Strategies and cost-effectiveness evaluation of persistent albuminuria screening among high-risk population of chronic kidney disease

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Abstract

Background: Screening for persistent albuminuria among the high-risk population is important for early detection of CKD while studies regarding screening protocol and related cost-effectiveness analysis are limited. This study explored a feasible and cost-efficient screening strategy for detecting persistent albuminuria among the high-risk population.

Methods: A cohort study including 157 clinically stable outpatients with a risk factor of CKD and whose laboratory tests revealed an albumin-creatinine-ratio (ACR) between 30 and 300 mg/g of creatinine during the previous 12 months was conducted to assess the validity of alternative screening strategies. Each participant collected three first morning urine samples in three consecutive months. These samples were labeled as DAY-1, MONTH-2 and MONTH-3. In the first month, a random spot sample in the afternoon of the first day and another morning sample on the second day were collected and labeled as Random and DAY-2. Persistent albuminuria was defined by abnormal ACR (≥30 mg/g creatinine) for DAY-1, MONTH-2 and MONTH-3. Alternative strategies were DAY-1; Random; DAY-1 + Random; DAY-1 + DAY-2; and DAY-1 + Random + DAY-2. To evaluate the economic performance of those alternative strategies, a hybrid decision tree/Markov model was developed based on the cohort study to simulate both clinical and cost-effectiveness outcomes. Sensitivity analyses were conducted to investigate assumptions of the model and to examine the model’s robustness.

Results: Altogether, 82 patients exhibited persistent albuminuria. All of the five strategies had sensitivity higher than 90%. Of these strategies, Random had the lowest specificity (46.7%), and DAY-1 + Random + DAY-2 had the highest specificity (81.3%). Estimated cost for each quality adjusted life year (QALYs) gained were ¥112,335.88 for DAY-1 + Random, ¥81,346.99 for Random and ¥10,327.99 for DAY-1 + Random + DAY-2. When compared with DAY-1 strategy, the sensitivity and specificity of which were 100.0 and 69.3%, respectively. DAY-1 + Random + DAY-2 had the highest effectiveness and incremental effectiveness of 11.87 and 0.73 QALYs. At a willingness-to-pay threshold of ¥100,000 per QALY, DAY-1 + Random + DAY-2 had the highest acceptability of 91.0%. Sensitivity analyses demonstrated the robustness of the results.

Conclusions: In order to make a quick diagnosis of persistent albuminuria among high-risk population, the strategy of combining two first morning urine samples and one randomized spot urine sample in two consecutive days is accurate, saves time, and is cost-effective.

Keywords: Persistent albuminuria, Screening strategy, Chronic kidney disease, Cost-effectiveness analysis
Background

Chronic Kidney Disease (CKD) is a worldwide public health problem associated with multiple adverse outcomes and a heavy burden on the healthcare system [1, 2]. Early diagnosis and intervention have been demonstrated to be effective to improve prognosis and reduce the burden of CKD [3, 4]. Among indicators of kidney damage, albuminuria has been proven to be an independent risk factor of end-stage renal disease, cardiovascular disease, and mortality in the CKD population. Hence, the current clinical guidelines advocate that persistent albuminuria (lasting for 3 months or longer) is an essential criterion for CKD diagnosis [2, 5–7].

However, most of the current large-scaled studies were based on single-measurement of albuminuria [8–11]. Previous studies show that the fluctuation of albuminuria is substantial. The repetition positive rate based on an initial positive measurement is only 50% in CKD 1–2 stage [12, 13]. The day-to-day variations of normo- and micro-albuminuria are ±467 and ±170%, respectively [14]. Therefore, relying on a single measurement of albuminuria might lead to over-estimation of disease prevalence, as well as unnecessary treatments. The CKD screening process should start with an accurate identification of persistent albuminuria.

Previous studies had investigated the population, testing items, and frequency of CKD screening [2, 9, 15, 16]. Cost-effectiveness analyses showed that it is more effective in screening high risk populations including diabetes, hypertension, and coronary heart disease, as compared to the general population [9, 15, 16]. Patients who had been diagnosed as persistent mild-to-moderate albuminuria or early stage CKD were recommended to re-assess ACR and eGFR once a year, but few of the previous studies explored or evaluated the screening protocol for persistent albuminuria [2].

