Correlation between the protein/creatinine ratio in spot urine and 24-hour urine protein

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ABSTRACT

INTRODUCTION: Measurement of the protein content in a 24-hour urine sample is the definitive method of establishing the presence of abnormal proteinuria. However, the urine collection is cumbersome. The spot urine protein to creatinine ratio seems to be a reliable diagnostic tool for urine protein measurement. Objective: Our aim was to evaluate the spot urine protein/creatinine ratio against 24-h urine total protein excretion in different proteinuria ranges by comparing samples collected simultaneously in patients of Hospital del Mar during the last year. Material and method: Observational, cross-sectional study of 159 consecutive paired determinations of 24-h urine total protein excretion and the spot urine protein/creatinine ratio (P/C) in renal patients. The strength of the correlation was determined by calculating the intraclass correlation coefficient (ICC) and the Spearman correlation coefficient (SCC). Results: Among all groups, there was a significant correlation between 24-hour proteinuria and the P/C ratio (SCC: r = 0.91, P < 0.001). The correlation in different levels of proteinuria were: <300mg (SCC: r = 0.498, P < 0.001; ICC: 0.46), 300-3499mg (SCC: r = 0.828, P < 0.001; ICC: 0.66) and > 3500mg (SCC: r = 0.181, P = NS; ICC: 0.18). Conclusions: In summary, a strong correlation was observed between spot urine protein/creatinine ratio and 24-h urine total protein excretion in proteinuria levels from 300mg/day to 3499mg/day. A lower correlation was also maintained in 24-h urine total protein <300mg. In our experience, there is no relevant correlation between spot urine protein/creatinine ratio and 24-h urine total protein excretion in nephrotic-range proteinuria.

Keywords: Renal insufficiency. Proteinuria. Nephrotic syndrome. Urine protein-creatinine ratio.

INTRODUCTION

Protein excretion varies in the course of the day; for this reason 24-hour proteinuria (Prot/24) has been considered as the classic reference method for protein determination.¹ The co-
lection of urine for 24 hours is tedious, and errors may occur during the process. For this reason, the protein/creatinine ratio (P/C) in spot urine was developed as a diagnostic alternative.1-3 Spot urine is collected during the second urination of the morning, and the first 20-25ml are discarded. After this, without disrupting urination, the middle urine is collected in the receptacle, and the last portion is also discarded.4 The potential error in determining proteins in a spot urine sample as a result of daily variation does not exceed the error in collecting a 24-hour urine sample.5 Although the correlation between P/C ratio and Prot/24 has been established, previous studies suggest that this correlation varies in accordance with different levels of proteinuria.4-6 Some of the most important studies are included in Table 1, with comments below in the Discussion section.

The aim of our study was to clarify whether the P/C ratio in spot urine, a sample easy to obtain and handle, is correlated with Prot/24 in samples collected consecutively from patients at our hospital.

METHODS

Patients

Prospective observational study between October 2010 and March 2011. A total of 159 patients over 18 years of age with different degrees of kidney failure. Proteinuria was determined in all patients as part of their examinations or clinical study. The study of Prot/24 and P/C ratio was undertaken simultaneously. All patients were recruited at an outpatient clinic or during an in-patient stay in the Nephrology Department of Hospital del Mar.

Method for determining proteinuria and creatinine

The patients were given instructions to collect urine for 24h and spot urine. After discarding the first urine of the day (which was included in the 24-hour urine sample), 3-5ml of urine were collected in the second sample for calculating the P/C ratio, which was calculated by dividing the proteinuria (mg/dl) of urine creatinine (g/dl). The 24-hour proteinuria was expressed in milligrams.

The creatinine concentration in urine was determined by Jaffé’s method for colorimetric and kinetic determination of creatinine, and the protein concentration in urine by the turbidimetric method, using in both cases an Auto Analyzer Hitachi Modular DPP (Roche).

Statistical analysis

The quantitative variables were expressed as mean and standard deviation, and the qualitative variables as percentages.

