS276-283, diluted 1:500, from Dr. Muller-Esterl, Mainz, Germany) (26) overnight at 4°C followed by extensive washes with PBS that contained 1% BSA. The cells were subsequently incubated with secondary antibody (Alexa Fluor 594; Molecular Probes, Eugene, OR; 1:200) for 2 h at room temperature and subjected to extensive washes thereafter. Cells were then incubated with the AT,R antibody (sc-1173; Santa Cruz Biotechnology, Santa Cruz, CA; 1:200) (27) overnight at 4°C followed by extensive washes with PBS that contained 1% BSA. Subsequently, the cells were incubated with secondary antibody (Alexa Fluor 488; Molecular Probes; 1:2000) for 2 h at room temperature. After extensive washes, the sections were covered with fluorescence mounting medium that contained a nuclear stain, DAPI. Immunofluorescence images were obtained with an Olympus BX51TRF microscope and an integrated Magnafire SP Digital “Firewire” Camera System (Olympus American Inc., Center Valley, PA). Negative controls included cells that were incubated in the absence of primary antibodies (single or both).

Western Blot Analysis

Immunoblotting was performed as described previously (10). Primary antibodies were a polyclonal anti-B2R (diluted 1:1500), CREB (Cell Signaling, Danvers, MA; 48H2, 9197; 1:1000), and p-CREB (Upstate Biotecnology, Lake Placid, NY; 06-519; 1:500). After three washes in PBS/Tween, the nitrocellulose membrane was exposed for 1 h at room temperature to the secondary antibody (horseradish peroxidase–linked goat anti-rabbit IgG). Immunoreactive bands were visualized using the ECL detection system (Amersham, Piscataway, NJ). The immunoblots were exposed to Hyperfilm-ECL films. The blots were then reprobed with a β-actin antibody (Sigma, St. Louis, MO; 1:4000) as a loading control.

Reverse Transcriptase–PCR

After DNase 1 digestion, 2 to 3 μg of total RNA was reverse-transcribed using SuperScript II Reverse Transcriptase (Invitrogen) in a 22-μl volume. One to three microliters of the reaction volume was used in a PCR volume of 25 μl. The primer sequences for BdkrB2 were as follows: Upstream 5′-AGA ACC TCT TTT CTC GCC-3′; downstream 5′-CGT GAC CTC CTT GAA CT-3′. PCR was performed at 94°C for 50 s, 60°C for 60 s, and 72°C for 90 s, 1.0 mM MgCl2, 30 cycles. The product size was 572 bp. The primer sequences for Agtr1A were as follows: Upstream 5′-GCA TCA TTT TGG TGG C-3′; downstream 5′-GAA GAA CAC CAC AAT CGC C-3′. PCR was performed at 94°C for 1 min, 55°C for 1 min, and 72°C for 2 min, 2.0 mM MgCl2, 35 cycles. The product size was 641 bp. The primer sequences for Agtr2 were as follows: Upstream 5′-AGT GCA TCG GCC AGG AGC TG-3′; downstream 5′-GAC AAA ACA GTG AG-3′. PCR was performed at 94°C for 1 min, 53°C for 1 min, and 72°C for 2 min, 1.5 mM MgCl2, 35 cycles. The product size was 309 bp. The primer sequences for glyceraldehyde-3-phosphate dehydrogenase (GAPDH) were as follows: Upstream 5′-CCC CTG GCC AAG GTC ATC CAT GAC AAC TT-3′; downstream 5′-GCA CAT GAG GTC CAC CCT GCT GCT GTA-3′. PCR was performed at 94°C for 50 s, 55°C for 50 s, and 72°C for 90 s, 1.5 mM MgCl2, 27 cycles. The product size was 516 bp. PCR reactions were visualized on a 1% agarose gel, and densitometric analysis was performed with the ChemiImager 4400 program (Alpha Innotech, San Leandro, CA).

