In early diabetes, the kidney grows and the glomerular filtration rate (GFR) increases. This growth is linked to ornithine decarboxylase (ODC). The study of hyperfiltration has focused on microvascular abnormalities, but hyperfiltration may actually result from a prior increase in capacity for proximal reabsorption which reduces the signal for tubuloglomerular feedback (TGF). Experiments were performed in Wistar rats after 1 week of streptozotocin diabetes. Kidney weight, ODC activity, and GFR were correlated in diabetic and control rats given difluoromethylornithine (DFMO; Marion Merrell Dow, Cincinnati, Ohio, USA) to inhibit ODC. We assessed proximal reabsorption by micropuncture, using TGF as a tool for manipulating single-nephron GFR (SNGFR), then plotting proximal reabsorption versus SNGFR. ODC activity was elevated 15-fold in diabetic kidneys and normalized by DFMO, which also attenuated hyperfiltration and hypertrophy. Micropuncture data revealed an overall increase in proximal reabsorption in diabetic rats too great to be accounted for by glomerulotubular balance. DFMO prevented the overall increase in proximal reabsorption. These data confirm that ODC is required for the full effect of diabetes on kidney size and proximal reabsorption in early streptozotocin diabetes and are consistent with the hypothesis that diabetic hyperfiltration results from normal physiologic actions of TGF operating in a larger kidney, independent of any primary malfunction of the glomerular microvasculature.
Ornithine decarboxylase, kidney size, and the tubular hypothesis of glomerular hyperfiltration in experimental diabetes

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Received for publication August 4, 2000, and accepted in revised form December 12, 2000.

In early diabetes, the kidney grows and the glomerular filtration rate (GFR) increases. This growth is linked to ornithine decarboxylase (ODC). The study of hyperfiltration has focused on microvascular abnormalities, but hyperfiltration may actually result from a prior increase in capacity for proximal reabsorption which reduces the signal for tubuloglomerular feedback (TGF). Experiments were performed in Wistar rats after 1 week of streptozotocin diabetes. Kidney weight, ODC activity, and GFR were correlated in diabetic and control rats given difluoromethylornithine (DFMO; Marion Merrel Dow, Cincinnati, Ohio, USA) to inhibit ODC. We assessed proximal reabsorption by micropuncture, using TGF as a tool for manipulating single-nephron GFR (SNGFR), then plotting proximal reabsorption versus SNGFR. ODC activity was elevated 15-fold in diabetic kidneys and normalized by DFMO, which also attenuated hyperfiltration and hypertrophy. Micropunctured data revealed an overall increase in proximal reabsorption in diabetic rats too great to be accounted for by glomerulotubular balance. DFMO prevented the overall increase in proximal reabsorption. These data confirm that ODC is required for the full effect of diabetes on kidney size and proximal reabsorption in early streptozotocin diabetes and are consistent with the hypothesis that diabetic hyperfiltration results from normal physiologic actions of TGF operating in a larger kidney, independent of any primary malfunction of the glomerular microvasculature.

standing a concomitant decrease in plasma volume which would otherwise oppose an increase in GFR (7).

It is also not fully understood what mediates renal enlargement at the onset of diabetes or how growth of the tubule and tubular reabsorption might be linked in diabetes. Nonetheless, an arginine–ornithine–polyamine pathway has been implicated in enlargement of the diabetic kidney. For instance, it has been shown that the activity of ornithine decarboxylase (ODC) in the kidney increases after only 1 day of streptozotocin diabetes, and that the ODC inhibitor, difluoromethylornithine (DFMO), attenuates renal hypertrophy during the first week of streptozotocin diabetes (8). In the present study, ODC activity was manipulated in order to demonstrate that renal hypertrophy, increased proximal reabsorption, and glomerular hyperfiltration are all linked to increased renal ODC activity in rats with early diabetes. We also studied the activity of arginine decarboxylase (ADC), an enzyme whose product, agmatine, has been shown to deactivate ODC through induction of ODC antizyme (9).

