Adaptation to sustained high plasma vasopressin in water and electrolyte homeostasis in the rat transgenic for the metallothionein–vasopressin fusion gene

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Abstract

Prolonged exposure of tissues to a receptor agonist often leads to adaptive changes that limit the subsequent responsiveness of the tissue to the same agonist. Recently, we have generated rats transgenic for the metallothionein I–human arginine vasopressin (AVP) fusion gene (Tg), which produced high plasma AVP with relatively preserved renal water excretion, suggesting that there might be adaptive mechanism(s) for maintaining water and electrolyte homeostasis against chronic AVP oversecretion from the earliest stage of life. In this study, to investigate whether down-regulation of AVP V2 receptor (V2R), which could possibly be caused by long-standing high plasma AVP, participates in this adaptive mechanism(s), non-peptidic V2R antagonist OPC31260 was administered to reverse the down-regulation, and water loading was performed after V2R antagonist treatment had been withdrawn. Additionally, to confirm the down-regulation, Northern blotting analysis for V2R mRNA was carried out. Tg rats showed slightly decreased urine volume and water intake with an equivalent plasma [Na+] level (Tg 140·4 ± 0·6 mEq/l; control 139·3 ± 0·6 mEq/l) under basal conditions. After water loading using a liquid diet containing zinc, which stimulates the promoter region in the transgene, the urine increase showed only limited suppression with a dramatically increased plasma AVP level and mild hyponatremia (135·8 ± 1·8 mEq/l) in Tg rats. When diet containing OPC31260 had been provided for 4 days until the day before the start of water loading, antidiuresis and hyponatremia (125·4 ± 1·4 mEq/l) were significantly potentiated. V2R mRNA expression in kidney was significantly less in Tg rats than in control rats under basal conditions, and this suppression was restored by OPC31260 treatment to levels comparable with those of control rats. These results suggest that long-standing high plasma AVP causes V2R down-regulation, and it may play an important role in the adaptive mechanism(s) for maintaining water and electrolyte homeostasis in chronically AVP-overexpressing rats.

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Introduction

The neurohypophyseal nonapeptide arginine vasopressin (AVP) is the most potent antidiuretic hormone that regulates water balance. It acts on the adenylate cyclase–coupled vasopressin V2 receptor (V2R) located in the renal collecting ducts (Brownstein 1983) and increases water reabsorption from primary urine through the aquaporin-2 (AQP2) water channel by stimulating its shunting from intracellular vesicles into the apical plasma membrane and its synthesis (Knepper 1997, Marples et al. 1999).

AVP deficiency results in polyuria and polydipsia, symptomatic of diabetes insipidus. There is a genetic animal model, the Brattleboro rat, which fails to produce sufficient AVP due to a single base pair deletion in its AVP gene (Schmle & Richter 1984). Since its discovery about 40 years ago (Vaitin et al. 1962), the Brattleboro rat has proved to be very useful for studying the physiological consequence of AVP deficiency (Woods & Johnston 1983, Stoehr et al. 1993, Walter et al. 1996). On the other hand, in some clinical circumstances, inappropriately high plasma AVP results in free water retention and hyponatremia (Barter & Schwartz 1967). After the first report from Murphy et al. (1987), several investigators have developed genetic models of AVP oversecretion using the transgenic technique. However, these animals did not demonstrate apparent water and electrolyte disorders (Habener et al. 1988, 1989, Miller et al. 1993, Waller et al. 1996). Recently, we have generated a rat transgenic for the fusion gene consisting of the heavy metal-inducible...
promoter region of the mouse metallothionein I gene and human AVP gene (Nagasaki et al. 2002). We have revealed that in the rat (1) plasma AVP immunoreactivity was remarkably elevated and increased further by administering zinc sulfate, (2) AVP protein was processed correctly and revealed almost equivalent bioactivity with authentic AVP, and (3) the plasma [Na⁺] concentration was not different from control rats under basal conditions, while it declined compared with control rats when a liquid diet containing zinc was provided. Interestingly, the change in water and electrolyte balance was relatively mild in spite of a markedly elevated plasma AVP level. Considering other previous studies of AVP-overexpressing animals, our data suggest there might be some adaptive mechanism(s) maintaining the plasma [Na⁺] levels in the case of chronically high plasma AVP.

