High Urine Volume and Low Urine Osmolality Are Risk Factors for Faster Progression of Renal Disease

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Background: Increased fluid intake slows renal disease progression in animal models. The relevance of these findings to human renal disease is not clear, although increased fluid intake often is recommended to patients with chronic renal insufficiency. This study tested the hypothesis that urine volume, urine osmolality (Uosm), or both are significantly associated with glomerular filtration rate (GFR) decline in patients with chronic renal insufficiency.

Methods: This is a retrospective analysis of Modification of Diet in Renal Disease (MDRD) study A patients with (N = 139) and without polycystic kidney disease (PKD; N = 442). The key outcome measure was GFR slope in relation to mean 24-hour urine volume and Uosm during follow-up in study A (mean, 2.3 years).

Results: The regression of GFR slope on mean follow-up 24-hour urine volume (adjusted for body surface area and MDRD diet and blood pressure group) showed that the greater the urine volume, the faster the GFR decline in patients both with and without PKD. For example, the difference in GFR slope for those with a mean follow-up 24-hour urine volume of 2.4 versus 1.4 L was −1.01 mL/min/y (confidence interval, −0.27 to −1.75) for patients without PKD and −1.20 mL/min/y (confidence interval, −0.06 to −2.34) for those with PKD. A similar but inverse relationship was shown between GFR decline and mean 24-hour Uosm in patients with (P = 0.01) and without PKD (P = 0.001). These associations remained significant after adjustment for 13 relevant baseline and follow-up covariates.

Conclusion: Sustained high urine volume and low Uosm are independent risk factors for faster GFR decline in patients with chronic renal insufficiency. Thus, high fluid intake does not appear to slow renal disease progression in humans. We suggest that until better evidence becomes available, patients with chronic renal insufficiency should generally let their thirst guide fluid intake. The advice to avoid “pushing fluids” might be particularly important for patients with PKD.

INDEX WORDS: Urine volume; renal disease progression; polycystic kidney disease (PKD).

Patients with chronic renal insufficiency commonly are advised to maintain a generous fluid intake. Two recent authoritative publications by nephrologists and for nephrologists recommend “increased” fluid intake in the management of chronic renal disease.1,2 The origins of this advice are both historic and contemporary. In the early years of renal physiology, it was shown that urinary urea clearance increased sharply as urine flow rates increased from 1 to 2 mL/min.3 Lower blood urea nitrogen (BUN) levels from chronic high fluid intake appear to be the basis for the historic recommendation to maintain urine volume at 3.0 L/d in chronic renal insufficiency.3

Contemporary objective evidence of benefit from a high fluid intake in chronic renal insufficiency is provided by studies of experimental ablative nephropathy in rats.4-7 Increasing fluid intake to partially decrease the normally high urine osmolality (Uosm) suppressed the maladaptive renal hypertrophy4-6 and renal interstitial fibrosis7 that are characteristic of that model.

Genetic models of polycystic kidney disease (PKD) in rodents also have been used to indirectly assess the effect of fluid intake on progression of renal disease. In Han:Sprague-Dawley rats, high water intake induced by increased sodium chloride intake increased renal cyst growth.8 Conversely, increased sodium bicarbonate intake, which also increased fluid intake, retarded renal cyst growth.9 However, in the latter experiments, renal wasting of sodium bicarbonate might have resulted in volume depletion that decreased cyst hydrostatic pressure and growth.

In humans, there has been no prospective study of the effect of fluid intake on renal disease
progression. However, the Modification of Diet in Renal Disease (MDRD) Study, with its extensive database, provided a unique opportunity to retrospectively determine the association between glomerular filtration rate (GFR) decline and daily urine volume. Patients with and without PKD were analyzed separately to assess whether an association of fluid intake and renal disease progression (GFR decline) might be different between patients with and without PKD.

METHODS

The patient population consisted of the study A cohort of the MDRD study (baseline GFR, 25 to 55 mL/min/1.73 m²). Study A was a 2 × 2 factorial design in which patients were randomly assigned to two different levels of blood pressure control; the usual goal (mean arterial pressure [MAP] of 102 to 107 mm Hg) or a low goal (MAP ≤ 92 mm Hg, if tolerated), and two different levels of dietary protein intake: the usual intake (1.3 g/kg of ideal body weight per day) or a low intake (0.6 g/kg of ideal body weight per day). Dietary instructions included avoidance of excess salt intake. Advice on fluid intake was left to the discretion of the physician. GFR decline was estimated from iothalamate clearances performed at 4-month intervals throughout the entire follow-up period. Blood pressure and other clinical testing were performed at 2-month intervals during follow-up. Patients with (N = 139) and without PKD (N = 444) were analyzed separately. Mean length of follow-up in MDRD study A was 2.3 years. MDRD study B patients (baseline GFR, 13 to 24 mL/min/1.73 m²) were not assessed because the small number of patients with PKD in study B limited the precision of the analyses.