Current clinical guidelines suggest that single-positive albuminuria results should be rechecked twice in the following two months to identify whether it is persistent [2]. However, for practical reasons, the time period of 2 months is not ideal in both clinical and research settings. A short time period of diagnosis is essential for maintaining a satisfactory response rate in both clinical and research practices. Developing a practical, time-saving, and cost-effective screening strategy with acceptable accuracy is definitely necessary.

This cohort study was conducted to establish such an initial, accurate, and time-saving screening strategy. Using the results of this study, a cost-effectiveness evaluation was completed. The purpose was to evaluate its economic performance from a societal perspective.

Methods

Participants

In the cohort study, outpatients admitted to Peking University First Hospital from January 2013 to January 2014, and who met the following inclusion criteria were included: 1) laboratory tests revealed an elevated ACR (30–300 mg/g creatinine) during the previous 12 months; 2) prior diagnosis of hypertension, diabetes, or coronary heart disease, but without pre-diagnosed kidney disease (including polycystic kidney disease, kidney tumor, transplanted kidney, primary or secondary glomerulonephropy, renal tubular interstitial disease, acute kidney injury, and CKD). Exclusion criteria included: age younger than 45 years; urinary tract infection; being hospitalized within the past 2 weeks; being pregnant or less than 6 months postpartum; congestive heart failure; receiving chemotherapy and/or radiotherapy in the past 12 months. Altogether 160 patients who met the inclusion criteria were willing to participate. This group did not meet any of the exclusion criteria.

Each participant received face-to-face training and an instruction book on sample collecting. All subjects completed questionnaires to document their sociodemographic status, personal/family medical history (e.g., hypertension, diabetes, and kidney disease) and lifestyle (e.g., smoking, exercise) with assistance from nurses familiar with the study. A history of medications used (e.g., nephrotoxic medications, angiotensin-converting enzyme inhibitors, angiotensin receptor blockers, and statins) was also obtained.

In the subsequent cost-effectiveness study, the target population was patients with high risk factors of CKD including diabetes, hypertension, and coronary heart disease. It is well recognized that this population possesses higher detectable rates of persistent albuminuria and it is also the target population for CKD screening [9, 15–17]. The starting age was 45 years which is the threshold of higher prevalence of chronic diseases including diabetes, hypertension, and coronary heart disease as well as CKD, reflecting the commonly accepted starting age of screening for persistent albuminuria [18].

Specimens and measurements

The complete process of the present study including Part I Cohort Study and Part II Cost-effectiveness Study is shown in Fig. 1. The process of sample collection and strategy design is shown in Fig. 2. Each participant was asked to collect five urine samples in three consecutive months. Three first morning urine samples on the same day in three consecutive months were collected and labeled as DAY-1, MONTH-2 and MONTH-3, respectively. Because of the day-to-day variations of albuminuria, one additional first morning urine sample on the second day in the first month was required, which was labeled as DAY-2. Due to the intraday variations caused by factors such as hydration status and physical activity, a random spot urine sample in the afternoon of the first day was required, which was labeled as Random (Fig. 2).
In the first month, these two first morning samples and one randomized spot sample were to be stored in 4 °C refrigerators until participants sent them to the hospital on the morning of the second day. All the samples were cryopreserved in the same −80 °C refrigerator. The same test item was completed within two consecutive batches in order to avoid biases caused by the test itself.

Albuminuria was measured by immunoturbidimetric methods. Urinary creatinine was measured by Jaffe’s method on a Hitachi 7170 autoanalyzer (Hitachi, Tokyo, Japan). Albumin to creatinine ratio (ACR; mg/g creatinine) was calculated. A positive result was defined as an ACR greater than 30 mg/g creatinine. The term of “albuminuria” was used to describe the presence of either microalbuminuria or macroalbuminuria.

For eGFR, overnight fasting blood samples were collected by venipuncture on interview days in the first and the third months. Serum creatinine (SCr) was measured by Jaffe’s method on a Hitachi 7170 autoanalyzer (Hitachi, Tokyo, Japan). Estimated GFR was calculated using an equation developed by an adaption of the Modification of Diet in Renal Disease (MDRD) study based on data of Chinese CKD patients [19].

\[
eGFR = 175 \times \text{SCr}^{-1.234} \times \text{age}^{-0.179} \times \text{if female, } \times 0.79
\]

In the equation, SCr is the serum creatinine concentration in mg/dL, age is in years.

