Monitoring kidney size to interpret MRI-based assessment of renal oxygenation in acute pathophysiological scenarios

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Abstract

Aim: Tissue hypoxia is an early key feature of acute kidney injury. Assessment of renal oxygenation using magnetic resonance imaging (MRI) markers $T_2$ and $T_2^*$ enables insights into renal pathophysiology. This assessment can be confounded by changes in the blood and tubular volume fractions, occurring upon pathological insults. These changes are mirrored by changes in kidney size (KS). Here, we used dynamic MRI to monitor KS for physiological interpretation of $T_2^*$ and $T_2$ changes in acute pathophysiological scenarios.

Methods: KS was determined from $T_2^*$, $T_2$ mapping in rats. Six interventions that acutely alter renal tissue oxygenation were performed directly within the scanner, including interventions that change the blood and/or tubular volume. A biophysical model was used to estimate changes in $O_2$ saturation of hemoglobin from changes in $T_2^*$ and KS.

Results: Upon aortic occlusion KS decreased; this correlated with a decrease in $T_2^*$, $T_2$. Upon renal vein occlusion KS increased; this negatively correlated with a decrease in $T_2^*$, $T_2$. Upon simultaneous occlusion of both vessels KS remained unchanged; there was no correlation with decreased $T_2^*$, $T_2$. Hypoxemia induced mild reductions in KS and $T_2^*$, $T_2$. Administration of an X-ray contrast medium induced sustained KS increase, with an initial increase in $T_2^*$, $T_2$ followed by a decrease. Furosemide caused $T_2^*$, $T_2$ elevation and a minor increase in KS. Model calculations yielded physiologically plausible calibration ratios for $T_2^*$.

Conclusion: Monitoring KS allows physiological interpretation of acute renal oxygenation changes obtained by $T_2^*$, $T_2$. KS monitoring should accompany MRI-oximetry, for new insights into renal pathophysiology and swift translation into human studies.

KEYWORDS

acute kidney injury, BOLD-MRI, hypoxia, kidney size, renal oxygenation
Current options for the treatment of acute kidney injury (AKI) are not adequate. Major obstacles include the limitations of serum creatinine-based metrics. To overcome this, several alternative blood- or urine-based markers reflecting renal injury, inflammation, fibrosis, or repair have been proposed. Despite the promise, the clinical performance of these markers has been modest, and none has advanced to provide a point-of-care diagnosis for AKI. In general, these markers fail to reveal early events in AKI pathophysiology, such as tissue hypoxia. Recognizing these limitations, synergistic approaches that include magnetic resonance imaging (MRI) are called for. MRI facilitates the non-invasive assessment of several structural and functional kidney features. Among these, kidney size (KS) has gained substantial interest as a marker to diagnose and stage kidney disorders since KS changes are associated with several renal pathologies.

What has not been considered, however, is that KS is also crucial for the interpretation of MRI-based assessments of renal tissue oxygenation obtained by blood oxygenation level-dependent (BOLD) MRI techniques. This approach relies on the fact that deoxygenated hemoglobin (deoxygenHb) is paramagnetic and, therefore, impacts the MRI relaxation volume; therefore the renal T2*, T2 – tissue O2 relationship is less sensitive to macrovessels. T2*, T2 reflect the amount of deoxygenHb per tissue volume; therefore the renal T2*, T2 – tissue O2 relationship is also dependent on the blood volume and the tubular volume fractions. Because acute changes in these fractions are often accompanied by KS changes, simultaneous measurements of changes in KS and T2*, T2 are essential for the accurate physiological interpretation of MR-based assessments of renal oxygenation. Since tissue hypoxia is a common early feature in the pathophysiology of AKI and progression to chronic kidney disease (CKD), physiological interpretation of MR probing of oxygenation could render non-invasive MR-oximetry a vital assay for research into renal (patho-)physiology and for clinical application.

Events leading to acute renal hypoxia are often associated with KS changes. Studies emulating clinical procedures such as clamping of the suprarenal aorta or renal artery during surgery, or the low arterial target pressure during cardiopulmonary bypass, revealed KS reductions. Conversely, anecdotal evidence indicates opposite effects for obstructing the renal vein, such as in partial nephrectomy or thrombus formation in renal cell carcinoma. Studies emulating administration of X-ray contrast media (CM) for cardiac procedures showed renal hypoxia and increased intratubular pressure. Due to the relatively rigid renal capsule, the latter results in an “intrarenal compartment syndrome”: as intrarenal pressure increases, intrarenal blood vessels become compressed, leading to tissue hypoxia. Accordingly, KS should significantly increase.

