Pathogenesis and Management of Acute Kidney Injury in Patients with Nephrotic Syndrome Due to Primary Glomerulopathies

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Abstract: Acute kidney injury in the context of nephrotic syndrome is a serious and alarming clinical problem. Largely, acute kidney injury is a relatively frequent complication among patients with comorbidities while it has been independently associated with an increased risk of adverse outcomes, including death and chronic kidney disease. Nephrotic syndrome, without hematuria or with minimal hematuria, includes a list of certain glomerulopathies; minimal change disease, focal segmental glomerulosclerosis and membranous nephropathy. In the light of primary nephrotic syndrome, pathophysiology of acute kidney injury is differentiated by the nature of the primary disease and the severity of the nephrotic state. This review aims to explore the clinical circumstances and pathogenetic mechanisms of acute kidney injury in patients with nephrotic syndrome due to primary glomerulopathies, focusing on newer perceptions regarding the pathogenesis and management of this complicated condition, for the prompt recognition and timely initiation of appropriate treatment in order to restore renal function to its baseline level. Prompt recognition of the precise cause of acute kidney injury is crucial for renal recovery. Clinical characteristics, laboratory and serological findings along with histopathological findings, if required, will reveal the implicated pathway leading to individualized approach and management.

Keywords: glomerular diseases; acute kidney injury; nephrotic syndrome

1. Introduction

Acute kidney injury (AKI), in the context of nephrotic syndrome (NS), is a serious and alarming clinical problem. Largely, AKI is frequent among patients with comorbidities, while it has been correlated with an increased frequency of adverse outcomes, including death [1–3], and chronic renal failure [4]. Specifically, death rates have been reported to increase with AKI severity from 15.9% in stage I, to 49.3% in dialysis-dependent patients [3]. Of those who become dialysis-dependent during hospitalization but come off dialysis and are discharged subsequently, 5%–20% will remain dialysis-dependent [4]. The risk of developing end stage kidney disease has been reported to increase 2-fold by one year for patients who recover from AKI within 10 days [5], while the risk further increases with the number of AKI episodes [6–8]. Patients with glomerular diseases are prone to develop AKI, either as a result of the aggressive form of the disease, or in relation with factors surrounding the pathophysiology of the glomerular lesion and its treatment. Nephrotic syndrome (without hematuria or with minimal hematuria) includes a list of many diseases; with respect to primary glomerulopathies, the commonest are minimal change disease, focal segmental glomerulosclerosis and membranous
nephropathy. Acute kidney injury is classified by the main pathophysiologic mechanism involved in prerenal, intrinsic renal and postrenal causes. Theoretically, any of these scenarios may occur in patients with NS due to primary glomerulopathy, although the prerenal and intrinsic ones are the most frequent seen in this setting.

This review aims to explore the clinical circumstances and pathogenetic mechanisms of AKI in patients with NS, focusing on newer perceptions regarding the pathogenesis and management of this complicated condition, for the prompt recognition and timely initiation of appropriate treatment in order to restore renal function to its baseline level.

2. Definitions and Epidemiology of AKI

Acute kidney injury (AKI) is defined by an unexpected decrease in the glomerular filtration rate (GFR), leading to the retention of nitrogenous waste products, as well as the dysregulation of extracellular volume and electrolytes. According to the Kidney Disease Improving Global Outcomes (KDIGO) [9] definition, AKI is characterized by an increase in serum creatinine by ≥0.3 mg/dL within 48 h or to ≥1.5 times baseline, which is known or presumed to have occurred within the past seven days, or urine volume <0.5 mL/kg/h for six hours [9]. Correction of volume depletion and exclusion of obstructive uropathy by ultrasonography are typically preceded. Depending on the time and degree of creatinine increase and urine output volume, AKI is staged in three grades [9]. However, this criteria system does not distinguish between the multiple etiologies that cause AKI while the criterion which is referring to, the urine volume, is controversial as it is not based on robust evidence [10–20].

The precise frequency of AKI in the general population is not known, probably due to differences in the definitions and cohorts of patients enrolled across studies. A meta-analysis of 143 studies, in more than 3.5 million hospitalized patients, applying the KDIGO definitions, found that 22% of the patients experience this complication [3], increasing to 57.3% for the intensive care units [21]. The incidence of AKI has been shown to increase in recent years [22] with standard risk factors including pre-existing kidney disease, diabetic nephropathy, volume depletion, effective arterial volume depletion.

