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NOVEL HAEMODYNAMIC STRUCTURES IN THE HUMAN GLOMERULUS

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KPA - Reconstruction imaging, software assistance, multi-photon microscopy,
manuscript
JSB - Multiphoton microscopy and software, manuscript
KBB - Confocal microscopy and software, manuscript
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Abstract

To investigate human glomerular structure under conditions of physiological perfusion we have analysed fresh and perfusion fixed normal human glomeruli at physiological hydrostatic and oncotic pressures using serial resin section reconstruction, confocal, multiphoton and electron microscope imaging.

Afferent and efferent arterioles (21.5±1.2µm and 15.9±1.2µm diameter), recognised from vascular origins, lead into previously undescribed wider regions (43.2±2.8 µm and 38.4±4.9 µm diameter) we have termed vascular chambers (VCs) embedded in the mesangium of the vascular pole. Afferent VC(AVC) volume was 1.6 fold greater than Efferent VC(EVC) volume. From the AVC long non-branching high capacity conduit vessels (n=7) (Con; 15.9±0.7µm diameter) led to the glomerular edge where branching was more frequent. Conduit vessels have fewer podocytes than filtration capillaries. VCs were confirmed in fixed and unfixed specimens with a layer of banded collagen identified in AVC walls by multiphoton and electron microscopy. Thirteen highly branched efferent first order vessels (E1;9.9±0.4µm diam.) converge on the EVC draining into the efferent arteriole (15.9±1.2µm diam.). Banded collagen was scarce around EVC.

This previously undescribed branching topology does not conform to the branching of minimum energy expenditure (Murray’s law), suggesting even distribution of pressure/flow to the filtration capillaries is more important than maintaining the minimum work required for blood flow. We propose that AVCs act as plenum manifolds possibly aided by vortical flow in distributing and balancing blood flow/pressure to conduit vessels supplying glomerular lobules. These major adaptations to glomerular capillary structure could regulate haemodynamic pressure and flow in human glomerular capillaries.

Introduction

The control of glomerular blood flow is crucial for maintaining efficient ultrafiltration across the glomerular filtration barrier (GFB). Glomerular disease is characterised by molecular and physiological perturbations and altered glomerular haemodynamics (intraglomerular pressure and hyper-perfusion), however, most of the models of glomerular haemodynamics in humans are based on experimental animals with small glomeruli. A few studies have attempted to reconstruct the human glomerular vascular network; a wax model of a human neonate glomerulus was reconstructed by Johnston in 1899 (21) and in 1956 plastic glomerular vessels were reconstructed from wax moulded outlines (6). These and later casting techniques render impressions of the glomerular surface capillaries with deeper vessels remaining largely hidden.

More recent computational methods have revealed nodes and branching in rat and human glomerular vasculature (33, 34, 47, 48, 53). The human reconstructions were performed on 5µm sections and/or on immersion fixed sources or only on small glomerular regions and the few studies of the vascular pole of the human glomerulus have used biopsy or cadaver recovered material (33, 56). To date, only one reconstructive study has been published using perfusion-fixation of a human transplant kidney but at elevated hydrostatic pressure (140mm Hg) where the authors chose a stereological approach for vessel analysis rather than reconstruction (4).
Glomerular capillaries operate at relatively high pressure in life which in turn sets urinary driving pressure in the Bowman’s capsule and tubules producing tubular flow. For instance, the human glomerular capillary hydrostatic pressure of 60 to 65 mmHg at the afferent end (43) falls only 2-3mmHg to the efferent end. Countering this filtration pressure is an afferent plasma colloid osmotic pressure of 25mmHg rising to 32mmHg at the efferent end (1). As a result of filtration, urinary space hydrostatic pressure is 20-25mmHg (61) pressurizing the proximal convoluted tubule producing flow through to the collecting duct and the renal hilus. Thus, the function and structure of the whole nephron relies upon the glomerular perfusion of an oncotically appropriate fluid at the correct hydrostatic pressure to raise the right physiological pressures and flows in the tubules. In biopsy/necropsy kidney specimens the absence of pressure during immersion fixation results in the collapse of both the glomeruli and tubules. Fixing at the correct physiological pressures (oncotic and hydrostatic) is therefore essential in investigating the true ‘functionally inflated’ architecture of the glomerulus.

We have previously shown that 3D ultrastructural reconstruction of animal and human glomeruli fixed under hydrostatic and oncotic physiological conditions allow the detailed analysis of the GFB and the identification of novel structural features such as the subpodocyte space (SPS)(39) One unexpected feature of light microscopic sections from these resin embedded human glomeruli was the frequency of wide vessel regions at the vascular pole when compared with rodent vascular poles implying different vascular structure. No mention of any such difference could be found in any recent study of human glomerular structure.

The haemodynamic requirements of rat and human glomeruli could shed light on any differing evolved morphologies. For instance, if glomerular volume is assumed to estimate perfused glomerular volume, this parameter does not scale in size with the increase in afferent arteriolar conductivity between rodents and humans. The human afferent arteriole has a conductivity 13 fold greater than that of the mouse (14000µm$^4$ vs 1100µm$^4$) but supplies a 23 fold larger glomerular volume [see Footnote 1]. Similarly, it is 3 times as conductive as that of the rat (4600µm$^4$), while supplying a 5 fold larger glomerular volume. If human glomerular morphology was simply scaled up from a small rodent pattern, then afferent arterioles should be closer to 26µm in diameter instead of 21µm.

This study therefore aimed to investigate these novel wide vascular regions of human glomeruli. How big were these regions? What was the wall structure and dimensions and were there any other associated features? Did the region constitute a wider region at the base of the afferent arteriole or a region of a thin walled capillary? Could these structural differences be involved in compensating for a high glomerular volume relative to the vascular input in human glomeruli? To address such questions, human kidneys were perfuse fixed (at physiological hydrostatic and oncotic pressures) and processed in such a way to reduce any accompanying tissue volume changes. Glomerular vasculature was observed and reconstructions made from fresh or fixed human kidney cortex using conventional light microscopy, confocal microscopy, multiphoton microscopy and transmission electron microscopy.

**Methods**
Fixation techniques

Human kidney tissue was sourced (with full ethical approval and consent of next of kin) from transplant kidneys (n=9) unused for technical reasons (eg poor major vessel condition, damage at retrieval, tumour in the contralateral kidney). The transport solution perfused through the kidney was Soltran (Potassium Citrate 0.86% w/v, Sodium Citrate 0.82% w/v, Mannitol 3.38% w/v; Magnesium Sulphate 1.0% w/v; Baxter Healthcare, UK). Approximately 2-3 litres of the solution was perfused through the kidney (200ml/minute, 120-140mmHg, 4°C) and then stored on ice. All other chemicals were sourced from Sigma-Aldrich, UK.

Kidneys were transported in ice-cold flush media. Centimetre diameter fresh cortical tissue was sampled from one pole for confocal and multiphoton microscopy and stored in chilled (4°C) HEPES buffered Ringers solution. Smaller 1mm diameter tissue pieces were taken from the cut surface and fixed in 2.5% glutaraldehyde in HEPES buffer to serve as immersion fixed samples for TEM. At 4-10°C kidneys were debrided of excess fat preserving the hilar components (renal artery, vein and ureter) and the sampled polar area of the kidney was clamped off with a large locking forceps. The renal artery was cannulated and the renal vein was cleared of any debris to allow outflow of perfusion fluid.

