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Randall’s plaque as the origin of calcium oxalate kidney stones

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Abstract

Eight decades ago, Alexander Randall identified calcium phosphate deposits at the tip of renal papillae as the origin of renal calculi. The awareness that these “Randall’s plaque” promote renal stone formation has been amplified during the past years by the development of endoscopic procedures allowing the in situ visualization of these plaques. Recent studies based upon kidney biopsies evidenced that apatite deposits at the origin of these plaque originate from the basement membranes of thin loops of Henle and then spread in the surrounding interstitium. In addition, scanning electron microscopy examination of calcium oxalate stones developed on Randall’s plaque evidenced that plaque may also be made of tubules obstructed by calcium phosphate plugs. Hypercalciuria has been associated to Randall’s plaque formation. However, several additional mechanisms may be involved resulting in increased tissular calcium phosphate supersaturation and the role of macromolecules in plaque formation remains elusive. At last, apatite crystals are the main mineral phase identified in plaques, but other calcium phosphates and various chemical species such as purines have been evidenced, revealing thereby that several mechanisms may be responsible for plaque formation.

Keywords Randall’s plaque · Kidney stones · Calcium · Oxalate · Apatite

Introduction

Calcium oxalate (CaOx) stones are now identified as the main type of urinary calculi in most countries throughout the world. They account for at least 70 % of all stones developed in the upper urinary tract and even more in male patients [1]. In some parts of the world, CaOx was found as the main component of about 90 % of all kidney stones [2]. Such a progression in CaOx stones accounts for the increase in the prevalence of kidney stone disease frequently reported during the past decades from countries where longitudinal epidemiological surveys are available as in the United States [3], in Europe [4, 5] or in Japan [6]. Among the risk factors involved in stone formation, low diuresis and imbalance in diet are well documented [7–9]. Although a lot of inherited or acquired diseases may induce kidney stone formation, all together account for less than 10–15 % of calcium stones in the general population. Thus, most of calcium stones result from biological disorders in urine such as hypercalciuria, hyperoxaluria and/or hypocitraturia without any “identified” pathology. Moreover, these disorders are inconstantly found when urine biochemistry is performed, suggesting the anomalies are intermittent or transient, or simply related to low diuresis, which results in an increased concentration of poorly soluble salts in urine. Thus, the so-called idiopathic nephrolithiasis is the bulk of calcium stones. However, during the last decade, a series of investigations by the group of Evan, Lingeman and Coe, highlighted the role of calcium phosphate deposits in the papilla tip as a starting point of CaOx stone formation [10–12]. Such deposits, termed Randall’s plaque because they were first described by Alexander Randall in the 1930s, initiate in the basement membrane of the long Henle’s loops and spread out in the interstitium around the Bellini’s ducts close to the papillary epithelium. When the epithelium is disrupted, the calcium phosphate deposits are in contact with the supersaturated urine issued from the vicinal nephrons, thus favouring CaOx crystalization and trapping of the crystals at the surface of the plaque. Thereafter, the stone growth is driven by urine supersaturation.

Original description of papillary calcified plaques by Randall

More than 75 years ago, Alexander Randall reported a study on “The origin and growth of renal calculi”, which appeared first in the New England Journal of Medicine [13] and the following year in the Annals of Surgery [14]. In this seminal paper, he described a hitherto unrecognized papillary lesion consisting in calcium phosphate deposits in the intertubular spaces of the renal papilla, observed in 19.6 % of 1,154 pairs of kidneys examined at autopsy. He made three observations:

- first, this papillary lesion which appeared as a cream coloured area near the tip of the papilla was not on its surface, but was subepithelial in location;
- second, the chemical composition of papillary plaques was shown to be made of calcium phosphate and carbonate;
- third, small calculi were found to be attached to plaque at the tip of papillae in 65 kidneys (2.8 %), their composition differing from that of the plaque. The stones were made of calcium oxalate;
- fourth, CaOx stones collected in routine practice often exhibited a portion which was smooth and somewhat depressed, resembling a facet corresponding with the morphology of the papilla.

Examining by light microscopy some calculi actually attached on a papillary plaque, Randall found that the stone was continuous with the interstitial plaque and that the epithelial covering of papilla was lost. Such aspect of calcium phosphate deposits in the interstitium was termed papillary lesion type 1 by Randall who proposed that papillary calcium deposits could act as a nidus for urinary salt accretion if epithelial covering of the papilla is lost. Randall also found in some cases Bellini’s ducts filled by calcium phosphate plugs (termed papillary lesion type 2). He suggested the phosphate plugs may be another way for CaOx stone formation.