For the analysis of correlation between Prot/24 and the P/C ratio, the Spearman correlation coefficient was calculated (SCC). The analysis of diagnostic concordance was performed using the intraclass correlation coefficient (ICC) between the values of urinary proteinuria obtained using Prot/24 and P/C ratio on the same patient. The degree of concordance between both measurements was also analysed using the Bland and Altman method.7

The statistical analysis of the data was performed with the program SPSS version 18.0 for Windows. Statistically significance was considered when P<0.05.

RESULTS

The basic characteristics of the study population are shown in Table 2, both for the total population and for the population stratified according to degrees of Prot/24. Statistically significant differences were not observed in age and distribution by sexes among the 3 subgroups. The kidney function measured by serum creatinine was worse the degree of proteinuria was greater. The underlying kidney disease of the 159 patients is detailed in Table 2.

As shown in Table 3, a direct and statistically significant correlation was observed between Prot/24 and P/C in the whole group studied (n=159): with an ICC of 0.756 (confidence interval [CI] 95%, 0.680-0.816) and a SCC of r=0.91 (P<0.05) (Figure 1A). After stratifying by degrees of proteinuria, the correlation between Prot/24 and P/C ratio was maintained in the proteinuria range lower than 3500mg/24 hours, but this correlation was not observed in Prot/24 in the nephrotic range. In the Prot/24 lower than 300mg/24hours, the correlation with P/C was maintained, although it was of lesser intensity, with an ICC of 0.456 (CI 95%, 0.230-0.635) and SCC of r=0.498 (P<0.001) (Figure 1B). The patients with Prot/24 hours of 300-3449 showed a very good correlation with P/C, with ICC of 0.656 (CI 95%, 0.508-0.766) and SCC with r=0.828 (P<0.001). In patients with proteinuria in the nephrotic range, with Prot/24 ≥3500mg/24h, no correlation was observed with the P/C, showing an ICC of 0.340 (CI 95%, -0.041-0.650) and a SCC of r=0.181 (P= not significant [NS]) (Figure 1C).

As shown in Table 4, when stratifying according to the degree of kidney failure determined by estimated glomerular filtration rate (eGFR) calculated by the MDRD method in 3 subgroups (>60, 30-59 and <30ml/min/1.73 m²), we did not observe differences between Prot/24–P/C correlations calculated for each sub-group.

The degree of concordance between both measurements calculated using the Bland and Altman method4 showed that for proteinuria levels in the lower ranges the concordance was good, while for higher ranges the concordance practically disappeared.
Table 1. Main studies comparing 24-hour proteinuria with protein/creatinine ratio in spot urine

