Relationship between Fractional Excretion of Magnesium and Renal Histology in Glomerulonephritis

Abstract
Progressive pediatric glomerulonephrases can lead to tubulointerstitial fibrosis (TIF) and chronic kidney disease (CKD). Fractional excretion of magnesium (FEMg) is a sensitive tubular function test (Ref). FEMg correlates positively with TIF (Ref). The purpose of the study was to correlate FEMg with renal histopathology in pediatric glomerular nephrosis (GN). Sixty three cases, between 1-12 years, of biopsy diagnosed GN were included purposively in this cross sectional study from November 2008 to April 2010. Cases were taken from Department of Paediatric Nephrology, Dhaka Shishu (Children) Hospital and Department of Paediatric Nephropathy, National Institute of Kidney Diseases & Urology, Sher-e-Bangla Nagar, Dhaka, Bangladesh. Minimal change disease (MCD) was most common at 27% and remaining 73% were other than MCD. The mean FEMg in MCD was normal (1.81 ± 0.34%) but in other than MCD as a group it was significantly elevated (7.60 ± 19.23%; p<0.001), and significantly elevated separately in diffuse mesangial proliferative glomerulonephritis (3.96 ± 1.06%; p<0.001), focal segmental glomerulosclerosis (9.99 ± 4.45%; p<0.01), focal segmental proliferative glomerulonephritis (4.63 ± 2.29%; p<0.05) and crescentic glomerulonephritis (17.74 ± 4.93%; p<0.01).

Keywords: Fractional excretion of magnesium; Glomerulonephritis; Tubular interstitial disease

Introduction
Numerous inflammatory and non inflammatory diseases affect the glomerulus and lead to alteration in glomerular permeability, structure & function. Glomerulonephritis (GN) is a generic term for several diseases and a histopathologic term signifying inflammation & proliferation of cells within the glomerulus [1]. It may be primary, restricted to kidney or may be part of a multisystem diseases. Clinical evaluation includes assessment of proteinuria, hematuria, presence or absence of renal insufficiency and presence or absence of hypertension and ultimately diagnosed with renal histology [2,3]. Glomerular diseases may have an indolent course or begin abruptly leading to acute or rapidly progressive glomerulonephritis. Chronic glomerulonephritis implies continuing glomerular injury that frequently leads to glomerular destruction and end stage renal failure. Glomeruli may be injured by several mechanism in different diseases but produce limited number even same histopathologic changes. The full assessment of a renal histology requires light microscopy, electron microscopy and immunofluorescence [4]. Proliferation of glomerular cells occur in most form of glomerulonephritis, commonly involves endothelial and mesangial cells and sometimes there may have crescent and sclerosis [5]. Abnormalities are not confined only to glomerulus, tubulointerstitial changes attend almost all forms of progressive glomerular and vascular injury. In most cases of glomerulonephritis long term course lead to chronic renal failure and characterized by tubular atrophy, interstitial fibrosis [6].

In children the most common presentation of glomerulonephritis is nephrotic syndrome [7]. Histologically minimal change disease (MCD) is the commonest 76.4%, other histological pattern are mesangial proliferative glomerulonephritis (MesPGN) 2.3%, membranoproliferative glomerulo nephritis (MPGN)7.5%, focal segmental glomerulosclerosis (FSGS) 6.9%, membranous nephropathy(MN) 1.5%. Other pattern of glomerulonephritis in pediatric patients are lupus nephritis, IgA nephropathy, Henoch - Schonlein purpura nephritis or isolated hematuria and or proteinuria. In NS, 30-40% steroid resistant MCD develops end stage renal disease by 5 years. Similarly MesPGNsz (50%), MPGN (20-30%), FSGS (21%) and MN (7.3%) progress to end stage renal disease within 5 years of age [9-13].

