An Unusual Case of Cholestatic Hepatitis due to Light-Chain Deposition Disease

Alessandro Grembiale a, Elena Garlatti a, Anna Ermacora a, Silvia Grazioli a, Massimiliano Balbi b, Maurizio Tonizzo a

aDepartment of Internal Medicine, ASFO – Pordenone, Pordenone, Italy; bDepartment of Internal Medicine, ASFO – San Vito al Tagliamento (PN), San Vito al Tagliamento, Italy

Keywords
Light-chain deposition disease · Monoclonal light chains · Amyloidosis · Cholestatic hepatitis · Bortezomib

Abstract
Light-chain deposition disease (LCDD) is a rare paraproteinaemia characterized by the deposition of monoclonal immunoglobulins with a non-fibrillar structure and hence Congo red negative deposits. Kidney disease is the more frequent manifestation, but other organs may also be involved. A 70-year-old man with hypertension and mild chronic renal failure showed a hepatomegaly without splenomegaly. His renal and liver test rapidly got worse. A serum electrophoresis and immunofixation isolated monoclonal kappa light-chain gammopathy, with serum free kappa light chain excess. The bone marrow biopsy showed the presence of interstitial infiltration of plasma cells like multiple myeloma type at initial phase. Periumbilical fat biopsy was negative. Echocardiography demonstrated an infiltrative cardiac disease. The biopsies of the duodenum small intestine mucosa showed flaps with eosinophil material (Mason’s staining) with atrophic crypts and chronic inflammation at chorion level. Amyloid substance was negative. There was a strong positivity for light chains kappa compatible with LCDD. A liver biopsy confirmed this finding. Therapy with dexamethasone and bortezomib improved clinical state and hepatic and renal laboratory tests. Chemotherapy based on novel anti-myeloma agents should be rapidly considered in LCDD patients with severe organ involvement.

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Published by S. Karger AG, Basel
Introduction

Light-chain deposition disease (LCDD), heavy-chain deposition disease, and light- and heavy-chain deposition disease are a rare group of paraproteinaemias characterized by the deposition of monoclonal immunoglobulins with a non-fibrillar structure and hence Congo red negative deposits [1].

The diagnosis of LCDD requires histological demonstration of monotypic light-chain (LC) deposition on immunofluorescence microscopy and ultrastructural analysis of the involved organs or tissues. LCDD can occur in the context of isolated monoclonal gammapathy or of symptomatic multiple myeloma and Waldenström’s macroglobulinemia. Light chain deposits are usually the κ (kappa) isotype and can affect almost all organs [2]. Kidney disease is the more frequent manifestation, resulting in chronic kidney failure with glomerular proteinuria, and sometimes nephrotic syndrome [3] but heart, liver [4], gastrointestinal tract, and peripheral nerves may also be involved. Liver involvement has been rarely reported in LCDD in asymptomatic patients, but in symptomatic ones, LCDD-associated liver involvement mainly manifests as cholestatic hepatitis and is associated with high mortality [5].

We report in this paper a patient with myeloma-associated LCDD who developed rapidly progressive liver and renal failure secondary to κ-light chain deposition, which rapidly recovered after chemotherapy. Patient has given his written informed consent to publish his case.

Case Report

A 70-year-old man with hypertension, kidney stones disease and mild chronic renal failure was admitted to our department with asthenia and sudden weight loss. Physical examinations showed hepatomegaly without splenomegaly. A liver ultrasound confirmed hepatomegaly with mild hepatic steatosis and a non-homogeneous echostructure with a starry sky appearance. There was no evidence of biliary obstruction, and the kidneys had a normal size without urinary tract obstruction. There was liver stiffness (Fibroscan©: 53.3 kPa with IRQ 18).

Blood tests showed serum creatinine: 2.3 mg/dL, ESR: 120 mm/h, γGT: 2003 IU/L, P-ALC: 732 IU/L, fibrinogen: 700 mg/dL, presence of monoclonal component IgA k: 14 g/L. Baseline liver tests, serum calcium, and blood coagulation parameters were normal. There was no history of alcohol abuse. Serological tests for hepatitis A, B and C, Epstein-Barr virus, cytomegalic virus, and herpes simplex virus were negative.

A few weeks later, renal and liver test rapidly got worse (serum creatinine: 6.7 mg/dL, total bilirubin: 4.8 mg/dL, direct bilirubin: 3.9 mg/dL, AST: 647 U/L, ALT: 485 U/L, LDH: 780 U/L), probably due to a concomitant septic state.

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A serum electrophoresis and immunofixation isolated monoclonal kappa LC gammapathy, with serum free kappa light chain excess of 47 mg/L, with a kappa/lambda ratio of 2.76. 24-h proteinuria was 1.71 g. Bence-Jones proteinuria was negative.

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The whole body radiological evaluation did not demonstrate osteolytic lesions. The bone marrow biopsy showed the presence of interstitial infiltration (between 10 and 20%) of plasma cells like a plasmacellular dyscrasia preferentially multiple myeloma type at initial phase.