<table>
<thead>
<tr>
<th>Author, year, reference</th>
<th>No.</th>
<th>Population</th>
<th>Samples compared</th>
<th>Main results</th>
</tr>
</thead>
</table>
| Shaw AB, et al. 1983 | 81  | - Patients with kidney disease  
- Healthy individuals | - Prot/24  
- P/C  
- Albustix | - P/C offers a good estimate of Prot/24, and it is indicated for screening in patients who probably do not adequately collect 24-h urine. |
| Villafruela JJ, et al. 1990 | 452 | - Patients with kidney failure stratified according to eGFR  
- Sub-classification by: interstitial nephritis, glomerulonephritis, acute kidney failure and functioning kidney transplant | - Prot/24  
- P/C | - Correlation influenced by types of renal disease, degree of deterioration in kidney function and degree of proteinuria  
- Weak correlation in cases of severe kidney failure, interstitial nephritis or severe proteinuria. |
- Persistent proteinuria (>1g/d during >3 months).  
- eGFR 20-70ml/min/1.73 m² | - P/C  
- Prot/24  
- eGFR | - First urine P/C is correlated with Prot/24.  
- P/C has greater predictive power for progression of kidney disease. |
| Price CP, et al. 2005 | Systematic review of 16 studies | - Pregnant women  
- Kidney disease  
- Kidney transplants  
- From Rheumatology visits | - P/C  
- Prot/24 | - High degree of correlation between P/C and Prot/24.  
- P/C can be used for proteinuria screening. |
| Antunes VV, et al. 2008 | 41, (246 urine samples, 6 per patient) | - Glomerular disease with:  
- Nephrotic syndrome due to focal and segmental glomerulonephritis or membranous nephropathy  
- Non-nephrotic proteinuria | - Prot/24  
- P/C  
- Plasma creatinine | - P/C is a precise method for determining proteinuria.  
- The greater the degree of proteinuria (determined by the Pearson correlation coefficient) the less the correlation between Prot/24 and P/C. |
| Guy M, et al. 2009 | 86  | - Diabetic nephropathy or  
- Damage due to AHT or  
- Renovascular disease or  
- Glomerulonephritis/glomerulosclerosis | - Prot/24  
- P/C urine, 1st hour, 2nd, 3rd, 4th sample  
- A/C urine, 1st hour, 2nd, 3rd, 4th sample | - Strong correlation between Prot/24 and P/C of 2nd and 3rd sample.  
- A/C presents strong correlation with Prot/24 when this is >1g/d. |
| Lambers Heerspink HJ, et al. 2010 | 701 | - Diabetic nephropathy, participating in RENAAL (Reduction of Endpoints in Non Insulin Dependent Diabetes Mellitus with Losartan) | - Prot/24  
- P/C  
- A/C | - A/C showed the strongest association with the risk of renal events.  
- Prot/24 has a weaker association; after calculation of P/C the predictive value increases. |
| Methven S, et al. 2010 | 6842 | - Interstitial disease  
- Multisystemic disease  
- Diabetic nephropathy  
- Primary glomerulonephritis  
- Chronic kidney disease of unknown cause | - P/C  
- A/C  
- Prot/24 | - P/C is more sensitive and more specific for the detection of proteinuria >0.5-1g/d  
- Total proteinuria is not accurately determined by A/C due to the variable elimination of other type of proteins.  
- A/C and P/C vary substantially according to age, gender, eGFR (effect related with muscle mass). |
| Methven S, et al. 2011 | 5586 | - >18 years  
- Without beginning kidney replacement therapy (including transplant)  
- Follow-up during >1 year | - P/C  
- A/C  
- Prot/24 | - Prot/24 predicts mortality and vascular events as equally as P/C and A/C.  
- A/C and P/C can stratify cardiovascular risk in kidney disease patients equally.  
- P/C is strongly correlated with low levels of Prot/24 (where it was traditionally considered that A/C was better). |
| Methven S, et al. 2011 | 5586 | - >18 years  
- Without beginning kidney replacement therapy (including transplant)  
- Follow-up during >1 year | - P/C  
- A/C | - Screening with P/C identifies additional 16% of patients with significant proteinuria, not identified with A/C. |

A/C: albumin-creatinine ratio in spot urine; eGFR: estimated glomerular filtration; AHT: arterial hypertension; P/C: protein/creatinine ratio in spot urine sample; Prot/24: 24-hour proteinuria.
DISCUSSION

Our study demonstrated that the P/C ratio has a strong correlation with the Prot/24 for values lower than 3500mg; nevertheless, we did not observe a correlation for proteinuria in the nephrotic range. Hence, in the group studied at our hospital, as the degree of Prot/24 increases, the degree of correlation both using the ICC and the SCC analyses decreases.

The adequate detection and quantification of proteinuria is of great importance in the management of patients with kidney disease. The collection of 24h urine is complicated, and it is not always performed correctly; an easy and reliable method is the once-only collection of urine for the measurement of the P/C ratio. Different studies have analysed the correlation between Prot/24 and the P/C ratio in spot urine.1,2,4-6,10-12,15 Our observational study confirmed the results obtained in other cross-sectional, observational studies, supporting the use of the P/C ratio in clinical practice due to the simplicity of collecting the sample and its low cost for Prot/24 lower than 3500mg.3,4,6,7,11,12

The usefulness of this ratio for proteinuria within the nephrotic range has been studied previously, but this

