Title
The origins of urinary stone disease: upstream mineral formations initiate downstream Randall's plaque

Permalink
https://escholarship.org/uc/item/0sk8n6vb

Journal
BJU INTERNATIONAL, 119(1)

ISSN
1464-4096

Authors
Hsi, RS
Ramaswamy, K
Ho, SP
et al.

Publication Date
2017

DOI
10.1111/bju.13555

Peer reviewed
The origins of urinary stone disease: upstream mineral formations initiate downstream Randall’s plaque

Ryan S. Hsi*, Krishna Ramaswamy*, Sunita P. Ho† and Marshall L. Stoller*

*Department of Urology, and †Division of Biomaterials and Bioengineering, Department of Preventive and Restorative Dental Sciences, School of Dentistry, University of California San Francisco, San Francisco, CA, USA

S.P.H. and M.L.S contributed equally to this work as co-senior authors.

Objectives
To describe a new hypothesis for the initial events leading to urinary stones. A biomechanical perspective on Randall’s plaque formation through form and function relationships is applied to functional units within the kidney, we have termed the ‘medullo-papillary complex’ – a dynamic relationship between intratubular and interstitial mineral aggregates.

Methods
A complete MEDLINE search was performed to examine the existing literature on the anatomical and physiological relationships in the renal medulla and papilla. Sectioned human renal medulla with papilla from radical nephrectomy specimens were imaged using a high resolution micro X-ray computed tomography. The location, distribution, and density of mineral aggregates within the medullo-papillary complex were identified.

Results
Mineral aggregates were seen proximally in all specimens within the outer medulla of the medullary complex and were intratubular. Distal interstitial mineralisation at the papillary tip corresponding to Randall’s plaque was not seen until a threshold of proximal mineralisation was observed. Mineral density measurements suggest varied chemical compositions between the proximal intratubular (330 mg/cm³) and distal interstitial (270 mg/cm³) deposits. A review of the literature revealed distinct anatomical compartments and gradients across the medullo-papillary complex that supports the empirical observations that proximal mineralisation triggers distal Randall’s plaque formation.

Conclusion
The early stone event is initiated by intratubular mineralisation of the renal medullary tissue leading to the interstitial mineralisation that is observed as Randall’s plaque. We base this novel hypothesis on a multiscale biomechanics perspective involving form and function relationships, and empirical observations. Additional studies are needed to validate this hypothesis.

Keywords
kidney, urolithiasis, calcification, physiological, urinary tract physiology

Introduction
Whale and dolphin kidneys are shaped in the likeness of a lung consisting of many smaller kidney bean-shaped alveoli-like renal units that resemble a small human kidney [1]. Desert rats compared to their counterparts that live in damp environments have nearly half as many nephrons and reduced filtration surface area for water economy, yet the overall shape and compartments of the kidney remain preserved [2]. The shape and size of the kidney among species varies depending on the overall size of the animal and its functional needs, yet the archetype of the renal functional unit is conserved. These and other form–function relationships provide insight into the mechanisms of physiological and pathological states within organ systems such as the kidney.

At the smallest scale, the functional unit of the kidney is considered to be the nephron. However, for urologists, the most clinically relevant functional units are the 8–12 paraboloid structures within a kidney visualised during endoscopy consisting of the renal papilla and medulla, which we term the ‘medullo-papillary complex’. This structure is observed in majority of mammalian renal systems, with some kidneys being uni-papillary systems, while others contain multiple papillae.

Regardless of the number of papilla, the paraboloid shape guides the various lengths of the tubules along its axis that
are responsible for adjusting the final urinary filtrate. We propose that this shape is a critical element and is predisposed to form biominerals. While the absolute number and density of tubules is uncertain, these functional units can be impaired due to changes in environmental stimuli. The shifts in stimuli can in turn cause functional impairment and favour stone formation while maintaining adequate urinary excretion. It is this process that we would like to apply to the kidney: environmental stimuli help define form, and that form guides function [3,4].

Current thought holds that the Randall’s plaque serves as a nidus on which stones form for idiopathic calcium oxalate stone-formers [5]. However, the formation of Randall’s plaques, and subsequent renal stones, is not well understood. We propose that Randall’s plaque formation at the tip of the medullo-papillary complex is not the headwaters but a downstream event. We hypothesised that mineral formation begins within the tubules in the proximal renal medulla and prompts a cascade of events that culminate to the distally observed interstitial mineralisation leading to Randall’s plaque formation in the papilla. We will demonstrate this by coupling the unique form of the human renal medullo-papillary complex to its function and providing evidence from high-resolution X-ray microscopy.

