Abstract. A specific anatomical lesion sharply localized to the cells of the medullary thick ascending limbs (mTAL) and characterized by mitochondrial swelling progressing to nuclear pyknosis and cell death is elicited reproducibly in isolated rat kidneys perfused for 15 or 90 min with cell-free albumin-Ringer's medium gassed with 5% CO₂, 95% O₂ (O₂ content, 1.5 vol/100 ml). The lesion, involving about half of mTALs, appears first in mTALs removed from vascular bundles and near the inner medulla, areas most likely to be anoxic. Hypoxic perfusion (O₂ content 0.12 vol/100 ml) exaggerates the lesion, wiping out gradations of damage and extending it to all mTALs. O₂-enriched perfusions using rat erythrocytes (O₂ content 7.1 vol/100 ml) completely eliminates the lesion (unless gassed with carbon monoxide). Similarly, supplementation of the perfusion medium with a purified hemoglobin (O₂ content 5.8 vol/100 ml) prevents mTAL injury. Perfusion with a fluorinated hydrocarbon blood substitute, Oxyferol (O₂ content 4.3 vol/100 ml) also attenuates the lesion.

These findings suggest that the mTAL is exquisitely susceptible to anoxic damage because of low O₂ supply imposed by the medullary vascular system and the high rate of metabolism mandated by active reabsorption of sodium chloride. The vulnerability of the mTAL to anoxic injury could play a key role in the pathogenesis of ischemic renal injury.

Introduction

During isolated perfusion of the rat kidney, a specific lesion is consistently observed, occurring within 15 min, confined to the medullary thick ascending limb (mTAL) of Henle's loop, which progresses from mitochondrial swelling to nuclear pyknosis and complete cellular disruption (1). Failure of urine-concentrating ability, defective distal sodium reabsorption with reduced glomerular filtration rate (GFR) and negative tubuloglomerular feedback, and increased potassium excretion all appear to be functional consequences of this injury (2). The addition of amino acids to the perfusate has been shown to attenuate the anatomical damage and to improve kidney function (2), but the basis for the mTAL lesion has remained obscure.

Recent observations by Epstein et al. (3) using organ spectrophotometry in whole isolated perfused kidneys, have indicated that a significant fraction (20–40%) of cytochrome a,a₃ (cytochrome c oxidase) as sensed by this technique is reduced but becomes more oxidized when mTAL transport is decreased by furosemide (3). Cytochrome a,a₃ is normally 95–98% oxidized in the presence of oxygen at partial pressures that fulfill the needs of oxidative phosphorylation (4). The authors concluded that substantial portions of the renal medulla may be operating on the verge of anoxia and therefore predisposed to anoxic damage. The present work, designed to test the possible relationship between the mTAL lesion and abnormal O₂ delivery, confirms this prediction and shows that anoxia is the basis for mTAL injury in the isolated perfused kidney.

Methods

Male Sprague-Dawley rats, weighing 370–470 g, fed on Purina rat chow (Ralston Purina Co., Chicago, IL) and allowed free access to water, were used for all experiments. Perfusion of the right kidney was performed according to the technique described by Ross et al. (5). Regular perfusion medium, as used in control rats, consisted of a Krebs-Ringer's-Henseleit solution with bovine serum albumin at a concentration of 6.7 g/100 ml and glucose at 5 mM, gassed with 5% CO₂, 95% O₂.

1. Abbreviations used in this paper: GFR, glomerular filtration rate; mTAL, medullary thick ascending limb.
Experimental groups

A. CONTROL GROUPS (n = 22). Kidneys were perfused with regular perfusion medium for either 15 (n = 8) or 90 min (n = 7). Control kidneys were fixed in situ to document normal mTAL morphology (n = 7).

B. HYPOXIC PERFUSIONS (n = 15). Kidneys were perfused with the same perfusion medium as control rats but gassed with 5% CO₂, 95% N₂. The histology was observed after 15 (n = 8) or 90 min (n = 7) of perfusion.

15 min of perfusion are required in this model to allow hemodynamic parameters to stabilize and insulin to equilibrate. Therefore, the 15-min perfusion experiments were not used for the study of kidney function, but for the morphologic study of the earliest phase of the mTAL lesion and its time sequence under different conditions of regular and low oxygenation.

