Acute Tubular Necrosis Following Interferon-Based Therapy for Hepatitis C: Case Study with Literature Review

Fabrizio Fabrizi a Alessio Aghemo b G. Battista Fogazzi a Gabriella Moroni a Patrizia Passerini a Roberta D’Ambrosio b Piergiorgio Messa a

a Division of Nephrology and b Division of Digestive Diseases, Maggiore Policlinico Hospital, IRCCS Foundation, Milan, Italy

Key Words
Hepatitis C • Acute Tubular Necrosis • Interferon • Ribavirin • Nephrotoxicity

Abstract

Background/Aims: Interferon treatment of malignant or viral diseases can be accompanied by various side-effects including nephro-toxicity. Methods: We report on a 68-year-old Caucasian male who received dual therapy with pegylated interferon 2a plus ribavirin for chronic hepatitis C. Results: After three months of antiviral therapy, the patient developed acute kidney failure (serum creatinine up to 6 mg/dL) with mild proteinuria (500 mg daily) and haematuria. Immediate immunosuppressive therapy with high-dose intravenous steroids did not improve kidney function. Kidney biopsy was consistent with acute tubular necrosis without glomerular abnormalities. He started long-term peritoneal dialysis (four regular exchanges) to provide both dialysis adequacy and ascites removal. Kidney function gradually improved over the following months (serum creatinine around 2 mg/dL) and peritoneal dialysis was continued with two exchanges daily. The temporal relationship between the administration of the drug and the occurrence of nephro-toxicity, and the absence of other obvious reasons for acute tubular necrosis support a causative role for pegylated interferon; benefit on kidney disease was noted after withdrawal of antiviral agents. An extensive review of the literature on acute tubular necrosis associated with interferon-based therapy, based on in vitro data and earlier case-reports, has been made. The proposed pathogenic mechanisms are reviewed. Conclusions: Our case emphasizes the importance of monitoring renal function during treatment of chronic hepatitis C with antiviral combination therapy as treatment may precipitate kidney damage at tubular level.
Introduction

Hepatitis C virus (HCV) is a blood-borne pathogen that appears to be endemic in many countries and is the main cause of chronic liver disease worldwide. The major long-term complications of chronic HCV infection are liver fibrosis, cirrhosis, and a high risk for hepatocellular carcinoma; the development of cirrhosis in published studies in the general population has ranged from 2% to 42% [1]. Antiviral treatment with pegylated interferon (peg-IFN) and ribavirin eradicates HCV infection in many patients (around 50%), depending on host and viral characteristics, especially the viral genotype [2]. However, interferon (IFN)-based therapy is complicated by frequent and, at times, serious adverse effects which represent an important barrier to treatment delivery. Information accumulated in the last decade suggests a rate of premature discontinuation of peg-IFNα and ribavirin treatment by patients with chronic HCV infection of around 9%-12% [3]. One of the reasons for the low SVR rates is the high frequency of adverse events related to peg-IFNα and ribavirin; therefore, the adherence to antiviral treatment would have a favorable effect on the sustained virological response (SVR) rate. In fact, patients who discontinued peg-IFNα and ribavirin treatment prematurely for whatever reason had an SVR rate of 12% compared with 65% of those who continued antiviral treatment despite dose reduction [4]. Combined antiviral therapy (conventional or peg-IFN plus ribavirin) impacts most, if not all, organ systems [4]. This darker side of IFN activity has been recognized by various investigators and was summarized in an editorial by Tang, who had asked the important question 'Interferon: friend or foe?'[5].

In addition to the widely recognized side-effects, nephro-toxicity is an uncommon but potentially severe side-effect of antiviral therapy [6]. Nephro-toxicity of combined antiviral treatment appears more related to IFN than ribavirin in view of the lack of reports on side-effects with ribavirin mono-therapy [7]. We report here the case of a patient who developed acute renal insufficiency due to acute tubular necrosis (ATN) during treatment with combined antiviral treatment (pegIFN α-2b plus RBV) for chronic hepatitis C. After discontinuation of antiviral agents, he showed partial recovery of renal function and remission of urinary abnormalities. A systematic review on the renal side effects of IFN-based therapy, based on laboratory investigations data and earlier case reports, was also performed.

