Value of antibodies to free light chains in immunoperoxidase studies of renal biopsies

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ABSTRACT

Aims Because immunoglobulin abnormalities may affect the kidney, investigation of renal biopsies requires immunohistological study of light chains. A problem is that most antibodies to light chains react with whole immunoglobulins as well as free light chains, and there are generally many more whole immunoglobulins than free light chains. The usefulness of antibodies that only detected free light chains was investigated.

Methods Antibodies to free light chains were used in an immunoperoxidase method on paraffin sections of 198 renal biopsies, and compared with conventional antibodies against light chains examined by immunofluorescence on 13 frozen sections and by immunoperoxidase on 46 paraffin sections.

Results Immunofluorescence and immunoperoxidase were concordant on 10 of 13 biopsies. Immunofluorescence detected slight deposition of light chains in three biopsies not shown by immunoperoxidase, of undetermined clinical significance. Using immunoperoxidase, the free light chain antibodies were more sensitive than conventional antibodies, giving much cleaner staining and better detection of deposits in AL amyloid, light chain deposition disease and cryoglobulinaemic glomerulonephritis. The free light chain antibodies showed discordance or ambiguity between immunohistological and clinical findings in seven (4%) of 185 patients with known immunoglobulin status. These included two of 28 cases of AL amyloid that showed no light chain deposition. The method was not designed for detection of light chain restriction in neoplastic plasma or lymphoplasmacytic cells.

Conclusions Polyclonal antibodies to free light chains are an improvement on conventional antibodies in immunohistological study of paraffin sections of renal biopsies and are useful in everyday practice.

INTRODUCTION

Abnormal immunoglobulin light chains may have various effects on the kidney, especially cast nephropathy, AL amyloidosis or deposition disease, although they may have no detectable effects.1–4 Any study of light chains in renal biopsies, by immunofluorescence on frozen or paraffin sections or immunoperoxidase on paraffin sections, has disadvantages, including the specially collected material required for frozen sections, impermanence of fluorescence, and pretreatment usually needed for paraffin sections, but most antibodies give comparable findings between frozen and paraffin sections.5–9 A problem is the specificity of antibodies against light chains. Most polyclonal antibodies react with light chains both free and bound to heavy chains, but usually react preferentially with whole immunoglobulins, because normal mean plasma concentrations of free light chains and whole immunoglobulins are about 23 mg/l10 and 19 g/L11 one thousand times different. Monoclonal antibodies may be specific for a free light chain, but the epitope recognised may not be expressed or accessible in all light chains.12

Polyclonal antibodies specific for free light chains are available,10 with applications in measurement of plasma light chains and diagnosis and management of clinical immunoglobulin disorders.13 Our study assessed the value of these antibodies in immunohistology of paraffin sections of renal biopsies, including comparison with conventional antibodies.

MATERIALS AND METHODS

Antibodies

Sheep antibodies to human free κ and λ light chains were supplied by The Binding Site, Birmingham, UK. Rabbit antibodies to κ and λ (A0191, A0193), mouse antibody to serum amyloid A (M0759) and fluorescein-conjugated antibodies to κ and λ (F0198, F0199) were obtained from Dako.

Immunostaining

For immunoperoxidase, endogenous peroxidase was blocked on dewaxed 3 μm paraffin sections by hydrogen peroxide. After preliminary experiments, optimum antigen retrieval for sheep antibodies was digestion with protease type 24 (Sigma, P8038), 0.05% w/v in phosphate buffered saline pH 7.2, at 37°C, initially for 45 min, with microscopic study of completeness of digestion and further digestion if necessary.5 Antibodies were applied at 1:400 (κ) or 1:200 (λ) for 45 min, then rabbit anti-sheep immunoglobulins, peroxidase-conjugated (Dako, P0163), at 1:100, for 45 min, followed by diamino-benzidine and hydrogen peroxide, and haematoxylin. Various methods of antigen retrieval for rabbit antibodies were tried, including no pretreatment, protease digestion, trypsin digestion, microwave heating, pressure cooking, and heating for 30 min in antigen retrieval solution pH 9.0 provided in a kit for an automated method (Bondmax: Leica Microsystems, Milton Keynes, UK). Most pretreatments matched automated antigen retrieval, but protease abolished immunostaining and no pretreatment gave weak immunostaining. Antibodies were applied at 1:20 000, followed by Bondmax anti-rabbit Poly-HRP-IgG, peroxidase substrate and counterstain. Amyloid A antibody (1:500) was used without pretreatment followed by Bondmax antimouse antibody. Frozen sections were covered with...
fluoresceinated antibodies and examined by fluorescence microscopy.