There could be several pathways for this change. Phosphorylation or internalization of the receptor plays a prominent role in the acute, short-term (seconds to minutes) adaptive response, and down-regulation of receptor mRNA or protein is concerned with the long-term (hours to days) response (Haddock & Malbon 1993). Since our AVP-overexpressing transgenic (Tg) rat is persistently exposed to high plasma AVP, we postulate that a long-term adaptive response, including down-regulation of V2R in the kidney, contributes to the maintenance of water and electrolyte homeostasis.

To test this hypothesis, we performed the following five experiments: (1) basal values of water and electrolyte homeostasis were checked; (2) renal reactivity to exogenously administered AVP was tested; (3) the effect of water loading with further plasma AVP elevation on water and electrolyte balance was more extensively examined; (4) to reverse the down-regulation of V2R, orally effective non-peptidic selective V2R antagonist OPC31260 (Yamamura et al. 1992) was administered, and water loading was performed after withdrawal of the antagonist; and (5) to confirm the down-regulation, V2R mRNA expression was evaluated.

Materials and Methods

Animals

Male rats homozygous for the transgene and age-matched control (Sprague–Dawley), 9–13 weeks of age, were analyzed. Details of the preparation of the rat transgenic for the human AVP gene used in this study have been described (Nagasaki et al. 2002). Rats were maintained under controlled conditions (23 °C; lights on 0900–2100 h), and all procedures were performed in accordance with the institutional guidelines for animal care at Nagoya University School of Medicine, which conform with the NIH animal care guidelines.

Materials

Zinc liquid diet

Commercial, nutritionally balanced liquid diet (Isocal plus, Mead-Johnson, Evansville, IN, USA) was diluted with an equal volume of 12.5 mM zinc sulfate in order to stimulate the metal-responsive element of the metallothionein promoter in the transgene. The formula of this liquid diet supplies 0.75 kcal/ml consisting of 45% carbohydrate, 15% protein, and 40% fat, along with 7.3 mEq/l sodium. The zinc liquid diet was provided at a volume of 30% body weight (BW) daily.

Diet containing V2 antagonist

Orally effective, non-peptidic selective V2R antagonist OPC31260, which was a gift from Otsuka Pharmaceutical Co. Ltd (Tokyo, Japan), was mixed with standard rat chow at a concentration of 0.05%.

Basal measurements

For the measurement of urine excretion, age-matched Tg and control rats were housed in metabolic cages individually and provided with standard rat chow and water ad libitum. After 2 days of habituation to the metabolic cage, basal 24-h urine volume, urine osmolality and 24-h water intake were measured. All rats were decapitated for the measurement of plasma [Na⁺] and AVP. The left kidney was removed for immunoblotting using anti-AQP2 antibody. There were five Tg and five control rats.

Renal reactivity to exogenously administered AVP

Renal reactivity to AVP was tested essentially as described in a previous report (Musabayane et al. 1985). In brief, rats were anesthetized with 2.1 M ethanol in distilled water (5 ml/100 g BW) administered by stomach tube and then placed on a continuous jugular infusion of 0.075 M NaCl and 0.44 M ethanol at 150 µl/min. The urinary bladder was cannulated via an incision in the abdominal wall and urine was collected into a preweighed tube every 3 min. After an equilibration period, 200 pg AVP (Peptide Institute, Osaka, Japan) dissolved in 200 µl 0.15 M NaCl was injected i.v. Urine volume was determined gravimetrically. There were three Tg and four control rats.

Effect of water loading

After 2 days of habituation to the metabolic cage, basal 24-h urine excretion and 24-h water intake were measured, and then the zinc liquid diet was provided once daily at a volume of 30% BW without the Chow to induce excess water intake. During the experiment, tap water was removed. Daily total urine volume, osmolality and [Na⁺] were measured. Rats were killed 12 h after the last zinc liquid diet had been provided at each of five different time-points (1 day prior to water loading (day 0), and 1, 2, 4, and 7 days after the start of water loading (days 1, 2, 4,
and 7 respectively) for measuring plasma [Na⁺] and AVP. There were six Tg and six control rats except for day 0 and day 1 (12 Tg and 12 control rats each). Plasma total protein (TP) was measured on day 0 and day 1 (n=6 each).