3. Frequency of AKI in Patients with NS

A modest reduction of GFR of about 30% is frequent among patients with NS, as shown in a series of 89 adults with biopsy-proven minimal change disease. In this series, diminished renal function was recorded in 22.5% of patients, typically aged over 60 years [23]. Similar findings have been found in other studies in adults and children [24–27]. Studies in children with NS have shown a close correlation of GFR alteration with both the reduced filtration fraction and the decreased serum albumin concentration that are relative to the severity of the nephrotic state. However, the frequency of AKI across patients with NS varies depending on the histopathological type and surrounding factors related to the patient and the management of NS itself. Smith et al. studied a series of cases with severe AKI which occurred in the clinical setting of idiopathic NS and found that, in 60% of them, certain abnormalities in the tubule-interstitial compartment were suggestive of acute tubular necrosis [28]. In children, hospitalized with NS, an incidence rate of 8.5% AKI has been reported [29,30]. More recently, there have been reported rates of AKI of 58.6% for 336 children and 50.9% for 615 hospitalizations [30]. After adjustments, factors such as infection and nephrotoxic medication exposure, including duration and intensity of exposure, were shown to be significantly associated with AKI in those patients. In adults, AKI rates have been reported to occur in 25%–35% of patients with minimal change disease, which in many occasions occur as part of the initial clinical presentation [28,31–34]. Adults with minimal change disease, who develop AKI, are more likely than those who do not develop AKI to be older, male, hypertensive, and to have more severe NS [35,36]. Waldman et al. [31], reported that AKI occurred in 25% of patients. In 70% of them, it was concurrent with the initial diagnosis of minimal change disease, while one third of them followed a disease relapse. Likewise, a study of 125 adults and adolescents from the Netherlands recorded AKI in 40% of them [36]. Patients with AKI in this context usually present with significant proteinuria, hypoalbuminemia, and edema. Acute kidney injury, in this
context, occurred a few months after the NS diagnosis and the majority of patients had hypertension and massive edema at presentation [28]. Remarkably, most patients were also oliguric, and nearly 20% of them required dialysis or died from intercurrent complications. In general, GFR has been reported to be well-maintained during the follow-up, although serum creatinine was higher in those who had presented with AKI [34]. There are less reports from patients with membranous nephropathy [37] or focal segmental glomerulosclerosis [38], probably due to the lower frequency of this complication among these patients. Notably, among 235 elderly patients, in whom the most frequent indication for renal biopsy was AKI (46%), 9.4% of them presented with NS and AKI [37]. The most frequent diagnosis was minimal change disease, which was found in one-quarter of these patients.

4. Pathophysiology of NS

The historic definition of nephrotic syndrome requires the presence of triad proteinuria, typically 3.5 g per day or more, lipiduria and hyperlipidemia, and edema. A proportion of patients may also have microscopic hematuria, although typically the urine sediment is bland. The main abnormality in NS is the loss of protein in urine. Inability to reabsorb the filtered protein may result in proteinuria, i.e., 0.5–2.5 g/day, in patients with a normal GFR [39]. If proteinuria is more than 2–2.5 g/day it implies that at least part of the lost protein is due to increased glomerular permeability [39]. The type of urinary protein may provide some information regarding the stage of the glomerular lesion. For instance, highly selective proteinuria (only very small molecules are filtered) is associated with better histopathological findings. As a result of the significant protein excretion, comes hypoalbuminemia [40]. In response to the low serum albumin level, the liver increases not only albumin formation but also lipid production. Thus, low density lipoprotein cholesterol, low-density lipoproteins and apolipoproteins B, CII, and CIII, are found to be increased in NS [41–45], primarily as a consequence of altered colloid oncotic pressure and triglycerides being over-produced [43–45]. Edema formation in patients with NS is explained by the “underfill hypothesis” in which the kidney is retaining sodium and water in order to fix intravascular hypovolemia, a condition which ends up in the development of edema, due to the forces of Starling [46–48]. Conversely, the “overfill hypothesis” pertains to renal sodium retention, which occurs as a primary phenomenon which increases blood volume [49,50]. However, the primary defect, as noted by Rostoker et al., is the abnormal glomerular permeability, a problem which is often remitted after treatment with glucocorticoids [51].

Hypercoagulability is another major complication of NS, which is occurs secondarily to impaired coagulation and fibrinolysis [44]. Plasma viscosity increases in conditions with reduced blood flow, increased interstitial pressure and endothelial injury [52]. Loss of antithrombin III is also particularly important since patients who lack this factor have a 50%–70% risk of experiencing thromboembolic events [53]. Citak et al. [53] showed that pediatric nephrotic patients who experienced thrombotic events had very decreased antithrombin III and increased fibrinogen when compared with those without any thrombosis. Moreover, factors XI, XII, and plasminogen [52,54–56] are lost in urine in the cases with massive proteinuria. Yet, procoagulant factors, including factors V, VII, VIII, and the von Willebrand factor, have been reported to increase due to hypoalbuminemia, because they are normally bound to albumin [52,55]. Additional problems in nephrotic patients include increased susceptibility to infection and atherosclerosis, malnutrition, hormone dysregulation and a vitamins and metals deficiency [42].

