Value of repeat biopsy in lupus nephritis flares

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ABSTRACT

Objectives: Renal flares are common in lupus nephritis (LN), and class switch is thought to be characteristic. There is no agreement on indications for performing a repeat renal biopsy. Our objective was to retrospectively review patients who had more than one renal biopsy performed on clinical indications, and analyse clinical, pathological and treatment changes after successive biopsies.

Methods: Forty-five patients with LN and one or more repeat renal biopsies were included, with a total of 116 biopsies.

Results: Of the 71 repeat biopsies, pathological transition occurred in 39 (54.9%). When having a previous biopsy with a proliferative lesion, class switch occurred in 55.6%, with 24.4% evolving into non-proliferative classes. When previous biopsy was class V, transition to other classes occurred in 58.3% and changes were all into proliferative classes. Conversion from one pure proliferative form to another (class III to class IV or vice versa) happened in 11.3% of the rebiopsies, with 62 rebiopsies (87.3%) leading to a change in the treatment regimen.

Conclusions: Histological transformations were common, and they occurred when the previous biopsy had non-proliferative lesions as well as when lesions were proliferative. Treatments were modified after repeat renal biopsy in the majority of patients. In this experience, kidney repeat biopsies were useful in guiding treatment of LN flares.

INTRODUCTION

Renal biopsy has a paramount role in the diagnosis of lupus nephritis (LN). Histologic findings provide the basis for treatment recommendations. Flares represent a significant problem because of the potential for cumulative damage that may lead to deterioration of renal function even after a successful treatment. The pathological class of LN may change to a different class during a disease flare.

The clinical relevance of the repeat renal biopsy remains in debate. Some authors suggest performing it after induction or maintenance treatment in order to assess treatment efficacy. Others propose a repeat biopsy only on clinical indications (worsening of renal function, persistent proteinuria or haematuria, suspicion of renal flare or class change, etc.). Even in this setting, the importance of a second biopsy may not be uniform, since some authors have proposed that patients with proliferative lesions on their original biopsy rarely switch to a pure non-proliferative nephritis during a flare, and in these cases, appropriate induction treatment can be initiated without additional biopsies.

Some features in the second renal biopsy have been associated with bad renal prognosis: persisting inflammatory lesions at 6 months of treatment, the presence of subendothelial deposits after 2 years of treatment and higher chronicity indexes. These findings suggest that repeat biopsies could have a prognostic value besides the immediate clinical relevance in taking therapeutic decisions.

Our objective was to retrospectively review patients who had more than one renal biopsy performed on clinical indications and analyse clinical, pathological and treatment changes after successive biopsies.
**METHODS**

**Patients**

Patients were recruited at the Hospital Italiano de Buenos Aires, Argentina. Systemic lupus erythematosus patients (fulfilling American College of Rheumatology (ACR)\(^{10}\) or The Systemic Lupus International Collaborating Clinics (SLICC)\(^{11}\) criteria) who had a diagnosis of LN and two or more renal biopsies after year 2001 were included. Electronic medical records were reviewed and clinical, laboratory and treatment data were obtained from each patient.

Laboratory values, such as serum creatinine, albumin, urea, proteinuria, complement levels (C3 and C4), antidouble-stranded DNA antibody (anti-dsDNA) titre, were selected during the month before and the month after the renal biopsy was performed. Antibodies were detected by the hospital laboratory using standard methods: indirect immunofluorescence for antinuclear antibodies on HEP 2 cells, immunofluorescence on criithidia luciliae for anti-dsDNA antibodies, ELISA for Sm antibodies, anticardiolipin antibodies, antibeta2glicoprotein I antibodies and lupus anticoagulant test according to the International Society of Thrombosis and Haemostasis (ISTH) guidelines.\(^{12}\) Patients were considered to have antiphospholipid syndrome if fulfilling modified Sapporo criteria (2006).\(^{13}\)

Follow-up time was calculated since first renal biopsy until last hospital visit.

**Renal biopsy**

Renal biopsy was performed on clinical indications: improvement of renal disease but persistence of non-nephrotic proteinuria and/or haematuria, persistence of or relapsing nephrotic syndrome or worsening of renal function. No patients were scheduled to receive the renal biopsy to assess the efficacy of treatment.