**Effect of water loading after V2 antagonist pretreatment**

As shown in Fig. 4A, to reverse desensitization of AVP action caused by long-standing high plasma AVP, diet containing V2 antagonist was provided ad libitum for 4 days (from day -4 to day -1) and withdrawn. During the V2 antagonist pretreatment, rats were allowed free access to water. Half of the V2 antagonist-pretreated group were decapitated to measure plasma [Na⁺] and AVP. Twelve hours after removing the diet containing V2R antagonist, the remaining rats were provided with the zinc liquid diet, and daily total urine excretion and urine osmolality were measured using the metabolic cage (day 1). Some other Tg and control rats were provided with the zinc liquid diet without V2 antagonist pretreatment. Twenty-four hours after the zinc liquid diet had been provided, all rats were decapitated to measure plasma [Na⁺] and AVP. There were six Tg and six control rats in each group.

**Immunoblotting**

The inner medulla was dissected from the left kidney which was sliced along the corticomedullary axis and
homogenized in 1 ml chilled isolation solution containing 250 mM sucrose, 10 mM triethanolamin, and 1 µg/ml leupeptin (Sigma, St Louis, MO, USA) and 0·1 mg/ml phenylmethylsulfonyl fluoride adjusted to pH 7·6. Protein concentration was measured with a Micro BCA Protein Assay Reagent Kit (Pierce, Rockford, IL, USA). All samples were diluted to 2 µg protein/µl with sample buffer containing 125 mM Tris, 2% SDS, and 10% glycerol adjusted to pH 6·8, then solubilized at 60 °C for 15 min. SDS-PAGE was carried out, and protein was electrophoretically transferred from gel to a nitrocellulose membrane (Hybond ECL; Amersham Pharmacia Biotech, Amersham, Bucks, UK). The membrane was blocked for an hour in 5% skimmed milk and was then probed for 48 h with affinity-purified rabbit polyclonal antibodies directed against AQP2 (kindly provided by Dr M A Knepper) diluted at 1:500 with a solution containing 137 mM NaCl, 4·0 mM KCl, 25 mM Tris adjusted to pH 7·4, 50 µl/dl Tween 20, and 0·1 g/dl bovine serum albumin. The second antibody was goat anti-rabbit IgG conjugated to horseradish peroxidase (New England Bio Labs Inc., Beverly, CA, USA) diluted at 1:1000. Antibody–antigen reaction was visualized by chemiluminescence reaction using ECL Western blotting detection reagents (Amersham Pharmacia Biotech). Relative quantitation of the band density from the immunoblots was performed by densitometry using NIH image-analysis software.

**Preparation of RNA and Northern blotting**

Age-matched Tg and control rats were fed a diet containing V2 antagonist for 4 days or fed continuously with standard rat chow. They were killed by decapitation, and the left kidney was rapidly removed, wrapped in aluminium foil, and frozen by burying it in dry ice. The kidneys, which had been frozen at −80 °C, were warmed very briefly to near 0 °C, and sliced along the cortico-medullary axis to dissect the inner medulla. Total RNA was extracted from the kidney inner medulla using TRIzol Reagent (Gibco-BRL, Life Technologies Inc., Rockville, MD, USA) according to the manufacturer’s recommendation. The amount of RNA was determined by the spectrophotometric method. Each 10 µg total RNA was size-fractionated and transferred to a nylon membrane (Gene Screen Plus Hybridization Membrane; Biotechnology Systems, NEN Research Products, Boston, MA, USA) for Northern blot hybridization using a 32P-labeled 602 bp cDNA probe designed specifically for rat V2R cDNA. The probe was generated by PCR (Terashima et al. 1999). To ensure even loading, the membrane was washed and then rehybridized with rat glyceraldehyde-3-phosphate dehydrogenase (GAPDH) cDNA probe (Fort et al. 1985). The integrated radioactivity of the mRNA band for V2R and GAPDH was quantified with a Bio-Image Analyzer (BAS2000; Fuji Photo Film Co. Ltd, Tokyo, Japan). The changes in V2R mRNA expression are presented as the percentage changes from non-treated control rats after correction for GAPDH mRNA expression. There were four Tg and four control rats in each group.

