

1	Absence of renal hypoxia in the subacute phase of severe renal ischemia
2	reperfusion injury
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26 Abstract

27 Tissue hypoxia has been proposed as an important event in renal ischemia reperfusion injury (IRI), particularly during the period of ischemia and in the immediate hours 28 29 following reperfusion. However, little is known about renal oxygenation during the subacute 30 phase of IRI. We employed four different methods to assess the temporal and spatial changes 31 in tissue oxygenation during the subacute phase (24 h and 5 days after reperfusion) of a severe 32 form of renal IRI in rats. We hypothesized that the kidney is hypoxic 24 h and 5 days after an 33 hour of bilateral renal ischemia, driven by a disturbed balance between renal oxygen delivery 34 (DO_2) and oxygen consumption (VO_2) . Renal DO_2 was not significantly reduced in the 35 subacute phase of IRI. In contrast, renal VO2 was 55% less 24 h, and 49% less 5 days after 36 reperfusion than after sham-ischemia. Inner medullary tissue PO₂, measured by 37 radiotelemetry was $25 \pm 12\%$ greater 24 h after ischemia than after sham-ischemia. By 5 days 38 after reperfusion, tissue PO₂ was similar to that in rats subjected to sham-ischemia. Tissue 39 PO₂ measured by Clark electrode was consistently greater 24 h, but not 5 days, after ischemia 40 after sham-ischemia. Cellular hypoxia, assessed by pimonidazole adduct than 41 immunohistochemistry, was largely absent at both time-points and tissue levels of hypoxia 42 inducible factors were downregulated following renal ischemia. Thus, in this model of severe 43 IRI, tissue hypoxia does not appear to be an obligatory event during the subacute phase, likely 44 due to the markedly reduced oxygen consumption.

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48 Introduction

Acute kidney injury (AKI) is a major cause of death and disability globally and places a major acute burden on healthcare systems (26). It also renders patients more susceptible to later development of chronic kidney disease (CKD) (2). For example, a diagnosis of AKI was found to be associated with an 8.8 fold excess risk of later development of CKD (8). Furthermore, the risk of later development of CKD increases with the severity of AKI (8). Tissue hypoxia has been proposed as an important driver in the pathogenesis of both AKI and CKD, although this proposition remains to be definitively tested (33).

Ischemia reperfusion injury (IRI) sustained from medical interventions often arises 56 57 from the obligatory need to restrict or completely prevent blood flow to the kidney, resulting 58 in a period of severe hypoxia or complete anoxia (15). Cellular damage such as acute tubular 59 necrosis and tubular apoptosis is evident during the reperfusion period and is likely driven in 60 part by the presence of tissue hypoxia during the period of ischemia. In experimental IRI, 61 cortical (27, 28, 41) and medullary (27, 28, 34) tissue hypoxia has also been observed during 62 the first few hours of reperfusion after complete renal ischemia. Importantly, the kidney was 63 observed to be hypoxic even with some level of, albeit incomplete, structural and functional 64 recovery (3, 4). However, there are little available data regarding renal tissue oxygenation 65 beyond the first few hours of reperfusion during the extension and recovery phase of IRI. This 66 information is required if we are to understand the role of tissue hypoxia in the natural history 67 of AKI, either as it progresses to end-stage renal disease or renal function recovers but the 68 risk of later CKD is increased.

The chief aim of the current study was to assess the time-course of changes in, and the spatial distribution of, tissue oxygen tension (PO₂) during the subacute phase of severe IRI (the first 5 days of reperfusion after 60 min of bilateral renal ischemia). We chose severe IRI in an attempt to model the clinical situation of severe AKI leading to end-stage renal disease, 73 cognizant of the possibility that renal oxygenation in this scenario might differ considerably 74 from that in milder forms of renal IRI. We tested the hypothesis that renal tissue is hypoxic 75 during the subacute phase of IRI. Four approaches were used for assessment of renal tissue 76 oxygenation, each with varying temporal and spatial resolution. Radiotelemetry was used to examine the time-course of changes in inner medullary tissue PO_2 in freely-moving rats (22, 77 78 23). Clark-type electrodes were used to characterize the spatial variations in renal tissue PO_2 79 in the renal cortex and medulla of anesthetized rats at both 24 h and 5 days after reperfusion. 80 This experiment also provided an opportunity to determine the contribution of changes in 81 renal oxygen delivery (DO_2) and oxygen consumption (VO_2) to alterations in renal tissue PO_2 82 24 h and 5 days after reperfusion. Pimonidazole adduct immunohistochemistry was used to 83 characterize the spatial distribution of cellular hypoxia 24 h and 5 days after reperfusion. We 84 also measured the expression of hypoxia-inducible factors (HIF-1 α and HIF-2 α) and some of 85 their downstream gene targets.

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87 Methods

88 Experimental Animals

Ten to twelve week old male, Sprague-Dawley rats (n=70) were obtained from the Animal Resources Centre (Perth, Western Australia). They were housed in a room maintained at 21–23 °C with a 12 h light/dark cycle. The rats were allowed free access to water and standard laboratory rat chow. All procedures were approved in advance by the Animal Ethics Committee of the School of Biomedical Sciences, Monash University as being in accordance with the Australian Code of Practice for the Care and Use of Animals for Scientific Purposes.

95

96 Induction of bilateral renal ischemia

Rats were anesthetized with isoflurane (IsoFlo[™], 05260-05, Abbott Laboratories,
USA), using a vaporizer and maintained at 2.5 – 3.0% v/v. A midline incision was made to

99 expose the left and right renal arteries. To induce bilateral renal ischemia (n = 36), blood flow 100 to both kidneys was prevented by the application of microvascular clamps (00398, S&T AG, 101 Switzerland) placed on both the left and right renal arteries and veins. Complete ischemia was 102 confirmed by observing the blanching of the kidneys. After an hour, the microvascular clamps 103 were removed, so blood flow to both kidneys was restored. Wounds were closed in layers 104 with sutures and each rat was then allowed to recover from the surgery on a heated pad for an 105 hour. A separate cohort of rats (n = 34) underwent the same procedure with the exception of 106 the application of the microvascular clamps and so served as controls (sham-ischemia). Rats 107 received subcutaneous injections of an analgesic (carprofen, 1.25 mg, Pfizer, Australia) for 108 two consecutive days following recovery from surgery.

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110 Protocol 1: Temporal changes in renal tissue oxygenation following renal ischemia

111 We employed a radiotelemetric method (22, 23) to characterize the temporal profile 112 of changes in renal tissue PO₂ after renal ischemia and reperfusion. Briefly, the oxygen 113 telemeter was implanted under isoflurane anesthesia so that the tip of the oxygen-sensing 114 carbon paste electrode was in the inner medulla of the left kidney (5 mm below the renal 115 capsule). One week after implantation of the telemetric probe, the rats underwent a second 116 surgical procedure for the induction of either bilateral renal ischemia (n = 7, body weight = 117 501 ± 20 g) or sham-ischemia (n = 5, body weight = 491 \pm 21 g). Renal tissue PO₂ was 118 recorded continuously for 1 day before and for 5 days after recovery from surgery. Rats 119 received subcutaneous injections of an analgesic (carprofen, 1.25 mg, Pfizer, Australia) prior 120 to laparotomy and for two consecutive days following recovery from surgery.

Measurements & Calculations: Current measured by the telemeters was filtered with a 25 Hz low-pass filter and artifactual measurements were removed when the 1st order derivative of the measured current exceeded the threshold of 5-500 nA/s. The zero-offset current, acquired when the rat was killed at the end of the study via induction of cardiac arrest under anesthesia (22), was determined and subtracted. Data are presented as a percentage of the average valueon the day before surgery to induce ischemia or sham-ischemia.

