Multiphoton Imaging of Renal Regulatory Mechanisms*

Most physiological functions of the kidneys, including the clearance of metabolic waste products, maintenance of body fluid, electrolyte homeostasis, and blood pressure, are achieved by complex interactions between multiple renal cell types and previously inaccessible structures in many organ parts that have been difficult to study. Multiphoton fluorescence microscopy offers a state-of-the-art imaging technique for deep optical sectioning of living tissues and organs with minimal deleterious effects. Dynamic regulatory processes and multiple functions in the intact kidney can be quantitatively visualized in real time, noninvasively, and with submicron resolution. This article reviews innovative multiphoton imaging technologies and their applications that provided the most complex, immediate, and dynamic portrayal of renal function—clearly depicting as well as analyzing the components and mechanisms involved in renal (patho)physiology.

The old saying “seeing is believing” has driven many optical imaging applications in various fields of biomedical research, including renal physiology. Multiphoton microscopy (a term that denotes two- and three-photon excitation fluorescence microscopy) is an advanced confocal laser scanning fluorescence imaging technique that has proven especially useful for deep optical sectioning of living tissues. Multiphoton imaging can provide ultra-sensitive detection of fluorescent signals with submicron resolution, meaning that even intracellular organelles can be visualized in intact organs. The technology is based on the concept that the simultaneous absorption of two photons of equal energy can cause excitation of a fluorophore equivalent to the absorption of a single photon of double the energy (7, 74). Accordingly, multiphoton excitation uses infrared light in the 680- to 1,080-nm range in current commercial systems. In contrast, conventional confocal (one-photon) fluorescence imaging uses UV or visible lasers (193–694 nm). With multiphoton imaging, these longer (double or triple) wavelength photons (one-half or one-third of the energy) allow for deeper penetration into tissues with much less scattering and deleterious effects. Since multiphoton excitation occurs mainly at the focal plane, 100% of emitted (already confocal) fluorescence can be detected. In most multiphoton applications, there is no need for descanning and filtering the emitted fluorescence through pinholes as with conventional, one-photon confocal imaging. Detailed descriptions of the technology and its advantages and disadvantages can be found elsewhere (7, 8, 15, 32, 45, 74).

Clear tissues like the brain can be easily visualized with multiphoton techniques in depths in the millimeter range (8, 15, 22, 74). In contrast, the use of this technology in the renal cortex has been limited to the initial 100–200 µm under the kidney surface mainly due to the highly light absorbing and scattering renal tissue structures (32, 45). Continuous improvements in optics, fluorescence detection, and laser technology, however, are helping us to go deeper in the intact kidney even with noninvasive or minimally invasive whole animal techniques. One example is the dispersion compensation (i.e., keeping laser pulses short, in the 100-femtosecond range) that is now available in many commercial systems (for example, the Mai Tai DeepSee from Spectra-Physics/Newport and the Chameleon Vision from Coherent). Another improvement that facilitates deeper imaging is the significantly increased output power in new systems (>4,000 mW in Chameleon Ultra-II or Vision from Coherent in contrast to 350 mW in the first Tsunami lasers from Spectra-Physics about 10 years ago), although one must be cognizant of potential damage to the sample due to heating effects of these high power lasers. As a result of these technical improvements and an important milestone discussed below, multiphoton visualization of mouse glomeruli is now possible in normal C57BL6 mice and does not require special animal strains with superficial glomeruli like the Munich-Wistar rat (9, 20, 28).

This review focuses on the applications of multiphoton imaging in the intact kidney. We discuss recent advances in renal physiology and the imaging of complex renal regulatory mechanisms in or between the nephron and the vasculature, which were made possible by this imaging approach. We recommend this work for not only the basic or clinical research scientist but also as a visual aid for teaching graduate and medical students.