All biopsies were assessed by experienced pathologists by light microscopy and immunofluorescence.

Renal biopsy was evaluated according to the WHO classification of LN\(^ {14}\) when the biopsy was performed before the year 2003, and according to the international society of nephology/Renal Pathology Society (ISN/RPS) classification of LN\(^ {15}\) after that date. If biopsy specimens were classified according to WHO classification (class I, normal, class II, mesangial proliferation, class III, focal and segmental proliferative glomerulonephritis, class IV, diffuse proliferative glomerulonephritis, class V, membranous glomerulonephritis and class VI, sclerosing glomerulonephritis), they were reassessed according to the ISN/RPS classifications, and the new classification was compared between successive biopsies.

Activity and chronicity indices were scored according to the 1983 proposal by Austin et al.\(^ {16}\) Class III, Class IV and combinations between III/IV plus V were considered proliferative classes. All the rest were considered non-proliferative.

**Treatment**

The choices of treatment regimens were up to the individual nephrologist and rheumatologist. We considered a treatment change when the immunosuppressive treatment was modified after the renal biopsy (drug change, drug addition, drug suspension).

**Statistical analysis**

Data were analysed using SPSS V.20.0 version for Windows. The continuous variables were presented as the mean±SD, or the median and IQRs where appropriate. Categorical variables are presented as percentages. For the univariate analysis we compared two groups using Student t test when normally distributed, and Mann–Whitney test when not. Two-sided p values <0.05 were considered statistically significant.

Cox regression analysis was performed using poor outcome as dependent variable and adjusting by age, activity and chronicity indexes, and creatinine levels.

Receiver operating characteristics (ROC) curves were used to determine the best cut-off points to predict adverse outcomes.

**Ethical approval**

We received confirmation from our ethical committee that ethics approval was not required for this retrospective study.

**RESULTS**

We identified 45 lupus patients (40 women) with at least two renal biopsies. These patients had a total of 116 biopsies. Clinical characteristics of the patients are provided in table 1.

Regarding histopathological analysis, proliferative classes were the most frequent (56.9%), followed by class V (16.4%) (table 2).

Clinical indications for biopsies were persistent non-nephrotic proteinuria and/or haematuria (n=59, 50.9%), persistence of or relapsing nephrotic syndrome (n=30, 25.9%), and worsening of renal function (n=21, 18.1%). We do not have data on the cause that motivated rebiopsy in the remaining six cases.

Of the 71 repeat biopsies, pathological transition occurred related to the previous biopsy in 39 (54.9%) and did not occur in 26 (36.6%) (table 5). We could not assess change in six of the rebiopsies because of unknown previous biopsies.

When the previous biopsies were classified as proliferative (class III, IV or combinations between III/IV+V) (n=45), histological change occurred in 25 (55.6%), and 11 of them changed to non-proliferative classes (24.4%). When previous biopsy was class V (n=12), transition to other classes occurred in seven (58.3%), and all changed to proliferative classes. Conversion from one pure proliferative form to another (class III to class IV or vice versa) occurred in eight of the repeat biopsies (11.3%).
Sixty-two repeat biopsies (87.3%) led to a change in treatment. The choices of treatment regimens were up to the individual nephrologist and rheumatologist. Treatments more frequently used for induction were cyclophosphamide (≥ 6 pulses in 34.3% and less than 6 pulses in 13.3%), mycophenolate (25.7%) and rituximab (5.7%). For maintenance, the more frequently used drugs were mycophenolate (58.7%), azathioprine (20.7%) and cyclophosphamide (9.8%).

At the end of the follow-up, seven patients were in dialysis and other four patients had duplicated their serum creatinine. This subgroup with adverse renal outcome represented 24.4% of this cohort. Characteristics of second renal biopsy in this subgroup are shown in Table 4. Median time to enter dialysis was 4.4 years (IQR 5.9) since first renal biopsy. Three patients died during follow-up, causes of deaths were ischaemic colitis, acute abdominal pain and infectious endocarditis. Five of the patients that entered dialysis had a class VI biopsy before and two had a class IV biopsy as the last one.