Recognizing the potential of MR-based probing of renal oxygenation as a meaningful tool for research into renal physiology and disorders, this study provides systematic tests for acute changes in T2* and T2 as markers of oxygenation, and in KS determined from the T2*, T2 maps. Serial in vivo parametric MRI mapping of T2*, T2 was performed in rats during interventions that alter renal tissue oxygenation reversibly (a brief occlusion of the suprarenal aorta, the left renal vein or both; hypoxemia), and with lasting-effects (injection of CM or furosemide). We hypothesize that monitoring KS will allow physiological interpretation of acute changes in renal oxygenation measured by T2*, T2.

### RESULTS

#### 2.1 | Aortic occlusion, renal venous occlusion, combined occlusion

By inflation of MR-safe remotely-controlled occluders, short-term occlusions of the suprarenal aorta, the left renal vein, and simultaneous occlusions of both vessels were performed.

Upon occlusion of the suprarenal aorta, the coronal mid-slice cross-sectional area of the kidney (hereafter referred to as “kidney size,” KS) determined from T2 maps (Figure 1A) decreased by 6 ± 1% (mean ± SEM; Figure 1B). Upon occlusion release, KS returned to baseline. Aortic occlusion resulted in decreases in T2 in the renal cortex (CO), outer medulla (OM), and inner medulla (IM) of 22 ± 2%, 27 ± 2%, and 13 ± 3% (Figure 1C–E). Upon release, T2 in the IM initially decreased further, yet T2 in all layers quickly returned to baseline. Similar results were obtained for T2. Upon aortic occlusion, KS determined from T2 maps decreased by 6 ± 2% and returned to baseline upon release (Figure 1B). T2 changes during aortic occlusion were more pronounced than for T2, decreasing by 29 ± 3%, 39 ± 3%, and 22 ± 4% in the CO, OM, and IM (Figure 1C–E). Upon release, T2 in the IM initially decreased further, yet T2 for all layers returned to baseline. Changes in T2 showed a strong correlation with changes in KS; correlations with changes in T2 were more moderate (Table 1).

Upon occlusion of the renal vein, T2-derived KS increased by 5 ± 1% and returned to baseline upon release (Figure 2B). Venous occlusion led to T2 decreases in all
renal layers which exceeded those observed for aortic occlusion, with decreases of 33 ± 2%, 39 ± 2%, and 32 ± 6% in the CO, OM, and IM (Figure 2C–E). $T_2^*$-derived KS increased by 7 ± 1% and returned to baseline upon release (Figure 2B). The $T_2^*$ decreases in CO, OM, and IM were larger than those in $T_2$ (60 ± 3%, 60 ± 2%, and 58 ± 4%; Figure 2C–E), and much larger than the $T_2^*$ decreases during aortic occlusion. Restoration of $T_2$ and $T_2^*$ toward baseline upon release of venous occlusion occurred somewhat slower than following aortic occlusion. Changes in $T_2$ showed moderate negative correlations with changes in KS, while negative correlations with changes in $T_2^*$ were moderate in the CO, and weak in the OM and IM (Table 1).

Simultaneous occlusion of both the suprarenal aorta and the renal vein did not affect KS; yet there was a small drop in $T_2^*$-derived KS about 1 min after release of the occlusion (Figure 3B). Simultaneous aortic and venous occlusion resulted in decreases in $T_2$ of 23 ± 1%, 28 ± 1%, and 20 ± 3% in the CO, OM, and IM (Figure 3C–E). Changes in $T_2^*$ were more pronounced than $T_2$ changes, though less dramatic than those observed with venous occlusion alone, with reductions of 46 ± 4%, 55 ± 2%, and 44 ± 4% in the CO, OM, and IM (Figure 3C–E). The $T_2$ and $T_2^*$ return to baseline were comparable to that upon release of aortic occlusion. Changes in $T_2$ and $T_2^*$ did not show any significant correlation with changes in KS (Table 1).