Intracellular Calcium Measurements

Peak increases in [Ca2+]i after activation of B2R in IMCD3 cells were evaluated using fluorescence microscopy. IMCD3 cells that were maintained in DMEM/F-12 medium that contained 10% FBS were plated on coverslips in 5-cm-diameter cell culture dishes, mounted in 200-μl perfusion chambers (Warner Instruments, Eugene, OR), and examined using a Nikon Diaphot (Melville, NY) inverted microscope with an attached Photon Technology International (Birmingham, NJ) deltascan fluorescence-based spectrophotometry system with excitation wavelengths set at 340 and 380 nm and emission collected at 510 nm. Cells were continually superfused with PBS that contained 1.8 mM Ca2+ at 37°C. Five fluorescence measurements per second were taken with appropriate background elimination. Fluorescence experiments were calibrated in vitro using the methods described by Grynkiewicz et al. (28).

For testing the effect of AngII on B2R-induced [Ca2+]i signaling, cells that were plated on coverslips were changed to serum-free medium for...
24 h and treated with $10^{-7}$ M AngII for the subsequent 24 h. Calcium fluorescence was measured in cells that were exposed to PBS for 0 to 100 s, followed by $10^{-7}$ M bradykinin (BK; 400 s) before being returned to PBS. Untreated cells were used as control. For evaluation of the effect of icatibant (formerly known as HOE-140), a selective B2R antagonist, on AngII-stimulated B2R signaling, calcium fluorescence was measured in AngII-treated cells ($>24$ h) that were exposed to icatibant ($10^{-6}$ M) for 100 s, followed by exposure to $10^{-7}$ M BK in the continued presence of icatibant (300 s) before being returned to PBS. Fluorescence intensity measurements were converted to calcium concentrations on the basis of calibration procedures that were previously described (29). Baseline $[Ca^{2+}]_i$ was calculated by taking the average value of the 5-s period before addition of BK to the chamber. Peak change in $[Ca^{2+}]_i$, in response to BK, was determined as the peak value of $[Ca^{2+}]_i$, reached on addition of BK minus the baseline. Mean values ± SE are presented for the peak time point value.

Promoter Constructs, Transient Transfection, and Reporter Assays

The BdkrB2 $-94/+55$-CAT promoter-reporter construct was previously described (8,9). The mutant BdkrB2 $-94$(P1mut)/CAT, $-94$(CRE-mut)/CAT, $-94$(P1mutCREmut)/CAT constructs were generated by introducing point mutations in the $-94/+55$-CAT construct at the p53-binding site (P1) and CREB binding site (CRE), singly or in combination, using the QuickChange site-directed mutagenesis system (Stratagene, La Jolla, CA) (10). All constructs were subjected to DNA sequencing to verify the sequence and orientation.

IMCD3 cells that were maintained in DMEM/F-12 medium that contained 10% FBS were plated in duplicate in 3.5-cm-diameter six-well plates at $2.0 \times 10^5$ cells/well 2 d before transfection, followed by 24 h of serum-free medium 1 d before transfection. Cells that were grown to 80% confluency were transfected with 1 μg of DNA per well of each promoter-reporter construct. One microgram of DNA per well pCAT3 Basic vector (empty vector) was transfected as a control. A β-galactosidase vector, pSVZ (0.4 μg of DNA/well; Promega, Madison, WI), was co-transfected to correct for transfection efficiency. Transfections were performed using Lipofectamine Plus Reagent (Life Technologies, Rockville, MD) according to the manufacturer’s recommendations. Four hours after transfection, the medium was replaced with fresh medium in the absence of serum. Twenty-four hours after transfection, cells were either treated with AngII ($10^{-7}$ M) or left untreated, and 6 h later, cell extracts were prepared using a reporter lysis reagent (Promega). Aliquots of cell lystate were analyzed for CAT activity after normalization for protein content or β-galactosidase activity as described previously (9).

