Strategies for preventing calcium oxalate stones

About 10% of people will experience nephrolithiasis in their lifetime, and about 70% of those will have recurrences. About 80% of stones are calcium based, and about 80% of those are calcium oxalate stones. We discuss here briefly the evidence for the prevention of calcium oxalate stones through dietary and pharmacologic measures.

Urinary risk factors

Both genetic and environmental factors contribute to stone formation, but the genes responsible for the heritable aspect of stone formation have not been delineated definitively. Mainstays of calcium stone prevention involve manipulation of urine chemistries (urine sodium, citrate, oxalate, uric acid and calcium levels as well as urine specific gravity). The urinary risk factors most often implicated in stone formation are listed in Table 1. Hypercalciuria is the most common trait associated with calcium stones, yet its cause in most patients remains unclear. It is still often referred to as “idiopathic hypercalciuria.” The efficacy of classifying hypercalciuria on the basis of cause remains controversial and of unproven value in clinical management.

Dietary modification

Fluid intake

The cornerstone of management is to increase urine volume. The effect appears to be linear, with a point of diminishing return reached at urine volumes of more than 2.5 L per day. We prescribe an intake of 2.5–3 L of fluids per day.

There are data to suggest that the type of fluid ingested matters. Epidemiologic studies show that drinking coffee and beer decreases the risk of stones. Drinking grapefruit juice consistently increases stone risk for unclear reasons, whereas lemon juice, higher in citrate content, might have beneficial effects on urinary chemistry but its ingestion has not been shown to prevent stones.

Calcium

Studies of dietary calcium’s effect on stone recurrence rates have led to major changes in nonpharmacologic manipulation. Epidemiologic evidence shows an inverse relation between dietary calcium intake and recurrence rates. This is probably best explained by calcium’s inhibition of intestinal oxalate absorption. A randomized controlled trial (RCT) assigned men with hypercalciuria to either a diet low in calcium (400 mg) and oxalate or a diet higher in calcium (1200 mg) with restricted intake of oxalate, protein and salt.1 At 5 years, the latter group had a 51% lower rate of stone recurrence than those following a low-calcium diet. Although controversy persists and replication of these findings under other clinical circumstances would be desirable, low-calcium diets are not recommended and can exacerbate the well-documented association of hypercalciuria with low bone mineral density and increased fracture rates.2 Whether calcium supplements could have a similar effect in lowering stone recurrence rates has not been tested. Taking cal-

| Risk factor           | Causes                                                                 | Treatment                                         | Level of evidence |
|-----------------------|------------------------------------------------------------------------|---------------------------------------------------|-------------------|
| Low urine volume      | Exercise, sweating, low fluid intake, heat, bowel disease              | Increase fluid intake to 2.5-3.0 L/d               | RCT (Borghi et al. J Urol 1996;155:839-43)           |
| Hypercalciuria        | Idiopathic, vitamin D intoxication                                     | Reduce sodium intake; begin thiazide diuretic therapy; do not reduce calcium intake except in extreme cases; increase dietary calcium intake to 1000 mg | RCT (Borghi et al. N Engl J Med 2002;346:77-84)      |
| Hyperoxaluria         | Dietary ingestion and endogenous metabolism; inflammatory bowel disease| Restrict oxalate intake; increase dairy intake accompanying oxalate-containing foods | No RCT (Holmes et al. Urol Res 2004;32:311-6)        |
| Hypocitraturia        | Renal tubular acidosis; other metabolic acidoses; chronic bowel disease; often idiopathic | Begin potassium citrate supplementation           | RCT (Barcelo et al. J Urol 1993;150:1761-4)         |
| Hyperuricosuria       | Excessive purine ingestion as animal protein                           | Reduce purine ingestion; begin allopurinol therapy | RCT (Ettinger et al. N Engl J Med 1986;315:1386-9)  |
| Increased sodium excretion | Excessive dietary sodium intake                                        | Restrict sodium intake                            | No RCT of sodium restriction alone (Borghi et al. N Engl J Med 2002;346:77-84) |

Note: RCT = randomized controlled trial.
Calcium carbonate supplements with meals reduces oxaluria, whereas taking them at bedtime increases calcium and has no effect on oxaluria. The preferred calcium supplement for people at risk of stone formation is calcium citrate because it helps to increase urinary citrate excretion. We recommend a dose of 200–400 mg if dietary calcium cannot be increased.