Fig. 1 Scanning electron microscopy photographs of calcium oxalate calculi initiated from a Randall’s plaque. a The plaque is made of a mixing of tubules with calcified walls (white arrow) and of tubules obstructed by calcium phosphate plugs (black arrow). b Randall’s plaque at the surface of another calcium oxalate stone with calcified tubules and a calcium phosphate plug partly obstructing the lumen of a collecting duct.
Scanning electron microscopy examination of CaOx stones developed from a Randall’s plaque provides evidence that plaque is made of a mixing of tubules with calcified walls (white arrow, Fig. 1a) and of tubules obstructed by calcium phosphate plugs (black arrow, Fig. 1a or white arrow, Fig. 1b). Please note in Fig. 1b, the calcium phosphate cluster within the tubular lumen (white arrow), while other tubules are empty with calcified walls. In other cases, much less frequent, the nidus of the stone appears only as a tubular plug (Fig. 2, black arrow). At the time Randall performed his studies, he could not identify the exact location and nature of calcium deposits within the papillary structure. Nor, due to the forensic nature of his study material, could he evaluate urinary solute excretion in the affected individuals.

Epidemiological considerations

Other authors confirmed the findings of Randall in autopsy studies during the same period: for example, Rosenow found plaques in 22.2 % of 239 kidneys, whereas Anderson observed plaques in 12 % of kidneys in a series of 1,500 necropsies [15, 16]. In South Africa, Vermooten [17] examined 1,060 pairs of kidneys. He found Randall’s plaques in 17.2 % of Caucasians and only 4.3 % of Bantus, a finding in keeping with his observation of the lower incidence of calcium nephrolithiasis in the local population of African descent than Caucasian descent. Papillary calcifications were seen in the subepithelial interstitial tissue together with collagen fibres, and were not found in the lumen of collecting ducts [18]. Forty years after the first description of papillary calcified deposits acting as a nidus for calcium stone formation, Luis Cifuentes-Delatte, a Spanish urologist enthusiastic for research on stone formation observed with his colleagues that 142 out of 500 consecutive calculi exhibited a concavity characteristic of a papillary origin [19]. In 61 (43 %) of these umbilicated calculi, plaques were typically made of apatite, some of them showing calcified collecting ducts [20].

Fig. 2 Scanning electron microscopy photograph of a tubular plug (black arrow) at the origin of a calcium oxalate monohydrate stone

Fig. 3 Whewellite stone initiated from a carbapatite Randall’s plaque (white arrow)
In our experience, based upon 45,774 calculi referred to the Necker hospital stone laboratory over the past three decades, we found by morphologic examination coupled with FTIR analysis that 8,916 (19.5 %) calculi exhibited a depressed surface (umbilication) suggestive of a papillary attachment [21]. Most of these stones (92.5 %) were made of CaOx monohydrate (whewellite) either pure (Fig. 3) or admixed with CaOx dihydrate (weddellite) as observed in Fig. 4. As previously reported, the composition of Randall’s plaque is carbapatite in 90 % of cases, but other crystalline phases may be found as main component of Randall’s plaque such as amorphous calcium phosphate, brushite or whitlockite, or trioxypurines like sodium urate [19, 21]. However, detailed endoscopic examination of renal cavities during percutaneous nephrolithotomy (PNL) or ureteroscopic procedures suggests a much higher prevalence of Randall’s plaques than that deduced from stone examination. An explanation could be that modern urological treatment to remove stones often use fragmentation techniques such as shockwaves, ultrasounds or laser. As a consequence, stone fragments collected for analysis may have lost both the part of Randall’s plaque fixed to the stone and the typical depressed face. In 1997, using endoscopic procedures, Low and Stoller reported the presence of Randall’s plaques on one or more papillae in 74 % of 57 stone formers, the prevalence being 88 % for CaOx stones [22]. In 2006, Matlaga et al. [23] found that 91 % of papillae in 23 patients with idiopathic CaOx nephrolithiasis harboured plaques and half of papillae had attached calculi. Digital imaging provided undisputable evidence of stone attachment to plaque. Indeed, epidemiological differences in the occurrence of Randall’s plaques may be observed according to the investigated population. Recent reports from European countries suggest that Randall’s plaque could be less frequent in Europe than in the United States. In Italy, Ruggera et al. reported the occurrence of Randall’s plaque in 44.4 % of 27 stone formers who underwent ureterorenoscopy or PNL and renal papillae biopsy [24]. In France, Olivier Traxer found Randall’s plaques in 57 % of 287 stone formers who underwent flexible ureterorenoscopy. They were observed on all papillae in 54 % of these patients. Such plaques were visible in only 27 % of 173 non-stone forming patients [25, 26].
Of note, in France, stones harbouring an umbilication were found to be three times more frequent in the recent years than at the beginning of the 1980s and patients were younger and younger [27]. Children may form CaOx stones from Randall’s plaque. As shown in Fig. 5, when considering only unfragmented stones spontaneously passed or removed by urological procedures without fragmentation, the part of stones developed from a Randall’s plaque is very high and dramatically increasing with age from childhood up to the age class 20–30 years, then slowly decreasing thereafter. In our experience, 39 % of all spontaneously passed stones had a typical umbilication on their surface and 5,728 among 8,662 (66.1 %) of spontaneously passed CaOx monohydrate calculi were umbilicated and were developed from a Randall’s plaque.