Uosm was calculated as follows:

\[
\text{Uosm (mOsm/L)} = \frac{[2 \times \text{24-hour urine (sodium [mEq]}}}{2} + \text{potassium [mEq]} + 35.7
\]

\[
\times \text{24-hour urine urea nitrogen (UUN; g)/urine volume (L)}
\]

In the absence of urinary glucose, this method estimates Uosm within 10% of directly measured Uosm. Urinary glucose was not measured in the present studies; however, only 3.8% of MDRD study A patients had diabetes. All had type 2 diabetes.

Plasma osmolality (Posm) was calculated as follows:

\[
\text{Posm (mOsm/L)} = 2 \times \left(\text{serum sodium [mEq/L]} + \text{serum potassium [mEq/L]} + \frac{\text{BUN (mg/dL)}}{2.8} + \frac{\text{blood glucose (mg/dL)}}{18}\right)
\]

Baseline daily urine volume and Uosm were calculated as the average of monthly measurements of these factors obtained throughout the follow-up period.

To control for differences in body size, all analyses relating GFR decline to urine volume or Uosm were performed using regression models that included body surface area (BSA) as a covariate. Separate regression analyses were used to relate GFR decline to baseline values and mean follow-up values of daily urine volume and Uosm. Daily urine volume was log transformed in the regression analyses because of skewness.

To assess the influence of potential confounding factors in the association of GFR decline with daily urine volume and Uosm, each of these regression analyses was conducted after controlling for each of the following sets of covariates: (1) BSA and randomized treatment group; (2) factors in (1) plus baseline GFR and five other baseline factors shown to independently predict GFR decline in the MDRD study: race, MAP, 24-hour urine protein-creatinine ratio, serum transferrin level, and serum high-density lipoprotein (HDL) cholesterol level; and (3) factors in (1) and (2) plus mean follow-up levels of MAP, protein intake estimated from 24-hour UUN, serum sodium concentration, use of diuretics, use of angiotensin-converting enzyme (ACE) inhibitors, use of β-blockers, use of calcium channel blockers, and 24-hour urine protein-creatinine ratio.

Regression analyses of GFR decline were conducted using two-slope models in which each patient was assumed to have an initial rate of GFR decline in the first 4 months and a possibly different slope thereafter. The association of baseline and follow-up factors with GFR decline was evaluated as a time-weighted average of regression coefficients relating these factors to GFR decline in the first 4 months and in the subsequent follow-up period. A mixed-effects model was used for these regressions to account for correlations among multiple GFR measurements for the same patients. Similar analyses were used to relate GFR decline to the use of diuretics, ACE inhibitors, β-blockers, and calcium channel blockers; mean follow-up Posm levels; and Uosm times daily urine volume (osmolar excretion). For graphic presentation, mixed-effects models were used to obtain mean GFR slopes for patients within quartiles of daily urine volume and Uosm after controlling for baseline BSA and randomized treatment group.

Simple linear regression analyses were used to relate mean follow-up values of log daily urine volume and Uosm to mean follow-up MAP without covariate adjustment. Logistic regression was used to relate the percentage of patients administered diuretics, ACE inhibitors, β-blockers, or calcium channel blockers at more than 50% of follow-up visits to mean follow-up values of log daily urine volume and Uosm. All regression analyses were conducted separately for patients with and without PKD. All mean values are shown as ± 1 SEM. P of 0.05 or less is considered statistically significant.

RESULTS

Table 1 lists baseline characteristics of patients in study A with and without PKD. Patients with PKD comprised approximately 25% of the cohort. Patients with PKD are assumed to have autosomal dominant PKD.
Figures 1 and 2 show 24-hour urine volume and Uosm in patients with and without PKD assigned to the usual or low-protein diets during follow-up in the MDRD study, respectively. As shown, the low-protein diet resulted in slightly lower daily urine volumes in patients with and without PKD, probably because urine volume is affected by solute excretion in chronic renal failure.\textsuperscript{13} Note that mean follow-up 24-hour urine volume and Uosm were similar between patients with and without PKD assigned to the same dietary group. To account for differences in urine volume and Uosm associated with the diet group, analyses of GFR slope on urine volume and Uosm were adjusted for effects of the diet group.

