Mechanisms of progression and regression of renal lesions of chronic nephropathies and diabetes

Giuseppe Remuzzi, … , Ariela Benigni, Andrea Remuzzi


The incidence of chronic kidney diseases is increasing worldwide, and these conditions are emerging as a major public health problem. While genetic factors contribute to susceptibility and progression of renal disease, proteinuria has been claimed as an independent predictor of outcome. Reduction of urinary protein levels by various medications and a low-protein diet limits renal function decline in individuals with nondiabetic and diabetic nephropathies to the point that remission of the disease and regression of renal lesions have been observed in experimental animals and even in humans. In animal models, regression of glomerular structural changes is associated with remodeling of the glomerular architecture. Instrumental to this discovery were 3D reconstruction studies of the glomerular capillary tuft, which allowed the quantification of sclerosis volume reduction and capillary regeneration upon treatment. Regeneration of capillary segments might result from the contribution of resident cells, but progenitor cells of renal or extrarenal origin may also have a role. This review describes recent advances in our understanding of the mechanisms and mediators underlying renal tissue repair ultimately responsible for regression of renal injury.

Find the latest version:

http://jci.me/27699-pdf
The incidence of chronic kidney diseases is increasing worldwide, and these conditions are emerging as a major public health problem. While genetic factors contribute to susceptibility and progression of renal disease, proteinuria has been claimed as an independent predictor of outcome. Reduction of urinary protein levels by various medications and a low-protein diet limits renal function decline in individuals with nondiabetic and diabetic nephropathies to the point that remission of the disease and regression of renal lesions have been observed in experimental animals and even in humans. In animal models, regression of glomerular structural changes is associated with remodeling of the glomerular architecture. Instrumental to this discovery were 3D reconstruction studies of the glomerular capillary tuft, which allowed the quantification of sclerosis volume reduction and capillary regeneration upon treatment. Regeneration of capillary segments might result from the contribution of resident cells, but progenitor cells of renal or extrarenal origin may also have a role. This review describes recent advances in our understanding of the mechanisms and mediators underlying renal tissue repair ultimately responsible for regression of renal injury.

Historical view
Progression to end-stage renal disease (ESRD) is common in chronic nephropathies, independent of the initial insult. Since 1830, disorders of the kidney with albuminuria and changes of blood chemistry were defined as Bright’s disease (1, 2). In his 1931 book The renal lesion in Bright’s disease (2), Thomas Addis indicated that study of the urine could be advantageous to the categorization of structural disease in the kidneys. By 1939, Addis (3) introduced the idea of “osmotic work” and calculated how this work would vary with the amount of protein in the diet. An important implication of those studies was that dietary protein restriction could be of help for patients with renal impairment. Meanwhile, in 1932 Alfred Chanutin and Eugene Ferris (4) observed that removal of three quarters of the total renal mass in the rat led to a slowly progressive deterioration in the function of the remaining nephrons, with progressive azotemia and glomerulosclerosis. The glomerular lesions of the remnant kidney were associated with abnormal glomerular permeability and proteinuria. At that time, proteinuria was considered a marker of the extent of glomerular damage, despite the fact that Franz Volhard and Theodor Fahr in 1914 (5) and Wilhelm von Mollendorf and Philipp Stohr in 1924 (6) had already found that renal damage was related to exuberant protein excretion in the urine. In 1954 Jean Oliver and colleagues (7) recognized protein droplets in the cytoplasm of tubular cells. They suggested that such findings were possibly the result of impairment in the process of reabsorption of plasma proteins normally carried out by the renal tubule and proposed that proteinuria could lead to structural and functional nephron damage. Robert Platt, during the second of the two Lumleian Lectures delivered to the Royal College of Physicians of London (8), reported that “the functional disturbances known to occur in human renal disease are precisely those which occur in animal experiments as a result of reduction in the amount of functioning renal substance, that is, loss of nephrons. Rats from which 80% of the renal tissue has been removed had hypertrophy of the remaining nephrons, as they take in a volume of work which they would never be called up to perform in normal kidney.” This was interpreted as a possible adaptation to overcome the handicap imposed by the loss of nephrons. Shimamura and Morrison (9) found hyalinization of the glomerular structure after partial five-sixths nephrectomy in animals. In the late 1960s Brenner had access to a unique strain of rat with glomeruli on the cortical surface and developed a micropuncture technique (10). By such means, Brenner and coworkers clarified the pathophysiology of renal adaptation to nephron loss. They found that after removal of renal mass, arteriolar resistance lowers and plasma flow increases in remnant glomeruli (11). The tone of afferent arterioles drops by a greater degree than that of efferent ones, which increases glomerular capillary hydraulic pressure, leading to more filtrate formed per nephron. “These changes serve to enhance the filtration capacity of the remaining nephron units, minimizing the functional consequences of nephron loss, but are ultimately detrimental” (12). Brenner also found (13) that therapies that attenuate such adaptive changes limit GFR decline and structural damage (14). A possible link between glomerular hypertension and proteinuria was not established formally at that time; nevertheless, Cameron had already found that patients with nephrotic syndrome did progress more rapidly than those who had never been nephrotic (15, 16). This was in harmony with previous findings by Habib (17) that in focal and segmental glomerulosclerosis those patients who
had their proteinuria lowered by corticosteroids did not develop renal failure. In 1986, studies in rats (18) renewed the old idea that urinary proteins may have intrinsic renal toxicity and contribute to the progression of damage. Later, Eddy and Michael (19), in an experimental model of nephrosis, found that proteinuria correlated with increased numbers of interstitial cell infiltrates.

