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The cardiotonic steroid hormone marinobufagenin induces renal fibrosis: implication of epithelial-to-mesenchymal transition

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Fedorova LV, Raju V, El-Okdi N, Shidyak A, Kennedy DJ, Vetteth S, Giovannucci DR, Bagrov AY, Fedorova OV, Shapiro JI, Malhotra D. The cardiotonic steroid hormone marinobufagenin induces renal fibrosis: implication of epithelial-to-mesenchymal transition. Am J Physiol Renal Physiol 296: F922–F934, 2009.—We recently demonstrated that the cardiotonic steroid hormone marinobufagenin (MBG) induced fibrosis in rat hearts through direct stimulation of collagen I secretion by cardiac fibroblasts. This stimulation was also responsible for the cardiac fibrosis seen in experimental renal failure. In this study, the effect of MBG on the development of renal fibrosis in rats was investigated. Four weeks of MBG infusion triggered mild periglomerular and peritubular fibrosis in the cortex and the appearance of fibrotic scars in the corticomedullary junction of the kidney. MBG also significantly increased the protein levels and nuclear localization of the transcription factor Snail in the tubular epithelia. It is known that activation of Snail is associated with epithelial-to-mesenchymal transition (EMT) during renal fibrosis. To examine whether MBG alone can trigger EMT, we used the porcine proximal tubular cell line LLC-PK1. MBG (100 nM) caused LLC-PK1 cells grown to confluence to acquire a fibroblast-like shape and have an invasive motility. The expressions of the mesenchymal proteins collagen I, fibronectin, and vimentin were increased twofold. However, the total level of E-cadherin remained unchanged. These alterations in LLC-PK1 cells in the presence of MBG were accompanied by elevated expression and nuclear translocation of Snail. During the time course of EMT, MBG did not have measurable inhibitory effects on the ion pumping activity of its natural ligand, Na+–K+–ATPase. Our data suggest that the MBG may be an important factor in inducing EMT and, through this mechanism, elevated levels of MBG in chronic renal failure may play a role in the progressive fibrosis.

cardiotonic steroid hormone; ouabain; Na+–K+–ATPase; transcription factor Snail

CARDIOTONIC STEROIDS (CTSs), also known as cardiac glycosides or digitalis-like compounds, have been used for centuries to treat congestive heart failure. CTSs such as digoxin and digitoxin are still an important component of the clinical treatment of cardiac diseases (26, 65). CTSs were discovered in digitalis plants and amphibian tissues (25, 75). Recently, CTSs were found in the body fluids of mammals (5, 23, 29, 73). Moreover, elevated levels of endogenous CTSs including marinobufagenin (MBG) have been associated with various pathological conditions: essential hypertension (4, 24, 73), preeclampsia (3, 45, 83), diabetes (7), uremic cardiomyopathy (33), and cardiac failure (20, 48, 49, 66, 74). MBG, like other CTSs, binds to the extracellular domain of the α-subunit of Na+–K+–ATPase (16, 21), which, in addition to its well-known function of maintaining cellular electrochemical balance through pumping sodium and potassium ions, can act as a typical membrane receptor (14, 28, 35, 56, 73, 74, 84, 85, 88).

We hypothesized that MBG could promote renal fibrosis through EMT as well. First, because the elevated levels of MBG, or its analogs, accompany the conditions at which EMT occurs, including end stage renal disease (34), normal pregnancy, and preeclampsia (3, 45, 68, 83). Second, the interaction of MBG with the Na+–K+–ATPase activates Src and ERK1/2 protein kinase pathways and increases the production of reac-
tive oxygen species (20, 33, 35, 80, 81). Finally, as it was shown for ouabain, MBG possibly can trigger phosphoryosid
tide 3-kinase/protein kinase B (Akt) axis, stimulate NF-κB, and elevate the concentration of intracellular Ca\(^{2+}\) concentration (44, 55, 84, 88). Activation of these signaling pathways has been found to promote EMT in various animal and cellular models (11, 70, 78).

To test this hypothesis, MBG was administered to rats through a minipump for 4 wk. At the same time, we treated the porcine renal proximal tubular cell line LLC-PK1 with different concentrations of MBG and ouabain for up to 96 h. Several different aspects of EMT were analyzed including accumulation of collagen I in the kidney, expression of epithelial/ mesenchymal marker proteins, and transcription factors/regu-
lation of collagen I in the kidney, expression of epithelial/

**MATERIALS AND METHODS**

Chemicals of the highest purity available were obtained from Sigma (St. Louis, MO). Radioactive rubidium (\(^{86}\)Rb) was obtained from DuPont NEN Life Science Products (Boston, MA). MBG was isolated from toad venom (Bufo Marines) as described previously (6).

**Animals and animal model.** The animal studies were approved by the University of Toledo Health Science Campus Institutional Animal Use and Care Committee. Male, Sprague-Dawley rats were used for all of the studies. Rats weighing \(\geq 250\) g were subjected either to sham surgery or to osmotic minipump placement. MBG (10 μg/kg) was infused for period of 4 wk as described previously (20, 33).