Two patterns of clinicopathologic subsets have been generally delineated in nephrosis namely 1) a benign clinicopathologic entity associated with no tubulointerstitial disease such as minimal change lesion or mild mesangial proliferative glomerulonephritis 2) A progressive clinicopathologic disease associated with tubulointerstitial lesion such as that observed in focal segmental glomerulosclerosis, a moderate to diffuse mesangial proliferative glomerulonephritis and membranoproliferative glomerulonephritis. The clinical course in the former is usually associated with a self limited disease without renal diseases...
progression where as in the latter progressive renal disease is destined for chronic renal insufficiency and end stage renal disease. Tubular function testing i.e. may predict end stage renal disease. FEMg can reflect tubular function FEMg < 2.2% indicates intact tubular function whereas elevated FEMg indicates tubular damage. The greater the FEMg the greater the magnitude of TIF.

Materials and Methods

In this cross sectional study 63 patients, between 1-12 years of age, of histologically diagnosed glomerulonephritis were included. The study was performed in Department of Pediatric Nephrology, Dhaka Shishu Hospital & National Institute of Kidney Diseases & Urology, Sher-e-Bangla Nagar, Dhaka in the period of November 2008 to April 2010. Informed written consent was obtained from the legal guardian. Reassurance was given to the guardian regarding investigation. Clearance has been taken from the ethical committee. No study was cost was paid by study subjects. Biochemical indices and renal biopsy report were noted. Diuretics and aminoglycosides, they were stopped 24 hours and 48 hours prior to collection of serum and urine [Q: is this long enough?]. 10 ml urine samples were centrifuged and stored in refrigerator. Serum creatinine and magnesium, spot urinary creatinine and magnesium were measured. The MG method was used on the Dimension ® (Siemens, address) Clinical Chemistry System. FEMg was then calculated using the following formula:

\[ FEMg = \frac{\text{Urine magnesium} \div \text{plasma magnesium}}{\text{Urine creatinine} \div \text{plasma creatinine}} \times 100 \]

(Normal value of FEMg: 2.2% or less)

Light microscopy and direct immunofluorescent examinations were reviewed by a histopathologist blinded to FEMg (Table 1). Data were collected, compiled and analyzed by using SPSS version 12. Independent -t test was used as the test for significance. P value of <0.05 was considered statistically significant.

Results

FEMg for each diagnosis is given in Table 1. FEMg in MCD was normal and taken as standard. FEMg in MesPGN and MPGN was found increased but not significant (Figure 1, Table 2&3). In case of D-MesPGN, FSGN, FSGS and crescentic glomerulonephritis FEMg was increased and statistically significant. Chronic sclerosing GN and IgA Nephropathy were one each, so these are not mentioned in the table.

Table 1: Distribution of patients with glomerulonephritis by clinical diagnosis.

| Clinical diagnosis of the patient                           | Frequency | Percentage (%) |
|------------------------------------------------------------|-----------|----------------|
| Nephrotic syndrome (NS) 1st attack                          | 22        | 34.92          |
| Frequent relapse nephrotic syndrome (FRNS)                  | 17        | 26.98          |
| Steroid dependent nephrotic syndrome (SDNS)                 | 8         | 12.70          |
| Steroid resistant nephrotic syndrome (SRNS)                 | 1         | 1.59           |
| Nephrotic syndrome with renal failure                       | 2         | 3.17           |
| HSP with nephritic-nephrotic syndrome                       | 1         | 1.59           |
| SLE with Nephritis                                          | 4         | 6.35           |
| AGN with ARF                                                | 6         | 9.52           |
| Acute renal failure                                         | 1         | 1.59           |
| Chronic kidney disease                                      | 1         | 1.59           |
| Total                                                       | 63        | 100.00         |

Table 2: Relation of FEMg with renal histopathology of the patient.