Excessive proteinuria was also induced in rats by intraperitoneal injections of albumin (20, 21) or by transplanting a pituitary tumor (22). In both models proteinuria was followed by tubular damage and interstitial inflammation of macrophages and T lymphocytes. The availability of cultured cells with features of differentiated glomerular epithelial cells has recently prompted investigation into the effects of plasma proteins on the function of podocytes (23), currently thought to play a key role in the progression of renal lesions.

Epidemiology of chronic renal diseases

Chronic kidney disease (CKD) is a worldwide threat to public health, but the true dimension of this problem is not fully appreciated. Approximately 1.8 million people are currently treated with renal replacement therapy (RRT), which consists primarily of kidney transplantation, hemodialysis, and peritoneal dialysis (24, 25). More than 90% of these individuals live in industrialized countries, while available RRT in developing countries is scarce, and null in underdeveloped areas.

The treatment of a relatively limited number of patients represents a major societal commitment, since RRT absorbs a significant proportion of the health care budget (25). A forecast analysis based on data from the US Renal Data System and Medicare predicts that by the year 2010 the total number of patients on RRT will be double the current number and will exceed 650,000, which is expected to increase public expenditure for dialysis to $28 million per year (26).

Diabetes is the most common cause of ESRD in the US and in many other countries (27). Patients that have diabetes and are on RRT have a worse outcome, and their management costs a great deal more compared with patients that are on dialysis for other diseases.

A relevant question for health care planning is how many patients at an early stage of renal dysfunction will progress to ESRD? Recently the National Kidney Foundation has adopted a stratification of patients with CKD into 5 stages (28). Stage 1 includes patients with normal GFR but with urinary abnormalities; stages 2–4, individuals with increasing severity of disease; and stage 5, patients with ESRD (28). This staging system has been used to estimate the prevalence of CKD in the US. A survey was conducted in the frame of the Third National Health and Nutrition Examination Survey (NHANES III). A sample of 15,625 adults aged 20 years and older was analyzed. Kidney function, kidney damage, and stages of CKD were estimated from calibrated serum creatinine levels, spot urine albumin levels, age, sex, and race. The prevalence of CKD in the US adult population was estimated to be 10.8% (approximately 19.2 million people) (29). In Europe, a similar screening program was conducted in the frame of the Prevention of Renal and Vascular End-Stage Disease (PREVEND) study (30). Eighty thousand people in Groningen, The Netherlands, were evaluated for renal function and urinary abnormalities. It was found that up to 12% of the adult population had some degree of renal damage. If these data were to be extrapolated to the world population, the number of people with CKD could be estimated to be in the hundreds of millions.

