Comparison of Spot Urine Protein/Creatinine Ratio, Spot Urine Protein/Osmolality Ratio with Measured 24-Hour Urine Protein in HIV Subjects in Nigeria

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Abstract

Background and Objectives: Urine protein examination is a veritable tool in the management of renal diseases. Proteinuria estimation from 24-hour urine collection is the gold standard. Prompt urine protein assessment from spot urine sample has become necessary to eliminate inaccuracies inherent in timed urine collection. This study aims at comparing spot urine protein/creatinine ratio (SUPCR) and measured 24-hour urine protein (24 HUP), and also spot urine protein/osmolality ratio (SUPOR) and 24 HUP against measured 24-hour urine protein (24 HUP) for assessment of proteinuria in human immunodeficiency virus (HIV) subjects.

Methodology: Three hundred and ninety three HIV subjects from the HIV/AIDS clinic and 136 age- and sex-matched non-HIV subjects as Control. Investigations performed included 24-hour urine protein (24 HUP), spot urine protein (SUP), spot urine creatinine (SUCr), spot urine osmolality (SUOsm), fasting blood sugar (FBS), urinalysis, HIV screening and confirmatory test, electrolyte, urea and creatinine. SUPCR and SUPOR were calculated. Correlation statistics, 2 × 2 contingency table analysis, receiver operator characteristics (ROC) Curve analysis and Bland Altman plots were used to compare SUPCR and 24 HUP, and also SUPOR and 24 HUP in HIV subjects and control. SPSS version 17 and Medical statistical software were used to analyze the data. P<0.050 was taken as statistically significant.

Results: Using the 2 × 2 contingency table in the HIV subjects, the Sensitivity for SUPCR and SUPOR with 24 HUP was 43.4% and 11.5% respectively. Specificity for SUPCR and SUPOR with 24 HUP was 92.9% and 99.2% respectively. The SUPCR had a correlation (r) of 0.734 (p<0.001) with 24 HUP. In addition, SUPOR had a correlation coefficient of 0.417 (p<0.001) with 24 HUP. Using the Bland Altman plots, SUPCR compared with 24HUP limits of agreement were +0.361 g/day to -0.248 g/day in HIV subjects. In addition for SUPOR the limits of agreements were +0.440 g/day to -0.180 g/day in HIV subjects. The bias was 0.060 g/day and 0.130 g/day for SUPCR and SUPOR respectively. The receiver operating characteristics (ROC) curve showed that SUPCR randomly chosen value of 0.042 mg/mg and SUPOR chosen value of 0.010 mg/dl/mOsm/kg H₂O predicted 24 HUP at urinary excretion threshold of 0.150 g.

Conclusion: The SUPCR and SUPOR are reliable tests, for quantifying proteinuria in HIV subjects, and should be used in assessment of proteinuria in HIV subjects in Sub Saharan African countries.

Keywords: Proteinuria; HIV; 24 HUP; SUPCR; SUPOR; Bland Altman plot; ROC curve

Introduction

Chronic kidney disease (CKD) is defined by the presence of kidney damage or glomerular filtration rate (GFR) less than 60 ml/min/1.73 m² (1.0 ml/s/1.73 m²) for three or more months, irrespective of cause [1]. HIV infection is commonly associated with chronic kidney disease [2]. Accurate and fast diagnostic tools are necessary to diagnose and monitor HIV patients who may have CKD. In majority of subjects with CKD, the presence of CKD can be detected with 2 tests. These tests include a urine test for the detection of proteinuria and a blood test to estimate the GFR. These two tests are commonly used by Nephrologists for detection of CKD in Sub Saharan Africa and other parts of the world.