| Renal histopathology of the patient                         | No.   | Mean FEMg(%) | ±SD FEMg | t-value | P       |
|------------------------------------------------------------|-------|--------------|----------|---------|---------|
| Minimal change disease (MCD)                               | 17    | 1.81         | 0.34     | -       | -       |
| Mesangial proliferative glomerulonephritis (MesPGN)        | 4     | 3.50         | 1.08     | 3.07    | >0.05, ns|
| Diffuse mesangial proliferative glomerulonephritis (D-MesPGN) | 13    | 3.96         | 1.06     | 3.96    | <0.001, S|
| Focal segmental proliferative glomerulonephritis (FSGN)   | 10    | 4.63         | 2.29     | 3.86    | <0.05, s|
| Focal segmental glomerulosclerosis (FSGS)                  | 9     | 9.99         | 4.45     | 5.49    | <0.01, S|
| Crescent glomerulosclerosis (CSG)                          | 5     | 17.74        | 4.93     | 7.21    | <0.01, S|
| Membranoproliferative glomerulonephritis (MPGN)            | 3     | 13.97        | 4.28     | 4.91    | ns      |

SD: Standard Deviation; S: Significant; NS: Not Significant.
**Discussion**

Disease incidence was similar to the study by Madani A et al. [7] (66.26%) and Al-Rasheed et al. [14] (77%). In this study FEMg was normal in MCD (1.81 ±0.34%). In D-MesPGN (3.96±1.06), FSGS (9.99±4.45), crescentic GN (17.74±4.93), FEMg found abnormally elevated and statistically highly significant (p<0.01). In FSGN (4.63±2.29) FEMg also found elevated and significant (p<0.05). FSGS showed abnormally elevated FEMg in studies by Futrakul P; Deekajorndech T similar to this study. FSGN, crescentic GN and MPGN were not included in other studies; which are important cause of CKD [11-18]. (Please discuss other studies which reference FEMg- the purpose of the study was not so much important cause of CKD [11-18]. Tubulointerstitial injury is an invariant finding in the chronically diseased kidney, irrespective of the type of disease [19]. FEMg appeared as a sensitive indicator of severity of renal insufficiency as showed in chronic renal failure patients of Nigeria [20-22].

( I don't think the purpose of the study was to compare markers) Glomerular filtration rate, peritubular capillary blood flow, afferent arteriolar resistance were not assessed in this study which are important factor for tubular dysfunction or TIF due to limited resources but assessed in different study [23]. FEMg is normal in minimal change disease and abnormally elevated in other than minimal change disease which is statistically significant.

**Conclusion**

Fractional excretion of magnesium increases in other than minimal change disease and may be of value as a simple investigation in glomerulonephritis to assess disease severity.

**Table 3:** Comparison of FEMg between minimal change disease (MCD) and other than minimal change disease.

| Variable                        | Subject | Number | Mean   | ±SD     | t-value | P      |
|---------------------------------|---------|--------|--------|---------|---------|--------|
| Fractional excretion of magnesium | MCD     | 17     | 1.81   | 0.34793 | 6.95    | <0.001 |
|                                 | Other than MCD     | 46     | 7.6    | 19.234  |         | S      |

**References**

1. Watson R, Taylor MC, McGraw M (2003) Disorders of the urinary system. In: Mclntosh N & Smyth LR (Eds.), Forfar & Arneil’s *Textbook of Pediatrics*: (6th edn), Churchill Livingstone, Spain, pp. 613-620.

2. Falk JR, Chaules J, Nachman HP (2004) Primary Glomerular Disease. In: Brenner MB (Ed.), *Brenner and Rector’s The Kidney*: (7th edn), Elsevier, Philadelphia, USA, pp. 1293.

3. Johnson JR (2007) Introduction to Glomerular disease. Pathogenesis and classification. In: Feethalily J & Richard JJ (Eds.), Comprehensive clinical Nephrology. (3rd edn), Elsevier, Philadelphia, USA pp. 181-191.

4. Charles E, Alpers (2005) The Kidney. In: Kumar, Abbas & Fausto (Eds.), *Robbins & Cortan Pathologic basis of disease*: (6th edn), Elsevier Saunders, Philadelphia, USA, pp. 955-959.