5. Pathophysiology of AKI in Patients with NS

5.1. Acute Tubular Necrosis

Acute tubular necrosis in patients with NS is a sporadic but significant complication. It occurs due to hypovolemia in combination with hypoalbuminemia, often related to excessive diuresis. Hypovolemia may be the result of aggressive diuresis and the finding in physical examination of generalized edema. Undoubtedly, aggressive diuresis is necessary in cases with massive edema,
especially if it is related with inability to walk, dyspnea, pleural effusions, and/or ascites. Interestingly, despite the marked expansion of the extracellular fluid volume, such patients may report symptoms of decreased effective circulating volume, including tachycardia, low blood pressure and oliguria. In most patients with NS and acute tubular necrosis, the pathophysiology of acute tubular injury and renal impairment are explained through the changed hemodynamics. Although the blood volumes may be preserved, the oncotic pressure remains significantly low in many patients with NS and thus, intravascular volume contraction is a key factor [32,57–59]. Other mechanisms which may cause renal dysfunction include low renal perfusion pressure, cast nephropathy, i.e., tubular obstruction with protein cast and interstitial edema [28,60]. In cases with minimal change disease and AKI it has been shown that the administration of albumin solutions inclined renal plasma flow although this finding has been disputed by others [25,61,62]. Furthermore, reductions of the epithelial slit pore length may be critical for the glomerular capillary permeability to water and small solutes while, in these patients, acute tubular injury is manifested by the dilation of tubular lumens with flattening of epithelium, sloughing of epithelial cells into tubular lumens, and interstitial edema with a few scattered inflammatory cells. Exposure to non-steroidal anti-inflammatory drugs (NSAIDS) also contribute to the development of glomerular lesions in minimal change disease. Diffuse foot process effacement of podocytes is a typical finding with the absence of brush border and basolateral infoldings in the proximal tubules and negative immunofluorescence or scattered IgM mesangial staining [63–65] (Figure 1).

Figure 1. Segmental glomerular sclerosis with hyalinosis, as well as tubular dilation, tubular cell loss and detached epithelial cells into tubular lumens, in a patient with severe nephrotic syndrome due to focal segmental glomerulosclerosis (H&E X400).
Contrast Nephropathy

Nephrotic patients may also develop AKI after the administration of iodinated contrast media-enhanced radiographic procedures. The precise mechanism of acute tubular necrosis in cases with contrast nephropathy remains not well understood. Renal vasoconstriction and thus hypoxia in the renal tissue due to impaired viscosity, as well as a straight cytotoxic consequence of the contrast itself on the tubular cells [63–65] are considered as the primary factors. It is also unclear why the duration of acute tubular necrosis is shorter in cases with contrast induced nephropathy compared to acute tubular necrosis from other causes. It has been postulated that acute renal dysfunction comes as a result of functional changes rather than necrosis and thus, recovery occurs quickly. Tao et al., in studying a large cohort of matched patients, found that the risk of AKI, after the administration of intravenous contrast during computed tomography, was not higher in nephrotic patients [66]. Diagnosis of AKI in such cases is based upon the characteristic rise decrease in GFR within 24–48 h after the initiation of the contrast agent. Exclusion of other causes of AKI, using microscopic urinalysis and rarely a renal biopsy, is always required, while it should be differentiated from thrombotic complications of NS. Despite the fact that it is a reversible condition, it may occasionally be related to significant morbidity.

5.2. Acute Interstitial Nephritis

Acute interstitial nephritis is another setting of AKI, which is clinically characterized by the unexpected decline in renal function in combination with inflammation and edema of the renal interstitium. Suspicion and/or histopathologic documentation of acute interstitial nephritis in patients with NS should lead to a careful search for an etiologic agent, including infections, previous or concurrent administration of medicines such as antibiotics, NSAIDS, immune or neoplastic disorders. The most common antibiotics which may cause acute interstitial nephritis include cephalosporins, ciprofloxacin, ethambutol, isoniazid, macrolides, penicillin, rifampin, sulfonamides, tetracycline and vancomycin. Histologically, interstitial nephritis associated with NSAIDS is characterized by diffuse inflammatory infiltrates in the interstitium. The clinical picture typically includes renal insufficiency, low grade fever, rash and arthralgias. Often, some of the clinical sighs/symptoms are not present. NSAIDS have been known for their nephrotoxic potentials including their correlation with minimal change disease and with interstitial nephritis. They act through the inhibition of cyclooxygenase, a prostaglandin synthase, which is involved in the transformation of arachidonic acid to prostaglandins, prostacyclin, and thromboxanes [67,68]. More recent agents, such as selective cyclooxygenase-2 inhibitors, have anti-inflammatory and analgesic effects similar to classic NSAIDs with fewer adverse events. However, the risk of AKI appears not different to the use of different NSAIDs [69]. Risk factors for AKI induced by NSAIDs include history of chronic kidney disease, age, volume depletion and decreased effective arterial volume of any cause, other medications, i.e., diuretics and angiotensin-converting enzyme inhibitors or angiotensin receptor blocker, dosage and duration of therapy with NSAIDS [70].

