A path analysis to investigate the interaction between serum, urinary and demographic factors influencing urine calcium in kidney stone formers

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Abstract

Background Hypercalciuria is one of the most important urinary risk factors in kidney stone formers. This study aimed to delineate the interaction of some demographic, serum, and urinary risk factors influencing 24-h urinary (24-U) calcium excretion.

Methods This study was secondary data analysis, using data from 593 kidney stone patients referred to the Labbafinejad kidney stone prevention clinic from March 2015 to May 2019. The study considered serum, urinary and demographic factors that interact to influence 24-U calcium using path analysis. In addition to the direct impact of predictors on the 24-U calcium, this analysis considered the effects of the predictors on the 24-U calcium transmitted by a mediating variable named indirect effects.

Results The results showed that age indirectly affected on 24-U calcium through 25-hydroxy vitamin D (25(OH)D), serum and 24-U creatinine. As well, weight had an indirect effect through 24-urine metabolites (creatinine, citrate, urea, and sodium). Among serum variables, PTH and creatinine significantly directly affected on 24-U calcium. In comparison, 25(OH)D and phosphorus appeared to influence 24-U calcium indirectly through serum parathormone. Regarding 24-U metabolites, sodium, urea, and citrate had a significant direct effect on 24-U calcium. Moreover, 24-U creatinine has a significant direct and indirect effect on 24-U calcium through citrate and urea as mediator variables.

Conclusion Serum 25(OH)D and phosphorus, along with age and weight, indirectly affected urinary calcium through a third variable. Other variables (PTH, serum creatinine, and 24-U sodium, urea, and citrate) showed a direct effect on 24-U calcium excretion.

Keywords Kidney stones · Hypercalciuria · Path analysis · Vitamin D · Urine calcium

Background

In recent decades, the incidence and prevalence of kidney stones have increased worldwide. Urolithiasis is a common cause of morbidity with a lifetime risk of 5–12% in Europe and the USA [1, 2]. Since kidney stones are a potential risk factor for irreversible renal function loss with a high disease burden, many studies have been conducted to explore kidney stone formation's pathophysiology [3].

The most common component of nearly 80% of kidney stones is made of calcium in the form of calcium oxalate and, to a lesser extent, calcium phosphate [4]. Therefore, controlling excess calcium excretion in the urine (hypercalciuria) is very important for kidney stone prevention [4]. Hypercalciuria is one of the most common urine metabolic abnormalities among calcium stone formers, with a prevalence of 35–65%. Excess calcium excretion in the urine could be responsible for the supersaturated status of urinary calcium salts and lead to crystallization [5].

Two hormones play a crucial role in calcium homeostasis: parathyroid hormone (PTH) and vitamin D. It is noteworthy that urine calcium excretion is affected directly or indirectly by various factors (such as demographic or metabolic) that...
are not entirely known [6–8]. An in-depth understanding of this complex interaction between different variables that regulate calcium homeostasis could help to understand the pathophysiology of idiopathic hypercalciuria and improve its management. However, population-based data on this area is scarce, and the complex direct and indirect effects (through third variables) of serum and urinary metabolites on urine calcium have surprisingly not been fully explored. Thus, this study aims to assess the interrelationship between serum, urinary and demographic factors that may affect urine calcium in a population of calcium stone formers. Using pathway analysis, we hypothesize and test how urine metabolites and serum variables may, directly and indirectly, influence urine calcium.

Methods

This study was a secondary data analysis using the baseline data of our previous study [9]. This retrospective study included data from kidney stone patients referred to the Labbafinejad kidney stone prevention clinic from March 2015 to May 2019. The participants included in the study were patients ≥ 18 years old who had undergone complete metabolic workup of serum metabolites and valid 24-h urine analysis for kidney stone prevention. For path analysis, we encompass all factors that are directly or indirectly involved in calcium metabolism, including age and weight, the twenty-four-hour urine metabolites in terms of creatinine, urea, citrate, sodium, as well as serum variables including phosphorus (P), PTH, calcium (Ca), 25(OH)D, and creatinine (Cr). Patients who had a known history of primary hyperparathyroidism or any chronic diseases that could affect urine calcium (such as sarcoidosis), had a history of chronic kidney disease, consuming any medication interfering with calcium hemostasis three months before the assessment (such as thiazides, steroids, or calcium supplements), were pregnant or lactating, or had urinary tract infection at the time of assessment were excluded.