Median creatinine values at time of second biopsy were significantly higher in patients who had an adverse renal outcome (2.56 mg/dL (IQR: 1.7–3.64) vs 0.8 mg/dL (IQR: 0.7–1.03); p<0.0001). When performing a ROC curve using creatinine at the second biopsy to predict adverse renal outcome, the area under ROC curve was 0.986 (95% CI 0.957 to 1, p=0.015). A cut-off value of creatinine greater than or equal to 1.3 mg/dL had 100% sensitivity and 93.7% specificity for adverse renal outcome. In a similar way, mean chronicity index at the second biopsy (data missing from 12 patients) was significantly higher in patients with poor renal outcome (6.6±1.6 vs 2.9±1.7; p<0.0001). Area under ROC curve for adverse renal outcome was 0.937 (95% CI 0.817 to 1, p=0.06) and a chronicity index greater than or equal to 6.5 had 87.5% sensibility and 100% specificity for that outcome.

Table 1 Characteristics of lupus patients with more than one kidney biopsy during follow-up

| Lupus patients with at least 2 renal biopsies (n=45) | Female, n (%) | Age at first renal biopsy, median (IQR) years | Fulfilment of ACR 1997 lupus criteria, n (%) | Fulfilment of SLICC 2012 lupus criteria, n (%) | ANA positive, n (%) | Sm antibody positive, n (%) | Anti-dsDNA antibodies positive, n (%) | Antiphospholipid antibodies positive, n (%) | Time between first and second renal biopsy, median (IQR) years | Follow-up, median (IQR) years |
|---|---|---|---|---|---|---|---|---|---|---|---|
| n (%) | 40 (88.9) | 29.6 (17.1) | 37 (82.2) | 45 (100) | 42 (93.3) | 12 (26.7) | 29 (64.4) | 15 (33.3) | 3.4 (4.4) | 8.7 (7.5) |

*and **: insufficient data from 4 and 3 patients, respectively.
ACR, American College of Rheumatology; ANA, antinuclear antibodies; anti-dsDNA antibodies, antidouble-stranded DNA antibody; SLICC, The Systemic Lupus International Collaborating Clinics; Sm, anti-Smith antibodies.

Table 2 Classification of first and successive renal biopsies according to the International Society of Nephrology/Renal Pathology Society (ISN/RPS) 2003 classification of lupus nephritis

| Lupus nephritis classification | 1st biopsy (n=45) | 2nd biopsy (n=45) | 3rd biopsy (n=18) | 4th biopsy (n=7) | 5th biopsy (n=1) |
|---|---|---|---|---|---|
| Class II, n | 5 | 3 | 0 | 0 | 0 |
| Class III, n | 4 | 2 | 2 | 0 | 1 |
| Class IV, n | 22 | 25 | 6 | 4 | 0 |
| Class V, n | 7 | 8 | 3 | 1 | 0 |
| Class VI, n | 0 | 2 | 2 | 1 | 0 |
| Combination between III/IV + V, n | 1 | 3 | 5 | 1 | 0 |
| Vasculopathy, n | 1 | 0 | 0 | 0 | 0 |
| Unknown, n | 5 | 2 | 0 | 0 | 0 |

Table 3 Class changes in rebiopsies

| Initial biopsy | Rebiopsies with class change (n=39) |
|---|---|
| Class II | Class III 1 |
| Class III | Class IV 2 |
| Class IV | Class V 2 |
| Class V | Class III 5 |
| Vasculopathy (APS) | Class IV 1 |
| Unknown Class | Class VI 1 |

APS, antiphospholipid syndrome.
In multivariable Cox analysis, after adjusting by age at time of second biopsy and activity index, creatinine levels and chronicity index were still associated with poor renal outcome (HR: 1.14 (95% CI 1.01 to 1.29), and 1.58 (95% CI 1.1 to 2.3), respectively).