5.3. Renal Vein Thrombosis

Thromboembolic events and renal vein thrombosis represent one of the most serious complications of NS and thus, AKI may occur, especially in cases with bilateral thrombosis. Estimates of the frequency of renal vein thrombosis among nephrotic patients range from 5%–60%, with an overall incidence of 35% [71]. This wide-ranging variability in the frequency of thrombotic events in NS is probably related to the characteristics of the patients included in the studies, i.e., symptomatic or clinically silent thrombotic events, degree of proteinuria, as well as the methods used for detection of thrombosis. In studies of consecutive or unselected patients without membranous nephropathy, who underwent venography, the prevalence of renal vein thrombosis ranges from 10% to 50% [72]. In a prospective study of 151 nephrotic patients, renal vein thrombosis was diagnosed in 22% of them, while one
third of them had membranous nephropathy [73]. Overall, patients with membranous nephropathy and NS appear to be in a higher risk for thrombosis, when compared with nephrotic patients with other glomerular diseases [72–79]. In this regard, in a large cohort of 1313 patients with primary glomerular diseases, it was shown that the incidence of venous thromboembolic events was much higher among patients with membranous nephropathy (7.9% vs. 3% and 0.4% respectively). As a result, the histopathologic diagnosis was shown to be predictive for thrombosis occurrence after controlling for the 24 h proteinuria and serum albumin levels [74]. The risk of thrombosis has also been found to increase by the severity of hypoalbuminemia, a phenomenon, which is significant in patients with membranous nephropathy, as shown in a cohort of 898 patients with biopsy-proven disease [75]. The precise pathogenic mechanism is not known. The hypercoagulability disturbances and decreased fibrinolysis have been noted to be greater in patients with membranous glomerulopathy vs. those with other types of lesions [80,81]. In addition, the reduction in plasma volume may be more significant in membranous nephropathy [71]. A study which studied nephrotic patients specifically, who carried the diagnosis of membranous nephropathy and also experienced renal vein thrombosis, found immune complexes in their circulation which were not found in patients with membranous nephropathy without renal vein thrombosis [82]. It is speculated that these complexes may trigger the activation of the coagulation process directly or may activate factor XII [83]. Clinically, renal vein thrombosis may follow different scenarios, including abrupt onset of pain, renal dysfunction and macroscopic hematuria [84], although it may also quite often be silent [85]. In a prospective design, patients with renal vein thrombosis in the context of NS were noted to have two distinct types of clinical presentation; acute and chronic [73]. The setting of acute renal vein thrombosis includes a history of acute flank pain, hematuria, abnormal intravenous venogram and decline of GFR. Chronic renal vein thrombosis is more often asymptomatic. The significant incidence of renal vein thrombosis in nephrotic patients and especially with membranous nephropathy, together with the lack of symptoms, which is common, raises the question of how to handle these patients diagnostically, given the absence of information in asymptomatic patients and the fact that renal venography is an invasive procedure, not free of complications [71]. The use of ultrasonography, spiral computerized tomography with contrast and images by magnetic tomography may be the first step of the workup, as non-invasive procedures. However, the gold standard diagnostic test remains selective renal venography and thus, if any of the above reveals a high suspicion of thrombosis, then a renal venogram should be performed. Routine screening for renal vein thrombosis is not suggested for patients with NS, but selective high-risk patients, i.e., who have full blown NS with severe hypoalbuminemia (i.e., below 2 g/day) with symptoms and/or renal dysfunction should be investigated [86]. If a nephrotic patient experiences signs or symptoms of a renal infarct accompanied by acute renal failure, then acute complete renal vein thrombosis is suspected and thus, a selective renal venogram should be performed, which might be combined with a simultaneous therapeutic procedure. Renal biopsy is of great importance to determine the cause of renal vein thrombosis, if not known. Membranous nephropathy, membranoproliferative glomerulonephritis and minimal change disease are the leading causes, although any other diseases with NS can also present with renal vein thrombosis. In some instances, fibrin thrombi in blood vessels or glomerular capillaries may be seen while prominent congestion of glomerular or interstitial capillaries and a disproportionate degree of interstitial edema should always raise the possibility of renal vein thrombosis. Patients who receive a diagnosis of renal vein thrombosis, should be started on treatment with anticoagulatives immediately. Patients with symptomatic renal vein thrombosis are treated with unfractionated or low molecular weight heparin and then warfarin with a goal international normalized ratio of 2–3 [87–90]. Occasional patients with NS appear somewhat unaffected by heparin therapy, a fact which is related to severe anti-thrombin deficiency. The use of oral and parenteral direct thrombin inhibitors and factor Xa inhibitors have not been studied in patients with NS or impaired renal function, since such patients have been excluded from the related trials. Therapy with warfarin is given for a minimum of 6–12 months and should be continued as long as the patient remains nephrotic [71,86]. The use of oral and parenteral direct
thrombin inhibitors and factor Xa inhibitors have not been studied in patients [91] with NS while patients with impaired renal function have been excluded from the studies of these agents. Systemic fibrinolytic therapy is not recommended in patients with AKI due to renal vein thrombosis, since it carries significant risk for bleeding, especially intracranial [92–94]. Anticoagulation for asymptomatic renal vein thrombosis may be identified through screening at the time of diagnosis of NS, which is not recommended without any clinical and/or laboratory indication or in case of other imaging studies. Besides, there is no data regarding the role of anticoagulation in patients with NS with asymptomatic renal vein thrombosis except from limited data reported in case series [73,88]. The need of prophylactic anticoagulation in cases with NS depends upon the incidence of thrombotic events in the patients, the effectiveness of anticoagulative therapy to prevent them and the associated risk of bleeding with this kind of treatment. In this regard, the data from prospective or controlled studies, comparing the risks associated with undiagnosed venous thrombosis with the risk of long-term anticoagulation, are limited. However, we estimated [95] the frequency of venous thromboembolic events in an inception cohort of 898 patients with primary membranous nephropathy versus the risk of bleeding using data from a systematic review of the literature. The probability of benefiting from a prophylactic regimen was evaluated by the risk of bleeding and the level of serum albumin. Patients who were scored with a high risk of bleeding, were not able to receive prophylactic anticoagulation and benefit of it irrespective of the level of serum albumin. In light of clinical practice and in order to assist the decision for prophylactic coagulation initiation in these patients, a tool was created to estimate the probability of benefit based on an individual’s bleeding risk, serum albumin level, and benefit-to-risk ratio (http://www.gntools.com) [95].