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128 Protocol 2: Renal tissue oxygenation and its determinants after renal ischemia

Either 24 h or 5 days following recovery from renal ischemia or sham-ischemia, rats were anesthetized and prepared for the assessment of regional tissue PO_2 using a Clark electrode (50 µm tip, OX-50, Unisense, Denmark). We assessed i) cortical tissue PO_2 across a range of sites on the dorsal surface of the kidney and ii) a profile of tissue PO_2 with depth from the cortical surface. In this set of studies, we also determined the major determinants of tissue PO_2 , renal DO_2 and VO_2 .

135 Rats (n = 6-11 per group) were anesthetized with sodium thiobutabarbital (100) 136 mg/kg *i.p.*, Inactin; Sigma, St Louis, MO, USA). A tracheostomy was performed to facilitate 137 artificial ventilation with 40% inspired oxygen at a ventilation rate of 90-100 breaths/min and 138 a tidal volume of 3.5 ml (Ugo Basile, Model 7025, SDR Clinical Technology, NSW, 139 Australia) as previously described (1). The left carotid artery was catheterized to facilitate 140 arterial blood sampling and blood pressure measurement. The right jugular vein was 141 catheterized to facilitate infusion of maintenance fluid (154 mM NaCl) at a rate of 6 ml/h 142 during the period of surgical preparation. The bladder was catheterized, for collection of urine 143 from the left kidney, for assessment of renal function using standard clearance methods. The 144 degree of saturation of hemoglobin with oxygen was measured continuously using a sensor 145 placed on the foot (Mouse Ox, Starr Life Sciences, Oakmont, PA, USA).

The right renal artery and vein were ligated and a catheter was passed from the right renal vein through the vena cava and into the left renal vein for the sampling of renal venous blood. Total renal blood flow (RBF) was measured using a transit-time ultrasound flow probe (Type 0.7 VB, Transonic Systems Inc., NY, USA) placed around the left renal artery. Following completion of the surgical preparations, rats received bolus doses of [³H]-inulin (10 μCi in 50 μl, Perkin Elmer Australia, Melbourne, Australia) and pancuronium bromide (2
mg/kg, Astra Zeneca Pty Ltd, NSW, Australia) intravenously. A maintenance infusion of 2%
w/v bovine serum albumin (Sigma Aldrich, St Loius, MO, USA) in 154 mM sodium chloride
delivered 676 nCi/h [³H] inulin and 0.1 mg/kg/h pancuronium bromide through the jugular
vein at a rate of 2 ml/h. The infusion commenced once all surgical preparations were
completed and was maintained throughout the rest of the protocol.

157 After a 1 h equilibration period, a 0.5 ml sample of arterial blood was taken for blood 158 oximetry. The plasma component of the sample was later used for assessment of the concentrations of [³H]-inulin and sodium. A 0.1 ml sample of renal venous blood was also 159 160 collected for blood oximetry. Renal tissue PO2 was then assessed using a Clark electrode 161 attached to a micromanipulator. Two series of measurements were taken. In the first series, 162 the electrode was advanced 2 mm from the renal surface, into the cortex, at 6 randomly 163 chosen sites across the left kidney. The second series established a profile of tissue PO₂ with 164 depth below the cortical surface. The electrode was moved to the mid-point of the cortical 165 surface of the kidney and advanced into the kidney at 1 mm increments up to a depth of 10 166 mm from the renal surface as previously described (32). Once all measurements were taken, a 167 second set of blood samples, from the carotid artery and the renal vein, was taken as before. Urine made by the left kidney, during the period of measurement of tissue PO₂, was collected 168 169 for measurement of the concentrations of $[^{3}H]$ -inulin and sodium.

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Measurements & Calculations: Arterial pressure, heart rate (triggered by arterial pressure), RBF, core body and tissue temperature and renal tissue PO₂ measured by Clark electrode were digitized as previously described (32). Urinary and plasma concentrations of sodium were determined using ion-sensitive electrodes (EasyElectrolytes, Medica Corporation, Bedford USA). Glomerular filtration rate (GFR) was determined by the clearance of [³H]inulin. Blood chemistry was assessed using a point-of-care device (iSTAT®, CG8+ 177 cartridges, Abbott laboratories, Abbott Park, IL, USA). Arterial and venous blood oxygen178 content was calculated as previously described (1).

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180 Protocol 3: Cellular hypoxia and hypoxic signaling after renal ischemia

Either after 24 h or 5 days of recovery from bilateral renal ischemia or sham ischemia (n = 6 per group), rats were prepared for perfusion-fixation of the right kidney. In this set of studies, the chief aim was to assess cellular hypoxia using pimonidazole adduct immunohistochemistry. Pimonidazole chloride (HP1-1000Kit, Hydroxyprobe Inc., USA) was administered, at a dose of 60 mg/kg *i.p.* three hours before perfusion-fixation of the kidney.

186 Three hours after the injection of pimonidazole, rats were anesthetized with sodium 187 pentobarbital (60 mg/kg, i.p., Sigma Aldrich, MO, USA). The left carotid artery was 188 catheterized to facilitate arterial blood sampling. A midline incision was then made exposing 189 both kidneys and the bladder. A urine sample was taken by puncturing the bladder wall and 190 was frozen at -20 °C for later analysis. The left renal artery and vein were isolated and freed 191 from surrounding connective tissue and fat. Lidocaine (2% w/v; Xylocaine®, Astra Zeneca, 192 NSW, Australia) was applied onto both vessels to prevent spasm of the renal artery. Silk 193 ligatures (3/0 Dysilk, Dynek Pty Ltd, SA, Australia) were placed around the vena cava above 194 the level of the right kidney, around the left renal artery and vein and around the abdominal 195 aorta. An incision was made in the abdominal aorta below the level of the left kidney and a 196 polyurethane catheter connected to the perfusion apparatus was advanced into the aorta, 197 facing upstream, thereby facilitating retrograde perfusion. A 1 ml blood sample was taken 198 from the carotid artery for later analysis. The left renal artery and vein were then ligated and 199 the left kidney removed, decapsulated and snap frozen in liquid nitrogen for later analysis of 200 HIF-1 α and HIF-2 α protein and gene expression of HIF-1 α , HIF-2 α , VEGF- α and HO-1. 201 Prior to freezing, the left kidney was sectioned in the coronal plane into 4-5 slices of 202 approximately 1–2 mm thickness.

The ligatures surrounding the vena cava and abdominal aorta were tied off and the right kidney perfused with 100–150 ml of 4% w/v paraformaldehyde (PFA, paraformaldehyde powder, no. 158127, Sigma-Aldrich) at room temperature and a pressure of 150 mmHg. The inferior vena cava was incised to vent perfusate. The perfused kidney was removed, decapsulated and stored in 4% PFA for 48 h before it was processed for embedding and staining at the Monash Histology Platform.

Blood chemistry was assessed using a point-of-care device (iSTAT®, CHEM8+
cartridges, Abbott laboratories). Urinary albumin concentration was determined using direct
competitive enzyme linked immunosorbent assay (Nephrat II, NR 002, Exocell Inc., PA,
USA). Urinary creatinine concentration was determined using an assay based on Jaffe's
reaction of alkaline picrate solution with creatinine (Creatinine Companion, 1012 Strip Plate,
Exocell Inc.).

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216 *Quantification of fibrosis.* The right kidney was processed, embedded in paraffin and 217 sectioned at a thickness of 5 μm in the coronal plane. Collagen deposition was assessed by 218 staining with 1% w/v picrosirius red. The cortical, outer and inner medullary region of the 219 kidney in each section was identified using Aperio Imagescope (Leica Biosystems Imaging 220 Inc., Australia). The amount of collagen deposited was quantified as a percentage of the entire 221 area in each region.

