Pathogenesis of nephrolithiasis
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Introduction
Urinary stone disease is a common clinical problem as the lifetime risk of having a stone is 10% in the population [1]. Furthermore the incidence of nephrolithiasis is increasing with dramatic increases noted in females and children [2,3]. In addition there are changes in the frequency and composition of urinary calculi. For example the incidence of staghorn calculi is decreasing in the European countries and the proportion of struvite calculi among staghorn calculi is decreasing both in Europe and USA [4]. These trends are related to changes in lifestyle, dietary patterns and improving medical services [5]. Renal stones are commoner in men but the ratio varies in different parts of the world. In the West it is 2.7:1 while in Sri Lanka it is 3.6:1 [6,7]. Although staghorn calculi are commoner in women, in Sri Lanka there is a male preponderance with a ratio of 4:1 [8]. These differences suggest that patterns of stones differ in different geographical localities prompting scientists to believe in different aetio-pathogenesis as well. However the composition and distribution of renal stones in Sri Lanka do not differ in different parts of the country studied [9].

Fortunately profound advances in endoscopic design, durability and accessories have revolutionised the minimally invasive therapy of nephrolithiasias. Vast improvements in imaging techniques have been made during the recent past. These advances have made treatment of nephrolithiasis, patient friendly with minimal morbidity and short hospital stay. However treatment of nephrolithiasis does not end with successful removal of the offending stone. There is a high tendency for these stones to recur and within 5 years 35% to 50% of patients develop another stone [10]. Therefore prevention of a future recurrence of stones is an important necessity to avoid long term morbidity and improve the health of these people. In order to formulate preventive strategies it is imperative to understand the aetiopathology of nephrolithiasis.

Pathogenesis
There is no consensus on the aetiopathology of nephrolithiasis. This alone is sufficient indirect evidence for its multifactorial nature. However it is clear that pathogenesis of renal stones constitutes of two major steps. Firstly the immediate changes that occur at the site of stone formation. Secondly the general factors which induce and promote these changes at the site.

Level I (at the tissue level)
Supersaturation – Supersaturation occurs when the concentration of a stone-forming salt exceeds its solubility in a solution. This leads to formation of foci of its solid phase. Urine is a medium which is more complex than pure solution [11]. Therefore the required critical concentration for supersaturation may rise up to eightfold and new solids will not form during this period. This concentration range is called the metastable zone. Though new stones may not form in this range, existing stones may aggregate and grow in the metastable zone. This has clinical importance as any tiny fragments of stones left after treatment may lead to bigger stone formation easily though new stone formation would be unlikely if the patient remains in the metastable zone. Exceeding supersaturation is the cause of uric acid and cystine calculi [12].

Plaque theory - In 1930s Randall described a pre-calculus lesion (plaque) in the renal papilla and proposed that a sub-epithelial calcification of renal papilla becomes the nidus of calcium oxalate monohydrate papillary calculi, as a consequence of the disruption of the papillary epithelial layer by the hydroxyapatite (calcium phosphate) plaque [13]. Such plaques are present in all idiopathic calcium oxalate stone formers but not in healthy controls [14]. Recently, it was found that in patients susceptible to
the development of calcium oxalate stones, the plaque is initiated in thin-loop of Henle basement membranes, basement membranes of collecting tubules, and the vasa recta [15]. However the initial trigger for this crystallization is not clear yet. A multifactorial process seems to be likely. An increased urinary calcium excretion and an acidic urinary pH have been incriminated as contributory factors that initiate the crystallization process by promoting apatite (calcium phosphate) depletion [16].

Recently there is another interesting hypothesis suggested as the mechanism for formation of plaques. Accordingly vasa recta play a major role in the process which is aptly termed the vascular theory [17]. The hypoxic and hyperosmolar environment in the papillary tip makes descending and ascending vasa recta vulnerable for changes which lead to atherosclerotic–like lesions and calcifications in its wall. These calcifications erode into papillary interstitium which enlarges supported by cellular promoters. Proponents of this vascular theory cite the higher incidence of stones in individuals with metabolic syndrome, diabetes mellitus and coronary artery disease as supporting evidence [18,19]. Plaque formation is the most plausible theory so far to explain the formation of idiopathic calcium oxalate stones.

Level II (general factors)
Although these changes described above at the site of stone formation have been well described their importance to clinicians who manage patients with renal stones in real life is minimal. It is the general factors which induce and modify these pathological processes that are important for practicing clinicians. Interplay of three sets of general factors lead to formation of stones.
(i) Genetic/metabolic
(ii) Habits
(iii) Intrinsic factors related to the urinary tract
Genetic and metabolic factors that contribute to nephrolithiasis include hypercalciuria, hypercalcaemia, hyperoxaluria, hyperuricosuria, hypocitraturia, hypomagnesaemia and renal tubular acidosis [20]. Some of these conditions when severe can cause stones on their own by exceeding supersaturation. But the majority leads to stone formation in the presence of other factors too. Genetic and metabolic factors can be divided into two broad categories. Diseases with generalised system involvement like hyperparathyroidism, sarcoidosis and gout constitute the first group. The second group include diseases with urinary tract manifestations only e.g. hypercalciuria, hyperoxaluria and hypocitraturia. Habits of individuals make a significant contribution to stone formation especially when the individuals have another predisposing factor categorised under genetic/metabolic or urinary tract factors. An average adult should produce about 2 litres of urine a day [20]. To achieve this, a person living in a tropical country like Sri Lanka should drink about 3 litres of water a day. Those who work outdoors and engage in certain occupations are at risk of developing nephrolithiasis, e.g. labourers, drivers, masons and carpenters [8]. Such people should drink adequate amounts of water to avoid stone formation. Out of all possible measures of prevention this is the only intervention that has been proven consistently to be of value in prevention of nephrolithiasis [21]. It is cheap and realistic too when compared to other dietary interventions since these are life time changes. The quality of water in causation of stones is a contentious issue. Hard water when boiled is free of excess calcium salts and theoretically safe to be consumed [22]. Some have queried whether high fluoride content in water potentiates renal calculi formation [1,23]. Although it is possible theoretically there is no convincing evidence to prove it.

