Spectrum of glomerular diseases in native kidneys in patients attending Nepal Medical College Teaching Hospital

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ABSTRACT

Background and Aims: Glomerular disease is one of the major cause of end stage kidney disease in Nepal. We have undertaken this study to know the spectrum of glomerular disease in native kidneys in patients attending department of nephrology of Nepal Medical College Teaching Hospital.

Methods: This is a retrospective analysis of patients who have undergone native kidney biopsy from July 2013 to June 2014. Seventy five cases were included. We reviewed the biopsy requisition and where available admission forms for symptoms, indications and complications of the biopsy, urine analysis report and final biopsy report. Kidney biopsy was done under USG guidance or assistance. Automated biopsy gun of 16 or 18G was used. Kidney tissues were sent for light microscopy and immunofluorescence examinations. After 24 hrs of biopsy, USG-KUB and urine RME were repeated for complications

Results: Majority of our study population (62%) was from age group of 13 to 30 years. Sixty five percent of the patients (n= 49) were admitted with a diagnosis of nephrotic syndrome. Most frequent histopathological finding was Minimal Change Disease (26.7%) followed by LN (21.3%), Membranous Nephropathy (18.6%), Focal Segmental Glomerulosclerosis (13.4%) and IgA nephropathy (9.3%). Patients with the histopathological diagnosis of LN and IgAN presented with hematuria in 81.2% and 71.4% respectively. MCD patients presented with proteinuria only. Total no. of glomeruli in the sample was 23.41±11.55.

Conclusion: Nephrotic syndrome was the most common indication for the renal biopsy. MCD was the most common histological finding in our series and Lupus nephritis is not uncommon in our series.

INTRODUCTION

The prevalence of renal diseases and etiologies of Chronic Kidney Disease (CKD) varies greatly with age, sex, race, geographical distribution and indication of kidney biopsy. Glomerular diseases are one of the major cause for end stage kidney disease. To know the distribution of renal disease, we should take help of kidney biopsy which is a gold standard for the diagnosis. Kidney biopsy is a relatively safe and an essential procedure to prognosticate and plan the treatment. The procedure has become safer with life-threatening complications occurring in less than 0.1% of biopsies and the yield of obtaining the specimen has increased greatly with

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the introduction of automated biopsy gun and the procedure
being done under the ultrasound guidance. The prevalence
of the glomerular disease would differ from one place to
another. The prevalence of the kidney disease changes
with the time. We undertook this study to know the spectrum of
the glomerular diseases in native kidneys in patients attending
department of nephrology of Nepal Medical College Teaching
Hospital.

METHODS:
This is a retrospective analysis of our patients who have
undergone native kidney biopsy from July 2013 to June 2014.
There were altogether 82 kidney biopsies done during this
period. Out of which only 75 cases were included for the
study. Seven cases were excluded in the analysis because of
inadequate information. We reviewed the biopsy requisition
and where available admission forms for the symptoms,
indications and complications of the biopsy, urine analysis
report and final biopsy report. For the quantification of
proteinuria either 24hrs urinary protein estimation or spot
urine for protein creatinine (P/C) ratio was taken. Indications of
kidney biopsy were nephrotic syndrome, nephritic syndrome,
unexplained Acute Kidney Injury of more than 2 weeks duration
and asymptomatic urinary abnormality. In patients with clinical
diagnosis of SLE, kidney biopsy was done when urine had
proteinuria of more than 500mg with or without hematuria.
Kidney biopsy was done under local anesthesia and under
aseptic precautions by a nephrologist or a resident posted in
the department. Kidney biopsy was done under USG guidance
or assistance with the help of a radiologist. Automated biopsy
gun of 16 or 18G was used. Two pieces of kidney tissues were
taken and were sent for histopathological examination for Light
microscopy (LM) and Immunofluorescence microscopy (IF).
Sample for LM and IF was sent in formalin and normal saline
respectively. All the samples were sent to India for histological
examination. Patients were admitted for observation and
kept in complete bed rest for atleast 24hrs. After 24 hrs of
biopsy, USG-KUB and urine RME were repeated for perinephric
collection or hematoma and microscopic hematuria. If there
was no complication then the patient was discharged after
24hrs of the procedure.
Statistical Analysis: All the data were compiled in Excel
worksheet and Mean, Standard deviations were calculated.