### Table 2. Characteristics of patients undergoing proteinuria determinations

<table>
<thead>
<tr>
<th>Variables</th>
<th>Proteinuria/24 hours (mg/24 h)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All &lt;300 300-3499 &gt; _3500</td>
</tr>
<tr>
<td>n</td>
<td>159 60 77 22</td>
</tr>
<tr>
<td>Men, no. (%)</td>
<td>93 (58.4) 30 (50) 49 (59.7) 17 (77.3)</td>
</tr>
<tr>
<td>Age (average ±SD, years)</td>
<td>59.7 ± 14.7 58.45 ± 15.5 61.14 ± 14.1 57.77 ± 14.46</td>
</tr>
<tr>
<td>Proteinuria (average and range, mg/24 h)</td>
<td>(33-17168) (33-294) (300-3332) (3503-17168)</td>
</tr>
<tr>
<td>Spot urine protein/creatinine (average and range)</td>
<td>(36.35-11652.41) (36.35-686.58) (99.05-5590.26) (1204.5-11652.41)</td>
</tr>
<tr>
<td>Serum creatinine (average ±SD, mg/dl)</td>
<td>1.96 ± 1.59 1.43 ± 1 2.12 ± 1.77 2.73 ± 1.72</td>
</tr>
</tbody>
</table>

SD: standard deviation.

### Table 3. Intraclass correlation coefficient and Spearman correlation coefficient between the values for 24-hour proteinuria and the protein/creatinine ratio in sporadic urine

<table>
<thead>
<tr>
<th>Proteinuria 24h (mg)</th>
<th>All &lt;300 300-3499 &gt; _3500</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nº</td>
<td>159 60 77 22</td>
</tr>
<tr>
<td>ICC</td>
<td>0.756 0.456 0.656 0.340</td>
</tr>
<tr>
<td>(CI 95%)</td>
<td>(0.680-0.816) (0.230-0.635) (0.508-0.766) (-0.041-0.650)</td>
</tr>
<tr>
<td>SCC (r)</td>
<td>0.91 0.498 0.828 0.181</td>
</tr>
<tr>
<td>P</td>
<td>&lt; 0.05 &lt; 0.001 &lt; 0.001 0.420</td>
</tr>
</tbody>
</table>

ICC: intraclass correlation coefficient; SCC: Spearman correlation coefficient; CI: confidence interval; n: sample.
was not the main objective of our study. In our population, for proteinuria in the nephrotic range we did not observe a correlation between the P/C ratio and Prot/24. In this respect, Antunes et al. have also demonstrated that, the greater the proteinuria, the lesser the correlation and adjustment between the different methods. Using the Bland and Altman method, it was observed that the Prot/24 closest to the nephrotic range proteinuria, there is less concordance between the methods (Figure 2).

Ruggenenti et al.11, in their study designed to compare different factors for predicting the risk of progression of kidney failure in non-diabetic patients, observed a correlation between P/C and Prot/24 in all degrees of proteinuria, which decreased as the P/C increased. However, they did not define the degree of proteinuria in which the differences appeared. Their study concluded that the P/C ratio predicted the risk of progression of kidney failure with more precision than the Prot/24. In patients with primary glomerulopathies, Morales et al. also detected this decrease in correlation, and they defined the best cut-off point for determining nephrotic range proteinuria as P/C >2.6 (considering the Prot/24 as standard reference). These authors found good correlation and agreement between P/C and Prot/24 for all renal function levels, but demonstrated more marked differences as urinary protein excretion increased.11 In our study, when stratifying by kidney function calculated using MDRD-4, both determinations presented a very strong correlation with the different models divided into three tertiles (>60, 59-30 and <30 ml/min/1.73 m²).