**DISCUSSION**

The objective of this retrospective study was to analyse the value of repeat renal biopsy in patients with LN flares. Flares represent a significant problem because of the potential for cumulative damage that may lead to deterioration of renal function as well as toxicity due to additional immunosuppression. There is enough evidence showing that relapsing nephritis has a worse renal prognosis. Clinical expression of renal flares differs. Some authors have postulated that nephritic flares (mainly when having an active urine sediment, proteinuria greater than 2 g/day and worsening of renal function) are more important than nephrotic flares (characterised only by proteinuria greater than 2 g/day). However, no clinical or biological feature uniformly predicts renal morphology, particularly the extent of renal lesions. Moreover, the histological picture in a patient with impaired renal function is unpredictable.

Relapse in LN has been reported in different studies from 27% to 66%, depending on patients’ characteristics, treatments and definitions of flare used in each cohort. Histological transformation from one class to another is very well recognised in LN, and it can be observed in 49–75% of repeat biopsies. Nevertheless, value of a second renal biopsy has been disputed.

There are two possible scenarios for repeat renal biopsy: (1) as a strategy for assessing response after induction or maintenance treatment. In these cases, the objective is to evaluate histological markers that may indicate efficacy of the treatment received and/or may predict patient’s renal course. (2) on the other hand, some authors have suggested repeat biopsies on clinical indications only, mainly when suspecting a relapse.

Some guidelines suggest a repeat renal biopsy when clinical response to treatment is not as good as expected, in case of flare or if suspecting a histological transformation, and some others only when anticipating a therapeutic change after the repeat biopsy.

In the last few years, prognosis and treatment of histological class III and IV has been homologated, leading to guidelines that do not differentiate the treatment of class III and IV LN. Considering this, some authors have suggested that transitions between proliferative classes have no additive value on treatment decisions and they suggest that when a patient with proliferative lesions on a previous biopsy presents with a renal flare, appropriate induction treatment can be initiated without additional biopsies, since a repeat biopsy will show similar lesions in most cases.

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**Table 4** Characteristics of the second renal biopsy in patients with duplication of creatinine and/or end stage renal disease at the end of follow-up

| Patient | Class in 1st biopsy | Age at 1st biopsy | Class in 2nd biopsy | Activity index (0–24) | Chronicity index (0–12) | Creatinine at the time of 2nd biopsy (mg/dL) | Renal outcome at the end of follow-up |
|---------|---------------------|------------------|---------------------|----------------------|------------------------|------------------------------------------|-------------------------------------|
| Patient 1 | IV                  | 64               | VI                  | 2                    | 8                      | 1.6                                      | Duplication of creatinine             |
| Patient 2 | V                  | 50               | VI                  | NR                   | NR                     | 3.6                                      | Dialysis                             |
| Patient 3 | V                  | 41               | V                   | 0                    | 3                      | 2.2                                      | Dialysis                             |
| Patient 4 | IV                  | 31               | IV                  | 5                    | 7                      | 1.7                                      | Duplication of creatinine             |
| Patient 5 | IV                  | 44               | IV                  | 2                    | 7                      | 1.4                                      | Dialysis                             |
| Patient 6 | IV                  | 48               | IV                  | 7                    | 7                      | 4.6                                      | Dialysis                             |
| Patient 7 | unknown             | 34               | III                 | 5                    | 7                      | 1.3                                      | Dialysis                             |
| Patient 8 | V                  | 41               | V                   | NR                   | NR                     | 2.5                                      | Dialysis                             |
| Patient 9 | V                  | 45               | V                   | NR                   | NR                     | 2.1                                      | Dialysis                             |

APL, antiphospholipid syndrome; NR, not reported.
In our cohort, histological transformations were common (54.9%) in agreement with previous reports, and frequencies of changes were similar between those with proliferative and non-proliferative lesions on the previous biopsy (55.6% and 58.3%, respectively, p=1). However, when having a previous biopsy with a proliferative lesion, class switch happened in 55.6%, with 24.4% transforming into non-proliferative classes. This finding differs from the study by Daleboudt et al., mentioned above.

When previous biopsy was class V, transition to other classes occurred in 58.3% and changes were all into proliferative classes. Only 11.3% of repeat biopsies showed conversion from one pure proliferative form to another (class III to class IV or vice versa). This is lower than in other cohorts.