### 2.2 Hypoxemia

A brief period of hypoxemia was induced by lowering the inspiratory oxygen fraction (FiO$_2$) from 21% (normoxia)
Induction of hypoxemia reduced T₂-derived KS by 2 ± 1%, and T₂ in CO, OM, and IM by 18 ± 3%, 15 ± 3%, and 8 ± 3% (Figure 4). T₂*-derived KS was reduced by 3 ± 1%, and T₂* was reduced by 23 ± 4%, 29 ± 4%, and 25 ± 5% in the CO, OM, and IM. T₂ changes correlated with changes in KS moderately in the OM, and weakly in the CO and IM; T₂* changes correlated weakly only in the CO (Table 1).

### 2.3 | X-ray CM iodixanol

Bolus injection of the CM, iodixanol, into the thoracic aorta induced a sustained increase in KS, as determined from T₂ and T₂* maps, with peaks of 10 ± 2% and 8 ± 2% about 6 min after the injection (Figure 5B). T₂ increased by 14 ± 3% and 17 ± 5% in the CO and OM immediately after CM injection, then normalized (Figure 5C–E). In the IM, T₂ decreased by 17 ± 7% about 17 min after CM and remained below baseline for the duration of the observation. T₂* showed an initial increase of 21 ± 3% and 24 ± 5% in the CO and OM immediately after CM, then decreased below baseline levels within about 16 min. In the IM, T₂* decreased by 33 ± 11% by about 16 min and remained below baseline. T₂ changes in the CO and OM were strongly correlated with changes in KS; T₂* changes in the CO showed a weak correlation (Table 1).

### 2.4 | Furosemide

Furosemide induced an increase of up to 4 ± 1% in T₂-derived KS (Figure 6B) and increased T₂ in the CO and OM by up to 11 ± 2% and 19 ± 2%; changes in the IM were not significant (Figure 6C–E). T₂*-derived KS changes did not reach statistical significance (Figure 6B), T₂* in the CO and OM increased by up to 6 ± 1% and 28 ± 4%; T₂* changes in the IM were not significant (Figure 6C–E). T₂ changes in the CO and OM showed strong correlations with changes in KS; T₂ changes in the IM and T₂* changes in the OM showed weak correlations (Table 1).

Urethane anesthesia provided stable systemic hemodynamics throughout all experiments as monitored by arterial pressure (Table S1).49

### 2.5 | Biophysical model

Using the biophysical model described in the “Methods” section, we estimated changes in O₂ saturation of Hb (Sat)
from measured changes in T₂* and in kidney size for the three vascular occlusions. According to Equation (4) of the biophysical model, these changes are expressed by the ratio of (1 – Sat)_{occlusion}/(1 – Sat)_{before occlusion}. The average (1 – Sat)/(1 – Sat₀) ratio in all renal layers, for all three occlusions, overall occlusion time points was approximately 2.1 (range 1.9–2.9, Table 2). On average, a relative 2.1-fold increase in the proportion of deoxyHb was found.

3 | DISCUSSION

Renal tissue hypoxia occurs very early in most forms of AKI and is a key feature in the progression to CKD and in diabetic kidney disease. MRI offers non-invasive full coverage of the kidney, and the MRI relaxation times T₂*, T₂ appear to be ideal surrogate markers of renal oxygenation. However, the relationship between renal tissue pO₂ and T₂*, T₂ is confounded by changes in hemocrit, the O₂ affinity of hemoglobin, and crucially, the blood and tubular volume fractions. Here, we performed serial MR-based measurements of kidney size and T₂*, T₂ during clinically realistic interventions in rats, directly while they were in the MR scanner, to examine this relationship. Our results demonstrate that monitoring of KS allows physiological interpretation of acute renal oxygenation changes obtained by T₂*, T₂.