**Methods**

Overview. Studies were performed in male Wistar rats after 7–8 days of insulin-treated diabetes and in weight-matched controls. First, the role of ODC in renal hypertrophy and hyperfiltration was assessed by measuring ODC activity in kidney cortex and by measuring the effects of the ODC inhibitor, DFMO (Marion Merrell Dow, Cincinnati, Ohio, USA), on kidney weight and GFR. Then micropuncture experiments were performed, in which the TGF signal was artificially manipulated as a means for changing single-nephron GFR (SNGFR) so that proximal tubular reabsorption could be characterized over a range of SNGFR in individual nephrons. The effects of diabetes and/or DFMO on this relationship between kidney size, proximal reabsorption, and SNGFR were analyzed as a direct test of the tubular hypothesis. All animal experiments were conducted in accordance with the NIH Guide for the Care and Use of Laboratory Animals.

The TGF paradigm and the tubular hypothesis of hyperfiltration. This “tubular hypothesis” of hyperfiltration is easily understood in terms of a familiar framework for describing interactions between the glomerulus and tubule which comprise the TGF system (shown in Figure 1). According to this framework, glomerular filtration and tubular reabsorption are linked by two physiologic processes. The first of these is glomerulotubular balance (GTB), which confers a forward effect of SNGFR on the fluid delivered to the distal nephron. The second process is TGF. TGF operates in the juxtaglomerular apparatus of each nephron and causes SNGFR to vary inversely with changes in the amount of salt reaching the macula densa at the end of Henle’s loop. Together, GTB and TGF comprise a closed-loop negative feedback control system which stabilizes nephron function.

The nephron must operate at the point where curves representing GTB and TGF intersect, and it follows that diabetic hyperfiltration cannot occur unless diabetes affects either the TGF curve or the GTB curve. Changes in glomerular hemodynamics that are not mediated by movement along the TGF curve are represented by a shift in the TGF curve itself, and changes in reabsorption which are not explained by the direct action of GTB are represented by a shift in the GTB curve. Therefore, hyperfiltration might be explained by a shift in either the TGF curve (a primary vascular event) or the GTB curve (a primary tubular event). If hyperfiltration is initiated by a primary vascular event, then the operating point must initially move down and to the right as well as upward, and hyperfiltration must be accompanied by increased late proximal flow (VLP) (shown in Figure 1, left panel). On the other hand, if hyperfiltration is initiated by a primary increase in proximal reabsorption, then the operating point must initially move leftward as well as upward, and hyperfiltration should be accompanied by a reduction in VLP (shown in Figure 1, center panel). Therefore, by testing for changes in VLP, one should be able to determine whether diabetic hyperfiltration is mainly the result of defective tone in the glomerular microvasculature or the consequence of increased proximal reabsorption. However, to actually measure VLP at the operating point is not trivial, since fluid cannot be collected from the proximal tubule without interrupting TGF, thereby rendering the operating point indeterminate. Another difficulty in establishing whether a change in reabsorption has led to a change in SNGFR or vice versa is that a primary shift in GTB will elicit a secondary resetting of TGF that maintains the operating point on the steep portion of the TGF curve (10, 11). In other words, a primary across-the-board increase in proximal reabsorption is expected to elicit a secondary shift in baseline arteriolar resistance, making it difficult to discern which came first (shown in Figure 1, right panel). In the present study, we applied micropuncture methods designed to overcome these difficulties in order to test the hypothesis that diabetic hyperfiltration results from a primary increase in proximal reabsorption.

The diabetic model. Male Wistar rats were made diabetic with streptozotocin (65 mg/kg, injected intraperitoneally; Sigma Chemical Co., St. Louis, Missouri, USA). Blood glucose concentration was measured after 24 hours and rats with levels greater than 300 mg/dl were carried on into studies. Insulin (PZI; Blue Ridge Pharmaceuticals, Memphis, Tennessee, USA) was administered daily by subcutaneous injection and titrated to maintain blood glucose concentrations of 200–400 mg/dl. To inhibit ODC, 200 mg/kg DFMO was administered intraperitoneally 12 hours in Ringer saline beginning 12 hours after the streptozotocin. Animals not receiving DFMO were injected with Ringer saline placebo.

