Glomerular nodules and long-spacing collagen in kidneys of patients with multiple myeloma

G. E. SCHUBERT1 AND A. ADAM

From the Institute of Pathology, University of Tübingen, Tübingen, West Germany

SYNOPSIS Light microscopic examination of the kidneys in 170 patients with multiple myeloma revealed three from patients without diabetes with glomerular nodules that were indistinguishable from those of diabetic glomerulosclerosis. Electron microscopic studies showed the nodules to be deposits of fine-grained material that corresponded to precipitations of protein but lacked the basement-membrane-like material characteristic of nodular glomerulosclerosis in diabetics. What is more, a unique collagen was observed in the marginal area of the nodules and in the periglomerular interstitium. The possible causes of the origin and distribution of this collagen not so far described in the human kidney are discussed.

In recent years, glomerular nodules which could not be differentiated by light microscopy from a nodular diabetic glomerulosclerosis have been observed in kidneys of patients with multiple myeloma who showed no clinical indications of diabetes mellitus. So far no suitable explanation for the pathogenesis and morphogenesis of these nodules has been given (Kenis, Cauchie, Potvliege, Smulders, Gompel, and Lambert, 1961; Kobernick and Whiteside, 1957; Martin, 1972; Olsen, 1972). We have recently seen three cases with these glomerular nodules and performed electron microscopic studies. We found that the ultrastructure of the mesangial nodules was different from the classical diabetic glomerulosclerosis. Furthermore, an extraordinary collagen was found which, to our knowledge, has not yet been described in the human kidney. We consider, therefore, a detailed description of these cases is warranted.

Materials and Methods

The kidneys from 170 patients with multiple myeloma were evaluated histologically. Nodular widenings of the glomerular mesangium, which strongly resembled those of diabetic glomerulosclerosis, were observed in three cases. Clinically, however, diabetes mellitus could be ruled out in these patients, and the kidneys were subjected to further examination.

The renal tissue examined was secured by needle biopsy in case 1 and by necropsy eight and about 20 hours after death in cases 2 and 3 respectively. The light microscopic investigations were carried out on paraffin sections 8μ thick using the following stains: haematoxylin-eosin, Goldner-Trichrom, PAS, Elastica van Gieson, and Congo red according to Puchtel (1962). The tissue in the second and third cases was examined on methachrylate-embedded Movat-silvered semithin sections 0.5μ thick. Electron microscopic studies were performed on formol-fixed necropsy specimens, postfixed in cold 2% OsO4 at a pH of 7.2. After embedding in methachrylate stain sections were stained with saturated uranyl acetate for 45 minutes; contrast staining followed in 0.4% lead acetate for two minutes.

Results

Case 1

A 62-year-old man had an IGA plasmocytoma, the existence of which had been known for about a year. Because of renal unsufficiency with oliguria and the increase of the serum creatinine to 17 mg% and of the serum urea to 220 mg%, a needle biopsy of the kidney was performed. The blood pressure at the time of the puncture was 160/80 mm Hg; the erythrocyte sedimentation rate was 75/127.

Histologically, four of the seven glomeruli in the
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specimen were completely hyalinized. One glomerulus showed a slight, another a pronounced, nodular widening of the mesangium without any distinct increase in mesangial cells; the capillary walls were relatively delicate. The nodules took on a yellowish-red colour with van Gieson stain, Congo red was negative and fibrin was not present. Atrophic tubules with isolated hyaline casts were present in the interstitium which was broadened by fibrosis and infiltrated by round cells. The interlobular arteries showed a significant hyalinosis of the intima and media.

The smooth light medium-red surfaced kidneys together weighed 240 g. The microscopic examination showed several fully hyalinized glomeruli in the subcapsular region. Atrophic tubules in this region, with conspicuously widened hyalinized basement membranes, were surrounded by fibrotic interstitium. In nearly all of the remaining glomeruli marked

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**Fig 1** Light microscopic appearance (25×) of a haematoxylin-and-eosin-stained kidney section from a 72-year-old non-diabetic man with Bence-Jones-K-myeloma (case 2). Note the numerous cell-deficient glomerular nodules.

**CASE 2**
A 72-year-old man, had a Bence-Jones-K-plasmacytoma which had been diagnosed one and a half years before death. The patient was oliguric during the last two weeks of life. Twelve days before death the serum creatinine rose to 6.3 mg% and the daily urine volume decreased to 400 ml. The blood pressure varied between 110/80 and 150/80 mm Hg. There were no clinical indications of diabetes mellitus. The titre of insulin antibody in the serum (Dr Wehner, Tübingen) was not increased. The patient died of uraemia and a severe pancreatitis.