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Pimonidazole adduct immunohistochemistry. Antigen retrieval was carried out by incubating the sections in citrate buffer (Target Retrieval Solution, DAKO, Australia) at 90 °C for 30 min. Sections were then washed in tris-buffered saline (154 mM NaCl) with Tween 20 (DAKO Australia) (TBST) once they had cooled to 80 °C. Excessive tissue peroxidase activity was then quenched using 0.03% v/v hydrogen peroxide containing sodium azide (DAKO, Denmark) for 10 min. Sections were then incubated in a protein block serum

229 (Protein Block Serum-free, DAKO, Australia) for 10 min, in order to remove non-specific 230 binding, and washed twice more in TBST. Sections were then treated with an affinity purified 231 polyclonal anti-pimonidazole antibody raised in the rabbit (1:200 dilution, PAb2627AP, 232 Hydroxyprobe Inc.) for 1 h at room temperature before incubation in goat anti-rabbit 233 secondary antibody conjugated with horseradish peroxidase (polyclonal goat EnVision, 234 DAKO, Denmark) for 30 min at room temperature. Sections were washed twice with TBST 235 before incubation with 3-diaminobenzidine (DAKO, Denmark) for 10 min and then 236 counterstained with haematoxylin (DAKO, Automations Hematoxylin, USA) before being 237 cover-slipped.

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239 Western blot analysis of HIF-1 α and HIF-2 α proteins. The snap frozen kidney was thawed 240 and the cortex, outer and inner medulla inclusive of the papilla were rapidly dissected. To 241 stop further enzymatic reactions, the tissue samples were placed in 8 µl per mg of radioimmuno-precipitation assay (RIPA) buffer (consisting of 50 mM Tris-Hcl, 150 mM 242 243 NaCl, 0.1% Triton X-100, 0.5% sodium deoxycholate, 0.1% sodium dodecylsulphate (SDS), 244 1 mM sodium orthovanadate, 1 mM NaF, 1:25 of 25x phosphatase inhibitor, 1:10 of 10x 245 phosphostop and 1:1000 dithiothreitol). The tissues were then homogenized at 14,000 RPM at 246 4 °C for 20 min and equal amounts of protein (30 µg, determined by a Bradford protein assay) 247 were loaded into a 7.5% pre-cast gel (7.5% Mini-Protean® TGX[™] Precast Protein Gels, 248 4561025, **Bio-Rad** Laboratories, USA) and fractionated electrophoretically in 249 Tris/Glycine/SDS running buffer at 300 V for 20 min. The fractionated protein in the gel was 250 then transferred onto a nitrocellulose membrane (Bio-Rad Laboratories). Non-specific binding 251 was blocked with 5% skim milk in TBST buffer. As the primary antibodies for HIF-1a 252 (NB100-479, Novus Biologicals, LLC, CO, USA) and HIF-2a (NB100-122, Novus 253 Biological LLC) are similar in molecular weight (115 and 118 kDa), we carried out the immunoblot analysis of each protein of interest on separate gels. The nitrocellulose 254

255 membranes were incubated overnight at 4 °C in the primary antibody (1:1000, raised in 256 rabbit) made up in a solution of 2.5% w/v bovine serum albumin. The membranes were then 257 incubated with 1:4000 secondary antibody (ECLTM anti-rabbit IgG, HRP-linked whole 258 antibody, GE Healthcare UK Limited, UK) and 1:15,000 conjugate (Precision Protein[™] 259 StrepTactin-HRP conjugate, Bio-Rad Laboratories) for an hour at room temperature. The 260 nitrocellulose membrane was developed using equal parts of ClarityTM Western Peroxide 261 Reagent (Bio-Rad Laboratories) and Clarity[™] Western Luminol/Enhancer Reagent (Bio-Rad 262 Laboratories) for 3 min before imaging. The intensity of the bands observed on the membrane 263 was quantified and corrected for variability in protein migration down the gel and for total 264 protein content loaded into the wells. Comparisons were made between treatment groups 265 across the two time points within each region (*i.e.* cortex, outer and inner medulla).

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267 *Quantitative real-time PCR*. The tissue samples were homogenized and total RNA was 268 isolated using the RNeasy Mini Kit (74104, Qiagen Inc., Australia). Pre-designed assays for 269 primers of the 18s house keeping gene (Rn03928990_g1), *HIF-1a* (Rn01472831_m1), *HIF-*270 2α (Rn00576515_m1), *VEGF-a* (Rn01511602_m1) and *HO-1* (Rn00561387_m1) genes were 271 obtained from ThermoFisher Scientific Inc. Real-time PCR was performed on ABI 7900 HT 272 (ThermoFisher Scientific Inc., Australia). Data were calculated by the 2^{- $\Delta\Delta$ Ct} method.

- 273
- 274 Statistical analysis

Statistical analyses were performed using the software package SYSTAT (Version 13, Systat Software, San Jose, CA). Two-sided $P \le 0.05$ was considered statistically significant. Normality was assessed using the Shapiro-Wilk test (40). Data that did not violate normality are presented as mean \pm standard error of the mean (SEM) while data that violated normality are presented as median (25th percentile, 75th percentile). Analysis of variance (ANOVA) was used to assess the independent effects of treatment and time and their interaction. For data the violated normality, an ANOVA on ranking (9) was performed instead. Dichotomous comparisons of continuous variables were made using Student's t-test for data that did not violate normality and the Mann-Whitney U-test was performed for data that violated normality. To protect against the risk of type I error arising from multiple comparisons, P-values were conservatively adjusted using the Dunn-Sidak procedure (30). Pvalues derived from within-subjects factors in repeated measures ANOVA were conservatively adjusted using the method of Greenhouse and Geisser (31).

288 Results

289 Protocol 1: Temporal changes in renal tissue oxygenation following renal ischemia

On the first day after reperfusion, inner medullary tissue PO_2 measured by telemetry was $25 \pm 12\%$ greater than its control level (Day -1) (Fig. 1). Tissue PO_2 then gradually fell to be close to its control level by the fifth day after reperfusion of the kidney. After shamischemia, inner medullary tissue PO_2 tended to gradually fall, so was $22 \pm 11\%$ less than its control level by day five after surgery.

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296 Protocol 2: Renal tissue oxygenation and its determinants after renal ischemia

Systemic parameters. Twenty-four hours after reperfusion, body weight did not differ significantly from that of rats that underwent sham-ischemia. By 5 days after renal ischemia, rats had lost 39.2 ± 6.1 g of their body weight. Left kidney weight 24 h after renal ischemia was similar to that after sham-ischemia. In contrast, left kidney weight was 56% greater 5 days following renal ischemia than after sham-ischemia (Table 1). Mean arterial pressure was similar in the two groups of rats at both 24 h and 5 days after surgery.

303 *Renal tissue oxygenation.* Tissue PO_2 in the renal cortex was highly heterogenous, both 24 h 304 and 5 days after either ischemia or sham-ischemia (Fig. 2A). Cortical PO_2 was, on average, 305 40% greater 24 h following renal ischemia than after sham-ischemia. By five days after renal 306 ischemia, cortical tissue PO_2 was 39% less than 24 h after ischemia and similar to that in rats subjected to sham-ischemia five days previously (Fig. 2B). Tissue PO_2 varied little with depth from the cortical surface. At 24 h after reperfusion, tissue PO_2 tended to be greater in rats subjected to ischemia than those subjected to sham-ischemia, the difference reaching statistical significance at depths of 5 mm (inner medulla), and 9 and 10 mm (cortex) (Fig. 2C). Five days after renal ischemia tissue PO_2 did not differ significantly from its level in rats subjected to sham-ischemia, at any depth below the cortical surface (Fig. 2D).

313 Renal hemodynamics and function. Renal blood flow was not significantly different in rats 314 subjected to ischemia, compared with rats subjected to sham-ischemia, both 24 h and 5 days 315 after surgery (Table 2). Twenty-four hours after ischemia, mean GFR (-99%), urine flow (-316 82%) and sodium excretion (-85%) were less than in after sham-ischemia (Table 2). 317 Fractional excretion of sodium did not differ significantly 24 h after ischemia compared to 318 sham-ischemia. By five days after ischemia, renal function was highly variable between rats, 319 with some rats having recovered relatively normal GFR while others remained in apparent 320 renal failure. Consequently, none of these variables differed significantly from their level in rats subjected to sham-ischemia. We were unable to detect a significant correlation ($r^2 = 0.03$, 321 322 n=8), in rats subjected to ischemia, between GFR and tissue PO₂ at day 5 after surgery.