**Oxalate**

The intake of foods high in oxalate should be limited. Although there are many such foods, some more frequently than others have been shown to be most at fault for raising urine oxalate levels (Box 1). Their impact might be mitigated by accompanying them with additional fluids and dietary sources of calcium to diminish oxalate absorption. No pharmacologic intervention has definitively been shown to be effective. Of note, since vitamin C can convert to oxalate, the use of vitamin C supplements may increase oxaluria and be associated with an increased risk of stone formation; therefore, the dose of these supplements should be limited to less than 1000 mg/d.

**Sodium**

Calcium excretion is directly linked to sodium excretion. Reductions in dietary sodium reduce calciuria, but no RCT of sodium restriction alone has been performed. The successful diet mentioned earlier that was higher in calcium and restricted intake of oxalate, protein and salt achieved a reduction in calcium excretion despite the higher calcium intake. This effect was attributed to the reduction in dietary sodium intake to less than 2 g/d.

**Protein**

The ingestion of animal protein has adverse effects on urine chemistries: it lowers citrate excretion and increases calcium and uric acid excretion. Epidemiologic data show that it correlates well with the prevalence of stone formation. The recent popularity of diets low in carbohydrates and high in animal protein have refocused attention on protein intake as a risk factor for stones. Patients with recurrent stones should minimize their protein intake to less than 80 g/d.

**Role of diabetes and obesity in stone formation**

Recent data have suggested an increased prevalence of stones among people with diabetes and among obese people. Insulin resistance may reduce urinary citrate excretion and increase calcium excretion, and a greater body mass index is associated with increases in urinary oxalate excretion. Low urine pH is associated with increased risk of uric acid stones as well. Weight loss (through a mechanism other than a diet high in animal protein) and improved diabetes control may help to prevent stone formation in appropriate patients, although this approach has yet to be tested in an RCT.

**Drug therapy**

**Thiazide diuretics**

Thiazide diuretics (e.g., hydrochlorothiazide, chlorthalidone and indapamide) have been proven in RCTs to be effective in reducing calciuria and stone recurrence. These drugs also induce positive calcium balance and thereby increase bone mineral density. Sodium restriction must be employed, and hypokalemia must be avoided since it can lead to hypocitraturia. The usual starting dose is 12.5–25 mg of chlorthalidone or hydrochlorothiazide. Most patients should receive supplementation with potassium citrate. Amiloride may help avoid hypokalemia, but triamterene should be avoided because of its low solubility.

**Alkalinizing agents**

Citrates inhibits the formation and growth of calcium crystals. Its use has been advocated both in patients with low urinary citrate excretion and in patients who have had calcium oxalate stones but who do not have this urinary abnormality. Use of the potassium salt is preferred, since sodium citrate supplementation will lead to increased calciuria.

**Allopurinol**

A minority of patients with calcium oxalate stones have hyperuricosuria as their only identifiable metabolic risk factor. Uric acid facilitates the precipitation of calcium oxalate crystals. A single RCT demonstrated the efficacy of allopurinol, 100 mg 3 times daily (although we prescribe 300 mg once daily), to reduce calcium stone formation. No benefit of allopurinol in patients with hypercalciuria was identified.

**Fish oil**

Interest in fish oil supplementation arose from the observation that kidney stones were virtually absent in the Inuits of Greenland, a population with a dietary mainstay of fresh fish. Supplementation with eicosapentaenoic acid leads to reduced calciuria but has not been tested in an RCT.

**Summary**

A limited number of RCTs have shown that a good strategy for preventing the recurrence of calcium oxalate stones includes increasing fluid intake, increasing the dietary intake of calcium and restricting the intake of salt, animal protein and oxalate-rich foods. Treatment with a thiazide diuretic, allopurinol and citrate has a role in se-

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**Box 1: Foods most commonly responsible for raising urine oxalate levels**

- Rhubarb
- Dark green leafy vegetables (e.g., spinach, chard, arugula)
- Beets
- Wheat bran
- Nuts, seeds
- Soy products
- Chocolate
- Tea
- Strawberries

Source: Massey L et al. J Am Diet Assoc 1993;93:901-6. See also www.ohf.org/docs/Oxalate2004.pdf and www.litholink.com/patientsDietInfo.htm
lected cases. Several other means of manipulating urinary chemistries have not been adequately tested in RCTs directed at stone prevention.

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