**Renal morphology and immunohistochemistry.** Kidneys for light microscopy and immunoperoxidase staining were fixed in formalin and embedded in paraffin. Kidney sections (4 μm) were stained with saturated picric acid containing 0.1% Sirius red (Sigma) for 1 h in the dark. For immunoperoxidase staining, deparaffinized and rehydrated renal sections were blocked with 10% goat serum and 3% BSA (both from Sigma) in PBS for 2 h at room temperature. Primary antibodies against Snail (Abcam, Cambridge, MA) were diluted 1:400 in 5% goat serum and 3% BSA and applied to renal sections overnight at 4°C. After being washed, the sections were treated as suggested by ABC protocol (ABC Elite kit; Vector Laboratories, Burlingame, CA) and counterstained with methyl green. Immunoperoxidase staining of kidney sections for α-smooth muscle actin (α-SMA) was performed with α-SMA kit (Sigma) with NovaRed chromophore (Vector Laboratories) and counterstained with methyl green. Bright light and polarized images of picosirius red and α-SMA were taken on a Nikon Eclipse 80i microscope equipped with a Nikon camera head DS-Fi1 (Nikon, Tokyo, Japan) or Olympus B microscope (Olympus Optical, Hamburg, Germany) with Evolution MP color digital camera (Media Cybernetics, Bethesda, MD) for Snail. For quantitative morphometric analysis, eight randomly chosen cortical fields (at least 6 from each animal from experimental group) lacking major blood vessels were digitized and the collagen volume determined using the Imager J software (National Institutes of Health) as previously described (33).

**Cell culture.** The porcine kidney proximal tubule cell line LLC-

PK1 was obtained from the American Tissue Culture Collection (Manassas, VA). Cells were grown either on plastic culture plates for protein extraction or on glass slides for immunocytochemistry. LLC-

PK1 cells were grown in DMEM (Sigma) containing 10% FBS (Hyclone, Logan, UT) and 1% penicillin streptomycin for 1–2 days until they reached 90–100% confluence. Before treatment, cells were serum starved for 12–18 h with serum free medium and then treated with different concentrations of MBG or ouabain ranging from 0.1 to 100 nM for up to 96 h. In control cells, the vehicle DMSO (\(\leq 0.01\%\)) was added. In some of the experiments, cells were also treated with 5 to 10 ng/ml of TGF-β1, (for up to 96 h). Fresh MBG and TGF-β1 (Sigma) were added daily for the entire duration of treatment.

**Preparation of whole tissue/cell lysates.** Frozen (−80°C) tissues were grinded in liquid nitrogen and placed immediately into ice-cold RIPA containing 50 mM Tris·HCl pH 7.5, 150 mM NaCl, 1% Nonidet P-40 substitute, 5% sodium deoxycholate, 0.1% SDS, protease inhibitors (0.2 mM 2-aminooethylbenzenesulfonyl fluoride, 8 μM bestatin, 2.8 μM E-64, 3 μM pepstatin, 4 μM leupeptin, and 0.16 μM aprotinin; Protease Inhibitor Cocktail, Sigma), and 1 mM EDTA. Cells were washed twice with ice-cold PBS and proteins were imme-
diately extracted in RIPA buffer. Protein concentration was estimated by Bio-Rad protein assay (Hercules, CA).

**SDS-PAGE, Western blotting, and autoradiography.** SDS-PAGE was carried out using standard protocols. Proteins were transferred onto PVDF membrane by semidry method (38) using trans-Blot SD semidry transfer cell (Bio-Rad). Membranes were blocked in 5% nonfat dry milk in Tris-buffered saline, containing 0.1% Tween-20. The following antibodies were used at concentrations recommended by the respective manufacturer: mouse anti-E-cadherin from BD Biosciences, clone 36; rabbit anti-occludin and anti-claudin I from Zymed Laboratories (South San Francisco, CA), rabbit anti-β-catenin from Sigma, rabbit anti-Snail from Abcam (Cambridge, MA), mouse anti-vimentin from Serotec (Oxford, UK), rabbit anti-fibronectin from Chemicon (Temecula, CA), and goat anti-actin clone I-19 from Santa Cruz Biotechnology (Santa Cruz, CA). For all Western blots, secondary antibodies from Santa Cruz Biotechnology were used.

**Cell invasion assay.** To assess the invasive potential of the trans-
formed cells, LLC-PK1 cells were grown on collagen gels (2.5 mg/ml, 2.5-mm thick). Gels were made from rat tail collagen Type I from BD Biosciences (Bedford, MA) in sixwell plates according to company protocol. After the gel was polymerized, the surface was washed once with sterile PBS and once with 10% DMEM medium and then the cell suspension was added. Cells were grown up to 90–100% confluence before they were serum starved and treated with 100 nM MBG for 72 and 96 h. After treatment, the collagen gels were separated from the wells and the cells on the collagen gels were washed twice with cold PBS and fixed in 4% formaldehyde for 30 min at room temperature. Gels were incubated with 0.1% Coomassie (R250) in 10% methanol and 10% acetic acid until sufficient staining was achieved. The gels were then destained with 50% methanol and 10% acetic acid with frequent washings. The gels were later mounted on glass slides using gelatin mount from Sigma. Differential interference contrast images of cells on the gel surface as well as the invading collagen were taken on IX71 inverted research microscope (Olympus) focused on different planes.

