Kidney biopsy for renal tubular acidosis: when tissue diagnosis makes a difference

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Abstract. Renal tubular acidosis (RTA) is a disorder with variable presentations and oftentimes a nebulous underlying primary diagnosis. We describe a rare cause of RTA as an unusual complication of proton pump inhibitor (PPI) therapy. We report a case of a 33-year-old male with history of hypertension, acid reflux, allergic rhinitis, and low testosterone admitted with complaints of fatigue, weight loss, and unexplained acidosis for ~2 months. His medications prior to admission included losartan, omeprazole, potassium chloride, sildenafil, and testosterone propionate injections. His physical exam was unremarkable with a blood pressure of 120/80 mmHg. Initial lab work showed a nonanion gap metabolic acidosis with serum bicarbonate level of 16 mM/L and potassium 3 mM/L. Urine studies showed urine pH of 6.5 and a positive urine anion gap. The serum creatinine was within normal range (1.13 mg/dL). He required massive doses of bicarbonate and potassium supplementation with minimal improvement of serum chemistries achieved. The cause of apparent distal RTA remained elusive despite extensive blood, urine, and imaging testing. Ultimately, a renal biopsy was obtained showing mild to moderate tubule-interstitial inflammation with 5% fibrosis. PPI therapy (omeprazole) was stopped, and he was started on prednisone 60 mg per day. Two weeks later, his RTA findings resolved, and he no longer required bicarbonate and potassium supplementation. Our case highlights the importance of recognizing a unique complication of RTA following PPI therapy. It also underscores the possible need for considering a kidney biopsy in the setting of nondiagnostic laboratory work up to uncover the underlying etiology of RTA and suspected allergic interstitial nephritis (AIN).

Clinical presentation

A 33-year-old male with history of hypertension, acid reflux, allergic rhinitis, and low testosterone was admitted with complaints of fatigue, weight loss, and unexplained acidosis for ~2 months. He had no exposure to lead, gasoline, paints, adhesives, or glues. His medications prior to admission included losartan, omeprazole, potassium chloride, sildenafil, and testosterone propionate injections. His physical exam was unremarkable, with a blood pressure of 120/80 mmHg. Initial lab work showed a nonanion gap metabolic acidosis with serum bicarbonate level of 16 mM/L and potassium 3 mM/L. Urine studies showed urine pH of 6.5 and a positive urine anion gap. The serum creatinine was within normal range (1.13 mg/dL). He required massive doses of bicarbonate and potassium supplementation with minimal improvement of serum chemistries achieved. The cause of apparent distal RTA remained elusive despite extensive blood, urine, and imaging testing. Ultimately, a renal biopsy was obtained showing mild to moderate tubule-interstitial inflammation with 5% fibrosis. PPI therapy (omeprazole) was stopped, and he was started on prednisone 60 mg per day. Two weeks later, his RTA findings resolved, and he no longer required bicarbonate and potassium supplementation. Our case highlights the importance of recognizing a unique complication of RTA following PPI therapy. It also underscores the possible need for considering a kidney biopsy in the setting of nondiagnostic laboratory work up to uncover the underlying etiology of RTA and suspected allergic interstitial nephritis (AIN).

Introduction

We describe a rare cause of renal tubular acidosis (RTA) as an unusual complication of proton pump inhibitor therapy. The patient presented with primary disturbance of RTA with seemingly normal kidney function. Despite the traditional diagnostic strategies focusing on serum and urine studies, the underlying etiology remained elusive. Ultimately, the patient required a percutaneous kidney biopsy (PKB) to secure the diagnosis of an occult acute interstitial nephritis (AIN) and RTA, likely a consequence of proton pump inhibitor (PPI) exposure.
ies, hepatitis-C antibodies, antinuclear antibodies, Anti-Ro, and Anti-La antibodies were also negative. He had no complaints of fatigue, arthralgias, dry eyes/dry mouth, or rashes, suggesting an underlying seronegative rheumatologic disorder.

He needed large doses of potassium and sodium citrate supplementation and per os amiloride to restore near-normal serum electrolyte values. Due to diagnostic uncertainty, a renal biopsy was performed, showing findings consistent with AIN and mild to moderate interstitial lymphocytic infiltration with evidence of tubulitis within the cortex. There was also 5% interstitial fibrosis, signifying a chronic component as well (Figure 1 and 2).

