Acute tubulointerstitial nephritis associated with atezolizumab, an anti-programmed death-ligand 1 (pd-L1) antibody therapy

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ABSTRACT
Direct stimulation of the antitumor activity of immune system through checkpoint inhibitors (ICIs) has demonstrated efficacy in the treatment of different cancer types. The activity of these antibodies takes place in the immunological synapse blocking the binding of the negative immunoregulatory proteins, thus leading to the finalization of the immune response. Despite having a favorable toxicity profile, its mechanism of action impedes the negative regulation of the immune activity which can potentially favor autoimmune attacks to normal tissues. Renal toxicity has been described in several ICI but not with atezolizumab, an IgG1 monoclonal antibody targeting PD-L1, approved by FDA as a second-line therapy for advanced urothelial carcinoma. Here we present a patient with a single kidney and metastatic renal cell carcinoma treated with atezolizumab and bevacizumab combination, with biopsy-proven acute interstitial nephritis, who had a complete resolution of renal dysfunction after steroid therapy.

Introduction
Despite immune system is able to eradicate cancer, malignant cells manage to escape immune attacks through different strategies. The capacity to exploit the host immune system to treat cancer is known as immunotherapy. Among the different treatments that this concept encompasses, immune checkpoint inhibitors (ICI), such as antibodies anti-cytotoxic T lymphocyte-associated antigen-4 (CTLA-4), anti-programmed cell death protein 1 (PD-1) and anti-programmed cell death ligand 1 (PD-L1), have demonstrated efficacy in the treatment of different cancer types. Roughly, the activity of these antibodies takes place in the immunological synapse blocking the binding of the negative immunoregulatory proteins, thus leading to the finalization of the immune response. The physiological function of these proteins is to protect the host against autoimmune attacks. Unfortunately, one of the strategies that cancer cells use to evade immune system is precisely through the overexpression of these immunoregulatory molecules, such as PD-L1.

Atezolizumab is a fully humanized IgG1 monoclonal antibody targeting PD-L1 that has been approved by FDA as second-line therapy for advanced renal carcinoma. Although anti-PD-1/PD-L1 antibodies have a favorable toxicity profile, its mechanism of action impedes the negative regulation of the immune activity which can potentially favor autoimmune attacks to normal tissues, known as immune-related adverse events (irAE). Among them, infrequent renal toxicity has been previously described with the use of other ICI such as nivolumab or pembrolizumab, but not with atezolizumab. Here we present, to our knowledge, the first case of immune-mediated acute tubulointerstitial nephritis (ATIN) associated with atezolizumab.

Case report
A 54-year-old Caucasian male, heavy smoker (accumulated dose: 40 packs year), with history of essential hypertension and depressive syndrome, was diagnosed with renal cancer in the context of hematuria. Primary tumor was located in left kidney and was associated with left renal vein invasion; additionally, he had synchronous lung metastases [American Joint Committee of Cancer (AJCC) Stage IV; T3aN0M1]. The patient underwent left radical nephrectomy; histopathological assessment confirmed renal cell carcinoma (RCC) with 20% of sarcomatoid differentiation foci. The patient accepted to participate in the clinical trial NCT02420821, and he was randomized in the treatment arm with atezolizumab-bevacizumab. Subsequently, it was initiated an intravenous infusion combination of atezolizumab (1200 mg) and bevacizumab (15 mg/kg) on days 1 and 22 of each 42-day cycle. Treatment was well tolerated with just few mild adverse events such as grade 1 fatigue, arthralgia, and myalgia, as per CTCAE. Disease evaluation performed at the end of the 3rd cycle showed partial response (45% decrease in the sum of target lesions).