C. OXYGEN-ENRICHED PERFUSIONS (n = 22). (These were 90-min perfusions, gassed with 95% O₂, 5% CO₂, unless specified otherwise.)

I. Erythrocytes (n = 10). 20–25 ml of fresh blood from two heparinized donor rats was added to a regular perfusion medium within 1 min from the start of the perfusion at a final hematocrit of 13.6±0.4. No filters were used for these experiments. A subset of two kidneys were perfused under the same conditions but gassed with 5% CO₂, 5% CO₂, and 90% N₂ to reduce O₂ carrying capacity back to low levels.

II. Hemoglobin (n = 8). A purified, stable, unmodified hemoglobin solution (P₅₀ = 16 Torr) supplied by Dr. R. Bolin (Department of the Army, Presidio of San Francisco, CA) was dialyzed against Krebs-Ringer's-Henseleit solution, concentrated by ultrafiltration, filtered, and added to the regular perfusion medium within 1 min from the start of the perfusion, at a final concentration of 3–5 g/100 ml.

III. Oxypherol (n = 4). A fluorinated carbon (FC-43 emulsion, Oxypherol, Alpha-Therapeutic, Los Angeles, CA) was used as a blood substitute instead of the regular perfusion medium.

Monitoring of oxygenation

In all experimental groups, the PO₂ of the perfusate was monitored using a blood gas monitor (PHM 73, Radiometer, Copenhagen). In groups A and B, the O₂ content was calculated using the constant of solubility of O₂ in plasma, and confirmed by direct measurement in 10 randomly selected samples. In group C the O₂ content was measured directly by volumetry using a Lex O₂ can (Lexington Instrument Corp., Waltham, MA). As shown in Fig. 1 there was good concordance between the expected O₂ contents and their actual measurements in the different groups.

Morphologic techniques

All kidneys of each control and experimental group were studied as separate groups for morphology. A three-way stopcock was incorporated into the circuit 5 cm from the arterial cannula to allow perfusion with the fixation solution, at the same pressure applied during the functional study, for an additional 5–8 min. The fixative solution contained 1.25% glutaraldehyde in 0.1 M phosphate buffer (pH 7.4). A 3-mm section of the outer medulla was removed, postfixed in 2% OsO₄, dehydrated and embedded in an araldite-epon 812 mixture. Large 1 μm sections were cut to include the entire length of the mTAL. Selected blocks were examined by electron microscopy. In two additional kidneys perfused under regular conditions, special transverse sections (perpendicular to the axis of the vasa recta) were obtained to study the relationship of the mTAL lesion to vascular bundles.

The histological evaluation was completed by one of us in a “blinded” fashion; that is, without knowledge of experimental conditions. Three zones of the inner stripe were analyzed: upper third: all mTALs intersecting a line immediately adjacent to outer stripe (within ~0.2 mm); middle third: all mTALs intersecting a line drawn midway between the borders of the inner stripe; and lower third: all mTALs intersecting a line immediately adjacent to the inner medulla (within ~0.2 mm).

These points were chosen for analysis because they provided areas in which topographical landmarks were easily ascertained. A percentage score was used to indicate the fraction of tubules involved with minimal to mild (chromatin margination, minor degrees of mitochondrial swelling), moderate (blatant mitochondrial swelling with limited nuclear pyknosis), or severe (blatant mitochondrial swelling with extensive nuclear pyknosis and cell fragmentation) changes. Between 68 and 272 tubules (mean 137) were evaluated per kidney in a total of 61 kidneys.

The data are presented as mean±SEM. Statistical analysis was by the t test, unless specified otherwise.

Results

Renal functions in the different groups

The functional parameters are summarized in Table I. In group B (hypoxic perfusion) a marked decrease in oxygen content was associated with low GFR and reduced fractional sodium reabsorption. The oxygen content of the perfusate was substantially increased in all C subgroups and most pronounced with erythrocytes (C₅). This group demonstrated improved conservation of sodium and potassium, as compared with perfusates containing albumin alone (group A).