**Historical and Current Perspectives on Kidney Stone Formation: Homage to Randall’s Theory**

Randall originally proposed that calcium deposition at the tips of renal papillae was the precursor to renal stones and described the characteristic ‘plaque’ as calcium phosphate, not within the tubular lumen, but ‘invading and replacing interstitial tissue’ [6,7]. He also observed that Randall’s plaques increase with age. He theorised that upon these plaques, calcium phosphate and calcium oxalate stones grow and eventually break off and subsequently become urinary tract stones that often are symptomatic. This implies that the Randall’s plaque is at the intersection between papillary mineralisation and urinary tract stone *in vivo*.

Developments in endoscopic technology have demonstrated Randall’s plaque in a majority of calcium stone-formers [8]. The fractional plaque coverage in idiopathic calcium oxalate stone-formers is higher than that in non-stone-formers [9], and the percentage of plaque coverage correlates with the number of stones formed [10]. Furthermore, endoscopically observed stones are often attached to renal papilla at sites of plaque [11].

Advanced imaging methods have been used to characterise the origins of Randall’s plaque from biopsy specimens from the papillary tip [12]. The identified mineral deposits are located within the interstitium of the renal papillary tip and often are thought to originate from the basement membrane of the thin loops of Henle. Plaques are thought to originate as spherical deposits of apatite as small as 50 nm in the interstitial matrix and are proposed as the beginnings of later plaque development that Randall had observed [13]. However, because these observations have been obtained from papillary biopsies, the upstream events that lead to Randall’s plaque formations have not been examined and are not understood.

**Medullo-Papillary Complex Form and Function**

The medullo-papillary complex is shaped like a paraboloid (a parabola rotated about its long axis), extending into the minor calyx of the collecting system (Fig. 1). Each of the 8–12 medullo-papillary complexes within the kidney is comprised of three anatomically distinct medullary zones based on the segments of the nephron: Zone 1, consists of the outer stripe of the outer medulla; Zone 2, the inner stripe of the outer medulla; and Zone 3, the inner medulla (Fig. 1) [2,14].

*Longitudinally*, nephrons can be divided into two major populations of long and short nephrons based on the length of the loops of Henle (Fig. 2) [2,15]. Compared to long-loop nephrons, short-loop nephrons are limited to Zones 1 and 2, and have no thin ascending limb segments. The long-loop nephrons form 180° bends predominantly in Zones 2 and 3 as they travel deeper towards the tip of papilla and become fewer in number, which affects urine concentrating ability [16]. The paraboloid shape mandates that there are short peripherally located nephrons and long centrally located nephrons in the medullo-papillary complex.

Blood vessels lie in close proximity to these loops to facilitate countercurrent exchange of ions and water in the transverse direction [17]. As the descending vasa recta enter Zone 1 at various levels, they dive longitudinally with little or no transverse branching towards the tip of the papilla, where they each make a 180° U-bend and split into two to three ascending vasa recta. From the base to the tip of the medullo-papillary complex, both the lumen diameter and the pressure within the vasa recta decrease as they approach the tip of the papilla [18]. Surrounding the descending vasa recta are one to three layers of smooth muscle cells in Zone 2, which disappear as they approach the tip of the papilla where pericytes replace them [17–19]. Pericytes are known to have mesenchymal stem cell-like properties including osteogenic phenotype [20]. Within Zone 3, the smooth muscle cells disappear, suggesting that deeper blood flow is not regulated by contractile mechanisms [21]. Non-fenestrated endothelium of the descending vasa recta becomes fenestrated, and after the U-bend converts to ascending vasa recta affecting water and small solute...
reabsorption along the length of the medullo-papillary complex [17].

Transversely, the medullo-papillary complex is divided into anatomically specific functional compartments that generate concentration gradients (Fig. 2) [14]. Zone 2 is divided into vascular bundles and interbundle regions [17,22]. The vascular bundles contain the descending vasa recta and ascending vasa recta that travel exclusively together with no branching [22]. The longest descending vasa recta emerge at the tip of the papilla into Zone 3, where there is higher interstitial osmolarity, and are located more centrally within these vascular bundles [23]. Within the interbundle regions are the thin limbs and collecting ducts. The vascular bundles and interbundle regions have minimal communication because the descending thin limbs and collecting ducts are located within the centre of the interbundle region, and the water-impermeable thick ascending limbs lie at the margin of the vascular bundles [22].

In contrast, Zone 3 is divided into collecting duct clusters with large diameters and vascular bundles of narrow...
diameters (Fig. 2) [21]. The collecting duct clusters predominately contain the collecting ducts, ascending thin limbs, and possibly water-impermeable descending thin limbs. The vascular clusters are dominated by ascending and descending vasa recta, and contain water-permeable descending thin limbs. Whereas the descending thin limbs associate closely with the collecting ducts in Zone 2, they anatomically transition to lie distant from collecting ducts in Zone 3. The physical separation of these anatomically specific compartments contribute to radial and axial concentration gradients along the course of the complex.