**Immunocytochemistry.** Cells were grown on eighteen well Lab-Tek II chamber slides under the conditions described above. After a specified time of treatment, cells were washed twice in PBS pH 7.4 and fixed in ice cold methanol for 10 min at 4°C. Cell walls were permeabilized with 4 mM sodium deoxycholate in PBS for 10 min at room temper-
ature and then washed in PBS containing 0.025% Triton for 5 min. Non-specific binding of the antibodies was blocked by incubation in PBS containing 1.5% normal horse serum from Vector Laboratories for 30–60 min depending on the antibody used. Cells were then incubated at room temperature for 1 h with primary antibodies. For immunocytochemistry, the same antibodies as for Western blotting were used except for E-cadherin and β-catenin staining. Prediluted mouse anti-E-cadherin and anti-β-catenin antibodies from Cell Marque (Rocklin, CA) were used as a manufacturer’s suggestion. Immunoperoxidase staining of E-cadherin was done using an ABC Elite kit from Vector Laboratories. For fluorescent immunostaining, fluorescein-conjugated goat anti-mouse IgG and Oregon Green 488 goat anti-rabbit IgG from Invitrogen (Eugene, AR) diluted 1:200 were applied for 30 min in the dark.

Confocal images were captured by a spectral confocal scanner (model TCS SP2; Leica, Mannheim, Germany) and microscope.
MG induces renal fibrosis and EMT

Analysis of collagen I synthesis. Analysis of collagen I synthesis was performed as described by Strutz et al. (76) with some modifications. Cells were grown to total confluence in 12-well plates in the presence of 50 μg/ml ascorbic acid and 50 μg/ml propionitrile (both from Sigma) and treated with 100 nM MBG for 72 and 96 h as described in Cell culture. Supernatants (100 μl) were collected, transferred to 96 well plates (Nunc), and incubated overnight at room temperature. Plates were dried the following day for 2 h and blocked in incubation buffer, containing 1.5% normal horse serum (Vector Laboratories) in PBS at 37°C for 10 min. Rabbit polyclonal antibody against collagen I was added at dilution 1:2,000 (Abcam, ab292) in incubation buffer and probed for 2 h room temperature. The plates were then washed five times with a washing solution containing PBS and 0.02% Triton X-100 followed by incubation with secondary antibodies (biotinylated goat-anti-rabbit; Bio-Rad) diluted 1:8,000 in incubation buffer for 1 h. The plates were washed five times, 5 min each, with washing buffer and left overnight for binding of streptavidin-horse-radish peroxidase conjugate (BD Bioscience) diluted 1:4,000 in incubation buffer in a cold room on a shaking apparatus. Finally, the plates were washed three times in washing buffer and incubated with o-phenylenediamine. When sufficient color was developed, the absorbance was measured at 490 nm at Spectra Max 250 plate reader (Molecular Devices). Assays were run in triplicates and repeated three times. Standardization was obtained using rat tail collagen, type I (BD Bioscience). Working range for the detection was from 100 ng to 2,500 ng/ml.

Preparation of nuclear fractions. The preparations of nuclear fractions were done as described previously (43). LLC-PK1 cells were incubated for 15 min on ice in hypotonic buffer containing 10 mM Tris·HCl, pH 7.2, and protease inhibitors cocktail (Sigma) diluted as described for RIPA buffer. After homogenization, the cell extracts were diluted with equal volume of 0.5 M sucrose in hypotonic buffer and centrifuged at 800 g for 10 min at 4°C. The pellet was resuspended, homogenized, and centrifuged at 430,000 g for 2 h to pass through a 2-M sucrose cushion. The nuclear pellets were resuspended in RIPA buffer and applied for Western blot analysis.

3-(4,5-dimethylthiazol-2-yl)-2,5-Diphenyltetrazolium bromide and lactate dehydrogenase viability assays. LLC-PK1 cells were grown to confluence in 96-well plates in DMEM containing 10% FBS as described above. Cells were serum starved overnight before treatment with MBG or ouabain. The viability of the cells was evaluated every 24 h, as suggested by protocols for 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT)-based and lactate dehydrogenase (LDH)-based toxicity assay kits (Sigma).

Measurement of Na⁺-K⁺-ATPase activity. Na⁺-K⁺-ATPase activity was assessed by 86Rb⁺ uptake assay as described previously (42, 43). LLC-PK1 cells were grown to confluence and treated with MBG or ouabain as described in Cell culture. Monensin (20 μM) was added to the medium before initiation of the 86Rb⁺ uptake to assure that maximal capacity of active uptake was measured. The uptake was initiated by addition of radioactive 86Rb⁺ and stopped after 10 min by addition of 0.1 M ice-cold MgCl₂. Intracellular 86Rb⁺ was precipitated with TCA and radioactive signal counted. The 86Rb⁺ uptake was calibrated with TCA-precipitated protein content for each treatment.