5.4. Warfarin (Anticoagulant) Associated Nephropathy

Another type of kidney injury, which comes as a result of overdosed anticoagulative therapy, has been described as anticoagulant related nephropathy. It represents a type of AKI which is a consequence of warfarin overdose [96,97]. Histologically it is characterized by acute tubular lesions and obstructed tubules with red blood cells casts [98]. This finding was documented in a series of nine biopsy proven cases who developed unexplained AKI following warfarin overdose. The precise incidence of this disorder is unknown, but it has been estimated that 17% of patients with international normalized ratio above 3 and no underlying kidney disease may experience an elevation in serum creatinine [99,100]. However, no histological data were revealed in this study and other causes of AKI might have been involved. Reasonably, nephrologists are happy with the risk which comes with a kidney biopsy in patients who need anticoagulation. One concern is the risk of thrombosis during the period anticoagulation is held, and the other the risk of hemorrhage from the kidney biopsy site after the systemic anticoagulation is resumed. The main diagnostic criterion for anticoagulant related nephropathy is glomerular hemorrhage following excessive anticoagulation. Notably, patients with abnormal glomerular membranes have been shown to be more vulnerable to this condition [100,101]. Glomerular bleeding leads to the formation of red blood cells casts, which cause obstruction in the renal tubules. This is the most striking histologic feature, as shown from specimens obtained from patients [98] and animal models [102–104]. Yet, the percentage of completely obstructed tubules seen by microscopy cannot always explain the severe deterioration of GFR [100,101,104]. It is possible that the small kidney specimen taken by a biopsy may not be representative of the real lesion. Besides, it has been hypothesized that another type of tubular epithelial cell injury may be involved, similar to the one which has been observed in patients with immunoglobulin A nephropathy or paroxysmal nocturnal hemoglobinuria [105,106]. Oxidative activity of heme and iron may be implicated in this type of tubular injury [107] or the anticoagulative agent itself may be directly toxic to the renal epithelial cell [104]. The major risk factor for developing anticoagulant related nephropathy is coagulopathy from warfarin or other anticoagulants use, although it is probably a multifactorial condition, since factors other than the degree of coagulopathy may be present and play a role in the development of AKI [98], including chronic renal impairment, diabetes mellitus, heart failure, arterial hypertension and
glomerular disorders [98]. The clinical presentation is characterized by hematuria and subsequent renal dysfunction which usually occurs within 8 weeks from warfarin initiation [98]. From observational studies and animal models it has been shown that a few days with warfarin induced coagulopathy is enough to cause anticoagulant related nephropathy. Gross hematuria is less common than microscopic hematuria [97, 98]. Absence of hematuria has also been reported, which is explained by the hypothesis that hemorrhaging glomeruli are shut down by the red blood cells casts and AKI, by the time these patients seek medical attention.

