Blood flow synchronization in renal microcirculation - a high-resolution imaging study.

Dmitry D. Postnov¹,²*, Donald J. Marsh³, Will A. Cupples⁴, Niels-Henrik Holstein-Rathlou², and Olga Sosnovtseva²

¹Department of Clinical Medicine, Faculty of Health, Aarhus University, Aarhus, Denmark
²Biomedical Sciences Institute, Faculty of Health and Medical Sciences, Copenhagen University, Copenhagen, Denmark
³Division of Biology and Medicine, Brown University, Providence, Rhode Island, USA
⁴Department of Biomedical Physiology and Kinesiology, Simon Fraser University, Burnaby, British Columbia, Canada
* dpostnov@cfin.au.dk

ABSTRACT

Aims
Internephron signalling and interaction are fundamental for kidney function. Earlier studies have shown that nephrons signal to each other over short distances and adjust their activity accordingly. Micropuncture experiments revealed synchronous clusters of 2-3 nephrons formed from such interactions, while imaging and modelling results suggested the possibility of larger clusters. Such clusters are expected to play an important role in renal autoregulation, but their presence has not been confirmed and their size has not been estimated. In this study, we present methodology for high resolution renal blood flow imaging and apply it to estimate frequency and phase angle differences in kidney blood vessels under normal conditions and after administration of the vasoactive agents angiotensin II and acetylcholine.

Methods and results
To resolve signals from separate arterioles in a sufficiently large field of view, we developed a method for renal laser speckle contrast imaging. Our setup provides imaging of blood flow in the kidney cortex with a limit of image resolution at 0.8 µm per pixel and imaging frequency of 160 Hz. We used the method to record from 1.5x1.5 mm² sections of the renal surface in anaesthetised Sprague-Dawley rats in unstimulated conditions and during IV infusion of the vasoconstrictor angiotensin II or the vasodilator acetylcholine. In each section, we resolved and segmented 94.8 ± 15.66 individual arterioles and venules, and analyzed blood flow using wavelet spectral analysis to identify clusters of synchronized blood vessels.

Conclusions
We observed spatial and temporal evolution of blood vessel clusters of various sizes, including the formation of large (>90 vessels) long-lived clusters (>10 periods) locked at the frequency of the tubular glomerular feedback (TGF) mechanism. The analysis showed that synchronization patterns and thus the co-operative dynamics of nephrons change significantly when either of the vasoactive agents is administered. On average, synchronization was stronger (larger clusters, longer duration) with angiotensin II administration than in the unstimulated state or with acetylcholine. While it weakens with distance, increased synchronization duration spanned the whole field of view, and likely, beyond it. Neighbouring vessels tend to demonstrate in-phase synchronization, especially in the vasoconstricted condition, which is expected to cause locally increased pressure variation. Our results confirm both the presence of the local synchronization in the renal microcirculatory blood flow and the fact that it changes depending on the condition of the vascular network and the blood pressure, which might have further implications for the role of such synchronization in pathologies development.

1 Introduction

The kidney represents a unique demand-driven, interconnected resource distribution network that is responsible for body homeostasis maintenance over a broad range of conditions, including variations in blood pressure and fluid intake and loss. With blood serving as the resource, single-nephron autoregulation mechanisms provide and regulate the demand. Based on measurements of tubule pressure responses to step changes in arterial pressure¹, vascular transfer functions²–⁷, and renal blood flow response to arterial pressure forcing⁸, two critical mechanisms in renal pressure autoregulation, the myogenic mechanism and tubuloglomerular feedback (TGF), have emerged as the critical components. Their actions combine to regulate blood flow, serving to maintain the delivery of water and solutes to various regions of the nephron at levels appropriate to their dynamic ranges. The two mechanisms operate at different time scales, generating spontaneous blood flow and pressure oscillations at different frequencies: 5-10 seconds (0.1-0.2 Hz) for the myogenic response and 30-50 seconds (0.02-0.033 Hz) for the
TGF\(^4,9\text{-}12\).