To offset any hyperfiltration and hyperperfusion during fixation normal hydrostatic and oncotic pressures were re-established by perfusing with an oncotically balanced (25mmHg oncotic pressure) flush solution (50ml, 20°C). Colloid osmotic pressures were measured using a modified Hanson osmometer. The flush solution temperature was kept low to minimise autolytic/proteolytic activity. The hydrostatic pressure in the renal artery was set at 100mmHg (similar to human mean arterial pressure). After the flush bolus, 400ml of fixative was perfused through the kidney at the same pressures and temperature. Flush solution concentration was (mM); NaCl(132), KCl (4.6), MgSO₄ (1.3), CaCl₂ (2), HEPES (5), NaHCO₃ (25), D-glucose (5.5), 6.5% (w/v) Ficoll 400. Fixative was the same as the flush solution but with 1.25% (w/v) glutaraldehyde. The glycocalyx stain 0.5% lanthanum nitrate and 0.5% dysprosium chloride was incorporated into the solutions in 2 kidneys.

1mm diameter samples of perfusion fixed kidney were taken from a medial subcapsular position and together with subcapsular immersion fixed samples were post-fixed in osmium tetroxide, dehydrated with ethanol and processed into Araldite resin using standard procedures.

To promote consistency in structural comparisons, measurement and observations were limited on the glomeruli of the outer (subcapsular) cortex of kidneys in a medial location half way between the poles (unless otherwise stated).

Reconstruction of vascular poles from perfusion fixed kidneys

Seven areas of resin embedded kidneys (n=4) which contained a high density of glomeruli were identified in Toluidine Blue stained sections. These areas were serially sectioned on a Reichert Ultracut microtome at 1µm thickness (2,095 sections approximately 300 sections per area). From these serial section runs, 3 or 4 fully sectioned glomeruli from each kidney were selected that clearly showed a vascular pole. The afferent arterioles of each of the 14 glomeruli were identified by tracing to a larger artery and/or the efferent arteriole traced to a peritubular position.
Digital micrographs (1,834) of serial sections of glomeruli (n=14) were made using a x40 objective on a Nikon E400 microscope. Digital images were repositioned, aligned, calibrated and measured using Image J software (NIH opensource ImageJ 1.46r & 1.47o) and compiled into image stacks. Topological maps were made of the route and diameter of the blood vessels coursing through the afferent and efferent parts of the vascular pole.

**Resin section thickness calibration and glomerular diameter**

Measurement and reconstruction in the sectioning direction is reliant upon the precision of the ultramicrotome mechanism controlling section thickness. To test the accuracy of the ultramicrotome, glomeruli were assumed to be spherical and of similar diameter in all directions. Glomerular diameter was measured in the sectioning direction (z) as well as in the section plane (x,y). An ellipse was fitted over the largest glomerular profile of a section image (x,y) and maximum and minimum diameters measured from this, the results were pooled (194.4±5.1µm n=28). In the image stacks of a glomerulus the first and last sections to contain the edge of glomerular blood vessels were found and the number of intervening sections counted (202.4±5.0 n=14). Assuming 1µm section thickness there was no significant difference between the estimates of glomerular diameter from either method (t-test; P=0.325) and no correction was needed for section thickness or measurements of length in the sectioning direction (z).

The glomerular diameters (2r_x 2r_y 2r_z) measured during the calibration of section thickness were used to calculate glomerular volume (V_G = 1.33 π r_x r_y r_z).

**Glomerular and vascular orientation in resin section reconstruction**

Vascular pole recognition was most easily achieved in 1µm serial resin sections where the section plane was par-axial with the *vascular pole - urinary pole* axis of the glomerulus, as a result reconstructed glomeruli were sectioned close to a paraxial plane. The true diameters of any vessel profile was measured by searching the sequential images for the appropriate vessel section and measuring vessel width (x,y). Section depth diameter was taken from the limits of vessel walls in the sectioning direction (z). Vessel lengths (between branch points for example) through the image stack were measured on section if possible or by triangulating through the stack using sectioning depth and horizontal ‘on section’ distance.

The three diameters of VCs (x,y and z) used to calculate the means in table 1 and 2 were further used to calculate afferent and efferent vascular chamber volume (V_AVC = 1.33 π r_AVC r''_AVC r'''_AVC ; V_EVC = 1.33 π r_EVC r''_EVC r'''_EVC).

Bends between arterioles and VCs were assessed in resin section image stacks of 10 glomeruli by assessing the afferent and efferent arteriole axis vector and measuring the change in angle into the VC axis vector (Fig.3A). This included measurements on section and in the sectioning direction and triangulation in vessels moving at angles to the section plane.

**Afferent first order (conduit) vessel ballooning in resin sections**

Any ballooning or hyperinflation of first order afferent (conduit) vessels was estimated initially by comparing conduit diameters in areas of potentially high transmural pressure gradient (conduit vessels with large areas of GFB, 0-60% mesangial cover) with
conduit diameters in areas of potentially low transmural pressure gradient (conduit vessels with 80-100% mesangial cover). These data were further dissected in each conduit vessel by subdividing the initial 0-60% mesangial cover group into 4 groups and using the 80-100% mesangial cover group as a baseline to calculate the fold change in diameter.

Podocyte cell body coverage of conduit vessels

Podocyte cell body (PCB) coverage on the urinary side of conduit vessels was estimated by measuring length of GFB in a vessel image covered by a visible podocyte cytoplasmic region and the accompanying areas where no cell body was apparent. This was compared with similar measurements from filtration capillaries.

VC recognition in single resin sections

To test whether evidence of VCs could be seen in single sections of glomeruli (being the more common way of looking at human biopsy glomeruli) the occurrence of widened vasculature at the vascular poles was assessed in single sections of renal cortex. In an additional 13 resin-embedded human kidneys, immersion and perfusion fixed single cortical sections (1µm thick) were stained with Toluidine Blue. Glomerular sections showing vascular poles were assessed for the frequency of vascular widening around the poles. Width was assessed by placing an ellipse around widened vascular profiles and taking the minimum diameter to eliminate oblique vascular diameter measurements.