5.5. Aggressive Disease

Of all primary glomerulopathies, only the collapsing variant of focal segmental glomerulosclerosis may follow a course of rapid progression due to the disease itself. Focal segmental glomerulosclerosis is manifested by segmental glomerular sclerosis, usually associated with hyalinosis. Four distinct variants have been described, with the collapsing variant to be more ominous. It is seen in human immunodeficiency virus (HIV) patients and in cases with idiopathic focal segmental glomerulosclerosis. It is characterized by hyperplastic and hypertrophied podocytes with vacuoles and periodic acid shiff positive cytoplasmic droplets in association with glomerular tuft collapse and thus, often is exhibiting severe acute tubular injury and deteriorating renal function. Focal or more commonly widespread changes in the tubules can also be seen. Tubules are dilated and filled with pale-staining luminal precipitate. It has been historically associated with a significant and quick decline of GFR [108–110]. In a study of 62 patients, with this type of lesion, the collapsing group comprised of more African Americans (82.8%) and more women [111]. Patients with collapsing focal segmental glomerulosclerosis presented with more severe renal dysfunction and lower serum albumin level [111, 112]. Primary collapsing focal segmental glomerulosclerosis represents an active, aggressive form of podocyte injury. Thomas et al. used a scoring system to describe specific pathologic findings where patients with collapsing focal segmental glomerulosclerosis had the highest total injury score [113]. Collapsing focal segmental glomerulosclerosis is most often seen in adults, with more males are affected, although this is not different from patients with focal segmental glomerulosclerosis without the collapsing lesion [114]. Patients with collapsing focal segmental glomerulosclerosis experience a short course from onset of symptoms to biopsy [109]. At presentation, patients have significantly higher levels of protein excretion with many patients excreting more than 10 grams per day. In addition, the time from onset of symptoms to renal biopsy has been shown to be significantly shorter, suggesting that a more fulminant disease course has been preceded prior to kidney biopsy. Weiss et al. have reported on six African Americans with renal insufficiency, massive proteinuria and renal biopsy showing glomerular collapse [115]. Laurin et al. however [111], in a retrospective study of patients, who received a diagnosis of collapsing focal segmental glomerulosclerosis between 1989 and 2012, assessed the rates of treatment response after adjusting for baseline characteristics and the type of immunosuppressive therapy. According to their results, the collapsing variant was not associated with a significantly worse renal survival, and probably the most important factor was shown to be the prompt diagnosis and initiation of immunosuppressive therapy in these patients [111].

Nevertheless, treatment of these cases is a challenge for clinicians as there are no prospective controlled trials and thus any approach is based principally upon clinical experience. The reported number of HIV-negative patients with collapsing focal segmental glomerulosclerosis, who have undergone complete remission in response to therapy is relatively small. A course of oral glucocorticoids for six months may be given as first line therapy (prednisone 1 mg/kg BW every day or 120 mg every other day for the first two months), then subsequently tapered over a minimum period of 6 months. We also administer antimicrobial prophylaxis against Pneumocystis pneumonia for the duration of glucocorticoid treatment. In cases with significant amounts of proteinuria, severe hypoalbuminemia and conserved GFR, a calcineurin inhibitor is added to the glucocorticoid regimen to accelerate reversal of the hypoalbuminemia and proteinuria. Notably, a combined treatment with glucocorticoids and other immunosuppressive agents was not related with better outcomes than therapy with glucocorticoids.
alone. A study which was performed retrospectively, and included 275 patients with biopsy proven focal segmental glomerulosclerosis, treated with immunosuppressive regimens, showed that for the collapsing variant there was no difference in the adjusted risk of end-stage renal disease between treatment with calcineurin inhibitors (with or without glucocorticoids) compared to treatment with glucocorticoids alone [111]. Besides, regimens including cyclophosphamide or chlorambucil were associated with unsatisfactory effectiveness and significant side effects [116].