**Randall’s plaques begin in basement membranes of thin Henle’s loops**

In an elegant study, Evan and co-workers recently provided substantial advances in the pathogenesis and structure of Randall’s plaques in patients with CaOx stones [10]. They took superficial biopsy specimens from papillae in the upper, middle and lower poles of a kidney during PNL performed for stone removal in 15 patients with recurrent, idiopathic CaOx nephrolithiasis, all of whom were hypercalciuric, and in 4 patients with CaOx stones and hyperoxaluria due to jejuno-ileal bypass for obesity.

By light microscopy, they confirmed Randall’s plaques to be in subepithelial position at the papillary tip, surrounding the openings of the ducts of Bellini. Calcium-containing deposits were found surrounding primarily the thin loops of Henle. Electron microscopy revealed these deposits to be located within the basement membrane of Henle’s loops and around adjacent vasa recta, but not within the cytoplasm of the cells. Infrared microspectroscopy of deposits identified spectral bands characteristics of hydroxyapatite, associated with calcium carbonate in about 20 % of cases. In summary, papillary plaques are interstitial and composed of apatite, originate in the basement membrane of the thin loops of Henle and subsequently spread into the surrounding interstitium, in the vasa recta and in the basement membranes of collecting ducts, without affecting collecting duct cells themselves. As said by the authors, “the overall impression is that the basement membrane of thin limbs is indeed the origin of plaque”.

In contrast, none of the four patients with enteric hyperoxaluria due to jejuno-ileal bypass exhibited such papillary plaques. Instead, they exhibited intratubular hydroxyapatite crystals in the lumen of some collecting ducts, attached to their apical surface. Thus, the site of initial crystallization differed according to the underlying metabolic abnormality. Of note, four non-stone-formers subjected to nephrectomy, taken as controls, had neither papillary plaques nor tubular crystals [10]. Looking for risk factors associated with Randall’s plaque formation,
Kuo et al. found that low urine output, high calcium content and low urine pH were significantly more frequent in calcium stone formers than without Randall’s plaque [28]. Moreover, Kim and coworkers reported that stone episodes were proportional to the coverage of papilla by Randall’s plaques [29].

**Role of hypercalciuria in the pathogenesis of Randall’s plaques**

Evan and co-workers speculate that hypercalciuria should be the main explanation at the origin of Randall’s plaques [10]. Before examining their arguments, one may remember that an epidemiologic case/control study identified high urinary calcium concentration to be the major risk factor for idiopathic calcium nephrolithiasis [30], whereas oxalate, uric acid and citrate concentrations did not differ between incident stone formers and contemporary controls. In the study of Evan et al. [10], the mean calcium excretion was 312 mg/day (7.8 mmol/day) in the 15 CaOx stone Fig. 5 Occurrence (%) of Randall’s plaque at the origin of spontaneously passed calcium oxalate stones according to the patient’s age formers, compared to 113 mg/day (2.8 mmol/day, p < 0.01) in the 4 non-lithiasic controls.