Pressure gradients exist along the medullo-papillary complex (Fig. 3) [19]. Longitudinally, pressure within Zone 1 is higher than in Zone 3 and drives perfusion and tubular flow. Among the descending vasa recta, the larger diameter vessels are located within the centres of vascular bundles and travel towards the tip of the renal papilla [19]. From proximal to distal, the long-looped nephrons associated with the descending vasa recta centrally have higher pressure gradients than the short-looped nephrons located peripherally in the medullo-papillary complex.

Chemical gradients within the medullo-papillary complex facilitate the formation of concentrated and dilute urine (Fig. 3). The medullary osmotic gradient is generated by active NaCl reabsorption from the thick ascending limb in Zones 1 and 2, maintained by the descending thin limb (water permeable but low permeability to Na and urea) and ascending thin limb (high Na permeability, low urea permeability, and impermeable to water) segments in Zone 3, and augmented by passive urea and water reabsorption in the collecting ducts [19,26]. Adding to the complexity of these gradients are the spatial relationships of the tubules and vessels within the compartments, generating local concentration gradients [17,21]. Finally, tissue oxygen gradients present within the medullo-papillary complex create areas of relative hypoxemia in the thick ascending limbs and medullary collecting ducts due to the relatively high metabolic demands of the channels in these zones [23,27,28].

Proposed Mechanism of Biominalisation from Proximal to Distal Regions in the Medullo-Papillary Complex

The rate of flow within the medullo-papillary complex can be described by Poiseuille’s equation, where the flow rate is
directly proportional to the pressure gradient along the tube, fourth power of the radius, and is inversely proportional to the fluid viscosity and tube length. The fluid velocity near the wall is minimal compared to the centreline velocity due to the resistance of flow from the tubular wall [29].

These fundamental principles can be applied at two levels: (i) at the level of the medullo-papillary complex, and (ii) at the level of the tubules within the complex. Taken together, pressure gradients within the peripherally oriented, shorter tubules within the paraboloid complex are lower than the centrally located tubules (Fig. 4). Given, the lower pressure gradients along with lower velocities at the tubular walls, over a lifespan, particulates can accumulate along the walls of these shorter tubules in Zones 1 and 2, akin to a stream where the floating leaves accumulate at the edges of the bank of a stream. As these shorter, peripheral tubules become progressively obstructed, there is a decrease in the functional volume of the overall complex. In other words, the functional ‘volume’ of tubules is reduced once the peripheral tubules are devoid of flow. As such, more flow is diverted centrally, which changes the internal pressure and subsequently the ‘stiffness’ of the medullo-papillary complex specifically at the interface where the shorter tubules end and the longer tubules continue to traverse into Zone 3. At this interface between Zones 2 and 3 could be a mechanoresponsive switch that is turned ‘on’ due to a shift in form (as the functional radius decreases to the fourth power, the pressure within the medullo-papillary complex increases significantly), and fluid is shunted to the centrally located tubules. Once this mechanoresponsive switch is turned on, a cascade of biological events is triggered distally. The centrally located smaller diameter yet pressurised vasa recta and loops of Henle develop a differential in the ‘hoop’ or circumferential strain at the interface between Zones 2 and 3 and plausibly could leak into Zone 3. At this interface, we hypothesise that integrins facilitate transfer of change in stiffness of the extracellular matrix to intracellular compartments, and that this differential in hoop strain could predominantly impact
pericytes located near the tip of the medullo-papillary complex. Pericytes are known to have progenitor-like phenotype and can differentiate into an osteoblastic lineage making the environment conducive to sequester inorganic ions – the early stages of cell-based biomineralisation [20]. It is through this postulate that we explain the formation of the apatite substrate: the Randall’s plaque. A second proposed mechanism involves elevated hydrostatic pressure in the vasa recta driving solutes into the interstitium through the fenestrated endothelium, which overwhelms resorption capability of the tubular cells in Zone 3, leading to accumulation of interstitial biominerals.

**Evidence for Mineralisation in the Medullo-Papillary Complex: Materials and Methods**

In an Institutional Review Board-approved study, radical nephrectomy specimens for renal tumour ($n=12$) were sectioned to isolate the medullo-papillary complex in grossly normal appearing kidney tissue, sufficiently distant from the malignancy in question. No kidneys with urothelial tumours were examined. All patients had no prior history of stone disease. The specimens were placed in a centrifuge tube in 50% ethanol and were mounted inside a calibrated micro-CT unit (MicroXCT-200; Carl Zeiss X-ray Microscopy, Pleasanton, CA, USA). The specimens were imaged at $4\times$ magnification, 40 kV, 8 W, 5 s exposure time, pixel size 5.01 μm, and with 2,000 image projections. Reconstructed digital volumes (using XMReconstructor software) were processed to determine mineral densities in milligrams per cubic centimetre and were related to surrounding tubular structures.