Nephrons, however, do not operate as stand-alone units. Within a single kidney all nephrons are linked via the renal vascular tree, which provides connections of different proximity - from few hundred microns for neighbouring nephrons separated only by their respective afferent arterioles; to nephrons only connected at the level of the renal artery, which plays a role of a single supply source for all the nephrons. Nephrons nested in such a network are bound to communicate and affect each other to some degree. In addition to interaction through the blood flow and pressure, nephrons were found to communicate via electrical signalling. Such interactions are proven to play a critical role in kidney function and can lead to complex co-operative dynamics and synchronization between nephrons.

Micropuncture experiments\(^13\text{-}16\) showed that neighbouring nephrons (originating from a common artery) adjust their TGF-mediated tubular pressure oscillations to attain a synchronised regime. Although these experiments confirmed the existence of synchronisation, only pairs or triplets of nephrons could be sampled at any one time. Assessment of cooperative efforts of a larger number of nephrons required a different approach. To address this challenge, several groups adopted laser Speckle Contrast Imaging (LSCI)\(^17\). In LSCI, media with moving light scattering particles, e.g. red blood cells, are illuminated with a near-infrared laser. The backscattered light, recorded by a camera, forms an interference pattern, which appears more or less blurred depending on the speed of the particles. This pattern is then analysed to obtain qualitative maps of particle velocity and thus a blood flow estimate\(^18,19\). First applied by Holstein-Rathlou et al. to map TGF oscillations over a large field of view, this method was later used to explore periodic activity in the myogenic frequency band\(^20\), analyse spatial correlations in the renal blood flow\(^21\), and study intra-renal drug distribution\(^22,23\). Although these results encouraged the large-scale synchronisation hypothesis, the method lacked resolution, both spatial and temporal, as well as signal-to-noise ratio, to confirm it convincingly.

In this paper, we further advance renal blood flow imaging methodology and confirm the presence of synchronised clusters spanning multiple nephrons for the first time at the level of individual arterioles and venules. Our LSCI setup and data processing approach allow imaging renal microcirculation with at 0.8 \(\mu\)m per pixel spatial resolution and imaging frequency to 160 Hz for 1024x1024 pixels. We apply it to study synchronous cluster formation.

2 Methods

2.1 Animal preparation
All experimental protocols were approved by the Danish National Animal Experiments Inspectorate and were conducted according to the American Physiological Society guidelines. Male Sprague Dawley rats (Taconic, Denmark) with average weight \(\approx 290\) g (n=5) were used. Before starting surgical procedures, animals were anaesthetized in a chamber with 8% sevoflurane. During the surgery, sevoflurane concentration was reduced to a final concentration of \(\approx 2\%\). Two catheters were inserted in the right jugular vein to allow continuous systemic infusion of drugs and saline. Another catheter was inserted in the carotid artery to measure mean arterial pressure with a pressure transducer (Statham P23-dB, Gould, Oxnard, CA). Then tracheotomy was performed, after which the rat was placed on a servo-controlled heating table maintaining body temperature at 37 \(\degree\)C and connected to a mechanical animal ventilator (60 breaths/min; 8 ml/kg bodyweight). To avoid secondary heartbeat and breathing artefacts, Nimbex (muscle relaxant, Sigma) was administered in a concentration of 0.85 mg/ml, first as a bolus injection of 0.5 ml, followed by a continuous intravenous infusion at a rate of 20 \(\mu\)l/min. The left kidney was then exposed, and the left ureter was catheterized to ensure free urine flow. To reduce motion artifacts and avoid drying the kidney surface during the experiment, we placed the kidney in a plastic fixation holder, covered it with warm agarose solution (1% Agarose, Sigma, 99% saline) and put a thin (0.1mm) cover glass on top of the kidney. Metal thread (40 micrometres in diameter) was bent in a “U” shape and positioned on top of the cover glass at the flattest location of the kidney surface, marking the region of interest for imaging procedures. Following the surgical procedures, the animal was left to stabilize for 20 minutes. Experiments were continued only if the mean arterial pressure remained within 100–120 mmHg during the control period. At the end of the experiment, animals were euthanized by overdose of sevoflurane, followed by cervical dislocation.
Figure 1. Schematics of the imaging setup and blood flow data example. (A),(B) Top and side view of the experimental setup and its key components. (C) Normalized flow map with microcirculatory vessels. Yellow and blue colours correspond to high and low blood flow, respectively. (D) Relative blood flow dynamics in four vessels, which are outlined in (C) with corresponding colours. (E) Dominant frequency in the TGF band of the blood flow oscillations is shown in (D). (F) Phase value at the dominant frequency. One can see that there are two vessels pairs in which blood flow is synchronized for more than 90% of time. Synchronization between the pairs is also well observed but breaks down when TGF activity change its frequency at 850–1050 s). These exemplary graphs reflect the data.