Bioassay of ODC/ADC activity. ODC and ADC activity were assayed in kidney cortex as previously described (9). After 7 days of diabetes, and 2–4 hours after the final dose of DFMO, animals were anesthetized with Brevital (75 mg/kg), and the kidneys were removed. A slice of renal cortex was placed in ice-cold reaction buffer (10...
mM Tris [pH 7.4], 2.5 mM dithiothreitol [DTT], 0.3 mM pyridoxyl-5-phosphate, 0.1 mM EDTA, and 0.5 mM Pefabloc [Roche, Chicago, Illinois, USA], lysed (polys- 
tron for 5 seconds), and centrifuged (30,000 g for 40 minutes). Supernatant was collected for ODC assay. 
The pellet was resuspended in ADC reaction buffer (10 
M TRIS, 2.5 mM DTT, 0.8 mM MgSO4, 0.5 mM Pefa-
bloc [pH 8.5]) for ADC assay. 250 µl was added to a glass 
reaction tube and incubated for 10 minutes with or 
without DFMO to establish specificity in ODC assay 
and in α-fluoromethyl arginine (DFMA, a specific ADC 
blocker) to establish specificity in ADC assay. 100 µM 
14C-ornithine or 14C-L-arginine (Dupont NEN 
Research Products, Boston, Massachusetts, USA) were 
added (0.1 µCi) for ODC or ADC assays, respectively. 
Tubes were vortexed and capped with trapping wells 
(Kontes, Vineland, New Jersey, USA) containing 250 µl 
CO2 trapping agent (Solvall; Dupont NEN Research 
Products). After incubation for 1 hour at 37°C, 
trichloroacetic acid was added to stop the reaction. 
Radioactivity in the trapping well was counted after 
night equilibration of 14CO2 gas between the solu-
tion and trapping well. ODC and ADC activities were 
normalized to total protein by the Lowry method (12). 
Measurement of kidney weight. To determination wet 
weight, kidneys were removed and decapsulated. The 
decapsulated kidney was placed on tissue paper for 1 
minute, then transferred to a scale and weighed. In 22 
of these kidneys, the mass of fat-free dry solids was also 
determined after drying overnight in an oven at 50°C. 
Measurement of GFR. Experiments were performed 4–6 
hours after the final dose of DFMO. Rats were anes-
ethetized with Inactin (100 mg/kg injected intraperi-
toneally) and body temperature maintained on a servo-
controlled heating table. Tracheostomy was performed 
(PE240) and catheters placed in the left femoral artery, 
right internal jugular vein, and urinary bladder (PE50). 
Ringer saline containing 5–10 µCi/ml 3H-inulin was 
infused as a marker of GFR. Diabetic animals received 3.5 
ml/h, and nondiabetics received 2 ml/h to account for the 
effect of hyperglycemia on urine output. One hour was 
allowed for equilibration, then GFR was measured by 3H-
inulin clearance in two 30-minute periods. The two val-
ues for each rat were averaged. Subsequently, animals 
were euthanized and kidneys removed for weighing. 
Assessment of proximal reabsorption. Micropuncture 
experiments were performed to assess the impact of 
diabetes ± DFMO on overall reabsorption and GTB in 
the proximal tubule. Because of GTB, reabsorption will 
varies with SNGFR, and hyperfiltration will confound 
other effects of diabetes on proximal reabsorption. To 
correct for this, we manipulated the TGF signal to 
determine the relationship of proximal reabsorption to 
SNGFR in individual nephrons, then tested for the 
effects of diabetes ± DFMO on this relationship. Experi-
ments were performed 3–7 hours after the last dose of 
DFMO. Animals were anesthetized and catheters 
placed as described above. The left kidney was exposed 
and prepared for micropuncture according to standard 
protocols employed in this laboratory (13). 3H-inulin 
(80–100 µCi/ml in Ringer saline) was infused as a 
marker of SNGFR. Diabetic animals received Ringer 
saline at 3.5 cm3/h. Nondiabetic animals received 
Ringer saline at 2 cm3/h. After 1 hour for equilibration, 
late proximal nephrons were localized on the kidney 
surface, and an obstructing wax block was inserted 
immediately upstream from the most downstream 
accessible segment. A microperfusion pipette contain-
ing artificial tubular fluid (ATF) was inserted down-
stream from the wax block to perfuse Henle's loop. 
Controlled perfusion of Henle's loop was performed in 
order to activate TGF and thereby cause SNGFR to 
change. During perfusion, timed collections of tubular 
fluid were taken upstream from the wax block in order 
to measure SNGFR and VLP. Collections were taken 
from each nephron during minimal TGF activation 
(Henle's loop microperfusion at 0 nI/min) and maxi-
tal TGF activation (Henle's loop microperfusion at 40 
ml/min). Perfusions were in random order. Nephrons 
were vented upstream from the wax block before each 
collection to prevent pressure from building up in the 
proximal tubule. Tubular fluid samples were assayed 
for volume by transfer to a constant-bore glass capil-
lar, then counted for radioactivity to determine 
SNGFR. Data from these paired collections were 
employed to characterize the dependence of proximal 
reabsorption on SNGFR. 