*This article contains supplementary files available online at the Physiology website.
Gomó Structure and Function

High-resolution optical sectioning through an entire living, intact glomerulus (diameter of ~100 μm) has become possible with multiphoton microscopy, with the first in vitro (42) and in vivo applications (9) having been published in 2002. The initial studies provided impressive, high-quality images of the complex cell structure of the glomerulus and the juxtaglomerular apparatus (JGA) (42), as well as direct visualization of the glomerular filtration of iv-injected fluorescent markers (9, 10, 32). Real-time changes in cell volume and cytosolic parameters (e.g., [Ca2+]i) of otherwise inaccessible cell types, including the extra- and intraglomerular mesangium, endothelium, and podocytes, were observed with this technology (42, 44–45). A few years later, the focus has shifted to quantitative imaging of basic renal functions, including the measurement of glomerular filtration rate (GFR). Using a number of water soluble, freely filtered, low molecular weight fluorescent markers, including a 10-kDa FITC-dextran (9, 10), Cascade blue (32), and Lucifer yellow (20), the process of glomerular filtration was visualized with high temporal resolution. This allowed the calculation of single nephron GFR (SNGFR) (20) using principles and techniques that were established earlier (4). A supplementary video file shows the renal clearance of the extracellular fluid marker Lucifer yellow (injected in a single iv bolus into the femoral vein) by glomerular filtration and its downstream movement in the proximal tubule (available online at the Physiology website) (from Ref. 20). Using two different molecular weight markers simultaneously injected iv and an optical ratiometric imaging approach, a technique has been developed by Yu and Molitoris for the rapid determination of GFR (71, 72) that has great potential for future use in clinical diagnostics.

Multiphoton imaging allowed both functional and morphological discoveries concerning the glomerular filtration barrier (GFB). FIGURE 1 demonstrates various applications both in vivo (FIGURE 1A, B, AND D) and in vitro (FIGURE 1C) in various species. Importantly, with the development of multiphoton technology, the imaging of glomeruli is now possible not only in Munich-Wistar rats that are renowned for their surface glomeruli (9, 10, 20, 28) but also in wild-type C57BL6 mice that have glomeruli deeper under the surface. The imaging of multi-
of glomeruli in the intact mouse kidney. -150-200 µm under the surface is demonstrated in Figure 1A using previously described techniques (20) and a Leica TCS SPS AOBS MP confocal microscope system (Leica Microsystems, Heidelberg, Germany) powered by a Chameleon Ultra-II multiphoton laser (Coherent, Santa Clara, CA). This significant development opens up new possibilities for imaging kidney function in genetically modified small animals.

A recent, potentially paradigm-shifting finding in work by Russo et al. using multiphoton imaging was the observation of mass filtration of fluorescein-conjugated albumin in normal kidneys and its rapid endocytosis in the proximal tubule (55). This has led to the authors’ assertion that the GFB normally leaks albumin at nephrotic levels and that this filtered albumin load is avidly bound and retrieved by cells of the proximal tubule (55). The issue is, however, highly debated (5), and multiphoton-based determination of the glomerular sieving coefficient for albumin that resulted in a value (0.02–0.04) that is 50 times greater than previously measured warrants future, careful review of the methodology. In contrast, another study found that glomerular permeability of rhodamine-conjugated 70-kDa dextran (only slightly larger than albumin) was minimal even in STZ-diabetic, hyperfiltering glomeruli as opposed to the highly dextran-permeable sclerotic glomeruli, the likely source of diabetic proteinuria (20).

Ultrastructural reconstruction of the podocyte and the glomerulus by Neal and colleagues (34) resulted in the exciting anatomical discovery of the subpodocyte space (SPS). The SPS was identified as the labyrinthine space between the underside of podocyte cell body/primary processes and the foot processes and was proposed to be a new layer of the GFB (34). Morphometric measurements suggested highly significant restriction to flow by the SPS (34, 35), a finding that was later confirmed functionally in the intact glomerulus and kidney by multiphoton imaging studies (56). Using methods demonstrated in Figure 1, C and D, it was found that rhodamine-conjugated-10-kDa dextran, but not the low-weight molecule Lucifer yellow, accumulated in SPS-covered regions of the GFB (56). The SPS-mediated resistance to flow adds more complexity to the glomerular filtration process and further suggests the important physiological role of podocytes in the regulation of GFR.

