THE EFFECT OF FEEDING PROTEIN AND UREA ON THE
RENAL CONCENTRATING PROCESS

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(Submitted for publication December 7, 1956; accepted December 28, 1956)

The role of dietary protein in the treatment of
renal disease remains a controversial one, in part
because the influence of diets high or low in pro-
tein upon normal renal function is not completely
understood. In the rat much evidence has ac-
cumulated that high-protein intakes lead to renal
hyper trophy (1–3) and that low-protein diets may
be associated with impairment of some renal func-
tions (4). Pullman, Alving, Dern, and Landowne
(5) have recently reviewed this subject and dem-
onstrated that in normal men a high-protein diet
slightly increases and a low-protein diet slightly
depresses glomerular filtration rate, renal plasma
flow and TmPAH.

The present experiments were undertaken to in-
vestigate the effects of alterations in dietary pro-
tein on the ability of normal kidneys to concen-
trate the urine. Maximum renal concentrating
capacity was altered strikingly by changing the
dietary intake of nitrogen, both as protein and (an
unexpected finding) as preformed urea.

METHODOLOGY

Normal young men abstained from fluids for 12
hours prior to coming to the laboratory at 8 a.m. on
the day of the experiment. In order to assure a constant
and maximal antidiuretic stimulus throughout all ex-
periments, Pitressin® was infused at the rate of 200
milliliters in 50 cc. of normal saline per hour. Subjects
remained in the supine position, standing only to void.
Urine was collected at intervals until its osmolar con-
centration (freezing point) had reached a constant max-
imum. In some subjects mannitol was then infused, so
that the maximum ability of renal tubules to reabsorb
water free of solute (Tm°H₂O) could be measured in 5
to 16 separate determinations within a range of urine
flows from 10 to 35 cc. per min. Freezing points of se-
rum and urine were determined using the Fiske osmo-
 meter. Clearances of inulin (6), creatinine (7), urea (8),
sodium and potassium (9) were also determined. The
“t” test of “Student” was used in analyzing the data.

Each subject was studied after three days on a Low-
Protein diet, containing approximately 20 grams of pro-
tein per day, and a High-Protein diet, containing 150 to
200 grams of protein per day. The latter was achieved
by the use of 100 to 200 grams per day of a high-protein,
low-sodium supplement. Both diets were approximately
isocaloric; in some subjects the low-protein diet consisted
chiefly of carbohydrate, in others fat predominated. Two
subjects were studied on a Regular diet, containing about
90 grams of protein per day. Other variations in dietary
regimen will be described under Results.

RESULTS (Table I)

Effect of low- vs. high-protein diets on renal con-
centrating ability, inulin clearance and urea
clearance (Figure 1)

Maximum urinary solute concentration and
maximum osmolar U/P ratio were invariably
higher after three days of a high-protein intake
than after a diet low or normal with respect to
protein. The increases in maximum urinary
osmolality with ingestion of large amounts of pro-
tein varied from 87 to 332 mOsm. per K. This
occurred in association with an increased basal
urinary flow containing from 1.2 to 5 times the
quantity of solutes excreted on a low-protein diet.
The maximum capacity of the kidneys to reabsorb
water free of solute (Tm°H₂O) was likewise in-
creased by 15 to 50 per cent in five subjects by
feeding protein.

Inulin clearance was slightly higher in each of
five subjects on a high-protein regimen than on a
diet low in protein. At the high urine flows ob-
tained during mannitol diuresis, the clearance of

6 Melactin, supplied by E. R. Squibb and Co.
<table>
<thead>
<tr>
<th>Subject and Surface area</th>
<th>Regimen</th>
<th>Before infusion of mannitol</th>
<th>During infusion of mannitol</th>
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<td>U_{max} mOsm./K</td>
<td>V cc./min.</td>
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<td>F. E. 1.8 (m^2)</td>
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<td>717</td>
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<td>658</td>
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urea averaged 70 per cent of inulin clearance. Protein feeding did not change this ratio.

Failure of sodium chloride to improve renal concentrating ability (Figure 2)

In contrast to the effects of protein ingestion on renal concentrating ability, maximum urinary concentration, maximum U/P ratio and $T_m^{H_2O}$ were unchanged or slightly diminished when a low-protein diet was supplemented by 20 grams of sodium chloride per day (equivalent in osmolar terms to the urea from 100 grams of protein) for three days prior to the test. An increased excretory burden of solute supplied as electrolyte did not, therefore, produce the same increase in renal concentrating capacity as did solute derived from protein.
**Effect of adding urea to the diet (Figure 3)**

When a low-protein diet was supplemented for three days, not by 140 grams of extra protein but by an equivalent amount of nitrogen in the form of 48 grams of urea daily, a marked increase occurred in maximum urinary concentration, maximum U/P ratio and T$_{ur}$H$_2$O. This effect, observed in six instances, was in all respects indistinguishable from that seen in subjects eating a diet high in protein.

**Effects of protein and urea in overhydrated subjects (Figure 4)**

It seemed possible that the effects of protein and urea might be secondary to some slight or inapparent dehydration consequent to solute diuresis and insufficiently compensated for by the subject's sense of thirst. Two subjects (F. E. and C. K.) were studied repeatedly on diets low and high in protein, with and without supplements of urea, taken during a period of three days during which
5 to 6 liters of water were imbibed daily, an amount sufficient to keep the measured concentration of each specimen of urine voided during this time below 150 mOsm. per K. With the possibility of dehydration thus eliminated, supplementary feedings of protein and urea consistently increased maximum urinary concentration and $Tm^+H_2O$.

