Effect of Hydrocortisone on Angiotensinogen (AGT) Mutation–Causing Autosomal Recessive Renal Tubular Dysgenesis

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Abstract: We have identified a founder homozygous E3_E4 del: 2870 bp deletion + 9 bp insertion in AGT gene encoding angiotensinogen responsible for autosomal recessive renal tubular dysgenesis (ARRTD) with nearly-fatal outcome. High-dose hydrocortisone therapy successfully rescued one patient with an increased serum Angiotensinogen (AGT), Ang I, and Ang II levels. The pathogenesis of ARRTD caused by this AGT mutation and the potential therapeutic effect of hydrocortisone were examined by in vitro functional studies. The expression of this truncated AGT protein was relatively low with a dose-dependent manner. This truncated mutation diminished the interaction between mutant AGT and renin. The truncated AGT also altered the glucocorticoid receptor (GR)-dependent transactivation, indicating that AGT may affect the development of proximal convoluted tubule by alteration of glucocorticoid-dependent transactivation. In hepatocytes, hydrocortisone increased the AGT level by accentuating the stability of mutant AGT and increasing its binding with renin. Therefore, hydrocortisone may exert the therapeutic effect through the enhanced stability and interaction with renin of truncated AGT in patients carrying this AGT mutation.

Keywords: renal tubular dysgenesis; angiotensinogen; rescue therapy; founder effect
1. Introduction

Autosomal recessive renal tubular dysgenesis (ARRTD) featured by the absence or poor differentiation of proximal convoluted tubules on histology, maternal oligohydramnios, pulmonary hypoplasia, profound hypotension, and anuria after birth is a rare inherited disorder caused by the inactivating mutations of genes responsible for renin-angiotensin system (RAS) [1–6]. To date, more than 80 different mutations in genes encoding proteins of RAS have been identified in patients with ARRTD. The majority of mutated genes are ACE followed by REN, AGT and AGTR1 [6,7]. Almost all affected fetuses die either in uterus or after birth with refractory hypotension and/or respiratory failure [7–10]. Although the compromised renal perfusion caused by the defect in RAS has been proposed to be responsible for the development of renal tubular dysgenesis, the exact pathogenesis remains to be elucidated [11–13]. Until now, there was no definitive treatment for patients with ARRTD. It is crucial to develop specific and rescue therapy.

Recently, we have reported six patients with ARRTD caused by AGT (angiotensinogen) mutation from six unrelated families [14]. This homozygous E3_E4 del: 2870 bp deletion + 9 bp insertion in AGT resulted in the skipping of exons and the generation of truncated protein (1–292 amino acids) as well as diminished serum AGT, Ang I, and Ang II. The rapidly fatal course even with aggressive therapy including inotropic agents, plasma infusion, and peritoneal dialysis was notable in 5 of 6 patients. Based on the previous studies showing that the glucocorticoid acted as a transcription regulator of AGT, the administration of persistently higher dose of hydrocortisone achieved a better blood pressure response and rescued one patient without the need for dialysis. Since no ARRTD patients with the truncating variant have survived to date, the spontaneous recovery of hypotension and renal hypoperfusion is unlikely. To the best of our knowledge, this is the first successful rescue therapy of ARRTD. Nevertheless, the molecular mechanism of the potential rescue effect of hydrocortisone on ARRTD remains elusive. In this study, we explored the pathogenesis of this AGT mutation and elucidated the potential rescue role of hydrocortisone in vitro functional analysis.

2. Materials and Methods

2.1. Index Case

This study was approved by the ethics committee on human studies at Chang Gung Memorial Hospital in Taiwan (IRB 201902035A3). An index neonate with ARRTD caused by homozygous E3_E4 del: 2870 bp deletion + 9 bp insertion in AGT was successfully rescued by systemic hydrocortisone. This mutation led to the exclusion of exon 3 and 4, and generated the truncated AGT (1–292 amino acids). A potentially candidate binding motif (LQDLL) of nuclear receptor was excluded by this mutation (Figure 1) [14].