2.2 Laser Speckle Contrast Imaging data acquisition
To assess microcirculation in the kidney cortex, we built a high resolution laser speckle contrast imaging setup. A single-mode fibre-coupled laser diode (785nm, LP785-SF100, Thorlabs, USA) controlled with a laser driver, and temperature controller (CLD1011LP, Thorlabs, USA) was used to deliver coherent light onto kidney surface with a power density of approximately...
10mW/cm², providing an optimal signal to noise ratio. Backscattered light was collected by a zoom imaging lens (VZM 1000i, Edmund Optics) at 5x magnification and recorded with a CMOS camera (Basler acA2000-165umNIR, 2048x1088 pixels, 5.5 µm pixel size) at an exposure time of 5 ms. A subset of 1024x1024 pixels was used for recording, from a 1.5x1.5mm region of the renal surface. In addition, a linear polarizing filter was placed in front of the objective to reduce artefacts from reflected light. While the imaging set up in this configuration allows imaging resolution of ≈ 0.8µm/px at the frame-rate of 160 frames per second (fps), in our experiments, we found it optimal, in terms of the field-of-view and data storage, to acquire images at 50 fps and resolution of ≈ 1.5µm/px.

Imaging was performed in a non-stimulated state (control) and following administration of angiotensin II (AngII), and acetylcholine (ACh), respectively. After collecting 20 min (2400 frames) of the baseline data, we initiated continuous administration of AngII at a concentration of 4 ng/ml and an infusion rate of 20 µl/min to cause systemic vasoconstriction. The infusion lasted for 30 min, out of which the first 10 min were allocated for blood flow to stabilize, and the following 20 minutes were recorded. Fifteen min after completion of the AngII infusion, we infused ACh at a concentration of 0.0375mg/ml and a rate of 20 µl/min, first permitting 10 minutes for blood flow to stabilize, and then recorded subsequent 20 minutes. Across all experiments, average arterial pressure was 112±2, 127±6 and 100±4 during the control, vasoconstricted (AngII) and vasodilated (ACh) conditions respectively.

2.3 Data analysis

Image registration. To allow high-resolution laser speckle contrast imaging of the renal microcirculation, we needed to reduce motion artefacts, as any lateral motion larger than 5-10 micrometres will prevent accurate estimation of the blood flow and further segmentation of microcirculatory vessels. Unlike brain imaging, when working with the kidney, there is no bone tissue that can be fixed to reduce respiratory motion, and applying even slight pressure on the kidney might result in abnormal blood flow due to the occlusion of the small vessels on its surface. At the same time, raw laser speckle images, or contrast images without temporal averaging, are not suitable for automated registration due to the absence of clear intensity landmarks. To resolve this issue, we placed a "U" shaped metal marker on the cover glass, which moves along with the kidney, as described above. As the first step of analyzing the data, the marker is segmented in all frames via thresholding and then used to estimate the translation type geometrical transformation required to register images. Estimated geometrical transformation is then applied to the raw laser speckle images prior to performing the contrast analysis.

Contrast analysis. Registered laser speckle images were processed to calculate temporal contrast \( K = \frac{\sigma(I)}{<I>} \), where \( \sigma(I) \) and \( <I> \) are standard deviation and mean of pixel intensity over 25 frames. Contrast values were then converted to the blood flow index as \( BFI = 1/K^2 \), which are then used in the ensuing analysis.

Vessels segmentation. To segment individual microcirculatory vessels, we calculated averaged in time BFI images and applied adaptive thresholding (MATLAB) to them. Automated segmentation was followed by manual clean-up, where we removed artefacts and occasional large surface vessels. We then calculated blood flow dynamics for each segmented individual microcirculatory vessel by averaging BFI values in pixels belonging to this vessel.