Path Analysis, an extension of the regression model, is a statistical technique used to examine causal relationships between two or more variables, and its pathways represent the hypotheses of researchers. This analysis provides a comprehensive picture of the associations between the predictors and dependent variable. Here, path analysis was used to determine the pathways by which urine metabolites and serum variables interact to influence the primary outcome variable, 24-U Ca. The path model, presented in Fig. 1, was developed by review the articles [8], textbooks [4] and the consensus of experts who specialize in various aspects of kidney stone prevention [10].

In this model, the direct effect represents the direct impact of one predictor on the outcome while not transmitted through a third variable. The indirect effect is the impact of a predictor on the outcome transmitted by a third variable. The total effect of a predictor on the response is obtained by summing direct and indirect effects related to different pathways. The size of the effects is reported both in standardized and unstandardized terms. Unstandardized coefficients used the original variables’ measurement scale and indicated the strength of the relationship between variables. Standardized coefficients were values transformed into the same unit and used to compare the effects of variables on the response. The study was carried out in accordance with the guidelines of the Helsinki Declaration (Fortaleza, Brazil, October 2013).

Statistical methods

Descriptive statistics including frequency and percentage for qualitative variables and means with standard deviations for quantitative ones were used. The mean of quantitative variables between patients with 24-U Ca less than 200 mg and 24-U Ca more than 200 mg was compared using independent samples t-test. Pearson correlation was calculated to determine the relationship between urine metabolites and serum variables. The main analytic plan, path analysis, involved the primary outcome variable, 24-U Ca, and the predictor variables of interest, including age, weight, 24-urine metabolites and serum variable. The adequacy of the path model was tested using the Relative Chi-Square (CMIN/DF), and the Root Mean Squared Error of Approximation (RMSEA). The RMSEA < 0.1, and CMIN/DF of 1–5, indicate an acceptable fit to the data. Path analyses were conducted using the Amos 24.0.0, an IBM SPSS Statistics module. Other statistical analyses were carried out with SPSS 25.0. The significance level was set at 0.05.

Results

Of the 1021 evaluated individuals, 593 (58%) had complete data and were included in the analyses. Table 1 summarizes the characteristics of the study participants, and compares patients with 24-U Ca less than 200 mg/day and 24-U Ca more than 200 mg/day.

As results showed, the mean of age, weight, serum PTH, serum Cr, 24-U Urea, 24-U Na, and 24-U Cit were significantly different between groups (P-value < 0.05 for all analyses). Besides, bivariate analysis was used to assess the existing correlations between variables. Table 2 presents Pearson’s correlation coefficients. The 24-U Ca had a significant positive correlation with weight, 24-U Urea, 24-U Cr, 24-U Na, and 24-U Cit, and had a significant negative correlation with age and serum Cr.
Path analysis result

The path analysis model is depicted in Fig. 1, which presents hypotheses about the relationships between variables. This model had a good fit with a CMIN/DF = 187.807/44 = 4.26, RMSEA = 0.086 (CI: 0.076, 0.097). The parameter estimates derived from the path analysis model are reported in Table 3. Furthermore, the total, direct and indirect effects of variables on the main outcome (24-U Ca) are provided in Table 4.

The results showed that age and weight had an indirect effect on 24-U Ca ($\beta = -0.394$ and $\beta = 0.780$, respectively, $P < 0.05$). The negative coefficient suggested that 24-U Ca is lowered with an increase in age. Furthermore, the increasing weight will indirectly increase 24-U Ca. Regarding serum variables, 25(OH)D did not significantly affect 24-U Ca. However, it had an indirect effect on 24-U Ca with serum PTH participation ($\beta = 0.028$, $P < 0.05$). Conversely, serum PTH directly affected 24-U Ca ($\beta = -0.254$, $P = 0.023$) without an indirect effect. Serum Ca did not have any direct
or indirect effect; however, Serum P appeared to influence 24-U Ca indirectly ($\beta = 1.907, P < 0.05$). Serum Cr had a significant direct effect on 24-U Ca ($\beta = -34.065, P < 0.001$).

