Diet-induced oxalate nephropathy from excessive nut and seed consumption

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SUMMARY
Oxalate is a metabolite consumed in nuts, beans and leaves, and excreted in urine. Oxalosis can cause nephropathy. We describe a rare case of a high-oxalate diet intended for irritable bowel syndrome (IBS) treatment causing oxalate nephropathy. A 59-year-old woman with a history of controlled hypertension presented with creatinine 1.8 mg/dL, increased from baseline 1.3 mg/dL. She denied recent illness, urinary stones, medication adjustments, herbal supplements and non-steroidal anti-inflammatory drugs use. Diet included six tablespoons of chia seeds and five handfuls of almonds daily to manage IBS symptoms. Renal biopsy showed glomerulosclerosis, fibrosis and calcium oxalate deposition. She switched to a low-oxalate diet, with improvement in laboratory markers. An earlier dietary history could have raised concern for oxalosis prior to renal biopsy. Providers should be trained to identify at-risk patients and provide appropriate dietary counselling.

BACKGROUND
Oxalate is an organic acid and end product of ascorbic acid metabolism in humans and plants. In humans, dietary free oxalate is absorbed in the stomach, distal small intestine and colon; oxalate bound to calcium, iron and magnesium is excreted in faeces.1 Serum oxalate is excreted in urine and excessive filtration can cause calcium oxalate stones or nephropathy. Oxalate nephropathy (ON), is a rare form of kidney injury from calcium oxalate crystal deposition in the kidney parenchyma. It can lead to tubular damage, interstitial inflammation and fibrosis that can cause acute or chronic renal failure, and may progress to end-stage renal disease. Oxalosis is most commonly caused by fat malabsorption, but other factors include mineral deficiencies, fasting states, loss of oxalate, degrading microbes, genetic defects in metabolism and dietary intake of 1 g of soluble oxalate daily compared with the normal western diet of <200 mg/day.1

TREATMENT
Based on the high oxalate content of her urine collection and calcium oxalate deposition on kidney biopsy, she was counselled to switch to a low-oxalate diet with high fluid intake. She started calcium carbonate with meals and worked with a nutritionist.

OUTCOME AND FOLLOW-UP
Follow-up labs 2 months later showed improved creatinine to 1.57 mg/dL, and it has remained at her baseline 1.3–1.6 mg/dL. Repeat 24-hour urine testing showed normalised oxalate (33 mg/day) and citrate (624 mg/day) excretion, with pH 7.2 (nl 5.8–6.2). Since cutting down on nuts and changing her diet, she has lost weight. She exercises daily and her blood pressures have remained with systolics at

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sprue, pancreatic deficiency and medications leave more free absorption, such as gastric bypass surgery, Crohn’s disease, coeliac disease or secondary from increase in oxalate absorption from the gastrointestinal tract and kidney health. Oxalosis can be either primary from genetic enzyme deficiencies or secondary from increase in oxalate absorption from the gastrointestinal tract. Normally, small amounts of free oxalate are absorbed by the stomach, distal small intestines and colon in humans. Oxalate also binds calcium, iron and magnesium, and in the gut this gets excreted with faeces. Any host or dietary factor that alters the amount of free oxalate influences absorption. The most common cause of secondary oxalosis is from fat malabsorption, responsible for 88% of cases in a systematic review by Lumlertgul et al. Conditions that cause fat malabsorption, such as gastric bypass surgery, Crohn’s disease, coeliac sprue, pancreatic deficiency and medications leave more free oxalate available for absorption. This is because fat has a greater affinity for calcium than oxalate. Calcium, iron and magnesium deficiencies as well as fasting states can also increase oxalate absorption. The microbiome, including Oxalobacter formigenes and Lactobacillus acidophilus, plays an important role in degrading oxalate. The diversity and extent of these gut bacteria fluctuates in response to dietary oxalate and antibiotics, further influencing oxalosis. Diets high in free oxalate also result in increased absorption.

Dietary oxalate comes primarily from plants. It is found in higher concentrations in the leaves and seeds. Foods known to have high levels of oxalate include nuts, beans, tea, spinach and kale, rhubarb, beets, potato skins, soy and cocoa. Vitamin C is also a potent source of oxalate as it is an end product of metabolism. Absorption rates vary between foods depending on the content of soluble oxalate, which is more bioavailable, and tea has a significantly higher percentage of oxalate absorption compared with spinach and rhubarb. Almonds, Brazil and pine nuts contain nearly twice as much soluble oxalate, compared with peanuts, pistachios and chestnuts.

Our patient consumed a high-oxalate diet as a result of following a popular diet for digestive health. The virgin diet, promoted to alleviate food intolerances and IBS, advocates elimination of certain foods and consumption of greens, nuts and seeds. In response, our patient consumed an estimated five times the typical quantity of oxalate daily. She ingested approximately 150 g of almonds daily (147–250 mg of gastric soluble oxalate/100 g and 216–305 mg of intestinal insoluble oxalate/100 g) and six tablespoons (1/8 cup) of chia seeds (380 mg oxalate/one-quarter cup), which ultimately caused kidney injury. While high-oxalate foods are nutritional, oxalosis can be harmful. Case reports have documented diet-induced ON secondary to spinach, kale, berries, nuts and seed milk, cocoa powder, wheat germ, green smoothies, chagas mushrooms and vitamin C supplements. The majority followed this diet for presumed health benefits and had partial recovery with restriction of oxalate consumption, increased oral hydration and calcium acetate supplementation to bind gut oxalate.

This case proves the importance of taking a thorough dietary history. In our case, a dietary history could have raised a concern for oxalosis earlier. While hypertension may have also contributed to the development of interstitial fibrosis, the patient’s blood pressure was controlled at the time of biopsy, thus the major intervention after completing the biopsy and urine collection was dietary counselling. At-risk patients with pre-existing gastrointestinal disease, kidney disease or mineral deficiencies should be counselled on the effect of high-oxalate diets on the kidneys. Vegetarians, who may consume greater quantities of foods rich in oxalate may also be vulnerable.

**Learning points**

- While oxalate nephropathy is a less common cause of kidney disease, it should be considered when patients present with chronic kidney disease or acute-on-chronic kidney injury, particularly in patients with history of pre-existing gastrointestinal, kidney or mineral disease.
- A thorough dietary history is extremely important in disease prevention, intervention and management.
- Curriculum in physician training should include nutrition education so that providers are comfortable discussing diet and can provide appropriate counselling.

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**Figure 1** Trichrome taken at 40x magnification showing the background fibrosis and glomerulosclerosis.

**Figure 2** H&E stain taken at 600x magnification with polarisation showing a couple of calcium oxalate deposits within renal tubules.
cooking techniques, such as soaking and cooking plants to help leach oxalate, fluid hydration, balancing high-oxalate diets with foods rich in free calcium, magnesium and iron, and limiting consumption of tea and nuts is important.2

Given the popularity of fad diets, the curriculum in physician training should include nutrition education. Providers must learn to take a dietary history and be prepared to provide appropriate counselling.

Contributors VG is the corresponding and first author. She is a third-year Internal Medicine resident (PGY3) at the George Washington University. She performed the literature review, wrote and revised the manuscript under consideration. LH was the pathologist on the case and prepared the slides. The senior author, RR is the patient’s current nephrologist. She gave guidance for the paper, and helped with editing and revising the manuscript. All authors have accountability and responsibility for the work and have approved of the final product.

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