Proteinuria is the commonest marker of CKD in adults and contributes to its progression by several mechanisms; and appropriate interventions that reduce proteinuria also improve patient outcome [3]. Urine protein estimation is one of the key parameters in the diagnosis and monitoring of renal functions in HIV-related renal disease states [4]. Evaluation of protein from 24-hour urine collection collection is the traditional and gold standard diagnostic test for quantification of proteinuria in the general population including in HIV subjects. The 24 HUP sample collection is often difficult, tedious, time-consuming, and is also riddled with errors [5,6]. Due to the problems associated with 24

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HUP collection, some alternatives for the quantification of proteinuria in HIV seropositive subjects have been considered. The alternatives include urinary dipsticks protein estimation, SUPCR, and SUPOR.

Dipstick urinalysis protein estimation is known to have a poor sensitivity and specificity profile for establishing significant proteinuria or quantitative change in proteinuria [7]. Measurement of albumin or total protein concentration in a spot sample by tests like SUPCR and SUPOR avoids the need for collection of a timed urine specimen but is affected by some factors including the state of hydration, sex, and age [4]. However, some studies have demonstrated that SUPCR in a random, single voided urine sample reflects 24-hour urine protein [8-10]. Random and morning urinary protein/osmolality ratio has also been found to reflect 24 HUP excretion [11].

There is paucity of data on HIV subjects comparing SUPGR against 24 HUP and SUPOR against 24 HUP, emanating from Sub Saharan Africa. In addition, with the increasing incidence of HIV infection in the region, it is pertinent that faster, cheaper, simpler, and evidence-based reliable methods of quantifying proteinuria in HIV patients be adopted. This will help in no small measure to improve the management of this group of patients in Sub Saharan African countries. Bearing this in mind we decided to carry out this study to validate the use of SUPCR, and SUPOR in quantifying proteinuria in HIV subjects, by comparing each of them respectively, with 24 HUP.

Materials and Methods

This was a cross-sectional comparative study of SUPCR and SUPOR against the “gold Standard,” 24 HUP, in newly-diagnosed HIV-seropositive subjects. The study was carried out between March, 2011 and August, 2011 in Federal Medical Centre, Owerri (FMC), Imo State, Nigeria. The subjects consisted of 393 newly-diagnosed, drug-naïve, HIV-seropositive adult subjects within the age range of 18-65 years, and 136 age- and sex-matched HIV-sero-negative subjects as control. Exclusion criteria included subject below 18 years or above 65 years, febrile illnesses, evidence of heart failure, urinary tract infection, diabetes mellitus, hypertension, evidence of urological disease, and use of drugs that could interfere with urinary creatinine excretion. The same exclusion criteria applied to the control group. An interviewer structured questionnaire was administered and relevant data collected. These included patient’s age, sex, diagnosis and co-morbidities (diabetic mellitus, hypertension etc.).

Approval for this study was obtained from the Ethical and Research Committee of FMC Owerri. Investigations performed by the subjects included: HIV screening and confirmatory tests, SUP, SUCr, SUOsm, serum creatinine, 24 HUP, FBS, and urinalysis.

Clear instructions were given to all the subjects on how to collect 24-hour urine sample. 24-hour urine protein was measured in the urine samples thus collected. A day-time random spot urine sample was collected. A 24-hour random spot urine sample was used in calculating SUPCR and SUPOR. This was to facilitate rapid analysis in our laboratory [12]. Urine protein was measured by photometric method, urine osmolality by freezing point depression method (using Precision System Osmette 5002 osmometer) and creatinine by modified Jaffe method. All the laboratory tests were carried out in the laboratory of FMC, Owerri, while urine osmolality tests were done by E.N Anyabolu. FMC, Owerri, is one of two tertiary health institutions in the State, and has a good number of laboratory scientists. SUPCR and SUPOR were determined.