Confocal and multiphoton light microscopy on fresh kidney slices

Aqueous fresh and fixed kidney was observed using confocal and multiphoton microscope techniques. A Nikon confocal microscope (Nikon Eclipse Ti) was set to image fixation induced autofluorescence (FIA). Millimetre and sub millimetre thick fixed renal cortical slices were washed in HEPES Ringer solution and the autofluorescent signal (FIA) at 488nm wavelength was used to image and obtain z stacks from glomerular vascular poles of up to 100µm depth from the cut surface. Using a multiphoton microscope, two fresh and two fixed unstained slices of renal cortex, were imaged as previously described (2). Two imaging modes were applied, fibrous collagen was visualised using second harmonic generation (SHG) and elastin from its intrinsic two photon fluorescence (TPF) along with any background fluorescence. TPF and SHG images were obtained using a modified confocal microscope (FluoView IX71 and F300, Olympus). Signal was produced using the 800 nm output of a mode-locked Ti:sapphire laser (Mira 900-D, Coherent Inc) pumped by a 532 nm solid state laser (Verdi V10, Coherent Inc.). The pulsed laser had a pulse width of 100 fs and a repetition rate of 76 MHz. The light was focused on to the sample using a 60X 1.2 NA water immersion objective (UPlanS Apo; Olympus). Signal was collected in the epi-direction using the objective lens and separated from the laser fundamental using a long pass dichroic mirror (670dcrx; Chroma Technologies). The signal was then passed through two filters (for TPF: CG-BG-39 and F70-500-3-PFU; and for SHG: CG-BG-39 and F10-400-5-QBL; CVI Laser) before being focused on a photomultiplier tube (R3896, Hamamatsu). Each 1024×1024 pixel image took 29 seconds to acquire, meaning a stack
of 100 images, each separated in the z-direction by 1 μm, took approximately 50 minutes to complete.

**Electron microscopy**

From 1μm Resin sections of renal cortex showing identifiable VCs, further sections were cut at 70-100nm thickness and stained with 10% Phosphotungstic acid (10 minutes). Sections were viewed and digital images taken on a Tecnai T12 (FEI UK Ltd).

**Calculation of vascular resistance**

The resistance to flow along the terminal part of the arterioles will change as blood enters AVCs and conduits, and exits EVCs. Assessing such resistances may give a better understanding of how blood flow will be affected by VCs and conduits, a correlate of total conduit resistance per unit length (R'\text{Con}) was derived from the Poiseuille equation (see Appendix 1)

\[
R'\text{Con} = \frac{1}{r^{4}\text{Con} \cdot n_{\text{Con}}} \tag{Eq.1}
\]

Where \(r_{\text{Con}}\) is the mean conduit vessel radius and \(n_{\text{Con}}\) is the number of conduit vessels merging from an AVC. \(R'\text{Con}\) provides a value that scales proportionately with total vascular resistance per unit length. Similarly a correlate of first order efferent (E1) resistance per unit length (R'\text{E1}) was estimated from \(1/\text{r}_{\text{E1}}^{4} \cdot n_{\text{E1}}\) and arteriole resistance per unit length (R'\text{AA}, R'\text{EA}) was calculated from \(1/\text{r}_{\text{AA}}^{4}\) and \(1/\text{r}_{\text{EA}}^{4}\).

**Statistics**

Data were represented throughout as either mean ± standard error of the mean or as median (interquartile range). Excel was used for collating data and initial statistics, Prism software (Graphpad Software Inc.) was used for statistical analysis generating histograms, correlations, parametric and non-parametric tests.

**Results**

**Glomerular structure from resin serial section image stacks**

**Glomerular Arterioles**

Assigning afferent and efferent labels to arterioles was accomplished by tracing the origin of these vessels in the serial section image stacks. Branches of cortical radial or interlobular arteries (38, 58) were traced to the afferent arterioles (Fig.1) and efferent vessels showed a characteristic peritubular branching course on emerging from glomeruli.

Afferent and efferent arteriole wall thickness were significantly different (6.6±0.3μm, 3.0±0.1μm respectively, paired t-test \(p<0.0001\ n=7\)) as were afferent and efferent luminal diameter (23.2±1.8μm, 17.6±2.0μm respectively, paired t-test \(p=0.02\ n=7\)). Wall thickness being a better predictor of arteriole type than luminal diameter. No correlation was found between the afferent (R'\text{AA}) and efferent (R'\text{EA}) arteriole resistance measure (Tab.3 \(R^{2} = 0.033\ P=0.53\)).
The efferent picture was confused by multiple efferent arterioles in 4 out of 14 glomeruli. Major efferent arterioles have been shown in table 2, the extra 1 to 3 minor efferents were in series or parallel with EVC and were 4.6-8µm diameter with one 11.5µm in series with an efferent VC. No extra afferent arterioles were seen.

Reconstruction of VCs and 1st order vessels

All 14 glomeruli (4 kidneys) analysed from image stacks of 1µm resin serial sections showed afferent and efferent widening of the arterioles, resulting in vascular chambers (VCs) embedded in the mesangium of the vascular pole (Fig. 2, see supplemental video 2a and 2b for full image stacks). Some afferent VCs (AVCs) protruded into a hilar or juxta-glomerular position (sections 198 & 209, Fig. 2B).

Vascular width and connectivity is illustrated in a scale diagram in figure 3A, (measurements from tables 1&2). To summarize, the 21µm diameter afferent arteriole (AA) leads into an ellipsoidal afferent vascular chamber (AVC; 49x48x32µm) which branches into on average 7 first order afferent vessels of 16µm diameter we have termed conduit vessels (Con; Fig. 2A, B, Fig. 3A, Tab. 1). These vessels had secondary vessels (AA) emerging at spacings of 32.8µm (median), with 41% of branches intervals between A2 greater than 40µm with a quarter of these above 100µm (Fig. 4A). Conduit branches into A2 were more frequent distal to the AVC at the glomerular edge (Fig. 2B. & supplemental video 2a and 2b). Conduit vessels coursed through mesangium and then either through the centre of the glomerulus or peripherally over the glomerular surface before branching into capillary networks (Fig. 2A & B) [Supplementary videos S2c and S2d (Fig. 2B as a reconstructions)].

At the efferent end of the filtration capillary network first order efferent vessels (E1) were more numerous (13 v. 7) and narrower than conduits (10µm v. 16µm diameter; Fig. 4B; Tab. 1&2). Secondary efferents (E2) merged at 15µm intervals into 13 first order vessels (E1) (Figs. 2A, B & 3A). Only 4% of E2 branch intervals on E1 vessels were above 40µm - (Fig. 4A). E1s converged into an efferent vascular chamber (EVC; 46x43x26µm) in turn disgorging into a 16µm diameter efferent arteriole (EA; Figs. 2A, B & 3A, Tab. 2).

In 10 of the 14 glomeruli where the orientation of afferent and efferent arterioles on entry into the VCs could be easily assessed, the AA bent 60º off its straight track into the AVC (AA/AVC angle = 120±6º), similarly, the EA bent 71º off track into EVC (EA/EVC angle = 109±7º Fig. 3A).

VC and glomerular size

AVC volume (V_{AVC} = 41±5x10^3µm^3) was 1.6 fold greater than EVC volume (V_{EVC} = 28±7 x 10^3µm^3), with no correlation between them (R^2 = 0.164 P=0.152). V_{AVC} varied over a greater size range (15-70x10^3 µm^3) with V_{EVC} more conserved (12 out of 14 between 10-40x10^3µm^3). Both V_{AVC} and V_{EVC} correlated significantly with V_G (Fig. 4C, D, Tab. 4), V_G being 100 fold larger than V_{AVC} and 150 fold larger than V_{EVC} 150. This implies a relationship of both the input the output manifold with the magnitude of the perfused volume.