**Blood pressure measurement**

Systolic blood pressure (SBP) was measured by tail-cuff sphygmomanometer (soft ran BP-98A; Omron, Tokyo, Japan) in conscious Tg and control rats, which were fed with standard chow, zinc liquid or diet containing V2 antagonist. The SBP value was the mean of more than three successive measurements. There were six Tg and six control rats in each group.

**Plasma AVP and sodium measurements**

Plasma samples were collected in chilled tubes containing EDTA (potassium salt) for AVP assay. After immediate separation, plasma AVP was extracted through a Sep-Pak C18 Cartridge (Waters Associates Inc., Milford, MA, USA) and measured using a highly sensitive radioimmunoassay (RIA) kit (AVP-RIA kit, kindly provided by Mitsubishi Chemical Co. Ltd, Tokyo, Japan) (Oiso et al. 1999).
Figure 3 Effect of water loading. After measurement of basal daily urine excretion, a zinc liquid diet was provided at a volume of 30% BW/day without the chow so as to induce excess water intake. (A) Daily total urine volume was continuously measured. Rats were killed at each of five different time-points (1 day prior to water loading (day 0), and 1, 2, 4, and 7 days after the start of water loading (day 1, day 2, day 4, and day 7 respectively)) for measuring (B) plasma $[\text{Na}^+]$ and (C) AVP. There were six Tg and six control rats except for day 0 and day 1 (12 Tg and 12 control rats each). (D) Plasma TP was measured on day 0 and day 1 ($n=6$ each). *$P<0.05$ vs control rat, **$P<0.01$ vs control rat, ***$P<0.001$ vs control rat. $$$P<0.001$ vs basal condition of Tg rats.
et al. 1988). Plasma [Na$^+$] was measured using an auto-analyzer (Hitachi Ltd, Tokyo, Japan).

Statistics

Results are expressed as the means ± S.E.M. Statistical analyses were performed using one-way ANOVA followed by Fisher’s PLSD test for multiple comparisons. Differences were considered statistically significant at P<0.05.

Results

Basal measurements

Under basal conditions, daily water intake (Fig. 1A) and urine volume (Fig. 1B) of Tg rats had slightly decreased compared with control rats; however, urine osmolality (Fig. 1C) failed to demonstrate significant difference. Plasma [Na$^+$] showed no significant difference between Tg and control rats (Fig. 1D) under basal conditions despite plasma AVP being extremely elevated in Tg rats (Fig. 1E). AQ1P2 protein in kidney inner medulla did not show a significant difference between Tg and control rats (Fig. 1F).

Renal reactivity to exogenously administered AVP

In control rats, AVP rapidly revealed an antidiuretic effect, and urine excretion decreased to 40% of the basal value 9 min after the injection. In Tg rats, its antidiuretic effect was attenuated (70% of the basal value at 9 min after the injection) compared with control rats (Fig. 2).