Statistical analysis. Data were analyzed by ANOVA 
or by Student’s t test with Bonferroni correction as 
appropriate. Where new variables were created by mathematically 
combining measured variables, standard errors 
were determined based on the principle that 

**Equation 1**

If \( \psi = f(x_1, \ldots, x_n) \), Then \( \sigma^2 = \sum_{i=1}^{n} \frac{\partial \psi}{\partial x_i} \sigma_i \sigma_i \)

Where \( \sigma_i \) are random variables, \( \sigma^2 \) refers to the mean 
square error, and \( r_{x_i, x_j} \) the correlation between \( x_i \) and \( x_j \).
Results

GFR, kidney weight, and ODC activity. Body weight, blood pressure, blood glucose concentration, hematocrit, and urine flow rate are depicted in Table 1, n = 6–8 animals per group. Rats with diabetes had elevated blood sugar, greater urine output, and weighed slightly less than controls. Body weight, blood pressure, blood glucose concentration, hematocrit, and urine flow were not affected by DFMO.

Although body weight was somewhat less in the diabetic rats, diabetic kidneys weighed 50% more than control kidneys (P < 0.00005), and GFR in diabetes was 25% greater than in controls (P < 0.0002). Diabetic rats receiving DFMO had less kidney mass (P = 0.0004) and lower GFR (P = 0.0012). In contrast, control rats receiving DFMO tended to manifest greater kidney weight and GFR than untreated controls, but these effects did not achieve statistical significance. Kidney weight was a strong predictor of GFR (r = 0.998 for the group means, Figure 2). For the subset of kidneys in which both wet and dry weights were measured, the two were tightly correlated (r = 0.962). Therefore, changes in wet weight corresponded to changes in the mass of fat-free dry solids.

ODC activity in kidney cortex was elevated 15-fold in diabetes. In diabetic rats treated with DFMO and studied 2–4 hours after the final dose of DFMO, ODC activity was not different from control rats (Figure 3). When compared to control kidneys, ADC activity was lower by 20% in rats with early diabetes. In diabetic rats receiving DFMO, the kidney ADC activity level was between that of control and diabetic rats not receiving DFMO (Figure 3).

Micropuncture results. Micropuncture data were obtained from 177 nephrons in 29 rats (control n = 11, other groups n = 6) weighing 295 ± 26 grams (mean ± SD, P = not significant between groups). SN GFR and VLP were measured in each nephron during orthograde perfusion of Henle’s loop at both 0 and 40 nL/min. In this way, TGF was used as a tool for manipulating SN GFR so that proximal reabsorption could be characterized at more than one delivered load. The results are depicted in Table 2 and Figure 4. The value of SN GFR obtained during zero TGF stimulus is referred to as SN GFR0. The value for SN GFR obtained during maximum TGF stimulus is referred to as SN GFR40. SN GFR0 was greater in diabetics than controls (P = 0.016), consistent with typical diabetic hyperfiltration. In diabetic rats receiving DFMO, SN GFR0 was significantly less than in non-treated diabetics (P = 0.001) and not different from control rats (P = 0.65). In contrast, DFMO had no significant effect on SN GFR0 in control rats (P = 0.5). By two-way ANOVA the effect of DFMO on SN GFR0 differed between diabetic and control rats (P < 0.0003). SN GFR40 was less than SN GFR0 in all nephrons indicating successful activation of TGF.