If the glomerular and VC volume (Fig. 4C, D) correlation is extrapolated back from larger glomeruli then a minimal VC volume can be reached where the volume describes a mere continuation of the attached arteriole (Fig. 3B). Accordingly, a
cylindrical minimum VC volume was calculated using average VC length (L) and arteriole radius (r), a minimum AVC volume of 1.57x10^4 µm^3 would occur at a V_G of 2.2x10^6 µm^3 (Fig.4c). Similarly, a minimum EVC volume of 0.75x10^4 µm^3 would occur at a V_G of 2.9x10^6 µm^3 (Fig.4D). Translating V_G into glomerular diameter, VCs would be minimal (a continuation of the arteriole) in human glomeruli below 160-180µm diameter (i.e. V_G = 2 - 3x10^6 µm^3).

**Conduit podocytes**

In resin section image stacks spanning a conduit vessel, we noted a significant lack of coverage of podocyte cell bodies (PCB) over the GFB surface (e.g. Fig.2A Con in sections 312 & 329; Supplemental 2a & 2b). Narrower, shorter first order efferent vessels (E1) were embedded in mesangium adjacent to the EVC and so had zero podocyte coverage (e.g. Fig.2B E1 in sections 249, 258 & 266). PCB area coverage was estimated in GFB conduit regions (n=10, i.e. Fig.2; GFB.Con Fig.3) and small filtration capillary regions (n=22) from 4 human glomeruli. Conduit vessel PCB area coverage was halved compared with small filtration capillaries (29±3% v. 55±3%; t-test - P<0.0001; Fig.5B).

**Conduit mesangial support**

Conduit vessels proceed from a central glomerular region with mesangium on all sides (Fig.3A Mes.Con) to regions with less mesangial attachment and areas of filtration barrier (Fig.3A GFB.Con). Appendix 2 shows that moving from mesangial supported regions of conduit to regions where this support is replaced by GFB more than doubles the hoop stress tendency to inflate or expand the vessel wall. To test if the GFB conduit regions showed any ballooning due to lack of mesangial support, conduit vessel diameters measured in 13 glomeruli (resin reconstruction method) were the same in high (80-100% mesangial cover) and low mesangial cover regions overall (17.7±0.8µm, 17.9±0.4µm respectively, paired t-test, n=61, p=0.28). However, after further division of the low mesangial cover data set, conduit vessels with the lowest mesangial cover (<15% mesangium, >85% GFB) showed significant inflation of 7% compared to high mesangial cover regions close to the AVC (paired t-test P=0.04, Wilcoxon P=0.04; Fig.5A).

**Vascular Resistance and volume relationships**

Since Poiseuille flow conditions do not apply to an ellipsoidal chamber manifold with many branches, the vascular resistance per unit length could not be calculated for VCs, therefore their capacity, V_AVG or V_EVC, was compared with glomerular vessel resistance parameters. Glomerular volume (V_G) was used as a correlate of perfusion volume and compared with the resistance parameters.

R'_{AA} did not correlate with any of the other R' parameters or V values, no correlation was found between R'_{AA} for afferent arterioles and V_AVG (R^2 = 0.014, P=0.68) or V_G which it supplies (R^2 = 0.065, P=0.38) (Tab.3). From the afferent VC there was a significant negative correlation between V_AVG and R'_{Con} (R^2 = 0.327, P=0.033; Fig.5C; Tab.3) showing that as the input manifold gets larger the supply conduits to the filtration capillary regions get proportionally more conductive (wider).

On the efferent side there was no similar correlation between efferent first order vessels R'_{E1} and V_EVC (R^2 = 0.088, P=0.303) though both of these correlated with V_G.
implying a link with perfusion volume. No correlation was found between $R'_{EA}$ and $V_{EVC}$ ($R^2 = 0.22, P=0.094$) but $R'_{EA}$ does correlate inversely with $V_G$ ($R^2 = 0.47, P=0.007$; Fig.5D, Tab.3) and directly with both first order afferents ($R'_{Con}$) and efferents ($R'_{E1}$).

Table 3 summarizes the capacity and resistance parameter correlations in the human glomerulus; strikingly, $R'_{AA}$ remains independent of all glomerular parameters but all other glomerular vascular entities appear fluid dynamically tied together.

**VC in single resin sections.**

Single sections of immersion and perfusion fixed kidney (n=13) revealed randomly orientated profiles of glomeruli with vascular poles (n=177). There was no significant difference in the occurrence of vascular widening at the vascular poles between immersion and perfusion fixed glomeruli or between juxta medullary (JM) and subcapsular (SC) glomeruli (Fig.6A). Analysis of all glomeruli together where no description was made in glomerular position (JMSC) in 8 immersion fixed tissues revealed vascular widening in 53±5% of vascular pole sections. Overall frequency was 60±4% for vascular widening in single sections of glomerular vascular poles.

The widened vascular regions found at SC vascular poles were 28.5±3µm and 30.7±2.1µm (minimal diameter) after immersion or perfusion fixation respectively and represented randomly oriented sections of presumably both vascular chambers. This lack of collapse shows that VCs appear to remain open even when the vascular pressure is reduced during fixation. The full morphology of JM vascular chambers remains to be investigated with serial sections.

**VC imaged by confocal and multiphoton microscopy**

Using a combination of fixation induced autofluorescence (FIA), two photon fluorescence (TPF) and second harmonic generation (SHG) modes, AVC could be seen with attached wide conduit vessels and AA in both fixed and fresh kidney slices (Fig.7). EVC was more difficult to observe with narrower blood vessels (E1) emerging from them. Measurements of recognised structures show similar dimensions using these optical sectioning methods and resin section reconstruction methods (Tab.4).

In addition to morphology SHG can detect collagen without the need for fixation or labelling. Coherent emission in SHG mode in unfixed glomeruli revealed a signal consistent with banded collagen which when overlaid with co-registered TPF images was positioned in the AVC walls (Fig.7, Supplemental video S3). The collagen sheath extended throughout the AVC and a short distance along the attached conduit vessel walls. A similar banded collagen signal was also seen in fixed tissues. TPF imaging showed fresh glomeruli with extensive vessel collapse in the filtration networks but VCs appeared resistant to collapse as was found with resin section reconstruction and resin single sections.

**VC wall appearance under electron microscopy**

No visible sign of collagen fibres could be seen in the 1µm light microscopy resin sections. Electron microscopy sections of AVC showed regions of banded collagen fibres in the surrounding mesangial matrix. The banding was sparse and poorly stained (30±1nm band spacing) and width of the fibres (30±2nm) in this partial sheath was consistent with Collagen I and III (Fig.8a-d). The collagen bundles extended to a depth of
4µm from the VC surface (Fig.8C; Tab.4). The endothelial lining of AVC contained few fenestrations together with cellular distortions and membrane blebs (Fig.8C), unlike the abundant fenestral density of the filtration capillaries.

**Discussion**

**Vascular chambers**

Human glomerular microvascular architecture is not as depicted in current texts. The vascular layout developed over the last 170 years since William Bowman (5) is of a single afferent arteriole which branches until filtration capillaries are reached. These filtration capillaries converge to form a single efferent arteriole conveying blood to the peri-tubular vasculature. This classic picture has been built up from biopsies or necropsies of mammalian kidney.