The rate of flow of perfusate to the kidney was significantly lower (and calculated renal resistance correspondingly higher) in the experiments of group C than in the control group (group A).

Morphological observations

Fig. 2 illustrates the injury to the mTAL as consistently seen in isolated kidneys perfused with regular medium (Fig. 2 B–
Table I. Oxygenation and Function of the Kidneys in the Different Groups

<table>
<thead>
<tr>
<th></th>
<th>Arterial PO2</th>
<th>Arterial O2 content</th>
<th>GFR</th>
<th>TR, m</th>
<th>FEK</th>
<th>Renal perfusion flow</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mmHg</td>
<td>vol/100 ml</td>
<td>ml/min</td>
<td>%</td>
<td>%</td>
<td>m/min</td>
</tr>
<tr>
<td>A (controls) (n = 7)</td>
<td>495±12</td>
<td>1.5±0.04</td>
<td>0.52±0.06</td>
<td>93.9±1.3</td>
<td>58.2±11.5</td>
<td>45.7±2.6</td>
</tr>
<tr>
<td>B (low oxygen perfusions) (n = 7)</td>
<td>38±1*</td>
<td>0.12±0.004*</td>
<td>0.12±0.02‡</td>
<td>45.7±1.2*</td>
<td>76.9±4.1</td>
<td>37.6±1.9</td>
</tr>
<tr>
<td>C1 (erythrocytes) (n = 8)</td>
<td>463±18</td>
<td>7.1±0.3*</td>
<td>0.54±0.1</td>
<td>97.4±0.5§</td>
<td>26.9±2.9§</td>
<td>10.5±2.0*</td>
</tr>
<tr>
<td>C1 (erythrocytes with 5% CO) (n = 2)</td>
<td>36.5</td>
<td>0.11</td>
<td>0.17</td>
<td>44.5</td>
<td>72.3</td>
<td>10.0</td>
</tr>
<tr>
<td>CII (hemoglobin) (n = 8)</td>
<td>460±10</td>
<td>5.8±0.3*</td>
<td>0.76±0.2</td>
<td>96.5±0.4</td>
<td>22.6±3.6§</td>
<td>24.4±1.4*</td>
</tr>
<tr>
<td>CI (Oxypherol) (n = 4)</td>
<td>513±8</td>
<td>4.3±0.1*</td>
<td>0.25±0.1</td>
<td>97.9±0.8</td>
<td>10.6±4.5§</td>
<td>13.4±1.4*</td>
</tr>
</tbody>
</table>

Data are given at 60 min. TRNA, tubular reabsorption of sodium; FEK, fractional excretion of potassium.

* P < 0.001, vs. group A.
‡ P < 0.005, vs. group A.
§ P < 0.02, vs. group A.

E), as compared with kidneys fixed in situ (Fig. 2 A, 2 E inset). As damage proceeds, nuclei become more obvious and chromatin margination occurs. Later, mitochondrial swelling appears, and finally the nucleus condenses (pyknosis) and the luminal membrane becomes fragmented. These changes correspond to the well-described sequence that is characteristic of ischemic cell injury (6). They can all be perceived by light microscopy in 1-μm sections, and are easily apparent in electron microphotographs. When limited detail was perceived in the cell in such sections (nuclei and mitochondria not easily visible, and the cell border was intact; e.g., Figs. 4, 7A, electron microscopy invariably revealed minimal or no damage.

Group A: gradients of mTAL damage

Evaluation of the severity of the lesion in the control group at both 15 and 90 min showed that it was distributed along two types of anatomical gradients, corresponding to the expected gradients of oxygen tension in the renal medulla.

Axial gradient. The most severe and consistent damage was located in the deepest portions of the mTAL, close to the inner medulla. The least severe damage, frequently absent, was found in the most superficial third of the mTAL, near its transition to the cortical thick ascending limb (Fig. 3). The damage increased mostly in its severity from 15 to 90 min (Fig. 3).

Transverse gradient and polarity. The lesion was consistently more advanced in the mTALs located most distant from the vasa recta bundles. Indeed, epithelium immediately adjacent to the vascular bundle was frequently preserved, while opposed cells in the same tubules were markedly affected (Fig. 4). These results suggest that proximity to oxygen supply influences the development of the lesion.