Figure 1
The progressive nature of chronic kidney disease. The progression to ESRD, as underlined by the progressive decline of GFR, is highly variable. Here is reported the natural history of autosomal-dominant polycystic kidney disease (ADPKD) in patients with PKD1 mutation in the PKD1 gene as an example of genetic renal disease. Progressive renal disease occurs in 20–40% of patients with type 2 diabetes. Progression to renal failure occurs in 30% of patients with IgA nephropathy after a follow-up of 25 years. Similarly, 30% of patients with membranous nephropathy reach ESRD within the 30-year follow-up period. A more rapid course is observed for patients with mesangial capillary glomerulonephritis or primary focal and segmental glomerulosclerosis, who possess persistently high urinary protein excretion rates.

Figure 2
Progression of nephropathy in type 2 diabetes. Following 10 years of stable renal function and normal UAE rate (<20 mg/min or <30 mg/d), UAE increases in 20–40% of type 2 diabetic patients. UAE persistently in the range of 20–200 mg/min or 30–300 mg/d (microalbuminuria) heralds the onset of incipient nephropathy. If left untreated, 20–40% of patients progress to overt nephropathy, a syndrome of macroalbuminuria (UAE rate >200 mg/min or >300 mg/d), declining glomerular filtration rate, and increased cardiovascular morbidity. With the onset of macroalbuminuria renal function progressively declines, and ESRDs eventually develop, requiring RRT with dialysis or transplantation. Diabetics with overt proteinuria have a higher risk of dying from cardiovascular disease (122).
ESRD is the most visible outcome of CKD. However, cardiovascular disease (CVD) is frequently associated with CKD, and individuals with CKD are likely to die of CVD before they develop renal insufficiency (54, 55, S12). Non-nephrotic patients with heavy albuminuria may progress to overt nephropathy, a syndrome of macroalbuminuria (UAE >200 µg/min or >300 mg/d), declining GFR, and increased cardiovascular morbidity (44, 45, 47, S9). In the case of macroalbuminuria, GFR relentlessly declines at an average of 10–12 ml/min/yr (48) (Figure 2). At least two-thirds of patients with overt nephropathy will die from CVD before they progress to ESRD, a mortality rate 5- to 8-fold higher than in the average population (49). While on dialysis, 21% of these patients will die within 1 year (50).

Nondiabetic glomerulopathies include IgA nephropathy. Progression to ESRD occurs in 30% of patients after a follow-up of 25 years (51). A further 20% will have impaired renal function and will progress eventually. Membranous nephropathy has a variable course (52, 53, S10, S11) with an insidious onset and increasing proteinuria up to nephrotic ranges. In the long term, spontaneous remission occurs in up to 30% of individuals, while the remaining two-thirds experience either equally persistent proteinuria of variable degrees — although usually of declining severity, and with normal or impaired but stable renal function — or progressive disease eventually leading to ESRD. Overall, approximately 30–40% of patients develop significant renal failure 10–15 years after the diagnosis of nephropathy (52, 53, S10, S11).

Most patients with mesangial proliferative glomerulonephritis and isolated hematuria maintain normal renal function for years. Cases with low-grade proteinuria also have a good long-term prognosis, while patients with heavy proteinuria may progress to renal insufficiency (54, 55, S12). Non-nephrotic patients with primary focal and segmental glomerulosclerosis have a benign disease. Actually more than 80% of these patients still retain normal renal function 10 years after the diagnosis of nephropathy (56). When proteinuria is within the nephrotic range, the course of the disease is rather malignant, and 50% of patients reach ESRD within 6–8 years (57).