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**Fig 2a** Semithin section of a glomerulus (420×) from case 2. Granular or grainy-appearing masses are visible among the numerous cell-deficient nodular widenings of the mesangium. The basement membranes are inconspicuous.

**Fig 2b** Electron microscopic appearance (77 000×) of a section of a nodular area from the mesangium in case 2. Note the fine granular deposits of the precipitated protein type.
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Fig 3a  Electron micrograph (14 000 ×) of the 'banded bodies' or 'long-spacing collagen' in the region of the capillary wall (arrows) and in the periglomerular interstitium (arrows) in case 2.

Fig 3b  Enlarged sections from fig 3a (77 000 ×) showing the delicate longitudinal striations of the long-spacing collagen.

Fig 4  Long-spacing collagen (77 000 ×) attached or in close apposition to normal collagen fibres in the glomerular mesangium of case 2. The periodicity of the broad cross striations is 88 nm. The lower margin is the basement membrane.
nodular widening of the mesangium was recognizable (fig 1), which was especially pronounced in non-atrophic areas of the inner and medial cortical zone. The cell-deficient, often cell-free, nodules were yellowish-red on van Gieson staining. They reacted negatively to amyloid staining as did the other regions. In semithin sections, the nodules appeared to have a fine granular or plaque-like structure (fig 2a). The basement membranes were not thickened, and subepithelial and subendothelial deposits could be found.

Electron microscopic examination showed these nodular mesangial thickenings to be fine granular structures of the precipitated protein type (fig 2b). Irregularly arranged precollagen and collagen fibres were observed in marginal sections of the nodules (fig 4). Spindle-shaped or long, ribbon-like structures with an average width at the widest part of 68.7 ± 25.5 nm and a total length of 375.4 ± 101.7 nm were seen in the immediate vicinity of the irregular fibres. These structures, marked by broad, dark, transverse striations with an average periodicity of 77.4 ± 15.5 nm, were seen under higher magnification to have delicate longitudinally orientated filaments (fig 3) and strongly resembled 'banded bodies' (Pillai, 1964) or 'curled collagen' (Garron, Feeney, Hogan, and McEwen, 1958). They occurred not only in the marginal areas but also occasionally within the fine granular glomerular nodules (fig 4), especially near the edges of mesangial cell nuclei. They were frequently observed in the interstitium surrounding the glomeruli and particularly often in the immediate vicinity of the basement membrane. Fibrin was not present in the glomeruli nor was amyloid found in the kidneys, although the necropsy showed an amyloidosis of the prostate.

**CASE 3**

In a 70-year-old man with gradually increasing renal insufficiency, the serum urea rose shortly before death to 365 mg%. The haemoglobin had fallen to 55%, the serum calcium to 3.2 m-equiv/l, and the serum phosphorus had increased to 7.5 m-equiv/l. The blood sedimentation rate was 123/143 mm and the globulin fraction in the serum increased to 20%. The blood pressure was 215/110 mm Hg. The patient had a heavy proteinuria with a protein excretion of 10 g/24 hr. A plasmacytoma was first diagnosed by sternal puncture only four weeks before death.

Histological examination of the enlarged, pale kidneys showed nodular widenings of the mesangium in practically all glomeruli. Under the light microscope these widenings appeared as fairly homogeneous deposits (fig 5a) between which lay several mesangial cells. The interstitium was slightly fibrotic; the renal tubules were wide. Under the electron microscope the glomerular deposits in the widened mesangium proved to be of the protein precipitate type. Embedded in these masses were numerous long, ribbon-like structures (fig 5b), which strongly resembled those of case 2. The periodicity of the broad, dark transverse striations was, on average, 78.0 nm. Often these structures were found...
in the vicinity of an apparently normal basement membrane or on the surface of mesangial cells. In addition, isolated collagen fibres were seen in the region of the mesangium.