Several surgical procedures (e.g., partial nephrectomy) require cross-clamping of the renal artery or suprarenal aorta, the renal vein, or simultaneous occlusions of both. If maintained for too long, such occlusions risk renal ischemia–reperfusion injury. At the onset of occlusions, renal tissue perfusion and O₂ delivery rapidly diminish, but O₂ consumption by FIGURE 2 Time courses during occlusion of the renal vein and recovery. (A) Exemplary T₂,T₂* maps obtained for a rat kidney in vivo. Time course of relative changes for (B) kidney size and T₂,T₂* for (C) CO, (D) OM, and (E) IM. Colors, absolute baseline values, and significance signs as in Figure 1.
active tubular transports continues, leading to a rapid and massive decline in tissue pO₂ and O₂ saturation of hemoglobin in intrarenal blood.45 Our previous studies with invasive probes (the gold standard) showed an equivalent decrease in tissue pO₂ and O₂ saturation of hemoglobin following both venous occlusion and aortic occlusion.27,45,53 However, in the present study, we observed that decreases in T₂*, T₂ following venous occlusion were much more pronounced than for aortic occlusion. The reason for this discrepancy is that the changes in T₂*, T₂ reflect changes in the blood volume fraction in response to the occlusions, that is, changes in the amount of deoxyHb per tissue volume, rather than directly mirroring O₂ saturation of hemoglobin. Upon aortic occlusion, the inflow of blood into the kidney is abruptly stopped while outflow via the renal vein continues, until pressures in intrarenal vessels and the vena cava are equalized. This reduces intrarenal blood volume and deoxyHb.45,53 Conversely, upon venous occlusion, the outflow of blood is abruptly stopped, while the inflow via the renal artery continues until the distension of intrarenal vessels is counterbalanced by the resistance of the renal tissue and the relatively rigid capsule, leading to an increase in intrarenal blood volume and deoxyHb.45,53 Consequently, renal oxygenation assessments by T₂*, T₂ alone will overestimate tissue hypoxia during venous occlusion and underestimate it during aortic occlusion. Simultaneous aortic and venous occlusion lowers tissue pO₂ and O₂ saturation of hemoglobin by a comparable degree to aortic occlusion and venous occlusion but does not change the blood volume fraction; accordingly, the decrease in T₂*, T₂ was less than for venous occlusion and more than for aortic occlusion, as illustrated qualitatively in Figure 7.
Changes in renal blood volume in response to these interventions were associated with parallel changes in KS. Upon aortic occlusion, KS decreased, and this change was correlated with changes in T2*, T2 in all renal layers. Conversely, upon venous occlusion, KS increased, and this was negatively correlated with changes in T2*, T2. With simultaneous aortic and venous occlusion, there was no KS change, and no correlation with T2*, T2 changes (Figure 7). Thus, physiological interpretation of T2*, T2 as surrogate markers for renal tissue oxygenation must take into account changes in KS. If T2*, T2 decrease and KS remains unchanged, tissue oxygenation is reduced. If T2*, T2 decreases, and KS also decreases, the reduction in tissue oxygenation is more severe than if KS is unchanged; if T2*, T2 decreases, and KS increases, the reduction in tissue oxygenation is less severe.

Clinical scenarios with decreased hematocrit or reduced pulmonary O2 diffusion lead to arterial hypoxemia, and the risk of AKI. We induced arterial hypoxemia, thus reducing renal O2 supply. The decrease in blood pO2 is attenuated by arterial chemoreceptor-actuated increase in ventilation. On the other hand, enhanced breathing reduces blood pCO2 and increases blood pH, which increases the O2 affinity of hemoglobin so that it releases less O2 in the microcirculation. The ensuing decrease in renal tissue pO2 is milder than that during the vascular occlusions, and the observed reduction in T2*, T2 are more subtle. The T2*, T2 decrease was accompanied by a KS reduction, which is related to hypoxemia-induced extra-renal vasodilation, resulting in a drop in arterial pressure and an ensuing decrease in renal arterial inflow. Thus, T2*, T2 and KS measurements reveal even subtle changes in renal tissue oxygenation.
X-ray CM can induce AKI, especially at large doses administered for cardiac interventions. This is the result of several mechanisms that lead to renal hypoperfusion and hypoxia, including fluid viscosity-induced increase in intratubular pressure, resulting in intrarenal compartment syndrome. We observed an initial increase in $T_2^*$, $T_2$ upon iodixanol injection in the CO and OM, followed by a decrease toward baseline levels for $T_2$ and even lower levels for $T_2^*$. While these $T_2^*$, $T_2$ changes are consistent with a previous MRI study, they differ from results we previously obtained with invasive pO2 probes. Using the same experimental paradigm as in the present study, we observed an immediate and massive drop in pO2 upon CM, that was sustained for 60 min. The explanation for this apparent discrepancy becomes clear when we note that KS increased upon CM injection, peaking within 6 min, and remained enlarged throughout the observation period. This reflects the compartment syndrome that results in compression of the intrarenal vessels and thus decreased deoxyHb (Figure 7). While the initial $T_2^*$, $T_2$ increase in CO and OM appears to reflect improved tissue oxygenation, it in fact deteriorates, as demonstrated with the pO2 probes, and indicated here by the KS change. The sustained renal enlargement indicates continued compartment syndrome, and thus the apparent return of $T_2$ toward baseline does not in fact reflect normalization of tissue oxygenation, and the $T_2^*$ decrease below baseline greatly underestimates the degree of hypoxia. These results underscore how MR-based assessment of renal oxygenation by $T_2^*$, $T_2$ is crucially dependent on monitoring accompanying changes in KS.