**Figure 1. Correlation between the protein/creatinine ratio in sporadic urine and 24-hour proteinuria**

A) total study sample; B) 24-hour proteinuria (Prot/24) <300 mg; C) 24-hour proteinuria >3500 mg.
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Table 4. Intraclass correlation coefficient and Spearman correlation coefficient between the values for 24-hour proteinuria and the protein/creatinine ratio in spot urine according to estimated glomerular filtration rate

<table>
<thead>
<tr>
<th>eGFR by MDRD ml/min/1.73 m²</th>
<th>≥ 60</th>
<th>59-30</th>
<th>&lt; 30</th>
</tr>
</thead>
<tbody>
<tr>
<td>No.</td>
<td>62</td>
<td>49</td>
<td>39</td>
</tr>
<tr>
<td>ICC (CI 95%)</td>
<td>0.661 (0.494-0.781)</td>
<td>0.681 (0.497-0.807)</td>
<td>0.793 (0.639-0.886)</td>
</tr>
<tr>
<td>SCC (r)</td>
<td>0.88</td>
<td>0.899</td>
<td>0.901</td>
</tr>
<tr>
<td>P</td>
<td>&lt; 0.001</td>
<td>&lt; 0.001</td>
<td>&lt; 0.001</td>
</tr>
</tbody>
</table>

ICC: intraclass correlation coefficient; SCC: Spearman correlation coefficient; CI: confidence interval; eGFR: estimated glomerular filtration, n: sample.

less, Methyen et al., in a sub-analysis focused on studying the capacity of P/C ratio and albumin/creatinine ratio in urine for proteinuria of 1000mg/24 hours, demonstrated that the technical differences varied substantially according to age, gender and the eGFR, an effect that they related directly with the muscle mass. ²

Figure 2. Bland and Altman plot showing concordance between 24-hour proteinuria and the protein/creatinine ratio in sporadic urine

Difference between 24-hour proteinuria and the protein/creatinine (P/C) ratio in spot urine (x-axis) versus the average proteinuria (y-axis). In the event that there is no systematic error, the points are distributed randomly on both sides of the straight line corresponding to the difference 0 between measurements. The thick line represents the average of the differences (systematic error of the second method versus the first one.) The dotted line represents the 95% confidence limits for that difference.
Some of the current clinical practice guidelines such as the KDOQI, CARI, KDIGO, UK Renal Association, NICE, and CSN define the presence of proteinuria according to the P/C ratio (different value according to the Scientific Society). On the other hand, other guidelines such as the SEN-semFYC and the ADA define the appearance of proteinuria according to the urine albumin/creatinine ratio. Recently, a consensus document was published on the evaluation of proteinuria in the diagnosis and follow-up of chronic kidney disease patients that determined that in the detection and monitoring of proteinuria or albuminuria, it was not necessary to collect 24h urine. In our study, given the difficulty of finding uniformity in the different guidelines and scientific societies for the initial determination of P/C or urine albumin/creatinine, we focused on the analysis of the P/C ratio. Nevertheless, in patients with suspicion of incipient diabetic nephropathy, just as the guidelines unanimously recommend, we advise the determination of albumin, expressed as albumin/creatinine in urine, once a year.

This study focused on determining the correlation between the P/C ratio in spot urine and the Prot/24 and showed an acceptable correlation when Prot/24 values are less than 3500mg. Hence, the simplification of the collection and subsequent calculation of the ratio in patients with proteinuria within that range could result in lower health care costs. Studies focused on analysing the save in health care costs by substituting the Prot/24 for the P/C could be of great use in the future.

Our study had a series of limitations: the main limitation is the need to increase the number of patients studied, since it was too small, specifically in the sub-group of patients who showed proteinuria within the nephrotic range. Another limitation is that no cause of kidney disease was excluded, including also a sample of patients with kidney transplants and pre-eclampsia patients. Increasing the number of samples collected and perhaps stratifying by underlying kidney pathology would help acquire a better knowledge of the correlation between the two techniques studied.

In conclusion, our study shows that the P/C ratio in spot urine is useful for Prot/24 less than 3500mg. The usefulness of measuring this parameter in cases with proteinuria within the nephrotic range is yet to be confirmed. In the same manner, future studies focused on monitoring patients with proteinuria within the nephrotic range can be useful for evaluating the efficiency of the P/C ratio in detecting complete or partial remission of proteinuria during the monitoring of the underlying renal disease.

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Conflicts of interest
The authors affirm that they have no conflicts of interest related to the content of this article.

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