Early Chronic Kidney Disease Care Programme delays kidney function deterioration in patients with stage I–IIIa chronic kidney disease: an observational cohort study in Taiwan

Shu-Fen Niu, Chung-Kuan Wu, Nai-Chen Chuang, Ya-Bei Yang, Tzu-Hao Chang

INTRODUCTION

Chronic kidney disease (CKD) is recognised as a global health concern because its overall prevalence in the general population is >10%. CKD prevalence ranged from 11.7% to 15.1%, with stages I–V accounting for 2.8%–4.2%, 2.7%–5.3%, 6.4%–8.9%, 0.3%–0.5% and 0.1%, respectively. Therefore, most patients with CKD are in the early stages, in which they may have no symptoms or signs. Such early disease stages are not easily discovered and diagnosed by primary care physicians; therefore, most patients had not consulted a nephrologist, with fewer than 6% of the patients had consulting a nephrologist even for stage III CKD. Without appropriate response or management of early CKD, it progresses to advanced CKD and finally to end-stage renal disease (ESRD), which usually requires management with dialysis. Therefore, exponential growth in medical costs is expected. Despite treatment, patients with ESRD have poor quality of life and high mortality risk.

In CKD stages I–II, an optimal outcome can be achieved with adequate assessment, diagnosis and treatment. The US Centers for Disease Control and Prevention recommends that early CKD progression prevention should include testing for and controlling CKD risk factors.
factors as well as maintaining a healthy weight through a balanced diet and physical exercise. Moreover, early monitoring and treatment in conjunction with lifestyle adjustments can improve the revisit rate of patients with CKD and delay renal function reduction.

In Taiwan, >85 000 patients require dialysis and the related National Health Insurance (NHI) expenditure reached NT$44.9 billion in 2017. To reduce kidney function deterioration, improve the quality of life, reduce the burden on the NHI programme and achieve the goal of prioritising prevention over management, Taiwan’s Ministry of Health and Welfare launched the Early CKD Care Programme aimed at active management of stage I–IIIa CKD. However, the effectiveness of intervention in delaying kidney function deterioration warrants exploration. Therefore, this study explored the effects of an intervention-based Early CKD Care Programme in reducing kidney function deterioration in patients with stage I–IIIa CKD.

MATERIALS AND METHODS

Data source

This cohort study obtained information on patients with CKD stages I–IIIa in the institutional and clinical research database of Taipei Medical University (CRDB). This database contains the electronic health and medical records of >3 million patients from three affiliated hospitals, namely Taipei Medical University Hospital, Wan Fang Hospital and Shuang Ho Hospital.

Study design and cohort

Figure 1 illustrates the patient selection process for the study cohort. From the CRDB, we identified patients with CKD who had more than two medical return visits between 1 January 2012 and 31 August 2017. We enrolled those who met the following criteria of CKD stages I–IIIa: patients with normal renal function but who present signs of kidney damage such as proteinuria, haematuria and other conditions with estimated glomerular filtration rate (eGFR) ≥90 mL/min/1.73 m²; patients with kidney damage with eGFR 60–89.9 mL/min/1.73 m² and patients with eGFR 45–59.9 mL/min/1.73 m², respectively. We excluded patients aged <18 years and those who were pregnant. The remaining patients were divided into the case group, those who participated in the Early CKD Care Programme with P4301C, P4302C or P403603C treatment codes, and the control group, consisting of those who did not participate in the Early CKD Care Programme.

Outcome measures and comorbidity

Major comorbidities diagnosed before the index date, according to claims data, were defined as baseline comorbidities. The comorbidities were identified using the International Classification of Diseases, Ninth Revision, Clinical Modification codes for hypertension, diabetes mellitus (DM), gout, hyperlipidaemia, heart disease and cerebrovascular disease, as shown in online supplemental table S1. Other baseline demographic data included age, sex, eGFR, CKD stage and number of comorbidities. Here, the eGFR was calculated as 186×creatinine−1.154×age−0.203 (×0.742 for female), and the number of comorbidities was defined as the sum of the aforementioned comorbidities in the year prior to the enrolment date. The outcome of the study was patient progression to CKD stage IIIb during the study period.

