Frequency and clinical histological analysis of glomerular diseases in a tertiary hospital in southern Brazil

Frequência e avaliação clínico-histológica das doenças glomerulares em um hospital terciário da região Sul do Brasil

Abstract

Introduction: The glomerulopathies are the most common biopsy-proven kidney diseases. The epidemiological investigation of glomerulopathies allows the identification of their distribution and main causes and enables the development of prevention and treatment strategies. Objective: This study aims to identify the frequency and clinical-pathological correlation of glomerular diseases diagnosed at the HC-UFPR over the period of 5 years. Methods: 131 biopsies were performed between January 1, 2008 and December 31, 2012 and were analysed by light and immunofluorescence microscopy. Histopathological slides were reviewed by a pathologist. Clinical and laboratory data and the immunofluorescence microscopy results were extracted from medical records. The findings were tabulated and analysed. Results: 128 of 131 cases were reanalysed. 46.5% were obtained from men. Patients’ age averaged 43 years for men and 38 for women. In 99 cases, the indication of biopsy was identified; 49.5% cases presented nephrotic syndrome, 17.17%, acute renal failure and 15.15%, chronic renal failure; 8.08%, nephritic syndrome; 6.06%, isolated proteinuria and 4.04%, isolated hematuria. In 61.21% an underlying disease related to the glomerulopathy could be identified; 33.62% corresponded to primary disease and in 5.17% of cases the nature of the glomerulopathy could not be determined. Among secondary glomerulopathies, the most frequent was Lupus Nephritis (49.29%), and among the primary, Focal Segmental Glomerulosclerosis (30.77%) and Membranous Nephropathy (25.64%). Conclusion: The average patient with glomerulopathy in this service is an adult with nephrotic syndrome. Unlike other reports, secondary glomerulopathies were predominant. These findings may reflect the tertiary characteristic of the assistance at HC-UFPR.

Resumo

Introdução: As glomerulopatias são as doenças renais mais frequentemente diagnosticáveis por biópsia. O levantamento epidemiológico das glomerulopatias permite identificar sua distribuição e principais etiologias e serve de subsídio para definição de estratégias de prevenção e tratamento. Objetivo: O presente estudo pretende identificar a frequência e a correlação clínico-patológica das glomerulopatias diagnosticadas por biópsia no HC-UFPR durante 5 anos. Métodos: Foram realizadas 131 biópsias no período de 1 de janeiro de 2008 a 31 de dezembro de 2012, submetidas à microscopia óptica e de imunofluorescência. Todas as lâminas de microscopia óptica foram revistas por um patologista. Dados clínicos e laboratoriais foram obtidos por revisão dos prontuários. Resultados: 128 de 131 casos foram reanalisados. 46.5% foram obtidos em homens. A idade média de realização da biópsia foi 43 anos para os homens e 38 para as mulheres. Em 99 casos identificou-se a indicação da biópsia; 49.5% apresentavam síndrome nefrótica, 17,17%, insuficiência renal aguda e 15,15%, insuficiência renal crônica; 8,08%, síndrome nefrítica; 6,06%, proteinúria isolada e 4,04%, hematuria isolada. Em 61,21% uma doença subjacente relacionada à glomerulopatia podia ser identificada; 33,62% correspondiam a doença primária e em 5,17% da natureza do glomerulopatia não puderam ser classificados. Entre as glomerulopatias secundárias, a mais frequente foi a nefrite lúpica (49,29%), e, dentre as primárias, glomeruloesclerose segmentar e focal (30,77%) e nefropatia membranosa (25,64%). Conclusão: O paciente com glomerulopatia neste serviço é adulto e portador de síndrome nefrótica. A contrário de outros relatos, observamos predominio das glomerulopatias secundárias, refletindo possivelmente o perfil terciário de atendimento do HC-UFPR.
INTRODUCTION

Glomerulopathies are the kidney diseases most often diagnosed through biopsy.1 Cellular and humoral immune mechanism of primary or secondary nature are very much involved in the pathogenesis of this disease category.2 Its progression often results in the need for renal replacement therapy, procedure which, besides being inconvenient for patients, generates high costs for the healthcare system - in 2012 alone, the Ministry of Health invested R $ 1.8 billion in hemodialysis.3 

Urinalysis, the estimated glomerular filtration rate and individual patient characteristics may indicate the possible diagnosis of a glomerular disease, but only the pathological evaluation of material obtained from renal biopsy enables the physician to establish a precise diagnosis and possible prognostic indicators. The main primary glomerulopathies found in surveys carried out around the world are: focal segmental glomerulosclerosis (FSGS), IgA (IgAN) nephropathy, minimal change disease (MCD), membranous nephropathy (MN), rapidly progressive glomerulonephritis (RPGN), post-infectious glomerulonephritis, and undetermined chronic glomerulonephritis.4 

Secondary glomerulopathies include: diabetic nephropathy (DN), lupus nephritis (LN), renal amyloidosis, and crescent glomerulopathy, related and unrelated to neutrophyl anticytoplasmic antibodies (ANCA).5 The epidemiological survey of glomerulopathies enables the physician to identify its frequency and main causative factors, and it serves as an input for defining their clinical, laboratory and histological features in a particular region, assisting in the development of prevention and treatment strategies.6 This study aims to identify the frequency and clinic-pathological correlation of the glomerular diseases diagnosed by renal biopsy at our institution over a five-year period.