Case Study

A 68-year-old Caucasian male was diagnosed with hepatitis C (genotype 1b) in 2009. His past medical history included tonsillectomy, coronary artery bypass graft surgery (1984), and percutaneous transluminal coronary angioplasty (PTCA) due to myocardial infarction (1999). He underwent liver biopsy (2009) which showed steatosis, chronic active hepatitis with nodular changes and mild activity; a quantitative PCR assay for hepatitis C virus RNA (HCV RNA) gave a viral load of 589,919 IU/mL. At that time kidney function was normal with serum creatinine of 0.95 mg/dL. In November 2009, he started combination antiviral therapy with peg-IFN α2a (180 mcg/week) and ribavirin (1,200 mg daily).

Renal function was normal throughout the initial months of antiviral therapy, but acute kidney failure (serum creatinine of 1.97 mg/dL) occurred in February 2010 [Figure 1]. Antiviral therapy was immediately discontinued and the patient referred to our renal Division. On admission he was pale and apyrexial, peripheral edema and macroscopic haematuria were present. Blood pressure was normal. Serum biochemical tests were as follows: serum creatinine 5.19 mg/dL, urea 180 mg/dL, haemoglobin 8.8 g/dL; urine analysis showing proteinuria (500 mg daily) with granular and red cell casts [Table 1]. Serum cryocrit of 3% was found whereas other immunological tests (ANA, ANCA, ENA, and complement) tested negative. HCV RNA was 44 IU/mL. Liver biochemistries were as follows: GOT 32 IU/L, GPT 6 IU/L, γ-GT 40 I/L, cholinesterasis 3034 IU/L, total protein 5.8 gr/dL, and albumin 2.6 g/dL. Initially his urine output was low, and anuria occurred on the third hospital...
Acute nephritic syndrome was suspected and we immediately started intravenous corticosteroid therapy (methyl-prednisolone 500 mg and 300 mg on two consecutive days). Immunosuppressive therapy was complicated by diabetes mellitus and further deterioration of renal function was noted (serum creatinine up to 6.2 mg/dL, urea 264 mg/dL). We started intermittent haemodialysis; on the 6th hospital day, he underwent percutaneous renal biopsy showing acute tubular necrosis (five glomeruli included). The tubular damage was diffused and involved all nephron segments; damage to the proximal tubules was shown by the occurrence of coarse vacuolar degeneration, whereas the distal tubules and collecting ducts were dilated, attenuated, and partially obstructed by either hyaline or granular or red cell casts. The intracapsular space of two glomeruli was partially filled haemorrhagic substance. The interstitium surrounding the damaged tubules was edematous but without inflammatory infiltrate. Necrosis in many tubules occurred. Mild and aspecific glomerular abnormalities were present including thin glomerular walls, focal expansion of mesangial matrix, and mild segmental proliferation of mesangial cells. Immunofluorescence analysis was fully negative. The renal biopsy excluded uric acid crystals or myeloma casts causing...