In conclusion, and differing slightly from other previous reports, histological switch happened in non-proliferative classes as well as in proliferative ones, with a higher proportion of changes from non-proliferative to proliferative (58.3% vs 24.4%, p=0.04).

Regarding our patients’ renal outcomes, they do not differ from previous reports in LN repeat biopsy studies. A quarter of the patients had a poor renal outcome, seven patients with end-stage renal disease and other four with serum creatinine duplication; three patients died. When predictors were analysed, having a higher serum creatinine and a higher chronicity index at the second biopsy was associated with poor renal prognosis. These results are in conformity with what has been previously found by authors who have analysed clinical parameters and repeat biopsy parameters to predict renal prognosis.

Several limitations of the present study need to be addressed. Our study had a small sample size and we have missing data. Twelve of the biopsies were performed at other hospitals. We do not have the results of some, and in others we only know the class but we could not have access to the histopathological material. Of the biopsies (104) performed at our hospital, not all of them were evaluated with activity and chronicity indices, and they were analysed by different pathologists with different classifications according to the year when they were performed, as mentioned above. We were not able to compare activity and chronicity indices in successive biopsies, analysis that would have been interesting in order to assess efficacy of treatments and prognosis.

There is a reluctance to repeat an invasive procedure with potential complications, but taking into account immunosuppressive treatments’ side effects, when a renal flare occurs, it seems reasonable to try to assess kidney impact with accuracy in order to adjust medications to each individual patient. Facing the impossibility of predicting histological changes based on clinical data, performing a repeat renal biopsy appears as a reasonable option. In our patients, rebiopsies were an important guide for treating physicians, since 87.3% conducted to a treatment suppressive treatments when biopsies showed a class VI LN.

CONCLUSIONS

In our cohort, histological transformations were common in successive biopsies (54.9%). Coming from a previous biopsy with non-proliferative lesions, transformations into proliferative occurred in 58.3%. On the other hand, 24.4% of the initial proliferative biopsies changed into non-proliferative in the repeat biopsy. We believe, that in renal flares, our experience confirms that repeat renal biopsy allows for more precise therapeutic decisions and better establishing of long-term prognosis in the individual patient.

Contributors All authors made substantial contribution to the following: conception and design, acquisition of data or analysis and interpretation of data; drafting the article or revising it critically for important intellectual content; final approval of the version published.

Competing interests None.

Ethics approval CEPI.

Data sharing statement No additional data are available.

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REFERENCES

1. Esdaile JM, Joseph L, MacKenzie T, et al. The pathogenesis and prognosis of lupus nephritis: information from repeat renal biopsy. Semin Arthritis Rheum 1999;29:135–48.
2. Moroni G, Pasqualli S, Quaglini S, et al. Clinical and prognostic value of serial renal biopsies in lupus nephritis. Am J Kidney Dis 1999;34:530–9.
3. Alsuwaida A, Husain S, Alghonaim M, et al. Strategy for second kidney biopsy in patients with lupus nephritis. Nephrol Dial Transplant 2012;27:1472–8.
4. Hill GS, Delahousse M, Nochy D, et al. Predictive power of the second renal biopsy in lupus nephritis: significance of macrophages. Kidney Int 2001;59:306.
5. Bihl GR, Petri M, Fine DM. Kidney biopsy in lupus nephritis: look before you leap. Nephrol Dial Transplant 2006;21:1749–52.
6. Ponticelli C, Moroni G. Renal biopsy in lupus nephritis—what for, when and how often? Nephrol Dial Transplant 1998;13:2452–4.
7. Daleboudt GMN, Bajema IM, Goemaere NNT, et al. The clinical relevance of a repeat biopsy in lupus nephritis flares. Nephrol Dial Transplant 2009;24:3712–17.
8. Hill GS, Delahousse M, Nochy D, et al. Outcome of relapse in lupus nephritis: roles of reversal of renal fibrosis and response of inflammation to therapy. Kidney Int 2002;61:2176–86.
9. Mosca M, Pasquarriello A, Tavoni A, et al. Predictors of renal outcome in diffuse proliferative glomerulonephritis in systemic lupus erythematosus. Lupus 1997;6:371–8.
10. Hochberg M. Updating the American College of Rheumatology revised criteria for the classification of systemic lupus erythematosus. Arthritis Rheum 1997;40:1725.
11. Petri M, Orbai A-M, Alarcón GS, et al. Derivation and validation of the Systemic Lupus International Collaborating Clinics classification criteria for systemic lupus erythematosus. Arthritis Rheum 2012;64:2877–86.
12. Pengo V, Tripodi A, Reber G, et al. Update of the guidelines for lupus anticoagulant detection. Subcommittee on Lupus Anticoagulant/Antiphospholipid Antibody of the Scientific and Standardisation Committee of the International Society on Thrombosis and Haemostasis. J Thromb Haemost 2009;7:1737–40.
13. Miyakis S, Lockshin MD, Atsumi T, et al. International consensus statement on an update of the classification criteria for definite antiphospholipid syndrome (APS). J Thromb Haemost 2006;4:295–306.
14. Gollbus J, McCune WJ. Lupus nephritis. Classification, prognosis, immunopathogenesis, and treatment. Rheum Dis Clin North Am 1994;20:213–42.
15. Weening JJ, D’Agati VD, Schwartz MM, et al. The classification of glomerulonephritis in systemic lupus erythematosus revisited. *Kidney Int* 2004;65:521–30.