323 Blood oximetry and renal oxygen consumption and delivery. Arterial blood hematocrit 24 h 324 after renal ischemia was 12% less than after sham-ischemia (Table 1). We were unable to 325 detect a significant correlation ($r^2=0.034$, n=9), in rats subjected to ischemia, between 326 hematocrit and tissue PO₂ 24 h after reperfusion. By 5 days after renal ischemia, hematocrit 327 was similar in the two groups of rats. Arterial blood PO₂ was 22% less, and SO₂ 2.7% less, in 328 rats 24 h after renal ischemia than after sham surgery. Renal DO₂ tended to be (29%) less 24 h 329 after renal ischemia than sham-ischemia, although this apparent effect was not statistically 330 significant (P = 0.06). There was no significant difference in renal DO₂ 5 days after surgery. 331 When both time points were considered together (24 h and 5 days), renal VO₂ was 55% less in rats subjected to ischemia than in those subjected to sham surgery. The fractional extractionof oxygen did not differ significantly between the treatments at either time-point.

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335 Protocol 3: Cellular hypoxia and hypoxic signaling after renal ischemia

336 Pimonidazole adduct immunohistochemistry. No pimonidazole adducts were detected in 337 tissues from rats that did not receive pimonidazole chloride or in sections that were not 338 incubated with the primary antibody (data not shown). Kidney sections from sham operated rats appeared morphologically normal (Fig. 3 & 4). Pimonidazole adducts were largely absent 339 340 in the cortical region of rats 24 h following sham-ischemia. However, there was diffuse 341 staining of pimonidazole adducts in tubular elements of the outer and inner medulla following 342 sham-ischemia. Kidney sections from rats 24 h following recovery from renal ischemia 343 showed relatively little staining for pimonidazole adducts across all regions of the kidney, but 344 some diffuse staining was present 5 days following ischemia and reperfusion. However, 345 luminal aspects of tubules were often stained positive for pimonidazole adducts after renal 346 ischemia, suggestive of marked tubular obstruction. There was significant cellular sloughing 347 and disintegration of the brush border/apical membrane of tubules after renal ischemia. In 348 addition, there were considerable cellular debris in the luminal aspects of tubules at 24 h after 349 renal ischemia. Tubular profiles surrounding the debris-riddled tubules were often flattened. 350 In contrast, tubules appeared to be mostly dilated 5 days after renal ischemia. By 5 days after 351 ischemia, tubules in the cortex, outer and inner medulla appeared to be more dilated than after 352 sham-ischemia or 24 h after renal ischemia.

353 *HIF-1a and HIF-2a proteins.* When both the 24 h and 5 day time-points were considered 354 collectively, the expression of HIF-1a protein in the cortex, outer medulla and inner medulla 355 was less after renal ischemia than after sham-ischemia (Fig. 5A-C). However, not all 356 comparisons at individual time-points were statistically significant. HIF-1a levels in the 357 cortex were 88.3% less 5 days after renal ischemia than at the corresponding time-point after

358 sham-ischemia (Fig. 5A). Similarly, in the outer medulla, HIF-1α was 62.2% less 24 h after 359 renal ischemia and 79.7% less 5 days after renal ischemia than after sham surgery (Fig. 5B). 360 In contrast, in the inner medulla, levels of HIF-1 α protein did not differ significantly, between 361 rats subjected to ischemia compared to those subjected to sham-ischemia, at either the 24 h or 362 5 day time-point (Fig. 5C). When both the 24 h and 5 day time points were considered 363 collectively, the expression of HIF-2 α protein was markedly less, in rats subjected to ischemia 364 compared to those subjected to sham-ischemia, in the cortex and the outer medulla but not in 365 the inner medulla. The level of HIF-2 α in the cortex was 86.9% less 5 days after ischemia 366 than after sham-ischemia (Fig. 5D). In the outer medulla of rats subjected to renal ischemia, 367 HIF-2a expression was 55% less 24 h and 89.2% less 5 days after ischemia, than after sham-368 ischemia (Fig. 5E). The deficits in HIF-1 α and HIF-2 α in rats subjected to renal ischemia did 369 not diminish between the 24 h and 5 day time points, if anything, becoming more marked 370 (Fig. 5).

371 *Expression of genes for HIF-1a, HIF-2a, VEGF-a and HO-1.* There were no significant 372 differences in the expression of mRNA for *HIF-1a, HIF-2a* or *VEGF-a*, either 24 h or 5 days 373 following renal ischemia compared to after sham-ischemia (Fig. 6). The expression of *HO-1* 374 tended to be greater after ischemia than after sham-ischemia, although this apparent effect 375 was only statistically significant at the 5 day time-point.

Collagen deposition. Twenty-four hours after renal ischemia, picrosirius red staining did not differ significantly, from that seen in rats subjected to sham-ischemia, in either the cortex or the outer medulla. However, it was 43% less in the inner medulla (Fig. 7). By 5 days after renal ischemia, picrosirius red staining was 50% greater in the cortex of rats subjected to ischemia than those subjected to sham-ischemia. There was an apparent effect of the duration of recovery period on picrosirius red staining, which in the cortex and inner medulla was significantly greater 5 days after ischemia or sham-ischemia than at the 24 h time point. *Indices of renal dysfunction.* The plasma concentrations of urea and creatinine, and the urinary albumin-creatinine ratio were all greater in rats after ischemia than after shamischemia (Fig. 8). These effects were statistically significant at the individual time points with the exception of urinary albumin to creatinine ratio 24 h after ischemia, where sufficient urine for analysis could only be generated from two animals.

388 Discussion

389 We determined the time-course of changes in, and the spatial distribution of, renal 390 tissue PO₂ during the subacute phase of severe renal IRI. Using four different methods for 391 assessing renal tissue oxygenation, we could not detect tissue hypoxia during the 392 extension/recovery phase of IRI. Indeed, if anything, there was relative hyperoxia up to 48 h 393 after an hour of bilateral renal ischemia. We also observed downregulation of the abundance 394 of HIF-1α and HIF-2α protein, particularly in the cortex and outer medulla, both 24 h and 5 395 days after reperfusion. The apparent absence of renal hypoxia is consistent with the pattern of 396 changes in renal DO₂ and VO₂ after ischemia and reperfusion. That is, RBF was relatively 397 normal but there was a marked reduction in sodium reabsorption, and so presumably oxygen 398 utilization for sodium reabsorption, at both 24 h and 5 days after reperfusion. When both 399 time-points were considered together, renal VO₂ was significantly less, and DO₂ tended to be 400 less, in rats subjected to ischemia than in those subjected to sham-ischemia. Thus, tissue PO_2 401 appears to be well maintained during the extension/recovery phase of severe renal IRI 402 because changes in renal DO₂ and VO₂ are relatively balanced.