Serum total cholesterol, triglyceride, and uric acid which were measured in the previous 3 months while in Peking University First Hospital were documented. Results of the monthly hemoglobin A1c review of patients with diabetes were obtained from the hospital’s medical records system.

**Screening strategy**

In the cohort study, each participant collected 5 samples labeled as DAY-1, DAY-2, Random, MONTH-2 and MONTH-3, respectively. The gold standard for a persistent albuminuria diagnosis was defined as abnormal ACR for DAY-1, MONTH-2 and MONTH-3 based on the current clinical guidelines [2]. Since single-measurement of ACR is the most common choice for research, and randomized spot urine sample is the most convenient in clinical practice, this study designated DAY-1 Positive and Random Positive as two alternative strategies. Because of the fluctuation of albuminuria, we designated DAY-1 + Random both positive, DAY-1 + DAY-2 both positive, and DAY-1 + Random + DAY-2 all positive as the other three strategies (Fig. 2).

**Statistical analysis**

In the cohort study, data were presented as mean ± SD for continuous variables and proportion for categorical variables. Socio-demographic and clinical characteristics were described. The sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV), and accuracy of each strategy were then calculated using the following formulas:

\[
\text{Sensitivity} = \frac{\text{True Positive}}{\text{True Positive} + \text{False Negative}} \times 100\%
\]

\[
\text{Specificity} = \frac{\text{True Negative}}{\text{True Negative} + \text{False Positive}} \times 100\%
\]

\[
\text{Positive Predictive Value} = \frac{\text{True Positive}}{\text{True Positive} + \text{False Positive}} \times 100\%
\]
The study next conducted a hybrid decision tree and Markov models to simulate the clinical pathway and long-term outcome. The decision tree was comprised of four arms, representing alternative screening strategies DAY-1, Random, DAY-1 + Random, and DAY-1 + Random + DAY-2, respectively. The decision tree adopted prevalence of albuminuria in the general population in China, as well as sensitivity and specificity of each screening strategy to determine how many individuals were in each state of the Markov model [11]. There are four states in Markov model to simulate the pathways of patients’ screening: 1) Negative Urine Test; 2) Screening detected CKD (Positive Urine Test); 3) Symptomatic CKD; 4) Die (Fig. 3). Patients’ negative urine test results might be a true negative to persistent albuminuria which would continue in this state, or might be a false negative which would be identified correctly in the following months or continued being misdiagnosed until the patient went to hospital because of CKD symptoms. Patients with positive urine test results would be diagnosed as CKD and would receive treatment although part of the test results were false positive. The patients who recovered with proper treatment and those who were initially misdiagnosed might revert to the Negative Urine Test state. Death may occur in any health state.

Cost-effectiveness analysis
A cost-effectiveness analysis was conducted based on the cohort study (Fig. 1). Validity assessments of the cohort study revealed that the DAY-1 + DAY-2 strategy, when compared with the DAY-1+ Random strategy, did not show any advantages in sensitivity or specificity. Therefore, the DAY-1 + DAY-2 strategy was excluded from the cost-effectiveness analysis due to the increased time costs associated with it. The gold standard (DAY-1 + MONTH-2 + MONTH-3) was eliminated from the cost-effectiveness analysis for the following reasons: 1) The study initially defined the gold standard for purposes of validity assessment of various strategies and was not concerned with its practicality [8]. Secondly, both the sensitivity and specificity of a gold standard should be 100%. On the perspective of economic evaluation, the strategy, which has no false positive or false negative detection, is incomparable to alternative strategies in cost-effectiveness analysis. Therefore, only DAY-1, Random, DAY-1 + Random, and DAY-1 + Random + DAY-2 strategies were included in the cost-effectiveness analysis.

Descriptions of demographic characteristics and validity assessment of the alternative strategies including their sensitivity and specificity were analyzed by IBM SPSS Statistic 20 (IBM Corporation, New York, USA).