16. Austin HA, Muenz LR, Joyce KM, et al. Prognostic factors in lupus nephritis. Contribution of renal histologic data. *Am J Med* 1983;75:382–91.

17. Sidiropoulos PI, Kritikos HD, Boumpas DT. Lupus nephritis flares. *Lupus* 2005;14:49–52.

18. Moroni G, Guagliati S, Maccario M, et al. “Nephritic flares” are predictors of bad long-term renal outcome in lupus nephritis. *Kidney Int* 1996;50:2047–53.

19. Wang G-B, Xu Z-J, Liu H-F, et al. Changes in pathological pattern and treatment regimens based on repeat renal biopsy in lupus nephritis. *Chin Med J (Engl)* 2012;125:2890–4.

20. Lu J, Tam L-S, Lai FM-M, et al. Repeat renal biopsy in lupus nephritis: a change in histological pattern is common. *Am J Nephrol* 2011;34:220–5.

21. Kdigo G. KDIGO clinical practice guideline for glomerulonephritis. *Kidney Int* 2012;2:139–274.

22. Ruiz Irastorza G, Espinosa G, Frutos MA, et al. Diagnosis and treatment of lupus nephritis. Consensus document from the systemic autoimmune disease group (GEAS) of the Spanish Society of Internal Medicine (SEMI) and Spanish Society of Nephrology (S.E.N.). *Nefrologia* 2012;32(Suppl 1):1–35.

23. Van Tellingen A, Voskuyl AE, Vervloet MG, et al. Dutch guidelines for diagnosis and therapy of proliferative lupus nephritis. *Neth J Med* 2012;70:199–207.

24. Radhakrishnan J, Catran DC. The KDIGO practice guideline on glomerulonephritis: reading between the (guide)lines—application to the individual patient. *Kidney Int* 2012;82:840–56.

25. Isenberg D, Appel GB, Contreras G, et al. Influence of race/ethnicity on response to lupus nephritis treatment: the ALMS study. *Rheumatology (Oxford)* 2010;49:129–40.

26. Rovin BH, Furie R, Latini K, et al. Efficacy and safety of rituximab in patients with active proliferative lupus nephritis: the Lupus Nephritis Assessment with Rituximab study. *Arthritis Rheum* 2012;64:1215–26.

27. Ginzler EM, Dooley MA, Aranow C, et al. Mycophenolate mofetil or intravenous cyclophosphamide for lupus nephritis. *N Engl J Med* 2005;353:2219–28.

28. Contreras G, Pardo V, Leclercq B, et al. Sequential therapies for proliferative lupus nephritis. *N Engl J Med* 2004;350:971–80.

29. Gordon C, Jayne D, Pusey C, et al. European consensus statement on the terminology used in the management of lupus glomerulonephritis. *Lupus* 2009;18:257–63.