Fig. 1. Marinobufagenin (MBG) infusion to rats induced renal fibrosis. A–H: microphotograph images taken from control rats (left) and MBG-administered rats (right). MBG administration induced significant accumulation of tubulointerstitial collagen proximal to interlobar and arcuate blood vessels (A and B). In the cortex, MBG administration resulted in development of peritubular and periglomerular fibrosis (C–F), as demonstrated by picosirius red staining imaged in bright (C and D) and polarized light (E and F). Development of periglomerular and peritubular fibrosis after MBG administration was confirmed by appearance of α-smooth muscle actin (α-SMA)-positive cells around renal corpuscle and in the tubulointerstitium (G and H). I: representative Western blot for proteins extracted from whole kidneys and probed against procollagen I antibodies. J: densitometry analysis of Western blots (n = 3 from more than 3 independent experiments; *p < 0.05 vs. control). K: results of computer-assisted morphometry measurements of collagen content in kidney interstitium (n = 20 from 3 independent experiments; *p < 0.05 vs. control).
Statistical analysis. Data are presented as means ± SE. Unpaired Student’s t-test was used to evaluate the difference between groups. Statistical significance was reported at the $P < 0.05$ and $P < 0.001$ levels.

RESULTS

MBG induces alterations of physiological parameters and renal fibrosis in rats. MBG infusion to rats for 4 wk resulted in an increase of plasma levels of MBG from 359 ± 16 to 546 ± 36 pmol/l, aldosterone from 191 ± 55 to 322 ± 38 pg/ml, and systolic blood pressure from 102 ± 2 to 136 ± 4 mmHg. Kidney sections of MBG-supplemented and control rats were stained with collagen specific picrosirius red. Fibrotic lesions with significant accumulation of collagen I around tubules were found in the area proximal to the interlobar and the arcuate vessels (Fig. 1, A and B). Furthermore, MBG infusion induced periglomerular and peritubular accumulation of collagen I (although to a lesser extend) in renal cortex, as demonstrated in the photomicrographs of picrosirius red stained kidney sections taken in bright and polarized light (Fig. 1, C–F).

Accumulation of interstitial collagen I fibers in the kidney cortex estimated by imaging of picrosirius red stained sections revealed a twofold increase with MBG infusion (Fig. 1K). The development of glomerular and peritubular fibrosis in MBG-treated rats was further supported by the presence of the α-SMA-positive cells around the renal corpuscles and in the tubular interstitium (Fig. 1, G and H). Western blotting of whole kidney extracts showed that procollagen I expression was increased four times in rats supplemented with MBG (Fig. 1, I and J). Therefore, 4 wk of MBG infusion induced activation of matrix-producing perivascular and periglomerular mesenchymal cells and consequent secretion of collagen I fibers in the renal cortex, the corticomedullary junction, and the columns of Bertini. This effect of MBG could be due to renin-angiotensin-aldosterone system induced TGF-β1 upregulation. Indeed Western blotting of whole kidney extracts showed upregulation of TGF-β1 after MBG treatment (data not shown).

In contrast to mainly perivascular and periglomerular activation of mesenchymal cells, the profibrotic transcription fac-
tor Snail, a key regulator of EMT, was found de novo expressed in both cortical and medullary tubular epithelial cells in MBG-treated kidneys (Fig. 2). Moreover, in some epithelial cells the Snail protein was specifically located inside the nuclei. Western blotting of proteins extracted from whole kidneys showed a threefold increase of Snail protein expression after MBG infusion. The basal level of Snail present in the extracts from control kidneys reflects the expression of Snail protein in the muscularis layer of the renal calyx (not shown). Surprisingly, Western blotting of proteins from whole kidney tissues as well as immunohistochemical analysis of kidney sections did not reveal any changes in the expression and cellular localization of the epithelial markers E-cadherin and β-catenin with MBG administration (not shown).

Effect of MBG on LLC-PK1 cells morphology. The proximal tubular epithelial cell line LLC-PK1 was chosen for our in vitro studies because these cells have been intensively used in our laboratory for investigation of the mechanism of ouabain signaling through the Na+/K+-ATPase signalosome. We treated subconfluent (≈50%) and confluent cultures of LLC-PK1 cells with MBG or ouabain in concentrations of 0.1, 1.0, 10.0, 50, and 100 nM, and TGF-β1 (5–10 ng/ml) as a positive control for EMT. The morphological changes of the cells were monitored at 24, 48, 72, and 96 h. Control LLC-PK1 cells formed a strong epithelial sheet characterized by cobblestone-like morphology typical for epithelial cells (Fig. 3). Clear changes in cellular morphology to fibroblast-like cells were observed only in confluent cultures treated with MBG for 72 and 96 h at 100 nM and to some extent at 50 nM (not shown). Ouabain, another CTS, did not induce alterations in the LLC-PK1 cells that were noted with MBG in all our experiments. Fibroblast-like cells were absent in confluent culture of LLC-PK1 cells treated with TGF-β1 and in subconfluent cultures treated either with ouabain or MBG. It has been shown earlier by Masszi et al. (51) that the confluent cultures of LLC-PK1 cells are resistant to TGF-β1-induced EMT and that only subconfluent cultures showed EMT-like alterations in the presence of TGF-β1. The morphological features of the transformation of LLC-PK1 cells induced by 100 nM MBG for 96 h to cells resembling the mesenchymal phenotype occurred to a greater extent while the cells were at confluence rather than other experimental settings. Therefore, all further experiments were performed only at conditions in which the LLC-PK1 cells had achieved confluent monolayers.