Accuracy
\[
\text{Accuracy} = \frac{\text{True Positive}}{\text{True Positive + False Negative + True Negative + False Positive}} \times 100\%
\]

Negative Predictive Value
\[
\text{Negative Predictive Value} = \frac{\text{True Negative}}{\text{True Negative + False Negative}} \times 100\%
\]
although it is not an ideal setting. Mortality was exacted from the China Health and Family Planning Statistical Yearbook 2013 [18]. Treatment strategies and the following outcomes of ACEI/ARB for persistent albuminuria were approximately simulated based on literatures [22–24]. The study allowed for treatment discontinuation in the model due to noncompliance and side effects of ACEI/ARB. It was assumed that most side effects appeared in the first 3 months of treatment and 25% of patients receiving ACEI/ARB would discontinue. Additionally, the study assumed another 2% loss due to the gradual discontinuation caused by noncompliance and that patients who discontinued ACEI/ARB treatment would not restart [25]. The analysis also investigated a model with costs and benefits discounted at 5% [26]. The study assumed non-informative prior distributions for all model parameters.

One-way sensitivity analyses were conducted to investigate the impact on cost-effectiveness results from varying key parameters by plus or minus 25% [27]. We varied treatment efficacy by range of 95% confidence interval and varied the cost by plus or minus 25%. The annual discount rate was varied from 0-10%. Probability sensitivity analysis was also conducted.

The cost-effectiveness analysis and sensitivity analysis were conducted using TreeAge Pro 2011 (TreeAge Software, Inc. Massachusetts, USA).

**Results**

**Cohort study**

Altogether, 160 participants were included in the present study. There were 2 females and 1 male who did not complete the 2 months following because of hysterectomy under general anesthesia, myocardial infarction, and gastrointestinal bleeding, respectively. Their data and test results were then excluded. Therefore, 157 participants aged 45–90 years were included in our final analyses.

Demographic characteristics are summarized in Table 2. The minimum and maximum of baseline ACR tests were 30.3 mg/g and 295.7 mg/g creatinine. All of the patients were in microalbuminuria magnitude at baseline. The baseline ACR of patients who were diagnosed as persistent albuminuria was higher than the baseline of all participants (106.7 (IQR 50.1, 215.9) mg/g creatinine vs. 63.6 (IQR 40.8, 134.0) mg/g creatinine). The average age was 63.4 ± 9.0 years, and 53.5% were males. The percentage with hypertension and diabetes was 79 and 79.6%, respectively. The majority of patients who had hypertension and/or diabetes controlled their blood pressures and/or HbA1c levels well. The proportion of patients with coronary heart disease and stroke were 19.7 and 8%, respectively. Nearly 50% patients were using ARB/ACEI.

**Validity of screening strategies**

According to the golden standard, 82 of 157 patients had persistent albuminuria. Table 3 shows sensitivity, specificity, positive predicted value, negative predicted value, and accuracy of the five strategies. All strategies were with sensitivity higher than 90%. The sensitivity of DAY-1 strategy reached 100.0% while its specificity was 69.3%. Random strategy had the lowest specificity of 46.7% and accuracy of 72.6%. Adding another sample taken on DAY-1 to retest, adding a randomized spot specimen on the same day (DAY-1 + Random), or a first morning specimen on the consecutive day (DAY-1 + DAY-2) did not substantially improve the sensitivity or
specificity. Validity performances of DAY-1 + Random and DAY-1 + DAY-2 were quite similar to each other. Compared with other strategies, DAY-1 + Random + DAY-2 showed a relatively higher specificity, sensitivity and accuracy, which were 81.3, 93.9 and 87.9%, respectively.

Cost-effectiveness analysis for screening strategies
Results of the base case analysis of cost-effectiveness are shown in Table 3. DAY-1 + Random + DAY-2 had the highest effectiveness and costs of 11.87 QALYs and ¥18,652.73 per person. Compared with DAY-1, when 1) DAY-1 + Random strategy was applied, costs increased by ¥3867.70 and effectiveness by 0.03 QALYs; 2) When the Random strategy was applied, costs increased by ¥2028.30 and effectiveness by 0.25 QALYs; 3) When the DAY-1 + Random + DAY-2 strategy was applied, costs increased by ¥7589.32 and effectiveness by 0.73 QALYs. Model estimates of Incremental Cost-effectiveness Ratios (ICERs) were calculated as ¥112,335.88/QALYs for DAY-1 + Random; ¥8134.69/QALYs for Random and ¥10,327.99/QALYs for DAY-1 + Random + DAY-2. DAY-1 + Random and Random were absolutely dominated by DAY-1 and by DAY-1 + Random + DAY-2.