FIGURE 1, C AND D, provides examples of how the podocyte cell body, main processes, and intracellular vesicles can be visualized.

In vivo multiphoton imaging of renal tubular functions

A–D: Alexa Fluor 546-labeled siRNA uptake in rat proximal (PT) and distal (DT) tubule segments. Compared with background fluorescence at time 0 (A), significant levels of AF546-siRNA were detected (B–D) after injection into the carotid artery. Within 2 min of injection, robust siRNA uptake was visible in the PT, mainly at the apical brush border membrane (B), indicating uptake from the tubular lumen. Although lower amounts than that in the PT, siRNA uptake was detected in cells of the DT 15–20 min after injection (C). Distal nephron siRNA uptake included the macula densa (MD) region of the thick ascending limb (TAL) adjacent to the glomerulus (G) in D. E and F: pseudocolor images of intracellular calcium ([Ca²⁺]i) in PT, DT, and cortical collecting duct (CCD) in the intact mouse kidney using Rhod-2 AM injected under the renal capsule. Scale is 20 µm. G: Rhod-2 fluorescence recording from cells of the PT shows regular oscillations in [Ca²⁺]i with a slow (~20–40 mHz) and a fast (100–200 mHz) component, which appears to be in phase with oscillations in tubular diameter.
in the intact glomerulus and kidney. The drawings in FIGURE 1E (from Ref. 56) illustrate the two regions of GFB either covered by the SPS or not (naked region).

Real-time multiphoton imaging of the iv injected extracellular fluid marker Lucifer yellow was also instrumental in the direct visualization of the bulk fluid flow in the JGA interstitium (54) originally proposed and described by Rosivall (52, 53). Ultrafiltration of plasma through the fenestrated endothelium in the terminal renin-positive segment of the afferent arteriole (52, 53) was further confirmed by using this imaging approach (54). It has been suggested that this dynamic fluid flow in the JGA may help filter the released renin into the renal interstitium (endocrine function) or perhaps modulate the concentration of chemical mediators in the JGA interstitium that are important in the regulation of renal and glomerular hemodynamics (54).

Tubular Transport

Multiphoton microscopy has been used to study various tubular functions including proximal tubule endocytosis of gentamicin (9), folic acid (57), siRNA (33), and albumin (32, 55), as well as tubular secretion (63), protein expression (64), cyst development (9, 63), and apoptosis (9). FIGURE 2, A–D, shows time-lapse multiphoton imaging of iv-injected siRNA uptake in rat proximal and distal nephron segments, illustrating the feasibility of using the siRNA technique for efficient gene silencing in all nephron segments. With the help of low or intermediate molecular weight markers that were injected iv (Lucifer yellow, 10- to 70-kDa dextrans; Lucifer yellow, the GFB (56)), more complex and further insights of podocytes and podocyte cell processes and further studies (56).

FIGURE 3. Imaging the juxtaglomerular apparatus (JGA)