The viability of the LLC-PK1 cells was assessed after treatment with cardiotonic steroids at 24, 48, 72, and 96 h by MTT- and LDH-based toxicity assays. As determined by these two assays, there were no lethal effects on the LLC-PK1 cells by 100 nM MBG during the examined time points. The results of viability by the MTT assay for LLC-PK1 cells treated for 24 h are shown in Table 1. MBG even at a concentration of 250 nM did not have any toxic effect on LLC-PK1 cells at 24 h. At the same time, ouabain in concentrations 50 and 100 nM, at which MBG triggers EMT-like morphological alterations of LLC-PK1 cell, induced cell death by 15 and 25%, respectively, even after 24 h of treatment. EMT is generally characterized by acquisition of morphological features specific for fibroblasts and increased cells migration and invasion. Epithelial cells in confluent monolayer are not motile at all. To test whether MBG induces invasive motility, we seeded cells on collagen I gel and analyzed invasion of the cells into the collagen matrix (Fig. 4, A and B). We found that after 72 h of MBG treatment while many cells still are incorporated in patches of epithelial sheet, ~30% of LLC-PK1 cells are scattered. After 96 h >20% are scattered on the surface of the gel and ~40% of the total number of cells

Table 1. Effect of MBG and ouabain on viability of LLC-PK1 cells and pumping activity of Na+/K+-ATPase

<table>
<thead>
<tr>
<th>Condition</th>
<th>MTT Staining, %</th>
<th>Maximal Na+/K+ Pump Activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>100.0±3.8</td>
<td>100.0±4.8</td>
</tr>
<tr>
<td>100 nM MBG</td>
<td>106.7±2.3</td>
<td>91.8±7.8</td>
</tr>
<tr>
<td>250 nM MBG</td>
<td>105.5±3.7</td>
<td>76.3±5.6*</td>
</tr>
<tr>
<td>10 nM ouabain</td>
<td>106.0±3.1</td>
<td>NA</td>
</tr>
<tr>
<td>50 nM ouabain</td>
<td>86.6±3.9*</td>
<td>37.6±4.9†</td>
</tr>
<tr>
<td>100 nM ouabain</td>
<td>74.8±1.4†</td>
<td>30.0±0.6†</td>
</tr>
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Viability of LLC-PK1 cells in the presence of ouabain and marinobufagenin (MBG) was estimated by 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) staining, and pumping activity of Na+/K+ -ATPase was determined by 86Rb+ uptake after 24 h treatment with cardiotonic steroids (CTSs). MTT staining and the pumping activity at the absence of CTSs were taken as 100%. Means ± SE from more than 3 experiments performed in sextuplicates are shown. *P < 0.05, †P < 0.001 vs. control.
indeed invaded the collagen matrix (Fig. 4C). There were no cells found in the collagen matrices in control cultures or cultures treated with MBG for 72 h.

**Effect of MBG on the expression of epithelial proteins.** It has been shown previously that LLC-PK1 cells are resistant to TGF-β1-induced EMT in intact confluent monolayer and are susceptible to EMT if formation of cell-cell contacts is prevented by different experimental conditions including low density (51). In this so-called contact-dependent model of TGF-β1-induced EMT of LLC-PK1 cells, treatment with TGF-β1 triggered dramatic downregulation of E-cadherin, a major marker of epithelial phenotype after 72 h posttreatment. In our initial experiments when confluent and subconfluent cultures LLC-PK1 cells were treated with TGF-β1, these findings were confirmed. When we treated subconfluent (50–60% of cell density) cultures of LLC-PK1 cells with MBG in concentration from 0.1 to 100 nM up to 96 h, we documented downregulation of E-cadherin protein level as well (data are not shown).

Immunocytochemical staining of the MBG-treated confluent monolayer of LLC-PK1 cells showed that fibroblast-like cells, which are completely detached, did not express E-cadherin on
their surfaces (Fig. 5). At the same time, there were cells that demonstrated mesenchymal features as spindle-like shape and filopodias; however, these cells remained attached to the neighboring cells by their cellular membranes. These cells demonstrated intensive staining for E-cadherin at the sites of contacts. Western blotting analysis of cell lysates from MBG-treated cultures revealed that the total level of E-cadherin protein remained unchanged (Fig. 6). High levels of total E-cadherin protein expression in LLC-PK1 total cell extracts, despite massive detachment and scattering of cells induced by MBG treatment, can be explained by the long half-life of the E-cadherin protein. A similar discrepancy has been demonstrated in other cell systems (64, 76). The half-life of E-cadherin is estimated to be >40 h (46). Overall, these data suggest that in an MBG-induced model of EMT in LLC-PK1 cells, unlike a majority of EMT models, downregulation of E-cadherin does not preclude or induce EMT.