Furosemide has long been used in the clinic to increase urine flow rate and sodium excretion, and a furosemide “stress test” is suggested to predict renal
Furosemide decreases resorption in the thick ascending limb of the loop of Henle, resulting in increased tissue pO₂, particularly in the OM, with reported increases in T₂*, T₂. While our present T₂*, T₂ results agree with these studies, our observation of a small KS increase indicates that this is not solely the result of improved oxygenation. Rather, the increased KS is likely due to increased tubular fluid volume in the distal nephron, resulting in a mild form of compartment syndrome, again illustrating how T₂*, T₂ overestimates tissue oxygenation if not adjusted by KS assessments.
Measurements of acute changes in KS alone can not differentiate between changes induced by renal blood volume versus tubular volume changes. However, this distinction will often be clear from the specific intervention performed in preclinical experiments, and will also be obvious in many clinical scenarios with acute KS changes. Furthermore, advanced MR methodology supports monitoring of acute changes in the tubular or in the blood volume fraction using diffusion-weighted imaging. The quantitative correction factors obtained for T_2* and T_2 (slope and intercept of the linear regressions with KS) for the specific interventions are only valid for the present experimental setting. Although similar qualitative relationships between acute changes in T_2*, T_2 and KS will exist for comparable acute interventions in preclinical and clinical studies, specific quantitative correction factors will naturally depend on the particular experimental or clinical setting including the magnetic field strength, and species. Chronic kidney diseases, especially those with fibrotic alterations, will greatly affect the quantitative relationship between acute changes in T_2*, T_2 and KS. Our calculations for the three vascular occlusions demonstrate that the changes in O\textsubscript{2} saturation of Hb (Sat) can be extracted from measured changes in T_2* and in kidney size by use of a biophysical model.

Our data showed a (1 − Sat)/(1 − Sat\textsubscript{0}) ratio of ≃2.1, averaged overall renal layers, all three occlusions, and over all occlusion time points, indicating a relative 2.1-fold increase in the proportion of deoxyHb. To the best of our knowledge, to date the literature reports only Sat data for the cortex, but not for the outer and inner medulla of rats. Estimates based on invasive near-infrared spectroscopy suggest a baseline cortical blood Sat of approximately 65%. The average (1 − Sat)/(1 − Sat\textsubscript{0}) ratio of 2.1 derived from our model calculations corresponds to a Sat decrease of about 60%. Assuming a baseline cortical Sat of 65%, the occlusion-induced Sat could be as low as 26%, which according to the oxyHb dissociation curve of rats, would be equivalent to a blood pO\textsubscript{2} of about 22 mmHg. This is consistent with tissue pO\textsubscript{2} data we previously obtained using invasive probes. During aortic or venous occlusions, cortical tissue pO\textsubscript{2} decreased by 80–90%, reaching values <4 mmHg. This congruence indicates that the biophysical model yields physiologically plausible calibration ratios and Sat values.