Figure 3 shows the regression of GFR slope on mean 24-hour urine volume during follow-up in the MDRD study in patients with and without PKD. This analysis shows that the higher the mean 24-hour urine volume during follow-up, the greater the rate of GFR decline during follow-up. These associations are significant for patients with and without PKD.

Figure 4 shows the regression of GFR slope on mean 24-hour Uosm during follow-up in the MDRD study in patients with and without PKD. This analysis shows that, in general, the lower the Uosm, the more rapid the GFR decline. This relationship was expected because study A follow-up mean 24-hour urine volume and Uosm correlated significantly and inversely ($r = -0.57; P = 0.0001$).

To assess whether associations shown in Figs 3 and 4 could be explained by association of high daily urine volume and low Uosm with other risk factors for GFR decline, regressions shown in

\begin{table}
\centering
\caption{Baseline Demographics of the Study A MDRD Cohort}
\begin{tabular}{lll}
\hline
Renal Diagnosis & No. of Patients (\%)
\hline
PKD\textsuperscript{*} & 141 (24.1)
Glomerular diseases & 141 (24.1)
Hypertensive nephrosclerosis & 32 (5.5)
Tubulointerstitial diseases & 28 (4.8)
Urinary tract diseases & 22 (3.8)
Absence of 1 kidney & 19 (3.2)
Diabetic nephropathy & 17 (2.9)
Hereditary nephritis & 11 (1.9)
Unknown or other & 174 (29.7)
\hline
Sex & \\
& Men & 357 (61)
& Women & 228 (39)
Race & \\
& White & 495 (84.6)
& Black & 53 (9.1)
History of hypertension & \\
& Hypertensive & 499 (85.3)
& Nonhypertensive & 86 (14.7)
Age (y) & \\
& $<55$ & 325 (55.6)
& $\geq55$ & 260 (44.4)
\hline
\end{tabular}
\end{table}

\textsuperscript{*}Two patients with PKD were excluded from the present analysis because follow-up in the MDRD study was less than 1 year.
Figs 3 and 4 were adjusted for the baseline covariates GFR, BSA, race, urine protein-creatinine ratio, serum transferrin level, and serum HDL cholesterol level and the follow-up covariates of MAP, protein intake calculated from UUN, serum sodium level, and use of diuretics, ACE inhibitors, β-blockers, and calcium channel blockers. Results of these analyses are listed in Tables 2 and 3, in which associations of GFR decline with urine volume and Uosm are expressed as regression coefficients: GFR decline (milliliters per minute per year) per 1-L difference in mean follow-up urine volume (Table 2) or GFR decline per 100-mOsm/L difference in mean follow-up Uosm (Table 3).

As shown in the first data column of Table 2, at baseline the magnitude of the association of GFR decline with daily urine volume is relatively small and generally not significant. Conversely, during follow-up, the magnitude of the association of GFR decline with urine volume is relatively large and highly significant when adjusted only for BSA and diet group (large BSA and usual protein diet are associated with higher
urine volume). The stronger statistical association of GFR decline with follow-up daily urine volume compared with baseline daily urine volume may be explained by more precise ascertainment of follow-up values because they are the mean of 12 to 60 measurements/patient. Baseline values are the mean of only 2 measurements/patient.

As shown in the second and third data columns in Table 2, in patients without PKD, adjustment of the follow-up regression coefficient for baseline and follow-up covariates resulted in approximately a 50% reduction in the magnitude of association of GFR decline with follow-up urine volume, and statistical significance was lost for the association with urine volume with GFR decline. In patients with PKD, adjustment of follow-up regression coefficients for baseline and follow-up covariates had little effect on the magnitude of association of GFR decline with

Table 2. Regression of GFR Slope on Urine Volume: Effect of Adjustment for Baseline and Follow-up Covariates

