TINU syndrome: review of the literature and case report

Síndrome TINU: revisão de literatura à propósito de um caso

Abstract

Introduction: Tubulointerstitial nephritis and uveitis syndrome (TINU Syndrome) is an uncommon clinical entity, and the majority of patients are adolescents and young women. The case reported refers to an elderly patient with ophthalmologic symptoms detected earlier than kidney manifestations, being probably the first case described in Brazil. Case Report: Female patient, 60 years old, sought medical attention for complaints of "red eye". Three months after the first episode of eye manifestation, the patient presented with systemic symptoms and renal dysfunction. Renal biopsy showed tubulointerstitial nephritis with signs of activity. Discussion: The pathophysiology of TINU Syndrome remains poorly understood, probably involving both cellular and humoral immunity. This syndrome can be differentiated from systemic conditions associated with nephritis and uveitis, and prior exclusion of other diseases is necessary to confirm diagnosis, especially in the presence of ophthalmologic findings. Conclusion: The clinical suspicion and the knowledge of the management of the disease by nephrologists, internists and ophthalmologists is mandatory in the treatment of patients with TINU Syndrome.

Keywords: acute kidney injury; autoimmune diseases; interstitial; nephritis; uveitis.

Introduction

Tubulointerstitial nephritis (TIN) is a significant cause of kidney failure in children and adults, and it is characterized by renal histological and functional abnormalities.1 TIN is rarely accompanied by uveal inflammation, an uncommon clinical entity,2-4 first described in 1975 by Dobrin et al.,4 receiving the name tubulointerstitial nephritis and uveitis syndrome (TINU syndrome).7 TINU is characterized by favorable acute tubulointerstitial nephritis and recurrent chronic uveitis.8

There are about 200 cases reported in the literature,9 and most of the TINU patients are adolescents and young...
women.\textsuperscript{10} Its etiology is unknown, with several possible triggers that can start it, such as infections and drug use.\textsuperscript{2,11}

Diagnosis is mainly clinical, proving the presence of TIN and uveitis with no evidence of other systemic or infectious diseases. Symptoms are mostly nonspecific, including fever, headache, fatigue and anorexia,\textsuperscript{1,12,13} causing the clinical diagnosis to be difficult and often delaying treatment onset.\textsuperscript{1}

The case described below was found in an elderly patient, with pre-nephritis ophthalmic manifestations, and it appears to be the first report of its kind from Brazil.

**Case Report**

Female patient, 60 years old, with a past of systemic hypertension, sought medical treatment in September 2010, complaining of a “Red Eye” (Figure 1) of sudden onset, with pain on palpation and eye movement, associated with mild bilateral eyelid edema. She was diagnosed with non-bacterial conjunctivitis. Corticosteroid-based eye drops were prescribed for seven days, with symptom resolution. Seven days after interrupting the use of eye drops, the patient developed a new ophthalmic crisis, to which the prior approach was maintained, and symptoms subsided.

![Figure 1. Anterior uveitis in patients with TINU syndrome.](image)

Thirty days after the condition had started, the patient presented with low-intensity pain in her upper left quadrant, continuous, not ventilator-dependent, for 15 days, progressing to severe pain. A chest X-ray revealed mild pulmonary infiltrates in the left lung base. It was suggested the possibility of atelectasis secondary to pain or pneumonic disease, treated with clarithromycin for 7 days and 2 days with dexamethasone/indomethacin, to which she responded with clinical and radiological improvement. Thirty days after pain resolution, the patient had an episode similar to her initial ophthalmic condition. Three months after the first episode of her eye disease, she presented with systemic symptoms and lost 5 kg (7% of body weight) in one month without changes to her blood pressure.

Laboratory tests showed azotemia - creatinine: 2.5 mg/dL and urea: 75 mg/dl, while her baseline creatinine was 0.9 mg/dl. Urinalysis with urine glucose ++, without other changes; 24-hour proteinuria: 1.65g; Renal ultrasound: kidneys of normal size, slight increase in cortical echogenicity and good cortex-medulla differentiation. The other tests carried out are shown on Table 1. After admission, she was submitted to pulse therapy with methylprednisolone 500 mg for 3 days followed by prednisone 1 mg/kg/day for 1 month, and the medication was weaned during for 4 months, with progressive improvement in her tests. In January 2011 (post-pulse therapy), she was submitted to a renal biopsy, which showed glomeruli within normal limits (global sclerosis: 2/30), tubular epithelial degenerative changes with spotted atrophy and mild diffuse interstitial fibrosis and tubulointerstitial nephritis with signs of inflammatory activity. Direct immunofluorescence revealed no glomerular deposits and tubular cylinders with IgM and IgA (Figure 2).

Three months after discontinuing the immunosuppressive medication, she had a new episode of anterior uveitis, and was started on corticosteroid eye drops and ketorolac. As a side effect, she had an increase in intraocular pressure, and was started on brinzolamide/timolol and brimonidine tartrate. We tried to wean her off the corticosteroid eye drops at times, but always with worsening of uveitis.
Currently, after 52 months of follow up, the patient remains in use of cyclosporine ophthalmic emulsion because of the failure in interrupting the medicine. Urinalysis is within the normal range, and urea and creatinine with values of 36 mg/dL and 1.0 mg/dL, respectively.