Chromatin Immunoprecipitation

Chromatin immunoprecipitation (ChIP) assays were performed using reagents and protocols from Upstate Biotechnology as described previously (10,30). IMCD3 cells were plated in 10-cm culture dishes and starved for 24 h before AngII treatment. After AngII stimulation ($10^{-7}$ M) for the indicated time periods, cells were incubated with 1% formaldehyde solution in growth medium for cross-linking (10 min at room temperature) followed by addition of 1 ml of 10% glycine to quench unreacted formaldehyde for 5 min at room temperature. Cell pellets were resuspended in SDS lysis buffer that contained protease inhibitor and phosphatase inhibitor cocktails. DNA was sheared by sonicating the cell lystate to produce an average DNA fragment size between 200 and 1000 bp and diluted 10-fold in ChIP dilution buffer. Immunoprecipitation was performed with antibodies to p-CREB (Ser133; Upstate Biotechnology; 06-519; 20 μl), CREB-1 (Santa Cruz; 240, sc-58; 25 μl), acetylated H4 (Upstate Cell Signaling; 06-866; 1:150 dilution), or control normal Ig (IgG) antibodies overnight at 4°C. DNA-protein antibody complexes were captured on protein A/G-conjugated agarose beads. DNA-protein cross-links were reversed at 65°C overnight followed by proteinase K treatment to free DNA. Immunoprecipitated DNA was purified by spin column and subjected to PCR. Sequences of the primers that were used for PCR of the mouse B2R gene that flanked the p53-CRE enhancer are as follows: Forward primer 5′-AGG GCG GAG GTG CCC AGG AGA GTG ATG ACA-3′; reverse primer 5′-GCT TCT GTG TTG TAG GGA GT-3′.

Statistical Analyses

All data are expressed as means ± SEM and analyzed by t test or one-way ANOVA. All experiments were performed at least three times in duplicate. Differences between experimental groups were considered statistically significant at $P < 0.05$.

Results

AngII Upregulates BdkrB2 Gene Expression in Renal Collecting Duct Cells

Previous studies suggested the presence of AT1R and B2R in several renal cell types, including vascular (smooth muscle, juxtaglomerular, endothelial) and collecting duct cells (31,32). Using double-immunofluorescence staining, we show that AT1R and B2R are coexpressed on the cell membrane of mouse IMCD3 cells (Figure 1A). Control experiments that were conducted in the absence of primary antibodies revealed no specific staining (data not shown). AngII signals via two receptors, AT1R and AT2R, which are coupled to different intracellular signaling pathways. Reverse transcriptase–PCR (RT-PCR) assays were therefore performed to determine the AngII receptor type(s) expressed by IMCD3 cells. The results revealed that

Figure 1. Expression of bradykinin (B2R) and angiotensin II (AngII) type 1 receptor (AT1R) in inner medullary collecting duct (IMCD3) cells. (A) Immunofluorescence staining of B2R (red) and AT1R (green) in IMCD3 cells. Merged image (yellow). (B) Reverse transcriptase–PCR (RT-PCR) showing expression of AT1R mRNA in IMCD3 cells. In contrast, there is little if any expression of AT2R mRNA in these cells. RNA from embryonic mouse kidneys (E13.5) was used as a positive control, and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was used as a loading control. Magnification, ×100.
IMCD3 cells express predominantly AT1R transcripts; in contrast, only minimal amounts of AT2R mRNA were detectable (Figure 1B). Positive control RNA samples from E13.5 mouse embryonic kidneys showed abundant AT1R and AT2R transcripts. These results indicate that IMCD3 cells express predominantly the AT1R.

To test whether AngII regulates BdkrB2 gene expression, we stimulated quiescent IMCD3 cells with AngII (10^{-7} M), in the presence or absence of the AT1R antagonist candesartan (10^{-6} M). Total cellular RNA was extracted 3 or 24 h later. Untreated cells served as controls. RT-PCR showed a significant increase in BdkrB2 mRNA levels in AngII-treated as compared with controls (190% at 3 h and 70% at 24 h; P < 0.05; Figure 2). Candesartan abrogated the stimulatory effect of AngII on BdkrB2 gene expression, indicating that the AngII-induced increase in BdkrB2 mRNA is mediated by the AT1R (Figure 2).