Discussion

Examination of the kidneys from patients with multiple myeloma has failed to demonstrate characteristic changes in the glomeruli that can account for the frequent occurrence of renal insufficiency associated with this illness (Boulet, Mirouze, Pages, Barjon, Fabre, and Jaffiol, 1962; Hamburger, Richet, Crosnier, and Antoine, 1966; Kenis et al, 1966; Reubi, 1970; Schubert, 1974). The repeatedly described thickenings of the basement membrane, the increases in the numbers of epithelial, endothelial, and mesangial cells, as well as subepithelial deposits (Abrahams, Pirani, and Pollak, 1966; Costanza and Smoller, 1963; Fries, Bansilion, Brunat, Molinie, and Traeger, 1967; Kobernick and Whiteside, 1957; Rosen, Cortell, Adner, Papadopoulos, and Barry, 1967), are, according to Fisher, Perez-Stable, and Zawadzki (1964), more likely to be the result of an altered protein supply than the cause of renal disturbances. Widening of the mesangium (Allen, 1962; Fisher et al, 1964) also cannot account for all cases of renal disturbances in patients with multiple myeloma and renal insufficiency, since Wehner, Feurer, and Wehner (1971) by exact measurements found decreased mesangial areas in some cases.

Our findings agree with reports of nodular widenings of the mesangium in some patients with plasmacytoma (Kobernick and Whiteside, 1957; Kenis et al, 1961; Martin, 1972; Olsen, 1972). The kidney morphology in these patients could not be differentiated from a diabetic glomerulosclerosis, even though there were no clinical signs of diabetes mellitus. Olsen (1972) suggested that the nodular widenings might be produced by abnormal mesangial deposits such as immunoproteins that provoke an increased synthesis and/or reduced catabolism of basement membrane-like substance. However, deposits of basement membrane-like material typical for long-term diabetic patients with nodular glomerulosclerosis were not apparent in the kidneys of our myeloma patients. Consequently, we were unable to verify Olsen's theory by electron microscopic examination of the glomerular nodules. Nevertheless, our findings show that the fine granular deposits in the mesangium correspond to protein precipitates which, according to Olsen, could precede the increase of basement membrane-like substance in this region. Amyloid deposits (Tellum and Lindahl, 1954; Heptinstall and Joekes, 1960; Olsen, 1972; Lendrum, Slidders, and Fraser, 1972) could neither be demonstrated by light nor by electron microscopy in our cases. Thus, although we have not completely clarified the nature of these nodules of plasmacytoma, our electron microscopic results point strongly toward a structure consisting of protein deposits and a scanty collagen fibre synthesis in the marginal sections of the glomerulus.

The kidneys of our patients examined by electron microscopy showed longish, frequently spindle-shaped bodies with coarse cross striations, which strongly resemble the 'banded bodies' of Pillai (1964), the 'curly collagen' of Garron et al. (1958), and the 'long spacing collagen' which Luse, Zopf, and Cox (1963) could reconstitute in vitro. To the best of our knowledge, these structures have not been previously described in the human kidney. On the other hand, this type of collagen has been described, for example, in the walls of cerebral vessels in old age (Schloot, 1966); in acoustic neurinomas and peripheral Schwannomas, where this structure was found to be absolutely diagnostic of these tumours (Raimondi, Mullan, and Evans, 1962; Luse, 1960); in other brain tumours (Ramsey, 1964); in Meissner's tactile corpules of the human finger (Cauna and Ross, 1960); within degenerating sensory cells of the human membranous labyrinth in Ménière's disease (Friedman, Cawthorne, McClay, and Bird, 1963); in human nodes (Luse et al, 1963) in Descemet's membrane of man and several animals (Feeney et al, 1961; Jakus, 1962; Van der Zypen, 1971); in retinal arterioles (Gärtnert, 1966); around capillaries in the subcommissural organ of the brain of rats (Wetzstein, Schwink, and Stranka, 1963; Naumann and Wolfe, 1963); in the perineurium of tibial nerves of rats after constriction (Pillai, 1964); and in the region of the iridocorneal angle of the eye in patients with clinically manifest or latent systemic disease of the connective tissue (Rohen, 1962).

Information is still too incomplete to warrant any definite conclusions regarding the cause of the origin and distribution of this particular collagen in the human kidney. We presume that changes in the local milieu favour its synthesis, since the dependence of the collagen macromolecule (tropocollagen) aggregation on environmental conditions is well known. Factors such as the amount and kind of salt, pH, temperature, and the concentration of collagen determine the formation of different aggregates with different band patterns. It is interesting to note that Pappas (1960) produced collagen with a periodicity of 120 nm amid normal filaments by adding chondroitin sulphate to the medium in tissue cultures of fibroblasts. The lower periodicity of the 'banded bodies' or 'long-spacing collagen' in our cases could be due to the existence
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of unique local conditions prevailing in their kidneys. The spatial relationships between the glomerular nodules, the basement membranes, and the 'banded bodies' is consistent with this notion. In subsequent investigations it will be necessary to examine whether particularly favourable conditions occur for the synthesis of this collagen in kidneys of patients with plasmacytoma.

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References