2.2. Construction of Plasmids

Various AGT coding regions were synthesized by polymerase chain reaction and subcloned into EcoRI and XhoI sites of vector pSG5.HA [15]. Three pSG.HA.AGTs, wild type, truncated 1–292, and 1–375 (R375Q), were created. The pSG.HA.AGT (1–375) was used for negative control due to the recurrent and truncating mutation. Plasmid quality has been checked by GE NanoVue spectrophotometer (GE Healthcare Systems, Chicago, IL, USA). A A260/A280 value of 1.80–1.90 was an indication for pure plasmid DNA. All plasmid DNAs were verified by sequencing analysis (Mission Biotech, Taipei, Taiwan).

2.3. Cell Culture, Transfection and Protein Stability Assay

HK-2 (human kidney-2) cells purchased from the Bioresource Collection and Research Center were grown in Dulbecco’s Mod. of Eagle’s Medium/Ham’s F-12 medium with 10% fetal bovine serum. L02 cells were grown in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum. HK-2 and L02 cells (Cell Bank of China Science, Shanghai, China) were transfected with indicated amount of pSG5.HA.AGT–wild type and pSG5.HA.AGT (1–292 and 1–375/R375Q) constructs. For protein stability analysis,
transfected cells were treated with 50 µg/mL cycloheximide (CHX) for 20, 40, 80, 180, and 360 min. The cells were treated with vehicle or respective hormones for 24 h and were harvested for Western blotting analysis.

2.4. Western Blotting

The cell lysates were prepared in lysis buffer (100 mM Tris-HCl (pH 8.0), 150 mM NaCl, 0.1% SDS, 1% Triton X-100) at 4 °C, separated by sodium dodecyl sulfate polyacrylamide gel electrophoresis, and electrotransferred onto polyvinylidene difluoride membranes (Millipore, Bedford, MA). The following proteins were detected using specific antibodies: Anti-HA (clone 3F10, Roche, 1:5000 dilution), Anti-α-actinin (H-2, sc-17829, Santa Cruz Biotechnology, 1:5000 dilution), Anti-AGT (11992-1-AP, Proteintech Group, 1:1000 dilution), Anti-GR (H-300, sc-8992, Santa Cruz Biotechnology, 1:1000 dilution), and Anti-p53 (DO-1, sc-126, Santa Cruz Biotechnology, 1:1000 dilution).

2.5. Proximity Ligation Assay (PLA)

Cells in situ PLA signal were measured by Duolink® In Situ Fluorescence Kit (Sigma-Aldrich, Saint Louis, MO, USA) according to the manufacturer’s instructions. In brief, HK-2 and L02 cells were cultured on coverslips in DMEM/F-12 with 10% FBS and DMEM medium with 10% FBS, respectively. These cells were transfected with pSG5.HA vector, HA-tag hAGT (full-length, a.a. 1–485), HA-tag hAGT (truncated, a.a. 1–292), and HA-tag hAGT mutant (R375Q) constructs. Primary antibodies, anti-Renin (rabbit polyclonal, HPA005131, Sigma), and anti-AGT (mouse monoclonal, 60126–1-Ig, Proteintech Group)
were mix and dilute at 1:100 in the Duolink® Antibody Diluent (1×). For obtaining the
digital images, the samples were analyzed by fluorescence microscope.

2.6. Luciferase Reporter Analysis of Glucocorticoid Receptor (GR)-Dependent Transactivation

The expression pKSX vectors for human GR and reporter gene for the MMTV-LUC
have been described previously [16]. HK-2 cells were plated in 24-well plates and trans-
fected using jetPEI (Polyplus Transfection Inc., New York, NY, USA) following the manu-
facturer’s protocol (Promega luciferase assay kit and DLR2 model). The total DNA was
adjusted to 1.0 µg using empty pSG5.HA vector. Cells were harvested for luciferase reporter
assays using a Promega Luciferase Assay Kit (Madison, WI, USA). Values are expressed
numerically as relative light units. Luciferase activity is presented as the mean ± SD of
two transfected wells and is representative of at least three independent experiments. The
results shown are representative of at least three independent experiments.