The main point of these micropuncture experiments was to evaluate proximal reabsorption as a function of SN GFR. This was done in order to overcome the confounding effect of differences in SN GFR as the apparent cause of differences in proximal reabsorption. In Figure 4 these results are expressed as absolute proximal reabsorption (APR), fractional proximal reabsorption, or late proximal TF/Pinulin. (TF/Pinulin = 1/fractional delivery, and is a standard index for tracking water flux along the nephron). When comparing diabetic to control rats (Figure 4a), each of these indices suggests that proximal reabsorption is increased in diabetes. From these data it is also clear that fractional reabsorption was not constant, but declined with increasing SN GFR. The rate at which fractional reabsorption declined with increasing SN GFR was greater in diabetes, such that regression lines drawn for the data in each group converged near the SN GFR0 values for diabetes. In other words, the impact of diabetes on proximal reabsorption was obscured when SN GFR was maximized by eliminating flow past the macula densa.

Figure 2
GFR vs. kidney weight in diabetic and control rats ± treatment with DFMO. GFR is for two kidneys. Kidney weight is wet-weight of left kidney. Dashed lines are 95% confidence intervals for linear regression. r² = 0.996.

Figure 3
Activities of ODC and ADC in control and diabetic rats. AP < 0.05 vs. control.
Intergroup comparisons for APR, late proximal TF/Pinulin, and proximal fractional reabsorption (FPR) were made as follows: First, the range of overlapping SNGFR was determined for the two groups being compared. Second, values for the index being examined and its SE were calculated as functions of SNGFR by linear interpolation between the values at SNGFR₀ and SNGFR₄₀. Third, intergroup comparisons were made by Student’s t test at each end of the overlapping range with Bonferroni correction for multiple group comparisons. The results are apparent in Figure 4. Comparing diabetics with controls, APR was greater in diabetics at the lower end of the SNGFR overlap (P = 0.02), while TF/Pinulin and FPR were greater in diabetes at both ends of the overlap region (P < 0.02 for each relevant comparison, Figure 4a). Among diabetics, treatment with DFMO reduced APR, TF/Pinulin, and FPR throughout the entire region of SNGFR overlap (P <0.0003 for each relevant comparison, Figure 4b). In control rats, DFMO had no effect on APR, TF/Pinulin, or FPR anywhere within the region of overlapping SNGFR (Figure 4c). Comparing control + DFMO with diabetes + DFMO, APR, TF/Pinulin, and FPR were each less in the diabetes + DFMO group throughout the region of overlapping SNGFR (P <0.01 for each relevant comparison, Figure 4d).

These data also permit assessment of proximal tubular function in terms of the efficiency of GTB. An index of autoregulatory efficiency for GTB was calculated analogous to the index used to describe autoregulatory efficiency of blood flow in an organ. For GTB, this index is given by:

\[
\frac{\Delta APR}{\Delta SNGFR} \times \frac{SNGFR}{SNGFR}
\]

This dimensionless index will equal unity when fractional reabsorption is constant over the range tested and will equal zero when absolute reabsorption is independent of SNGFR. This index also reduces the noise inherent in the simpler index, \(\Delta APR/\Delta SNGFR\), arising from variability in the location along the nephron where the micropuncture samples are obtained. Group mean indices for GTB efficiency were calculated from the data and variances determined as described under Methods. The results are depicted in Figure 5 and illustrate that, while overall proximal reabsorption is increased in diabetes, GTB efficiency is reduced in diabetes and not restored by DFMO.

![Image of Figure 4](image_url)

**Figure 4**
Pairwise intergroup comparisons of different indices of proximal reabsorption, each shown as a function of SNGFR. Line segments connect data obtained at minimum (higher SNGFR) and maximum (lower SNGFR) TGF activation. Group mean ± SEM is also shown in Table 2. Statistical significance of intergroup differences are described in the text for the regions of overlapping SNGFR. (a) Effects of diabetes. (b) Effects of DFMO in diabetes. (c) Effects of DFMO in controls. (d) Diabetes + DFMO vs. control + DFMO. DM, diabetes mellitus; Con, control.