<table>
<thead>
<tr>
<th></th>
<th>Adjusted for Diet, BP Group, and BSA</th>
<th>Adjusted for Diet, BP Group, and Baseline Covariates*</th>
<th>Adjusted for Diet, BP Group, and Baseline and Follow-Up Covariates†</th>
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<tbody>
<tr>
<td></td>
<td>Regression Coefficient‡ ± SE</td>
<td>Regression Coefficient‡ ± SE</td>
<td>Regression Coefficient‡ ± SE</td>
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<td></td>
<td></td>
<td>P</td>
<td>P</td>
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<tr>
<td>Baseline urine volume</td>
<td></td>
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</tr>
<tr>
<td>Non-PKD</td>
<td>$-0.27 \pm 0.34$</td>
<td>$-0.12 \pm 0.31$</td>
<td>$-0.13 \pm 0.31$</td>
</tr>
<tr>
<td>PKD</td>
<td>$-0.15 \pm 0.50$</td>
<td>$-0.09 \pm 0.50$</td>
<td>$-0.18 \pm 0.51$</td>
</tr>
<tr>
<td>Follow-up urine volume</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Non-PKD</td>
<td>$-1.01 \pm 0.38$</td>
<td>$-0.46 \pm 0.35$</td>
<td>$-0.52 \pm 0.35$</td>
</tr>
<tr>
<td>PKD</td>
<td>$-1.20 \pm 0.58$</td>
<td>$-1.15 \pm 0.58$</td>
<td>$-1.22 \pm 0.69$</td>
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</tbody>
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Abbreviation: BP, blood pressure.

*Baseline covariates: GFR, BSA, race, MAP, urine protein-creatinine ratio, serum transferrin level, and serum HDL cholesterol level.

†Follow-up covariates: MAP, protein intake calculated from UUN, serum sodium level, and use of diuretics, ACE inhibitors, $\beta$-blockers, and calcium channel blockers.

‡Regression conducted on log-transformed urine volume because of positive skewness. Regression coefficients expressed as milliliters per minute per year of GFR slope per 1 L of urine volume when urine volume is equal to follow-up median of 2.4 L.
follow-up urine volume, but the strength of the statistical association was diminished.

Table 3 regression coefficients are positive values because of the inverse relationship between urine volume and Uosm. As shown in the first data column of Table 3, at baseline, the magnitude of association of GFR decline with Uosm is relatively small and significant only for patients without PKD. However, during follow-up, the magnitude of the association of GFR decline with Uosm is relatively large and highly significant when adjusted for MDRD diet and blood pressure group and BSA. As shown in the second and third columns of Table 3, adjustment of follow-up regression coefficients for baseline and follow-up covariates resulted in approximately a 50% reduction in the magnitude of the association of GFR decline with follow-up Uosm. However, in patients with and without PKD, statistical significance was maintained.

The statistical association of GFR decline with Uosm was more robust than that of urine volume. This may be explained by greater precision of Uosm than urine volume determination. That is, 24-hour Uosm is calculated with the volume factor in both the numerator and denominator (see Methods) and therefore is affected less by urine collection errors than 24-hour urine volume measurement.

Proteinuria was the baseline and follow-up covariate that most influenced adjustments of regression coefficients (Tables 2 and 3). The effect of proteinuria may be explained because daily urine volume and urine protein-creatinine ratio correlated both at baseline ($P = 0.08$) and during follow-up ($P = 0.002$; data not shown), and proteinuria is a risk factor for faster GFR decline. The greater effect of the adjustment for proteinuria in patients without PKD may be explained by the greater baseline 24-hour proteinuria in patients without PKD (1.14 ± 0.05 g) compared with that of patients with PKD (0.29 ± 0.04 g). Although the magnitude of the association of GFR decline with follow-up urine volume and Uosm was greater in patients with than without PKD, these differences did not achieve statistical significance ($P = 0.29$ for differences in urine volume association; $P = 0.36$ for differences in Uosm association).

We examined whether high daily urine volumes were explained better by excess fluid intake or renal sodium and/or water wasting. This analysis showed that patients with high urine volumes tended to have lower serum sodium concentrations (patients with PKD, $P = 0.07$; patients without PKD, $P = 0.01$; data not shown). Furthermore, in those with the highest urine volumes, urine was frankly hypotonic to plasma (mean Uosm and Posm for those in the lowest quartile of Uosm were 212 ± 3 and 301 ± 1 mOsm/L, respectively; $P < 0.0001$). Thus, the