Upon further retrieval and review of past records, his baseline creatinine was 0.6 – 0.8 mg/dL ~ 3 months prior to his hospital presentation. He was also started on omeprazole around that time, suggesting a close temporal relationship between rise in serum creatinine and PPI initiation. In light of the above, his PPI therapy was immediately stopped. Based on retrospective studies [1] and relative safety of short-term steroid therapy, we felt it was reasonable to proceed with a short course of glucocorticoids (oral prednisone 60 mg daily, weaned over 6 weeks). Two weeks later, during his outpatient follow-up appointment, the RTA was resolved and he no longer required bicarbonate and potassium supplementation.

**Discussion**

Renal tubular acidosis is a relatively uncommon disorder in adults, which is defined based on different renal pathophysiologic causes of a hyperchloremic (normal anion gap) metabolic acidosis [2]. It is generally classified based on location of injury (proximal tubule or distal tubule), which signifies the primary defect unique to the function of that segment of the nephron. The corollary is that proximal tubule defect (type 2) consists of the inability to reclaim bicarbonate, and the distal tubule has the inability to secrete acid (type 1) or has a hypoaldosterone state (type 4) that drive the nonanion gap acidosis [3]. The etiology of these disorders is very heterogeneous, ranging from infectious, autoimmune, malignancy, metabolic, and structural to drug-induced disorders [2]. Accordingly, clinicians often have to work backwards to determine the primary cause of the problem, and oftentimes the inciting event remains a
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mystery. This was best exemplified by a retrospective case analysis of 58 distal renal tubular acidosis (dRTA) cases done in Baltimore showing that 55% of them were classified as idiopathic [4]. The work up of dRTA often relies of blood, urine, and noninvasive imaging testing; tissue sampling customarily is not pursued for making the diagnosis.

While dRTA is a well-described phenomenon of chronic tubulointerstitial disease, it is rarely seen as the sole pathologic finding of AIN. Classically, AIN is often defined as an acute decline in renal function with an inflammatory infiltrate within the kidney interstitium, but it can have more subtle presentations as well. Furthermore, it is known that the “classic” triad of fever, rash, eosinophilia has an insufficient sensitivity and specificity for the diagnosis [5]. Nonetheless, clinicians still use the “classic triad” as the primary means of diagnosis of AIN. As a result, AIN is underreported, and renal biopsies only account for roughly 15% of all acute kidney injuries [5]. In our patient, the cause of dRTA remained obscure despite extensive blood, urine, and imaging testing. It can be concluded that there is insufficient data to describe the full spectrum of renal injury due to either missed diagnosis or lack of histopathologic correlation. Given his normal baseline, we had a high index of suspicion for an anatomic (histological) process to explain this marked and seemingly sudden emergence of biochemical abnormalities. There have been a few case reports describing RTA in settings of acute kidney injury (AKI) secondary to AIN due to medications [6]. There is strong evidence suggesting PPIs, in particular omeprazole, being associated with AIN-related injury. This was best demonstrated by a single-center retrospective study showing biopsy-proven AIN in a setting of PPI therapy [7]. There is little to no data with regards to metabolic acidosis and hypokalemia as presenting cause of PPI-associated AIN. We found one case report suggesting that PPI can worsen acidosis and hypokalemia [8]. The PPIs mechanism of action is inhibiting the H, K ATPase in the stomach for treatment of gastroesophageal reflux disease. Although the kidney also has H, K ATPase channels, it is known to not be affected by the proton pump inhibitor action [9]. PPI are able to inhibit H, K ATPase only at very low PH (< 2), which only occurs in the canaliculi of the parietal cells, but not in the kidney. Therefore, a PPI is extremely unlikely to explain metabolic acidosis based on the “pharmacologic” effect of the medication. There is only a single case report describing a correlation of PPI with worsening metabolic acidosis and hypokalemia [8].