Transmitted light-DIC (A) and fluorescence images (B) of an in vitro isolated rabbit JGA consisting of the glomerulus (G) and attached, microperfused afferent arteriole (AA) and thick ascending limb containing the macula densa (MD). B: cell membranes of MD cells and the vascular endothelium in AA and G were labeled with R18 (red), renin granules with quinacrine (green), and cell nuclei with Hoechst 33342 (blue). Figure taken from Ref. 45. C: direct visualization of electrical signals in interlobular artery (ILA) and attached afferent arterioles (AA) in vitro using ANNINE-6, a new, highly sensitive, ultrafast voltage-sensing dye. ANNINE-6 fluorescence is highly cell membrane-specific and intense in vascular smooth muscle cells. Cell nuclei were labeled with Hoechst 33342 (blue). Scales in A–C are 20 μm. D: components of the TGF calcium wave include ATP release from MD cells in response to elevations in tubular NaCl and/or flow rate and paracrine, purinergic calcium signaling. Extracellular ATP-induced elevations in mesangial, vascular smooth muscle, and renin producing JG cell (Ca2+) are then propagated to distant cells of the AA, efferent arteriole (EA), and G away from the MD region, causing cell contractions.
Multiphoton microscopy allowed real-time visualization of cytosolic variables (e.g., intracellular ion concentrations) in intact tubular epithelia in vivo such as pH and [Ca\(^{2+}\)](61). One caveat when loading cells in vivo with the widely used acetoxy-methylester (AM) forms of fluorophores is the significant esterase activity of plasma. AM dyes (when used in vitro) can quickly enter the cell across the plasma membrane and get trapped after cleavage of the AM group by intracellular esterases. When injected in vivo, however, plasma enzymes can rapidly cleave the AM groups, resulting in the formation of the free, cell-impermeable form before reaching the target cells, thereby making cell loading very inefficient (16). One possible solution is the targeted bulk loading of fluorescent indicators by direct injection under the renal capsule, similar to the multicell bolus loading (MCBL) technique developed for multiphoton imaging of the brain in vivo (12). This method was successful for visualizing real-time changes in proximal tubule cell pH (important to study bicarbonate reabsorption) and [Ca\(^{2+}\)](61). Here, we show the loading of proximal and distal tubule cells with the calcium fluorophore Rhod-2 AM injected under the renal capsule (FIGURE 2E and F) using previously described methods (20, 61). The use of this red calcium fluorophore was preferred over green dyes (like Fluo-4) due to the low overlap of its emission with the significant renal tissue autofluorescence. In all nephron segments, regular oscillations in [Ca\(^{2+}\)] were observed with one slow (~20–40 mHz) and one fast (100–200 mHz) component (FIGURE 2G). [Ca\(^{2+}\)], oscillations were most likely functionally associated with variations in tubular fluid flow rate due to tubuloglomerular feedback and the myogenic mechanism of the afferent arteriole that have the same characteristics (16, 28, 30), since the proximal tubule diameter showed similar, phase-matched oscillations (FIGURE 2G). A supplementary video file shows proximal tubule [Ca\(^{2+}\)], oscillations in the intact kidney from the same in vivo study as shown in the still image of FIGURE 2F (see supplemental video at the Physiology website).

### Tubulo-vascular Interactions

One of the earliest applications of multiphoton imaging in renal tissues was the visualization of the juxtaglomerular apparatus (JGA) (42). The multiphoton imaging approach proved to be very useful for the study of this hardly accessible, highly heterogeneous cell complex that is one of the most important regulatory sites of renal salt and water conservation as well as blood pressure maintenance (19–21, 25–27, 40–42, 44, 45, 54, 56, 61). The main JGA functions include the high distal tubular [Na\(^{+}\)]-induced afferent arteriolar vasoconstriction that controls GFR, renal blood flow [tubuloglomerular feedback (TGF)], and the low tubular [Na\(^{+}\)]-induced renin release that activates the renin-angiotensin system (RAS) (59). Numerous details of these important physiological mechanisms were observed with multiphoton microscopy using the in vitro dissected and microperfused JGA model (FIGURE 3, A–D). Supplementary video files are available in the online version of this article that demonstrate in this living JGA preparation the TGF and renin release mechanisms (see supplemental files online at the Physiology website).