The biophysical model facilitates quantitative assessment of relative changes in Sat from relative changes in renal T_2* and KS. It provided physiologically plausible values for the specific setup used in our preclinical study as a mandatory precursor to clinical studies. Due to its non-invasive nature, our approach suits swift

![Diagram](https://example.com/diagram.png)
translation from pre-clinical research to human studies. It is very much conceivable that MR-based estimates of relative changes in Sat may ultimately become a diagnostic biomarker. However, a number of prerequisites must be fulfilled to meet this goal. While our study obtained serial MR data in the same rats (intraindividual time courses) before the intervention (baseline) and during the acute intervention, this will be barely routine or practical in a typical clinical setting. To address this difference between our preclinical study and clinical reality, it is essential to obtain age, BMI, sex, and magnetic field strength corrected normal reference values for renal $T_2$* and KS in healthy humans using standardized MRI protocols. A similar approach has been used for myocardial $T_2$* mapping, which is now very well established for the quantitative assessment of tissue iron content and for the therapy of iron overload disorders.\textsuperscript{63–67}

It stands to reason that the normal reference values of renal $T_2$* and KS can be deduced from large population imaging studies such as the German National Cohort or the UKBiobank.\textsuperscript{68,69} Using these standardized MR protocols in acute clinical scenarios, assessment of the deviation of the relationship between $T_2$* and KS of individual patients or patient groups from the normal reference obtained for healthy subjects would allow quantitative estimation of alterations in $O_2$ saturation of hemoglobin.

Our biophysical model assumes deoxyHb to be the dominating factor. This assumption applies very well to $T_2$* which reflects the amount of deoxyHb per tissue volume. Its reciprocal value $R_2$* is directly proportional to the fraction of deoxyHb ($=1 – \text{Sat}$) and the blood volume fraction ($= \text{blood volume/kidney volume}$).\textsuperscript{70} $T_2$ is a physical constant for perfused tissue. Its reciprocal value $R_2$ scales linearly with blood oxygenation.\textsuperscript{71} $R_2$ includes contributions other than magnetic susceptibility. Modeling and calibration involved in converting $T_2$ into Sat require further experimental studies.\textsuperscript{72} This calibration should include renal $T_2$ mapping during hyperoxia (100% inspiratory $O_2$) to distinguish $T_2$ contributions which are not related to magnetic susceptibility from those governed by the amount of deoxyHb per tissue volume.\textsuperscript{71} Upon successful calibration of renal $T_2$ versus Sat our biophysical model can be refined and applied for renal $T_2$ oximetry, which will be our next target.

Previous studies showed that MR-based assessment of KS complements other markers for diagnosis and staging of kidney disorders. Here, we demonstrate that KS monitoring is essential for the physiological interpretation of acute changes in renal tissue oxygenation derived from $T_2$*, $T_2$. As KS can be readily obtained from $T_2$*, $T_2$ maps without the need for additional scans, this should always accompany the assessment of MRI-derived oxygenation results. Driven by technical advances including simultaneous $T_2$* and $T_2$ mapping,\textsuperscript{61,73} renal MR oximetry can greatly support preclinical studies into the mechanisms of renal pathophysiology. Moreover, this non-invasive approach to probing renal oxygenation holds the promise of swift translation to human studies, for example, for the assessment of drug effects, and for clinically meaningful diagnosis. First steps toward this include adaptation of the MRI protocol for simultaneous KS and $T_2$, $T_2$ measurements, and reversible test interventions applicable to human beings.

4 | METHODS

4.1 | Animal preparation

Investigations were approved by the LaGeSo of Berlin in accordance with German Animal Protection Law and EU Directive 2010/63/EU. Male Wistar rats ($n = 37$, aged 12–13 weeks, 270–300 g, Harlan-Winkelmann, Borchen, Germany) were studied. An intraperitoneal dose of urethane (0.2 g/ml; 6 ml/kg, Sigma-Aldrich, Steinheim, Germany) was used as anesthesia throughout surgeries and MRI examinations. Urethane provides long-lasting anesthesia and has the least effects on cardiovascular and respiratory control compared to other anesthetics.\textsuperscript{49} Preparation included insertion of vascular catheters and probes for measurements of hemodynamics and oxygenation.\textsuperscript{41,46,53,74} In a subgroup ($n = 13$), two MR-safe remotely controlled inflatable occluders were applied around the suprarenal aorta and the left renal vein.\textsuperscript{45–47} Thereafter, rats were transferred into the MR scanner. They were spontaneously breathing and continuously provided with air (1 L/min). Body temperature was maintained at 37°. A balloon on the thorax was used for respiration-triggered MR data acquisition.\textsuperscript{45,46}