The diagnosis of AIN is generally made on clinical suspicion, and, thus, most of the time there is low sensitivity and specificity of making that diagnosis [10]. It is not uncommon for PPI’s to cause an AIN, but it is exceedingly rare for it to present solely as dRTA. As a result, it would be prudent to undertake an intervention that would provide the most benefit with the least potential for adverse effect, such as simply withholding medication. Percutaneous kidney biopsy (PKB) is generally a safe and effective procedure to secure histologic sampling of the renal cortex, included in our institutional experience in the context of graduate medical training [11, 12]. The results of the PKB showed us the definitive finding of AIN as well as some early scarring, making us concerned about potential long-term repercussions of this event (perhaps decades later) [7]. One of the larger retrospective studies done in patients with AIN due to medications showed improved renal recovery with no significant difference in adverse events in patients treated with steroids when compared to the control group [1]. Accordingly, given the potential for benefit and safety of short-term glucocorticoid therapy for AIN [1, 7], we felt it was reasonable to proceed with a short course of prednisone, as described in our report. Our case also underscored the importance of obtaining PKB during work-up of otherwise unexplained RTA [13].

Conclusion

RTA is condition that can have a variable and subtle presentation. Our case demonstrated that traditional laboratory work-up may not suffice to fully evaluate the underlying etiology. Furthermore, PKB may be warranted to rule out discrete histopathologic injury, such as AIN. Early diagnosis may be crucial in reversing morbidity of an ongoing subclinical injury.
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Conflict of interest

The authors have no conflict of interest to declare.

References

[1] González E, Gutiérrez E, Galeano C, Chevia C, de Sequera P, Bernis C, Parra EG, Delgado R, Sanz M, Ortiz M, Goicoechea M, Queveda C, Olea T, Bourich H, Hernández Y, Segovia B, Praga M; Grupo Madrileño De Nefritis Intersticiales. Early steroid treatment improves the recovery of renal function in patients with drug-induced acute interstitial nephritis. Kidney Int. 2008; 73: 940-946. CrossRef PubMed

[2] Rodríguez Soriano J. Renal tubular acidosis: the clinical entity. J Am Soc Nephrol. 2002; 13: 2160-2170. CrossRef PubMed

[3] Rossert JA, Fischer EA. Acute interstitial nephritis. In: Comprehensive Clinical Nephrology, 2, Johnson RJ, Feehally J. (Eds), Elsevier Limited, Philadelphia 2003. Vol 1, p.769.

[4] Caruana RJ, Buckalew VM Jr. The syndrome of distal (type 1) renal tubular acidosis. Clinical and laboratory findings in 58 cases. Medicine (Baltimore). 1988; 67: 84-99. CrossRef PubMed

[5] Praga M, González E. Acute interstitial nephritis. Kidney Int. 2010; 77: 956-961. CrossRef PubMed

[6] Neelakantappa K, Gaal GR, Lowenstein J. Renal and Fanconi syndrome. Am J Kidney Dis. 1993; 22: 333-336. CrossRef PubMed

[7] Murithi AK, Leung N, Valeri AM, Cornell LD, Sethi S, Fidler ME, Nasr SH. Biopsy-proven acute interstitial nephritis, 1993-2011: a case series. Am J Kidney Dis. 2014; 64: 558-566. CrossRef PubMed

[8] Okamoto N, Nambu T, Matsuda Y, Matsuo K, Osaki K, Kanai Y, Ogawa Y, Yonemitsu S, Iita R, Muro S, Sugawara A, Oki S. Distal renal tubular acidosis that became exacerbated by proton pump inhibitor use. Intern Med. 2012; 51: 2591-2595. CrossRef PubMed

[9] Howden CW, Reid JL. Omeprazole, a gastric “proton pump inhibitor”: lack of effect on renal handling of electrolytes and urinary acidification. Eur J Clin Pharmacol. 1984; 26: 639-640. PubMed

[10] Raghuwan R, Eknoyan G. Acute interstitial nephritis – a reappraisal and update. Clin Nephrol. 2014; 82: 149-162. CrossRef PubMed

[11] Fülop T, Alenu B, Dosabhoj NR, Bain JH, Pruett DE, Szombathelyi A, Dreisbach AW, Tapolyai M. The safety and efficacy of percutaneous renal biopsy by Physician-in-Training in an academic teaching setting. South Med J. 2014; 107: 520-525. CrossRef PubMed

[12] Islam N, Fülop T, Zsom L, Miller E, Mire CD, Lebrun CJ, Schmidt DW. Do platelet function analyzer-100 testing results correlate with bleeding events after percutaneous renal biopsy? Clin Nephrol. 2010; 73: 229-237. CrossRef PubMed

[13] Perazella MA. Diagnosing drug-induced AIN in the hospitalized patient: a challenge for the clinician. Clin Nephrol. 2014; 81: 381-388. PubMed