**DISCUSSION**

The TINU syndrome has an approximate prevalence of 3.5 cases/million people and an incidence of 0.2 cases/million/year,\(^3,7\) probably being underdiagnosed;\(^14,15\) without racial or ethnic predominance among the cases reported.\(^16\)

There is a higher incidence among women,\(^2,9\) in the ratio of 3:1 and, according to recent studies, the prevalence in men has increased.\(^3\)

The disease manifests earlier in men, ranging from 9-52 years; while in women it involves those ranging from 10-74 years of age.\(^2,7,10\) There are few cases described in the literature involving the elderly.\(^7,9\)

The main manifestations are nonspecific and associated with the renal disease, including, in decreasing order, fever, weight loss, fatigue and malaise, anorexia, abdominal pain, among others.\(^1,2,7,9,15-17\)

Uveitis may precede (21%), happen concurrently (15%) or succeed (65%) nephritis,\(^9,16,18\) with eye symptoms starting, in average, one month after the systemic effects. The main ocular disease is anterior uveitis, which occurs in 80% of patients, and the most common symptoms are pain and redness.\(^2\)

In our patient, the disease manifested initially by anterior uveitis, which happened three months before the renal involvement. After the onset of nephritis, there were new episodes of uveitis, showing a lack of temporality between the two events.

The pathophysiology of the TINU syndrome remains poorly understood, but reports suggest that

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<th>Table 1</th>
<th>Laboratory tests from the patient of the Case Reported Above</th>
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<td>23/12/10</td>
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<td>Hb (g/dl)</td>
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<td>/Ht (%)</td>
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<tr>
<td>VHS (mm)</td>
<td>71</td>
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<td>PCR (mg/dL)</td>
<td>0.28</td>
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<td>Ur (mg/dl)</td>
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<tr>
<td>Cr (mg/dl)</td>
<td>2.5</td>
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<tr>
<td>Proteinuria 24h</td>
<td>1.65 g</td>
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**Figure 2.** Kidney biopsy. A. Interstitial inflammation consisting of lymphoid cells that dissociate the tubular epithelium. (PAS - 400x); Renal biopsy B. Interstitial broadening by fibrosis and lymphocytic inflammatory infiltrate (Masson trichrome - 400x).
it is the result of an autoimmune process that involves cellular and humoral immunity. The renal and ocular injuries in the clinical course of TINU may suggest a common antigen to these two structures that could be the target of a crossed autoimmune reaction.

This syndrome can be differentiated from systemic conditions associated with uveitis and nephritis. TIN has a variety of etiologies and it would be no different with TINU syndrome. The role of infections and non-infectious causes is speculated in the immune response, leading to the development of the syndrome.

Kim et al. consider that the main causes of concomitant nephritis and uveitis, which can be mistaken for the TINU syndrome are: sarcoidosis, Sjogren’s syndrome, lupus, Wegener’s granulomatosis, Behcet’s disease, and it is imperative to order tests to rule these out.

In 2001, Mandeville et al. published the TINU syndrome diagnostic criteria, requiring the presence of acute interstitial nephritis and uveitis, with no other systemic disease as a cause, in addition to completing the 3 following criteria: 1. abnormal kidney function; 2. abnormal urinalysis; and 3. Systemic disease lasting ≥2 weeks.

Laboratory tests in the evaluation of TINU syndrome are not typical. Moreover, the ANCA, antinuclear antibody, anticardiolipin antibodies, anti-DNA antibody and rheumatoid factor may be positive and may be hypocomplementemia, and polyclonal hypergammaglobulinemia and circulating immune complexes.

The patient had mild anemia, elevated inflammatory markers, azotemia and increased proteinuria. C4 was high (60.8 mg/dL), and C3 dosage was normal (154 mg/dL). C-ANCA, P-ANCA and FAN were negative.

Histological evidence of TIN should be sought for the definitive diagnosis of the syndrome. However, an invasive procedure such as a kidney biopsy may not be suitable for all patients, and should be considered in a case-by-case basis.

TIN may resolve spontaneously, but the treatment of patients with progressive renal failure include the use of corticosteroids. The nephropathy becomes chronic in 11% of cases and only 5% of patients require dialysis. In this case, after pulse therapy, the patient experienced a reduction in urea and creatinine levels.

Topical steroids associated with cycloplegic agents are commonly recommended as a treatment option for anterior uveitis. If refractory, one should consider using oral steroids or immunosuppressive agents. Reports indicate that in 14% of the cases the symptoms persist for more than 3 months and uveitis recurrence occurs in about 50% of patients. In a series of 33 cases, the median value for disease persistence was 7 months, ranging from 1 to 147 months.

Although this patient of ours manifested the TINU syndrome at 60 years of age, she developed a favorable renal function outcome, presenting azotemic values within the normal range even after 52 months of follow-up. Due to the increase of intraocular pressure with use of topical corticosteroids, we decided to use cyclosporine eye drops, making continuous use without possibilities of discontinuing it, as it always resulted in uveitis recurrence.

**CONCLUSION**

This tubulointerstitial nephritis and uveitis case makes us suggest that this rare syndrome should be considered in the differential diagnosis of unexplained tubulointerstitial nephritis, especially in the presence of ocular disorders, even in older patients. There is evidence suggesting that TINU is underdiagnosed, especially in young patients who have mild and asymptomatic kidney disease. It is necessary to disclose this information among nephrologists, internists and ophthalmologists, because their interaction is paramount in the treatment of patients with the TINU syndrome.

**REFERENCES**


TINU syndrome