METHODS

This research project was approved by the Research Ethics Committee under number 329.124. 

The survey of patients was carried out by retrospective search in the biopsy records of the Pathology Service (SAP), dated January 1st, 2008 to December 31st of 2012. Using the keywords: kidney, glomerulopathy, nephritis, glomerulonephritis and nephritis, we found 396 cases, broken down into an initial spreadsheet.

After initial screening we consulted the pathology report files. We excluded the cases pertaining to: neoplasia; nephrectomy; pyelonephritis and transplantation. We included only those reports which really dealt with glomerulopathies and renal biopsies with inconclusive diagnoses pending further analysis. This selection resulted in 131 cases, whose medical records were reviewed. We extracted the following information: age; gender; underlying disease and/or comorbidities (diabetes mellitus, hypertension, systemic lupus erythematosus, other); serology (HIV, hepatitis B, hepatitis C and syphilis); clinical presentation (nephrotic syndrome, nephritic syndrome, isolated hematuria, isolated proteinuria, acute renal failure, chronic renal failure); 24-hour proteinuria; partial data from urinalysis (proteinuria, hematuria; cylinders; leukocyturia); serum creatinine; creatinine clearance; serum albumin; lipid profile (total cholesterol and fractions and triglycerides); hemodialysis or peritoneal dialysis at the time of biopsy. The corresponding histological slides, stained by HE, PAS, PAMS and Masson’s were reviewed.

The findings were tabulated in a Microsoft Excel spreadsheet 12.0. Descriptive statistics were performed using the same software, and the statistical parameters were expressed as mean, median and standard deviation. We used the Fisher’s exact test to compare data at the GraphPad InStat application. We considered a $p \leq 0.005$ as significant.

RESULTS

Of the 131 selected patients, 128 had material that could be reevaluated.

The indication of biopsy could be identified in 99 cases, of which 49 (49.5%) had nephrotic syndrome, two of them associated with renal insufficiency; 17 (17.17%) acute renal failure and 15 (15.15%)
chronic renal failure; 8 (8.08%) nephritic syndrome; 6 (6.06%), proteinuria alone and 4 (4.04%), hematuria alone (Table 1).

<table>
<thead>
<tr>
<th>Clinical Presentation</th>
<th>Number of cases</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nephrotic syndrome</td>
<td>49</td>
<td>49.5</td>
</tr>
<tr>
<td>Acute renal failure</td>
<td>17</td>
<td>17.17</td>
</tr>
<tr>
<td>Chronic renal failure</td>
<td>15</td>
<td>15.15</td>
</tr>
<tr>
<td>Nephritic syndrome</td>
<td>8</td>
<td>8.08</td>
</tr>
<tr>
<td>Proteinuria alone</td>
<td>6</td>
<td>6.06</td>
</tr>
<tr>
<td>Hematuria alone</td>
<td>4</td>
<td>4.04</td>
</tr>
</tbody>
</table>

In all the 128 cases, the biopsy technique adopted was the incisional biopsy through a cutting needle. Biopsies obtained an average of 13 glomeruli; one case had no glomeruli. In this and 11 other cases (9.38%) there were no signs of glomerular disease within the methods employed, so that glomerulopathy was confirmed in 116 cases.

Of the 116 confirmed cases of glomerulopathy, 54 (46.5%) were men (p = 0.0739). The mean patient age was 40 years; among men, the mean was 43 years, and for women, 38. Six subjects were aged 70 years or higher. In 72 cases (62.06%), the glomerulopathy was secondary; in 38 cases (32.75%) it was considered primary; and in 6 cases (5.17%) we were unable to classify it as primary or secondary. Among men, there were 30 cases of secondary glomerulopathy (58.82%); and among women, 42 (71.19%). The distribution between men and women can be seen on Figure 1. Among the most frequently identified comorbidities we found: hypertension (70 cases, 60.34%); systemic lupus erythematosus (43 cases, 37.06%); diabetes (20 cases; 17.24%); HIV (11 cases; 9.48%); hepatitis C (11 cases, 9.48%); hepatitis B (4 cases; 3.45%) and hepatitis B associated with hepatitis C (1 case; 0.86%). This data and its distribution between the genders can be found on Table 2.