**Table 1**
Baseline data in rats used to determine GFR, kidney weight, and ODC activity

<table>
<thead>
<tr>
<th>Group</th>
<th>Body wt (g)</th>
<th>Blood pressure (mmHg)</th>
<th>Blood glucose (mg/dl)</th>
<th>Hct (%)</th>
<th>Urine flow (μl/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>341 ± 11</td>
<td>111 ± 4</td>
<td>112 ± 3</td>
<td>46 ± 1</td>
<td>14 ± 2</td>
</tr>
<tr>
<td>Control + DFMO</td>
<td>348 ± 6</td>
<td>105 ± 4</td>
<td>119 ± 9</td>
<td>46 ± 1</td>
<td>16 ± 4</td>
</tr>
<tr>
<td>Diabetes</td>
<td>303 ± 6</td>
<td>96 ± 5</td>
<td>321 ± 24</td>
<td>47 ± 1</td>
<td>28 ± 5</td>
</tr>
<tr>
<td>Diabetes + DFMO</td>
<td>299 ± 6</td>
<td>102 ± 5</td>
<td>303 ± 10</td>
<td>47 ± 1</td>
<td>23 ± 3</td>
</tr>
</tbody>
</table>

*AP < 0.05 vs. control or control + DFMO.*
Discussion

These studies were performed to test the hypothesis that glomerular hyperfiltration in diabetes begins with growth of the kidney, which increases proximal reabsorption and reduces the signal for TGF. This departs from the usual notion that diabetic hyperfiltration results from intrinsic loss of tone in some element of the glomerular microvasculature. Normal GTB dictates that proximal reabsorption will change monotonically with SNGFR. Hence, there is no surprise in finding increased net tubular reabsorption in the hyperfiltering diabetic kidney. However, in the present case, proximal reabsorption remained elevated even after correcting for hyperfiltration, implying an increase in reabsorption beyond the amount attributable to GTB. Therefore, the difference in proximal reabsorption between control and diabetic rats was too great to have resulted from the difference in SNGFR. On the other hand, TGF-mediated vasodilation of the glomerular arterioles is the expected consequence of a primary increase in proximal reabsorption and likely accounts for hyperfiltration in this model of early diabetes. Based on prior work of others we anticipated that growth of the diabetic kidney would be linked to increased ODC activity (8). The present findings confirm this observation and extend it by demonstrating that chronic ODC blockade also mitigates the characteristic increases in proximal reabsorption and GFR in rats with diabetes, while bearing no significant effect on glomerular filtration or proximal reabsorption in nondiabetic rats. Since it is the tubule which mainly accounts for enlargement of the kidney in diabetes, it is likely that DFMO mitigates diabetic hyperfiltration by preventing the tubule from growing.

As illustrated in Figure 1, when a primary increment in tubular reabsorption leads to a TGF-mediated increase in SNGFR, there should be a telltale reduction in the remaining TGF stimulus. In other words, salt concentration at the macula densa, or ambient VLP, should be reduced in diabetes. Because the collection of late proximal fluid for the purpose of measuring VLP interrupts TGF and renders the operating point indeterminate, it is not possible to assess natural VLP by simple micropuncture of the late proximal tubule. Nonetheless, late proximal collections have been made by various investigators in the process of studying glomerular hemodynamics in diabetes which are analogous to those presently obtained during zero perfusion of Henle’s loop. Under these conditions, fractional reabsorption does not appear to be affected by diabetes. From this, one might simply conclude that diabetic and nondiabetic nephrons operate at different points along identical GTB curves and that net reabsorption is greater in the hyperfiltering diabetic due to the normal action of GTB. Misstatements in some textbooks which equate GTB with constant fractional reabsorption reinforce this impression. In fact, the present data confirm that fractional reabsorption increases with TGF activation, especially in diabetes. Manipulating TGF to adjust SNGFR reveals a major primary increase in proximal reabsorption attributable to diabetes. This is concealed when flow past the macula densa is reduced to zero and fractional reabsorption converges to the same value in diabetic and nondiabetic nephrons. Since nephrons do not naturally operate with zero distal flow, convergence at this particular point has no intrinsic physiological significance.

