

Causes

Amphotericin B Toxicity

[Amphibian-terminator-bee](#)

Impaired renal function is a relatively common complication of amphotericin B, including urinary potassium wasting and hypokalemia, urinary magnesium wasting and hypomagnesemia, and metabolic acidosis due to RTA type I. This is reversible with the suspension of the drug. The mechanism of RTA type I induced by amphotericin is due to an increase in membrane permeability caused by the drug which results in the reduction of ion concentration gradients between the cytoplasm of distal tubule cells and the tubule lumen. In this context, potassium leaks from the cytoplasm down a favorable concentration gradient into the lumen (causing hypokalemia due to increased urinary loss) and hydrogen ions diffuse down their gradient from the lumen into the cytoplasm of distal tubule cells. Therefore it can be said that the defect in acid excretion induced by amphotericin b is due to back-diffusion of secreted hydrogen ions.

Autoimmune Disease

[Auto-in-moon](#)

The mechanism of RTA type I in autoimmune diseases is mainly due to autoantibodies against the transporters in the intercalated cells. For example, in primary Sjogren's syndrome, renal involvement is one of the extraglandular manifestations and in most cases, it affects the renal tubules through tubular interstitial nephritis and occasionally autoantibodies against a certain transporter. All segments of the nephron can be involved but distal RTA is the most frequent tubular dysfunction in Sjogren's syndrome. A similar mechanism is involved in other autoimmune diseases.

Lithium

[Lithium-Battery](#)

Drugs causing type I RTA include: lithium, amphotericin B, NSAIDs, Ifosfamide. The mechanism of RTA type I induced by lithium is similar to the mechanism mentioned for amphotericin b. It is possible that lithium administration induces distal renal tubular acidosis by allowing excessive back-diffusion of acid.

Obstruction Of The Urinary Tract

[Obstructed kidney with tuba](#)

In obstructive uropathy, there has been shown to be a transport defect in the distal nephron, characterized by impaired sodium reabsorption and decreased secretion of hydrogen and potassium leading to distal tubular acidosis. Note that in obstructive uropathy there would be hyperkalemia and not hypokalemia. In the other causes of RTA type I there is hypokalemia.

Complication

Kidney Stones

[Kidney-throwing Stones](#)

Calcium stones are formed because calcium tends to precipitate in an alkaline solution. Other factors contribute to calcium stone formations such as reduced urinary citrate since acidemia enhances proximal citrate reabsorption. Additionally, the acidemia present in RTA causes increased calcium phosphate release from bone during bone buffering of retained acid. This increased bone turnover is also associated with an increased risk of calcium stones. Calcium phosphate stones may be seen bilaterally and this should raise suspicion for RTA type I. Patients with recurrent calcium stones (particularly calcium phosphate stones) and a urine pH that is persistently 5.5 or higher should be evaluated for distal RTA.

Treatment

Bicarbonate

[Bi-car-bombs](#)

Treatment consists of correcting metabolic acidosis with Alkali therapy with sodium bicarbonate or sodium citrate (Shohl's solution). If hypokalemia persists despite the correction of the acidosis, serum bicarbonate potassium citrate alone or combined with sodium citrate is indicated.