The Early CKD Care Programme

The Bureau of NHI in Taiwan launched the Early CKD Care Programme in 2011. Patients who participated in the programme constitute this study’s case group. The programme involved (i) referral to a nephrologist and provision of medication for hypertension, diabetes and hyperlipidaemia to delay kidney function deterioration, avoid damage caused by improper medication and prevent complications; (ii) CKD case managers enrolled these patients and provided nursing education and lifestyle consultations and routinely monitored disease.
progress and conducted renal function tests, urinalysis and urine albumin–creatinine or protein–creatinine ratio evaluations. The CKD case managers informed the doctors and patients’ families regarding medical practice and care-giving. The nursing education provided during the enrolment period included the following: (i) teaching the basic structure and functions of kidneys; (ii) introducing the common symptoms of kidney conditions as well as the examination values; (iii) explaining daily care and prevention of kidney conditions; (iv) communicating the importance of routine monitoring; (v) communicating the importance of consulting a doctor before using medication; (vi) introducing kidney needle biopsy; (vii) introducing hyperlipidaemia, hypertension, DM, kidney conditions and their complications and (viii) explaining dietary instructions. Lifestyle recommendations included the following: smoking cessation; weight loss, particularly for those with BMI >25 kg/m² or men and women with a waist circumference of >90 and >80 cm, respectively; daily protein intake <1.5 gm/kg; prevention of routine or excessive alcohol consumption; adequate exercise and daily salt intake <100 mEq. Routine physical examinations were conducted at least once every 6 months for CKD stages I–IIIa and urine protein, urine creatinine, serum creatinine, low-density lipoprotein and haemoglobin A1c were tested. The control group received routine care and was not enrolled or monitored by CKD case managers.

Statistical analysis
Descriptive statistics were used to summarise the demographic data. Continuous variables are presented as mean and SD, and categorical variables are presented as the number of enrollees and percentage (%). The models were matched by age, sex, eGFR and CKD stage with 1:2 propensity score to reduce bias between the case group and the control group. Considering that the number of participants in the case group was substantially smaller than those in the control group, we chose a greedy and nearest neighbour matching for propensity score matching (PSM) algorithm. Before PSM, we used Student’s t-test to assess age and eGFR; and the χ² test or Fisher’s exact test were used for sex, CKD stage, number of comorbidities, hypertension, DM, gout, heart disease, hyperlipidaemia and cerebrovascular disease. After PSM, we evaluated the differences between matched pairs using the signed rank test for continuous data and McNemar’s test for binary data. Multivariable Cox proportional hazards models were matched to all the candidate variables, including comorbidity numbers, hypertension, DM, gout, hyperlipidaemia, heart disease and cerebrovascular disease to determine the risk factors for patients progressing to CKD stage IIIb. Subgroup analysis was used to determine the risk factors for patients progressing to CKD stage IIIb from baseline CKD stage IIIa or the stages before it. A two-sided statistical test at 5% significance was used. Analyses were performed using SAS (V.7.11; SAS Institute, Cary, North Carolina, USA).

Patient and public involvement
The study used de-identified data from the institutional and Taipei Medical University Research Database. No patients were involved in developing the research question or in determining the outcome measures. Patients were not involved in designing the study. There are no plans to disseminate the results of this study to any participants.

RESULTS
Study population characteristics
Table 1 presents the characteristics of the study population. Before PSM, a total of 159 774 patients with stage I–IIIa CKD were enrolled from the participating hospitals, including 1038 in the case group and 158 736 in the control group. All the variables were significantly different between the two groups (all p<0.001). Age was significantly higher in the case group than in the control group. By contrast, eGFR was significantly lower in the case group than in the control group. The proportion of sex, CKD stage IIIa, hypertension, DM, gout, hyperlipidaemia, heart disease, cerebrovascular disease and proportion of number of comorbidity were significantly higher in the case group than in the control group. To reduce bias, 1:2 PSM was used to match the age, sex, eGFR and CKD stage. After PSM, 3114 patients with stage I–IIIa CKD from the participating hospitals during the study period were finally enrolled in the study, including 1038 in the case group and 2076 in the control group. The proportion of hypertension, DM, gout, heart disease, hyperlipidaemia, cerebrovascular disease and proportion of number of comorbidities remained significantly higher in the case group than in the control group (all p<0.001). Distribution of eGFR among cases and controls during the follow-up period is shown in online supplemental table S2.