Next, we investigated whether AngII upregulates B2R gene expression at the protein level. Quiescent IMCD3 cells were treated with varying concentrations of AngII (10^{-10} to 10^{-6} M), whereas untreated cells were used as control. Whole-cell lysates were collected 8, 24, and 48 h after AngII treatment and subjected to Western blot analysis. As shown in Figure 3, AngII stimulated dosage-dependent increases in B2R protein expression as early as 8 h after treatment and persisted for up to 48 h (8 h 2.6 ± 0.3-fold; 24 h 2.9 ± 0.4-fold; 48 h 2.2 ± 0.3-fold relative to control; P < 0.05 versus control). Collectively, these results demonstrate that AngII stimulates BdkrB2 gene expression in IMCD3 cells at mRNA and protein levels.

**AngII-Stimulated BdkrB2 Is Actinomycin D Sensitive**

To explore the molecular mechanisms by which AngII regulates B2R gene expression, we pretreated quiescent IMCD3 cells with the transcriptional inhibitor actinomycin D (10^{-6} M) for 2 h, followed by AngII (10^{-7} M) for 6 h. Untreated cells were used as a control. Actinomycin D (10^{-6} M) abrogated the increase of BdkrB2 mRNA levels induced by AngII (AngII 3.0 ± 0.69; AngII + actinomycin 0.45 ± 0.17 versus control; P < 0.05; n = 4; Figure 4A). These data suggest that AngII upregulates B2R gene expression at the transcriptional level.

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**Figure 2.** AngII upregulates BdkrB2 gene expression. Quiescent IMCD3 cells were treated with AngII (10^{-7} M) in the absence or presence of the AT1R antagonist candesartan (Cand; 10^{-6} M). Untreated cells served as time control. Total RNA was extracted 3 or 24 h after treatment, and RT-PCR was performed using mouse-specific BdkrB2 and gapdh primers. (A) Ethidium bromide–stained gel. (B) Bar graph representing the means ± SEM of the ratios of BdkrB2 and gapdh band intensities expressed relative to control. *P < 0.05 versus other groups (n = 3); †P < 0.05 versus AngII.

**Figure 3.** AngII upregulates B2R protein expression. Quiescent IMCD3 cells were treated with increasing concentrations of AngII (10^{-10} to 10^{-6} M). Untreated cells served as control. Whole-cell lysate was collected 8, 24, and 48 h after AngII treatment and was subjected to Western blot analysis. (A) Bar graph representing the ratios of B2R/β-actin band intensity expressed relative to control. (B) Immunoblots using anti-B2R and β-actin antibodies. B2R protein size is 43 kD. *P < 0.05 versus control; †P < 0.05 versus 10^{-8} M AngII.
AngII Stimulates BdkrB2 Promoter Activity

In previous studies, we demonstrated that a highly conserved BdkrB2 promoter fragment that extends from −94 to +55 bp, relative to the transcription start site at +1, is sufficient to drive reporter CAT activity in IMCD3 cells (10). This promoter region contains a composite cis-regulatory module for the transcription factors p53 and CREB, and cooperative interaction between the two proteins is essential for optimal BdkrB2 promoter activity (10). Accordingly, we tested the effects of AngII on the activity of BdkrB2 −94/+55-CAT promoter activity using transient transfection assays in IMCD3 cells (Figure 4B). The promoterless vector pCAT3Basic served as a control. Twenty-four hours after transfection, cells were treated with AngII (10−7 M) for 6 h and whole-cell lysates were harvested and assayed for CAT activity. As shown in Figure 4, C and D, AngII stimulated BdkrB2 promoter-driven reporter activity to 1.7 ± 0.08-fold (P < 0.05 versus untreated cells; n = 5), whereas the promoterless construct pCAT3Basic was not affected by AngII.

To determine the functional importance of p53 and CREB in AngII-mediated transcriptional activation, we introduced mutations in the −94/+55-CAT construct at the p53(P1)- and CRE cis-regulatory elements, either singly or in combination. These mutations have been shown to inhibit the binding of p53 and CREB to their respective sites (10). IMCD3 cells were transiently transfected with −94/+55-CAT construct, the single-mutant constructs −94P1mut/CAT and −94CREmut/CAT, and the double mutant construct −94P1mutCREmut/CAT. Twenty-four hours after transfection, the cells were treated with AngII (10−7 M) for 6 h. CAT assays revealed that mutations in CRE or p53 binding sites, either singly or in combination, abrogated AngII-stimulated BdkrB2 promoter activity (Figure 5).