Sensitivity analysis
A threshold to judge cost-effectiveness was also drawn, which is ¥100 thousand/QALY (three times gross domestic product (GDP) per capita). One-way sensitivity analysis showed that, compared DAY-1 + Random + DAY-2 to DAY-1 + Random, DAY-1 and Random respectively, variables including false negative rate of DAY-1 + Random + DAY-2, true positive rate of DAY-1 and Table 2 Demographic Characteristics

|                                | All Participants | Persistent Albuminuria |
|--------------------------------|------------------|------------------------|
| In Total (n)                   | 157              | 82                     |
| Age; years (SD)                | 63.4 (9.0)       | 62.0 (8.9)             |
| Male; n (%)                    | 84.0 (53.5)      | 52.0 (63.4)            |
| Current Smoker; n (%)          | 39.0 (24.8)      | 24.0 (46.2)            |
| Body-mass Index (kg/m²; SD)    | 24.7 (6.6)       | 26.1 (3.3)             |
| Primary Disease; n (%)         |                  |                        |
| Hypertension                   | 124 (79.0)       | 66 (80.5)              |
| Diabetes                       | 125 (79.6)       | 66 (80.5)              |
| Others                         | 25 (15.9)        | 13 (15.9)              |
| SBP ≤ 140 mmHg                 | 92/124 (74.2)    | 65/82 (79.3)           |
| HbA1c ≤ 8%                     | 92/125 (73.6)    | 59/82 (72.0)           |
| History; n (%)                 |                  |                        |
| Coronary heart disease         | 31 (19.7)        | 17 (20.7)              |
| Stroke                         | 13 (8.3)         | 5 (6.1)                |
| Cancer                         | 19 (12.1)        | 7 (8.5)                |
| Laboratory Tests; mean(SD)     |                  |                        |
| Uric acid (μmol/L)             | 324.1 (80.6)     | 335.0 (76.9)           |
| Triglyceride (mmol/L)          | 1.6 (1.0)        | 1.6 (0.9)              |
| LDL cholesterol (mmol/L)       | 2.5 (0.8)        | 2.4 (0.9)              |
| HDL cholesterol (mmol/L)       | 1.1 (0.3)        | 1.1 (0.3)              |
| Hemoglobin A1c (%)             | 5.7 (3.3)        | 5.9 (3.3)              |
| Creatinine (μmol/L)            | 92.0 (21.12)     | 96.0 (22.3)            |
| eGFR (mL/min per 1.73m²)       | 75.7 (20.6)      | 73.4 (18.1)            |
| ACR (mg/g creatinine; median[IQR]) | 63.6 (40.8–134.0) | 106.7 (50.1–215.9)    |
| Drugs                          |                  |                        |
| ACE (%)                        | 13 (8.3)         | 7 (8.5)                |
| ARB (%)                        | 75 (47.8)        | 41 (50.0)              |
| CCB (%)                        | 69 (72.0)        | /125 HT                |
| Statin (%)                     | 87 (55.4)        | 47 (57.3)              |
| /66 HT                         |                  |                        |
+ Random, true positive rate of Random, cost and utility of CKD, impact on incremental cost and effectiveness the most.

At a willingness-to-pay threshold of ¥100,000 per QALY, the probability of each strategy being cost effective was 91% for DAY-1 + Random + DAY-2, 8% for Random, 1% for DAY-1 and 0% for DAY-1 + Random. The DAY-1 + Random + DAY-2 strategy showed absolute superiority to other strategies.

**Discussion**

Our study focused on the very beginning step of CKD screening. It revealed, in order to diagnose persistent albuminuria correctly and quickly, and to guide clinical treatment, the strategy of combining two first morning urine samples and one randomized spot urine sample on two consecutive days is accurate, time-saved and cost-effective. Multi-time sample collection of this strategy adjusted for the variation of protein excretion of spot urine samples. It is much more convenient to operate and much easier to control specimen quality than 24 h urine collection, so this procedure could be generalized to both clinical and research settings. Diagnosis and treatment based on single morning or random urine sample might lead to overestimation of prevalence, as well as unnecessary treatment.