Association of Early CKD Care Programme and risk factors with early CKD progression
Table 2 lists the crude HRs and adjusted HRs (aHRs) of all variables for stage I–IIIa CKD that progressed to CKD stage IIIb during the study period. Compared with patients in the control group, the HR for progression to CKD stage IIIb was 0.72 (95% CI 0.61 to 0.85) for those participating in the Early CKD Care Programme. After adjustments for the variables listed in table 1, those in the control group still exhibited significant risk for progression to CKD stage IIIb (aHR 0.67; 95% CI 0.55 to 0.51). In addition, DM, heart disease or cerebrovascular disease in patients with stage I–IIIa CKD were significant risk factors for progression to CKD stage IIIb. The Kaplan-Meier curves for the cumulative incidence of progression to CKD stage IIIb was significantly higher in patients with stage I–IIIa CKD who did not participate in the Early CKD Care Programme (control group) than the curves in those who participated in the programme (case group) during the follow-up period (log-rank test, p=0.0025; figure 2).
The median follow-up duration was 3.0 (95% CI 1.0 to 4.7) years. Deterioration to CKD stage IIIb within 1, 3 and 5 years was respectively noted 374, 563 and 644 patients in the control group and 140, 217 and 234 patients in the case group.

**Association of Early CKD Care Programme and risk factors between CKD stage I–II and CKD stage IIIa with early CKD progression**

Of the 3114 patients with stage I–IIIa CKD in this study, 1382 patients with CKD stages I–II and the remaining 1732 patients were in stage IIIa. Table 3 lists the crude HRs and aHRs of all variables for the progression of CKD from stage I–IIIa to IIIb during the study period. In the CKD stages I–II subgroup, the Early CKD Care Programme, the number of comorbidities and comorbid hypertension, DM, gout, heart disease, hyperlipidaemia and cerebrovascular disease had no significant influence on the progression of CKD from stage I–IIa to IIIb even after adjustment for the variables. However, in the stage IIIa CKD subgroup, compared with those in the control group, the HR for progression to CKD stage IIIb in those with participated in the Early CKD Care Programme was 0.72 (95% CI 0.60 to 0.87). After adjustments for the variables listed in Table 1, participation in the programme remained a significant protective factor against progression to CKD stage IIIb (aHR 0.67; 95% CI 0.55 to 0.81). In addition, compared with patients with stage IIIa CKD but without DM, those with DM were at a greater risk of progression to CKD stage IIIb (HR 1.26; 95% CI 1.01 to 1.57 and aHR 1.69; 95% CI 1.16 to 2.47). Compared with patients without heart disease with CKD stage IIIa, those with heart disease with CKD stage IIIa were at a greater risk for progression to CKD stage IIIb after adjustment for the variables (aHR 1.65; 95% CI 1.12 to 2.45).

**DISCUSSION**

In this clinical observational study, we demonstrated that patients with stage I–IIIa CKD who participated in the Early CKD Care Programme exhibited significantly delayed deterioration of renal function to CKD stage IIIb compared with non-participants, particularly those patients in stage IIIa. We also observed that DM, heart disease and cerebrovascular disease are risk factors for deterioration of renal function in patients with stage I–IIIa CKD.
Compared with the control group, the case group had a higher mean age, a lower eGFR, a higher proportion of CKD stage IIIa and more comorbidities before PSM. In the real-life clinical scenario, these disparities are reasonable. First, patients with stage I and II CKD typically have no noticeable symptom; hence, they are usually not referred to a nephrologist. Second, patients with CKD stage IIIa are more likely to manifest clinical symptoms than patients with earlier stages of the disease and, therefore, consult a nephrologist or seek medical attention. Third, patients with CKD IIIa with more comorbidities are more likely to be referred to a nephrologist than are those with fewer comorbidities. Fourth, older patients with more comorbidities are also more likely to be referred to specialists than younger patients with same comorbidities. CKD managers frequently encourage patients with clinical symptoms and those who consulted a nephrologist, have more comorbidities or are older to participate in the Early CKD Care Programme. Therefore, PSM was used to match variables such as age, sex, eGFR and CKD stage to reduce the bias of basic characteristics between the two groups during further analysis.