We investigated the effects of MBG treatment on expression of the core tight junction proteins occludin and claudin I. At 72 h of treatment, cells were detached from each other, yet they still expressed considerable amount of claudin I and occludin (Fig. 5) around the cellular membranes. However, after 96 h of treatment, the transformed cells were characterized by more dispersed specific staining for tight junction proteins. Western blots run with extracts from LLC-PK1 cells showed that MBG decreased expression of claudin I by 30% at 96 h posttreatment (Fig. 6), while the level of occludin remained unchanged (Fig. 6).

Effect of MBG on the expression of mesenchymal proteins. To further investigate whether MBG indeed promotes EMT in LLC-PK1 cells, we analyzed the expression of the mesenchymal marker proteins fibronectin, vimentin, and collagen I.

Fig. 5. Indirect immunofluorescence staining for epithelial marker proteins E-cadherin (A and B), occludin (C and D), claudin I (E and F), and mesenchymal marker proteins vimentin (G and H), and fibronectin (I and J). Left: shows immunostaining of control LLC-PK1 cells grown for 96 h; right: LLC-PK1 cells grown in the presence of 100 nM of MBG for 96 h. Completely detached cells with fibroblast like morphology do not show E-cadherin specific staining, while transformed cells still attached to each other express E-cadherin at their contact sites. MBG treatment induced marked redistribution of tight junction proteins occludin I and claudin and increased staining for mesenchymal markers vimentin and fibronectin.

Fig. 6. MBG treatment did not induce any changes in the level of adherent junction protein E-cadherin, insignificantly reduced level of tight junction protein occludin, and decreased level of claudin I by 30%. Representative Western blot of proteins extracted from control and MBG-treated (100 nM) LLC-PK1 cells at 4 time points and probed with antibodies against E-cadherin, occludin, and claudin I, and densitometry analysis of Western blots (n = 15, from more than 3 independent experiments). *P < 0.05 vs. control.
MBG induced upregulation of vimentin expression with a maximal effect after 72 h of treatment (Fig. 5). MBG also stimulates the expression of fibronectin with a maximal effect at 72–96 h of treatment (Fig. 5). Immunoblotting of proteins extracted from LLC-PK1 cells at all time points showed that MBG induced upregulation of vimentin and fibronectin as early as 24 h of treatment (Fig. 7). We also assessed the effect of MBG on collagen I expression in LLC-PK1 cells. Collagen I secretion was increased twofold in cell culture supernatants treated with MBG (Fig. 8).

**MBG governs expression and subcellular localization of the transcription repressor Snail.** Several transcription factors govern EMT in tumor and benign epithelial cells including the zinc finger protein Snail (9, 13, 53, 63, 64). In untreated LLC-PK1 cells, the Snail protein was present in the cytoplasm of 39 ± 6% of the cells, homogeneously in the nuclei of 12% ± 6 of cells, and in 14 ± 3% of the cells in the nuclear periphery (Fig. 9). At 24 h posttreatment, the total level of Snail expression was increased and remained elevated during the experimental time span as demonstrated by Western blotting (Fig. 10B). At 24 h posttreatment, Snail was localized in the nuclei of 72 ± 4% of the cells and in the nuclear periphery in 14 ± 3% of the cells, and only 4 ± 2% of the cells displayed cytosolic Snail location. Although at 48 h, the total expression of Snail reached a maximal level, the protein presentation in nuclei was decreased to approximately one-half of that noted at 24 h (39 ± 6% homogenous and 20 ± 4% in periphery) and drastically increased in the cytoplasm (22 ± 4%). At 72 h of MBG treatment, a second wave of nuclear translocation of Snail protein was observed. At that time, 76 ± 5% of cells showed homogenous nuclear staining for Snail, and in only 6 ± 2% of the cells, Snail was present in the nuclear periphery. Redistribution of Snail protein made its next turn after 96 h of MBG treatment. At that time in the majority of cells (80 ± 3%), Snail was located in the nuclear periphery, only 8 ± 3% of the cells displayed homogenous nuclear staining, and 10 ± 2% of the cells showed cytosolic staining for Snail. Western blotting of proteins extracted from whole cells or from nuclear fractions confirmed that MBG not only upregulates expression of Snail protein but it also induces its relocation to nuclei (Fig. 10).

MBG-induced EMT of LLC-PK1 cells and signaling though Na+/K+-ATPase. We analyzed whether the MBG effects on the transformation of LLC-PK1 cells are due to inhibition of Na+/K+-ATPase (Table 1). At a concentration of 100 nM, MBG, as assessed by 86Rb⁺ uptake, does not inhibit pumping activity of Na+/K+-ATPase at 24 h posttreatment. Furthermore, inhibition was not recorded at 48, 72, or 96 h (not shown) of MBG treatment. At the same time, ouabain at a concentration of 50 nM inhibits more than half of the pumping activity. MBG treatment caused a slight but statistically significant decrease in the protein level of the α-subunit of Na+/K+-ATPase after 96 h of treatment with MBG (Fig. 11). This observation is an agreement with previous findings that ouabain and to a less extent MBG caused endocytosis of the α-subunit of Na+/K+-ATPase in LLC-PK1 cells (42, 43).