Genetic studies in rodents and humans
Kidney diseases are in part genetically determined; therefore, individuals with a familial history of renal failure have a 3- to 9-fold greater risk of ESRD (58). However, the search for kidney

## The progressive nature of kidney disease
Certain renal diseases, including rapidly progressive glomerulonephritis, although rare, have a very rapid course that quickly leads to irreversible ESRD. More common nephropathies do progress less rapidly, but still evolve to ESRD at different rates (Figure 1). When serum creatinine is elevated beyond a certain level, progression is inevitable (S1–S3, 38).

The rate of progression of renal disease in hereditary kidney diseases such as autosomal-dominant polycystic kidney disease is highly variable, due to genetic heterogeneity (S4–S6, 39). Individuals with mutations in polycystic kidney disease 1 (PKD1) experience a more severe disease course, ultimately progressing to ESRD by the average age of 54, while individuals with mutations in PKD2 experience loss of renal function approximately 20 years later (40–42, S7). When renal function falls below 75% of normal the decline is rapid, requiring RRT in a matter of 5–10 years (S8, 43).

After 10–15 years of stable renal function or even hyperfiltration, small amounts of albumin may appear in the urine of 20–40% of patients with type 1 or type 2 diabetes. UAE rate in the range of 20–200 µg/min or 30–300 mg/d (microalbuminuria) (44–46, S9) is an early marker of nephropathy. If left untreated, 80–100% of microalbuminuric patients with type 1 and 20–40% of patients with type 2 diabetes progress to overt nephropathy, a syndrome of macroalbuminuria (UAE >200 µg/min or >300 mg/d), declining GFR, and increased cardiovascular morbidity (44, 45, 47, S9). In the case of macroalbuminuria, GFR relentlessly declines at an average of 10–12 ml/min/yr (48) (Figure 2). At least two-thirds of patients with overt nephropathy will die from CVD before they progress to ESRD, a mortality rate 5- to 8-fold higher than in the average population (49). While on dialysis, 21% of these patients will die within 1 year (50).

Nondiabetic glomerulopathies include IgA nephropathy. Progression to ESRD occurs in 30% of patients after a follow-up of 25 years (51). A further 20% will have impaired renal function and will progress eventually. Membranous nephropathy has a variable course (52, 53, S10, S11) with an insidious onset and increasing proteinuria up to nephrotic ranges. In the long term, spontaneous remission occurs in up to 30% of individuals, while the remaining two-thirds experience either equally persistent proteinuria of variable degrees — although usually of declining severity, and with normal or impaired but stable renal function — or progressive disease eventually leading to ESRD. Overall, approximately 30–40% of patients develop significant renal failure 10–15 years after the diagnosis of nephropathy (52, 53, S10, S11).

Most patients with mesangial proliferative glomerulonephritis and isolated hematuria maintain normal renal function for years. Cases with low-grade proteinuria also have a good long-term prognosis, while patients with heavy proteinuria may progress to renal insufficiency (54, 55, S12). Non-nephrotic patients with primary focal and segmental glomerulosclerosis have a benign disease. Actually more than 80% of these patients still retain normal renal function 10 years after the diagnosis of nephropathy (56). When proteinuria is within the nephrotic range, the course of the disease is rather malignant, and 50% of patients reach ESRD within 6–8 years (57).