### Table 3. Regression of GFR Slope on Uosm: Effect of Adjustment for Baseline and Follow-Up Covariates

<table>
<thead>
<tr>
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<tr>
<td></td>
<td>Regression Coefficient‡ ± SE</td>
<td>$P$</td>
<td>Regression Coefficient‡ ± SE</td>
</tr>
<tr>
<td>Baseline Uosm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-PKD</td>
<td>0.62 ± 0.29</td>
<td>0.03</td>
<td>0.25 ± 0.26</td>
</tr>
<tr>
<td>PKD</td>
<td>0.07 ± 0.47</td>
<td>0.87</td>
<td>−0.03 ± 0.47</td>
</tr>
<tr>
<td>Follow-up Uosm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-PKD</td>
<td>1.25 ± 0.32</td>
<td>&lt;0.001</td>
<td>0.63 ± 0.30</td>
</tr>
<tr>
<td>PKD</td>
<td>1.55 ± 0.59</td>
<td>0.01</td>
<td>1.40 ± 0.60</td>
</tr>
</tbody>
</table>

Abbreviation: BP, blood pressure.
*Baseline covariates: GFR, BSA, race, MAP, urine protein-creatinine ratio, serum transferrin level, and serum HDL cholesterol level.
†Follow-up covariates: MAP, protein intake calculated from UUN, serum sodium level, and use of diuretics, ACE inhibitors, $\beta$-blockers, and calcium channel blockers.
‡Regression coefficients expressed as milliliters per minute per year of GFR slope 100 mOsm/L of Uosm.
combination of decreased serum sodium concentration and frankly hypotonic urine suggests that excess water intake was the cause of high daily urine volume, not a renal concentrating defect. Moreover, for patients with the highest daily urine volumes, MAP was either maintained (patients without PKD) or increased (patients with PKD; \( P < 0.001 \); Fig 5). The tendency toward maintained or increased MAP occurred despite a trend for greater diuretic (furosemide) use in those with the highest daily urine volumes (patients with PKD, \( P = 0.09 \); patients without PKD, \( P = 0.002 \); Fig 6). Thus, that blood pressure was maintained or increased despite increased diuretic use in those with high urine volumes suggests that primary renal salt wasting also was not the cause of high urine volumes.

Although daily urine volume and diuretic use were significantly associated, diuretic use was not significantly associated with GFR decline (\( P = 0.28 \) for patients without PKD; \( P = 0.54 \) for patients with PKD). For this analysis, diuretic use was a continuous variable (percentage of visits in which diuretics were administered) and
adjustment was made for all baseline and follow-up covariates.

Mean follow-up Posms were 303 ± 7 and 302 ± 6 mOsm/L in patients with and without PKD, respectively. Follow-up Uosm-Posm ratio in patients with and without PKD was significantly related to GFR decline in a manner similar to that shown for Uosm alone (Table 3). We also examined the association of GFR decline with baseline and follow-up total urinary osmoles (daily urine volume × Uosm), 24-hour urine sodium level, and 24-hour urine sodium plus potassium levels. None of these associations was significant (data not shown). Thus, the association of GFR decline with urine volume and Uosm was related to rate of urine water excretion, not solute excretion.

Median within-patient coefficient of variation of follow-up daily urine volume was 17.2% (10th percentile, 11.3%; 90th percentile, 24.7%). The between-patient coefficient of variation of the geometric mean of each patient’s follow-up daily urine volume values was larger at 30.2%. This indicates that variation between patients in urine volume was larger than variation in urine volume within patients over time. Thus, categorizing patients by geometric mean levels of follow-up daily urine volume is reasonable.

DISCUSSION

The present study used the MDRD study database to examine retrospectively the relationship between fluid intake (reflected by 24-hour urine volume and Uosm) and renal disease progression (decline in GFR). We tested the hypothesis that fluid intake is significantly associated with GFR decline during follow-up in the MDRD study. Patients with and without PKD were analyzed separately because of evidence from experimental models of renal disease that high fluid intake might increase progression in patients with PKD, but decrease progression in patients without PKD.

We found that for patients with and without PKD, there was a significant association between mean 24-hour urine volume and GFR decline during follow-up in the MDRD study. The higher the mean 24-hour urine volume, the greater the GFR decline. Mean 24-hour urine volume was significantly and inversely related to mean 24-hour Uosm. Thus, we also found a significant but inverse association between mean Uosm and GFR decline during follow-up in the MDRD study in patients with and without PKD.