Synchronization analysis. To study synchronization patterns between microcirculatory vessels and, thus, obtain insight into inter-nephron communication, we apply continuous wavelet transform analysis (Morse wavelet, MATLAB) to segmented vessels’ BFI. We identified the frequency and phase of dominant periodic activity in the 0.015-0.05Hz frequency band associated with the TGF mechanism. Vessels with less than 10% prominence of the activity peak were discarded and not used for synchronization analysis. In this study, we consider blood flow in different segmented vessels to be synchronized whenever their dominant frequencies match. To quantify blood flow synchronization over the field of view, we analyzed phase differences between synchronized vessels, average synchronization duration and its dependency on the distance between vessels, and the probability of the vessel’s blood flow to be synchronized with N% of the vessels in the field of view. To provide a "single-value" characterization of the synchronization at a given moment of time, we also introduced the synchronization degree parameter S:

\[
S(t) = \sqrt{L(t)/(N*(N-1))},
\]

where \( L \) is a number of frequency matching pairs of vessels, \( N \) is a total number of observed segmented vessels, and \( t \) is the time. \( S \) represents the relation of the observed number of frequency matching pairs to their maximum possible number. Thus \( S = 1 \) corresponds to all segmented blood vessels having the same dominant frequency, while \( S = 0 \) to all segmented blood vessels having a different dominant frequencies. However, it is important to notice that \( S = 0 \) is impossible to reach due to the discrete nature of the measured data and the analysis. In our case, the 0.015 to 0.05 Hz range is split into 18 fixed values, so that if there were more than 18 vessels, it became unavoidable for some of them to have an identical dominant frequencies. In practice, for 100 vessels with randomly chosen dominant frequencies, the minimum observed \( S \) would be \( \approx 0.25 \pm 0.06 \), which can be confirmed with a simple computational experiment.

Statistical analysis. Paired t-test was applied to compare results between control, AngII infusion and ACh infusion. P-values greater than 0.05 are reported as not significant. Results were expressed as mean ± standard deviation (SD) unless
Figure 2. Examples of instantaneous frequency patterns. Each circle location corresponds to a segmented vessel with color-coded frequency. Bottom panels show normalized blood flow in each segmented vessel over 5 minutes centered around the time corresponding to the frequency pattern snapshot. (A) High synchronization degree $S=0.98$ - large cluster covers the whole field of view. (B) Moderate synchronization degree $S=0.65$ - most of the vessels are split between two clusters. (C) Low synchronization degree $S=0.23$ - no clear clustering pattern, some vessels do not have pronounced TGF activity (frequency colour-coded as black). All data were captured in the same animal but under different conditions: control (A), AngII infusion (B), ACh infusion (C). Black-colored circles represent vessels where TGF activity was considered too weak (less than 10% prominence of the activity peak).
3 Results

3.1 Synchronization patterns
An example of a high-resolution blood flow map with segmented microcirculatory vessels is shown in Fig. 1.(c). Fig. 1.(d) shows RBF of four vessels outlined in (c) with corresponding colours (red, green, orange and black). From the corresponding dominant frequency and phase (Fig. 1.(e) and (f)), it can be seen that these vessels form two frequency-locked clusters (red-green and black-orange) that are synchronous most of the time. Moreover, there are long periods when these clusters synchronize, forming a larger cluster, interrupted with an asynchronous interval (from $\approx 850$ to $\approx 1050$ seconds).

We generalized this approach to the full-field blood flow imaging of the kidney surface. Figure 2 shows examples of high (A), moderate (B), and low (C) instantaneous synchronization degree, with $S = 0.97, 0.65,$ and $0.23$ respectively. In the first case, flow in almost all of the identified vessels oscillates at the same TGF frequency, forming a cluster of $>100$ vessels within the field of view. This cluster is likely to be even larger since it can spread outside the view field and in-depth in the kidney cortex. In the case of moderate synchronization degree - several frequency clusters with $\approx 10 − 60$ vessels can be identified, while for the low $S$ there are multiple groups of 2-3 vessels displaying the same frequency but no distinct pattern over the field of view. All three regimes were observed in the same animal during control, vasoconstricted (AngII infusion) and vasodilated (ACh infusion) conditions respectively. Averaged over the whole observation period and all animals ($N=5, 20$ min per condition) synchronization degree has moderate to high ($S = 0.54 \pm 0.09$) values during AngII infusion, low to moderate ($S = 0.36 \pm 0.1$) during ACh, and low to high in control ($S = 0.45 \pm 0.22$).