**DISCUSSION**

A number of recent studies (73, 74) revealed that endogenous CTSs control multiple physiological functions of mammalian organisms, e.g., blood pressure, sodium homeostasis, and cardiac activity. Presumably, all CTSs transmit their ex-
tracellular signal through the same molecule, the α-subunit of Na⁺-K⁺-ATPase (16, 21). MBG and ouabain belong to two different subclasses of CTSs: bufadienolides, containing a five-membered lactone ring, and cardienolides with six-membered lactone, respectively. The structure of lactone ring moiety of CTS has been shown to determine binding and inhibitory activities of CTSs (21).

We have shown that administration of MBG to rats induced kidney fibrosis. MBG likely promoted fibrosis by targeting different populations of renal cells. First, direct activation of interstitial fibroblasts could be accounted for roughly a twofold increase in collagen I content in the tubular interstitium. This level of collagen I increment by MBG, ouabain, and digoxin has recently been shown in several types of fibroblasts (19, 20). Second, MBG, presumably indirectly, through stimulation of the renin-angiotensin-aldosterone system induced TGF-β₁ secretion and consequent activation of perivascular mesenchymal cells and fibrotic scar formation. Recent investigations demonstrate that pericytes and bone marrow cells (41), as well as fibroblasts, emerged through EMT (89) contribute to the de-
velopment of the perivascular fibrosis in the kidney. Whether MBG alone could have any effect on these types of cells in triggering renal fibrosis needs to be studied. Third, de novo expression of the transcription factor Snail could initiate EMT in renal tubules thus promoting tubular degeneration and fibrotic scar formation.

The transcription factor Snail is an evolutionary conserved protein; its homologs have been associated with EMT across the animal kingdom (50). Activation of Snail in the adult kidney has been sufficient to induce fibrosis (8). In addition, the attenuation of renal fibrosis by the synthetic vitamin D paricalcitol and the chemokine receptor agonist BX471 has been correlated with downregulation of transcription factor Snail (36, 77). Renal fibrosis and upregulation of Snail in kidneys of the MBG-administered rats could be due to the activation of the renin-angiotensin-aldosterone system and consequent upregulation of the profibrotic TGF-β and its receptor system (10).

We have provided evidence that MBG alone can trigger EMT in renal epithelia. In particular, treatment of LLC-PK1 cells with 100 nM MBG upregulates the expression of the mesenchymal protein markers vimentin, fibronectin, and collagen I and induces cell scattering and invasive motility. A particular peculiarity of MBG-induced EMT was the finding that this steroid induces transformation of epithelial cells to the mesenchyme without significant downregulation of epithelial proteins. In a majority of the described EMT models, dramatic downregulation of E-cadherin and detachment of cells precede EMT (11, 30, 39, 53, 57, 78, 90, 92). When LLC-PK1 cells were treated with ouabain at concentration 10–100 nM, we did find the decrease in E-cadherin protein expression and cell detachment, but ouabain was unable to trigger mesenchymal gene programming. Cell detachment and β-catenin translocation to the nucleus in the presence of 1 μM of ouabain were shown recently in MDCK cells. The authors did not report whether MDCK cells in these experiments underwent EMT (12, 37). Another study showed that a concentration of 1 μM ouabain was toxic for MDCK cells (1).

EMT in LLC-PK1 cells seen in the presence of MBG is accompanied by upregulation of Snail protein. In a majority of described embryonic, carcinoma, and adult EMT, the expression of Snail protein is associated with 1) acquisition by epithelial cells a fibroblastoid, invasive phenotype; 2) downregulation of epithelial proteins, especially E-cadherin; and 3) upregulation of mesenchymal proteins. The set of all these changes is known as complete EMT (53, 63, 64). Nevertheless, in some developmental processes or during wound healing, migrating cells expressing high levels of Snail show only transient loss of polarity by redistribution of tight- or adherent-junction proteins (so called partial EMT; Refs. 27, 32). Because the MBG-treated LLC-PK1 cells had only mild downregulation of their tight/adherent junction proteins, we would rather classify this process as a partial EMT. In most epithelial cells, Snail is not detected in the absence of stimuli (9). Yet, there are reports in which E-cadherin and Snail are both detected in the same cell lines. In these cells, Snail function is controlled by its intracellular location through phosphorylation by cytosolic and nuclear protein kinase GSKβ3 (87, 94). Phosphorylated Snail protein is much less active as a repressor of E-cadherin and an inducer of the mesenchymal genes (15). We suggest that the partial EMT seen in LLC-PK1 could be a consequence of high basal levels of Snail protein. Since MBG treatment induced translocation of the transcription repressor Snail to the nuclei at 24 and 72 h, we presume that MBG could regulate Snail phosphorylation as well. After 96 h of MBG treatment, Snail localized in the periphery of the nuclei, suggesting its association with the inner nuclear membrane. Sequestering of transcription factors on the inner nuclear membrane restricts their access to target genes and temporarily limits their transactivation/transrepression abilities (31).