## Genetic studies in rodents and humans
Kidney diseases are in part genetically determined; therefore, individuals with a familial history of renal failure have a 3- to 9-fold greater risk of ESRD (58). However, the search for kidney

### Table 1
Definitions of renal disease outcomes according to functional and structural changes

<table>
<thead>
<tr>
<th>Proteinuria</th>
<th>Progression</th>
<th>Remission</th>
<th>Regression</th>
</tr>
</thead>
<tbody>
<tr>
<td>≥1 g/24 h</td>
<td>Decliningb</td>
<td>Stable</td>
<td>Increasing</td>
</tr>
<tr>
<td>&lt;1 g/24 h</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;0.3 g/24 h</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*bFaster than physiological decline associated with aging (1 ml/min/1.73 m² per year). Table modified with permission from The Lancet (91).
for both the onset and the progression of diabetic nephropathy in type 1 and type 2 diabetes mellitus (61, 62), although other include polycystic kidney disease, Finnish nephrotic syndrome, progressive nephropathies, the gene coding for angiotensin-converting enzyme (ACE) has been found to represent an independent risk factor in patients with the ACE DD allele of ACE II or ID genotype (63, 64). Similarly, the D allele has been found to represent an independent risk factor for both the onset and the progression of diabetic nephropathy in type 1 and type 2 diabetes mellitus (61, 62), although other studies failed to find any association between insertion/deletion polymorphisms and diabetic nephropathy (S13). Rather than simply a marker of damage (70), ultrafiltered proteins can be toxic to the kidney (18, 71). In humans, proteinuria predicts progression and renal outcomes (72, 73, S17) in diabetic nephropathy (S15). The predictive value of proteinuria and of its correction with RAS inhibitors

### Key role of proteinuria and of its correction with RAS inhibitors

Rather than simply a marker of damage (70), ultrafiltered proteins can be toxic to the kidney (18, 71). In humans, proteinuria predicts progression and renal outcomes (72, 73, S17) in diabetic (S18) and nondiabetic renal disease (74). In the 274 patients with nondiabetic chronic nephropathies and clinical proteinuria large cohort of Japanese patients with type 2 diabetes at more than 80,000 single nucleotide polymorphisms loci allowed identification of the engulfment and cell motility 1 (ELMO1) gene as a candidate for conferring susceptibility to diabetic nephropathy (S15).

In an effort to identify kidney disease quantitative trait loci (QTL), Fawn-Hooded hypertensive rats, Munich Wistar Frømter rats, and Dahl salt-sensitive rats have been crossed with kidney disease-resistant strains, and the progeny have been phenotyped for markers of renal impairment (67). Fifteen genomic regions were found to contribute to kidney disease in the rat, with 12 replicated in separate rat crosses using different parental strains (67). One QTL (named Rf-2) on rat chromosome 1 is concordant to human chromosome 19q13, which contains a locus for a monogenic form of segmental glomerulosclerosis (S16). The rat QTL on chromosomes 2 and 3 are concordant with a kidney disease QTL identified in Pima Indians (68), and the rat QTL on chromosome 11 is concordant with 2 QTL for creatinine clearance on human chromosome 3q27 found in African Americans and whites (69). Kidney disease QTL in rats may help predict the location of corresponding disease genes in humans.

### Table 2

Regression of renal damage in experimental nephropathies

<table>
<thead>
<tr>
<th>Model</th>
<th>Start of treatment (wk)</th>
<th>Treatment</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Puromycin aminonucleoside-</td>
<td>0</td>
<td>Low-protein diet + ACEi</td>
<td>(99)</td>
</tr>
<tr>
<td>induced nephropsis</td>
<td>0</td>
<td>ACEi + Ang IIRA (125)</td>
<td></td>
</tr>
<tr>
<td>Subtotal nephrectomy</td>
<td>8</td>
<td>High dose ACEi (100)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>High dose ACEi (103)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>ACEI (S22)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>ACEI + Ang IIRA (S23)</td>
<td></td>
</tr>
<tr>
<td>Spontaneous nephropathy</td>
<td>20</td>
<td>ACEI (101)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>ACEI + Ang IIRA (102)</td>
<td></td>
</tr>
<tr>
<td>Aging</td>
<td>72</td>
<td>Ang IIRA (104)</td>
<td></td>
</tr>
<tr>
<td>Diabetic nephropathy</td>
<td>16 or 32</td>
<td>HGF gene transfer (105)</td>
<td></td>
</tr>
</tbody>
</table>