Patient compliance is essential in both clinical and research settings. The time window of 2 months suggested by current guidelines is too long to apply in large-scale studies and clinical practice, especially for patients with mild albuminuria and risk factors of CKD. While they are more likely to ignore the follow-up because of the mild to moderate magnitude of albuminuria, these patients are at an extremely high risk of CKD. The present study focused on this group, phoned 1460 candidates who met the inclusion criteria and did not meet exclusion criteria, and invited them to take part in the three time points, two months follow-up study. Only 160 of them were willing to participate. Overlong duration of follow-up was the main reason for their rejection. Many of them indicated that they would like to take part in the 2-day scenario in the first month if it was allowed. In another population-based study in West Malaysia, only 52.2% of respondents with albuminuria agreed to be retested in the following weeks [8]. This research was a sub-study of another large-scale cohort. It is reasonable to deduce that the group who were willing to simultaneously participate in two trials is characterized by better

| Table 3 Performance of screening strategies |
|--------------------------------------------|
| DAY-1 | Random | DAY-1 + Random | DAY-1 + DAY-2 | DAY-1 + Random + DAY-2 |
|--------------------------------------------|
| Validity Assessments                        |
| True Positive No.                           | 82 | 79 | 79 | 79 | 77 |
| Sensitivity %                               | 100 | 96.34 | 96.34 | 96.34 | 93.9 |
| 95%CI                                       | 95.6–100.0 | 89.7–99.2 | 89.7–99.2 | 89.7–99.2 | 86.3–98.0 |
| True Negative No.                           | 52 | 35 | 57 | 58 | 61 |
| Specificity %                               | 69.3 | 46.7 | 76.0 | 77.3 | 81.3 |
| 95%CI                                       | 57.6–79.5 | 35.1–58.6 | 64.7–85.1 | 66.2–86.2 | 70.7–89.4 |
| False Positive No.                          | 23 | 40 | 18 | 17 | 14 |
| False Positive %                            | 30.7 | 53.3 | 24.0 | 22.7 | 18.7 |
| 95%CI                                       | 20.5–42.4 | 41.5–65.0 | 14.9–35.3 | 13.8–33.8 | 10.6–29.3 |
| False Negative No.                          | 0 | 3 | 3 | 3 | 5 |
| False Negative %                            | 0 | 3.66 | 3.66 | 3.66 | 6.1 |
| 95%CI                                       | 0–4.4 | 0.8–10.3 | 0.8–10.3 | 0.8–10.3 | 2.0–13.7 |
| PPV %                                       | 78.1 | 66.4 | 81.4 | 82.3 | 84.6 |
| NPV %                                       | 100.0 | 92.1 | 95.0 | 95.1 | 92.4 |
| Accuracy %                                  | 85.4 | 72.6 | 86.6 | 87.3 | 87.9 |
| Cost-effectiveness Analysis                 |
| Cost (¥)                                    | 5167.42 | 11063.42 | 9035.12 | – | 18652.73 |
| Incremental Cost (¥)                        | 0.00 | 2028.30 | 3867.70 | – | 7589.32 |
| Effectiveness (QALYs)                       | 10.85 | 11.13 | 10.88 | – | 11.87 |
| Incremental Effectiveness (QALYs)           | 0.00 | 0.25 | 0.03 | – | 0.73 |
| ICER (¥/QALYs)                              | 0.00 | 8134.69 | 112335.88 | – | 10327.99 |

PPV Positive Predict Value, NPV Negative Predict Value, QALY Quality-Adjusted of Life Year, ICER Incremental Cost-effectiveness Ratio
compliance than the general outpatient; however, their response rate was still low. Hence, shortening the diagnosis time window and simplifying the screening process would be effective to improve patient compliance. In doing so, the feasibility of the screening strategy would be improved in both clinical practice and population-based studies.