Moreover, we performed a periumbilical fat biopsy that was negative for the staining with the Congo red, and there were no aspects related to amyloid deposits.

Transthoracic echocardiography demonstrated moderate hypertrophic cardiomyopathy (no pulmonary hypertension), with mainly septal evidence of infiltrative cardiac disease (left
ventricle ejection fraction 60%) and organized pericarditis adherent to the right ventricle (thickness 14 mm), without signs of compression on the cardiac chambers.

The patient also underwent gastroscopy, and the biopsies of the duodenum small intestine mucosa showed flaps with eosinophil material at the level of the lamina propria (Masson’s staining) with atrophic crypts and chronic inflammation at the chorion level (Fig. 1 a, 1. b, 1. c). The Congo red staining for the research of amyloid substance was negative. The search for amyloid A and P was negative; but there was a strong positivity for light chains kappa +++ on immunohistochemistry, preferentially compatible with LCDD.

A liver biopsy was also performed which confirmed the presence of amorphous eosinophilic deposits at the sinusoidal level associated with atrophy moderate hepatic parenchyma (Fig. 2 a, 2. b, 2. c). The substance was Congo red negative, kappa +++ light chains, PAS−.
We concluded that it was LCDD with hepatic, gastrointestinal, probably renal and cardiac involvement, associated with IgA K myeloma. We started therapy with dexamethasone and bortezomib with improvement of hepatic and renal function laboratory tests.

We observed a progressive and complete recovery of cholestatic and cytolytic hepatitis over the next 6 months with mild improvement of renal failure.

**Discussion**

LCDD is a rare disease of non-fibrillar deposition of monoclonal light-chain immunoglobulins, usually manifesting in the fifth or sixth decade of life with a male predominance, but the incidence is unknown [6]. In LCDD, the monoclonal light chains are of the kappa subtype in 92% of cases, and it is typically associated with multiple myeloma or other lymphoproliferative disorders [6].

LCDD occurs in about 5% of patients with multiple myeloma, typically reported as the underlying disease in these patients [7, 8]. However, recent reports indicate that a high proportion of LCDD cases are now found in patients without symptomatic plasma cell disorder, who meet the new criteria for monoclonal gammopathy of renal significance [9]. Less frequently, LCDD may occur in the context of lymphoid hemopathy such as Waldenström disease. Unlike AL amyloidosis, LC deposits in LCDD, mostly the kappa isotype, do not stain with Congo red, and show a typical histological linear pattern along basement membranes on immunofluorescence microscopy, with a characteristic linear powdery punctate appearance on electron microscopy [10, 11].

Based on histological and postmortem studies, liver involvement has been considered to be frequent in LCDD, but severe liver complications have been rarely reported [12]. Principal presentation was intrahepatic cholestasis, associated with cytolysis in 33% of cases. The most frequent histological finding was linear deposits of monoclonal LC in the sinusoidal or perisinusoidal spaces with a predominance of kappa isotype, as we reported.

A cardiac origin was also unlikely to account for liver failure, given the absence of right ventricular heart failure at echocardiography. The transient septic state may have contributed to worsening liver and kidney function, but its resolution and the liver biopsy results undoubtedly showed evidence of LC deposition. We finally attributed liver and kidney failure as a specific manifestation of the disease in our patient.

Steroids and melphalan, high-dose melphalan, autologous stem cell transplantation and, more recently, bortezomib-based chemotherapy, are some of the treatment options [13]. Chemotherapeutic regimens containing the proteasome inhibitor bortezomib have shown efficacy and good tolerance profile in both AL amyloidosis [14] and Randall-type MIDD, but their effect on the clearance of LC tissular deposits, although possibly accepted, remains to be demonstrated. Another study reported the disappearance of nodular mesangial lesions and disappearance of kappa light chain deposits in a patient with kidney LCDD under long-term chemotherapy [15].

**Conclusion**

These data indicate that chemotherapy, based on novel anti-myeloma agents, should be rapidly considered in LCDD patients with severe organ involvement, in order to rapidly induce efficient and sustained suppression of the pathogenic monoclonal LC.

Our case confirms that liver involvement may be an important complication that requires prompt diagnosis and therapy.
Statement of Ethics

The patient has given his written informed consent to publish his case including publication of images.

Conflict of Interest Statement

The authors have no conflicts of interest to declare.

Funding Sources

None.

Author Contributions

A. Grembiale: acquisition, analysis, and interpretation of data for the work; drafting the work or revising it critically for important intellectual content; final approval of the version to be published. E. Garlatti, A. Ermacora: substantial contributions to the conception and interpretation of data for the work; drafting the work or revising it critically for important intellectual content, final approval of the version to be published. S. Grazioli, M. Balbi, M. Tonizzo: revising the work critically for important intellectual content and final approval of the version to be published.

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