*Failure of acute loading with urea to increase $U_{max}$ or $Tm^+H_2O$ (Figures 5 and 6)*

Four subjects who had eaten a diet low or normal in protein for the preceding three days ingested 30 grams of urea acutely while receiving an infusion of Pitressin®. A typical experiment is shown in Figure 5. The osmolality of the urine was essentially unaltered although the excretion of urea quadrupled. On the other hand, maximum urinary concentration was considerably increased in every case after the same subject supplemented his diet with urea for three days. $Tm^+H_2O$ was not changed by the acute ingestion of 30 grams of urea in two subjects in whom this was tested (Figure 6).

**DISCUSSION**

Data in the literature concerning the influence of dietary variations on renal concentrating ability are scanty and conflicting. Addis and Shevky (10) suggested that healthy persons deprived of water might excrete urine of subnormal specific gravity when their diets contained little salt and protein; however, the differences which they obtained were of questionable statistical significance and were not confirmed by Miller, Price, and Longley (11) or by Addis and Foster (12), who in addition pointed out the fallacy of using specific gravity as a measure of osmolar concentration. McCance (13) found that maximum urinary osmolarity after dehydration was not changed appreciably by omitting salt from the diet or by ingesting urea on the morning of the experiment. Hayman, Shumway, Dumke, and Miller (14) reported that two dogs when deprived of fluids excreted urine of lower specific gravity while on a cracker-meal diet than when eating a diet composed largely of meat.

The effect of protein in producing hypertrophy of the kidneys of rats is well known (1–3). When the diet of rats is supplemented by nitrogen in the form of urea, renal hypertrophy, less marked than with protein feeding, has been reported (2, 15, 16) and denied (1, 3). Other urinary solutes, with the possible exception of phosphate, do not enhance renal growth (1). Protein feeding greatly increases renal blood flow and glomerular filtration rate in the dog (17) but in man (5) it produces only a slight, though statistically significant, rise in the clearances of insulin and PAH and in $TmPAH$.

The idea that urea might obligate less water in the urine than an equimolar quantity of salt, or that maximum concentrations of urea could be attained in a urine already maximally concentrated with respect to salt was suggested by earlier workers (18, 19), but Hervey, McCance, and Taylor (20, 21) concluded from acute studies in healthy persons during hydropenia that at every urine flow there was a limiting osmotic pressure which did not discriminate between osmols of urea and osmols of salt. The present experiments indicate that in order for urea to raise maximum urinary osmolarity it must be administered over several days. It is therefore unlikely that its effect can be explained solely or chiefly by an unusual ability of the renal tubules to concentrate urea per se to a greater extent than other solutes. If that were the case, prior feeding of protein or urea would not increase urinary osmolarity measured during the determination of $Tm^+H_2O$, when mannitol and not urea is the chief osmotic constituent of the urine. Even if the assumption is made that urea obligates no water at all, only 20 to 50 per cent of the increases in $Tm^+H_2O$ observed in the present experiments after diets high in protein or urea could be accounted for by the presence of additional urea in the urine under mannitol diuresis.

The ability of renal tubules to secrete urea into the urine has been established for the kidney of the agglomerular fish (22), the frog (23, 24) and the kangaroo rat (25). It is conceivable that a

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*It should be pointed out, however, that there are no published data dealing intensively with the effects of urea vs. other solutes on the limiting osmotic pressure of the urine in man below a urine flow of about 3 cc. per min. Scattered observations suggest that during hydropenia a maximum urinary osmolar concentration may be maintained at somewhat higher levels of urine flow and solute excretion when urea is utilized as a loading solute than when sodium chloride is loaded (13).*
similar mechanism, stimulated by the necessity to excrete large amounts of urea, might operate in humans to add urea to tubular urine at or past the point where water is abstracted in the final concentrating process. Such a process should, however, be reflected in a rise in the urea/inulin clearance ratio. This did not occur in the present experiments in which $C_{\text{urea}}/C_{\text{inulin}}$, measured during osmotic diuresis, was remarkably constant on both high and low-protein diets. Secretion of urea by renal tubules is therefore not a likely explanation for the augmentation of renal concentrating ability which was observed during mannitol diuresis in persons fed protein or urea. These data do not rule out the possibility that increased tubular secretion of urea might operate at low urine flows to increase maximum urinary osmolality, an effect which might be submerged in the flood of proximal tubular urine accompanying mannitol diuresis. It is interesting in this connection that Schmidt-Nielsen observed that mannitol appeared to block tubular excretion of urea in the kangaroo rat (25).

Although the kidneys can apparently be "trained" to concentrate more or less efficiently by several days of water deprivation or of forced drinking (26), the explanation of the influence of protein and urea on the concentrating process does not lie with any tendency they might have to produce dehydration. Dietary supplements of protein and urea improved renal concentrating ability in continuously overhydrated subjects as well as in persons permitted to drink at will.

It seems clear that physiological variations in dietary protein and urea excretion, as well as in the state of bodily hydration (26), produce well-marked and separable adaptive responses on the part of the renal tubules which have an obvious utility in terms of the body's economy of water. Such effects upon renal concentrating ability cannot be neglected in future considerations of the effects of disease or therapy upon this important function of the kidneys.

**SUMMARY**

1. Changes in the dietary intake of nitrogen, either as protein or urea, over a period of three days, in normal subjects produced well-marked parallel changes in maximum urinary osmolar concentration and in $T_m^{\text{H}_2\text{O}}$.

2. Feeding of protein or urea increased the renal response to Pitressin® in continuously overhydrated subjects as well as in persons permitted to drink at will.

3. Although $U_{\text{max}}$ and $T_m^{{\text{H}}_2{\text{O}}}$ were increased by chronic administration of urea, they were not increased by acute urea loading.

4. The data suggest that both protein and urea, when administered chronically, promote an adaptive response by the renal tubules by which water is conserved more efficiently and renal concentrating ability is augmented.

**REFERENCES**