The methods we used to assess renal oxygenation have both strengths and weaknesses (11, 33). Radiotelemetry allows continuous measurement of renal tissue PO_2 in the absence of confounding effects of anesthesia (22, 23). However, tissue PO_2 can only be expressed in relative terms and can be measured at only one site in each animal. Clark electrodes allow generation of a spatial map of tissue PO_2 , but only in anesthetized animals

408 (11, 33). Furthermore, it is not possible to resolve tissue PO_2 to the level of specific vascular 409 and tubular elements, except in the superficial cortex (43). In addition, as we have found 410 previously with Clark electrodes inserted into renal tissue from the dorsal surface of the 411 kidney (32), the steep cortico-medullary gradient in tissue PO₂ generated in many previous 412 studies (6, 10, 29) is not obviously evident. We have no adequate explanation for this, 413 although it may relate to our use of relatively large electrodes (50 µm) or the angle of entry to 414 the renal tissue, from the dorsal surface of the kidney, as a consequence of which the tip of the 415 electrode does not enter the renal papilla. Pimonidazole adduct immunohistochemistry allows 416 detection of cells with $PO_2 < 10$ mmHg but does not provide a quantitative measure of tissue 417 PO_2 (37). Furthermore, as we found in the current study and previously (1), it is prone to 418 artifactual staining of cellular debris and casts within damaged tubules. Quantification of the 419 abundance of HIF-1 α and HIF-2 α protein provides information about the state of hypoxia 420 signaling pathways. However, factors other than tissue PO₂ contribute to the regulation of 421 HIF signaling (16). Thus, interpretation of our failure to detect hypoxia by any one of these 422 methods would merit caution. However, the fact that our observations were consistent across 423 the four methods provides compelling evidence that, at least in this severe form of IRI, tissue 424 hypoxia is not an obligatory characteristic of the period from 24 h to 5 days after severe renal 425 ischemia and reperfusion.

The most likely explanation for the absence of hypoxia 24 h and 5 days after reperfusion, and even increased tissue PO₂ at 24 h, is reduced sodium reabsorption and thus renal VO₂. In the rats we studied, the deficit in sodium reabsorption 24 h after ischemia and reperfusion could be attributed to the decreased filtered load of sodium. This appears to drive downregulation of Na⁺-K⁺-ATPase activity. For example, in response to severe renal ischemia (*i.e.* 60 min), the abundance (and activity) of basolateral Na⁺-K⁺-ATPase pumps, the apical Na-K-2Cl and the thiazide-sensitive Na⁺-Cl⁻ cotransporters was shown to be greatly

reduced (25). But the magnitude of the apparent reduction in renal VO₂ we observed was 433 434 considerably less than the magnitude of the reduction in sodium reabsorption. For example, 435 sodium reabsorption was less than 1% of rats subjected to sham-ischemia, while VO₂ was 436 34% that of rats subjected to sham-ischemia 24 h after reperfusion. These observations are 437 consistent with the concept that oxygen utilization for sodium reabsorption becomes less 438 efficient in AKI. In support of this concept, Redfors and colleagues studied renal oxygen 439 utilization in patients with AKI subsequent to cardiothoracic surgery (35). They found a 440 deficit in sodium reabsorption of 59% in patients with AKI after cardiothoracic surgery 441 compared to patients without AKI (35). In contrast, renal VO₂ was similar in the two groups 442 of patients. Furthermore, renal VO₂ per unit of reabsorbed sodium was 2.4 times greater in 443 patients with AKI than in those without AKI (35). The inefficiency of oxygen utilization for 444 sodium reabsorption in AKI appears to be driven by multiple factors, including loss of 445 polarity of Na⁺-K⁺-ATPase pumps, oxidative stress and reduced bioavailability of nitric oxide 446 (17, 24).

447 Renal tissue PO_2 is determined by the balance between local DO_2 and VO_2 (12). 448 Thus, tissue PO₂ during recovery from AKI is likely to be model-dependent. In a model of 449 severe AKI such as the one used in the current study, in which the filtered load of sodium 450 (and thus oxygen utilization for sodium reabsorption) is greatly reduced but renal blood flow 451 (and thus presumably local tissue DO_2) is well preserved, the absence of tissue hypoxia, and 452 even tissue hyperoxia, might be expected. On the other hand, tissue hypoxia might be 453 predicted in a model of less severe renal dysfunction, and thus better preserved GFR. This 454 concept is consistent with clinical observations in patients after renal transplantation. Using 455 blood oxygen level-dependent magnetic resonance imaging (BOLD-MRI), Sadowski and 456 colleagues observed greater renal medullary oxygenation in patients with acute allograft 457 rejection than in patients with normal functioning allografts, despite the former having a 458 deficit in renal medullary perfusion (38). Similarly, Rosenberger and colleagues observed low 459 HIF-1 α abundance in biopsies of patients with non-functional allografts, but induction of HIF-460 1 α in biopsies from functional grafts (36). Thus, there is a strong rationale for the methods 461 used in the current study to be applied to a less severe model of AKI, in which tissue hypoxia 462 might be more likely to occur.

463 It is noteworthy that HIF-1 α and HIF-2 α protein expression was downregulated not 464 just at 24 h after reperfusion, presumably driven in part by increased tissue oxygen 465 availability, but also 5 days after reperfusion, when tissue PO₂ was similar in rats exposed to 466 ischemia and sham-ischemia. Inhibition of HIF-1 α and HIF-2 α abundance appears to be 467 mediated by post-translational processes at both 24 h and 5 days after reperfusion, since the 468 expression of mRNA for these proteins was relatively normal at both time-points. The 469 bioavailability of HIFs is influenced by various factors, such as their phosphorylation (20), 470 and hydroxylation of proline and asparagine residues on HIFs (44) that target these protein for 471 ubiquitinylation. The levels of proline hydroxylases (PHDs) have been shown to be unaltered 472 following ischemia and reperfusion of the kidney (13, 39). A caveat to that is the post-473 translational modification of HIFs by PHDs in the kidney is likely complex given that the 474 expression patterns, and thus sensitivity, of PHDs varies in different regions of the kidney 475 likely because of the heterogeneity of renal tissue PO_2 under physiological conditions (39). It 476 is also noteworthy that mRNA for VEGF- α and HO-1, genes under the control of the HIF-1 α 477 and HIF-2 α promoter, were not downregulated at 24 h or 5 days after reperfusion. This 478 observation is consistent with the concept that factors other than HIFs regulate expression of 479 these genes in the subacute phase of severe IRI. The signaling pathway for VEGFs is complex 480 and is critical for neo-vascularization. A myriad of factors apart from HIFs, such as VEGF 481 receptor signaling complexes and neurolipin, are able to modulate the abundance and activity 482 of VEGFs (21). Kanellis and colleagues showed that expression of VEGF was unaltered in

response to ischemia reperfusion of the kidney (18). Interestingly, the expression of VEGF receptor 2 was increased following ischemia and VEGF was redistributed to the basolateral membrane, consistent with the established role of VEGF in the maintenance of an adequate blood supply, in remaining viable tissues, as evinced in the current study by relatively well maintained renal blood flow (19). Nevertheless, the permanent loss of peritubular capillaries, due to inadequate vascular reparation and/or neo-vascularization, appears to be an important event in the progression from ischemia-induced AKI to CKD (2, 4).

490 **Perspectives & Significance**

491 In models of AKI induced by complete renal ischemia, hypoxia during the period of 492 ischemia is obligatory and is likely one of the drivers of necrosis and apoptosis associated 493 with the development of AKI after reperfusion. Furthermore, other important factors during 494 reperfusion, such as oxidative stress (5, 7) and influx of immune-modulatory cells (14, 42) are 495 initiated, at least partly, by the hypoxia during ischemia. In the first few hours after 496 reperfusion (acute phase), reduced renal tissue or microvascular PO₂ has been observed in 497 some (27, 28) but not all (1) cases. To the best of our knowledge, our current report describes 498 the first detailed investigation of tissue oxygenation during the subacute phase of renal IRI. 499 We provide compelling evidence that, at least in severe IRI modeling subacute and end stage 500 renal disease, renal tissue hypoxia is not present 24 h and 5 days after reperfusion. It is 501 possible that the absence of hypoxia at these time-points in this experimental model of severe 502 IRI is a consequence of the degree of renal damage and the consequent deficit in renal oxygen 503 consumption. Thus, future studies should focus on less severe models of AKI and follow 504 animals for longer periods after reperfusion, to better characterize the natural history of renal 505 oxygenation during progression from AKI to CKD.

