Past Decline Versus Current eGFR and Subsequent Mortality Risk

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ABSTRACT

A single determination of eGFR associates with subsequent mortality risk. Prior decline in eGFR indicates loss of kidney function, but the relationship to mortality risk is uncertain. We conducted an individual–level meta-analysis of the risk of mortality associated with antecedent eGFR slope, adjusting for established risk factors, including last eGFR, among 1.2 million subjects from 12 CKD and 22 other cohorts within the CKD Prognosis Consortium. Over a 3-year antecedent period, 12% of participants in the CKD cohorts and 11% in the other cohorts had an eGFR slope < -5 ml/min per 1.73 m² per year, whereas 7% and 4% had a slope >5 ml/min per 1.73 m² per year, respectively. Compared with a slope of 0 ml/min per 1.73 m² per year, a slope of -6 ml/min per 1.73 m² per year associated with adjusted hazard ratios for all-cause mortality of 1.25 (95% confidence interval [95% CI], 1.09 to 1.44) among CKD cohorts and 1.15 (95% CI, 1.01 to 1.31) among other cohorts during a follow-up of 3.2 years. A slope of +6 ml/min per 1.73 m² per year also associated with higher all-cause mortality risk, with adjusted hazard ratios of 1.58 (95% CI, 1.29 to 1.95) among CKD cohorts and 1.43 (95% CI, 1.11 to 1.84) among other cohorts. Results were similar for cardiovascular and noncardiovascular causes of death and stronger for longer antecedent periods (3 versus <3 years). We conclude that prior decline or rise in eGFR associates with an increased risk of mortality, independent of current eGFR.


CKD affects 10%–16% of the global population.1,2 Numerous studies have reported the significant association of low eGFR at a single time point with mortality,3–9 a more frequent occurrence than ESRD, even among patients with late stages of CKD.10 Recently, there has been great interest in whether a decline in eGFR adds information to mortality risk assessment beyond eGFR at a single time point. Clinicians are often faced with a situation in which current eGFR is known along with its past trajectory. Thus, a clinically relevant question is whether past trajectory of eGFR can provide additional information beyond current eGFR.11,12

A surprising finding in previous studies was that an increase in eGFR was associated with an increased risk of mortality. Whether these observations are generalizable is uncertain, because they were on the basis of data from single centers13,14 and/or cohorts with mean baseline eGFR values of ≥50 ml/min per 1.73 m².11–16 Improvement in eGFR in a CKD population might show different...
associations with mortality than that in a general population cohort. In addition, the U-shaped association might be driven by confounding factors, such as weight loss or heart failure. Thus, a comprehensive investigation about eGFR increase and mortality risk is warranted.

The objective of the study was to use meta-analysis to address two clinically relevant questions: given patients presenting with a particular eGFR, does the prior eGFR trajectory provide additional prognostic information with respect to mortality risk beyond the present eGFR per se, and if so, what is the shape of this relationship?

RESULTS

Associations with eGFR Slope

Over a 3-year antecedent period, median (interquartile range [IQR] numbers of creatinine measurements were 7 (IQR, 7–7) in the CKD and 5 (IQR, 4–5) in the other (general population/high cardiovascular risk) cohorts; 12% of participants in the CKD and 11% of participants in the other cohorts had an eGFR slope <−5 ml/min per 1.73 m² per year, whereas 7% and 4% experienced an eGFR slope >+5 ml/min per 1.73 m² per year during the antecedent period, respectively. There were no consistent differences in the age or sex distribution between subjects with antecedent slopes of <−5, ≥−5 to ≤+5, and >+5 ml/min per 1.73 m² year; however, black subjects tended to be in the <−5 ml/min per 1.73 m² category (Supplemental Table 1, Table 1). Subjects with annual slopes <−5 ml/min per 1.73 m² per year had a higher prevalence of elevated albuminuria, were more often diabetic, and were more likely to have a history of cardiovascular disease (CVD) compared with subjects in the stable or increasing eGFR slope categories (Supplemental Table 2).