Effect of water loading

Both Tg and control rats drank all the zinc liquid diet within the first few hours of administration. As a consequence of water loading, urine volume increased remarkably in both control and Tg rats (Fig. 3A). In Tg rats, although AVP secretion was markedly stimulated by zinc sulfate (Fig. 3C), urine volume was decreased only slightly compared with control rats (Fig. 3A). This suppression was observed until day 4 (Fig. 3A). Urine osmolality decreased remarkably after initiation of water loading in both control rats (from 2304 ± 141 (basal) to 580 ± 19 mOsm/kg (day 1), P<0.001) and Tg rats (from 2368 ± 85 (basal) to 585 ± 72 mOsm/kg (day 1), P<0.001). Neither urine osmolality nor daily [Na$^+$] excretion (urinary [Na$^+$] × urine volume) showed a significant difference between Tg and control rats (data not shown). However, cumulative [Na$^+$] excretion from the start of water loading became slightly higher in Tg than in control rats after day 3 (day 3, control rats 189 ± 0.3 mEq/kg BW; Tg 205 ± 0.4 mEq/kg BW, P<0.01; day 7, control rats 367 ± 0.6 mEq/kg BW; Tg 398 ± 1.2 mEq/kg BW, P<0.05). In response to water loading, plasma [Na$^+$] decreased gradually and plateaued at 125 mEq/l 4 days after initiation of water loading in Tg rats, whereas it did not demonstrate any significant change in control rats (Fig. 3B). Plasma TP did not show a significant difference between Tg and control rats under basal conditions. After providing zinc liquid diet, TP significantly decreased compared with the basal value, whereas plasma TP of control rats did not change (Fig. 3D).

Effect of water loading after V2 antagonist pretreatment

V2 antagonist caused a significant increase in urine volume and a decrease in urine osmolality in both Tg and control rats (Fig. 4B and D respectively). Their effects were attenuated in Tg compared with control rats. After 4 days of V2 antagonist pretreatment, plasma AVP increased markedly in control rats, but was unaffected in Tg rats (Table 1). In addition, V2R antagonist had no effect on plasma [Na$^+$], which was not significantly different between Tg and control rats either before or after pretreatment (Table 1). BW was not altered by the diet containing

Table 1 Effect of V2 antagonist on plasma [Na$^+$] and AVP. Values are means ± S.E.M.; n=6

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<th>Standard chow</th>
<th>Diet containing V2 antagonist</th>
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<tr>
<td>Plasma [Na$^+$] (mEq/l)</td>
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<tr>
<td>Control</td>
<td>139.3 ± 0.6</td>
<td>141.0 ± 1.2</td>
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<td>Tg</td>
<td>141.2 ± 0.9</td>
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<td>Plasma AVP (pg/ml)</td>
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<td>Control</td>
<td>2.3 ± 0.2</td>
<td>13.8 ± 0.1***</td>
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<tr>
<td>Tg</td>
<td>15.6 ± 2.9***</td>
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Rats were fed, for 4 days, a diet containing V2 antagonist or 0.05% OPC126, or a normal diet, then decapitated to measure plasma [Na$^+$] and AVP.

**P<0.01 vs control rats. ***P<0.001 vs standard chow.

Figure 4 Effect of water loading after V2 antagonist pretreatment. To reverse desensitization of AVP action caused by long-standing high plasma AVP, a diet containing V2 antagonist was provided ad libitum for 4 days (from day −4 to day −1), and then animals were switched to the zinc liquid diet (day 1) (A). Daily total urine volume and urine osmolality were measured on day −4 (B and D respectively) and day 1 (E and F respectively). Water intake was measured on day −4 (C). Some other Tg and control rats were provided with the zinc liquid diet without V2 antagonist pretreatment. Twenty-four hours after the zinc liquid diet was provided, the rats were decapitated for measuring plasma [Na$^+$] (G) and AVP (H). There were six Tg and six control rats in V2 antagonist-pretreated group.

$^{*}P<0.05$ vs control rats. $^{**}P<0.01$ vs control rats. $^{***}P<0.001$ vs control rats. $^{****}P<0.001$ vs control rats. $^{****}P<0.01$ vs V2 antagonist (−) Tg rats. $^{***}P<0.001$ vs control rats. $^{****}P<0.01$ vs V2 antagonist (−) Tg rats.

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V2 antagonist. In Tg rats, withdrawal of V2 antagonist pretreatment significantly potentiated suppression of urine excretion in response to water loading (Fig. 4E) and significantly elevated urine osmolality (Fig. 4F) compared with non-pretreated Tg rats. However, in control rats, there was no significant effect on urine volume (Fig. 4E) or urine osmolality (Fig. 4F) after water loading. It was also found that more potent hyponatremia was induced by water loading in Tg rats with V2 antagonist pretreatment, which had been stopped before starting the zinc liquid diet (Fig. 4G) despite there being no difference in plasma AVP levels between the V2 antagonist-pretreated group and non-pretreated group (Fig. 4H). BW increased by 6.0% 24 h after water loading in V2 antagonist-pretreated Tg rats, but did not significantly increase in non-pretreated Tg rats or in either pretreated or non-pretreated control rats (data not shown).