In addition to the strategy suggested by current clinical guidelines, several alternative strategies were investigated in previous studies, including single randomized urine sample ACR, single first morning urine sample ACR, single spot urine sample ACR plus retest by a 24-h sample UAE [2, 6, 7, 28, 29]. The present cohort study assessed the performance of single randomized as well as the first morning urine sample ACR test. A high false positive rate is the major limitation of the single spot urine sample test. The specificity of DAY-1 and Random were 69.3 and 46.7%, respectively. In principle, measurement by 24-h urine sample should be the most accurate method, because it largely controls impacts of physiological factors including the circadian rhythm. However, due to the cumbersome collecting process, it is problematic to apply the 24-h urine test in large-scale populations and it is difficult to control sample quality. Convenience is the major advantage of spot urine sample. However, un-detectability of the variation of urine albumin excretion limits its reliability. With these considerations, the study designed three spot sample-retest strategies including DAY-1 + Random, DAY-1 + DAY-2 and DAY-1 + Random + DAY-2. Results of the diagnostic assessment demonstrate that retesting is beneficial for the recognition of false positives and improving specificity. In this study, DAY-1 + Random + DAY-2 showed the highest accuracy. It is reasonable to deduce that multi-timing spot sample measurements allows the detection of physiological variations in urine albumin excretion. Thus, with this strategy, more false positive cases were recognized and the specificity of the strategy was improved. Adding another spot sample to the first morning sample (DAY-1 + Random and DAY-1 + DAY-2) also increased the accuracy compared with the single test strategy (DAY-1). However, with both the additional random sample and the first morning sample, there was no significant difference in validity between these two-spot-sample strategies although the veracity of DAY-1 was definitely superior to Random.

Previous cost-effectiveness analyses on CKD screening strategies mainly focused on the population and/or test items [10, 16, 27]. Few studies investigated the scenario of persistent albuminuria diagnosis. This study’s model considered the whole pathway from screening to death and compared different strategies for CKD screening. By the approach of cost-effectiveness analysis, the study evaluated DAY-1 + Random + DAY-2 from a societal perspective and demonstrated that this is a cost-efficient and feasible strategy to generalize to large-scale screening.

The analysis results of this study showed, given similar sensitivity, superior specificity yields the highest cost-effectiveness. Evaluating the DAY-1 + Random + DAY-2 strategy, its higher validity rate led to greater performance on effectiveness, although its time and economic costs were also the highest. However, from the perspective of long-term financial expense, reduction of the false positive rate will maximize the economic value through avoiding over-treatment and waste of resources.

This study has limitations that deserve mention. First, a larger sample size would be helpful to make the validity assessment of alternative strategies more stable. One-way sensitivity analysis showed the veracity of alternative strategies was one of the most important factors which impacted the model. In conclusion, a larger sample size would improve not only the veracity of strategy evaluation, but also the robustness of the Markov model. Second, although patients follow various courses other than the states in this study, it was modeled in this way based on available data and literature. Finally, this study defined incidence of CKD as constant in the models although rates increased with age and were influenced by primary disease and complications. We may overestimate the effectiveness.

Conclusions
This study established a time-saving, accurate, and cost-effective screening protocol for persistent albuminuria. It explored the protocol (which is the combination of two consecutive morning urine samples and a random urine sample) and also evaluated its economic performance to ascertain whether it could be generalized to a large-scale population. Since the strategy is demonstrably practical and cost-effective, it could be generally applied in both clinical practice and researches.

Abbreviations
- ACR: Albumin to creatinine ratio; CKD: Chronic kidney disease; eGFR: Estimated glomerular filtration rate; ICER: Incremental cost-effectiveness ratio; IQR: Interquartile range; MDRD: Modification of diet in renal disease; QALY: Quality adjusted life year; SCr: Serum creatinine; UAE: Urinary albumin excretion

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Availability of data and materials
The present study was a part of the project “Establishment of early diagnosis pathway and model for evaluating progression of chronic kidney disease (DAY-131100004713007)”. Data from the cohort study will not be shared currently since the whole project is not finished. If there are any requests about details of the project, please contact with the corresponding author Luxia Zhang. Literature from which the data was extracted for the cost-effectiveness analysis is identified in the Reference section.

Authors’ contributions
WH and ZL formed the study concept, conducted the cohort study and cost-effectiveness evaluation, analyzed the data, interpreted the results, and drafted the manuscript. YL interpreted the results of the cost-effectiveness analysis and revised the manuscript. WF interpreted the results of the cohort study and revised the manuscript. ZL is the corresponding author and revised the manuscript for important intellectual content. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

Consent for publication
Not applicable.

Ethics and consent to participate
The ethics committee of Peking University First Hospital approved the study. Not applicable.

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