According to the authors and to the editorial comments of their work by Bushinsky [31], the sequence of events could be as follows. Hypercalciuria is associated with intestinal calcium absorption, which results in postprandial peaks in serum calcium, reflected in increased calcium concentration in glomerular filtrate and in the fluid presented to the thin limbs, then to the thick ascending limb. High calcium reabsorption at this site enriches the outer medulla with calcium, which is washed into the deep medulla by the descending vasa recta. The origin of calcium phosphate deposits in the interstitium may be the consequence of several mechanisms. As reported by Asplin and co-workers [32], the narrow part of the long loops of Henle is often supersaturated for calcium phosphate, which may precipitate. Despite the short transit time in that part of the nephron (20–30 s), where most particles remain free in the tubular lumen, some of them may be attached to the apical cell membrane, especially if urinary macromolecules and other inhibitors such as citrate are in low concentration or if an oxidative stress was able to damage the cell surface. Then, an endocytosis process may be initiated to eliminate them by intracellular dissolution [33]. Calcium and phosphate ions resulting from the apatite dissolution are transferred in the interstitium and may locally induce high CaP supersaturation favouring new crystallization of apatite in the basement membrane of thin limbs, which is unusually thick and is composed of collagen with abundant mucopolysaccharides. Apatite crystals, once initiated, are able to accumulate and extend in the interstitium. At the microscopic scale, Evan et al. have shown that apatite crystals are composed of successive layers of mineral and macromolecules such as osteopontin [11]. Extensive accumulation of interstitial calcium deposits up to the papillary epithelium may result in a disruption of the epithelium by a yet poorly understood mechanism, thus making the apatite crystals exposed to the supersaturated urine. In the first following steps, new layers of calcium phosphate and macromolecules may cover the plaque, after what CaOx crystals formed in the surrounding supersaturated urine may be attached on the plaque, thus initiating stone formation [11, 26].

As underlined by Tiselius [34], calcium phosphate precipitation requires high CaP supersaturation, which is favoured by high pH value, high concentration of calcium and/or phosphate ions. Indeed, the physicochemical environment in the interstitium of the inner medulla favours calcium phosphate supersaturation around the tip of Henle’s loops, especially in the presence of hypercalciuria and low urine output. In case of an excessive acid load, as observed in patients who eat large amounts of animal proteins, an increased excretion of H+ ions by the collecting duct cells results in a low urine pH and an increased production of bicarbonate ions transferred to the interstitium. High protein intake is also responsible for a high excretion of phosphate ions, which could be another favouring factor for explaining calcium phosphate supersaturation in tubular lumen and the papillary interstitium.

As a result of the high acid load (the same is observed in the case of antidiuresis), the pH increases in the inner medulla, inducing supersaturation of calcium phosphate that may precipitate. Moreover, in hypercalciuric states, which appear to be more frequent in patients with Randall’s plaques, a default of calcium reabsorption in the proximal part of the tubule results in an increased delivery of calcium in the distal part of the nephron, which may contribute to the supersaturation of calcium phosphate and/or CaOx [35]. In the case of low diuresis, the hyaluronic acid content in the inner medulla decreases [36] thus reducing its availability to act as an inhibitor of crystallization and results in increased concentration of solutes in the interstitium.
While endocytosis of CaP particles or secondary precipitation of CaP in the basement membrane of the cells may result from local high supersaturation, another phenomenon could explain why CaOx crystals may be secondary deposited on the CaP Randall’s plaque after the papillary epithelium has been lost by a yet unexplained mechanism. Indeed, it is well known that CaP particles are the most frequent mineral phases found in urine [37]. In the case of low diuresis or high acid load, the secretion of H+ in the distal part of the nephron results in a rapid decrease of the pH inducing dissolution of CaP particles. As a consequence, an increased calcium concentration may increase enough the CaOx supersaturation to induce nucleation of CaOx crystals that are excreted in the caliceal cavities [3, 38] and then may be fixed on the CaP particles at the surface of the papilla [26]. Then, the stone growth depends on the level of CaOx supersaturation, which is linked to the diuresis, and the concentration of both calcium and oxalate, and also of inhibitors such as citrate. When the stone is too large or too heavy for the mechanical resistance of the Randall’s plaque that plaque may be disrupted and the stone free in the caliceal space. Thereafter, it may be rapidly spontaneously passed (the most frequent situation) or retained in the calyx. In such case, new CaOx crystals may progressively cover the umbilication and the calcium phosphate plaque, which disappears at the surface of the stone. When removed or spontaneously expelled, the stone does not exhibit any Randall’s plaque which is found in the core of the stone as suggested by Miller et al. [39] and illustrated in Fig. 6.