### Table 1. Blood chemistries at presentation and over follow-up

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Creatinine (0.5-1.2, mg/dL)</td>
<td>1.07</td>
<td>5.19</td>
<td>1.89</td>
<td>2.04</td>
</tr>
<tr>
<td>Blood urea nitrogen (8-20, mg/dL)</td>
<td>35</td>
<td>180</td>
<td>98</td>
<td>112</td>
</tr>
<tr>
<td>AST (5-38, IU/L)</td>
<td>62</td>
<td>32</td>
<td>84</td>
<td>43</td>
</tr>
<tr>
<td>ALT (5-41, IU/L)</td>
<td>98</td>
<td>6</td>
<td>106</td>
<td>65</td>
</tr>
<tr>
<td>γGT (8-61, IU/L)</td>
<td>71</td>
<td>40</td>
<td>225</td>
<td>86</td>
</tr>
<tr>
<td>Cholinesterase (5,300-12,900, IU/L)</td>
<td>9.392</td>
<td>8.503</td>
<td>13.113</td>
<td>8.204</td>
</tr>
<tr>
<td>Total bilirubin (0.1-1.1, mg/dL)</td>
<td>0.85</td>
<td>1.2</td>
<td>0.8</td>
<td>0.58</td>
</tr>
<tr>
<td>Uric acid, mg/dL</td>
<td>6.2</td>
<td>5.3</td>
<td>4.6</td>
<td>6.1</td>
</tr>
<tr>
<td>Total protein (6.6-8.7, g/dL)</td>
<td>7.6</td>
<td>7.3</td>
<td>5.9</td>
<td>6.3</td>
</tr>
<tr>
<td>Albumin (3.4-4.8, g/dL)</td>
<td>4.0</td>
<td>3.8</td>
<td>3.3</td>
<td>3.3</td>
</tr>
<tr>
<td>Total cholesterol (&lt;200, mg/dL)</td>
<td>159</td>
<td>151</td>
<td>173</td>
<td>173</td>
</tr>
<tr>
<td>Triglycerides (&lt;170, mg/dL)</td>
<td>117</td>
<td>75</td>
<td>111</td>
<td>111</td>
</tr>
<tr>
<td>IgA (70-400, mg/dL)</td>
<td>360</td>
<td>343</td>
<td>320</td>
<td>405</td>
</tr>
<tr>
<td>IgG (700-1600, mg/dL)</td>
<td>1.758</td>
<td>1.087</td>
<td>1.122</td>
<td>915</td>
</tr>
<tr>
<td>IgM (40-230, mg/dL)</td>
<td>185</td>
<td>119</td>
<td>119</td>
<td>192</td>
</tr>
<tr>
<td>C3 (90-180, mg/dL)</td>
<td>106</td>
<td>99</td>
<td>102</td>
<td>111</td>
</tr>
<tr>
<td>C4 (10-40, mg/dL)</td>
<td>22</td>
<td>21</td>
<td>32</td>
<td>24</td>
</tr>
<tr>
<td>Rheumatoid factor (&lt;18, IU/mL)</td>
<td>7</td>
<td>9</td>
<td>12</td>
<td>12</td>
</tr>
<tr>
<td>PTH (15-65, pg/mL)</td>
<td>56.7</td>
<td>77.8</td>
<td>94</td>
<td>162</td>
</tr>
<tr>
<td>HCV RNA (IU/mL)</td>
<td>392.091</td>
<td>44</td>
<td>42.230</td>
<td>8.723</td>
</tr>
</tbody>
</table>

Abbreviations: AST, aspartate aminotransferase; ALT, alanine aminotransferase; γGT, glutamyl transpeptidase; HCV RNA, hepatitis C virus viraemia; PTH, parathormone.
tubular obstruction; the chronology of exposure to nephro-toxins such as non-steroidal anti-inflammatory agents or radio-contrast was not appropriate to implicate these etiologies to renal failure. No drug rashes or eosinophilia was detected. Abdominal computed tomography (CT) scan demonstrated abundant peri-hepatic and peri-splenic ascites, kidneys with regular size and increased echogenicity and no evidence of a stone or an obstruction; blood flow was regular without vascular lesions in the kidney and liver. Tenckhoff peritoneal catheter was implanted in March (2010) and the patient started peritoneal dialysis with the aim to provide both dialysis adequacy and ascites removal (four regular exchanges throughout the day). He was discharged with serum creatinine of 4.03 mg/dL and urea 107 mg/dL.

He continued peritoneal dialysis at home (continuous ambulatory peritoneal dialysis-CAPD) with normal blood pressure, and stable body weight. Urine output resumed on the following weeks and renal function gradually improved. At routine follow-up 96 days after the onset of acute renal failure, the patient’s creatinine level was 2.1 mg per deciliter. CAPD was continued with two exchanges daily.

Two months later, he was again in charge at the Hospital for mental confusion with altered level of consciousness- diagnosis of hepatic encephalopathy was made, and medical therapy with oral lactulose was initiated. He did well until February 2013 when he was again hospitalized at our Hospital due to symptomatic pneumonia; intravenous antibiotics were given with benefit. Important anemia was also observed (ten units of red blood cells administered). Gastrointestinal (GI) endoscopy showed esophageal varices without signs of active blood loss, and hemorrhoids were found by colonoscopy. Chest x ray showed a right pleural effusion; thus, a connection between pleural and peritoneal cavities was suspected and dialysis was stopped. Fourteen days later peritoneal dialysis was started again with serum creatinine and urea of 1.9 mg/dL and 132 mg/dL, respectively; body weight increased of around 5 Kg with edema refractory to oral diuretics. He received intravenous iron and folic acid supplementation, and the dose of subcutaneous erythropoietin was increased. Biochemical liver tests were stable (Child-Pugh B7), HCV RNA 8,723 IU/mL. Normal blood pressure was again recorded. Therapy with insulin was interrupted and oral medications were started with good control of blood sugar levels. His medications currently include: omeprazole 20 mg daily, folic acid 5 mg daily, lactulose 10 g daily, calcitriol 0.5 mcg three times weekly, cardioaspirin 100 mg daily, furosemide 250 mg twice daily, potassium canrenone 100 mg daily, darbepoetin alpha 60 mcg sc weekly, bisoprolol 2.5 mg twice daily, iron sulphate 329 mg daily, and repaglinide 0.5 mg three times daily.