In human glomeruli both arterioles exhibit vascular widenings more frequently associated with low pressure veins (venous sinuses of the brain) or with large arteries (carotid sinus). However, the glomerular VCs are high pressure arteriolar afferent and efferent chambers with multiple openings, the closest definition in physical terms is a plenum manifold (plenum - a chamber containing pressurized fluid to control distribution; manifold - a pipe or chamber branching into several openings).

Plenums and manifolds in industry stabilize, distribute or balance fluid flow through multiple inlets and outlets (i.e. inlet and exhaust manifolds on internal combustion engines). Therefore, our initial hypothesis for glomerular vascular chambers is that they function to balance the pressure and/or flow through the intervening filtration regions without the need for conventional branching within the confined space of the glomerulus. These haemodynamic considerations are not relevant in smaller rodent glomeruli with smaller perfusion volumes relative to arteriolar conductivity (see introduction).

These VC manifolds persist in the glomerulus despite pressure changes, VC walls are resistant to collapse during immersion fixation or when observed fresh at zero pressure. The VC position at the vascular pole allows mesangial structural support and Collagen I/III appears to provide (additional) structural integrity. The physiological significance of this collapse resistance is not yet clear.

Collagen III has been observed in glomeruli of collagen nephropathies (7, 14) with collagen III in mesangium and/or capillary walls. No report could be found of Collagen I or III in mesangium of normal glomeruli and this report is the first to find banded Collagen (I and/or III) in normal glomeruli close to the vascular pole. Banded collagen has previously been found in kidney cortex, where 30nm fibres showed hybrid labelling with Collagen I and III (13), however, the identity of VC wall banded collagen remains to be confirmed by immunohistochemistry.

VCs appear to be ubiquitous in the adult kidney. We confined resin section reconstructions in this study to subcapsular glomeruli to surmount any size difference between subcapsular and juxtaglomerular glomeruli seen in humans and other species.
Even so, the resin single section work shows a surprisingly similar occurrence of vascular widening in 50-60% of vascular pole glomerular profiles (Fig.6), implying that VCs exist in both cortical locations with similar sized VCs in both juxtamedullary and subcapsular glomeruli.

**Afferent and efferent arterioles**

No previous study has measured the diameter of fully opened human glomerular arterioles perfusion fixed at their operating pressures. Previous human AA diameters vary from 13-16µm (18) to diabetic biopsy diameters of 29µm for AA and 19µm for EA(44). Other than biological variability, this range of arteriolar diameter is likely due to: volume changes in tissue processing, oblique sections of vessel or low pressure fixation producing collapsed profiles (for example; Tab.4 fresh AA - 13.8µm; Fig.1 in ref.(45)). These problems appear minimized with the fixation and resin embedding techniques of this paper.

A correlation between afferent arteriolar diameter and mean glomerular capillary area has previously been seen as consistent with loss of autoregulation (18). Here a correlate of AA resistance per unit length (R'AA) did not scale with any other glomerular parameter measured including R'EA (Tab.3) preserving the independent autoregulatory control of AA. In contrast EA resistance per unit length (R'EA) was inversely correlated with V_G (Fig.5D; Tab.3), and correlating remarkably with R'_Con at the afferent end (Tab.3). Unlike AA, EA is linked in fluid dynamic terms with the Glomerulus it drains.

**Conduit vessels**

The first order afferent vessels or conduits were noted by Bowman in 1842, with 2 to 8 branches which visibly ‘subdivide only once or twice as they advance over the surface of the ball’ (5). The few buried deep inside the glomerulus unseen by Bowman may explains the result of 2 to 11 seen in this current study. We also confirm the luminal width of these first order afferent vessels as being as wide as the efferent arteriole (21).

Conduit vessels show fewer branches than their efferent counterparts but branch frequency increases at the start of perfusion regions often at some point on the glomerular periphery (Fig.2). No previous branch data exists for these vessels however, the interbranch length for all rat glomerular vessels at 26.3±24.9µm(SD) (48) is between the medians, 32.8µm (conduit afferent) and 15µm (efferent) of the skewed distributions found here.

Conduit vessels close to the AVC are embedded in mesangium, those distal to the AVC have a GFB. While detailed conduit ultrastructure remains to be confirmed, no aberrant GFB capillary morphology has been noted in all our studies of normal human glomeruli (data not shown). It appears that conduit GFB is similar to filtration capillary GFB except for the scarcity of podocyte cell bodies on the conduit GFB surface. It remains to be determined if conduit podocytes are just responding to local conditions or are a sub-population of conduit podocytes with the extra-long major processes necessary to cover the GFB area in foot processes.

The GFB is known to remain intact and expand under excess pressure (25, 27) and conduit vessels with a 86-100% GFB - or a sparse 0-14% mesangial attachment around the circumference showed diameter expansion by 7% compared to conduit vessels surrounded by and embedded in mesangium (Fig. 5A) - not enough GFB expansion to
explain podocyte cell body free areas on the conduit vessels but below the damaged
‘giant capillary’ inflation levels previously reported (25). Conduit inflation might be
expected considering the reduced podocyte coverage, thin walls and wide diameter and
estimates of wall forces show conduit vessels with a high proportion of GFB and low
mesangial attachment are the most susceptible to hoop stress of all glomerular vessels
(Appendix 2). This marks conduits as a target in hypertensive disease and hoop stress
failure has been observed in rat primary afferents (equivalent to conduits) due to
glomerular hypertension (with marking albuminuria and glomerulosclerosis) (26).

The subpodocyte space, identified under podocytes (39) should be present under
conduit podocyte cell bodies (awaiting EM confirmation). Incidentally, the light
microscopy derived filtration capillary podocyte cell body (PCB) area coverage of 55%
of the GFB fits well with the electron microscopy derived subpodocyte space coverage of
60% for filtration capillaries found previously (41, 50) suggesting most of human
subpodocyte space is under the podocyte cell body.

Other evidence for vascular chambers and conduits
Reconstructed rat glomeruli do not show vascular chambers (48). We confirmed
these findings by reconstructing rat glomeruli with Serial Block Face Scanning Electron
Microscopy (data not shown) and also found no evidence of VC.

Mammalian arterioles can widen pathologically (32), for instance, mesangiolysis
can remove mesangial support causing glomerular vessel aneurysms (35) but such
features would not be as highly conserved in shape or have an organized collagenous
support as seen in VC found here. Bowman also noted in the larger horse glomerulus that
afferent arterioles dilate on the surface prior to dividing but not in human glomeruli (5)
we show here that human glomerular vascular dilations are subsurface and would have
been invisible to Bowman. The modern conventional description merely reports that the
afferent arteriole branches into the glomerular capillary network (22).