2.7. Statistical Analysis

Resulting mean data were compared with a Student two-tailed t test. The percent
relative standard deviation (SD), was calculated as 100 multiplied by SD divided by
the mean.

3. Results

3.1. In Vitro Expression of Truncated AGT

The expressions of wild-type AGT (amino acids (aa) 1–485) and two truncated AGT
(AA 1–292 and 1–375/R375Q) proteins were examined after transient transfection of AGT
in an immortalized proximal tubule epithelial HK-2 cell line. HK-2 cells were transiently trans-
fected with the indicated amount of indicated pSG5.HA.AGT plasmid DNAs and incu-
bated for 46 h. The cell lysates were subjected to Western blotting with antibodies against
angiotensinogen (AGT). Alpha-actinin (ACTN) was used as the loading control. The ex-
pressions of wild-type AGT and two truncated AGT proteins were in a dose-dependent
manner with a relatively low expression of truncated AGT (AA 1–292) (Figure 2A). To
further examine their relative stabilities, the HK-2 cells, a proximal tubular cell line derived
from normal kidney, were transiently transfected with 0.4 µg pSG5.HA.AGT (wild-type
and amino acids 1–375/R375Q) and 0.8 µg pSG5.HA.AGT (amino acids 1–292) and trans-
fected cells were treated with 50 µg/mL cycloheximide (CHX), a de novo protein synthesis
inhibitor, for 20, 40, 80, 180, and 360 min and analyzed in the Western blot analysis for
the quantitation of these AGT proteins with ImageJ software. Compared to wild-type
AGT, the truncated AGT (AA 1–292) protein was more stable at the longer CHX treatment
time, but the truncated AGT (AA 1–375/R375Q) protein was less stable at the shorter CHX
treatment time (Figure 2B). The relative stability was calculated and compared with 0 h
CHX treatment (Figure 2C).