**Results**

In all specimens, micro-CT demonstrated streaks and aggregates of mineralised regions in Zones 1 and 2 within the medullo-papillary complex that correspond to more peripherally located tubules (Fig. 5A). Mineralisation was predominately intratubular in location. Only the specimens with greater volume of proximal, intratubular mineralisation were observed to have distal interstitial mineralisation near the papilla (Fig. 5B,C).

This suggests that proximal mineralisation could precede distal mineralisation. In other words, interstitial deposits were observed in association with the central nephrons in the inner medulla at the papillary tip, and never in isolation without the
more proximal peripheral intratubular mineralisation. Furthermore, the mean mineral densities of the proximal mineralisation were higher (330 mg/cm³) than that of the distal papillary region (270 mg/cm³). The volume fraction of mineralisation was distinct proximally compared to distal regions (0.2% vs 4.5%). These two mineralisation processes occur not only in separate locations, but their mineral densities and plausibly their chemical compositions could differ. It is with this preliminary evidence that we postulate the pathophysiological cascade with the earliest mineralisation being proximal and distant from the traditional Randall plaque.

Discussion
All filters over time become less efficient with the accumulation of minerals. The kidney is a biofilter undergoing constant reclamation of solutes and excretion of waste over decades. Here, we identify evidence that mineralisation within the medullo-papillary complex occurs in the absence of overt stone disease, and that greater amounts of proximal intratubular mineralisation is accompanied with greater amounts of distal interstitial mineralisation. Although visualisation of Randall’s plaque is a first sign of urinary stone disease endoscopically, the mineralisation process within the medullo-papillary complex is already well advanced when these plaques are apparent. For the first-time calcium stone-former, the expression of stone disease may represent the accumulation of lifestyle choices over prior decades. In a way, it is too late. Stones represent the end-product of a long cascade of events.

However, an appreciation that the aetiology of Randall’s plaque is initiated more proximally within the medullo-papillary complex may focus the development of therapeutics that would help modulate or slow the mineralisation process. For instance, based on Poiseuille’s equation, viscosity of the tubular or blood flow might be modulated to allow improved flow and decelerate the process of mineral deposition. It also has been observed that not all who have Randall’s plaques form kidney stones, as 19% of the normal population have a grossly visible plaque [6,7], which is higher than reported kidney stone prevalence [30]. Forming kidney stones may therefore require a ‘two-hit’ process, with Randall’s plaque as a predisposed insult.

Additional studies are needed to confirm these observations, but challenges remain. Age-related processes over the long-term are difficult to study. A suitable animal model to study renal mineralisation is yet to be identified, as there is no established animal model of Randall’s plaque formation, and most animal models have uni-papillary systems as opposed to the multi-papillary system found in humans. Despite these challenges, recent advances in micro- and nano-resolution imaging techniques and modelling microfluidics may provide more insights into the biomineralisation process.

Conclusion
Biomechanical approaches to renal physiology and pathophysiology can add insight to the current paradigm on the initiation of nephrolithiasis. The urinary tract was designed to transport fluids, not solids. We now realise that once a calculus becomes dislodged from the papilla, it is too late. This detached, free-floating stone now poses the threat of growth, obstruction, and causing symptoms to its host. While mechanisms within the urinary milieu, such as supersaturation and nucleation, contribute to stone maturation within the collecting system, it is the pathological changes that occur at the tissue level within the parenchyma that may be fundamentally more important.
Acknowledgements

This research was supported by contracts National Institutes of Health (NIH) National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK) - 1P20DK100863 (Marshall L. Stoller); NIH National Institute of Dental and Craniofacial Research (NIDCR) - R01DE022032 (Sunita P. Ho); NIH/ National Center for Research Resources (NCRR) – S10RR026645 (Sunita P. Ho); NIH/National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK) - R21DK109912 (Multiple PI: Sunita P. Ho (contact); Marshall L. Stoller). The authors also thank Justin Klein for concept-driven illustrations [Medical Illustration (www.JustinKleinCMI.com)] in this manuscript.

Conflicts of Interest

The authors of this manuscript certify that they have no affiliations with or involvement in any organisation or entity with any financial interest in the subject matter or materials discussed in this manuscript.

References


Correspondence: Sunita P. Ho, Division of Biomaterials and Bioengineering, Department of Preventive and Restorative Dental Sciences, School of Dentistry, 513 Parnassus Avenue, Health Sciences West 813, University of California San Francisco, HSW813, San Francisco, California 94143, USA.

e-mail: sunita.ho@ucsf.edu