VCs may not be present in all human glomeruli, during development, glomerular
capillaries arise from one dilated vessel (11) and neonate vascular widening has been
shown prior to the five first order afferent branches (21) although this has been ascribed
to a vessel remnant from the developing nephron (11). Interestingly, the glomerular
diameter increase in children from 112µm (birth) to 167µm (15years) (34) and VC
scaling with V_G shows that VCs may not exist in child glomeruli which are below 160-
180µm diameter, providing these glomeruli follow the adult glomerular correlation (Fig.
3B & 4C,D). Conduit vessel resistance (R'_con) also scales with V_G, whether this
correlation continues in smaller (child) glomeruli or whether the primary afferents in
children even constitute ‘conduit’ vessels needs evaluation.

Renal biopsies do occasionally show evidence of VCs and conduit vessels in
section, a survey of images in biomedical journals reveal light micrographs showing a
15µm conduit vessel and 20µm VC (20), a 30µm diameter VC (52) and VCs at both
efferent and afferent ends (44). However, without the context of a serial section stack
these micrographs remain as widened vascular profiles.

VCs could be artefacts of processing volume changes, however, glomerular
diameters (~200µm) derived here were between immersion fixed (160 - 170µm) (10, 31)
and autopsy diameters (260-270µm) (8) and closely match in vivo ultrasound values of
200µm (15, 23), suggesting glomerular volume changes during processing were minimal overall.

Wide profiles at the vascular pole in single sections can be dismissed as collapsed vessels. Put simply, an afferent arteriole terminus of 21µm diameter with a circumference of 66µm could conceivably collapse to a flattened squashed-circle profile approximately 30µm wide which if sectioned longitudinally would fit exactly with the 28-30µm wide profiles measured, however, 60% of randomly orientated single sections of vascular poles all showed these wide vascular regions - far too frequent for the collapse argument. Additionally, in this study vessel collapse was seen in filtration capillaries in fresh glomeruli (multiphoton microscope: Supplemental Fig. S3) but with VCs held open. VCs are not collapse artefacts but stiff walled vascular structures.

The Murray relationship

The relationship between branching vessel diameters was derived by Murray on the principle of minimum work for blood flow (36, 37) where the radius cubed of the parent vessel equals the sum of the cubes of the daughter vessel radii. The Murray relationship holds for arteries and venules of rat kidney down to the afferent arterioles and venules leading away from the tubular networks (42), but it is not known if it continues into the glomerulus. The Murray relationship in whole human kidney also remains to be assessed.

A Murray constant (K) was calculated for each set of vessels leading into and away from human glomerular VCs in all 14 glomeruli reconstructed from resin sections:

\[ K = r^3 n_V \]  \hspace{1cm} \text{eq.2}

Where \( n_V \) is the number of vessels and \( r \) is the radius. Using \( r_{AA}, r_{AVC}, r_{Con}, r_{E1}, r_{EVC}, r_{EA} \) and appropriate \( n \) to calculate \( K \), the Murray relationship breaks at the VCs and the first order vessels (conduit and E1 vessels; Fig.6B), where daughter vessels do not have the same Murray constant as parent vessels.

This is an exception to Murray’s Law – a plenum/manifold exception, where flow distribution from a single arteriole provides a high pressure distributive flow into many glomerular lobes in a short distance. An estimate of K values for second order afferent vessels (A2 in 2 glomeruli) showed that K may return to the value predicted by the afferent arteriolar radius after skipping the VC and conduit vessels (Fig.6B). Other Murray’s law exceptions occur where a higher surface area is required in the exchange vessels of an organ, for instance alveolar capillary networks (59).

The possible mechanisms producing a set of vessels following Murray’s law includes an endothelial transducer triggering remodelling after a shear force threshold was exceeded(46). Altering the threshold could induce the vessel diameter changes seen here. However, the Murray relationship requires laminar flow through vessels and the haemodynamic flow will be complex from an afferent arteriole into an ellipsoidal vascular chamber with several outlets.

VC haemodynamics

If glomerular volume is used as a measure of perfusion capacity, it rises and falls along with the size of the AVC and the EVC (Fig.4 C&D). Larger AVCs feed more
blood to larger glomerular filtration regions and thence to larger EVCs. As the size
increases the resistance of the conduit vessels, E1 and EA (not AA) falls to accommodate
the flow (vessels get wider in proportion to Poiseuille flow) (Fig.5 C&D). All of the
major vessels of the human glomerulus past the afferent arteriole are linked in some way
in terms of flow and capacity (Tab.3). How would flow progress from laminar flow in an
afferent arteriole through the AVC to the conduit vessels? And similarly from efferent E1
vessels through EVC to the efferent arterioles?

A clue to VC flow characteristics comes from the kinks and bends in AAs. One
constant feature of the glomeruli analysed is the bend as the afferent arteriole enters the
AVC. These bends can be readily seen in the glomeruli of figures 1, 2A and 2B
(supplemental videos 2a and 2b) and showed an average 60° deviation from a straight
path. The fluid flow at a bend in a channel is known to induce vortices
(49), we hypothesize that the summation of all bends in an afferent arteriole (i.e. see bend from
interlobular –AA junction in Fig.1) could induce a single major vortex in the AVC
possibly aiding distributive flow centrifugally into conduit vessels.

If such a vortex with its axis in the midline of the AVC adopts the properties of a
“rigid-body” or “rotational” vortex, then the pressure at the AVC edge at the conduit
vessel openings would depend both on the hydrostatic pressure and the dynamic pressure
(set by the angular momentum of the moving fluid – ½ρω² where ρ=density; ω = angular
velocity). Crucially however the dynamic pressures within this form of vortex are
uniform (3).

We speculate that in health the AVC and the complex (vortical) fluid movement
within it, may ensure a uniform driving pressure into the conduit vessels – maximising a
uniform distribution of flow to each of the glomerular lobules. The loss of this equalising
distributary mechanism through microvascular disease, mesangial proliferation occluding
the AVC, hyperperfusion or immunological injury, could potentially result in localised
hyperfiltration and excess shear stress in some glomerular segments with stasis in others.
This has implications for glomerular disease in which only some perfused regions of the
glomerulus appear to have sustained sclerotic/fibrotic damage (eg FSGS) while adjacent
lobules appear normal.

The structure of the efferent vascular chamber, with many microvessels
converging on a chamber, lends itself to the development of an irrotational vortex (plug
hole vortex) balancing EVC pressure gradients and promoting balanced removal of blood
from the glomerular tuft (3).

Conclusion

We show for the first time in human glomeruli that clearly defined afferent
arterioles lead into afferent vascular chambers of ellipsoid shape and structure embedded
in the mesangium of the glomerular vascular pole and ensheathed in collagen fibrils.
These chambers are plenum manifolds with many emergent relatively unbranched wide
blood vessels or conduits conveying blood to the periphery of the glomerulus. Branching
frequency increases at the end of the conduits leading to filtration capillary networks
which lead back to smaller efferent vascular chambers in the mesangium of the vascular
pole and then the efferent arteriole. The conduit vessels are sparsely covered with
podocytes, and conduit fluid resistance scales with the size of the afferent vascular
chambers. Both vascular chambers scale with glomerular capacity suggesting absence of
vascular chambers in glomeruli below 160µm diameter (the glomeruli of children). Resistance correlates of first order afferent (conduit) and efferent vessels and efferent arterioles (but not afferent arterioles) scale together and inversely with glomerular volume. We propose that all these structures represent a large glomerulus adaptation allowing even haemodynamic flow distribution and pressure balance across the many lobes of a human glomerulus.