After PSM, we observed that the case group still showed more comorbidities such as hypertension, DM, gout, dyslipidaemia, heart disease and cerebrovascular disease, than the control group. Hypertension and CKD are closely interlinked. Uncontrolled hypertension can accelerate CKD progression; thus, blood pressure control is essential to prevent CKD progression. DM is also a major cause of CKD and a risk factor for CKD progression. Compared with those without DM, patients with DM have a 3.8-fold higher risk of developing CKD. Among patients with type 2 DM, 42.3% have kidney injury. Compared with CKD patients without DM, those with DM developed CKD earlier and experienced more severe CKD complications. Intracellular hyperglycaemic leads to...
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In addition to hypertension and DM, gout is independently associated with CKD. Patients with hyperuricaemia are particularly susceptible to gout development. Hyperuricaemia is an independent risk factor for CKD, and hyperuricaemia treatment may delay CKD progression. Chronic hyperuricaemia is associated with hypertension, metabolic syndrome, CKD and cardiovascular disease. Dyslipidaemia is a risk factor for CKD, and CKD causes alterations in the lipoprotein profile. Therefore, the dyslipidaemia–CKD relationship is reciprocal. Hypertension, DM and dyslipidaemia are major causes of cardiovascular and cerebrovascular disease in patients with CKD. Treatment of hypertension, DM and dyslipidaemia in patients with CKD can reduce the occurrence of cardiovascular events and improve associated outcomes. Given the links between these diseases, the high proportion of heart and cerebrovascular disease observed in the case group may be expected. In theory, CKD in patients with many comorbidities should progress more rapidly from stage I–IIIa to IIIb than those with fewer comorbidities. However, in our study, despite having more comorbidities, the case group had better renal outcomes than the control group. Therefore, the Early CKD Care Programme may be assumed to be instrumental in delaying renal function deterioration.

The effect of the Early CKD Care Programme on the progression of CKD from early stages to stage IIIb was analysed. We found that participation in the programme significantly delayed the progression of CKD from stage IIIa to IIIb, however, we also observed no significant results for the progression of CKD from stages I–II to IIIb.

Table 3 Univariable and multivariable Cox regression analysis for the risk of baseline stage <3a progression to stage 3b and stage 3a progression to stage 3b among the Early Chronic Kidney Disease Care Programme and other risk factors

|                     | Baseline stage <3a n=1382 | Baseline stage=3a n=1732 |
|---------------------|---------------------------|--------------------------|
|                     | Univariable | Multivariable* | Univariable | Multivariable* |
|                     | HR (95% CI) | P value | aHR (95% CI) | P value |
| Group               |            |            |            |            |
| Control             | ref         | ref        | ref         | ref        |
| Case                | 0.75 (0.52 to 1.08) | 0.1244 | 0.75 (0.48 to 1.17) | 0.2059 |
| Comorbidity number  |            |            |            |            |
| 0                   | ref         | ref        | ref         | ref        |
| 1                   | 0.61 (0.33 to 1.12) | 0.109 | 0.50 (0.20 to 1.26) | 0.142 |
| 2                   | 0.82 (0.49 to 1.38) | 0.4528 | 0.59 (0.14 to 2.48) | 0.4753 |
| 3+                  | 0.99 (0.57 to 1.72) | 0.9617 | 0.64 (0.06 to 6.54) | 0.7025 |
| Hypertension        |            |            |            |            |
| No                  | ref         | ref        | ref         | ref        |
| Yes                 | 0.93 (0.63 to 1.36) | 0.6964 | 1.10 (0.45 to 2.66) | 0.7025 |
| DM                  |            |            |            |            |
| No                  | ref         | ref        | ref         | ref        |
| Yes                 | 1.48 (0.94 to 2.34) | 0.0932 | 1.99 (0.87 to 4.54) | 0.1032 |
| Gout                |            |            |            |            |
| No                  | ref         | ref        | ref         | ref        |
| Yes                 | 0.79 (0.43 to 1.46) | 0.4536 | 1.05 (0.45 to 2.43) | 0.9181 |
| Hyperlipidaemia     |            |            |            |            |
| No                  | ref         | ref        | ref         | ref        |
| Yes                 | 0.73 (0.48 to 1.11) | 0.1404 | 0.76 (0.34 to 1.70) | 0.5014 |
| Heart disease       |            |            |            |            |
| No                  | ref         | ref        | ref         | ref        |
| Yes                 | 1.20 (0.74 to 1.93) | 0.4599 | 1.47 (0.70 to 3.12) | 0.3093 |
| Cerebrovascular disease |            |            |            |            |
| No                  | ref         | ref        | ref         | ref        |
| Yes                 | 1.78 (0.97 to 3.28) | 0.0644 | 1.89 (0.84 to 4.26) | 0.1247 |

*The multivariable model was adjusted for all variables.

aHR, adjusted HR; DM, diabetes mellitus.