AngII Stimulates CREB Phosphorylation and Association with BdkrB2 Promoter

Multiple signal transduction pathways are known to converge on the transcription factor CREB, leading to its phosphorylation on serine 133 (33). This posttranslational modification promotes the interaction of CREB with its co-activator, CBP/p300, and enhances CREB transcriptional activity (11,13). In addition, CREB phosphorylation has been shown to facilitate the interaction of CREB with p53 and thus their co-recruitments to target promoters (15,33). For examination of the effect of AngII on CREB phosphorylation, IMCD3 cells were incubated with AngII (10−7 M) and cellular lysates were harvested at various time points ranging from 0 to 60 min. The lysate was subjected to Western blotting using p-ser133-CREB or CREB antibodies. The results showed that AngII significantly increased p-CREB/CREB ratio at 30 min after stimulation (Figure 6, A and B).

We next asked whether AngII stimulation promotes the binding of p-CREB to the endogenous BdkrB2 promoter using ChIP-PCR. As shown in Figure 6C, baseline occupancy of BdkrB2 promoter by p-CREB was minimal and comparable to background levels. This was not due to the lack of CREB binding to the promoter (Figure 6C). Exposure of IMCD3 cells to AngII for 15 min induced a considerable increase in p-CREB at the BdkrB2 promoter to 3.3-fold (n = 2) relative to control. Consistent with the ability of p-CREB to interact with and recruit CBP/p300, this effect was associated with hyperacetylation of promoter-associated histone H4 (Figure 6D). These results indicate that AngII stimulates association of p-CREB with the BdkrB2 pro-
peak response to BK to 16.10 ± 2.59 nM in AngII-treated cells ($n = 10$; $P < 0.05$; Figure 7B). The results indicate that AngII-treated cells respond by increasing B2R mRNA and protein synthesis as well as by enhanced membrane-associated B2R signaling.

**Reduced BdkrB2 Gene Expression in Agtr1A-Null Mice**

To determine whether the BdkrB2 gene is regulated by AngII in vivo, we compared $BdkrB2$ mRNA levels in wild-type versus $Agtr_1^{-/-}$, $Agtr_1^{-/-}$, or $Agtr_1^{-/-}/B_1^{-/-}$ mice. RT-PCR revealed that $BdkrB2$ mRNA levels, factored for GAPDH, were 53% lower in the renal medulla of $Agtr_1^{-/-}$ and $Agtr_1^{-/-}/B_1^{-/-}$ than in $Agtr_1^{-/-}$ and $Agtr_1^{-/-}/B_1^{-/-}$ mice (0.14 ± 0.01 versus 0.3 ± 0.06 units; $P < 0.05$; Figure 8A). In contrast, the renal cortex showed no differences in $BdkrB2$ gene expression among the genotypes (data not shown). $BdkrB2$ mRNA levels tended to be lower in the heart of $Agtr_1^{-/-}$ and $Agtr_1^{-/-}/B_1^{-/-}$ than in $Agtr_1^{-/-}$ and $Agtr_1^{-/-}/B_1^{-/-}$ mice (0.54 ± 0.04 versus 0.64 ± 0.02; factored for GAPDH; $P = 0.07$; Figure 8B). These results together with the cell culture data support the notion that $BdkrB2$ gene expression is under physiologic regulation by AngII.