In the visit corresponding to cycle 5 day 22, the patient referred to have had general malaise, 39°C fever and grade 1 diarrhea for the previous 2 weeks. Physical examination was
unremarkable except for temperature of 37.8°C and mild dehydration. Routine blood tests revealed acute kidney injury stage III (according to Acute Kidney Injury Network classification) with creatinine of 5.6 mg/dL (ULN: 1.3 mg/dL), which represented a sharp increase compared to his baseline value (1.2 mg/dL), and eosinophilia (7.2%, corresponding to 400 eosinophils/mm³). Urinalysis revealed 2+ protein (protein/creatinine ratio 782 mg/g), RBC 8/hpf and WBC 9/hpf. Urine sediments showed 2–3 eosinophils. There had not been recent exposure to nephrotoxic agents such as antibiotics, contrast or analgesics. He was taking, for more than three years, atenolol 50 mg q.d., enalapril 5 mg b.i.d., venlafaxine 75 mg q.d., and lorazepam 1 mg q.d. as needed. Renal ultrasound showed a normal right kidney with no evidence of hydronephrosis. Despite volume repletion and spontaneous remission of diarrhea, the patient persisted with AKI with preserved diuresis; for this reason, a kidney biopsy was performed in order to establish the etiology of the disorder.

In the renal biopsy were observed 18 glomeruli with a preserved architecture. No signs of necrotizing lesions or associated inflammation were observed in the glomerular tufts. An extensive diffuse inflammatory infiltrate consisting of both T-helper cells and cytotoxic T cells some of them with scattered PD1 deposits, plasma cells and eosinophils was observed in the interstitial component, also with tubulitis. Of note, there were some tubular cells that expressed PD-L1. No granulomatous lesions were observed. There were no remarkable findings in the vascular compartment. Immunofluorescence staining did not reveal glomerular immune deposits (Fig. 1).

The patient was treated with a steroid pulse [i.v. methylprednisolone 125 mg × 3 days], obtaining clinical and analytical improvement, characterized by resolution of fever and malaise, and a decrease of the creatinine to 1.45 mg/dL after 8–10 weeks of follow-up. Steroids were progressively tapered once periodic creatinine evaluations had shown no impairment of the renal function. Anticancer therapy has not been reintroduced since ulcerative radiological evaluations have shown partial response, which is currently maintained by the time our case is reported (PFS: 25 months).

**Discussion**

PD-L1 is a transmembrane protein widely expressed in antigen-presenting cells, including B cells, dendritic cells, macrophages and tumor cells. Its interaction with PD-1 of T cells downregulates the immune response maintaining the peripheral tolerance to self and external antigens, through the inhibition of proliferation and activation of T cells, and inhibiting the cytolytic activity of the immune system.

This mechanism of avoiding the action of immune system is used by many tumor cells by the expression of PD-L1. In fact, in many tumor types, the more overexpression of PD-L1, the worse cancer prognosis. Blocking the pathway of PD-L1/PD-1 activates T cells, and this tool has been used for therapeutic purposes, with the aim of releasing the immune system activity against the tumor tissue. But if there is an excessive response, it can also damage healthy tissue, producing irAE.

![Figure 1](image-url)

*Figure 1.* The renal biopsy showed interstitial inflammatory infiltrates with focal tubulitis (A; PAS original magnification × 20). Immunohistochemical stainings revealed a predominant T cell infiltrate (B; CD3, 20x), consisting of both T-helper cells (C; CD4, 20x) and cytotoxic T cells (D; CD8, 20x). There were scattered PD1 positive cells (E; PD1, 20x, the enlarged insert 40x) and some tubular cells expressing PD-L1 (F; PD-L1 80X).

is not only restricted to tumor cells, but it is also found in healthy tissue.

The renal tubule epithelial cells (TEC) are one of the few type of epithelial cells that also express major histocompatibility complex class II molecules, which allows TEC to function as antigen presenting cells (APC) for T cells, leading to the activation of the immune response. As APC, TEC also has the capacity to modulate T-cell responses negatively. PD-L1 has been described as a negative coestimulatory molecule, and its expression can be induced by activation with IFN-gamma. PD-L1 was not observed in renal glomerulus. Thus, it has been suggested that the TEC PD-L1/T cell PD-1 binding would have a protective role against immune-mediated tubulointerstitial injury. Therefore, if anti-PD-L1 antibodies block PD-L1 receptor, T-cells would not be deactivated, which propitiates their activity against antigens presented by TEC. Thus, T cells seems to play a major role in the pathogenesis of ATIN related to ICIs, both CD4+ and CD8+. This would be accompanied by an increase of cytokines (IL-18 and IFN-gamma, among others, as observed over the course treatment with atezolizumab), and consequently an increase in cytotoxic lymphocytes. These cytokines orchestrate the inflammatory reaction, resulting, as in others drug-induced ATIN, in an interstitial infiltrate...
of lymphocytes, macrophages, monocytes, eosinophils and/or polymorphonuclear neutrophils, in addition to interstitial edema and disruption of the tubular basement membrane.\textsuperscript{16}