**V2R mRNA expression**

In Tg rats, V2R mRNA expression in kidney inner medulla was significantly less than in control rats under the basal conditions, and this suppression was restored by V2 antagonist treatment to levels comparable with those of controls rats. In control rats, V2 antagonist treatment had no significant effect on V2R mRNA expression (Fig. 5).

**Blood pressure measurement**

There was no significant difference between Tg and control rats in SBP under basal condition (control rats 117 ± 3 mmHg; Tg 115 ± 3 mmHg). Neither zinc liquid diet nor diet containing V2 antagonist exerted a significant effect on SBP (zinc liquid diet: control rats 120 ± 4 mmHg, Tg 116 ± 3 mmHg; diet containing V2 antagonist: control rats 114 ± 4 mmHg, Tg 116 ± 2 mmHg).

**Discussion**

In this study, basal physiological measurements with regard to water and electrolyte homeostasis, as well as plasma AVP and AQP2 protein levels in the kidney inner medulla were performed in Tg rats. While plasma AVP was markedly elevated, urine volume and water intake did not show dramatic changes, with an almost equivalent AQP2 protein level in the kidney. Renal reactivity to exogenously administered AVP using overhydrated, alcohol-anesthetized Tg rats clearly revealed desensitization of its antidiuretic action. Providing the zinc liquid diet, which induces potent water loading and further plasma AVP elevation by stimulating metal responsive elements in the transgene, demonstrated disruption of renal water excretion and developed hyponatremia in response to it in Tg rats. However, the alteration in water and electrolyte balance was relatively mild in spite of the extremely high plasma AVP levels. We also showed that V2R mRNA was down-regulated in Tg rats, and that OPC31260 pretreatment reversed the down-regulation and potentiated the antidiuretic action of AVP.

There have been several AVP-overexpressing transgenic animals before ours (Murphy et al. 1987, Habener et al. 1989, Grant et al. 1993). However, to our knowledge, no group has demonstrated hyponatremia or water retention caused by AVP-induced antiureasis. Habener et al. (1989) reported that the renal plasma membrane

![Figure 5](image_url)
fraction from their AVP-overexpressing transgenic mice showed diminished activation of adenylate cyclase following in vitro exposure to AVP, and they concluded that the animals had a nephrogenic diabetes insipidus-like state due to AVP receptor desensitization. We have generated AVP-overexpressing transgenic animals using the rat as a host animal. Rats enable more precise physiological analyses, such as urine volume or water intake, than mice and provide blood samples large enough to measure plasma AVP, which needs extraction before being applied to RIA. Our Tg rats showed slightly decreased urine volume with a normal plasma [Na+] level and AQP2 protein amount in the kidney under basal conditions. AVP regulates the water permeability of the renal collecting duct, not only by increasing expression of AQP2 (DiGiovanni et al. 1994, Marples et al. 1999) but also by transport of vesicles containing AQP2 to the apical plasma membrane (Knepper 1997). With regard to the discrepancy in the data for urine volume and AQP2 protein in this study, it is possible that the distribution of AQP2 protein in the collecting duct cell might be altered in Tg rats. Next, we investigated the effect of water loading with further plasma AVP elevation with the zinc liquid diet. As expected, plasma AVP concentration potently increased by stimulating the transgene. Plasma [Na+] decreased gradually and plateaued at day 4. Water loading-induced diuresis was slightly suppressed until day 4, and plasma TP decreased in Tg rats compared with the basal value, suggesting water retention. Although daily urinary [Na+] excretion never showed a significant difference, cumulative urinary [Na+] excretion slightly increased in Tg rats after day 3. Considering that hyponatremia was already established at day 3, the natreuresis may contribute to the maintenance of hyponatremia rather than its induction. This is consistent with a previous report of Verbalis (1994), which elucidated the mechanism of prolonged hyponatremia in his rat model of the syndrome of inappropriate antidiuretic hormone secretion. Overall, we have confirmed the physiological action of AVP in Tg rats. Considering the remarkably high plasma AVP in Tg rats, disruption of water and electrolyte homeostasis was very mild.