RESULT:
Out of 82 biopsies, only 75 renal biopsy reports were included
in the study as complete information was not available in
the rest. There were 42 females in our study. Average age of
the study population was 31.32±11.95 years. Majority of our
study population was from age group of 13 to 30 years (Fig 1).
Sixty five percent of the patients (n= 49) were admitted with
a diagnosis of nephrotic syndrome and 18% (n=14) with the
diagnosis of lupus nephritis (LN). All patients had proteinuria
of variable degrees (mean proteinuria of the study sample:
7.23±5.62 g) and 44% (n=33) had microscopic hematuria at
presentation.

Most frequent histopathological finding was Minimal Change
Disease (MCD) (n=20; 26.7%) followed by LN (n=16; 21.3%),
Membranous Nephropathy (MN) (n=14; 18.6%), Focal
Segmental Glomerulosclerosis (FSGS) (n=10; 13.4%) and IgA
nephropathy (IgAN) (n=7; 9.3%). In others category (10.7%),
there were MPGN, C1q Nephropathy, ANCA associated
vasculitis and Chronic Kidney Disease (Fig 2). Out of 16 LN
cases, 14 were of Class IV and other were one each of class III
and combination of classes III and V. There were two samples
that had crescents in the biopsy samples and both of them
were of LN class IV.
Patients with the histopathological diagnosis of LN and IgAN presented with hematuria in 81.2% and 71.4% respectively. MCD patients did not have hematuria, they presented with proteinuria only. MN and FSGS had hematuria in 42.8% and 40% respectively (Table: 1).

Table 1. Distribution of patients according to histopathological diagnosis and pre biopsy microscopic hematuria and proteinuria

<table>
<thead>
<tr>
<th>Microscopic hematuria</th>
<th>% of patient with hematuria</th>
<th>Mean Proteinuria</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCD</td>
<td>0</td>
<td>8.73±6.18</td>
</tr>
<tr>
<td>LN</td>
<td>13</td>
<td>81.2</td>
</tr>
<tr>
<td>FSGS</td>
<td>4</td>
<td>40</td>
</tr>
<tr>
<td>MN</td>
<td>6</td>
<td>42.8</td>
</tr>
<tr>
<td>IgAN</td>
<td>5</td>
<td>71.4</td>
</tr>
<tr>
<td>Others</td>
<td>5</td>
<td>62.5</td>
</tr>
</tbody>
</table>

Total no. of glomeruli in the sample was 23.41±11.55 (15.77±7.66 in LM and 7.96±5.83 in IF). Complications following biopsy were as shown in the table 2.

Table 2. Post renal biopsy complications

<table>
<thead>
<tr>
<th>Complications</th>
<th>No.</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Macroscopic hematuria</td>
<td>6</td>
<td>8</td>
</tr>
<tr>
<td>Microscopic hematuria</td>
<td>8</td>
<td>10.7</td>
</tr>
<tr>
<td>Perinephric hematoma</td>
<td>2</td>
<td>2.67</td>
</tr>
<tr>
<td>Clot in bladder</td>
<td>1</td>
<td>1.34</td>
</tr>
<tr>
<td>Clot colic</td>
<td>1</td>
<td>1.34</td>
</tr>
</tbody>
</table>

We had total of 15 patients who had microscopic hematuria post biopsy, but seven had microscopic hematuria pre renal biopsy so we could not ascertain whether in these patients the appearance of hematuria was biopsy related. There were no cases that needed blood transfusion or had hypotension post renal biopsy. There was no death related with renal biopsy.

DISCUSSION:

Our biopsy series shows that MCD was a common histopathological finding followed by lupus nephritis. As our center is a referral center we may have had 20% of cases of LN. Our study population’s average age was 31 years showing a younger and productive age group of our country is suffering from kidney diseases. This is similar to other studies from Nepal. Average age was 30.6 years for males and 32.9 years for females in the study done by Aryal et al. In the study by Ghimire M et al., the age of the study population was 30yrs for females in the study done by Aryal et al.