Group B: effects of low oxygen perfusion

Low oxygen perfusion greatly enhanced the extent and severity of the mTAL lesion. Most mTALs in area B and C were affected by the injury, and showed moderate to severe damage both at 15 and 90 min (Figs. 3 and 5). Thus, hypoxic perfusion both exacerbated the damage and markedly attenuated its gradations (Fig. 3).

Group C: effects of O2 enriched perfusion

In all perfusions supplemented with erythrocytes the lesion was conspicuously absent (Fig. 6). In two kidneys gassed with carbon monoxide, O2 content was <1 vol/100 ml and the lesion reappeared, despite the presence of erythrocytes (Fig. 6). During perfusion with hemoglobin the mTAL lesion was again prevented (Figs. 7 and 8). With Oxypherol the injury was remarkably attenuated but still present (Figs. 7 and 8).

The relationship between the extent and severity of the mTAL lesion and the O2 content of the perfusion is summarized in Fig. 8. The typical lesion is found during regular perfusion at an O2 content ~1.5 vol/100 ml. A 10-fold decrease in the O2 content (in the low O2 perfusions) transforms the lesion to extensive and almost complete destruction of the mTAL. Conversely, increasing the O2 content to 4–5 vol/100 ml with Oxypherol affords significant protection, while the lesion is no longer detected above an O2 content of 6–7 vol/100 ml, with hemoglobin or erythrocytes.

Discussion

Leichtweiss et al. (7) have previously shown that the tissue partial pressure of O2, as measured by platinum microelectrodes inserted into the rat kidney from cortex to medulla, drops steeply in the cortico-medullary junction to values clustered around 10 mmHg in the medulla, both in vivo and during isolated perfusion (calculating from the depth of electrode penetration in Leichtweiss's studies, this PO2 was probably recorded at the level of the inner stripe of the outer medulla). It has been proposed (8) that the inner medulla functions adequately in this hypoxic environment because of its reliance on both aerobic and anaerobic glycolysis. However, recently, the concept of cellular hypoxia of the renal outer medulla has been suggested, using direct measurement.
Figure 2. A consistent sequence of cellular injury as seen during isolated kidney perfusion can be readily recognized in 1-μm plastic sections. In control material (A) (kidneys perfused with fixative in situ), mTAL mitochondria can be defined as rod-shaped elements and the normal nuclear chromatin distribution can be appreciated. In early phases of injury, chromatin margination occurs that later is associated with mitochondrial swelling. In B, earlier phases of mitochondrial swelling can be perceived especially in the tangential section of the mTAL (upper). Mitochondrial swelling is more evident in C (right) where chromatin margination is also marked. mTAL epithelium in C (extreme left) has a mild degree of chromatin margination associated with minimal mitochondrial swelling. The epithelium of the TAL in D shows the final stages of cell injury. The cells are fragmented and luminal membranes cannot be defined; nuclear pyknosis and mitochondrial swelling are present. Electron microscopy of such tubules as noted in (E) reveal nuclear pyknosis (arrow), marked mitochondrial swelling, and loss of luminal and basal membranes. In sharp contrast, elements contained within the interstitium (I) have relatively intact nuclear and membrane components. The inset shows, by comparison, an electron micrograph of a normal kidney fixed in situ. × 640 (A–D), × 3,000 (E and inset).

by optical spectroscopy of the redox state of cytochrome a,a3, the terminal electron carrier of the mitochondrial chain, which transports electrons directly to oxygen. The oxidative state of this enzyme in rat kidneys was estimated as at least 20% reduced in vivo (9) and 30–40% reduced during isolated perfusion (3), indicating that despite high Po2 in artery and vein, substantial portions of the kidney may exist in an hypoxic milieu. Since inhibitors of cortical metabolism did not affect the redox state of the cytochrome, whereas furosemide or bumetanide produced a significant further oxidation of cytochrome a,a3, it appeared that the mTAL was an important site of reduced cytochrome oxidase, so that cells of the mTAL might operate on the brink
of anoxia during isolated perfusion or eventually in vivo during ischemia.

The present study demonstrates that the selective and reproducible damage to the mTAL in isolated rat kidney perfusion (1) is a lesion of anoxia. The observation that the damage is most severe in areas removed from vascular bundles and near the inner medulla, areas likely to suffer from poor O\textsubscript{2} supply, itself suggests O\textsubscript{2} deficiency as a key factor. (Since the mTAL of the short-looped nephrons are most removed from the vascular bundle (10), these nephrons are singularly vulnerable to limited availability of O\textsubscript{2}.) Indeed, the demonstration of these gradients of damage provides a remarkable morphological counterpart to the axial (cortico-medullary) gradient of tissue O\textsubscript{2} shown by Leichtweiss et al. (7) and to the transverse (vasculo-cellular) gradient of PO\textsubscript{2} suggested by Epstein et al. (3).

The effacement of these gradients, and their replacement by a lesion involving all mTALs in their entire length as early as 15 min of hypoxic perfusion support the concept that the regular mTAL lesion seen consistently during perfusion with albumin is a lesion of anoxia, preferentially affecting cells that have high oxygen demand and are removed from oxygen supply. The high selectivity of this injury is emphasized by the findings

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Figure 3. (Upper panel) Axial gradient of mTAL damage at 15 and 90 min of regular perfusion. The extent of mTAL damage in the inner stripe of the outer medulla was evaluated in three regions: outer (A), mid (B), and inner (C) (see Methods). Increasing severity of damage is noted progressing from the most superficial aspect of the inner stripe to the zone located adjacent to the inner medulla. Increasing severity of damage is also noted from 15 (mostly moderate) to 90 min (mostly severe). (Lower panel) Attenuated axial gradient of mTAL damage at 15 and 90 min of low oxygen perfusion. Striking extension of the lesion to all nephrons in areas B and C and most nephrons of area A is noted. Increasing severity of damage is mainly seen in area A at 15 and 90 min, as compared with regular perfusions. n represents total number of tubules evaluated in each group; * refers to comparison of adjacent columns; P < 0.05; † refers to comparison of respective columns at 90 vs. 15 min, P < 0.001; ‡ and ††, refer to comparison of respective columns of low oxygen perfusions vs regular perfusions, P < 0.001 and <0.05, respectively. P was calculated by Mann-Whitney U test.

Figure 4. Transverse gradient of mTAL damage (as seen at 90 min). In A, low power (inset) reveals the tubular-vascular organization of the inner stripe in horizontal section. The collecting ducts (+) are centrally located between vascular bundles which are located in the upper right and lower left. The higher power (A), with vascular bundles at lateral aspects shows that the greatest mTAL changes (bars) generally occur near collecting ducts (C) and in zones most distant from vascular bundles. In B, the low power sagittal section (inset) also shows maintenance of mTAL epithelia adjacent to the vascular bundles (arrows). Higher power (B) of zone between arrows reveals that the polarity of the mTAL damage is such that epithelium (arrows) adjacent to the vascular bundle is preserved while opposed cells within the same tubule are severely injured. The most severe mTAL damage again is related to collecting ducts (C). A and B, × 250; insets, × 65.
at 90 min of hypoxic perfusion, where extensive and diffuse tubular destruction was still basically confined to the mTAL.

Measures to increase medullary O₂ delivery resulted in a marked improvement related to O₂ content of the perfusate. Supplementation of the albumin perfusate with fresh rat blood to a hematocrit of 14% resulted in complete preservation of the mTAL integrity. To minimize hemolysis, erythrocytes were not separated but supplementation with plasma per se did not provide any diminution of the mTAL lesion (data not shown). Moreover, carbon monoxide, which markedly reduced O₂ binding to hemoglobin, reversed the protection afforded during erythrocyte perfusions, underlining the dependence of this protection on improved oxygen-carrying capacity. Additional verification of this hypothesis was afforded by the hemoglobin experiments. Nearly complete protection was provided by supplementing the albumin perfusate with a purified hemoglobin solution possessing improved oxygen-carrying capacity. Finally, increased oxygen dissolution in the fluorinated carbon blood substitute FC 43 emulsion also led to significant protection of the mTAL, again suggesting improved oxygen delivery to the mTAL as the key factor in its protection.
Increased conservation of potassium seen in the three types of \( O_2 \)-enriched perfusions (Table I) may be related to a diminution of solute delivery to the distal tubule during isolated perfusion (11), owing to improved reabsorption of NaCl in the mTAL. Fractional reabsorption of sodium did in fact tend to be higher in the high oxygen perfusions of group C, significantly so in Group C. This tendency may well have been opposed by the obligatory urinary excretion of hemoglobin dimers in group C\(_{\text{II}}\) and of perfluorotributylamine\(^2\) in group C\(_{\text{III}}\). Similar improvements in sodium conservation have already been reported with erythrocytes or fluorinated carbon perfusions (12, 13). Normalization of the solute reabsorption is not expected when glucose is used as the only substrate (2). In several experiments where amino acids (2) were included in addition to erythrocytes or hemoglobin, a fractional reabsorption of 99% or higher was observed. Preliminary observations suggest that preserved mTAL morphology is also associated with improved concentrating ability (unpublished observations).

\(^2\) Perfluorotributylamine has a mol wt of 660, is present in FC 43 at a concentration of 300 mM, and may dissociate from emulsion particles to produce an osmotic diuresis.

The steep cortico-medullary gradient of \( O_2 \) in kidney tissue is the result of the medullary vascular structure and the conditions imposed by the counter-current exchange system (14). The hairpin conformation of the vasa-recta prevents the dissipation of a cortico-medullary gradient of osmolality, but at the cost of counter-current diffusion of \( O_2 \) between arterial and venous limbs. The high \( O_2 \) consumption of this zone keeps medullary \( O_2 \) low and almost undisturbed by axial flow. During isolated perfusion, the high arterial \( O_2 \) favors arteriovenous diffusion, leaving medullary \( O_2 \) in the same low range as found in vivo (7) (~10 mmHg). Blood or perfusate flowing through the vasa recta of the outer medulla, therefore, will have its \( O_2 \) rapidly lowered to this range. At this \( O_2 \) tension, because of the sigmoid shape of the \( O_2 \) dissociation curve, blood still has a significant \( O_2 \) content, whereas the amount of \( O_2 \) dissolved in an albumin solution is extremely low, as illustrated in Table II. As a result, oxygen delivery to the medulla is inevitably compromised during isolated perfusion without hemoglobin. The elevated rate of perfusion and high content of dissolved \( O_2 \) in the perfusate may serve to maintain delivery of oxygen to cortical tissues close to that in vivo, but such compensation cannot be achieved in the medulla, which consequently suffers anoxic damage.

It is conceivable that the marked increase in renal perfusion flow seen in the isolated kidney perfused with albumin is, at

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Figure 7. If hemoglobin is included in the perfusate, the mTAL is preserved (A and B). Oxypherol (C) also attenuates the injury, but mild to moderate damage can still be recognized in some mTALs (right). \( \times 400, \times 640; \times 400 \).

Figure 8. Relationship between degree of mTAL damage and oxygenation of perfusion medium. The quantitation of damage is for the whole inner stripe of the outer medulla (areas A, B, and C combined). *, refers to extent (moderate or severe); †, to severity of damage compared with regular perfusions; both to \( P < 0.001 \). **, \( P < 0.05 \).
Table II. Estimated Oxygen Delivery to Whole Rat Kidney and to Outer Medulla In Vivo and during Isolated Perfusion

<table>
<thead>
<tr>
<th></th>
<th>Arterial O2 content</th>
<th>Flow</th>
<th>O2 delivery</th>
<th>mTAL lesion</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mmHg</td>
<td>%</td>
<td>ml/min/g</td>
<td>ml/min/g</td>
</tr>
</tbody>
</table>

Whole kidney
- Kidney in vivo: 70–90* 18–19* 3–5‡ 0.5–0.9 Never seen§
- Isolated perfused kidney (albumin): 500–600 1.2–1.8 24–28 0.3–0.5 Always seen

Outer medulla
- Kidney in vivo: 25† 4** 2† 0.08 Never seen
- Isolated perfused kidney (albumin): 25‡ 0.06 13‡‡ 0.008 Always seen