ACEI, ACE inhibitor; Ang IIRA, Ang II receptor antagonist.
Evidence from clinical trials in type 1 and type 2 diabetics. Bjorck et al. (86) found that in type 1 diabetics with overt nephropathy at comparable blood pressure, enalapril reduced the rate of GFR decline more than did treatment with a beta blocker. In another trial of 409 type 1 diabetics (80), results documented less progression to the combined end point of doubling serum creatinine, ESRD, or death while on captopril compared with placebo. Similar data are available for type 2 diabetics. In the Reduction of Endpoints in NIDDM with the Angiotensin II Antagonist Losartan (RENAAL) study (87), fewer patients reached the composite end point of doubling serum creatinine, ESRD, or death in the losartan group compared with placebo. In the losartan group, proteinuria was reduced by greater than 30%, while for individuals on placebo, there was a slight increase by the end of the follow-up period. The renal and cardiovascular effects of losartan were fully driven by the effects of the drug on urinary proteins and largely dependent on the amount of residual proteinuria. In the Irbesartan Diabetic Nephropathy Trial (IDNT) (88), patients on irbesartan had a lower rate of progression to the primary endpoint (doubled serum creatinine, progression to ESRD, or death) compared to amlodipine or placebo. The irbesartan group had greater than 30% reduction in proteinuria, compared with 6% for amlodipine and 10% for placebo. The beneficial effect remained even after correction for the difference in blood pressure.

Evidence from meta-analyses of ACE inhibitor trials. The ACE Inhibition in Progressive Renal Disease (AIPRD) meta-analysis (89, 90) confirmed that proteinuria is a strong risk factor for progression of chronic renal disease and that patients with more severe renal disease benefit most from ACE inhibitor treatment. Moreover, a strong relationship between early changes in urinary proteins upon treatment and disease outcome has been found (91), again confirming previous findings (75).

Is glomerular sclerosis reversible?

Preliminary observations in humans

The case of a patient with systemic lupus. Multidrug treatment, titrated to urinary proteins, blunted proteinuria and stabilized GFR for years in a young girl with nephrotic proteinuria and systemic lupus (92). She was given an ACE inhibitor in order to maintain a diastolic blood pressure of 90 mmHg or less. Her proteinuria reduced remarkably. Three years later, proteinuria was again measured at 9 g/24 h, and she had severe renal failure. There was no sign of active lupus. In addition to dietary sodium and protein restriction, she was given an ACE inhibitor plus an angiotensin II (Ang II) receptor antagonist, and a statin (92) with up titrations as deemed appropriate. Full remission of proteinuria was achieved within 6 months. Seven years later, proteinuria averaged 0.1–0.2 g/24 h, and renal function improved. This case shows that it is now possible to stabilize or even reverse disease progression, even in an advanced phase of the disease (Figure 3).

The case of pancreas transplantation. Regression of diabetic renal disease with glomerular architecture remodeling has been observed in a few patients with type 1 diabetes after 10 years of normoglycemia induced by a pancreatic transplant (93). Kidney function, however, did not return to pretransplant values. These findings show that sustained normoglycemia induces regression of lesions of diabetic glomerulopathy, extending previous evidence obtained in experimental animals (94, 95) to humans.
The REIN follow-up study. Nephrotic patients of the REIN core study that continued on ramipril for another 2 years (as part of the REIN follow-up study) had a progressive amelioration in the rate of GFR decline up to 1 ml/min/yr (96, 97). Among the 78 patients treated with the ACE inhibitor in this study, 10 patients showed improvement of GFR and never reached ESRD (97). GFR slopes in 16 additional patients stabilized, or worsened so slowly that ESRD would be delayed beyond the patients’ life expectancy. Patients that switched to ramipril from conventional treatments continued to progress on follow-up. Thus, ESRD risk reduction went from 50% in the core study (18 months) to 300% in the follow-up study (3–4 years), a finding consistent with the time-dependent effects of ACE inhibitors (97) in inducing regression of the disease. The time-dependent amelioration of the GFR slopes (Figure 3) was paralleled by progressive reduction in urinary protein excretion.