**Randall’s plaque and stone composition**

First studies by Evan and co-workers suggest that apatite crystals are the mineral phase for interstitial plaque and that amorphous calcium phosphate (ACP) deposits may cover the plaque secondly after it outcrops at the surface of the epithelium [11]. However, as suggested by Tiselius [34] ACP formation is favoured by high CaP supersaturation, which may occur not only in the urine, but also in the interstitium. In fact, other works suggest that ACP may be more abundant in the interstitial part of the Randall’s plaque than at its surface [40]. As previously reported, Randall’s plaque is made of carbapatite in most cases, but other calcium phosphates and other chemical species, such as sodium urate may be identified within Randall’s plaque [21]. As shown in Table 1, about 15 different crystalline species were found among 11,016 calculi exhibiting a papillary umbilication. Carbapatite was identified in 97.6 % of cases, either pure or mixed with other calcium phosphates or other compounds such as purines. However, in some cases, carbapatite was not detected and then, another mechanism
than Randall’s plaque may be involved in stone formation. We cannot exclude that some of these compounds such as uric acid were only deposited at the surface of the papilla because of a local tissue lesion and then acted as a factor for heterogeneous nucleation of calcium oxalate. Of note, around the Randall’s plaque the first layers of the stone are made in most cases by CaOx monohydrate [21, 38].

### Table 1: Occurrence of crystalline species found as Randall’s plaque at the stone surface

<table>
<thead>
<tr>
<th>Components of papillary deposits</th>
<th>Main</th>
<th>Minor</th>
<th>Total occurrence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium phosphates</td>
<td>10716 (97.3)</td>
<td>–</td>
<td>10894 (98.9)</td>
</tr>
<tr>
<td>Carbobapatite</td>
<td>10404 (94.4)</td>
<td>351 (3.2)</td>
<td>10755 (97.6)</td>
</tr>
<tr>
<td>Amorphous carbonated calcium phosphate</td>
<td>264 (2.4)</td>
<td>629 (5.7)</td>
<td>893 (8.1)</td>
</tr>
<tr>
<td>Whitlockite</td>
<td>30 (0.3)</td>
<td>87 (0.8)</td>
<td>117 (1.0)</td>
</tr>
<tr>
<td>Octacalciun phosphate</td>
<td>3 (0.03)</td>
<td>13 (0.12)</td>
<td>16 (0.15)</td>
</tr>
<tr>
<td>Brushite</td>
<td>15 (0.14)</td>
<td>23 (0.2)</td>
<td>38 (0.34)</td>
</tr>
<tr>
<td>Other species</td>
<td>300 (2.7)</td>
<td>–</td>
<td>503 (4.6)</td>
</tr>
<tr>
<td>Struvite</td>
<td>3 (0.03)</td>
<td>9 (0.08)</td>
<td>12 (0.1)</td>
</tr>
<tr>
<td>Prodibrite</td>
<td>2 (0.02)</td>
<td>0</td>
<td>2 (0.02)</td>
</tr>
<tr>
<td>Calcite</td>
<td>9 (0.08)</td>
<td>12 (0.1)</td>
<td>21 (0.2)</td>
</tr>
<tr>
<td>Sodium hydrogen urate</td>
<td>232 (2.1)</td>
<td>138 (1.25)</td>
<td>370 (3.4)</td>
</tr>
<tr>
<td>Uric acid</td>
<td>43 (0.4)</td>
<td>24 (0.2)</td>
<td>67 (0.6)</td>
</tr>
<tr>
<td>Ammonium urate</td>
<td>4 (0.04)</td>
<td>3 (0.03)</td>
<td>7 (0.06)</td>
</tr>
<tr>
<td>Potassium urate</td>
<td>4 (0.04)</td>
<td>0</td>
<td>4 (0.04)</td>
</tr>
<tr>
<td>Opaline silica</td>
<td>2 (0.02)</td>
<td>1 (0.01)</td>
<td>3 (0.03)</td>
</tr>
<tr>
<td>Porphyrines</td>
<td>1 (0.01)</td>
<td>16 (0.15)</td>
<td>17 (0.15)</td>
</tr>
<tr>
<td><strong>Total number of samples</strong></td>
<td><strong>11016 (100)</strong></td>
<td><strong>1291 (11.7)</strong></td>
<td><strong>–</strong></td>
</tr>
</tbody>
</table>

### Conclusion

Idiopathic CaOx stones, the occurrence of which has been frequently reported as increasing during the past half century, mainly from a Randall’s plaque made of carbapatite. In parallel, the progress in endoscopic material and optics, making easy the visual inspection of kidney papillae provides evidence that Randall’s plaques are become very common in stone formers as in the general population. Such changes are in line with in-depth modifications in dietary habits in industrialized countries over the past decades. A better knowledge regarding the mechanisms involved in the formation of calcified deposits within the inner medulla of the kidney is required in order to provide efficient advices able to reduce the occurrence of Randall’s plaque and stone formation.

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