3.2. Measurement of Renin-AGT Interaction in Kidney

Due to this deletion mutation of AGT resulting in skipping of exons, we further
examined whether this truncated AGT impaired its interaction with renin in kidney. HK-2
cells were transiently transfected with 0.5 µg pSG5.HA vector and indicated pSG5.HA.AGT
plasmid DNAs. Using the proximity ligation assay (PLA), we examined the possible
interaction defect between renin and wild-type and truncated AGT proteins. We analyzed
the samples with a fluorescence microscope and obtained digital images. As shown in
Figure 3, there was a significant decrease in number of PLA of two truncated AGT proteins
(AA 1–292 and 1–375/R375Q), but not wild-type protein, indicating that this truncated
AGT protein might diminish the AGT-Renin interaction.
Figure 2. The expression level and stability of various AGT proteins. (A) Human kidney-2 (HK-2) cells (5 × 10^4 cells/well) were transiently transfected with 0.4 μg pSG5.HA.AGT wild-type (AA 1–485) and pSG5.HA.AGT (AA 1–375/R375Q) and 0.8 μg pSG5.HA.AGT truncated (AA 1–292). (B, C) HK-2 cells (5 × 10^4 cells/well) were transiently transfected with indicated amount of hydrocortisone and dexamethasone for 24 h. The cell lysates were subjected to Western blotting with antibodies against AGT and GR. ACTN was used as the loading control. By increasing concentrations of hydrocortisone and dexamethasone, a decrease in glucocorticoid receptor (GR) with both drugs was observed (Figure 4A). The ubiquitin-proteasome pathway in regulating GR protein turnover was a classic mechanism to terminate glucocorticoid responses [17]. There was no apparent change of endogenous AGT proteins treated with hydrocortisone and dexamethasone in L02 cells. The Western blotting data showed that there was no apparent change of these three exogenous AGT proteins treated with
either dexamethasone (100 nM) or hydrocortisone (10 μM) for 24 h in L02 cells transiently transfected by 0.4 μg pSG5.HA.AGT wild-type (AA 1–485) and pSG5.HA.AGT (AA 1–375/R375Q) and 0.8 μg pSG5.HA.AGT truncated (AA 1–292) (Figure 4B). Lastly, we checked whether hydrocortisone could differentially affect the protein stability of wild-type AGT and truncated AGT using the CHX pulse-chase experiment. The truncated AGT (AA 1–292) was more stable than wild-type AGT in L02 cells. With a 46-h hydrocortisone treatment, our quantitative data demonstrated that hydrocortisone increased the amount of exogenous wild-type and truncated AGT (AA 1–292) proteins (Figure 4C,D compare the absence and presence hydrocortisone at 0 h CHX treatment time) and the stability of truncated AGT (AA 1–292) was better than wild-type AGT protein after the 90-min CHX treatment (Figure 4C,D). In Figure 4E, pretreated 2 h proteasome inhibitor MG132 demonstrated that exogenous HA.AGT (AA 1–485 and 1–292) or endogenous p53 could be rescued from the protein degradation pathway in the 120-min CHX treatment of L02 cells. This indicates that both wild-type and truncated AGT could be degraded by proteasome. These findings implicated that the increased serum AGT by hydrocortisone did not result from the direct increase of its production, but the stabilization of cytosolic truncated AGT.

Figure 3. Proximity ligation assay between AGT and renin in kidney. (A) Illustration of deleted region of truncated AGT protein involving serpin domain and cleavage sites of renin and angiotensin-converting enzyme (ACE); (B,C) HK-2 cells (5 × 10^4 cells/well) were transiently transfected with 0.4 μg pSG5.HA.AGT wild-type (AA 1–485) and pSG5.HA.AGT (AA 1–375/R375Q) and 0.8 μg pSG5.HA.AGT truncated (AA 1–292). Measurement of Renin-AGT interaction by proximity ligation assay. The results (C) are representative of three independent experiments.
Figure 4. The effect of hydrocortisone on mutant AGT proteins. (A) L02 cells were treated with indicted amount of hydrocortisone and dexamethasone for 24 h; (B) L02 cells were transiently transfected with 0.4 µg pSG5.HA.AGT wild-type (AA 1–485) and pSG5.HA.AGT (AA 1–375/R375Q) and 0.8 µg pSG5.HA.AGT truncated (AA 1–292) and transfected cells were treated with dexamethasone and hydrocortisone for 24 h; (C,D) L02 cells were transiently transfected with pSG5.HA.AGT (1–485 and 1–292) and transfected cells treated with 10 µM hydrocortisone for 46 h and 50 µg/mL CHX for 20, 40, 80, 180, and 360 min. The relative stabilities of wild-type and mutant AGT proteins were calculated and compared with 0 h CHX treatment. Both wild-type and truncated AGTs increased with a decreased glucocorticoid receptor (GR) after hydrocortisone treatment. (E) Effect of proteasome inhibitor on AGT degradation. L02 cells were transiently transfected with 0.4 µg pSG5.HA.AGT wild-type (AA 1–485) and 0.8 µg pSG5.HA.AGT truncated (AA 1–292) and transfected cells treated with 0, 1, 3, 5, and 10 µM MG132 for 2 h and then treated with 10 µg/mL CHX for 120 min. p53 is for a control MG132 treatment and alpha-actinin (ACTN) is for a loading control. The results (A–E) are representative of three independent experiments.
3.4. Effect of Hydrocortisone on the Interaction between Renin and Mutant AGT in Liver

Liver is the major organ for the AGT synthesis. The PLA study showed that hydrocortisone enhanced AGT-Renin interaction in both wild type and truncated AGT in a normal human hepatic L02 cell line (Figure 5A,B). This pinpointed that hydrocortisone could enhance the interaction between renin and AGT by increasing the stability of mutant AGT and may contribute the subsequent increase of Ang I and II.