5.6. Crescentic Glomerulonephritis Superimposition

Occasionally, patients with NS may develop AKI as a result of disease transformation to rapidly progressive glomerulonephritis with crescent formation. This infrequent clinical scenario has been reported in patients with already diagnosed membranous nephropathy who presented with a rapid decline in renal function and switch from the nephrotic syndrome clinical setting to nephritic syndrome. In these cases, a new biopsy revealed that anti-glomerular basement membrane or anti-neutrophil cytoplasmic antibodies (ANCA) associated glomerulonephritis was superimposed on membranous nephropathy [117]. The first case of combined membranous and crescentic glomerulonephritis was reported by Klassen et al. [118] and was associated with anti-glomerular basement membrane antibodies. The author suggested that membranous nephropathy primarily damaged the glomerular basement membrane and released antigens that incited the anti-glomerular basement disease. Pettersson et al. reported a series of 17 patients with this kind of mixed lesion. Of those patients with follow up, 12 ended up in end stage renal disease, and the remaining responded to therapy with variable degrees of renal dysfunction [115]. Another mechanism of crescent formation in the context of membranous nephropathy has been associated with ANCA disease. In additional to other cases reported in the literature, Nasr et al. [119] described 14 cases with simultaneous diagnosis of ANCA-associated glomerulonephritis and membranous nephropathy. In 13 of them, patients with this mixed lesion was revealed at the first clinical presentation by histopathology while in 1 patient, membranous nephropathy preceded ANCA associated glomerulonephritis by 7 months. The majority of patients in this series presented with rapidly progressive renal dysfunction and nephrotic range proteinuria. The mean 24 h urine protein was 6.5 g, and 9 had full blown NS [119]. Thus, the development of crescentic glomerulonephritis in a background of a primary glomerulopathy should be included in the differential diagnosis of a patient with AKI. A second kidney biopsy is mandatory for these cases in order to explain AKI and make the final diagnosis. Besides, speedy diagnosis and therapy initiation is fundamental in all cases with rapidly progressive glomerulonephritis. Immunosuppressive therapy pertains to induction of remission with glucocorticoids and cyclophosphamide [120,121], given as intravenous pulses of methyl-prednisolone (7 mg/kg for 3 consecutive days) followed by oral prednisone (1 mg/kg for a minimum of 4 weeks), subsequently reduced over the next 3–5 months [122]. Cyclophosphamide is administrated, either intravenously as monthly pulses (0.5–1) g/m² body surface are, or orally (2 mg/kg/day), adjusted by the leukocyte count. Additional interventions, such as plasma exchange [123] and/or administration of the anti-CD20 agent rituximab depends on the severity of renal impairment, the immunological environment surrounding this complication i.e., ANCA positive, anti-glomerular basement membrane antibody positive or pulmonary hemorrhage. Importantly, speed in diagnosis and early induction of immunosuppressive therapy are fundamental, because the type of lesion may follow a catastrophic course, leading to end stage renal histopathology within a few weeks (Figure 2).
While AKI from calcineurin inhibitors is a reversible condition following a prompt decrease in blood pressure, with cyclosporine is a fact. However, usually it is a reversible effect in case of long-term therapy. Factors predisposing to it include high dosing, older age, co-administration of NSAIDs, hypovolemia or excessive use of diuretics, exposure to medications which inhibit cytochrome P-450 3A4/5 (CYP3A4/5), thereby increasing exposure to calcineurin inhibitors metabolites, or drugs that inhibit P-glycoprotein-mediated efflux of calcineurin inhibitors from tubular epithelial cells, thereby increasing local renal exposure to calcineurin inhibitors. Also, genetic polymorphisms in the genes encoding CYPS4A/5 and P-glycoprotein have been identified to play a role in calcineurin inhibitor metabolism [130]. Patients with AKI related to calcineurin inhibitors toxicity should be checked for all the above risk factors in order to help quick recovery. While AKI from calcineurin inhibitors is a reversible condition following a prompt decrease in blood level, if it remains long-term it may turn to a chronic progressive renal disorder, which is a permanent condition [131,132]. Although most come from the use of cyclosporine in kidney transplant recipients, a similar pattern of renal injury has been documented for tacrolimus as well. A retrospective study in
16 children treated with calcineurin inhibitors for steroid resistant NS found that [133] AKI was very common (81.3%) counting for 0.7 AKI episodes per patient year of follow-up [133]. Yet, when the authors restricted the rate to patients who have achieved remission, the number was 0.34 episodes of AKI per patient year. Among those, 71% of AKI episodes recovered to the baseline serum creatinine value, while in 29% a new baseline serum creatinine was set. Another study of 119 children with idiopathic NS and AKI [10] reported drug toxicity as the second most frequent cause [134]. In contrast, [135] a retrospective study of 20 patients with steroid-resistant NS, who received cyclosporine A for 5 years, showed that, although there was an initial drop in GFR, renal function remained stable afterwards and no one developed chronic renal failure. Several studies with cyclosporine and tacrolimus in kidney transplantation demonstrated a relationship between whole blood or plasma cyclosporine concentrations and acute nephrotoxicity that was recovering with dose reduction [136,137]. Thus, patients with AKI attributed to calcineurin toxicity should be managed with close monitoring and diminishing of the daily dosage. Finally, identification of the genetic polymorphisms related with the drug-metabolizing enzymes might be useful in the assessment of individualized dosage algorithms for calcineurin inhibitors, as are substrates for CYP3A5 and P-glycoprotein [138], in order to avoid AKI renal dysfunction episodes.

6. Management of Patients with AKI in Patients with NS

General measures for the management of AKI include fluid management in order to restore extracellular volume and increase the effective circulating volume. The initial approach however is to assess the volume of the patient and determine if he is euvolemic, hypervolemic or hypovolemic. This assessment, which is based on accurate records of fluid input and output, daily body weights, physical examination, blood pressure and pulse measurements, will subsequently guide clinical practice in terms of fluid administration or not.