A monotonous diet has been incriminated with formation of renal stones [1]. Animal proteins, by providing a high acid load may promote stones. An acidic urinary pH leads to an increased bicarbonate resorption into the renal medulla and a consequent increase in interstitial pH that causes calcium phosphate depletion [16]. Though many advise restriction of calcium rich food, it may be useful only in a small number of patients with absorptive hypercalciuria [24]. On the other hand a low calcium diet is proven to cause renal stones and is harmful to bone and dental health [25]. Effect of calcium supplementation on known stone formers appears to be counter-productive [26]. If calcium supplements are essential due to other medical reasons, it will be a wise move to assess the 24 hour urinary calcium level from time to time before deciding to continue supplementation. If calcium supplements are to be taken they should be taken after meals to minimise hypercalciuria. Those with intestinal hyperoxaluria could avoid oxalate rich food like cocoa based products, spinach, asparagus, tomatoes, rhubarb and parsley. Restriction of red meat and nuts is useful in patients with raised uric acid levels in blood and urine.

In the absence of sophisticated laboratory facilities to identify individual metabolic derangements, a sensible
approach to adjust the diet would be more practical and have a better compliance in preventing urolithiasis in susceptible individuals. A diet consisting of fruits, vegetables, low salt, low protein and a balanced intake of calcium, fats and carbohydrate should be the recommended diet for stone formers. Citrate supplementation with lemon or orange juice would be useful [27]. Citrate is one of the inhibitors of urolithiasis. The others are magnesium, pyrophosphate, protease inhibitors and glycosaminoglycan. Indigenous herbal products may help in expulsion of small stones or may prevent stone formation by contributing to the fluid intake if taken regularly [28,29].

The possible urinary tract factors that cause renal stone formation include infections with urease-producing organisms leading to alkaline urine, urinary stasis due to obstruction and foreign bodies lodged in the urinary tract. Alkaline urine is known to produce struvite stones. This was believed to be the main cause of staghorn calculi traditionally. However it has been shown now that most staghorn calculi are not struvite stones and are not associated with urease-producing organisms [30]. Stasis due to obstruction (e.g. congenital pelvi-ureteric junction obstruction) is well known as an underlying cause of nephrolithiasis though every patient with obstruction does not produce stones even when longstanding. Foreign bodies like forgotten ureteric stents and shrapnel lodged inside the kidney can be the focus of stone formation.

Preventive measures
Measures to prevent stone recurrence can be delivered at three levels (Panel 1).
The first constitutes general advice about dietary and drinking habits. Such advice is useful to all patients with stone disease. A diet consisting of fruits, vegetables, low salt, low protein and a balanced intake of calcium, fats and carbohydrate is recommended for stone formers. They should drink 2.5-3 litres of water a day. The second level includes specific dietary recommendations based on comprehensive analytical tests to identify any underlying metabolic derangement and composition of removed stones. The third level is pharmacological agents. Pharmacological agents like thiazides, allopurinol and potassium citrate have been shown to reduce the occurrence of stones in certain subgroups of patients (Panel 2). Adverse effects and poor compliance are

Panel 2 – Recommended pharmacological treatments for aggressive, recurrent stone formers

<table>
<thead>
<tr>
<th>Urine abnormality</th>
<th>Therapeutic agent</th>
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<tbody>
<tr>
<td>High excretion of calcium</td>
<td>Thiazide, Orthophosphate</td>
</tr>
<tr>
<td>High excretion of oxalate</td>
<td>Dietary restriction of oxalate, potassium citrate</td>
</tr>
<tr>
<td>Low excretion of citrate</td>
<td>Potassium citrate</td>
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<tr>
<td>Low excretion of magnesium</td>
<td>Thiazide+Mg</td>
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<tr>
<td>Renal tubular acidosis</td>
<td>Potassium citrate</td>
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<tr>
<td>Low urinary pH</td>
<td>Potassium citrate</td>
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<tr>
<td>High excretion of urate</td>
<td>Allopurinol</td>
</tr>
<tr>
<td>Brushite stone</td>
<td>Thiazide+Mg</td>
</tr>
<tr>
<td>Uric acid stone</td>
<td>Allopurinol, Potassium citrate</td>
</tr>
<tr>
<td>Cystine stone</td>
<td>Potassium citrate, Thiopronin, Captopril</td>
</tr>
<tr>
<td>Infection stone</td>
<td>Ammonium chloride, Antibiotics</td>
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obstacles for these drugs as these should be taken over very long periods to prevent a single stone event. In addition sophisticated laboratory facilities with dedicated staff are required for detection and diagnosis of underlying metabolic derangements in order to identify the subgroups. Hence second and third level preventive measures are generally reserved for aggressive stone formers who develop recurrent stones that grow fast despite above mentioned changes in habits and life style.

Too many dietary and life style restrictions during the entire life purely to prevent stones may not be practical and realistic in most stone formers. One may rather identify small stones early and treat with minimally invasive procedures like shock wave lithotripsy. Hence an annual ultrasonography of the urinary tract is a sensible way of following up all stone formers. It would also be an opportunity to remind them regarding the necessity to drink adequate amounts of liquids and to have a balanced diet while maintaining a healthy BMI.

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