Figure 5. The effect of hydrocortisone on the interaction between AGT proteins and renin in liver. (A) L02 cells (8 × 10^4 cells/well) were transiently transfected with 0.4 µg pSG5.HA.AGT wild-type (AA 1–485) and pSG5.HA.AGT (AA 1–375/R375Q) and 0.8 µg pSG5.HA.AGT truncated (AA 1–292) and transfected cells were treated with hydrocortisone for 24 h. (B) Measurement of Renin-AGT interaction by proximity ligation assay. The results are representative of three independent experiments.

3.5. Mutant AGT Impaired the Glucocorticoid Receptor (GR)-Dependent Transactivation

Previous studies showed that GR might play important roles in the development of proximal convoluted tubule [18,19]. In addition, the truncated AGT identified in our case excluded the potential binding motif of nuclear receptor. Here, we addressed whether full-length AGT could be involved in the GR-dependent transactivation and mutant AGT could alter this GR-dependent transactivation. HK-2 cells were transfected with 0.25 µg of the MMTV-LUC reporter plasmid and pKSX.GR (0.15 µg) and/or indicated amounts of various AGT expression vectors in the absence or presence of 100 nM Dexamethasone or 1 µM hydrocortisone for 46 h. We first observed that dexamethasone and hydrocortisone were able to activate GR-dependent MMTV-Luc reporter activities. Wild-type AGT enhanced GR-dependent MMTV-Luc reporter activities in below 180 ng for dexamethasone stimulation and 40 ng for hydrocortisone stimulation, whereas suppressed GR-dependent MMTV-Luc reporter activities over these critical amounts. Both truncated AGTs (1–292 and 1–375/R375Q) suppressed this GR-dependent MMTV-Luc reporter activity in all tested dosages (Figure 6A).
These results suggested that truncated AGT may affect the development of proximal convoluted tubule by alteration of GR-dependent transactivation (Figure 6B).

Figure 6. The functional status of various AGTs examined by the modulation of GR-dependent MMTV-LUC activity in HK-2 cells. (A) HK-2 cells were transfected with 0.25 µg of the MMTV-LUC reporter plasmid and pKSX.GR (0.15 µg) and/or indicated amounts of various AGT expression vectors in the absence or presence of 100 nM Dexamethasone or 1 µM hydrocortisone for 46 h. The presented data are the means of three experiments (mean ± S.D.; n = 3); (B) The mutant AGT exerts different glucocorticoid receptor-dependent transactivation than wild AGT.
4. Discussion

In this study, we performed the functional analysis of a homozygous deletion mutation of AGT responsible for ARRTD. In vitro studies, this AGT mutant had a lower expression level than wild type AGT. PLA study demonstrated the attenuated interaction of this truncated protein with renin. Hydrocortisone could not only enhance the stability of truncated AGT but also accentuate its interaction with renin (Figure 7).

Figure 7. Proposed Pathogenesis of Angiotensin II deficiency due to large deletion of AGT in Renal Tubular Dysgenesis. Normally, wild-type AGTs are converted to angiotensin II through enzymes of renin and ACE. Mutation in AGT gene yields a shortened and abnormal AGT with decreased cleavage by renin and subsequent low serum angiotensin I and II. Hydrocortisone increases the production of angiotensin II by enhancing the interaction between renin and truncated AGT.