**Discussion**

AngII, acting via AT$_1$R, stimulates renal tubular sodium and water reabsorption and causes renal vasoconstriction. In contrast, kinins promote natriuresis and diuresis and increase renal blood flow via stimulation of B2R. Therefore, the balance between the RAS and KKS affects salt and water handling, volume homeostasis, and BP control (2). This is well illustrated in $BdkrB2^{-/-}$ mice, which are susceptible to AngII-induced hypertension (22). AT$_1$R and B2R belong to the seven-transmembrane GPCR family of receptors and are coexpressed in several cell types, including vascular and renal epithelial cells, including the renal collecting duct (31,32). Both short- and long-loop interactive pathways exist between the AT$_1$R and B2R. In the short-loop pathway, AT$_1$R and B2R communicate directly with each other. The two receptors form stable heterodimers, resulting in enhanced activation of AT$_1$R signaling (4). The AT$_1$R-B2R complex seems to signal as a “super AT$_1$R.” It is interesting that a significant increase in AT$_1$R–B2R heterodimerization occurs in hypertensive preeclamptic women and is associated with a five-fold upregulation in B2R levels (5). Our findings in this study support the concept that $BdkrB2$ gene expression is positively regulated by AngII and that signaling via AT$_1$R–B2R activates B2R expression.

The long-loop pathway that connects AT$_1$R and B2R receptors involves gene regulation. This cross-regulatory pathway seems to operate in cardiac/vascular smooth muscle cells and, as shown in this study, in renal collecting duct cells. Infusion of subpressor doses of AngII into wild-type mice upregulates cardiac myocyte $BdkrB2$ mRNA by 47% (6). Furthermore, treatment of vascular smooth muscle cells in culture with AngII produces a time-dependent induction of $BdkrB2$ mRNA that is maximal at 3 h and declines to baseline by 24 h. This effect is blocked by AT$_1$R antagonism (6). This study provides new evidence that the $BdkrB2$ gene is a downstream target of the AT$_1$R in vivo. This conclusion is based on the following find-
ings: (1) Agtr1\textsuperscript{A−/−} or Agtr1\textsuperscript{B−/−} mice have reduced levels of BdkrB2 mRNA in the renal medulla as compared with wild-type or Agtr1\textsuperscript{B−/−} mice; (2) exogenous AngII upregulates BdkrB2 mRNA and protein expression in IMCD3 cells, and this effect can be blocked by AT1R blocker; (3) AngII stimulates BdkrB2 gene transcription and promoter-driven reporter activity; (4) AngII promotes occupancy of the BdkrB2 promoter by activated CREB; and (5) AngII-treated cells exhibit augmented B2R-mediated intracellular calcium signaling in response to acute bradykinin stimulation. The AT1R–B2R feed-forward mechanism creates a new level of cross-talk between two GPCR with wide-ranging implications on renal salt and water handling, BP regulation, and growth and differentiation.

In the kidney, AT1R and B2R are expressed in a developmentally regulated manner, increasing from relatively low levels in the embryo to peak levels perinatally (34). Moreover, immuno-
histochemical analysis has shown that AT1R are more widely distributed along the nephron than previously recognized, including renal vascular smooth muscle and proximal and distal epithelial sites. B2R is expressed primarily in collecting duct cells but also in the vascular and interstitial elements within the kidney. In rodents, there are two subtypes of AT1R: AT1A and AT1B. AT1A is the predominant subtype expressed in most tissues and mediates the known physiologic functions of AngII. This study demonstrated that BdkrB2 mRNA levels were 53% lower in the renal medulla of Agtr1A−/− mice than in Agtr1+/+ or Agtr1B−/− mice. In contrast, the renal cortex showed no differences, indicating that renal medullary BdkrB2 gene expression is regulated by AngII via the AT1A-R subtype. This study also showed that cardiac BdkrB2 mRNA levels tended to be lower in Agtr1A−/− than in Agtr1A+/+ mice, consistent with previous data showing that AngII induces BdkrB2 gene expression in cardiac myocytes and vascular smooth muscle cells. It is important to note here that because our studies were performed in adult mouse kidneys and IMCD cells, in which AT1A is the predominant AngII receptor, it is not surprising that AngII effects were mediated by AT1A, rather than AT1B or AT2. These findings do not, however, preclude similar interactions between these receptors and B2R in relevant tissues (adrenal, AT1B) or developmental stages (embryos, AT2).