4.2 | MRI experiments

MRI data were acquired on a 9.4 Tesla small animal MR system (Bruker Biospec 94/20, Bruker Biospin, Ettlingen, Germany) using a linear radiofrequency volume resonator and a 4-channel surface coil array tailored for rats (Bruker Biospin).\textsuperscript{37} For geometrical planning and slice positioning, $T_2$-weighted pilot scans were acquired. Local volume selective magnetic field shimming was done on an ellipsoid accommodating the left kidney using an automatic optimization algorithm based on free induction decay length. $T_2$* and $T_2$ mapping were performed with respiration-gated protocols. Details of the MRI parameters are listed in Table 3.
4.3 | Image analysis

Parametric maps of absolute $T_2^*$ and $T_2$ were calculated by pixel-wise mono-exponential fitting to the signal intensities of the $T_2^*$- and $T_2$-weighted images acquired at different echo times.\textsuperscript{37} Median $T_2^*$ and $T_2$ values for regions-of-interest (ROI) within the renal cortex (CO), outer medulla (OM), and inner medulla (IM) were calculated from the parameter maps. ROI placement was done with a standardized semi-automatic method, as previously described.\textsuperscript{75} This procedure positions the ROIs (5 for CO and OM each, 3 for IM) such that they exclude the transition regions between renal layers to avoid partial volume effects. For $T_2^*$, $T_2$ mapping-based determination of KS, segmentation of the coronal mid-slice cross-sectional area of the kidney (here referred to as “kidney size,” KS) was done using a previously described automatic bean-shaped model.\textsuperscript{37}

4.4 | Longitudinal quantification of changes in kidney size and oxygenation upon pathophysiological interventions

To investigate the relationship between changes in $T_2^*$, $T_2$ and KS, six pathophysiologically relevant interventions that alter renal tissue oxygenation reversibly (brief occlusion of the suprarenal aorta, the left renal vein or both; hypoxemia), and with longer-lasting effects (injection of CM or furosemide) were used. In addition to their effects on oxygenation, occlusion of the aorta results in decreased renal blood volume, occlusion of the renal vein induces an increase in renal blood volume, and simultaneous occlusion of both vessels does not affect renal blood volume. Administration of CM is expected to increase the tubular volume fraction concomitant with its effect on oxygenation.

Rats equipped with vascular occluders underwent a series of interleaved $T_2^*$ and $T_2$ mappings (short: MR scans) prior to the occlusions (control period), during occlusions, and following the release of occlusions. The aorta was occluded for 3.8±0.3 min ($n = 13$ rats; time depending on respiration gating). The occluder was then deflated, and the animals were allowed to recover for at least 7 min to ensure complete restoration of pre-occlusion hemodynamics and oxygenation. Time of flight-based MR angiography was performed immediately after inflation/deflation of the occluder to confirm occlusion/reperfusion of the vessels.\textsuperscript{45-47} After recovery from the aortic occlusion, the same procedure was performed for renal venous occlusion ($n = 12$), and subsequently for simultaneous combined aortic and venous occlusion ($n = 10$).

In the second subgroup ($n = 11$), rats underwent MR scans during a control period of normoxia with an FiO$_2$ of 21%, during a short period (3.8±0.1 min) of hypoxia (FiO$_2$ = 10%) and 10 min of recovery (FiO$_2$ = 21%). The FiO$_2$ was monitored as previously described.\textsuperscript{45,48}

In the third subgroup ($n = 8$), rats underwent MR scans before (control) and following a 1.5 ml bolus of iodoxanol solution (320 mg/ml iodine, GE Healthcare Buchler, Braunschweig, Germany) injected into the thoracic aorta, followed by 0.2 ml saline chaser, as previously described.\textsuperscript{40,41}

In the fourth subgroup ($n = 5$), rats underwent MR scans before (control) and following an i.v. bolus of furosemide (5 mg/kg, ratiopharm GmbH, Ulm, Germany) followed by a 0.2 ml saline chaser.