Statistical analysis

SPSS version 17.0 (SPSS Inc., Chicago, IL, USA) statistical software was used to analyze the data. The 2 × 2 contingency tables were used to calculate the sensitivity, specificity, positive predictive value, negative predictive value and accuracy obtained. The cut-off value for the 2 by 2 table was 24 HUP 0.150 g, SUPCR 0.150 mg/mg, SUPOR 0.150 mg/dL/mOsm/kg H2O. Predictive indices of SUPCR and SUPOR for 24 HUP were determined. The strength of association between the SUPCR and against 24 HUP, and also SUPOR and against 24 HUP was determined by correlation statistics. P<0.05 was taken as statistically significant. Bland Altman’s plot was used to compare SUPCR, SUPOR, each with 24 HUP (Medcalc statistical software) BLAND ALTMAN [13]. Cutoff values, sensitivity and specificity of SUPCR and SUPOR were assessed for predicting 24 HUP excretion “threshold” of 0.150 g/day by ROC curve. These four instruments, namely 2 × 2 contingency table analysis, correlation statistics, receiver operator characteristics (ROC) curve analysis and Bland Altman plots were used to compare SUPCR, SUPOR with 24 HUP in HIV subjects and Control. P<0.05 was taken as statistically significant.

Results

The mean age of the HIV subjects was 39 ± 11 years, and their age ranged between 18 and 65 years. Out of 393 HIV subjects, 283 (72.0%) were females, while 110 (28.0%) were males. The control consisted 136 subjects, of these 98 (72.1%) were females, while 38 (27.9%) were males.

The female/male ratio was approximately 3:1 in both the HIV subjects and control subjects. However, 22 subjects were excluded from the study on account of inadequate 24-hour urine collection.

In the HIV subjects, the mean 24HUP, SUPCR, and SUPOR were 0.187 ± 0.290 g, 0.133 ± 0.371 mg/mg, and 0.035 ± 0.050 mg/dl/mOsm/kg H2O respectively. In the Control, the mean 24 HUP, SUPCR, and SUPOR were: 0.095 ± 0.087 g, 0.082 ± 0.163 mg/mg, 0.042 ± 0.135 mg/dl/mOsm/kg H2O respectively. There was statistically significant difference using 24 HUP in the HIV subjects and the Controls, p<0.001. In contrast, SUPCR and SUPOR showed no statistically significant difference between the HIV subjects and the Control, p=0.120 and p=0.357 respectively (Table 1).

Using the 2 × 2 contingency table, in HIV subjects, the sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV), and accuracy for SUPCR were 43%, 92.9%, 74.6%, 92.8% and 76.8% respectively. In SUPOR, the sensitivity, specificity, PPV, NPV, and accuracy were 11%, 99.2%, 87.5%, 69.9%, and 70.6% respectively. In addition, in the control the sensitivity, specificity, PPV, NPV, and

<table>
<thead>
<tr>
<th>Variable (mean)</th>
<th>All subjects (n=529)</th>
<th>HIV subjects (n=393)</th>
<th>Controls (n=136)</th>
<th>p value (n=136)</th>
</tr>
</thead>
<tbody>
<tr>
<td>24HUP (g ± SD)</td>
<td>0.162 ± 0.256</td>
<td>0.187 ± 0.290</td>
<td>0.095 ± 0.087</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>SUPCR (mg/mg ± SD)</td>
<td>0.120 ± 0.330</td>
<td>0.133 ± 0.371</td>
<td>0.082 ± 0.163</td>
<td>0.120</td>
</tr>
<tr>
<td>SUPOR (mg/dl/mOsm/Kg H2O ± SD)</td>
<td>0.037 ± 0.081</td>
<td>0.035 ± 0.050</td>
<td>0.042 ± 0.135</td>
<td>0.357</td>
</tr>
</tbody>
</table>

Table 1: Daily urine protein estimation of the study population.
accuracy for SUPCR were 20.0%, 95.6%, 44.4%, 87.4%, and 84.5% respectively. The values for SUPOR for sensitivity, specificity, PPV, NPV, and accuracy were; 15.0% 98.2%, 60.0%, 87.0%, and 86.0% respectively.