The limitations of micropuncture notwithstanding, there are published data consistent with the present finding that diabetes causes a shift toward greater proximal reabsorption. In a prior study, we employed a noninvasive optical technique to measure late proximal flow in rats after 5 weeks of diabetes. Under those conditions, which allowed for normal operation of TGF during the measurements, VLP was not increased in

![Figure 5](image_url)

**Figure 5**

Autoregulation of tubular reabsorption by GTB. Refer to text for formula of index. Index is unity when fractional reabsorption is constant across SNGFR. Index is zero when net reabsorption is constant across SNGFR. $^aP < 0.05$ vs. control. $^bP < 0.05$ vs. diabetes.

![Figure 6](image_url)

**Figure 6**

A working model to account for the effects of DFMO in the present study. The effects denoted with question marks have been demonstrated in cell culture (see text for references), but remain to be confirmed in diabetic animals.
Table 2
Micropuncture data (mean ± SEM)

<table>
<thead>
<tr>
<th>Group</th>
<th>SNGFR&lt;sub&gt;40&lt;/sub&gt; (nl/min)</th>
<th>SNGFR&lt;sub&gt;0&lt;/sub&gt; (nl/min)</th>
<th>APR&lt;sub&gt;40&lt;/sub&gt;</th>
<th>APR&lt;sub&gt;0&lt;/sub&gt;</th>
<th>TF/Pin&lt;sub&gt;40&lt;/sub&gt; (nl/min)</th>
<th>TF/Pin&lt;sub&gt;0&lt;/sub&gt; (nl/min)</th>
<th>FPR&lt;sub&gt;40&lt;/sub&gt; (nl/min)</th>
<th>FPR&lt;sub&gt;0&lt;/sub&gt; (nl/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>26.9 ± 1.1</td>
<td>39.4 ± 1.3</td>
<td>14.2 ± 0.7</td>
<td>18.1 ± 0.8</td>
<td>2.25 ± 0.08</td>
<td>1.89 ± 0.04</td>
<td>0.52 ± 0.02</td>
<td>0.46 ± 0.01</td>
</tr>
<tr>
<td>Control + DFMO</td>
<td>28.9 ± 1.5</td>
<td>42.7 ± 1.6</td>
<td>14.8 ± 0.8</td>
<td>18.3 ± 1.1</td>
<td>2.32 ± 0.12</td>
<td>1.82 ± 0.06</td>
<td>0.53 ± 0.02</td>
<td>0.43 ± 0.02</td>
</tr>
<tr>
<td>Diabetes</td>
<td>25.5 ± 1.7</td>
<td>46.3 ± 2.1</td>
<td>16.1 ± 0.9</td>
<td>20.1 ± 1.3</td>
<td>3.63 ± 0.32</td>
<td>1.85 ± 0.06</td>
<td>0.65 ± 0.03</td>
<td>0.43 ± 0.02</td>
</tr>
<tr>
<td>Diabetes + DFMO</td>
<td>24.2 ± 1.3</td>
<td>36.8 ± 1.9</td>
<td>11.1 ± 0.9</td>
<td>12.3 ± 0.9</td>
<td>2.80 ± 0.13</td>
<td>1.54 ± 0.05</td>
<td>0.45 ± 0.03</td>
<td>0.33 ± 0.02</td>
</tr>
</tbody>
</table>

P values from two-way ANOVA for effects of diabetes, DFMO, and diabetes × DFMO

- DM NS <0.0005 0.001 <0.0005 <0.0005 0.001 <0.0005 <0.0005
- DFMO NS 0.072 0.013 <0.0005 <0.0005 0.001 <0.0005 <0.0005
- DM NS <0.0005 0.001 <0.0005 <0.0005 0.003 NS <0.0005

Subscripts 0 and 40 refer to the rate at which Henle's loop was perfused (nl/min) as the data were collected. Perfusion at 40 nl/min was intended to maximally stimulate TGF. APR, net proximal reabsorption; TF/Pin, late proximal TF/Pinulin; FPR, fractional proximal reabsorption; DM, diabetes mellitus; NS, not significant.

In the present study, TGF was used as a tool to manipulate SNGFR. The main purpose was not to study TGF itself. However, as a condition of being able to manipulate SNGFR, it was necessary for there to be a TGF response. As it turned out, the range over which SNGFR could be manipulated by activating TGF with ATF was extended beyond normal in diabetes. This is somewhat surprising, since we previously demonstrated that diabetes reduces the sensitivity with which the TGF system compensates for small perturbations in ambient tubular flow (14). However, the sensitivity to small perturbations depends on the slope of the TGF curve at the operating point rather than on the maximum range of the TGF response that was tested in the present study. With caveats regarding the perfusate composition, and noting that the prior study was conducted in 5-week diabetics, the combined studies suggest a TGF curve in diabetes which is flatter at its steepest point, but which operates over a wider range.