The Physiology visualization content and/or the point of view (POV) of cell [Ca\(^{2+}\)], etc. The TGF calcic signaling axis is involved in feedback control of both the afferent arteriole and the glomerulus (48, 62). Using the ultrastable and intravascular calcium sensing dye (11, 12) much faster release and reuptake of calcium is suggested as a mechanism leading to TGF oscillations (14). In the distal convoluted tubule (CD), a wave of calcium is transmitted from the macula densa to the juxtaglomerular apparatus (JGA) by the dilatory function of GABAergic interneurons (44, 45, 54, 56, 61). The main JGA functions include the high distal tubular [Na\(^{+}\)]-induced afferent arteriolar vasoconstriction that controls GFR, renal blood flow, tubuloglomerular feedback (TGF), and the low tubular [Na\(^{+}\)]-induced renin release that activates the renin-angiotensin system (RAS) (59). Numerous details of these important physiological mechanisms were observed with multiphoton microscopy using the in vitro dissected and microperfused JGA model (FIGURE 3, A–D). Supplementary video files are available in the online version of this article that demonstrate in this living JGA preparation the TGF and renin release mechanisms (see supplemental files online at the Physiology website).
brane and get into the intracellular organelle enzymes that are important for the intracellular oscillation (20, 51). The cell loading of the renin enzyme is the target for the multiphoton imaging (20). A supplementary file (available at the Physiology website) (from Refs. 40, 44). Real-time visualization of the JGA revealed that increased salt content and/or tubular fluid flow in the distal tubule at the point of the macula densa (the sensory element of the JGA) triggers mesangial and vascular smooth muscle cell (Ca^{2+}) elevations and cell contractions (44, 45). The TGF calcium wave illustrated in FIGURE 3D and qualitatively approximated from the MD region toward both the proximal afferent and efferent arterioles and all cells of the glomerulus at a rate of ~12 μm/s (44). The TGF calcium wave was mediated by extracellular ATP (44), supporting earlier findings (1, 17, 38) that ATP itself is directly involved in TGF and not simply through its breakdown to adenosine (2, 3, 44, 62). Using ANINE-6 [a new, highly sensitive and highly cell membrane-specific voltage-sensing dye (FIGURE 3C), a recent study observed a much faster propagating electronic vascular TGF signal as well (31), a mechanism that has been implicated in nephron synchronization (31). These vascular conducting cells demonstrate the importance of intercellular communication in the JGA, which involves not only extracellular chemical mediators (24, 49, 50) but gap junctions as well (44, 49, 69, 70). An interesting new development concerning connexins, the building blocks of gap junctions, is the possibility of connexin hemichannels mediating the release of ATP (44, 67).

Multiphoton imaging has also helped to visualize morphological changes of the JGA in real-time and in four dimensions. For example, cell volume of the macula densa was found to be a good indicator of tubular salt-dependent macula densa cell swelling or shrinkage was also dependent on simultaneous changes in tubular fluid osmolality (25). Also, a recent elegant study by Komlosi et al. found regular oscillations in [Ca^{2+}] in the cortical thick ascending limb around the macula densa (26), suggesting that, in addition to the classic tubulo-vascular contact mechanism existing between the connecting tubule segment and the proximal part of the afferent arteriole (50). The complexity of tubulo-vascular interactions can be visualized on the intact, whole organ level using multiphoton imaging (20). A supplementary file in the online version of this article demonstrates the nonlinear nature of glomerular and tubular functions (available at the Physiology website) (from Ref. 20). Due to the myogenic and TGF mechanisms, regular periods of glomerular contraction-relaxation were observed, resulting in oscillations of GFR and tubular fluid flow rate. The oscillations in proximal and distal tubular fluid flow showed similar characteristics (slow and fast components) as described earlier (FIGURE 2C) (16, 30). Alterations in urine concentrating and diluting mechanisms and the effects of furosemide causing diuresis, enlargement of collecting ducts that resulted in the compression of peritubular capillaries were also directly visualized (20, 46). The latter is consistent with the paradoxical, preglomerular vasodilator but renal blood flow-reducing effect of furosemide (38).
The Intra-renal Angiotensin System