Table 2 shows the correlation of 24 HUP with SUPCR and SUPOR respectively, in HIV subjects. SUPCR was found to had a correlation coefficient of 0.734 (p<0.001). SUPOR was found to have a correlation coefficient of 0.417 (p=0.001) in HIV subjects. In addition, in the Control, SUPCR was found to have a correlation of 0.518 (p<0.001), SUPOR also had a correlation 0.336 (p<0.001).

Figures 1A-1D showed the Bland Altman plots for SUPCR, and SUPOR when compared individually with 24 HUP in HIV and control subjects. Tables 3,4 summarize the findings.

Figures 1E,1G showed ROC of SUPCR and SUPOR test result variables against 24 HUP as standard in study population. The positive actual state or cutoff was ≥0.150 g. In HIV subjects, SUPCR had an AUROC (area under receiver operator curve) of 0.678, p<0.001, confidence interval 95% (0.617 to 0.738), while SUPOR had an AUROC of 0.678, p<0.001, confidence interval 95% (0.619 to 0.736). In the controls, SUPCR had an AUROC of 0.753, p<0.001, confidence interval 95% (0.633 to 0.872), while SUPOR had an AUROC of 0.614, p<0.001, confidence interval 95% (0.485 to 0.744) The ROC analysis showed that SUPCR of 0.042 mg/mg (sensitivity 91% and specificity 90%) and SUPOR value of 0.010 mg/dl/mOsm/kg H₂O (sensitivity 91%, and specificity 90%) predicted 24 HUP at 0.150 g.
Discussion

This study compared the SUPCR and SUPOR against "gold standard" 24 HUP in HIV subjects. In the 2 by 2 table analysis in HIV subjects at cut off of 0.150 for the three methods, the sensitivity for SUPCR, and SUPOR was, 43%, and 11% respectively, while the specificity was 92%, and 99.2% respectively in predicting 24 HUP. These shows that both spot urine tests had low sensitivity and high specificity. This tends to suggest that they are both poor screening tests, good diagnostic tests. Similar results were also obtained in the control. We did not, however, evaluate these predictive indices of SUPCR and SUPOR for 24 HUP at higher cut-off values of proteinuria.

The correlation of SUPCR and SUPOR, with 24 HUP, respectively, in the HIV subjects, showed This study showed that SUPCR had a high correlation coefficient (0.734) with 24 HUP (r=0.734, p<0.001) , while SUPOR had a moderate correlation (0.417) with 24 HUP (r=0.417, p<0.001) in HIV subjects. The p values were significant in both cases. In the Control the SUPCR had a moderate correlation (0.518), while SUPOR had a mild (0.336) correlation. This correlation values, of SUPCR with 24 HUP, in HIV subjects, where slightly lower than those reported in some studies [14-18]. None the less one of these studies was conducted in an HIV population, while the other was in a general population. This study also showed lower correlation values for SUPCR and 24 HUP, compared with those of two previous studies [11,12].

Bland-Altman analysis is typically used to compare measurement techniques against a reference value, usually an accepted gold standard.

Table 3: Summary of Bland Altman Plots in HIV subjects.

<table>
<thead>
<tr>
<th>g/day</th>
<th>SUPCR v 24HUP</th>
<th>SUPOR v 24HUP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bias</td>
<td>0.060</td>
<td>0.130</td>
</tr>
<tr>
<td>Precision (+)</td>
<td>0.214</td>
<td>0.289</td>
</tr>
<tr>
<td>Precision (-)</td>
<td>-0.098</td>
<td>0.164</td>
</tr>
<tr>
<td>Limits of Agreement (+)</td>
<td>0.361</td>
<td>0.440</td>
</tr>
<tr>
<td>Limits of Agreement (-)</td>
<td>0.248</td>
<td>0.180</td>
</tr>
</tbody>
</table>

24HUP: 24-Hour Urine Protein; SUPCR: Spot Urine Protein/Creatinine Ratio; SUPOR: Spot Urine Protein/Osmolality Ratio; SD: Standard Deviation.