4.5 | Statistical analysis

Data were evaluated for Gaussian distribution using the Shapiro–Wilk test. Relative intervention-mediated

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**Table 3** Details of MRI protocols used for $T_2^*$ and $T_2$ mapping

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<th>Method</th>
<th>$T_2^*$-mapping</th>
<th>$T_2$-mapping</th>
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<td>Repetition time TR (ms)</td>
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<td>Multi spin-echo MRI</td>
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<td>Number of echoes</td>
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<td>Inter-echo time ΔTE (ms)</td>
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<td>Excitation flip angle ($^\circ$)</td>
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<tr>
<td>Acquisition time $t_{eq}$ (s)</td>
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<tr>
<td>In-plane spatial resolution w/o zero filling ($\mu$m$^2$)</td>
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<td>Field of view (mm$^2$)</td>
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<tr>
<td>Matrix size</td>
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<td>Slice thickness (mm)</td>
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changes in KS and $T_2^*$, $T_2$ were analyzed by the non-parametric repeated-measures Friedman test, followed by Dunn’s post hoc test with the Benjamini-Hochberg correction for multiple comparisons. Correlations between relative changes in KS and $T_2^*$, $T_2$ were assessed using repeated-measures correlation. Data were analyzed using R v.3.6.3 with the packages “rstatix,” “dunn.test,” and “rmcorr.” $p < 0.05$ was considered significant.

### 4.6 Biophysical model

To evaluate the quantitative features of the observed $T_2^*$-based signal changes and the relative changes in KS, we used a model to extract changes in O$_2$ saturation of Hb (Sat) from measured changes in $T_2^*$ and in KS for the three interventions involving vascular occlusions.

$T_2^*$ reflects the amount of deoxygenated Hb per tissue volume. Its reciprocal value $R_2^*$ is proportional to the fraction of deoxygenated Hb ($=1-\text{Sat}$) and the blood volume fraction ($BVF = \text{blood volume } [BV]/\text{kidney volume } [KV]$).

$$R_2^* \sim (1 - \text{Sat}) \frac{BV}{KV}$$  (1)

For the model, we assume that all changes in KV ($\Delta KV$) during the vascular occlusions are caused by blood volume changes ($\Delta BV$).

$$\Delta BV = \Delta KV$$  (2)

With this assumption, the ratio of $R_2^*$ obtained during the occlusions versus $R_2^*$ observed for baseline conditions ($R_2^*_{0}$) prior to the occlusion can be expressed as:

$$\frac{R_2^*}{R_2^*_{0}} = \frac{(1 - \text{Sat})_{0} KV_0 (BV_0 + \Delta KV)}{(1 - \text{Sat})_0 BV_0 (KV_0 + \Delta KV)}$$  (3)

Rearranging the ratio of the deoxygenated Hb fractions and substituting changes in the blood volume fraction ($BVF = BV/KV$) leads to:

$$\left(1 - \frac{\text{Sat}}{\text{Sat}_0}\right) = \frac{R_2^*}{R_2^*_{0}} \frac{BVF_0}{BVF_0 + \frac{\Delta KV}{KV}}$$  (4)

Equation (4) relates the $R_2^*$ ratio to the ratio of the deoxygenated Hb fractions using the baseline blood volume fraction and the relative kidney volume change as correction factors.

Assuming (i) that the O$_2$ saturation of Hb at baseline and the degree of its decrease during the occlusions do not differ among the three occlusions, and (ii) that the KV changes are uniform across the kidney, we can estimate the baseline $BVF_0$ for the three renal layers. This estimation permits the calibration of Equation (4) to convert $R_2^*$ ratios to ratios of $(1 - \text{Sat})$.

We measured kidney size by planimetry of the mid-slice cross-sectional area ($A$). We converted this into kidney volume under the assumption that changes in the third dimension are similar to changes in the two measured dimensions:

$$\frac{\Delta KV}{KV_0} = \left(\frac{\Delta A}{A_0} + 1\right)^{1/3} - 1$$  (5)

The calibration was done by minimizing the variance of the resulting $1 - \frac{\text{Sat}}{\text{Sat}_0}$ on the average deviation among the three occlusions. The calculated calibration factors $BVF_0$ were 0.268, 0.394, and 0.273 for CO, OM, and IM, respectively.

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### CONFLICT OF INTEREST

All authors declare no competing interests.

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### REFERENCES


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