5. Davis DL, Avner DL (2007) Glomerular Disease. In: *Kligman, Behrman & Stanton (Eds.), Nelson Text book of pediatrics*: (7th edn), Elsevier, Philadelphia, USA, pp. 2163-2167.

6. Isiakul-Piechocka, Krzywański M (1996) The role of tubulo interstitial changes in progression of kidney failure in patient with chronic glomerulonephritis. Przegl Lek 53(5): 443-453.

7. Madani A, Daryoush F, Esfahani TS, Mohseni Parvin, Atayee N, et al. (2003) Glomerular diseases in Iranian children: clinico-pathological correlation. Pediatr Nephrol 18(9): 925-928.

8. (1981) ISKDC the Primary Nephrotic Syndrome in children: Identification of patients with minimal change nephrotic syndrome from initial response to prednisone. J Pediatr 98(4): 561-564.

9. Bagga A, Srivastava RN, Nephrotic syndrome (2011) Pediatric Nephrology (5th edn) Jaypee Brothers Medical Publishers, New Delhi pp. 219-224.

10. Niaudut PBoyer O (2009) Idiopathic Nephrotic syndrome in children: Clinical Aspects. Pediatric Nephrology, 6th ed. Heidelberg:springer: pp 667-702.

11. Futrakul P, Yarudi S, Futrakul N, Seneviratana R, Pongsin P (1999) Tubular function and tubulointerstitial Disease. American Journal of Kidney Diseases 33(5): 886-891.

12. Deekajorndech T (2005) Fractional Excretion of Magnesium in Systemic Lupus Erythematosus. J Med Assoc Thai, 88(6): 743-745.

13. Futrakul N, Futrakul P (2008) Prevention of End-stage renal disease: An Innovative strategy. Thailand, chulalongkorn university printing house 2-31.

14. Al-Rasheed SA, Al-Mugeirem MM, Al-Salloum AA, Al-Sohaibani MO (1996) Childhood renal disease in Saudi Arabia. A clinico-pathological study of 167 cases. Int Urol Nephrol, 28(5): 607-613

15. Srivastava T, Simon SD(1999) Alon US. High incidence of FSGS in nephrotic syndrome of childhood. Pediatr Nephrol 13: 13-18.

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16. Gianoglio B (1998) From the Italian registry of Pediatric renal biopsies. Pediatr Nephrol 12: 225.

17. Deekajorndech TA (2007) Biomarker for Detecting Early Tubulointerstitial Disease and ischaemia in Glomerulonephropathy. Renal Failure 29: 1013-1017.

18. Futrakul N, Sila-asna M, Futrakul P (2007) Therapeutic Strategy towards renal restoration in chronic kidney disease. Asian Biomedicine 1(1):33-43.

19. Nath KA (1992) Tubulointerstitial changes as a major determinant in the progression of renal damage. Am J Kidney Dis 20(1): 1-17.

20. Oladipo OO, Onubi J, Awobusuyi O, A fonja OA (2003) Fractional excretion of magnesium of chronic renal failure patients in Lagos, Nigeria. Niger Postgrad Med J 10(3):131-134.

21. Futrakul N, Panichakul T, Sirisinha S, Futrakul P, Siriviriyakul P (2004) Glomerular endothelial dysfunction in chronic kidney disease. Renal Fail 26(3): 259-64.

22. Bhimma R, Adhikari M, Asharam K, Connolly C (2004) The spectrum of kidney disease (stage 2-5) in KwaZulu-Natal, South Africa. Pediatr Nephrol 23(10): 1841-1846.

23. Futrakul N, Yenrudi S, Sensirivatana R, Watana D, Laohapaibul A, et al. (2000) Peritubular capillary flow determines tubulointerstitial disease in idiopathic Nephrotic syndrome. Renal Fail 22(3): 329-335.