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	Sham, 24 h		Ischemia, 24 h		Sham, 5 days		Ischemia, 5 days	5	2-1	way ANO	VA		tomous arison
Parameter		Ν		Ν		Ν		Ν	P _{Tr}	PT	P _{Tr*T}	S1 Vs I1	S5 Vs I5
Systemic													
Body weight after													
ischemia or sham-	397.6 ± 23.8	10	378.2 ± 17.6	9	469.4 ± 14.9	7	364.5 ± 19.7	8	0.005	0.17	0.05	0.77	0.002
ischemia (g)													
Kidney weight (g)	1.5 (1.3, 1.5)	10	1.5 (1.4, 1.6)	9	1.4 (1.4, 1.7)	7	2.2 (2.2, 2.8)	8	< 0.001	< 0.001	0.04	0.92	< 0.001
Kidney weight (g/g BW)	3.6 (3.3, 4.1)	10	3.9 (3.7, 4.2)	9	3.3 (3.2, 3.5)	7	6.6 (5.4, 8.3)	8	< 0.001	0.16	< 0.001	0.55	0.002
Mean arterial blood pressure (mmHg)	118.3 (108.4, 140.4)	10	123.9 (108.7, 134.4)	9	116.9 (102.3, 118.7)	7	116.3 (87.2, 122.3)	8	0.59	0.07	0.79	0.86	0.93
Blood oximetry													
Arterial blood PO ₂ (mmHg)	113.7 ± 6.1	10	89.2 ± 4.7	9	97.4 ± 5.0	7	100.7 ± 7.7	8	0.09	0.70	0.03	0.01	0.93
Hematocrit (%)	44.8 (41.9, 45.6)	10	38.5 (35.4, 40.3)	9	43.0 (41, 44.5)	7	42.1 (41.4, 43.3)	8	0.01	0.35	0.06	0.04	0.07
Arterial blood SO ₂ (%)	98.3 (97.1, 99.0)	10	96.5 (92.3, 97.5)	9	96.5 (94, 97.5)	7	97.5 (96.6, 98.5)	8	0.36	0.80	0.003	0.02	0.22
Renal oxygen delivery (µmol/min)	30.5 ± 2.7	10	21.7 ± 2.4	9	29.7 ± 5.5	7	20.8 ± 2.9	8	0.03	0.95	0.99	0.06	0.28
Renal oxygen delivery (nmol/min/g BW)	80.5 ± 2.7	10	57.7 ± 5.6	9	64.3 ± 12.7	7	56.5 ± 6.8	8	0.20	0.60	0.60	0.10	0.82
Renal oxygen consumption (µmol/min)	2.9 (1.1, 6.0)	6	1.1 (0.7, 2.7)	7	2.6 (1.8, 4.1)	7	1.4 (0.9, 2.1)	8	0.04	0.99	0.99	0.30	0.06
Renal oxygen consumption (nmol/min/g BW)	8.3 ± 2.8	6	4.1 ± 1.1	7	6.1 ± 1.2	7	3.8 ± 0.6	8	0.08	0.70	0.80	0.30	0.20
Fractional extraction O ₂ (%)	10.1 (6.0, 17.8)	6	5.1 (3.3, 11.7)	7	7.9 (7.1, 14.4)	7	6.9 (4.7, 8.2)	8	0.11	0.95	0.70	0.44	0.51

639 Table 1: Systemic and blood oxygen parameters of rats 24 h or 5 days after ischemia or sham-ischemia

Normality of the data was assessed using the Shapiro-Wilk test. Data that did not violate normality are expressed as mean \pm standard error of the mean while data that violated normality are expressed as median (25th percentile, 75th percentile). P_{Tr}, P_T and P_{Tr*T} are the outcomes of 2 way analysis of variance (ANOVA) with factors treatment (Tr) and time (T) for data that did not violate normality. For data that violated normality, an ANOVA on ranking was performed instead. Dichotomous comparisons of continuous variables were made using Student's t-test for data that did not violate normality. For data that violated normality. For data that violated normality. For data that violated normality, a Mann-Whitney U-test was performed for dichotomous comparisons. P-values for dichotomous comparisons were conservatively adjusted using the Dunn-Sidak correction with k=2 to account for the fact that comparisons were made at 24 h and 5 days. BW: body weight, S1: 24 h after sham-ischemia, I1: 24 h after ischemia and reperfusion, S5: 5 days after sham-ischemia, I5: 5 days after ischemia and reperfusion.

642													
	Sham, 24 h		Ischemia, 24 h		Sham, 5 days		Ischemia, 5 days		2-way ANOVA			Dichotomous comparison	
Parameter		Ν		Ν		Ν		Ν	$\mathbf{P}_{\mathbf{Tr}}$	P _T	P _{Tr*T}	S1 Vs I1	S5 Vs I5
Renal blood flow (ml/min)	3.5 ± 2.9	10	3.0 ± 0.4	9	3.6 ± 0.7	7	2.4 ± 0.3	8	0.13	0.83	0.68	0.55	0.29
Renal blood flow (µl/min/g BW)	9.09 ± 0.98	10	7.93 ± 0.79	9	7.71 ± 1.59	7	6.65 ± 0.78	8	0.30	0.21	0.96	0.60	0.81
Renal plasma flow (ml/min)	1.7 (1.5, 2.1)	10	1.7 (1.4, 2.0)	9	1.6 (1.5, 3.0)	7	1.6 (0.9, 1.9)	8	0.42	0.65	0.87	0.36	0.51
Renal plasma flow (µl/min/g BW)	5.1 ± 0.6	10	4.9 ± 0.5	9	4.4 ± 0.9	7	3.8 ± 0.4	8	0.77	0.31	0.94	0.96	0.80
Glomerular filtration rate (ml/min)	0.8 (0.7, 1.2)	10	0.001 (0, 0.008)	9	1.0 (0.5, 1.8)	7	0.07 (0.01, 0.4)	9	< 0.001	0.17	0.11	< 0.001	0.14
Glomerular filtration rate (nl/min/g BW)	2400 (1520, 2960)	10	3.3 (0, 2.2)	9	2110 (1040, 3470)	7	190 (27, 1080)	8	< 0.001	0.18	0.03	< 0.001	0.20
Urine flow (µl/min)	7.0 ± 1.0	10	1.0 ± 0.5	9	10.0 ± 4.0	7	6.0 ± 2.0	8	0.01	0.09	0.99	0.001	0.36
Urine flow (nl/min/g BW)	16.0 (10.0, 28.0)	10	2.2 (0, 5.5)	9	17.0 (13.0, 25.0)	7	15 (1.1, 39.0)	8	0.04	0.05	0.06	0.002	0.66
Sodium excretion (µmol/min)	0.4 (0.2, 1.0)	9	0 (0, 0.18)	9	0.20 (0.2, 0.6)	7	0.20 (0.1, 0.3)	8	0.004	0.46	0.09	0.003	0.38
Sodium excretion (nmol/min/g BW)	1.0 (0.5, 2.1)	9	0 (0, 0.47)	9	0.5 (0.36, 1.3)	7	0.6 (0.35, 0.71)	8	0.02	0.53	0.05	0.01	0.90

640641 Table 2: Renal hemodynamic of rats 24 h or 5 days after ischemia or sham-ischemia

Sodium reabsorption (µmol/min)	111.9 (93.6, 170.2)	9	0.17 (0, 1.0)	9	138.3 (63.3, 250.4)	7	8.75 (1.5, 52.0)	8	< 0.001	0.2	0.15	< 0.001	0.14
Sodium reabsorption (nmol/min/g BW)	307.5 (204.7, 427.6)	9	0.46 (0, 2.7)	9	310 (143.6, 479.3)	7	25.7 (3.6, 153.1)	8	< 0.001	0.18	0.03	< 0.001	0.20
Filtration fraction (%)	46.9 (34.1, 69.8)	10	0.1 (0, 0.5)	9	48.5 (29.7, 60.5)	7	7.0 (0.6, 21.7)	8	< 0.001	0.13	0.02	< 0.001	0.20

Normality of the data was assessed using the Shapiro-Wilk test. Data that did not violate normality are expressed as mean \pm standard error of the mean while data that violated normality are expressed as median (25th percentile, 75th percentile). P_{Tr}, P_T and P_{Tr*T} are the outcomes of 2 way analysis of variance (ANOVA) with factors treatment (Tr) and time (T) for data that did not violate normality. For data that violated normality, an ANOVA on ranking was performed instead. Dichotomous comparisons of continuous variables were made using Student's t-test for data that did not violate normality. For data that violated normality, a Mann-Whitney U-test was performed for dichotomous comparisons. P-values for dichotomous comparisons were conservatively adjusted using the Dunn-Sidak correction with k=2 to account for the fact that comparisons were made at 24 h and 5 days. BW: body weight, S1: 24 h after sham-ischemia, I1: 24 h after ischemia and reperfusion, S5: 5 days after sham-ischemia, I5: 5 days after ischemia and reperfusion.