Different hypotheses have been raised about the role of the interaction among lymphocytes and potential antigens in this particular ATIN. It has been reported that 14 of 19 patients\textsuperscript{17,18} had associated drugs potentially causing tubulointerstitial nephritis. Based on this observation, Izzedine and colleagues\textsuperscript{19} suggested that ICIs can reactivate drug-specific inactive T cells, previously sensitized by nephritogenic antigens. This consequently could produce a response of memory T cells against drugs. Cortazar also suggests that there could be a “reprogramming” of immune system, losing the acquired tolerance against own endogenous antigens. The latter would allow to explain the long latency observed sometimes from the administration of the drug to the development of nephritis.\textsuperscript{17}

Moreover, the combination of atezolizumab and bevacizumab, an inhibitor of the vascular endothelial growth factor (VEGF), seems to be associated with further increase in intra-tumoral CD8$^+$ cells due to enhanced trafficking, and with an increase of intra-tumoral MHC-I, Th1 and T-effector markers.\textsuperscript{20} In addition, VEGF is known to produce immunosuppressive effects. It has been reported that the VEGF produced in the tumor microenvironment enhances expression of PD-1 and other inhibitory checkpoints involved in CD8$^+$ T cell exhaustion,\textsuperscript{21} as well as the inhibition of lymphocyte maturation.\textsuperscript{22} Moreover, VEGF would have the capacity to increase the production of myeloid-derived suppressor cells (MDSC), which are able to decrease T cell function.\textsuperscript{23} Although the anti-PD-L1 and anti-VEGF association is increasingly supported due to its antitumoral effects, it can favor the development of irAE due to similar mechanisms.\textsuperscript{24}

Anti-VEGF therapies have significant renal implications, but ATIN had been described in very few anecdotic cases.\textsuperscript{25,26} Bevacizumab and other anti-VEGF drugs are frequently related with proteinuria and hypertension. The most common histopathologic kidney lesion is thrombotic microangiopathy, with other glomerular lesions occurring less frequently. The mechanism for glomerular injury may develop from loss of VEGF effect on maintaining the filtration barrier.\textsuperscript{27}

This case, with no evidence of glomerular injury on renal biopsy, reinforces the hypothesis of the synergistic role of the bevacizumab and atezolizumab combination in inducing an antitumor immune response,\textsuperscript{20} but also producing an event of ATIN as a side effect. To our knowledge, this is the first case report of biopsy-proven ATIN caused by anti-PD-L1 antibody, confirming the role of this drug as trigger of autoimmunity. The presence of an infiltrate primarily composed of CD4/CD8, PD-1 positive T cells, and some tubular cells expressing PD-L1 and acting as an antigen presenting cell; support the hypothesis of autoimmune disorder associated with this case of ATIN.

This patient has shown susceptibility to atezolizumab, since a partial response was reached in the first radiological evaluation and is currently maintained after 25 months at the expense of a severe irAE. A relationship between developing irAEs and clinical benefit from ICIs has been hypothesized and reported through the last years, firstly with ipilimumab. Beck et al.\textsuperscript{28} and Downey et al.\textsuperscript{29} found better response rates in tumor regression in those patients who developed colitis and any irAE, respectively. Contrarily, subsequent ipilimumab data from an expanded access programme\textsuperscript{30} and a retrospective study\textsuperscript{31} suggest the opposite. In the case of the anti-PD-1 antibodies, there has been reported a relationship between cutaneous irAE and better outcomes in patients with melanoma.\textsuperscript{32-36} However, there is still no general agreement between the correlation of clinical benefit from ICIs and outcomes. It is a priority to explore this relationship in the future.

In summary, our case highlights the aggressive T cell-mediated pathophysiology of ATIN in the setting of ICI therapy with atezolizumab. These adverse effects are potentially serious, especially in patients with decreased renal mass, so common in RCC, despite being the incidence of ATIN ICIs-associated in the general population low (≈1%).\textsuperscript{37} Close follow-up and early treatment with steroids are the key for the management of this complication.

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