Table 3

Univariable and multivariable Cox regression analysis for the risk of baseline stage <3a progression to stage 3b and stage 3a progression to stage 3b among the Early Chronic Kidney Disease Care Programme and other risk factors.
Although the case group had low HRs for stage IIIb CKD compared with the control group, this difference was non-significant. CKD progression from stage I–II to IIIb may require some time, which could explain why few patients in the control group with stage I–II CKD progressed to stage IIIb during the follow-up period. Although some studies have developed clinical prediction models for CKD, the study groups in these investigations generally had stage III–IV CKD and ESRD was defined as the outcome. No clinical prediction model has yet been designed for stages I–II or IIIa–IIIb CKD. Further investigation employing clinical prediction models for early-to-advanced CKD are warranted. Figure 2 illustrates that the protective effect of the Early CKD Care Programme was sustained over the follow-up period, although the difference in cumulative incidence rate between the two groups gradually increased. The decrease of the slope over time may be attributed to the fact that patients who overcame the decline of their eGFR to <45 for over 1 year had good compliance or few comorbidities.

In our clinical study, patients with stage I–IIIa CKD with DM, heart disease or cerebrovascular disease exhibited considerable risk of progression to stage IIIb CKD. These results are similar to the findings of the Kidney Early Evaluation Programme (KEEP) and a population-level cohort study by Tonelli et al. Besides other conditions, DM and heart disease are also significant risk factors for the progression of CKD from stage IIIa to IIIb. Therefore, in addition to the Early CKD Care Programme, the Diabetes Shared Care Programme, which has been proven to reduce cardiovascular and cerebrovascular events and mortality risks, may be implemented.

The current study had some limitations that may affect the interpretation of the results. First, the CRDB only included data from three educational medical institutions located in New Taipei City and Taipei City in Taiwan. The greater Taipei area has adequate medical resources and, thus, may not be representative of all clinical situations in Taiwan on account of the urban–rural medical disparity. Second, our study cannot completely eliminate concerns related to selection bias because this phenomenon may be attributed to multiple reasons, including differential rates of death, and cause-specific models could feature assumptions that do not necessarily resolve competing risk issues. Third, the clinical outcome of our study was limited to the progression of early CKD; this work does not provide a comprehensive assessment of cardiovascular events and mortality. Fourth, the study did not take the potential effects of reversible kidney injury into account. Finally, the ethnicity of most of Taiwan’s population is Chinese; thus, the results may not be generalisable to populations of other ethnic backgrounds.

In conclusion, the results of this study revealed that patients with stage I–IIa CKD who participated in the Early CKD Care Programme benefit from a reduction in renal function deterioration. As such, this programme should be promoted and implemented, especially among those with stage IIIa CKD. More research is needed to understand what type of participants in the Early CKD Care Programme and which aspects of the programme yield the more effective results.

**Author affiliations**

1. Department of Nursing, Shin Kong Wu Ho-Su Memorial Hospital, Taipei, Taiwan
2. Department of Nursing, Fu Jen Catholic University, New Taipei, Taiwan
3. College of Nursing, Taipei Medical University, Taipei, Taiwan
4. Division of Nephrology, Department of Internal Medicine, Shin Kong Wu Ho-Su Memorial Hospital, Taipei, Taiwan
5. School of Medicine, Fu Jen Catholic University, New Taipei, Taiwan
6. Clinical Data Center, Office of Data Science, Taipei Medical University, Taipei, Taiwan
7. Division of Cardiovascular Surgery, Shin Kong Wu Ho-Su Memorial Hospital, Taipei, Taiwan
8. Clinical Big Data Research Center, Taipei Medical University Hospital, Taipei, Taiwan
9. Graduate Institute of Biomedical Informatics, College of Medical Science and Technology, Taipei Medical University, Taipei, Taiwan

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**ORCID iD** Chung-Kuan Wu http://orcid.org/0000-0003-4446-0167

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