Currently, we cannot exclude that other transcription regulators, besides Snail, could be involved in activation of mesenchymal genes in the response to MBG treatment. However, our preliminary studies revealed that quenching reactive oxygen species, which was shown to upregulate the expression of Snail protein (70), completely inhibited MBG-induced EMT of LLC-PK1 cells. Since MBG treatment did not significantly decrease the amount of E-cadherin in LLC-PK1 cells, we, as expected, did not find any alterations in total β-catenin expression levels and its subcellular localization. Nevertheless, transcription factor TCF4/LEF-1, a master-gene of EMT, was de novo expressed in LLC-PK1 cells after 72–96 h of treatment.

Fig. 10. MBG upregulated and governed nuclear translocation of the transcription factor Snail. Western blots of whole cellular (A) and nuclear (B) extracts probed against anti-Snail antibody.

Fig. 11. Protein level of the α1-subunit of Na⁺-K⁺-ATPase was moderately but significantly decreased after treatment with MBG for 96 h, as shown by representative Western blot and densitometry analysis (n = 10). P < 0.05 vs. control.
with 100 nM MBG (not shown). In the absence of transcriptional coactivator β-catenin, TCF4/LEF-1 is characterized by a weak binding affinity to promoter regions (2). Since we did not detect β-catenin in nuclei, the transcriptional effectiveness of TCF4/LEF-1 in MBG-treated LLC-PK1 cells remains unclear.

Unlike most epithelial cells, LLC-PK1 cells are resistant to TGF-β1-induced EMT in intact confluent monolayers. These cells are susceptible to EMT induced by TGF-β1 only if cell-cell contacts are prevented by various experimental conditions including low cell density (51). In this contact-dependent model of EMT in LLC-PK1 cells, TGF-β1 has triggered dramatic downregulation of E-cadherin after 72 h of treatment. In our initial experiments, when confluent and subconfluent cultures of LLC-PK1 cells were treated with TGF-β1, these effects were confirmed. Moreover, when subconfluent (50–60% of cell density) cultures of LLC-PK1 cells were treated with MBG in concentrations from 0.1 to 100 nM for 96 h, we documented a concentration-dependent downregulation of E-cadherin protein levels. Yet, in these experiments, we did not find any fibroblast-like alteration of LLC-PK1 cells. This was a surprising observation, since Na+/K+-ATPase resides in the basolateral aspect of the established epithelial monolayer and it was reasonable to suggest that in subconfluent cultures Na+/K+-ATPase would be more accessible for the MBG ligand (58–60). We propose that a pool of Na+/K+-ATPase molecules, which specifically resides in the apical junctional complex in the polarized epithelia and connected through accessory proteins to the cytoskeleton, may play a crucial role in the MBG effect on transdifferentiation of LLC-PK1 cells (59).

The plasma concentration of MBG after 4 wk of administration to rats reached nearly 0.5 nM, while EMT in LLC-PK1 cells occurred at much higher concentrations (50–100 nM). These discrepancies may be attributed to the different sensitivities of MBG to the rodent and porcine Na+/K+-ATPase (22, 43). Concentrations of MBG used in our in vitro experiments were below levels at which changes of the pumping activity of Na+/K+-ATPase could be recorded by 86Rb uptake. On the other hand, ouabain concentrations of 50–100 nM inhibited Na+/K+-ATPase activity in the same cells substantially over time and eventually caused cell death. Therefore, the absence of significant EMT under these conditions for ouabain was not surprising. We have previously demonstrated that although equimolar amounts of ouabain and MBG have similar acute (30 min) effects on 86Rb uptake, equimolar amounts of MBG induces much smaller amounts of Na+/K+-ATPase endocytosis in LLC-PK1 cells than ouabain, and, over longer periods of time, MBG inhibits 86Rb uptake much less than ouabain (43). We would therefore speculate that long-term exposure to substantial amounts (e.g., 100 nM) of ouabain induces programmed cell death whereas smaller amounts of ouabain promote cell growth and proliferation (14, 84), processes shown to be incompatible with EMT (54). We also speculate that MBG, in addition to activation of proteins within the Na+/K+-ATPase signalosome complex, might also induce cell cycle arrest, a necessary prerequisite of the de-differentiation programs such as EMT (54, 79, 82). Notably, a similar difference between patterns of ouabain- and MBG-induced signaling have been previously reported in another renal epithelial cell line, MDCK, in which ouabain not MBG induced cell death (1).

In summary, we report for the first time that CTS, in the form of MBG, has the capacity in vivo and in vitro to induce the EMT process, which is directly involved in the development of organ fibrosis. Our findings further support the inflammatory discovery of the nonpumping, signaling function of Na+/K+-ATPase: furthermore, they broaden our understanding of endogenous CTS actions and provide important information for the prevention and treatment of fibrotic diseases.

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