We found 35 cases of lupus nephritis (30.17%), 23 cases of FSGS (19.83%) - of which 12 (10.34%) were identified as primary FSGS, membranous nephropathy (10 cases, 8.62%) and diabetic nephropathy (9 cases; 7.76%) (Figure 1). The other results can be found on Table 3. Of the 35 cases identified as lupus nephritis, 26 (74.28%) were women (p = 0.0024). Chronic tubulointerstitial changes were present in 50% or more of the kidney cortex from 20 patients. These patients had FSGS, diabetic nephropathy, hypertensive arterial-arteriolar nephrosclerosis and lupus nephritis.

**DISCUSSION**

Due to the high cost and relevant social and economic factors, for which kidney diseases are responsible, countries like Japan and Italy regularly record the nationally diagnosed glomerulopathies. In Brazil, data on the prevalence and epidemiological characteristics of glomerulopathies are scarce. A few Brazilian hospitals have proposed the creation of a local database. The latest national survey on glomerular diseases until this study was published in 2009, and involved 9,617 renal biopsies in the country. To date, there are no studies in the literature on the distribution of glomerulopathies in patients treated at our institution or even in our state, which assigns relevance to this study.

In Brazil, the main indication for biopsy was nephrotic syndrome, which coincides with data from other studies carried out in our country and the world. Indeed, nephrotic syndrome is considered one of the Absolute indications for renal biopsy in individuals older than 6 years of age.

The average obtained from 13 glomeruli per sample meets the bulk of its adequacy regarding the guidelines, which impose the appropriate number of 8-10 glomeruli per sample. This finding, coupled with the presence of only one sample without glomeruli shows that obtaining a histopathological sample has been satisfactory in our institution, which partly ensures diagnostic accuracy. Of the 128 selected patients, 12 showed no signs of glomerular disease through light microscopy or immunofluorescence; that is, during this period, in almost 10% of the suspected glomerulopathies, we were facing other illnesses, normal histopathological kidney or diseases which diagnosis could have been better elucidated through an ultrastructural investigation, not available in our country during the study period. However, it is noteworthy that 6 (50%) of these cases were from patients who were infected with H1N1 during the epidemic of July and August 2009, with samples obtained due to a specific and scientific epidemiological interest, which revealed mainly tubular degenerative changes, as previously published.

It was possible to trace the profile of patients seen because of glomerulopathy in our country. There was
In our study, secondary glomerulopathies were more common than the primary ones. This goes against the overwhelming majority of results presented in the literature, as well as national and regional results (Paulista Registration of Glomerulonephritis, Federal District, Belém and Amazonas), in agreement only with studies carried out in Belgium and Paraguay. Possibly, this is due to the fact that the series came from a tertiary hospital.

The prevalence of secondary glomerulopathies was more significant among females, a reflection of the high prevalence of lupus nephritis, since systemic lupus erythematosus (SLE) is classically predominant in women. In fact, lupus nephritis was more frequent among women than men (26 women and 9 men) and glomerulonephritis was the most frequently diagnosed in our series. This result is in agreement with the Paulista Registration of Glomerulonephritis and most of the series studied, in which it is first among secondary glomerulopathies. Because it is an illness with high morbidity and mortality, especially when associated with common risk factors in developing countries, such as unfavorable

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**Table 2** Distribution of comorbidities per gender

<table>
<thead>
<tr>
<th>Comorbidities</th>
<th>Total # of cases</th>
<th>Males</th>
<th>Females</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arterial hypertension</td>
<td>70</td>
<td>33</td>
<td>37</td>
</tr>
<tr>
<td>Systemic lupus erythematosus</td>
<td>43</td>
<td>12</td>
<td>31</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>20</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>HIV</td>
<td>11</td>
<td>8</td>
<td>3</td>
</tr>
<tr>
<td>Hepatitis C</td>
<td>11</td>
<td>7</td>
<td>4</td>
</tr>
<tr>
<td>Hepatitis B</td>
<td>4</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Hepatitis C + Hepatitis B</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