To visualize clustering tendencies, we calculate the probability of a randomly chosen vessel at any given moment of time to have the same dominant frequency as $X\%$ of the vessels in the field of view. From Fig. 3 it can be seen that infusion of AngII significantly (p<0.05) increases the prevalence of clusters covering 20-50$\%$ of the vessels - chances that a randomly chosen vessel belongs to such a cluster at any given moment are $\approx 50\%$, while for control and ACh infusion they are $\approx 20\%$. $N=5$ animals were used to create these graphs. Paired t-test was used to produce P-values. P-values smaller than 0.05 are considered to be significant and marked with "*".

3.2 Phase waves and spatial localization
Another distinct feature that we have observed is that the phase of oscillations within a cluster is space-dependent - it is mostly the same for the closely positioned vessels and gradually changes with distance. Figure 4 illustrates how phase within clusters evolves in space and time, forming the phase waves. Note that both (A) and (B) panels are from the same animal as was shown in Fig. 3 in control and vasoconstricted conditions, respectively, and that only vessels belonging to the largest cluster are shown. Change of the wave direction by $\approx 90^\circ$ in the same animal with constant vascular structure suggests different synchronization
centres where phase waves originate. Propagation speed is \( \approx 0.37 \) and \( 0.30 \text{ mm/s} \) for (A) and (B) respectively and spatial period of the phase waves is \( \approx 8.85 \) and \( 3.95 \text{ mm} \) (respective phase difference \( |\delta| = 0.71 \) and \( 1.59 \text{ rad} \) was observed over \( 1 \text{ mm} \) distance). The possibility of such phase-waves was previously hinted at in our earlier laser speckle contrast imaging study, where spatial correlations in renal blood flow were analysed\(^{21}\). See supplementary videos for more examples of phase waves dynamics.

**Figure 4.** Phase waves. (A) and (B) show the spatio-temporal evolution of TGF activity phase in vessels that belong to the largest cluster observed during control and AngII infusion. Both observations are from the same animal as shown in Fig. 2(A) and (B). Arrows indicate the wave direction.

Phase difference distribution (Fig. 5, A), calculated over all of the frequency matching vessels in all animals, shows the prevalence of in-phase synchronization (\( |\Delta| \leq \frac{\pi}{12} \), rad). This result is in good agreement with experimental micropuncture observations, where it is explained by the presence of fast electrical coupling acting over short distances. In all conditions, the larger phase differences are less prevalent, with the anti-phase synchronization (\( |\Delta| \geq \frac{11\pi}{12} \), rad) observed over just \( 1\% \) of time in control and during ACh infusion. However, AngII infusion increases this number to \( 5\% \), showing a statistically significant difference with other conditions. Phase difference grows with distance, as can be seen from Fig. 5 (B), reflecting the presence of phase waves. While change is relatively small in the control and vasodilated conditions (\( \approx 0.3 \) and \( 0.25 \text{ rad/mm} \)), it is strongly enhanced in the vasoconstricted condition, reaching, on average, \( \approx 1 \text{ rad} \) over \( 1 \text{ mm} \) distance. As we showed with mathematical modelling\(^{26}\), such difference can be explained by strengthened hemodynamic coupling, which the increased vascular tone should cause.
Figure 5. Localization of phase and synchronization in space. (A) Phase differences prevalence for synchronized vessels. (B) Phase differences distribution over the distance between vessels. (C) Synchronization duration normalized by the total observation time for different distances between vessels. N=5 animals were used to create these graphs. Paired t-test was used to produce P-values. P-values smaller than 0.05 are considered to be significant and marked with "*". Values between 0.05 and 0.1 are shown as $p<0.1$, where relevant, to highlight a trend in the data.