Table 4: Summary of Bland Altman plots in the control subjects.

<table>
<thead>
<tr>
<th>g/day</th>
<th>SUPCR v 24HUP</th>
<th>SUPOR v 24HUP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bias</td>
<td>0.030</td>
<td>0.070</td>
</tr>
<tr>
<td>Precision (+)</td>
<td>0.125</td>
<td>0.164</td>
</tr>
<tr>
<td>Precision (-)</td>
<td>0.067</td>
<td>0.30</td>
</tr>
<tr>
<td>Limits of Agreement (+)</td>
<td>0.217</td>
<td>0.258</td>
</tr>
<tr>
<td>Limits of Agreement (-)</td>
<td>0.159</td>
<td>0.124</td>
</tr>
</tbody>
</table>
We want to know by how much the new method is likely to differ from the old; if this is not enough to cause problems in clinical interpretation we can replace the old method by the new or use the two interchangeably. 

[13] Bland Altman plot was used in this study to compare SUPCR and SUPOR against 24HUP respectively. The bias and precisions were low, and though good correlation was observed between SUPCR, SUPOR with 24 HUP in this study, it is expedient that reasonable limits of agreement be demonstrated for them to be used interchangeably. The 95% limits of agreements values were within acceptable range.

We found, in this study, for both SUPCR and SUPOR, when each was compared with 24 HUP, that the results were similar within 95% limits of agreement. These findings, and correlation, strongly suggest that both tests may be used interchangeably with the 24 HUP, especially the SUPCR that had a high correlation with 24 HUP in HIV subjects. Similar results were reported by previous studies which also showed that SUPCR and 24 HUP could be used interchangeably in HIV subjects and in the general population [14,18]. This is in keeping with findings from some previous studies [14,18].

The predictive indices of SUPCR and SUPOR against measured 24 HUP in HIV were also evaluated using the ROC curves. Based on ROC comparison, our study showed low sensitivity and specificity for SUPCR (at values ≥0.150 mg/mg) and SUPOR (at values ≥0.150 mg/dl/mOsm/ Kg H2O) methods in predicting 24 HUP of ≥0.150 g in HIV subjects. There was no difference between SUPCR and SUPOR in predicting 24 HUP at this level. This is similar to the findings reported by two studies with slightly higher AUROC [19,20]. This is in keeping with some earlier studies [19,20]. However SUPCR was found to be better than SUPOR when compared with 24 HUP in another study [21]. In addition, with further evaluation in our study, we found that in SUPCR, of 0.042 mg/mg represented the best threshold to reliably predict 24 HUP of 0.150 g, with high sensitivity, and high specificity in this study. This is consistent with previously published reports [20-22]. In addition, SUPOR of 0.01 mg/dl/mOsm/kg H2O represented the best threshold to reliably predict 24 HUP of 0.150 g, with high sensitivity and high specificity. This is similar to the study by Wilson et al. [11] in which they found SUPCR of 0.120 with sensitivity of 95% and specificity of 93% to represent normal range proteinuria in 24-hour urine collection. This compared to a SUPCR of less than 0.05 (sensitivity 96%, specificity 90%).

**Conclusion**

SUPCR is a fast, convenient and reliable method of estimating proteinuria in both HIV and non HIV population from the correlation, Bland Altman plot analysis and ROC curves. The same applies to the SUPOR test. HIV clinics in Sub-Saharan African County may use the tests in quantifying proteinuria in this group of subjects. However, a conversion equation for SUPCR and SUPOR are needed for effective use of both methods in quantitative urine protein estimation.

**Limitations of the Study**

The sample size is relatively small. A larger sample size preferably involving many centers is desirable. Subjects that took part in the study were not on admission; as a result compliance with strict urine collection may have been compromised by some of the subjects. Ideally the urine should be stored in a fridge at a temperature 2-6°C during the collection of the urine, it was not certain that all patient complied strictly with this guideline.

**References**


