RENAL INVOLVEMENT IN MONOCLONAL GAMMOPATHIES

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Morphologic characteristics

AL amyloidosis

11.6±8.6

0%

1.1±1

5.5% (4)

26% (19)

30% (22)

13.7% (10)

30% (22)

30% (22)

33% (24)

19% (14)

18% (13)

41.1% (30)

Mveloma

kidney

8.7±5.3

1.4% (1)

1±1.9

100%

2.9% (2)

4.3% (3)

27.5% (19)

65% (45)

4.3% (3)

4.3% (3)

35% (24)

56.5% (39)

20.3% (14)

LCDD

9.5+4.9

50% (10)

1±1.5

50% (10)

0%

20% (4)

40% (8)

40% (8)

5% (1)

20% (4)

35% (7)

40% (8)

15% (3)

INTRODUCTION AND AIMS

Monoclonal gammopathies (MG) are a group of disorders characterized by the proliferation of a clone of plasma cells, frequently associated with organ dysfunction.

The most common MG are MG of undetermined significance (MGUS), smoldering multiple myeloma, multiple myeloma (MM), light-chain amyloidosis (AL) and monoclonal immunoglobulin deposition disease (light chain and heavy chain deposition disease).

The aim of this study was to evaluate the histological findings of kidney biopsies and clinical data in patients with MG, with or without MM criteria

Clinical characteristics

	AL amyloidosis (n=73)	Myeloma kidney (n=69)	LCDD (n=20)
Male gender (%)	42.5% (31)	42% (29)	55% (11)
Age (years)	64.6±10	66±9.7	60.2±10.5
Caucasian race (%)	90.9% (40/44)	100% (30/30)	100% (11/11)
Dialysis requirement (%)	6.3% (4/63)	90% (36/40)	56.3% (9/16)
Presence of known MM (%)	12.3% (9)	39.1% (27)	25% (5)
Presence of known MG (%)	27.4% (20)	62.3% (43)	40% (8)
Scr (mg/dl)	1.9±1.5	6.5±3.5	5.9±6
Proteinuria (g/24h)	7.3±5.8	2.4±4.4	5.2±3.2
Hematuria (%)	45.9% (17/37)	58.3% (14/24)	100% (5/5)
Reason for kidney biopsy (%)			
Nephrotic syndrome	68.5% (50)	5.8% (4)	30% (6)
Nephritic syndrome	4.1% (3)	2.9% (2)	0%
Non nephrotic proteinuria	9.6% (7)	4.3% (3)	10% (2)
Acute kidney injury	4.1% (3)	43.5% (30)	20% (4)
Rapidly progressive renal failure	2.7% (2)	26.1% (18)	30% (6)
Hematoproteinuria with/out CKD	4.1% (3)	15.9% (11)	10% (2)
Light chain classe	λ 82% (60)	Λ 27; k 25	K 55% (11)
Heavy chain classe (n)			
No	14	27	14
lgG / lgA / lgM	16/7/1/0	11/6/0/1	1/0/2/0

PATIENTS AND METHODS

This was a retrospective review of all kidney biopsies (KB) performed in our laboratory in patients with MG in the last 30 vears

Clinical data: age, gender, race, associated diseases, presence of MM criteria, reason for biopsy, dialysis requirement

Laboratorial data: serum creatinine (Scr), 24h proteinuria, presence of hematuria, serum and/or urinary monoclonal protein, light chain classe

Morphological data: type of renal disease, interstitial infiltrate, interstitial fibrosis and tubular atrophy degree, vessels status, immunofluorescence results

Mean glomeruli number

Mean lobulated glomeruli

Mean sclerosed glomeruli

Myeloma kidney (%)

Interstitial fibrosis

No

Mild

Moderate

Severe Tubular atrophy No

Mild

Moderate Severe

Immunofluorescence Other than k / λ



LCDD – Light chain deposition disease; HCDD – heavy chain deposition disease TI nephrities – tubulointerstitial nephrities; IgA neph – IgA nephropathy; ATN - acute tubular necrosis; MP GN - membranoproliferative glomerulonephrities; FSGS – focal and segmental glomerulosclerosis; Malignant HT – malignant hypertension DN – diabetic nephropathy

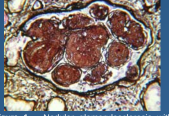
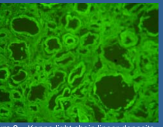


Figure 1 - Nodular glomerulosclerosis with thickness of glomerular and tubular basement membranes (Methenamine Silver x 400)



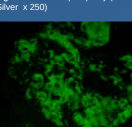


Figure 3 - Kappa light chain linear deposits along the tubular basement membrane (IF x250) Myeloma kidney - 83 ✓ 69 isolated myeloma kidney ✓ 2 with ATN ✓ 1 with minimal change disease ✓ 1 with crescentic glomerulonephr ✓ 10 associated with LCDD ✓ 4 associated with AL amyloidos View metadata, citation and similar papers at core.ac.uk

Of these patients, only

✓ 49.2% were known to have a M

✓ 24.5% were known to have MM

Figure 2 - Cast nephropathy (Mether Silver x 250)

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CORE

CONCLUSIONS

MG may be associated with a variety of renal disorders, some usually related to monoclonal chain, but others not typically related. The majority of patients with myeloma kidney didn't have a previous MM diagnosis and the majority of patients submitted to renal biopsy didn't have a MG previously identified. Although the reason for KB was different in the 3 major lesions identified, some patients with AL amyloidosis presented also myeloma kidney, as did half of the patients with LCDD. Therefore, KB constitutes the gold standard to establish diagnosis and ascertain therapeutic in patients with MG and suspicion of renal involvement.

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