Since synchronization in renal blood flow is extended far beyond a pair of nephrons and is unlike synchronization in relatively homogeneous media, one would expect some space localization due to the topological features of the vascular network. Figure 5(C) illustrates how average synchronization duration changes with distance. It is clear that synchronization is stronger in the vasoconstricted condition, with average duration reaching $\approx 40 \pm 13\%$ of the observation time for neighbouring vessels and gradually reducing to $24 \pm 6\%$ for vessels located at more than 1 mm distance. In control, synchronization duration varies greatly, but on average, it also reduces with distance, although at a slower rate than during AngII infusion - from $\approx 30\%$ to $\approx 21\%$. During the ACh infusion the synchronization duration is $\approx 15 - 13\%$, with only 2% reduction over 1 mm of distance. Higher synchronization duration in control and during AngII infusion compared to ACh infusion suggests a long-distance nephron-to-nephron communication or a common driving force.

4 Discussion

In this study, we have designed a methodology for high-resolution blood flow imaging in renal microcirculation and applied it to study the synchronization of TGF oscillations in control, vasoconstricted (AngII infusion) and vasodilated (ACh infusion) conditions. Our data confirm that blood flow in renal microcirculation tends to demonstrate clustered, frequency-locked activity, with the clustering size and tendencies changing depending on the animal condition. Synchronization tends to be stronger, and cluster size is larger during AngII infusion, with synchronization degree $S = 0.54 \pm 0.09$ and average synchronization duration ranging from $\approx 40 \pm 13\%$ to $24 \pm 6\%$ of the observed time depending on the distance between vessels. During the ACh infusion, on the contrary, synchronization seems to be disrupted, with $(S = 0.36 \pm 0.1)$ and average synchronization duration $\approx 15 \pm 7 - 13 \pm 7\%$ of the observed time. Finally, in the normotensive condition, we observed mixed behaviour with highly variable synchronization degree $S = 0.45 \pm 0.22$ and duration ranging from $\approx 30 \pm 27\%$ to $21 \pm 21\%$. A possible explanation for the stronger synchronization during the AngII infusion will be a stronger hemodynamic coupling due to increased vascular resistance. It is also supported by the increased number of vessels synchronized in anti-phase during the infusion, which, as we predicted using mathematical modelling$^{26}$, is a natural consequence of stronger hemodynamic coupling.

We also observed phase waves that travel over the TGF-frequency clustered vessels (Figs 4, and 5(B)). Interestingly, the direction of the phase waves is not predetermined by the vascular structure - depending on the flow dynamics and other,
yet unknown factors, it can change even within the same animal (see Fig. 4). Such behaviour might be related to diffusive interaction between nephrons or the formation of synchronization centres (pacemakers) at different locations. The presence of phase waves supports the deterministic nature of synchronization rather than random entrainment at the same dominant TGF frequency. Similar effects are well studied in excitable media with diffusion interaction mechanisms such as brain and heart tissues. Such similarity might suggest a presence of a diffusive mechanism in the inter-nephron interaction or a long-distance fast electrical signalling, e.g. via conducted vasoreactivity.

In our experiments, we imaged 1.5x1.5 mm² field of view in 5 animals with 94.8 ± 15.66 individual segmented vessels on average, each of which is 10 – 30µm in diameter. While it provides an estimate of synchronization in the nephrons activity, direct translation from individual segmented vessels to nephrons is challenging and requires further exploration. Factors that are critical to consider are (i) penetration depth of LSCI when applied to renal imaging, (ii) type of the vessels in the field of view, and (iii) topology of the nephro-vascular network. While in theory, LSCI can collect the blood flow signal from as deep as 300-400 µm, in practice, visually resolvable signal typically comes from top 50-150 µm of the vascular structure.

Considering high vascular density close to the renal surface, it would mean that LSCI is likely limited to imaging vessels originating from 10000 nephrons in outer 30% of rat renal cortex, which would result in 40 nephrons in the 1.5x1.5µm field of view. The larger number of segmented vessels can be explained by their mixed type - afferent and efferent arterioles as well as venules are likely to be segmented. Distinguishing vessels types in LSCI images will require further exploration and registration with high-resolution structural imaging. It, however, does not mean that the observed clusters were limited to, at the most, 40 nephrons. When considering renal vascular topology, it is to be expected that within the 1.5x1.5µm field of view, we observe arterioles that arise from different non-terminal arteries. Depending on the branching order, each of such arteries can branch into ten-several hundreds of nephrons, but only a small number of these nephrons will have arterioles reaching close enough to the surface to be segmented from LSCI images. Thus, when a synchronous cluster is observed with LSCI, it is likely to extend several branching orders in depth and reach the size of hundreds and even thousands of nephrons.