Review of the literature

In 1981, purified human leukocyte interferon (interferon-α) became available for clinical use [6]. IFNs are a family of natural cytokines provided with ability to interfere with viral replication, cell proliferation, and immuno-regulation. Three types of IFNs (α, β, and γ) exist, according to the major IFN peaks observed under high performance liquid chromatography (HPLC). The major cells of origin for α, β, and γ IFNs are leukocytes (other than lymphocytes), fibroblasts, and T cells and natural killer cells, respectively. Many other cell types are also capable of producing lower levels of IFN-α, and -β when virally infected. Type I IFNs include IFNα, β, and ω species. IFNγ activates different intracellular signaling events by the use of a distinct set of receptors, namely type II IFN receptors. Recombinant DNA technology has allowed cloning and mass production of IFNs for therapeutic use: IFNα which include IFNα-2a and IFNα-2b is currently approved for the treatment of multiple medical conditions such as hepatitis B and C, and some types of malignant diseases (malignant melanoma, chronic myeloid leukaemia, hairy cell leukaemia, mycosis fungoides, and non-Hodgkin follicular lymphoma) [8]. IFN-β is recommended for relapsing multiple sclerosis (MS) [9]. IFN-γ is indicated for the treatment of severe malignant osteopetrosis [10].

Although IFNs have been shown to be of benefit in many clinical situations, many retrograde aspects to IFNs therapy have been emphasized- acute toxicity (influenza-like
symptoms and fever), subacute and chronic toxicity (fatigue, anorexia, and general wasting) have been attributed to IFN-based therapy. Complications associated with the administration of IFNs can affect nervous (depression, lethargy, somnolence, mental slowing), hematopoietic (leukopenia, anemia, and thrombocytopenia), cardiovascular (myocardial infarction, arrhythmias, hypotension), musculoskeletal (myalgias, arthralgies), and endocrine system (thyroid dysfunction). Gastrointestinal toxicity consists mostly of anorexia, vomiting, diarrhea, and nausea.

Renal function is usually well preserved during IFN use and IFN-induced renal toxicity is considered quite unusual, although some evidence in the scientific literature exists. An important adverse effect of IFN on kidney is graft rejection after kidney transplantation—at least 12 published studies, although uncontrolled retrospective or observational studies, have shown kidney graft dysfunction during IFN-based therapy for chronic hepatitis B or C [11]. Reported rates of kidney graft dysfunction range from 9% to 100%, with most episodes occurring between 1 and 8 months after initiation of the therapy. In most cases, IFN-induced acute kidney injury appears to be predominantly related to its immuno-modulatory properties, leading to increased rates of both cell- and antibody-mediated rejection; acute rejection induced by IFN being frequently insensitive to corticosteroids and irreversible. On the other hand, the development of HCV-related fibrosing cholestatic hepatitis (FCH) may be an indication for IFN use after kidney transplantation as FCH has an ominous course. In these subjects, IFN-based therapy may be potentially lifesaving and should be given despite the risk of kidney graft rejection and graft loss. For HCV-infected kidney transplant recipients in whom the benefits of antiviral treatment clearly outweigh the risks, mono-therapy with standard IFN has been recommended, according to the Kidney Disease Improving Global Outcomes (KDIGO) guidelines. In any circumstance, this course of therapy should be undertaken only after the patient has been properly informed of the risks of both treatment and electing not to treat [11].

The most common adverse effect on the kidney during treatment of immuno-competent individuals with IFN is proteinuria. Early reports from phase II and III clinical trials on the use of IFN-α have documented the occurrence of proteinuria (15% to 25%) and abnormal urinalysis (14%) [6]. In many instances, these abnormalities were mild and reversible after stopping IFN. A variety of glomerular, interstitial, and tubular abnormalities sometimes seen in the same biopsy specimen, have been ascribed to IFN-associated nephro-toxicity. Some of these lesions have occurred after short-term (<16 weeks) administration of IFNs, and others with more chronic use (8 months to 6 years). In many reports, discontinuation of IFNs led to improvement of kidney disease.