Regarding 24-h urine metabolites, it was shown that 24-U Na, 24-U Urea, and 24-U Cit had a significant direct effect on 24-U Ca ($\beta = 0.220, \beta = 1.817, \beta = 0.066$, respectively, $P < 0.001$). 24-U Cr has a significant direct and indirect effect on 24-U Ca with participation of 24-U Cit and 24-U Urea as mediating variables ($\beta = 27.616$ for direct effect and $\beta = 30.256$ for indirect effect, $P < 0.05$).

According to standardized coefficients Table 4, 24-U Cit had the most effect on 24-U Ca, following 24-U Cr, 24-U Urea, 24-U Na, serum Cr, and weight. Serum Ca had the lowest effect on 24-U Ca.

### Discussion

According to the studies, several risk factors could contribute to calcium kidney stones. Among them, hypercalciuria is one of the most common urine metabolic abnormalities [5]. The excretion of calcium in the urine might be affected by serum variables, demographics, and other urinary metabolites. Our study revealed that 24-U Ca excretion decreases with increasing age, serum PTH and serum Cr. Conversely, 24-U Ca increases with weight gain, increasing 24-U Na, 24-U Cit, 24-U Urea, and 24-U Cr.

Among demographic factors, we included age and weight in our analysis. Previously, two large-sample studies found no association between age and hypercalciuria [6, 7]. However, Taylor et al. reported an inverse association between age and 24-U Ca; every five-year increase in age was associated with a 6 mg (95% CI 4 to 9 mg/day) decrease in 24-U Ca [8]. Our analyses showed that age indirectly affected urine calcium through serum 25(OH)D, serum Cr and 24-U Cr.

Previous studies showed that higher weight is associated with higher urinary calcium excretion [11]. Our findings showed that body weight indirectly affects 24-U Ca through 24-U Na, 24-U Urea, and 24-U Cr. Higher sodium [12, 13] and protein intake [14, 15] in obese persons was reported in some prior studies. Regarding urinary sodium and urea excretion corresponding to the amount of sodium and protein intake, it can be said that the indirect effects of weight on urinary calcium excretion might be partially due to higher salt and protein intake in patients with higher weight. However, matching with the dietary data is required to confirm these findings.

Vitamin D causes a positive balance of calcium in the body through stimulation of intestinal calcium absorption. However, since different studies’ results are inconsistent [7, 16–21], the relationship between serum vitamin D and 24-U calcium excretion is under debate. A recent meta-analysis, including twenty-two observational studies (3510 kidney stone formers and 19,718 controls), revealed that 25(OH)D was similar in both groups. Conversely, 1,25-dihydroxy vitamin D (1,25(OH)2D) was significantly higher in stone formers than controls. More detailed studies on kidney stone formers indicated that hypercalciuric patients had significantly higher 1,25(OH)2D and 25(OH)D compared with normocalciuric stone patients and non-stone forming controls [22]. Two population-based studies, InChianti Study [23] and Swiss Survey on Salt Intake Study [6], showed that the level of serum 25(OH)D was positively associated with urinary calcium excretion in men but not in women. Other studies failed to observe any correlations between 25(OH)D and urinary calcium excretion [19–21]. The results of our previous study [24] showed that the 24-U Ca increased in

| Variables                  | 24-U Ca > 200 mg (N=222) | 24-U Ca ≤ 200 mg (N=371) | Total (N=593) | p-value* |
|----------------------------|--------------------------|---------------------------|---------------|---------|
| Age                        | 46 (11)                  | 49 (13)                   | 48 (12)       | 0.001** |
| Weight                     | 81.92 (14.2)             | 79.3 (14.0)               | 80.3 (14.1)   | 0.028*  |
| Serum phosphorus           | 3.471 (0.62)             | 3.52 (0.67)               | 3.50 (0.65)   | 0.319   |
| Serum parathormone         | 47.89 (26.32)            | 56.01 (30.12)             | 53.00 (29.02) | 0.001** |
| Serum calcium              | 9.51 (0.51)              | 9.58 (0.53)               | 9.55 (0.52)   | 0.120   |
| Serum 25-hydroxy vitamin D | 24.43 (17.81)            | 23.90 (17.40)             | 24.10 (17.54) | 0.721   |
| Serum creatinine           | 1.09 (0.19)              | 1.20 (0.38)               | 1.16 (0.33)   | <0.001*** |
| 24-h urine urea            | 28.26 (11.17)            | 23.62 (9.32)              | 25.36 (10.29) | <0.001*** |
| 24-h urine sodium          | 172.31 (61.80)           | 146.84 (56.32)            | 156.38 (59.67) | <0.001*** |
| 24-h urine citrate         | 636.09 (334.61)          | 476.89 (261.20)           | 536.49 (300.64) | <0.001*** |