Long-term follow-up in diabetes. A post-hoc analysis of the Captopril study (S20) on 108 patients with type 1 diabetes and nephrotic proteinuria at study entry found that over a follow-up period lasting more than 3 years, 7 of 42 patients on captopril had full remission of proteinuria. In these patients, renal function stabilized. After an 8-year extended follow-up period, GFR remained stable in 6 patients who still had less than 1 g/24 h of proteinuria (98). These findings challenge the common belief that diabetics with nephrotic-range proteinuria can have inexorable progression to ESRD, showing that remission and even regression of the disease can occur (Table 1). This is one of the first indications that no further renal function loss is possible in these patients, provided that urinary protein excretion can be limited by the treatment (Figure 3).

Animals studies help clarify the significance of human findings
The finding that a low-protein diet or ACE inhibition can reverse proteinuria and glomerulosclerosis is well established in animals (Table 2) (S21, 99–102). When an ACE inhibitor and an Ang II receptor blocker were combined in a genetic model of progressive nephropathy, reduction of glomerular sclerosis was even more evident, particularly in those glomeruli that had less severe lesions to begin with (102). This shows that remodeling of glomerular architecture is possible, which would imply some form of regeneration of the capillary network (103, S22).

Among the mediators involved in the regression of sclerosis (104, 105, S23), plasminogen activator inhibitor-1 (PAI-1) is a plausible candidate, given the role of PAI-1 — which is potently induced by Ang II — in inhibiting matrix degradation (104). PAI-1 (as shown by immunostaining) progressively increases as sclerosis develops, while in rats receiving ACE inhibitors, PAI-1 expression decreased (S23). In all of the studies described above, quantification of sclerosis was based on semiquantitative scores. Yet 2D analysis of a single section of a biopsy specimen overestimates the number of normal glomeruli and underestimates the actual extent of glomerulosclerosis (106, 107). Thus one cannot currently tell what volume of glomerular capillaries is actually affected by sclerosis and to what extent the sclerosis volume is effectively reduced by treatment.

A possible answer from 3D reconstruction studies
To quantify the extension of sclerosis volume and capillary regeneration upon treatment, 3D reconstruction of the entire capillary tuft of hundreds of glomeruli has been performed in a rat model (108). The 3D analysis of the capillary tuft was instrumental to demonstrate reabsorption of sclerosis and regeneration of capillary tissue after administration of a high dose of ACE inhibitor to rats with very advanced nephropathy (Figure 4). The treatment remarkably reduced sclerosis volume in most glomeruli, unless they were already almost totally sclerosed. ACE inhibitors also enlarged the volume of intact capillaries by up to 40% (Figure 5) (108). These structural changes allowed the kidney to regain function with time.

Interpretation of existing data
Mechanisms responsible for sclerosis regression include the inhibition of TGF-β (102) and the decrease in PAI-1 levels (104, S23). However, precisely which cells are involved in the process of scar tissue removal is not yet known. While glomerular, endothelial, and mesangial cells seem to proliferate in some circumstances, it is generally accepted that more differentiated podocytes do not usually proliferate (109, S24), making it unlikely that new segment formation can occur simply by replication of resident cells. Podocytes, however, may promote capillary growth by stimulating proliferation and migration of glomerular endothelial cells (110).

Regeneration of capillary segments in the glomerular tuft may depend on other cells. Bone marrow cells act as a reservoir for glomerular mesangial cells in rodents (111), and cross-bone marrow transplantation from young to old mice allows a partial regression of structural lesions associated with aging (112). Regression of glomerulosclerosis and neoformation of glomerular tissue has indeed been linked to progenitor cells of renal or extrarenal origin (110, 112).