643 Figure 1: Temporal changes of inner medullary tissue PO₂ following renal ischemia or sham ischemia. Values are mean \pm SEM for rats subjected to either an hour of sham (n=5) or 644 645 bilateral renal ischemia (n=7). Tissue PO₂, assessed as current through the carbon paste 646 electrode, was recorded before (day -1) and after (days 0-5) surgery. Current was averaged over 24 h period and is expressed as a percentage of its mean value on to the day before the 647 648 surgery (day -1). P_{Treatment}, P_{Time} and P_{Treatment*Time} are the outcomes of a 2-way repeated 649 measures analysis of variance with factors treatment and time. * denotes $P \le 0.05$ for specific 650 comparisons between the two treatment groups at each time point using Student's unpaired t-651 test, without correction for multiple comparisons.

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653 Figure 2: Assessment of tissue PO₂ by Clark electrode. The electrode was first inserted 2 654 mm into the cortex at 6 random sites across the left kidney. In panel A, multiple 655 measurements are shown for each rat, with the various rats represented by different symbols. 656 Filled symbols represent rats subjected to renal ischemia (n=9 at 24 h and n=8 at 5 days) 657 while open symbols represent rats subjected to sham ischemia (n=10 at 24 h and n=7 at 5 658 days). Measurements of cortical tissue PO₂ for each rat were averaged and are presented as 659 the between rat mean \pm SEM in (B). In panel B, P_{Tr}, P_T and P_{Tr*T} are the outcomes of a 2-way 660 analysis of variance (ANOVA) with factors treatment and time. P-values above each pair of 661 columns and error bars show the outcomes of Student's unpaired t-test conservatively 662 adjusted using the Dunn-sidak correction with k = 2 to account for the fact that comparisons 663 were made at 24 h and 5 days. A tissue PO₂ profile with depth was established by advancing 664 the electrode from the cortical surface at 1 mm increments, up to 10 mm into the left kidney 665 either 24 h (C) or 5 days (D) following recovery from either ischemia or sham-ischemia. 666 Symbols and error bars are the mean \pm SEM for rats subjected to either an hour of sham (open 667 circles) or bilateral renal ischemia (closed circles). In panels C and D, P_D, P_{Tr} and P_{D*Tr} are the outcomes of 2-way repeated measures ANOVA with factors depth and treatment. * denotes P 668 669 ≤ 0.05 and is the outcome of Student's unpaired t-test without correction for multiple 670 comparisons 671

Figure 3: Pimonidazole adduct immunohistochemistry of renal sections 24 h following
 recovery from bilateral renal ischemia or sham-ischemia. Images are typical of the
 cortical, outer and inner medullary region of the 6 kidneys examined in each group.

Figure 4: Pimonidazole adduct immunohistochemistry of renal sections 5 days following
 recovery from bilateral renal ischemia or sham surgery. Images are typical of the cortical,
 outer and inner medullary region of the 6 kidneys examined in each group.

- 680 Figure 5: Expression of HIF proteins after bilateral renal ischemia or sham ischemia. 681 Immunoblots for HIF-1 α (A-C) and HIF-2 α (D-F) of tissue extracts from the cortex, outer and 682 inner medulla of the left kidneys of rats 24 h and 5 days following recovery from either sham-683 ischemia (open circles) or bilateral renal ischemia (closed circles); n = 6 per group. Panel G 684 shows a typical image of the gel following electrophoresis and panel H reflects a typical 685 image of the nitrocellulose membrane following transfer. Values are expressed as median (25th percentile, 75th percentile). Paired comparisons were performed using the Mann-686 Whitney U-test. Because paired comparisons were made at two time points, P-values were 687 conservatively adjusted using the Dunn-Sidak method with k = 2. P_{Tr} , P_T and P_{Tr*T} are the 688 689 outcomes of 2 way analysis of variance on ranking with the factors treatment and time. AU: 690 arbitrary unit, S1: 24 h after sham-ischemia, S5: 5 days after sham-ischemia, I1: 24 h after 691 ischemia, I5: 5 days after ischemia
- 692

693 **Figure 6**: **mRNA expression of** *HIF-1a*, *HIF-2a*, *VEGF-a* and *HO-1*. Expression of *HIF-*694 *1a*, *HIF-2a*, *VEGF-a* and *HO-1* are presented as relative to that of control animals. Values are 695 expressed as mean \pm SEM and * denotes P < 0.05 for specific comparisons between the two 696 treatment groups at each time point using Student's unpaired t-test.

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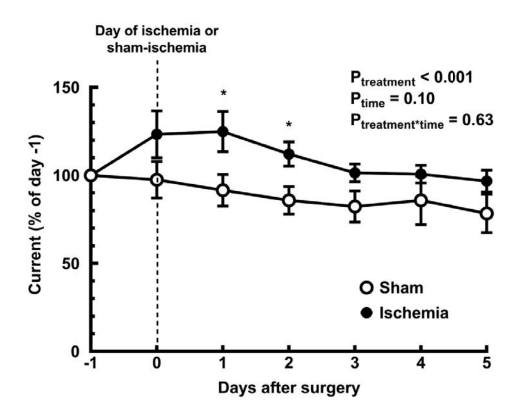
698 Figure 7: Collagen deposition in kidneys of rats. The percentage areas of interstitial fibrosis 699 relative to the areas of the cortex, outer and inner medulla are shown for rats 24 h and 5 days 700 after recovery from either sham-ischemia or bilateral renal ischemia (IR), n = 6 per group. 701 Values are expressed as mean \pm SEM. Paired comparisons were performed using Student's 702 unpaired t-test (*P ≤ 0.05). Because paired comparisons were made at two time points, P-703 values were conservatively adjusted using the Dunn-Sidak method with k = 2. P_{Tr} , P_{T} and 704 P_{Tr*T} are the outcomes of 2 way analysis of variance with the factors treatment (Tr) and time 705 (T).

706

707 Figure 8: Indicators of renal dysfunction. Plasma concentrations of urea (A) and creatinine 708 (B) and the urinary albumin to creatinine ratio (C) are shown for rats 24 h and 5 days after 709 sham-ischemia (open circles) or bilateral renal ischemia (closed circles), n = 6 per group. Values are expressed as median (25th percentile, 75th percentile). Paired comparisons were 710 performed using the Mann-Whitney U-test. Because paired comparisons were made at two 711 712 time points, P-values were conservatively adjusted using the Dunn-Sidak method with k = 2. 713 P_{Tr}, P_T and P_{Tr*T} are the outcomes of 2 way analysis of variance on ranking with the factors 714 treatment (Tr) and time (T).