IMCD3 cells are epithelial cells that retain many differentiated characteristics of the inner medullary collecting duct in vivo. As shown in this study, IMCD3 cells express AT1R and B2R mRNA and protein, whereas AT1R mRNA was difficult to detect. Therefore, IMCD3 cells express predominantly the AT1R subtype, which is compatible with the finding that AngII-induced B2R gene expression was completely abrogated by pharmacologic AT1R antagonism. As a GPCR, B2R generally signals through Gαq to stimulate phospholipase Cβ. In addition, B2R interacts with several other G proteins, including Gαs, Gαq, and Gα12/13. BK stimulates Gαq-sensitive phospholipase Cβ leading to phosphoinositide hydrolysis and intracellular free Ca2+ mobilization. Increased intracellular Ca2+ can in turn mediate the activation of cytosolic Ca2+-dependent endothelium nitric oxide synthase and Ca2+-sensitive phospholipase A2, thereby leading to nitric oxide and prostaglandin release. Our study showed that AngII treatment amplifies the intracellular Ca2+ response to bradykinin, whereas the B2R antagonist icatibant reduces the peak response to bradykinin in AngII-treated cells. Altogether, these data indicate that AngII stimulates B2R signaling in renal epithelial cells.

The BdkrB2 gene is located on human chromosome 14, mouse chromosome 12, and rat chromosome 6. The overall organization and structure of the BdkrB2 gene are generally similar in the three species: Three exons and two introns. The human and the rat but not the murine BdkrB2 genes have an alternatively spliced exon between exons 2 and 3, termed exon 2b. The human proximal BdkrB2 promoter regions suggest that these control factors exist in the human kidney. Previous analysis of the rat BdkrB2 gene identified a modular enhancer located at nucleotides −44 to −82 of rat BdkrB2 promoter, which is 100% conserved between rat and mouse and 85% conserved between rat and human genes. The BdkrB2 enhancer receives inputs from three transcription factors with overlapping expression in the collecting ducts, namely, p53, CREB, and KLF-4. CREB phosphorylation on Ser-133 is required for optimal stimulation of BdkrB2 promoter activity through recruitment of CBP and enhancement of p53–CREB interactions. Therefore, the renal collecting duct epithelial cells contain the necessary machinery for the regulation of BdkrB2 transcription. The sequence similarities between rodent and human proximal BdkrB2 promoter regions suggest that these control factors exist in the human kidney.

This study demonstrates that AngII-induced BdkrB2 gene expression is mediated transcriptionally because it is completely blocked by actinomycin D. Moreover, a promoter–reporter construct that harbors the modular BdkrB2 enhancer is responsive to AngII stimulation. Our investigations of the signaling inputs that link the AT1R to the BdkrB2 enhancer revealed several important findings. First, AngII stimulates the phosphorylation of CREB on Ser-133 within 15 to 30 min, a modification that is required for the interaction of CREB with its co-activator, CBP/p300, resulting in enhanced CREB tran-
scriptional activity (11). Second, as demonstrated by ChIP, AngII-stimulated occupancy of the BdkrB2 promoter by activated CREB, as well as induced promoter histone hyperacetylation. These events also occurred within 30 min. Acetylation of promoter-associated nucleosomes is thought to relax the chromatin structure, thereby facilitating access of transcriptional activators to their binding sites on DNA (38). Third, introduction of point mutations in the CRE- or p53-binding sites abrogated AngII-induced activation of the BdkrB2 promoter. Fourth, RT-PCR analysis revealed upregulated expression of BdkrB2 mRNA as early as 3 h after stimulation with AngII. On the basis of these findings, we propose a model in which AT(R) signaling stimulates sequential series of events that lead to phosphorylation of CREB and its binding to the p53-CRE modular enhancer in the BdkrB2 promoter. Phosphorylation of CREB promotes recruitment of CBP and p53 and stimulates localized histone hyperacetylation and activation of BdkrB2 transcription in renal epithelial cells. The enhanced transcription of BdkrB2 gene leads to augmented mRNA and protein levels, which results in enhanced intracellular calcium signaling and functional responses to bradykinin.

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Disclosures

None.

References