Stem cells also exist in the adult kidney and can theoretically act to repair lesions. A distinct population of progenitor-like cells exists that express vimentin (a marker typically expressed by metanephric mesenchyme during kidney development). These cells are localized in proximal and distal tubuli and peritubular capillaries and can retain mitogenic potential (113). Following ischemia (113), such cells enter the cell cycle, divide, and migrate to the site of damage (113). Renal papilla can also be a niche for kidney stem cells (114) that start proliferating after renal ischemia.

Stem cells of bone marrow origin, both hematopoietic and mesenchymal, also contribute to kidney regeneration (115, 116, S25, S26). Transplanting male bone marrow into female recipients yielded Y chromosome–positive cells that localized and differentiated in tubular epithelium and glomerular podocytes (115).

Mesenchymal stem cells, by promoting resident tubular cell proliferation, limited renal injury and improved renal function in mice with cisplatin-induced acute renal failure (117). That bone marrow or resident kidney cells repair tissue damage is plausible, given results in the heart (118). In the process of cardiac repair, stem cell migration and homing is facilitated by HGF (119). Whether this applies to the kidney has yet to be confirmed (S27). Ang II blockade also limits TGF-β expression. Since TGF-β suppresses HGF (S28), one might speculate that ACE inhibitors exert a beneficial effect by preserving the HGF-dependent pathway of renal repair. Evidence in animals has shown that ACE inhibition prevents glomerular and tubular injury by upregulating renal mRNA levels of HGF (120). HGF could therefore be pivotal in regenerating the kidney owing to its capacity for inducing renal cell proliferation and limiting apoptosis, which adds to its chemotactic effect on stem/progenitor cells.

Future perspectives
It has been found that the repair of renal tissue involves remodeling of the glomerular capillary network (S22). The process of glomerular restructuring can now be analyzed by geometrical computational models. For this type of investigation, last-generation
confocal and multiphoton microscopy would be most suitable. Computer programs can then be used to analyze serial sections of glomeruli and automatically compute capillary network topology. Using these theoretical models, one can estimate the distribution of blood flow and water filtration along the network and the pressure difference across the glomerular membrane in individual capillary segments (121). In addition, graph theory could be used to compare topological differences among individual capillary networks (S29). Identifying differences within glomerular network organization that emerge during sclerosis regression may also be a potential application of the above models and may help to identify changes in cell and gene expression occurring in newly formed capillary segments.

Another challenge for future research is to clarify which cells – whether of bone marrow origin or resident renal stem cells – are involved in sclerosis regression and to what extent, as well as the dynamics of intrarenal events involved in tissue repair. Greater knowledge regarding the regression of kidney lesions and a better understanding of how to influence this process will hopefully reveal new paths toward postponing the progression of renal disease in humans.

Acknowledgments

This manuscript is the collaborative result of the joint effort of a number of outstanding colleagues that reviewed the research activity of the last 20 years at the Mario Negri Institute and Bergamo Hospital. Those scientists to whom the present authors are profoundly indebted are Piero Ruggenenti, Arrigo Schieppati, Norberto Perico, Marina Noris, Marina Morigi, and Mauro Abbate. We thank them for their valuable discussions and review of the manuscript.

Note: References S1–S29 are available online with this article; doi:10.1172/JCI27699DS1.

Address correspondence to: Giuseppe Remuzzi, Mario Negri Institute for Pharmacological Research, Via Gavazzeni, 11, 24125 Bergamo, Italy. Phone: 39-035-319-888; Fax: 39-035-319-331; E-mail: gremuzzi@marionegri.it.


106. Remuzzi, A., Pergolizzi, R., Mauer, M.S., and Ber


120. Matsumoto, K., et al. 1999. Prevention of renal dam-
age by angiotensin II blockade, accompanied by increased renal hepatocyte growth factor in experimental hypertensive rats. Hypertension. 34:279–284.