* As determined in several control rats under anesthesia.
‡ See reference 10.
§ Normal mTAL morphology was observed in seven control kidneys fixed in situ by glutaraldehyde perfusion.
† From references 1 and 2, and present work.
* The upper range of tissue O2 in the outer medulla as measured by Leichtweiss et al. (7) was chosen as an estimate of arterial O2 in this region.
** Estimate from reference 15.
‡‡ Cortical and medullary flow are thought to be increased in roughly equal proportions during isolated perfusion (16, 17.

least in part, a physiological response to the low O2 supply. This vasodilation was indeed remarkably blunted in all O2-enriched perfusions (Table I) where normal renal innervation and renin substrate supply were absent. The effects of erythrocytes on perfusate viscosity are minimal below a hematocrit of 30 as measured either in vivo or in vitro (18) and are unlikely to be the explanation of these striking changes in flow. In other experimental models, acute anemia increases renal blood flow (19), whereas hyperbaric oxygenation decreases renal blood flow (20), supporting the concept that O2 availability might be a factor in the regulation of renal blood flow, as shown in other tissues (21). Prostaglandins could theoretically mediate this hypoxic renal vasodilation, but indomethacin does not prevent its occurrence during the first hour of perfusion (G. Kershaw and J. Stoff, unpublished observations), and their role remains to be proven. This singular sensitivity of the medullary portion of the kidney to O2 content, at constant arterial Po2, might also make it an ideal site for the regulation of erythropoietin synthesis.

We have recently shown that decreasing the work of transport and thereby reducing the demand for oxygen by mTAL cells can prevent the mTAL lesion. Perfusion with furosemide or ouabain or with hyperoncotic albumin medium (to halt glomerular filtration) consistently reduce or abolish the anoxic damage to the mTAL (22). Similarly, the addition of amino acids to the perfusate attenuates the mTAL lesion [2], probably by improving proximal tubular reabsorption ["unloading" the mTAL] and by increasing renal perfusion flow [reference 2, and unpublished observations] with the net result of a combined decrease in O2 consumption and increase in O2 delivery in the mTAL. Some of these experiments appear to be examples in the kidney of the general principle, so familiar in the pathophysiology of the heart, that cellular work accelerates anoxic injury and that an imbalance between oxygen supply and demand is a prelude to cell damage. It is not surprising that the cells of the mTAL, by virtue of their high rate of metabolism (23) and their special location with limited O2 supply, are so exquisitely susceptible to anoxia. They might therefore be expected to be especially vulnerable in vivo to ischemia as well.

Several reports have, in fact, indicated a defect in TAL function in experimental models of ischemic renal failure (24–26). Structural changes including mitochondrial swelling and nuclear pyknosis have been identified in the mTALs of kidneys examined from rats submitted to hemorrhagic hypotension (27) or to glycerol injections (28). Decreased concentrating ability (one of the most consistent defects in human acute tubular necrosis), polyuria (in nonoliguric renal failure), and activation of tubuloglomerular feedback (29) could all be direct consequences of a selective mTAL ischemia. Focal or diffuse TAL necrosis in acute renal failure has been reported by several observers (30–34). The protection afforded by the prophylactic administration of furosemide in several models of ischemic acute tubular necrosis has never been clearly explained (35) but evokes an intriguing parallel to the prevention of mTAL anoxic damage during isolated perfusion by decreasing O2 demand by cells. Indeed, the redistribution of cortical flow in favor of the medulla, an early event in hypotension (36), and the profound disproportionate fall in GFR in acute renal failure might be viewed, at least in part, as a regulatory mechanism developed to protect...
the mTAL from the dangerous and precarious equilibrium between \( O_2 \) demand and supply within the outer medulla of the kidney.

The selective vulnerability of mTAL cells to anoxic damage, so clearly demonstrable in isolated perfused kidneys, might influence their fragility in other experimental models (isolated TAL tubules, isolated TAL cells) where careful oxygenation may be crucial. Finally, the specific susceptibility of the mTAL to anoxia may play a key role in a variety of disease states characterized by renal medullary ischemia (37).

References