Discussion

This report describes a case of acute renal failure by acute tubular necrosis during antiviral therapy (peg-IFNα plus ribavirin) for chronic hepatitis C. The chronology of progressive rise in serum creatinine a few weeks after the initiation of antiviral therapy supports a possible relationship between cytokine therapy and deterioration of renal function. The improvement of renal function after discontinuation of peg-IFNα and the absence of other evident causes of ATN strongly suggest a link between ATN and antiviral therapy with peg-IFNα. In fact, there was no increase in plasma uric acid concentration suggestive of a tumour lysis syndrome, no underlying infection or hypotension, no use of radio-contrast or other medications, e.g. antibiotics or non-steroidal anti-inflammatory drugs, to account for the tubular necrosis.

IFN-α therapy has been implicated in various renal diseases such as renal thrombotic microangiopathy (in patients with chronic leukemia) [12], interstitial nephritis [13], and glomerular diseases including podocytopathies (focal segmental glomerulosclerosis and minimal change disease) or immune complex glomerulonephritis (membranous nephropathy, membranoproliferative glomerulonephritis, IgA nephropathy), as shown in
several case reports published over the last 20 years [14-30]. In addition, case series have been published [31]. Additional papers provided evidence of exacerbation of pre-existing glomerular disease during IFN [32-33]. The usual presentation of glomerular disease associated with IFN is isolated proteinuria, and nephrotic syndrome [14-31]. In a minority of patients, nephritic syndrome with renal failure has been recorded. Conversely, renal toxicity with acute renal failure due to ATN appears less frequent and it had been previously reported in a few patients [34-39]. As listed in Table 2, these reports concerned mostly the treatment of haematological or solid neoplastic disorders, with dosages of interferon frequently higher than in the current case. Three other case reports on IFN-associated renal failure described focal ATN in kidney biopsy in association with glomerular abnormalities [14, 30, 40].

How IFN therapy causes renal injury at tubular level is unknown, even if this is an area of intense research. An in vitro cell culture model system has been established to investigate the effects of IFNa on renal epithelial cells. It has been found a reversible, dose- and time-dependent decrease of trans-epithelial resistance (TER) of renal proximal tubular monolayer cells (LLC-PKI porcine proximal tubular cells). IFN can directly affect the epithelial barrier function of renal proximal tubular cells, increasing the leakiness of the epithelial layer after IFNa treatment. ‘Backleak of glomerular filtrate into the renal interstitium might thus be induced, generating edema with inflammatory changes; these in vitro observations could explain the severe and in part life threatening adverse effects of IFNa, such as renal dysfunction or the capillary leak syndrome [41-42]. A novel and independent action of IFNa on renal tubular epithelium is the induction of apoptosis in renal tubular epithelial cells; IFNa supports the activation of caspase-3, -8 and -9, DNA fragmentation, and nuclear condensation. IFNa also caused mitochondrial depolarization. In other words, IFNa induced apoptosis is directed by an extrinsic death receptor signaling pathway, amplified by an intrinsic mitochondrial pathway. The apoptotic signaling pathways activated in proximal tubular cells resemble the pathways induced by IFNa in melanoma and bladder carcinoma cells [43].

Recently, Satou et al [44] have observed in animal models (rodents) that interferon γ (IFNy) induces activation of the intrarenal renin-angiotensin system (RAS) by modulating