While the exact role of inter-nephron communication, co-operative dynamics and synchronization in kidney-related pathology development is still unclear and requires further exploration, it is evidently altered by the blood pressure and vascular tone. Strong local coupling and in-phase synchronization, while being not evident at the renal artery level, are likely to increase pressure variation at the level of afferent arterioles, thus increasing chances of local damage and aggravating pathological condition.

**Funding.** D.D.P. was supported by grant NNF17OC0025224 awarded by Novo Nordisk Foundation, Denmark and by grant R345-2020-1782 awarded by Lundbeck Foundation, Denmark.

**Conflict of Interest:** none declared.

**Data Availability Statement.** The data underlying this article will be shared on reasonable request to the corresponding author.

**References**

1. Young DK, Marsh DJ. Pulse wave propagation in rat renal tubules: implications for GFR autoregulation. American Journal of Physiology-Renal Physiology. 1981;240(5):F446–F458.
2. Chon K, Chen YM, Holstein-Rathlou NH, Marsh DJ, Marmarelis VZ. On the efficacy of linear system analysis of renal autoregulation in rats. IEEE transactions on biomedical engineering. 1993;40(1):8–20.
3. Chon KH, Raghavan R, Chen YM, Marsh DJ, Yip KP. Interactions of TGF-dependent and myogenic oscillations in tubular pressure. American Journal of Physiology-Renal Physiology. 2005;288(2):F298–F307.
4. Cupples WA, Loutzenhiser RD. Dynamic autoregulation in the in vitro perfused hydronephrotic rat kidney. American Journal of Physiology-Renal Physiology. 1998;275(1):F126–F130.
5. Marmarelis VZ, Chon KH, Holstein-Rathlou NH, Marsh DJ. Nonlinear analysis of renal autoregulation in rats using principal dynamic modes. Annals of biomedical engineering. 1999;27(1):23–31.
6. Sakai T, Hallman E, Marsh DJ. Frequency domain analysis of renal autoregulation in the rat. American Journal of Physiology-Renal Physiology. 1986;250(2):F364–F373.
7. Shi Y, Wang X, Chon KH, Cupples WA. Tubuloglomerular feedback-dependent modulation of renal myogenic autoregulation by nitric oxide. American Journal of Physiology-Regulatory, Integrative and Comparative Physiology. 2006;290(4):R982–R991.
8. Holstein-Rathlou N, Wagner AJ, Marsh DJ. Tubuloglomerular feedback dynamics and renal blood flow autoregulation in rats. American Journal of Physiology-Renal Physiology. 1991;260(1):F53–F68.
9. Holstein-Rathlou N, Marsh D. Oscillations of tubular pressure, flow, and distal chloride concentration in rats. American Journal of Physiology-Renal Physiology. 1989;256(6):F1007–F1014.

10. Yip K, Holstein-Rathlou N, Marsh DJ. Mechanisms of temporal variation in single-nephron blood flow in rats. American Journal of Physiology-Renal Physiology. 1993;264(3):F427–F434.

11. Sosnovtseva O, Pavlov A, Mosekilde E, Holstein-Rathlou NH. Bimodal oscillations in nephron autoregulation. Physical Review E. 2002;66(6):061909.

12. Just A. Mechanisms of renal blood flow autoregulation: dynamics and contributions. American Journal of Physiology-Regulatory, Integrative and Comparative Physiology. 2007.

13. Holstein-Rathlou NH. Synchronization of proximal intratubular pressure oscillations: evidence for interaction between nephrons. Pflügers Archiv. 1987;408(5):438–443.

14. Kallskog O, Marsh D. TGF-initiated vascular interactions between adjacent nephrons in the rat kidney. American Journal of Physiology-Renal Physiology. 1990;259(1):F60–F64.

15. Yip KP, Holstein-Rathlou N, Marsh DJ. Dynamics of TGF-initiated nephron-nephron interactions in normotensive rats and SHR. American Journal of Physiology-Renal Physiology. 1992;262(6):F980–F988.

16. Sosnovtseva OV, Pavlov AN, Mosekilde E, Yip KP, Holstein-Rathlou NH, Marsh DJ. Synchronization among mechanisms of renal autoregulation is reduced in hypertensive rats. American Journal of Physiology-Renal Physiology. 2007;293(5):F1545–F1555.