Appendix 1.

Vascular resistance

The vascular resistance to flow will change as blood flows along AA into AVCs and conduits and later pools in EVCs before flowing into EA. To better understand how blood flow is affected by the changing morphology a correlate of vascular resistance $R_{Con}$ was derived from the Poiseuille equation using vessel radii and vessel number. For VCs the flow will be complex and non-laminar in the spheroidal shape and so the Poiseuille equation could not be used so VC volume was used as a measure of VC capacity.

Resistance changes in arterioles and conduit vessels

For conduit vessel resistance ($\sum R_{Con}$) coming out of the afferent VC where $R_{Con3}$ is the resistance of the 3rd conduit vessel in parallel:

$$\frac{1}{\sum R_{Con}} = \frac{1}{R_{Con1}} + \frac{1}{R_{Con2}} + \frac{1}{R_{Con3}} + \ldots + \frac{1}{R_{Conn}} \quad \text{eqA1.1}$$

For $n_{Con}$ similar conduit vessel resistances $R_{ConX}$

$$\frac{1}{\sum R_{Con}} = \frac{n_{Con}}{R_{ConX}} \quad \text{eqA1.2}$$

For fluid of viscosity $\eta$, the resistance to flow through a tube of length $L$ is inversely proportional to the $4^{th}$ power of the radius (Poiseuille’s law), similarly:

$$R_{ConX} = \frac{8 \eta_{Con} L_{Con}}{\pi r_{Con}^4} \quad \text{eqA1.3}$$

Where $L_{Con}$ is conduit vessel length and $r_{Con}$ the mean conduit vessel radius. If the viscosity of the blood flowing through VC and attached vessels ($\eta_{Con}$) is assumed not to
change (low filtration into mesangium in these vessels) then $\eta_{Con}$ with $\pi$ and $\delta$ can be combined into a constant $k_{Con}$:

$$R_{ConX} = \frac{k_{Con} L_{Con}}{r_{Con}^4} \quad \text{eqA1.4}$$

Combining equation eqA1.2 and eqA1.4:

$$\frac{1}{\sum R_{Con}} = \frac{L_{Con}^{-4} n_{Con}}{k_{Con} L_{Con}} \quad \text{eqA1.5}$$

Inverting eqA1.5 and dividing by $L_{Con}$ and $K_{Con}$ yields a measure of the total conduit vessel resistance per unit length ($R'_{Con}$).

$$\frac{\sum R_{Con}}{L_{Con} k_{Con}} = \frac{1}{r_{Con}^{-4} n_{Con}} = R'_{Con} \quad \text{eqA1.6}$$

$1/r_{Con}^{-4} n_{Con}$ was used to estimate a correlate of vascular resistance per unit length of all conduit vessels in parallel ($R'_{Con}$). Similarly, 1st order efferents were assessed using $1/r_{El}^{-4} n_{El}$ ($R'_{El}$). Correlates of afferent and efferent arteriole resistance per unit length ($R'_{AA}$, $R'_{EA}$) were estimated with $1/r_{AA}^{-4}$ and $1/r_{EA}^{-4}$.

Appendix 2.

Vascular wall stress

The conduit vessel wall morphology appears similar to filtration capillaries; however, conduits are much wider. Greater diameter tubes or vessels of the same wall thickness are more susceptible to pressure damage or rupture. How might conduit vessel wall stress compare with other glomerular vessels?

VC and conduit vessel wall stress

The effective wall strength and compliance of systemic capillaries is largely due to basement membrane/basal lamina (40). Assuming that glomerular vascular wall strength is due to the glomerular basement membrane (GBM, 0.3µm and less than 1/10th of vessel radius) then the Laplace equation (60) can be used to derive the hoop stress ($S_h$) of the vascular wall (the force exerted circumferentially trying to pull the wall apart). For cylindrical conduit vessels:

$$S_{hCon} = \frac{\Delta P_{Con} L_{Con}}{t_{Con}} \quad \text{eq.A2.1}$$

Where $\Delta P_{Con}$ is the hydrostatic pressure difference across conduit vessel wall of radius $r_{Con}$, and effective wall thickness $t_{Con}$.

The equation for a near spherical VC is half that of an equivalent diameter cylinder:
\[ S_{hVC} = \frac{\Delta P_{VC} r_{VC}}{2 t_{VC}} \quad \text{eq.A2.2} \]

Where \( \Delta P_{VC} \) is the hydrostatic pressure difference across the VC wall of radius \( r_{VC} \), and effective wall thickness \( t_{VC} \). The effective strength of the arteriolar wall will be a composite of strengths of this thick multilayered structure, however, the arteriole smooth muscle wall thins as it transitions into the VC with only endothelium, basal lamina and collagen sheath surrounded by mesangial matrix.

**Parameters used in Calculations**

**\( S_{hAVC} \) for afferent VC (AVC):**

\( r_{AVC} = 22\mu m \) [mean of \( r'_{AVC}, r''_{AVC}, r'''_{AVC}; \text{Tab.1} \)],

\( t_{AVC} = 0.5-4\mu m \) [between the first mesangial lamina thickness \( \sim 0.5\mu m \) (see Fig.8d) and the collagen sheath dispersed over 4\( \mu m \) (Tab.4, Fig.8)]

\( \Delta P_{AVC} = 23\text{mmHg} \) [AVC luminal pressure of 63mmHg (43) minus mesangial pressure - a high proportion of capillary hydrostatic pressure (9) - likely 40mmHg since mesangial cells respond to 40mmHg and above (19, 30)].

\( S_{hAVC} = 8 - 66 \text{ kPa} \) (equation A2.2) \( \leq 8 \text{ kPa} \) if the effective \( \Delta P_{AVC} \) is lower due to pressure dissipating gradients and effective \( t_{AVC} \) thicker due to additional mesangial matrix support (7)

**\( S_{hMC} \) for mesangial conduit vessel (MC):**

The mesangial backed conduit vessels (Fig.3a, Mes.Con) adjacent to AVCs would share the same mesangial protection and possibly collagen sheath as the AVCs.