Despite more than 150 patients with ARRTD caused by different mutations in genes encoding proteins of RAS [1,2,18–20], only few patients with ARRTD from unrelated families caused by AGT mutations have been reported. Seven different mutations including 5 missense, 1 nonsense, and 1 large deletion mutations on AGT were identified. All of the mutants were located in the serpine domain [1,8,14,21]. Of note, the serpine domain is critical for the AGT cleavage by renin [18,22,23]. The present case with defect in AGT resulting in the deletion of serpine domain had low serum Ang I and II levels. By PLA analysis, we demonstrated this truncated AGT identified in our patient impaired the interaction between renin and mutant AGT. Thus, this genetic defect of AGT led to diminished serum AGT, and the low Ang I and Ang II results from, at least partially, the impaired AGT cleavage by renin. The vitro study revealed low expression of mutant AGT in line with the findings of our recent report that low amount of truncated AGT protein in liver and kidneys of patients with same AGT mutation [13]. Consistent with our finding, low serum AGT has been reported in cases with other truncated mutation, R375Q [24–26]. This attenuated expression of truncated AGT protein could be caused by either reduced hepatic production or enhanced hepatic degradation of truncated AGT.

Organic hypoperfusion, especially renal hypoperfusion in fetus, resulting from dysregulation of RAS is the main pathogenesis of renal tubular dysgenesis [6,9,12]. No correlation between clinical severity and mutated genes pinpointed that the functional RAS was critical for renal tubular development and the low angiotensin production was the bona fide cause of organic hypoperfusion. We found that this AGT mutation led to truncated AGT that excluded the amino acids that contained the potential binding motif (LQDLL) for nuclear receptor [27]. Since GR, one of nuclear receptors, is abundant in proximal convoluted tubule during the tubular development [18,19,28]. In mouse and rat, the AGT is primarily detected in proximal tubules at early gestation [29–32]. Therefore, the AGT may play an important role in acting as a growth factor during proximal convoluted tubule.
line with this speculation, we found that this mutant AGT could alter the GR-dependent transactivation and might provide another pathogenesis of renal tubular dysgenesis caused by this AGT mutation.

Glucocorticoid has been shown to be the regulators of generation of AGT in liver by increasing the abundance mRNA of AGT in vivo and in vitro studies [15,33,34]. Our in vitro study showed that the effect of hydrocortisone and dexamethasone did not affect the production of truncated AGT. However, hydrocortisone was shown to enhance the stability of the truncated AGT with the CHX treatment. This enhanced stability of truncated AGT may allow it more interaction with renin. As demonstrated by PLA, the hydrocortisone not only actually increased the interaction of the truncated AGT but also wild-type AGT with renin. This interesting finding may suggest that the beneficial effect of hydrocortisone is not limited to this specific mutation. In addition, hydrocortisone has been demonstrated to exert an effect on the expression of GR, which is abundant in proximal convoluted tubule during the development [19,31,35]. To elucidate this potential benefit of hydrocortisone on GR expression on the development of proximal convoluted tubule, further animal study is warranted.

5. Conclusions

This large deletion of AGT identified in the index case led to the truncated AGT and decreased its cleavage by renin with subsequent low Ang I and Ang II generation. This truncated AGT also altered the GR-dependent transactivation. High dose of hydrocortisone, at least in part through increasing the stability of the truncated AGT and enhancing the cleavage of AGT by renin, may be a potential therapy for ARRTD caused by AGT mutation.

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Institutional Review Board Statement: The study was conducted according to the guidelines of the Declaration of Helsinki, and approved by the Institutional Review Board at Chang Gung Memorial Hospital in Taiwan (IRB 201902035A3).

Informed Consent Statement: Informed consent was obtained from all subjects involved in the study.

Data Availability Statement: The data supporting reported results can be found at https://livendmctsghedu-my.sharepoint.com/:x:/g/personal/jamesdin1124_office365_nedmctsgh_edu_tw/ETfx47Rl7NFDtvvYpzhsf6B65GFg_MaIzEylQb0Vki1Dg?e=wp3049, accessed on 31 March 2021.

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