In a study done by Khakurel et al., where LN cases were few.

Regarding the pattern of the histological diagnosis, there were quite difference in distribution within and outside the country. In our study, MCD was the most common finding but Sharma A et al and Aryal G study showed MN as the common histological finding in their study. Khakurel et al reported as MCD being most common finding in those patients who had undergone both LM and IF examinations and in Kidney biopsy without IF, mesangial proliferative GN (MesPGN) was commonest. Increased no. of MCD in our series may have been due to absence of Electron microscopy examination of the samples.

IgAN was the most common histological finding in Korea (15%) and China (13%), MCD was second and third common disease respectively. Likewise, in a study done in Romania, MCD was the third commonest disease with 8.5%. In Thailand, MCD comprised of 45.8% of total primary glomerular diseases. IgAN was the most common histological finding in Korea and China. In contrary to the trend of prevalence of IgAN in Asian countries, our study IgAN was present in 9.3% only. The studies from other parts of Nepal also shows similar trend of prevalence of IgAN as ours. This may show that diagnosis of IgAN may be underdiagnosed in Nepal because of possible conservative practice in performing renal biopsy, absence of screening program and widespread unavailability of facilities. FSGS was the second most common disease in the study done by Khakurel S et al and Sharma A et al, while it was fourth common cause of glomerular disease in our study.

In the world, African countries, India and Brazil have more incidences of FSGS.

We had a good no. of glomeruli, it is said that for a good interpretation of renal tissue, a sample should have at least 10 glomeruli. Waldo B et al showed in their study that absence of perinephric bleeding within an hour of post biopsy was predictive of an uncomplicated course while the presence of a perinephric haematoma was not reliably predictive of a clinically significant complication post-renal biopsy.

We routinely did post renal biopsy USG to look for any complications but after 24 hrs of biopsy in our cases.
Complications in our series were few and comparable with other studies. Macroscopic hematuria was seen in 8% and microscopic hematuria was seen in 10.6%. However, microscopic hematuria was seen in 15 patients but as there were seven patients amongst these who had hematuria prior to the renal biopsy so we could not ascertain whether in these patients the appearance of hematuria was biopsy related. Perinephric hematoma was seen in only 2.6%. In the study of Ghimire M et al, the complications like macroscopic hematuria and perinephric hematoma were seen in 6.7% and 5.3% respectively but they did not look at the occurrence of microscopic hematuria. Mendelsohn and Cole found an overall complication rate of 5.3% in their series. Burnstein et al reported complications in 14.3% of 91 patients, out of which 6.6% were minor (microhematuria not requiring transfusion) and rest were major. Study from Romania showed that serious complications were observed most frequently when a large fragment was harvested (>20 glomeruli) and they had gross hematuria in 8.1%, hemorrhage with hypotension in 2.4% and hematoma with hypotension in 0.9%. Though in our study we had total no. of glomeruli of 23, we did not have major complications requiring intervention. The complications may also be related with no. of pricks for harvesting the biopsy tissue and as this was a retrospective study we haven’t documented in our study how many pricks were done on each patient. However, in our institute we do not recommend pricking for more than 3 to 4 times for harvesting the tissue.

This study further emphasizes that the histological pattern of the glomerular diseases depends on age, sex, geographical distribution and indication of kidney biopsy and it changes with the time. Our study has few limitations like it being a retrospective study and the biopsy samples not being examined with electron microscopy. At the time of this study our country did not have immunofluorescence test so all the samples were sent to India for examination in appropriate preservatives and we had to wait for more than 10 days on an average for the report. It is a very high time that we need a renal biopsy registry which is not available in Nepal and need to have more liberal indications for renal biopsy.

In conclusion, Nephrotic syndrome was the most common indication for the renal biopsy. MCD was the most common histological finding in our series and Lupus nephritis is not uncommon in our series. Complication of renal biopsy is uncommon and renal biopsy under USG guidance can be a safer procedure.

REFERENCE: