Nephrolithiasis: site of the initial solid phase

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Most cases of nephrolithiasis are associated with the relatively common metabolic abnormality of idiopathic hypercalciuria (1). These patients generally absorb an excess amount of dietary calcium leading to increased urine calcium excretion and supersaturation with respect to calcium oxalate and calcium phosphate; they subsequently form stones. Other patients with nephrolithiasis, who have had an intestinal bypass procedure, absorb oxalate in excess leading to increased urine oxalate excretion and supersaturation with respect to calcium oxalate; they also subsequently form stones. In these and other causes of nephrolithiasis, the site of the initial solid phase has long been the subject of debate. Over 65 years ago, A. Randall demonstrated that interstitial crystals located at, or adjacent to, the papillary tip, Randall’s plaques, were common in stone formers (2). He found that these crystals were composed not of calcium oxalate, the most common solid phase found in patients with nephrolithiasis, but of calcium phosphate (3). He believed that the calcium phosphate crystals formed in the papillary interstitium and then eroded into the urinary space, serving as a heterogeneous nucleation surface for calcium oxalate.

B. Finlayson later argued that, due to rapid flow of the renal ultrafiltrate through the tubule, there was insufficient time for formation of a lumen-obliterating solid phase (4), which also suggested that an intratubular site of stone formation was unlikely. However, other investigators found that calcium oxalate crystals adhered to cultured tubular cells (5), where they could either be endocytosed or remain on the cell surface, serving as a nidus for growth into larger, clinically significant, calculi.

Site of the initial solid phase

Where is the site of initial crystallization — the interstitium, the tubular lumen, or perhaps the renal calyx, where supersaturated fluid awaits excretion into the ureter? Knowing the site of initial crystallization would improve understanding of the pathogenesis of stone formation and allow investigators to propose and test more focused hypotheses. This would help them to devise effective therapy aimed at preventing recurrent nephrolithiasis, which affects approximately 50% of stone formers within five years of the initial stone (6). Yet until the elegant study by A.P. Evan et al. reported in this issue of the *JCI* (7), we did not have an answer to this rather elementary question. These investigators performed kidney biopsies on stone-forming patients to determine the anatomical site and composition of the initial solid phase. They sampled areas adjacent to Randall’s plaques in patients undergoing percutaneous nephrolithotomy. In hypercalciuric calcium oxalate stone formers, they found initial calcium phosphate (apatite) crystallization in the basement membrane of the thin limbs of the loop of Henle (Figure 1) with subsequent extension to the vasa recta, then to the interstitial tissue surrounding the terminal collecting ducts, and finally, in the most severe cases, to the papillae. Erosion of this solid phase into the urinary space, which is supersaturated with respect to calcium oxalate, may have promoted heterogeneous nucleation and formation of kidney stones. In patients with hyperoxaluria resulting from intestinal bypass, the initial crystals were again a calcium phosphate complex, but these arose within the tubule lumens of terminal collecting ducts (Figure 2). Contact of these crystals with urine, supersaturated with respect to calcium oxalate, may have promoted heterogeneous nucleation and formation of kidney stones. Nonstone formers, subjected to nephrectomy, had neither plaque nor crystals. Thus there are different sites of initial...
crystallization depending upon the metabolic abnormality leading to stone formation.

Potential mechanisms for stone formation

Why does the initial solid phase form in these distinct locations, and why are the initial crystals apparently only calcium phosphate? The basement membrane of the thin limb appears an unlikely site for initial crystallization in patients with idiopathic hypercalciuria. It is not the site of either vectorial calcium or phosphorus transport (8) and, since even the transtubular permeabilities of these ions are very low (8), it is difficult to link supersaturation with respect to calcium oxalate (CaOx), may have promoted heterogeneous nucleation and formation of calcium oxalate kidney stones (F).

Intraluminal crystal formation in the collecting duct appears a more likely site for initial crystallization in patients following intestinal bypass surgery. The collecting duct fluid can be hypertonic with elevated concentrations of calcium leading to supersaturation. Yet the urine from the patients in Evan's study was undersaturated with respect to calcium phosphate, indicating that, thermodynamically, a stone should not form. However, the lack of demonstrable supersaturation may be a function of the 24-hour urine collection; the maximal supersaturation, and thus the propensity for stone formation, is never detected. While a 24-hour urine collection is an important predictor of the likelihood of forming stones, it is not the sole predictor. It seems probable that supersaturation initiates crystal formation, but we still do not understand the relationship between the degree of urinary supersaturation and stone disease.

Future directions

Now that we know where the initial solid phase forms, what are the next questions? Investigators studying the kidney generally concentrate on the effects of transport on tubular fluid ion concentration; the current study will force us to look more carefully at the effects of basolateral membrane transport on interstitial ion concentrations. We know little about supersaturation in this critical region of the kidney, yet this is where the majority of stones originate. Unneeded calcium and oxalate must be excreted in a minimal amount of urine to rid the body of these potential crystallizing the interstitium. Vectorial proton transport into the collecting duct would alkalize the interstitium. The pH of the vasa recta would also increase following gastric proton secretion, the so-called alkaline tide, resulting in less bicarbonate removal from the medullary interstitium. The increased pH would decrease the solubility of calcium phosphate complexes. Perhaps an extracellular matrix protein, specific to the papillary interstitium, could provide a site promoting heterogeneous nucleation (12), which occurs with a lower degree of supersaturation than homogeneous nucleation. Future studies will be necessary to test these hypotheses.

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in the bypass patients there was not areas of dense crystal deposition, while evidence for crystal-induced cell injury in pathic hypercalciuria, there was evidence for a solid phase (15). In patients with idiopathic hypercalciuria, there appears to regulate growth of the solid phase rarely form because we produce inhibitors to initial nucleation and spontaneous aggregation (12). In rats, stone formation occurs when the magnitude of the supersaturation overcomes this potent inhibition (14). The current study should point us in the direction of investigating the relationship between supersaturation and inhibitor proteins not only in the urine but in the interstitium as well.

Crystals may stimulate production of proteins, such as osteopontin, which appear to regulate growth of the solid phase (15). In patients with idiopathic hypercalciuria, there was evidence for crystal-induced cell injury in areas of dense crystal deposition, while in the bypass patients there was not only cell injury but also cell death (7). Were levels of osteopontin increased in the medullary interstitium in either of these types of stone formers?

**An animal model of stone formation**

The genetic hypercalciuric stone-forming rat exhibits metabolic abnormalities similar to patients with idiopathic hypercalciuria in that these rats absorb excessive amounts of intestinal calcium, they fail to adequately reabsorb filtered calcium, and their bone resorption is uniquely sensitive to 1,25(OH)2D3. All of these characteristics are apparently due to an increase in the number of receptors for vitamin D (16, 17). The genetic hypercalciuric stone-forming rats spontaneously form calcium phosphate stones (18) similar to those found in this study unless their diet is augmented with an oxalate precursor (19). Both hypercalciuric rats and humans appear to be predisposed to initially form calcium phosphate stones and not the commonly observed calcium oxalate stones. In both rats (20) and humans (21) the upper limit of metastability, that level of supersaturation at which a solid phase forms, increases with increasing calcium oxalate, but not calcium phosphate, supersaturation. Thus rats and humans appear protected against calcium oxalate stone formation unless a nucleation site, such as the more easily formed calcium phosphate crystal, is present.

This study highlights the role of physician scientists working with basic scientists in medical research to jointly address important problems using sophisticated clinical and laboratory techniques and then applying these results to refine hypotheses for further testing. Agile movement between the bedside and the bench, as exemplified in this study, will provide insight into, and ultimately prevention of, disorders such as nephrolithiasis.

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**Figure 2**

Stone formation in patients following intestinal bypass. Initial calcium phosphate (apatite) crystallization was found within the tubule lumens of the terminal collecting ducts (A and B). Contact with the urine, supersaturated with respect to calcium oxalate, may have promoted heterogeneous nucleation and formation of calcium oxalate kidney stones (C).