All values stand for Mean (Standard deviation)

24-U Ca 24-h urine calcium

*p < 0.05, **p < 0.01, ***p < 0.001

*Independent sample t-test
vitamin D supplemented patients, which was not associated with serum 25(OH)D or PTH changes. Other results of that study suggested that the increase in 24-U Ca might be due to other factors such as dietary sodium and protein intake. In line with our previous findings, the current study results showed that serum 25(OH)D did not have a direct effect on 24-U Ca, and the indirect effect was through serum PTH. This effect is negligible; therefore, the treatment of hypovitaminosis D could be safe in kidney stone formers, in case other variables, such as sodium intake, are controlled. Further randomized clinical trials are needed in this regard.

PTH is responsible for minute-by-minute regulation of serum ionized calcium, through stimulation of renal calcium reabsorption and bone resorption. Furthermore, PTH stimulates the conversion of 25(OH)D to its active hormonal form (1,25(OH)2D) in the kidney, thereby promoting absorption of calcium in the small intestine [25, 26]. A negative correlation between serum PTH and urinary calcium excretion is shown in previous studies, which is considered as the normal process of serum calcium balance. Similarly, our current results showed that PTH had a significant inverse effect on vitamin D supplemented patients, which was not associated with serum 25(OH)D or PTH changes. Other results of that study suggested that the increase in 24-U Ca might be due to other factors such as dietary sodium and protein intake. In line with our previous findings, the current study results showed that serum 25(OH)D did not have a direct effect on 24-U Ca, and the indirect effect was through serum PTH. This effect is negligible; therefore, the treatment of hypovitaminosis D could be safe in kidney stone formers, in case other variables, such as sodium intake, are controlled. Further randomized clinical trials are needed in this regard.

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24-U Ca directly. However, excessive PTH secretion from a parathyroid adenoma (primary hyperparathyroidism) leads to excessive bone resorption and increased renal synthesis of 1,25(OH)2D, which in turn enhances intestinal absorption of calcium. Therefore, the net effect is elevated serum calcium levels, which leads to higher urine calcium. Conversely, the level of urine calcium is decreased in secondary hyperparathyroidism, due to increased calcium resorption from kidney tubules. One of the common causes of secondary hyperparathyroidism are vitamin D deficiency, therefore the level of serum PTH is decreased after vitamin D supplementation.

Previous observational findings of the association between serum Ca and excretion of Ca in 24-h urine are inconsistent. The study, which was conducted on 1293 participants (624 men and 669 women), found a positive association between serum Ca and 24-U Ca in women but not men [6], while the InChianti study (595 subjects including 302 men and 293 women) observed this association in men but not women [23]. In an interventional small-sized study by Peacock et al. [27] on 72 individuals, a positive association was observed between serum Ca and 24-U Ca in both men and women. Our results revealed that there is no significant direct or indirect association between serum Ca and 24-U Ca. Normally, serum calcium is tightly controlled in a narrow range during calcium hemostasis. Since we excluded patients with primary hyperparathyroidism or any other diseases affecting serum Ca, total serum Ca was in a narrow range in all patients, and we could not find any association between serum and urine Ca. However, serum P appears to influence 24-U Ca indirectly. It seems that the effect of serum P on urinary Ca levels may be more prominent, and further studies are needed in this field.

In line with previous studies [7, 8], our results revealed that higher excretion of Na was positively correlated with 24-U Ca. The positive effect of Na intake, measured through 24-U Na, on urinary Ca excretion has been demonstrated in previous studies [28, 29]. Because sodium and Ca are reabsorbed at common sites in the renal tubules, the direct correlation of sodium and Ca in the urine is expected [30].