17. Holstein-Rathlou NH, Sosnovtseva OV, Pavlov AN, Cupples WA, Sorensen CM, Marsh DJ. Nephron blood flow dynamics measured by laser speckle contrast imaging. American Journal of Physiology-Renal Physiology. 2011;300(2):F319–F329.

18. Boas DA, Dunn AK. Laser speckle contrast imaging in biomedical optics. Journal of biomedical optics. 2010;15(1):011109.

19. Briers D, Duncan DD, Hirst ER, Kirkpatrick SJ, Larsson M, Steenbergen W, et al. Laser speckle contrast imaging: theoretical and practical limitations. Journal of biomedical optics. 2013;18(6):066018.

20. Scully CG, Mitrou N, Braam B, Cupples WA, Chon KH. Detecting physiological systems with laser speckle perfusion imaging of the renal cortex. American Journal of Physiology-Regulatory, Integrative and Comparative Physiology. 2013;304(11):R929–R939.

21. Brazhe AR, Marsh DJ, Holstein-Rathlou NH, Sosnovtseva O. Synchronized renal blood flow dynamics mapped with wavelet analysis of laser speckle flowmetry data. PLoS one. 2014;9(9):e105879.

22. Postnov DD, Holstein-Rathlou NH, Sosnovtseva O. Laser speckle imaging of intra organ drug distribution. Biomedical optics express. 2015;6(12):5055–5062.

23. Postnov DD, Salomonsson M, Sorensen CM, Sosnovtseva O. A simple method to ensure homogeneous drug distribution during intrarenal infusion. American Journal of Physiology-Renal Physiology. 2017;312(3):F543–F548.

24. Postnov DD, Sosnovtseva O, Tuchin VV. Improved detectability of microcirculatory dynamics by laser speckle flowmetry. Journal of biophotonics. 2015;8(10):790–794.

25. Postnov DD, Tuchin VV, Sosnovtseva O. Estimation of vessel diameter and blood flow dynamics from laser speckle images. Biomedical optics express. 2016;7(7):2759–2768.

26. Postnov D, Postnov D, Marsh D, Holstein-Rathlou NH, Sosnovtseva O. Dynamics of nephron-vascular network. Bulletin of mathematical biology. 2012;74(12):2820–2841.

27. Duhlem MA, Graf R, Strong AJ, Dreier JP, Dahlem YA, Sieber M, et al. Two-dimensional wave patterns of spreading depolarization: retracting, re-entrant, and stationary waves. Physica D: Nonlinear Phenomena. 2010;239(11):889–903.

28. Shibata M, Bures J. Reverberation of cortical spreading depression along closed-loop pathways in rat cerebral cortex. Journal of neurophysiology. 1972;35(3):381–388.

29. Alonso S, Bär M, Echebarria B. Nonlinear physics of electrical wave propagation in the heart: a review. Reports on Progress in Physics. 2016;79(9):096601.

30. Davis MA, Kazmi SS, Dunn AK. Imaging depth and multiple scattering in laser speckle contrast imaging. Journal of biomedical optics. 2014;19(8):086001.

31. Zheng S, Xiao S, Kretsga L, Cruz-Martín A, Mertz J. Depth resolution in multifocus laser speckle contrast imaging. Optics Letters. 2021;46(19):5059–5062.
32. Letts RF, Zhai XY, Bhikha C, Grann BL, Blom NB, Thomsen JS, et al. Nephron morphometry in mice and rats using tomographic microscopy. American Journal of Physiology-Renal Physiology. 2017;312(1):F210–F229.

33. Marsh DJ, Postnov DD, Rowland DJ, Wexler AS, Sosnovtseva OV, Holstein-Rathlou NH. Architecture of the rat nephron-arterial network: analysis with micro-computed tomography. American Journal of Physiology-Renal Physiology. 2017;313(2):F351–F360.

34. Postnov DD, Marsh DJ, Postnov DE, Braunstein TH, Holstein-Rathlou NH, Martens EA, et al. Modeling of kidney hemodynamics: probability-based topology of an arterial network. PLoS computational biology. 2016;12(7):e1004922.