Thirty-two patients were referred to the National Amyloidosis Centre (NAC). Their renal biopsies were independently examined with the rabbit antibodies to light chains without antigen retrieval, and with antibodies to amyloid A and if necessary to other potential amyloid precursors.14

Renal biopsies
Thirteen biopsies examined by immunofluorescence on frozen sections were studied using the free light chain antibodies on paraffin sections, as were another 185 paraffin-embedded biopsies. These included 46 biopsies used to compare the free light chain and rabbit antibodies, 32 at NAC and 14 others. Immunostained paraffin sections were examined without knowledge of clinical features or original reports of light chain findings. Diagnoses were made in usual ways.15 Conventional summary statistics (sensitivity, specificity and CIs of differences in proportions) were calculated.

Evidence of immunoglobulin disorders
Apart from three cases studied by immunofluorescence, all others had evidence for or against a disorder of immunoglobulins. Evidence was any of these: a serum paraprotein; a cryoglobulin; an abnormal ratio of serum free κ to λ concentrations; or a monoclonal (Bence Jones) light chain in urine. The Results mostly concentrate on the 185 biopsies, selected because there was adequate information about immunoglobulin status of the patients, 110 with an immunoglobulin abnormality and 75 without. Free light chain immunostaining was more likely to be done if an abnormality was known or suspected or was in the differential diagnosis.

RESULTS
Comparison of immunofluorescence and immunoperoxidase
Ten of 13 biopsies showed agreement between the methods, with AL amyloid (5), no light chain deposition (3), and deposition disease (2). The other three showed no free light chain deposition, and only faint deposition on immunofluorescence, one of κ in tubular basement membranes with slight chronic damage, one of κ in glomerular basement membranes but no structural glomerular abnormality, and one of more λ than κ in tubular basement membranes but no clinical immunoglobulin abnormality.

Comparison of different antibodies using immunoperoxidase
The 14 non-NAC biopsies comparing free light chain and rabbit antibodies had cast nephropathy (3), amyloid (3, two AL, one AA), no immunoglobulin abnormality (3), renal infiltration by lymphoplasmacytic lymphoma or myeloma (2), cryoglobulinemic glomerulonephritis (2, both type II) and deposition disease (1). Rabbit antibodies after automated antigen retrieval showed heavy staining of tubular epithelium, interstitial tissues and blood, but background staining was much lighter without pretreatment (figures 1, 2). After automated antigen retrieval, deposits of light chains in AL amyloid, cryoglobulinemia and deposition disease were impossible to distinguish from the background, and without pretreatment, immunostaining for light chains was weak and only seen in one case, with AL amyloid. Deposits were easily detected with free light chain antibodies, with clean background and no blood in vessels (figures 3, 4).

Both types of antibody showed a preponderance of one light chain in casts and tubular epithelium in cast nephropathy. Light chain restriction was identified in the myelomatous infiltrate only with rabbit antibodies, but with neither antibody in the lymphoplasmacytic lymphoma. Comparison of the antibodies, including findings in NAC (below), is in table 1.

Amyloid
With free light chain antibodies, 25 of 41 biopsies showed AL amyloid (21 λ, four κ), with concordance between clinical and

Figure 1 Renal biopsy containing amyloid in glomeruli in a patient with a κ paraprotein. Immunostaining with the rabbit antibody to κ light chains after automated antigen retrieval shows extensive staining but no detectable selective staining of amyloid deposits.

Figure 2 The same renal biopsy as in figure 1. Immunostaining with the rabbit antibody to κ light chains without pretreatment shows light background staining but no detectable staining of amyloid deposits.
immunoperoxidase findings (tables 2, 3). NAC confirmed \( \lambda \) AL amyloid in 13 of 17 \( \lambda \) cases and \( \kappa \) AL amyloid in one of two \( \kappa \) cases by immunohistology, and in four and one, respectively, from overall evidence.

Eight biopsies showed AA amyloid and three nonAL, nonAA amyloid with no free light chain immunostaining, concordant with no immunoglobulin disorder. NAC confirmed five cases of AA amyloid by immunohistology, and identified two nonAL, nonAA cases as LECT2 amyloid and one as fibrinogen A \( \alpha \) amyloid. Another case had no clinical immunoglobulin abnormality, but amyloid deposits reacted with antibodies to amyloid \( A \) and free \( \kappa \). NAC diagnosed AA amyloid immunohistologically.

The other four cases had a clinical immunoglobulin abnormality. One with a \( \kappa \) paraprotein showed deposition of both free \( \kappa \) and amyloid \( A \). NAC showed no deposition on immunohistology but considered this \( \kappa \) AL amyloid on overall evidence. In the other three, no free light chain deposition was detected. One with a \( \lambda \) Bence Jones protein was confirmed \( \lambda \) AL amyloid immunohistologically at NAC. One with a \( \lambda \) paraprotein had no immunohistological deposition at NAC but was considered \( \lambda \) AL amyloid on overall evidence. In one case with a \( \kappa \) paraprotein, the amyloid reacted only with amyloid \( A \) antibody. AA amyloid was confirmed by NAC.