The relationship between urinary citrate and 24-U Ca may be challenging to discuss. Taylor et al. [8] discussed that since acid load could increase urinary Ca and decrease urinary citrate, it would be reasonable to expect an inverse association between urinary citrate and Ca. Moreover, some studies reported that potassium citrate consumption could reduce urinary Ca [31, 32]. However, we observed a marked direct effect of 24-U Cit on urinary Ca excretion, which was reported by other studies too [7, 8]. Other factors such as dietary Ca intake may play a part in such association. Further studies are needed to elucidate the topic.

Data about the relationship between 24-U Ca and other factors included in our analysis are limited. Taylor et al. did not find any associations between urinary Cr and Ca after multivariate adjustment. However, we observed a positive effect of both serum and urinary Cr on 24-U Ca. Besides, our study revealed a direct effect of 24-U Urea on 24-U Ca. Since 24-U Urea is a substitute for dietary protein intake, our results suggested an increase in 24-U Ca with high protein intake. This finding is in line with existing studies. However, since we did not evaluate the 24-U sulfate or dietary intake, we could not evaluate the effect of animal and vegetable protein intake on 24-U Ca separately.

This study has some limitations. First, we could not assess the effect of calcium on PTH since we did not test ionized calcium. In addition, Serum 1,25-dihydroxy VitD3 was not included in the path analysis. Third, the dietary intake of our participants was not available in the study. So, the additional factors affecting 24-U calcium may not be considered in our model.

### Table 4

| Predictor variables | Unstandardized coefficient | Standardized coefficient |
|---------------------|-----------------------------|-------------------------|
|                     | Direct effect   | Indirect effect | Total effect | Direct effect | Indirect effect | Total effect |
| Age                 | –              | – 0.394         | – 0.394      | –              | – 0.056         | – 0.056      |
| Weight              | –              | 0.780           | 0.780        | –              | 0.128           | 0.128        |
| S creatinine        | – 34.065       | –              | – 34.065     | –              | 0.131           | – 0.131      |
| S calcium           | – 0.669        | –              | – 0.669      | –              | 0.004           | – 0.004      |
| S 25(OH)D           | 0.019          | 0.028           | 0.047        | 0.004          | 0.006           | 0.010        |
| S PTH               | – 0.254        | –              | – 0.254      | – 0.086        | – 0.086         | – 0.086      |
| S phosphorus        | – 1.907        | –              | 1.907        | – 0.014        | – 0.014         | – 0.014      |
| 24-U citrate        | 0.066          | –              | 0.066        | 0.229          | – 0.229         | – 0.229      |
| 24-U sodium         | 0.220          | –              | 0.220        | 0.152          | – 0.152         | – 0.152      |
| 24-U urea           | 1.817          | –              | 1.817        | 0.217          | – 0.217         | – 0.217      |
| 24-U creatinine     | 27.616         | 30.256          | 57.872       | 0.108          | 0.118           | 0.226        |

All values stand for model coefficients.

S serum, 25(OH)D 25-hydroxy vitamin D, PTH parathormone, 24-U 24-h urine
Conclusion

In conclusion, except for serum calcium in the normocalcemic range (due to excluded hypercalcemic patients), our findings showed that some of the serum, urinary, and demographic variables affect 24-h urinary calcium indirectly apart from the direct effects. In this regard, age, weight, serum 25(OH)D, and phosphorus indirectly affected on urinary calcium through mediator variables. Meanwhile, PTH, serum creatinine, 24-U sodium, urea, and citrate showed a direct effect on 24-U calcium excretion.

Author contributions Conceptualization: NB, ST, MT, FT; Acquisition of Data: ST, and MT; Methodology, analysis, NB; Writing-original draft preparation, FT, and NB; Writing-review and editing, FT, ST, MT, NB; Administrative and Supervision, AB.

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Availability of data and material Data and materials can be made available upon request.

Declarations

Conflict of interest The authors declare that they have no competing interests.

Ethics approval and consent to participate The Ethics Committee of the Urology and Nephrology Research Center at Shahid Beheshti University of Medical Sciences approved the study (ethic code, IR. SBMU. UNRC.1395.15).

Consent for publication Not Applicable.

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