---

**Table 2. Acute renal failure (due to ATN) associated with IFN therapy**

<table>
<thead>
<tr>
<th>Publication year</th>
<th>Reference number</th>
<th>Country</th>
<th>IFN type</th>
<th>IFN dose</th>
<th>IFN duration</th>
<th>IFN route</th>
<th>Gender</th>
<th>Age, yrs</th>
<th>Race</th>
<th>Renal presentation</th>
<th>Creatinine (mg/dL)</th>
<th>Original disease</th>
<th>Outcome after IFN withdrawal</th>
<th>Duration of follow-up</th>
<th>Kidney biopsy</th>
</tr>
</thead>
<tbody>
<tr>
<td>1988</td>
<td>34</td>
<td>US</td>
<td>IFN-γ</td>
<td>1x10⁷ U/</td>
<td>19 days</td>
<td>sc</td>
<td>Male</td>
<td>9</td>
<td>NA</td>
<td>ARF</td>
<td>14.3</td>
<td>Lymphoblastic leukemia</td>
<td>Complete recovery</td>
<td>33 days</td>
<td>ATN, FSGS</td>
</tr>
<tr>
<td>1993</td>
<td>35</td>
<td>UK</td>
<td>IFN-α</td>
<td>5 MUIx3/</td>
<td>10 mo</td>
<td>sc</td>
<td>Female</td>
<td>49</td>
<td>NA</td>
<td>ARF</td>
<td>11.5</td>
<td>Multiple myeloma</td>
<td>Complete recovery</td>
<td>NA</td>
<td>ATN</td>
</tr>
<tr>
<td>1995</td>
<td>36</td>
<td>Spain</td>
<td>IFN-α</td>
<td>5 MUIx3/</td>
<td>5 days</td>
<td>sc</td>
<td>NA</td>
<td>26</td>
<td>NA</td>
<td>ARF</td>
<td>21.7</td>
<td>Hepatitis B</td>
<td>Complete recovery</td>
<td>25 days</td>
<td>ATN</td>
</tr>
<tr>
<td>1997</td>
<td>37</td>
<td>France</td>
<td>IFN-α</td>
<td>3 MUIx/</td>
<td>42 days</td>
<td>sc</td>
<td>Male</td>
<td>42</td>
<td>NA</td>
<td>ARF</td>
<td>5.5</td>
<td>Hepatitis A</td>
<td>Complete recovery</td>
<td>90 days</td>
<td>ATN</td>
</tr>
<tr>
<td>1998</td>
<td>38</td>
<td>Saudi Arabia</td>
<td>IFN-α</td>
<td>3 MUIx/</td>
<td>10 days</td>
<td>sc</td>
<td>Male</td>
<td>30</td>
<td>NA</td>
<td>ARF</td>
<td>13.4</td>
<td>HBV-related GN</td>
<td>Complete recovery</td>
<td>NA</td>
<td>ATN</td>
</tr>
<tr>
<td>2004</td>
<td>39</td>
<td>Australia</td>
<td>Peg-IFN-a2a</td>
<td>180 mcg/</td>
<td>10 days</td>
<td>sc</td>
<td>Male</td>
<td>54</td>
<td>NA</td>
<td>ARF</td>
<td>2.7</td>
<td>Hepatitis C</td>
<td>Complete recovery</td>
<td>80 days</td>
<td>ATN, IgA nephropathy</td>
</tr>
</tbody>
</table>

Abbreviations: ATN, acute tubular necrosis; ARF, acute renal failure; FSGS, focal segmental glomerulosclerosis; IFN, interferon; GN, glomerulonephritis; NA, not available; NS, nephrotic syndrome; sc, subcutaneously
the STAT1-SOCS1 axis. In the kidney, intra-renal angiotensin II (Ang II) is derived from intra-renal angiotensinogen (AGT) which is produced mainly in renal proximal tubular cells (RPCTs). IFNγ augments AGT expression in RPCTs which may lead to increases in intra-renal RAS activity. The local RAS functions are expressed in individual organs in a tissue-specific manner; intra-renal AGT is increased in several forms of clinical and experimental hypertension and exacerbation of renal inflammation. These in vitro data can contribute to explain the tubular injury associated with IFN administration in humans.

An alternative explanation implicates an apoptotic activity of ribavirin, that has been recently established in in vitro settings [45]. Ribavirin proved to be a strong induced of apoptosis using various human cancer cell lines, and these findings are in keeping with the rationale of the initiation of the ribavirin treatment trial that is in progress on acute myeloid leukemia (AML) patients. It remains to assess whether the apoptotic activity of ribavirin (or its metabolites) is evident in the renal tubular cells and plays a role in the exacerbation of tubular necrosis instigated by IFN.

**Conclusion**

This case report gives emphasis to the renal toxicity of IFN-based therapy at tubular level. Because our patient had no other obvious reason for ATN, we postulate that it may be secondary to peg-IFN plus ribavirin therapy. Further studies are in progress to understand better pathogenesis, outcome and management of IFN-associated NTA. Physicians treating chronic hepatitis C patients with IFNs should be advised to be vigilant and be aware of these potential renal complications.

**Conflict of Interests**

None of the authors have any conflict of interest to report.

**Acknowledgements**

This work was supported in part by the grant ‘Project Glomerulonephritis’; in memory of Pippo Neglia.

**References**