715

Figure 1





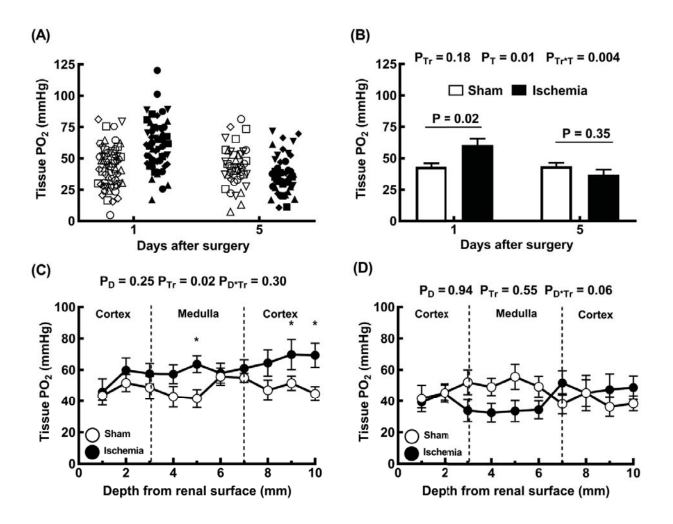
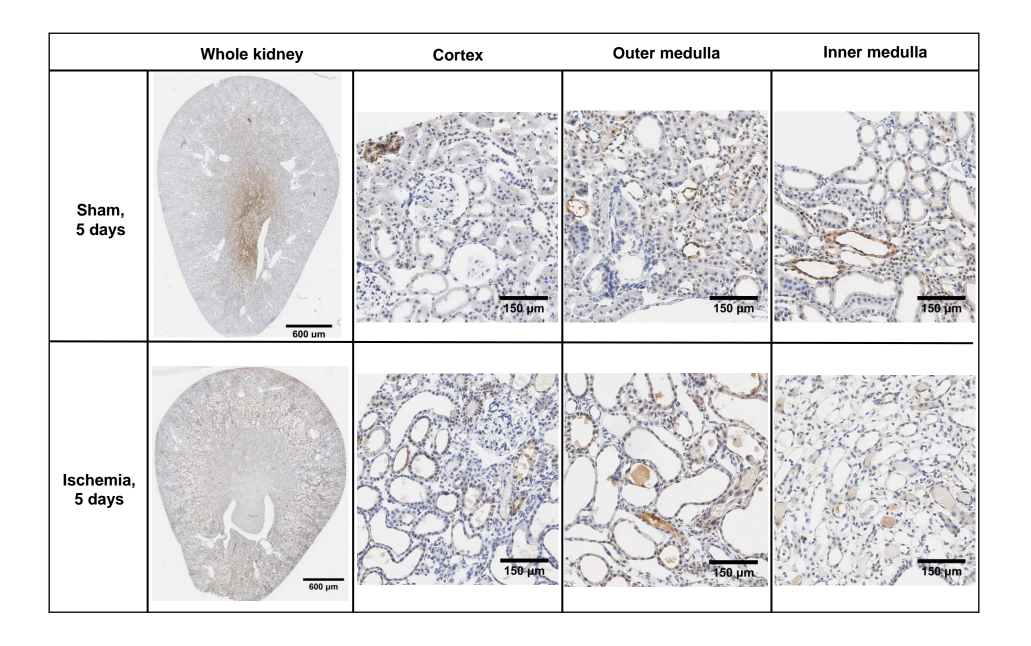
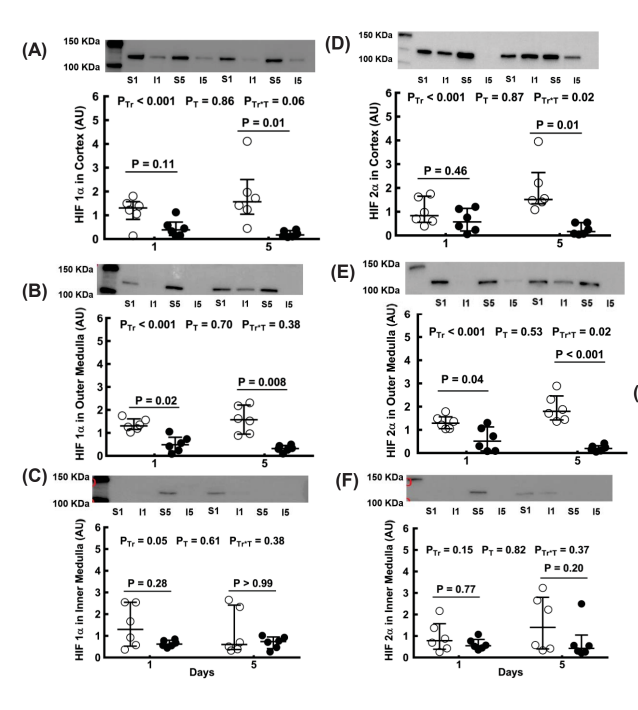


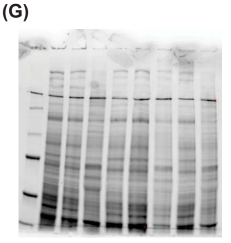
Figure 3

	Whole kidney	Cortex	Outer medulla	Inner medulla
Sham, 24 h	<u>б00 µm</u>	Т50 µm	<mark>Т150 µт</mark>	
lschemia, 24 h	Торияника Пория	То пр	Т <u>то, ит</u>	<u>150-µm</u>

Figure 4







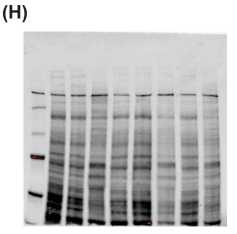
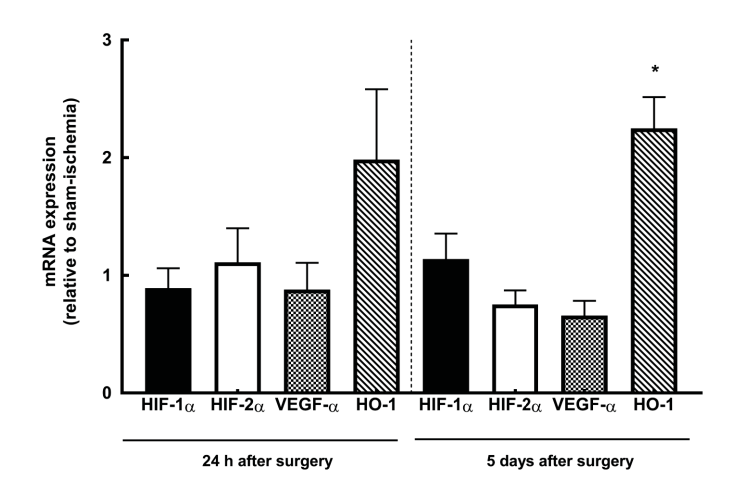


Figure 6



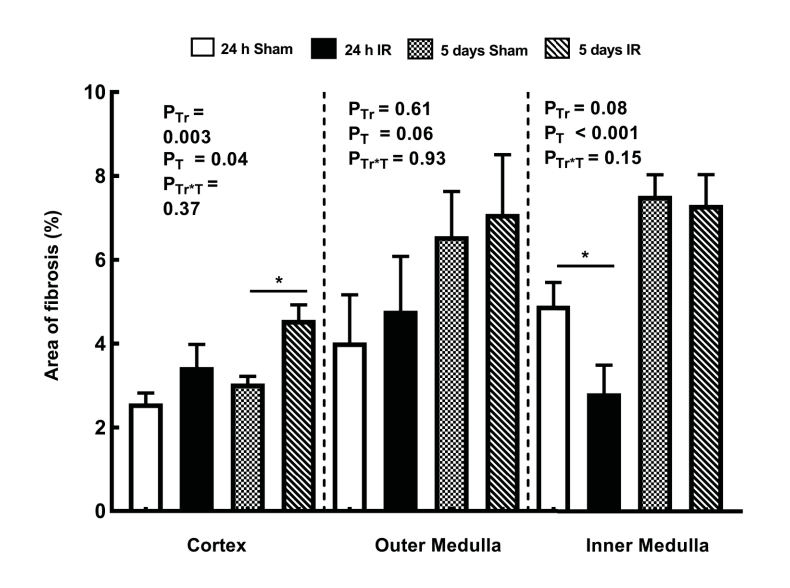


Figure 8