**Table 3** Distribution of glomerulopathies by order of frequency

<table>
<thead>
<tr>
<th>Histopathology Diagnosis</th>
<th>Number of Cases</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lupus nephritis</td>
<td>35</td>
<td>30.17</td>
</tr>
<tr>
<td>Primary FSGS</td>
<td>12</td>
<td>10.34</td>
</tr>
<tr>
<td>Secondary FSGS</td>
<td>11</td>
<td>9.48</td>
</tr>
<tr>
<td>Membranous nephropathy</td>
<td>10</td>
<td>8.6</td>
</tr>
<tr>
<td>Diabetic nephropathy</td>
<td>9</td>
<td>7.7</td>
</tr>
<tr>
<td>Membranous proliferative glomerulonephritis</td>
<td>6</td>
<td>5.17</td>
</tr>
<tr>
<td>Glomerulopathy secondary to vascular disease</td>
<td>5</td>
<td>4.31</td>
</tr>
<tr>
<td>Minimal lesion disease</td>
<td>4</td>
<td>3.44</td>
</tr>
<tr>
<td>Proliferative and exudative glomerulopathy</td>
<td>4</td>
<td>3.44</td>
</tr>
<tr>
<td>IgA Nephropathy</td>
<td>3</td>
<td>2.58</td>
</tr>
<tr>
<td>Necrotizing glomerulopathy</td>
<td>2</td>
<td>1.72</td>
</tr>
<tr>
<td>Terminal kidney</td>
<td>2</td>
<td>1.72</td>
</tr>
</tbody>
</table>

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**Figure 1.** Frequency of primary and secondary glomerulopathies, by gender

[Graph showing frequency]
socioeconomic conditions, it is important to pay attention to the kidney health of patients with LES.39

The second most found diagnosis on histopathologic analysis was focal segmental glomerulosclerosis, also highly frequent in national results6,14 and in results from Hispanic populations.16 is important to consider that frequent diagnosis of FSGS in our series and in the series above, can be attributed to population characteristics, i.e., selection of adults and nephrotic patients. However, FSGS is a morphological diagnosis, a common consequence of various renal diseases. In our series, we can distinguish the primary or secondary source of FSGS with some reliability, according to the analysis of 24-hour proteinuria, albumin and their own histopathology features.

The cases that progressed to proteinuria in the nephrotic range, hypoalbuminemia, diffuse podocyte disease signals and global sclerosis pattern, predominantly solidified were classified as primary, while those who had subnephrotic proteinuria, normoalbuminemia, focal podocyte disease and global sclerosis pattern predominantly obsolescent, with possible hyalinizing-sclerotic vascular changes, were secondary. As a result, we obtained a very similar proportion of cases of membranous nephropathy and primary FSGS, which differs from the Brazilian results, which put FSGS ahead of membranous nephropathy.6,14 These studies, however, did not make the distinction between primary or secondary FSGS. The significant finding of membranous nephropathy, comparable to the prevalence of primary FSGS in our sample, coincides with findings from Europe,20 a fact that can be attributed to the European heritage in our population.

Chronic tubulointerstitial changes (IFTA%) reflect the duration of kidney disease and the irreversibility of the alterations, and are directly related to serum creatinine levels. Twenty samples were found with IFTA below 50%, so that these cases were very likely diagnosed later. The glomerular diseases found in these cases were in similar proportion to each other - focal segmental glomerulosclerosis, diabetic nephropathy, lupus nephritis and hypertensive arteriolar nephrosclerosis. This information corroborates previous findings2 and highlights the fact that, in our country, such diseases are still diagnosed at a late stage.

Although this is a glomerulonephritis that had the highest increase in prevalence in the past 15 years in Brazil,6 IgA nephropathy was found in only three samples. This contradicts what was expected, because according to the 2009 survey, the South Region had the highest proportion of IgA nephropathy, among all regions of Brazil.6 Taking into account that the number of patients who had hematuria alone was small (4 cases), the low frequency of this disease in our study may be related to non-routine indication of biopsy in patients with this clinical condition,11 to the low identification or the low frequency of hospital admissions for glomerular hematuria in our center.

The limitation of this study was the small number of biopsies carried out in this center and the retrospective review. Still, this was an unprecedented study in our state, which enabled the recognition of the population affected by glomerulopathies and lays the foundation for a future Glomerulopathies Register in the State. It is important to point out that all records and biopsies were extensively revised, so that the conclusions were based on consistent and clinically relevant findings.

CONCLUSION

Through this study, we were able to trace the profile of patients seen because of glomerulonephritis in most tertiary institution of our state. We found equal distribution of men and women, in agreement with national studies6,14 and average age of 40 years. Most cases were secondary glomerulopathies, which was concordant with only two discordant previous studies34,35 and most published national6,10,13,14 and international studies.16,20,30,32,33 The most often diagnosed glomerular disease in our country was lupus nephritis, which was the main cause of secondary kidney disease in several studies.6,14,16,22,24,36-38 It is important to stress that the incidence of IgA nephropathy in our series was low, which may reflect the strategy of glomerular hematuria study in our country.

The profile of glomerulopathies found is a reflection not only of the tertiary profile of our hospital care, but also of our biopsy indications. The small number of samples in our series suggests that we should intensify some of the renal disease search strategies, to better identify them in our population.

REFERENCES


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