Verbalis & Drutarosky (1988) reported that continuous AVP infusion followed by extensive administration of water results in water retention and hyponatremia. However, daily water excretion began to increase in spite of sustained AVP infusion several days after initiation of water loading, namely the ‘vasopressin escape phenomenon’ (Verbalis & Drutarosky 1988, Murase et al. 1999, Tian et al. 2000). In our transgenic animal model, the escape from AVP-induced antidiuresis was already observed on the first day of water loading. Furthermore, the decrease in plasma [Na+] was much less even with more extensive water loading in rats with a plasma AVP level comparable with their study (Verbalis 1984). This evidence indicates that a potent adaptive mechanism for maintaining water and electrolyte homeostasis might be established in the case of chronic AVP oversecretion from the earliest stage of life in Tg rats.

The antidiuretic effect of vasopressin is mediated by V2R, which belongs to a large family of G protein-coupled receptors (GPCRs) with a typical seven transmembrane helix structure (Bichet 1994), located in the kidney collecting ducts (Ostrowski et al. 1992). Continuous or repeated exposure to an agonist promotes desensitization of GPCRs, and the attenuation of the hormonal response in spite of the presence of hormone, through short- and long-term regulation. Short-term regulation of transmembrane signaling appears to be largely independent of changes in gene expression such as phosphorylation or internalization of the receptor. Long-term regulation, in contrast, often reflects changes in the steady levels of mRNA or protein levels and/or changes in the turnover of receptor which causes down-regulation of receptor expression (Hadcock & Malbon 1993, Morris 1993).

Since we postulated that V2R down-regulation induced by long-standing high plasma AVP is responsible for the adaptive mechanism, we tried to reverse the down-regulation. β-Adrenoceptor, widely adopted as a prototype for the study of the regulation of GPCRs (Dohlman et al. 1991), increases in number when β-adrenoceptor blocking drugs are administered (Frishman 1987, van den Meiracker et al. 1989). Therefore, we used the non-peptidic antagonist OPC31260, which has been reported not to induce visible receptor internalization (Pfeiffer et al. 1998), so as to reverse the down-regulation for several days, after which water loading was performed. Interestingly, the diuretic effect of OPC31260 by blocking V2R was attenuated in Tg rats, in which V2R had already been partly blocked by the down-regulation. As expected, withdrawal of V2 antagonist pretreatment followed by water loading resulted in the potentiation of AVP-induced antidiuresis in Tg rats. Since blood pressure was not altered by the zinc liquid diet or V2 antagonist in both Tg and control rats, this effect is unlikely to be due to cardiovascular modification. Additionally, Northern blotting analysis for V2R in this study, which demonstrated suppressed V2R mRNA expression under basal conditions and its recovery after V2 antagonist treatment, was consistent with our hypothesis.

However, compared with a previous report from our laboratory (Terashima et al. 1998) regarding V2R mRNA down-regulation using dehydrated normal rats, the suppression of V2R mRNA in the basal state of Tg rats in the present study (75% of control rats) was less than the suppression in dehydrated Sprague–Dawley rats (65% of non-dehydrated), despite a higher plasma AVP level and a much longer duration in Tg rats than in the dehydrated normal rats in the previous study. Although the exact mechanism of agonist-induced desensitization is not yet fully understood, when considering the fact that AQP2 protein did not significantly increase under basal conditions in Tg rats, there may be a contribution from some
other mechanism(s), such as a change in the G protein or adenylate-cyclase regulation in the downstream signal transduction system of V2R prior to AQP2 expression (Laycock & Hanoune 1998). Our animal model will be useful to clarify the in vivo adaptive mechanism in the hormone excess state.

In conclusion, Tg rats showed attenuated AVP-induced antidiuresis and minimum disruption in water and electrolyte homeostasis. The down-regulation of V2R might, at least in part, be responsible for maintaining the homeostasis.

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