To assess whether the association of faster GFR decline with higher urine volume and lower Uosm could be explained by their association with other risk factors for GFR decline, regressions of GFR decline on urine volume and Uosm were adjusted for relevant baseline and follow-up covariates. In patients without PKD, adjustments resulted in approximately a 50% reduction in the magnitude of the association of GFR decline with urine volume and Uosm. In patients with PKD, these adjustments had little effect on the magnitude of the association of GFR decline with urine volume and Uosm. This difference between patients with and without PKD was the result of a significant positive correlation between urine volume and urine protein-creatinine ratio. A possible reason for the greater effect of this covariate adjustment in patients without PKD is that they showed greater proteinuria than patients with PKD.

There are two principal hypotheses that can explain the association of high daily urine volume/low Uosm with faster GFR decline. Hypothesis 1 states that high urine volume/low Uosm causes faster renal disease progression. In this scenario, excess fluid intake causes nephron damage (and cyst growth in patients with PKD). Hypothesis 2 states that high urine volume/low Uosm is the result of faster renal disease progression. This scenario can be explained in two ways: (1) faster progression of renal disease directly causes increased urine volume/decreased Uosm by causing greater tubular injury (urinary concentrating defect and/or salt wasting), or (2) faster progression of renal disease indirectly causes increased urine volume by directly increasing thirst.

Our analysis favors hypothesis 1 because patients with and without PKD had findings suggestive of excess water intake. That is, those with the highest daily urine volumes had frankly hypotonic urine, significantly decreased serum sodium concentrating, and maintained or increased blood pressure despite increased diuretic use. If high daily urine volumes had been the result of a urinary concentrating defect and/or salt wasting (hypothesis 2), one or more of the following conditions should have been present in those
with the highest urine volumes: greater serum sodium concentration, isotension, and lower blood pressure. None of these were present. As discussed, usually the opposite condition was present. Also, although PKD is associated with decreased urine concentrating ability, PKD is not known to cause persistent frankly hypotonic urine.

With respect to hypothesis 2b (high urine volume in patients with faster progression of renal disease is the result of a primary increase in thirst), there is no evidence for such a phenomenon in humans with chronic renal insufficiency. The thirst mechanism and antidiuretic hormone (ADH) release have been shown to be normal in patients with chronic renal insufficiency. Thus, together, our analysis suggests that the association of high urine volume/low Uosm with faster GFR decline can be explained by an adverse effect of excess fluid intake on renal function in patients with chronic renal insufficiency. A hypothesis that could explain how high urine volume might cause faster renal disease progression is that high urine volume increases intratubular volume and pressure, and these stretch forces could induce fibrogenic mechanisms. In patients with PKD, increased intratubular pressure caused by high urine volume also could promote cyst growth. This effect of urine volume on cyst growth might explain why in part the inability of strict blood pressure control, decreased protein intake, or ACE inhibitors to slow PKD progression.

The present work should not be construed as an endorsement of water restriction in patients with chronic renal insufficiency. This could incur risks. For example, excessively reducing urine volume in patients with PKD could increase the risk for urolithiasis. A recent report showed that mean 24-hour urine volume in patients with PKD with urolithiasis (1.75 L) was significantly less than that of patients with PKD without urolithiasis (2.25 L). Thus, if excess fluid intake is curtailed in patients with PKD, it might be appropriate to use dietary and/or pharmacological measures that reduce the likelihood of urolithiasis. Also, studies in animal models of renal disease showed that water restriction that results in high ADH levels promotes progression of renal disease, in part because ADH induces glomerular hyperfiltration. Thus, in patients with chronic renal insufficiency, water restriction that results in elevated ADH levels may be as undesirable as excess fluid intake. Bakir and Trinh-Trang-Tan suggested that the optimum fluid intake to slow progression of experimental renal disease is intake that produces a Uosm/Posm ratio slightly greater than 1.00. Ideally, this would be achieved by reducing excess solute intake and appropriately changing water intake.

In animal studies that showed slowed renal disease progression with increased fluid intake, the animals were able to form hypertonic urine. Thus, it is possible that increased fluid intake might be beneficial in early renal disease when renal concentration is relatively intact. However, we cannot test that hypothesis with MDRD data.

In summary, high fluid intake that results in increased urine volume and low Uosm is not associated with slower renal disease progression. Indeed, high fluid intake might promote progression of renal disease, although this cannot be proved from this retrospective analysis. We suggest the most prudent interpretation of our findings is that until better data become available, patients with chronic renal insufficiency should not be encouraged to ingest a high fluid intake unless it is needed to manage such specific problems as nephrogenic or central diabetes insipidus or urolithiasis. Avoidance of excess fluid intake might be particularly important for those with PKD.

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