**Other conditions**

Light chain cast nephropathy

Findings were concordant between immunostaining for free light chains and immunoglobulin abnormalities in 32 of 33 cases (26 \( \kappa \), six \( \lambda \)), with definite preponderance of one light chain in casts and tubular epithelial cells (figures 5, 6). In the

| Antibodies to free light chains | Deposition | No deposition |
|---------------------------------|------------|--------------|
| 29                              | 17         |              |
| Abnormality/no abnormality      | Abnormality/no abnormality |
| 28/1\*                          | 4/13       |              |

Sensitivity 88%, specificity 93%

| Conventional antibodies to light chains | Deposition | No deposition |
|----------------------------------------|------------|--------------|
| 20                                    | 26         |              |
| Abnormality/no abnormality             | Abnormality/no abnormality |
| 2010                                  | 124/14     |              |

Sensitivity 63%, specificity 100%

Difference in sensitivity (free light chain antibodies – conventional antibodies)=25% (95% CI 9% to 41%); difference in specificity (free light chain antibodies – conventional antibodies)=7% (95% CI –14% to 0%).

\* \( \kappa \) & AA deposition in AA amyloid.

\† No deposition in AL amyloid (2), neoplastic infiltration (2).

\‡ No deposition in AL amyloid (8), cryoglobulinaemic glomerulonephritis (2), light chain deposition disease (1), neoplastic infiltration (1).

**Table 2** Light chain immunohistological findings in 41 renal biopsies containing amyloid

| AL amyloid | AA amyloid | Other amyloid |
|------------|------------|--------------|
| Final diagnosis | 28 (23 \( \lambda \), 5 \( \kappa \)) | 10 | 3 |
| Free light chain antibodies n=41 | 21 \( \lambda \)+ve, 2 \( \lambda \) –ve, 4 \( \kappa \)+ve, 1 \( \kappa \) & AA+ve (25/28 (89%) unambiguously diagnostic) |
| Conventional light chain antibodies n=32 | 14 \( \lambda \)+ve, 5 \( \lambda \) –ve, 1 \( \kappa \) +ve, 2 \( \kappa \) –ve (15/22 (68%) diagnostic) |

For AL amyloid, difference in proportion unambiguously diagnostic (antibodies to free light chains – conventional antibodies to light chains)=21% (95% CI 2% to 39%).

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**Figure 3** The same renal biopsy as in figure 1. Immunostaining with the sheep antibody to free \( \kappa \) light chains shows selective staining of amyloid deposits in glomeruli.

**Figure 4** The same renal biopsy as in figure 1. Immunostaining with the sheep antibody to free \( \lambda \) light chains shows no staining.
In a discordant case, there was a κ paraprotein and a few casts showed anomalous colours with Congo red as found in some cases of cast nephropathy, without definite preponderance of κ on immunostaining.

**Light chain deposition disease**

Findings were concordant in all six cases of deposition disease (five κ, one λ), with glomerular deposition of the appropriate light chain, and often deposition in tubular basement membranes (figure 7).

**Other renal effects of immunoglobulin abnormalities**

In four of 11 cases, free light chain immunostaining did not show a preponderance of one light chain. Two were kidneys infiltrated by myeloma and lymphoplasmacytic lymphoma, and two were cases of cryoglobulinaemia (one type II, one type III). There was concordance in the other seven cases, all κ, four cryoglobulinaemic glomerulonephritis (one type I, three type II) (figure 8), two tubulointerstitial nephritis with light chains in tubular basement membranes, and one with crystalline deposits of light chains in podocytes.

**No evidence of renal effects of an immunoglobulin abnormality**

In 31 cases, there was no detectable effect of a paraprotein in the kidney and no immunohistological disproportion between κ and λ. The commonest diagnoses were diabetic glomerulopathy (6), membranous nephropathy (4) and chronic ischaemic damage (4).

### Table 3

| Light chain deposition and immunoglobulin abnormality | No light chain deposition and no immunoglobulin abnormality | No light chain deposition and immunoglobulin abnormality | Light chain deposition and no immunoglobulin abnormality |
|------------------------------------------------------|-------------------------------------------------------------|--------------------------------------------------------|--------------------------------------------------------|
| Amyloid n=41                                         | 26 (1*)                                                     | 11                                                     | 3 (1†, 2*)                                              |
| Light chain cast nephropathy n=33                    | 32                                                          | –                                                      | 1*                                                     |
| Light chain deposition disease n=6                   | 6                                                           | –                                                      | 0                                                      |
| Cryoglobulinaemia n=6                                | 4                                                           | –                                                      | 2 (1†, 1*)                                             |
| Miscellaneous n=3                                    | 3                                                           | –                                                      | 0                                                      |
| Neoplastic infiltration n=2                          | 0                                                           | –                                                      | 2†                                                     |
| No evidence of renal effects of immunoglobulin abnormality n=31 | 0                                                           | –                                                      | 3†                                                     |
| No immunoglobulin abnormality (excluding amyloid cases) n=63 | –                                                           | 62                                                     | –                                                      | 1*                                                     |

*Considered discordant.
†Not considered discordant, as discussed in text.