The present data reveal that, within the physiologic range over which SNGFR can be made to vary by TGF, diabetes causes a primary increase in proximal reabsorption. However, they also reveal that the efficiency of GTB is reduced in diabetes. This is suggested by contrasting the slopes in Figure 4a and is revealed more explicitly by the efficiency calculated for GTB and shown in Figure 5. Furthermore, DFMO prevented the overall increase in proximal reabsorption (presumably by preventing the tubule from growing), but did not normalize the efficiency of GTB. This combination is what one would expect if growth of the tubule leads to a generalized increase in the many transporters that contribute to proximal reabsorption, while diabetes, independent of growth, further increases transport by a saturable high-affinity transporter, i.e., the sodium-glucose cotransporter which is pressed into service by hyperglycemia. Indeed, it was recently reported that hyperfiltration can be postponed by aggressive insulin treatment in the first days of streptozotocin diabetes even though the kidney becomes larger (17).

Overall proximal reabsorption was lower in diabetics receiving DFMO than in controls receiving DFMO. This likely resulted from osmotically-mediated reductions in late proximal TF/P for Na and Cl, which normally drives passive reabsorption in this segment (18).

The effects of DFMO imply that ODC-polyamine pathways participate in diabetic renal hypertrophy. Biogenic polyamines are required for cell-cycle progression and proliferative growth (19, 20). However, the diabetic kidney enlarges through hypertrophy, not hyperplasia. Since induction of ODC is a mid-G1 cell-cycle event, the role of ODC in early diabetes might be to increase activity and fidelity of protein translation (20), consistent with the increased protein/DNA ratio in early diabetic kidneys (6).

In the present study, DFMO given to diabetic animals completely normalized ex vivo ODC activity, whereas its effects on renal hypertrophy and hyperfiltration were incomplete. The growth-inhibitory effects of DFMO in cancer trials have likewise been incomplete (21, 22) due to enhanced cellular uptake of polyamines which circumvent ODC (23, 24). The current results may be due to the bioassay for ODC having been performed 4 hours into a 12-hour DFMO dosing interval. However, it is also possible that the DFMO-treated diabetic kidney compensates by taking up exogenous polyamines, or that the dependence of hyperfiltration on polyamines is incomplete.
We previously reported that agmatine, which is made from L-arginine by ADC, is present in micromolar concentrations in the kidney (25) and that agmatine reduces the polyamine content of cultured cells by inducing ODC antizyme and by blocking polyamine transporters (9). However, the degree to which endogenous agmatine normally governs the polyamine content or size of the proximal tubule is not known. This issue is difficult to address in vivo where DFMA, the drug normally used to block ADC, is converted to DFMO (J. Satriano, unpublished observation). In the present study, ADC activity was reduced in diabetes although the magnitude of this effect was small relative to the increase in ODC activity and ADC would have to impact ODC with high gain in order for reduced endogenous ADC activity to have played a major role in growth of these diabetic kidneys.

A working model is presented in Figure 6 to summarize the main findings: Glomerular hyperfiltration in early diabetes correlates with increased kidney size. Growth of the diabetic kidney requires increased activity of ODC and correlates with reduced activity of ADC, although the physiological importance of the latter is unproven. Also linked to growth of the diabetic kidney is a primary increase in proximal reabsorption which is sufficient to reduce the signal for TGF, thereby causing SNGFR to increase. These findings favor growth of the proximal tubule as a cause, rather than a consequence, of diabetic hyperfiltration.

Acknowledgments

This work was performed with funds from the US Department of Health and Human Services, National Institute of Diabetes and Digestive and Kidney Diseases (DK56248) and from the Department of Veterans Affairs Research Service. V. Vallon was supported by grants provided by the Deutsche Forschungsgemeinschaft (Va 118/3-2) and the Interdisciplinary Clinical Research Center (IKFZ) Tübingen (BMF 01KS9602).