Heterogeneity of cortical renin synthesis in the kidney provides another good example of the advantages of this direct imaging approach. In addition to the JGA, which is the classic anatomical site of renin synthesis, significant amounts of renin were recently found in proximal tubules and in principal cells of the collecting tubule and the collecting duct (CD) (21, 46, 51). CD renin synthesis was particularly robust in pathophysiological states associated with high RAS activity, such as diabetes and renovascular hypertension (21, 46, 47). Also, the CD was identified as the major source of prorenin synthesized in diabetes (21) (FIGURE 4A). Interestingly, angiotensin II (ANGII) differentially regulates the synthesis of JGA (inhibition) and CD (stimulation) renin (21, 46, 47), making the conventional molecular studies (using whole kidney or whole cortical tissue samples) somewhat problematic. However, in vivo imaging of the two main intra-renal locations of (pro)renin synthesis is possible using multiphoton microscopy as shown in FIGURE 4, A AND B. Quinacrine, a nonspecific acidic fluorophore that labels renin granules very intensely has been used successfully to visualize JGA renin content both in vitro (FIGURE 38) (46, 45) and in vivo (FIGURE 4A) (20, 61, 65). It should be noted, however, that quinacrine is also a weak stain of all acidic organelles, including lysosomes and cell nuclei. Therefore, it must be used carefully and only with imaging applications that allow direct acquisition of renin granules (based on location), and separation from background fluorescence (in lysosomes, etc.) as shown in FIGURE 4, C AND D. According to the recent findings of immense CD (pro)renin synthesis in diabetes (21) (FIGURE 4B), CD prorenin may be released to cause systemic or local pathological actions at the recently identified and characterized (pro)renin receptor (6, 37, 38), either as a signaling molecule or as a source of locally produced ANGII. This would be consistent with the existence of a local distal tubular RAS (51) and with its possible regulatory effects on salt reabsorption (23, 43).

Using the multiphoton imaging approach, it is now possible to directly visualize not only renin granular content and release but also the enzyme activity of renin in intact renal tissues (FIGURE 5). Several fluorescence resonance energy transfer (FRET)-based fluorogenic renin substrates have been recently developed by various companies (Invitrogen, AnaSpec, Cayman Chemical, etc.) and contain a pair of donor-acceptor fluorophores connected by a sequence (decapetase) of rodent angiotensinogens, including the renin cleavage site at the Leu-Leu bond (Leu-Val in humans). Using the newest generation of highly sensitive protease FRET assays (FIGURE 5), in the absence of renin activity, the donor (5-FAM) fluorescence is quenched by the acceptor molecule (QXL 520) due to their close proximity and fluorescence resonance energy transfer (FRET) between them. However, upon cleavage into two separate fragments by rat or mouse renin, the fluorescence of 5-FAM is recovered and can be monitored. This fluorescence-based technique permits real-time measurement of renin activity, does not utilize radioactivity, and is conveniently performed within minutes, as opposed to conventional renin assays using radioimmuno methods.

For future applications, this imaging approach can be very useful to study the intra-renal RAS in pathophysiological conditions such as diabetes. Although it is well established that RAS activity is high in diabetes (13), it has been difficult to measure increased ANGII formation or activity in the diabetic kidney using conventional molecular techniques (73). This new fluorogenic renin substrate in conjunction with multiphoton imaging can help detect renin enzymatic activity and locally produced, quenched ANGII peptides, for example in the JGA as shown in FIGURE 5B. Also, the importance of RAS activation in several metabolic diseases, including diabetes, metabolic syndrome, and hyperuricemia, has been recognized (11, 13). The recent localization of the metabolic receptor GPR91 in the (juxta)glomerular vasculature and the macula densa (40, 45) and its possible role in JGA renin content was visualized in control and diabetic kidneys with multiphoton microscopy (60). This imaging approach can be used to monitor and study the complex mechanisms of RAS activation in many disease models.

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References
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