\( r_{MC} = 8\mu m \) [Tab.1]

\( t_{MC} = 0.5\mu m \) to 4\( \mu m \) [Tab.4, Fig.8]

\( \Delta P_{MC} = 23\text{mmHg} \) [see above]

\( S_{hMC} = 6 - 48 \text{ kPa} \) (equation A2.1) \( \leq 6 \text{ kPa} \), \( S_{hAVC} \) caveat as above)

**\( S_{hGC} \) for glomerular filtration barrier conduit vessel (GC):**

The conduit vessels away from the AVC are connected to mesangium only on a small part of their circumference the rest being normal GFB and GBM (Fig.3a, GFB Con)

\( r_{GC} = 8\mu m \) [Tab.1]

\( t_{GC} = 0.3\mu m \) [GBM thickness]

\( \Delta P_{GC} = 38\text{mmHg} \) [luminal \( P \) (63mmHg) minus urinary space \( P \) (25mmHg)]

\( S_{hGC} = 133 \text{ kPa} \) (equation A2.1)

**\( S_{hFC} \) for filtration capillaries (FC):**

\( r_{FC} = 3.5\mu m \)

\( t_{FC} = 0.3\mu m \) [GBM thickness]

\( \Delta P_{FC} = 38\text{mmHg} \) [luminal \( P \) (63mmHg) minus urinary space \( P \) (25mmHg)]
\[ S_{hFC} = 58 \text{ kPa} \quad (\text{equation A2.1}) \]

| Subscript abbreviation | ∆P (mmHg) | r (µm) | t (µm) | \( S_h \) (kPa) |
|------------------------|-----------|--------|--------|---------------|
| Afferent VC AVC        | 23        | 22     | 0.5-4.0| ≤ 8-66        |
| Mesangial conduit MC   | 23        | 8      | 0.5-4.0| ≤ 6-48        |
| GFB conduit GC         | 38        | 8      | 0.3    | 133 *         |
| Filtration Caps. FC    | 38        | 3.5    | 0.3    | 58            |

**Table A1.** Calculated vascular hoop stress \( S_h \). The peak is in the GFB conduit vessels (*).

\( S_h \) is difficult to estimate in the mesangial backed AVC and mesangial conduit vessels but our maximum estimate is less than half the value for the GFB Conduit. \( S_h \) falls in the filtration capillaries of the same wall thickness but these are protected by their small radius. At the efferent end the reduced radii and complete mesangial encasement of VC and of the short E1 vessels would result in lower \( S_h \) of the equivalent efferent vessels (not shown).

In conclusion, in human glomeruli, GFB conduit walls (GC) mark a peak of hoop stress caused by the relatively thin wall for the large diameter. While the AVC and the early conduit vessel are protected by mesangial backing, any mesangial disruption through immune-mediated damage, cell invasion or proliferation or disruption to the collagen sheath will change \( S_{hAVC} \) and \( S_{hMC} \) making AVC and mesangial conduit vessels vulnerable to pressure changes.

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Figure 5. Conduit diameter changes with mesangium; conduit podocyte attachment; resistance v capacity examples  A/ Conduit diameter changes relative to mesangial cover. Conduit vessel diameters adjacent to the afferent VC with mesangial cover of 80-100% (GFB coverage 0-20%) were compared with diameters of low mesangial covered (distal) regions of the same vessel. The fold change in diameter shows a significant diameter increase of 7.4% (*) when mesangial cover is minimal (0-14% i.e. GFB 86-100%). Paired t-tests and Wilcoxon matched pair test (P=0.04). B/ Histogram of podocyte cell body (PCB) area coverage of the filtration barrier of conduit vessels (filled bars) and small filtration capillaries (open bars). Conduits have significantly less PCB coverage of the GFB than filtration capillaries (ttest - P<0.0001). C/ Conduit resistance versus Afferent VC volume. A significant negative correlation exists between a correlate of conduit resistance (R’_{Con}) and afferent VC volume (V_{AVC}) (R^2 = 0.327, P=0.033). D/ Efferent arteriole resistance per unit length (R’_{EA}) reduces in line with increasing V_G (R^2 = 0.47, P=0.007).

Figure 6. Vascular widenings in single sections. Murray constant from vascular radii  A/ Observed occurrence of glomerular vascular widening in single sections. The frequency with which widening (implying VC presence) was observed at vascular poles in immersion and perfusion fixed glomeruli. SC - subcapsular glomeruli; JM - juxta-medullary glomeruli; JMSC - JM and SC glomeruli combined. (n = number of kidneys)  B/ In 14 glomeruli a Murray constant (K = r^3 n_v; where r is radius, n_v is vessel number; see text) was calculated for the afferent and efferent arteriolar tree leading through the VCs and thence into the 1st order vessels (Con and E1). In 2 glomeruli K was calculated for 2nd order vessels. The Murray relationship of equal K at each vessel level is absent in the AVC, EVC and conduit vessels.

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Supplemental

Supplemental video legends

Video S2a. Image stack for Fig 2a glomerulus.
Video to show the full image stack formed by Image J software from original 1µm serial section images. Stills in Fig.2a. Field of view 170 x 200µm approximately.

Video S2b, Image stack for Fig 2b glomerulus.
Video to show the full image stack formed by Image J software from original 1µm serial section images. Stills in Fig.2b. Field of view 190 x 220µm approximately.

Video S2c. Reconstruction x derived from Fig. S2b.
Red afferent arteriole derived vessels meeting with blue efferent arteriole derived vessels at purple points. Rotation around x axis. Not all vessels shown. Scale marks in µm.

Video S2d. Reconstruction y derived from Fig. S2b.
Red afferent arteriole derived vessels meeting with blue efferent arteriole derived vessels at purple points. Rotation around y axis. Not all vessels shown. Scale marks in µm.

Video S3. Reconstruction of an unfixed glomerulus from multiphoton microscope images. TPF and SHG modes were used to image the vessel walls (green) and banded collagen (blue) respectively. The intense blue signal from the collagen of Bowman’s capsule was covered by a circular black mask. The afferent arteriole opens into a VC at the 7 o’clock position, the banded collagen signal follows the walls of the VC and into the conduit vessels. Field width 200µm
Footnote 1

Afferent arteriole conductance estimated from the 4th power of vessel radii (mice, \( r = 5-6.5\mu m \), rats, \( r = 7-9.5\mu m \); human, \( r = 11\mu m \) [this article]) with human glomerular volume estimated from glomerular diameter (mouse=70\( \mu m \), rat=120\( \mu m \) and human=200\( \mu m \))
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|                     | AA       | AA       | AVC       | AVC Coll. | Conduit   | E1       | EVC      | EA       | EA       |
|---------------------|----------|----------|-----------|-----------|-----------|----------|----------|----------|----------|
|                     | Diam. (µm) | Wall t (µm) | Diam. (µm) | Wall t (µm) | Diam. (µm) | Diam. (µm) | Diam. (µm) | Diam. (µm) | Wall. t (µm) |
| Fixed Resin Recon.  | 14G, 4K  | 21.5±1.2 | 6.6±0.3  | 43.2±2.8  | *         | 15.9±0.7 | 9.9±0.4  | 38.4±4.9 | 15.9±1.2 | 3.0±0.1   |
| Fixed Aq. Confocal  | 4G*, 1K  | 28.4±1.9 | 6.3±0.8  | 35.8±3.5  | *         | 16.0±1.2 | 8.2      | 24.2     | 12.8     | *         |
| Fixed Aq. Multipho  | 3G*, 1K  | *        | *        | 50.2±3.7  | 4.2±0.8   | 12.8±1.6 | 6.9      | 28.1     | 7.4      | *         |
| Fresh Aq. Multipho  | 3G*, 2K  | 13.8     | 3.0      | 35.8±4.1  | 4.1±1.9   | 14.4±0.9 | *        | *        | *        | *         |
| Fresh Aq. Multipho  | Isolated I  G*, 1K | 23.0 | *        | 54.2      | 2.5       | 27.4     | *        | *        | *        | *         |