**Figure 5** Renal biopsy showing light chain cast nephropathy in a patient with an IgA λ paraprotein. Immunostaining with the sheep antibody to free λ light chains shows deposition in casts.

**Figure 6** The same renal biopsy as in figure 5. Immunostaining with the sheep antibody to free κ light chains shows less deposition of κ than λ (figure 5).
No immunoglobulin abnormality
Excluding 12 amyloid cases, 63 biopsies were from patients without an immunoglobulin abnormality as defined above, although many had immunoglobulin deposits, shown by antibodies to heavy chains. In 62 there was no disproportion between \( \kappa \) and \( \lambda \) on immunohistology. One showed subendothelial membranoproliferative glomerulonephritis with more \( \kappa \) than \( \lambda \) in glomerular deposits, without evidence of a monoclonal gammopathy.

**DISCUSSION**

Immunofluorescence on frozen sections and immunoperoxidase or immunofluorescence on paraffin sections rarely show complete concordance in detection of light chain deposition. Which method gives the correct or more clinically relevant finding is undetermined.\(^6\) The discrepancies in our study were minor and of uncertain clinical significance.

Comparison on paraffin sections between free light chain and conventional antibodies showed that differences were due to the antibodies, not pretreatments. The free light chain antibodies did not detect whole immunoglobulins and were more sensitive than conventional antibodies (table 1). Light chains in deposits were readily differentiated from background. The free light chain antibodies were better at typing AL amyloid, giving stronger and generally unequivocal staining (figures 3, 4; table 2). Two of 28 cases (7%) of AL amyloid were missed, although to different extents all antibodies to light chains miss cases of AL amyloid, up to 35% in one series.\(^14\)\(^\rightarrow\)\(^16\)\(^\rightarrow\)\(^17\)

Some cases of AL amyloid will still need other investigations for definitive diagnosis, such as laser microdissection of sections and mass spectrometry.\(^17\)\(^\rightarrow\)\(^18\) Reactivity of amyloid with multiple antibodies has been reported, for instance in 34% of biopsies with AA amyloid.\(^19\) and two of our 41 amyloid cases (5%) showed ambiguous immunostaining for both free \( \kappa \) and amyloid A. Four amyloid cases (10%) therefore showed discordance between clinical and immunohistological findings, overlooking one case with a paraprotein but confirmed to be AA amyloid.

The free light chain antibodies were useful in confirmation of deposition disease, with complete concordance, and usually of cryoglobulinaemic glomerulonephritis (figures 7, 8). The discordant case of type II cryoglobulinaemia, with mixed monoclonal IgM and polyclonal IgG, may have had insufficient excess of free light chains to be detectable. In type III cryoglobulinaemia, with polyclonal deposits, no excess of one free light chain would be expected, and was not considered discordant. There was little difference between free light chain and conventional antibodies in detection of light chains in cast nephropathy. The failure of free light chain antibodies to show an excess of one light chain in one of 33 cases (3%) may be because in virtually every case of cast nephropathy both light chains are found (figures 5, 6), and a small excess of one may be undetectable.

The free light chain method was developed to detect deposits rather than intracellular or surface immunoglobulins, and neoplastic infiltrates can be investigated in other ways.\(^20\) There was no deposition of free light chains in 32 cases with a monoclonal gammopathy but without diagnosable renal effects, including one case of AA amyloid (29% of those with immunoglobulin abnormalities). Similarly, 27% of patients with myeloma\(^3\) and 63% with a monoclonal gammopathy\(^4\) had no evidence of renal lesions related to a paraprotein. Accordingly, only seven of 185 cases (4%) were considered discordant: four amyloid, one cryoglobulinaemia, one light chain cast nephropathy, and one glomerulonephritis with excess \( \kappa \) deposition but no gammopathy.

Polyclonal free light chain antibodies are of practical use in the study of renal biopsies and have advantages over conventional light chain antibodies. Methods are similar and overall costs are also likely to be similar.
Take home messages

- Polyclonal antibodies to free light chains are an improvement on conventional antibodies to light chains in the immunoperoxidase study of paraffin sections of renal biopsy specimens, giving a cleaner background and sharper discrimination of sites of deposition.
- These antibodies are more sensitive than conventional antibodies in the detection of cases of immunological disorders in the kidney such as AL amyloid, light chain deposition disease and cryoglobulinaemic glomerulonephritis.

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