Replacement of albumin with or without diuretics in patients with NS has been a debate over the past decade. Nevertheless, long standing clinical experience has shown that treatment with diuretics as well as albumin administration results in diuresis and natriuresis in patients with edema and hypoalbuminemia. However, some patients do not respond to diuretics as expected, even with high doses of one drug or combined diuretic therapy. This phenomenon is probably due to the fact that the real intravascular compartment is relatively small while the neurohumoral systems, including the sympathetic nervous system, the antidiuretic hormone and renin-angiotensin II-aldosterone axons have been activated. In this regard, it has been shown that albumin infusion [139] increases the intravascular volume by witholding movement of liquids to the third space compartment. Yet, routine use of albumin solutions in NS with normal or increased plasma volume, is doubted. Furthermore, furosemide is highly bound to albumin (>90%), and consequently is not filtered at the glomerulus, while it is secreted in the proximal tubule. If the patient is severely hypoalbuminemic (<2 g/dL) [140], the proportion of the free drug which is available is bigger. Consequently, it diffuses into the tissues, with expansion in its volume of distribution, resulting in less delivery to the proximal tubule for secretion into the lumen. As a result, many clinicians end up combining loop diuretics with albumin to induce diuresis, natriuresis, and improve edema. The combination of furosemide and albumin is considered to restore, at least in part, the intravascular volume [141]. If furosemide is given in the same solution with albumin, it is considered even better for its secretion in the proximal tubule. However, even if furosemide is secreted in the proximal tubule, an amount of it is bound to the filtered albumin in the lumen. As a result, the free drug delivery to limb of Henle’s loop is further decreased, contributing to diuretic resistance. Hypertrophy of the distal tubular cells due to increased reabsorption of delivered NaCl have also been associated with resistance to diuretics [142]. Since the studies exploring the usefulness of combining furosemide and albumin to treat edema in patients with NS [142] are controversial, and the cost of albumin administration is too high, we suggest that the management of these patients should be individualized, and that combination therapy should be restricted to patients with resistance to therapy of more than one diuretic [142].
Serious and potentially life-threatening complications due to fluid overload, such as pulmonary edema, heart failure, hypertension, and being refractory to diuretic therapy, often require dehydration through dialysis sessions. In addition to hypervolemia, electrolyte abnormalities, metabolic acidosis and uremia may require renal replacement therapy temporarily, in cases with oliguria, especially in patients with acute tubular necrosis. In this setting, dialysis is continued until the regeneration of the tubular epithelium is evident. Pharmacologic management is also essential in these patients in order to avoid nephrotoxic medications and also adjust all renally excreted drugs to prevent the worsening of AKI. After the initial adjustment, which is made based on the calculated GFR the list of medications should be revisited regularly throughout the course of AKI and readjustments should be made as warranted if renal function improves or declines. In addition, levels should be routinely monitored for nephrotoxic medications with narrow therapeutic range such as aminoglycosides. Specific therapies depending on the etiology of AKI in patients with NS include anticoagulatives for renal vein thrombosis or immunosuppressive schemes for rapidly progressive glomerulonephritis as described above. In the latter case, early recognition and diagnosis by quick performance of kidney biopsy, is fundamental for its management with prompt initiation of immunosuppressive therapy (Figure 3).

Prophylactic measures for patients with NS in order to avoid AKI are generally limited and include the prevention of extreme restrictions in hydration or aggressive diuresis if not necessary, prophylactic anticoagulation, which may be given individually [95], according to the severity of NS, the specific features of each patient including age, coagulative disorders, hematocrit and propensity for hemorrhage and avoidance of nephrotoxic medications. Albumin replacement may be required in certain circumstances to avoid unwarranted intravascular hypovolemia.
Figure 3. Algorithm for patients with AKI in the context of nephrotic syndrome (NS) due to primary glomerulopathies.

AKI in patients with NS due to primary glomerulopathies

**Pathophysiologic mechanism**

- Acute tubular necrosis
- Acute interstitial nephritis
- Renal vein thrombosis
- Warfarin associated nephropathy
- Drug toxicity
- Aggressive disease
- Crescentic GN superimposition

**Clinical, laboratory and serological characteristics**

- Hypovolemia, hypoalbuminemia, low effective blood volume (h/o aggressive diuresis, contrast agent use etc)
- Fever, rash, arthritis, leukocyturia without bacteria
- Severe hypoalbuminemia, anemia
- Extra-glomerular hematuria, INR>4
- Specific medication exposure, i.e calcineurin inhibitors
- Worsening nephrotic syndrome
- New onset or worsening glomerular hematuria, (+) ANCA or anti-GBM abs.

**Diagnosis**

- Exclusion of other causes, urine sediment with brown casts
- Kidney biopsy
- Computed or classic angiography
- History of warfarin use, INR>4, urine sediment with RBC casts exclusion of other causes
- Kidney biopsy
- Exclusion of other causes, kidney biopsy
- Kidney biopsy
Author Contributions: Conceptualization, S.L.; methodology, S.L.; validation, S.L. and G.L.; investigation, S.L. and G.L.; resources S.L.; data curation, S.L. and G.L.; writing—review and editing, S.L., G.L., J.N.B.; writing S.L., G.L.; visualization, S.L., J.N.B.; supervision, J.N.B.

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