After adjustment, lower current eGFR, younger age, black race, higher total cholesterol, the presence of diabetes, and the presence of albuminuria (severely increased only in CKD cohorts; moderately increased and severely increased in the other cohorts) were associated with antecedent slope <−5 ml/min per 1.73 m² per year (Supplemental Table 3). Factors associated with an eGFR slope >+5 ml/min per 1.73 m² per year included higher current (last) eGFR, women, history of CVD, and the presence of albuminuria (severely increased only in CKD cohorts; moderately increased and severely increased in the other cohorts).

All-Cause Mortality

Among cohorts with 3-year antecedent data, 102,477 of 1,277,217 subjects died (8%) over a mean follow-up time of 3.2 years (Supplemental Table 4, Table 1). Among 12 CKD cohorts, 57,269 of 249,977 subjects died (23%), whereas among 22 other cohorts, 45,208 of 1,027,240 subjects died (4%). After antecedent intervals of 1 and 2 years, 223,979 of 1,765,589 (13%) and 158,617 of 1,597,849 (10%) subjects died, respectively (Supplemental Table 5).

| Table 1. Cohort characteristics and outcomes: characteristics of the CKD (n=12) and other (general population and high cardiovascular risk; n=22) cohorts that could provide data for a 3-year antecedent period |
|----------------|----------------|----------------|
| Variable       | Total Sample   | CKD Cohorts    | Other Cohorts |
| N              | 1,277,217      | 249,977        | 1,027,240     |
| Median no. SCr (IQR) | 5 (4–5)    | 7 (7–7)        | 5 (4–5)       |
| Slope <−5 ml/yr |                  |                |               |
| N, %           | 11               | 12             | 11            |
| Age (SD), yr   | 58 (17)          | 73 (11)        | 54 (17)       |
| Women, %       | 49               | 9              | 60            |
| Black, %       | 4                | 1              | 5             |
| Slope ≥−5 to ≤5 ml/yr |          |                |               |
| N, %           | 84              | 80             | 85            |
| Age (SD), yr   | 59 (17)          | 76 (10)        | 55 (16)       |
| Women, %       | 48              | 9              | 56            |
| Black, %       | 2               | 2              | 9             |
| Slope >+5 ml/yr |                |                |               |
| N, %           | 5               | 7              | 4             |
| Age (SD), yr   | 57 (19)          | 73 (10)        | 50 (17)       |
| Women, %       | 48              | 11             | 63            |
| Black, %       | 3               | 10             | 1             |
| Mean (SD) follow-up, yr | 3.2 (4.0) | 3 (1)          | 3 (4)         |
| ACM events     | 102,477         | 57,269         | 45,208        |
| CVM eventsb | 8231            | 340            | 7891          |

Risk of All-Cause Mortality Associated with a Decline in eGFR

Compared with subjects with no change in eGFR over the antecedent 3-year period, a slope of −6 ml/min per 1.73 m² per year was associated with hazard ratios (HRs) for all-cause mortality (ACM) of 1.25 (95% confidence interval [95% CI], 1.09 to 1.44) and 1.15 (95% CI, 1.01 to 1.31) among members of CKD and other cohorts, respectively (Figure 1, Supplemental Table 6). The risk of ACM associated with an annual eGFR decline was attenuated with shorter antecedent periods (corresponding to smaller absolute eGFR declines) (Supplemental Figure 1).

For both CKD and other cohorts, there was no statistically significant interaction of current eGFR and antecedent eGFR slope with ACM (P for interaction =0.17 and 0.19, respectively) (Figure 2). Higher current albuminuria was associated with higher ACM risk. Among albuminuria strata, the association between antecedent eGFR slope and ACM mortality overlapped only in the extremes of the eGFR slope distribution in the CKD cohorts and was roughly parallel by level of albuminuria in the other cohorts, suggesting a similar absence
of interaction between current albuminuria and antecedent eGFR decline (P for interaction = 0.67 [moderately increased albuminuria] and 0.45 [severely increased albuminuria] for CKD cohorts and P for interaction = 0.44 [moderately increased albuminuria] and 0.14 [severely increased albuminuria] for other cohorts) (Supplemental Figure 2).

The risk associated with an eGFR slope of $-6$ ml/min/1.73 m$^2$ per year over the 3-year antecedent period showed heterogeneity (Figure 3). Among CKD cohorts, metaregression suggested that differences in follow-up time (with higher HRs associated with shorter follow-up) and median age (with higher HRs associated with older age) may have accounted for some heterogeneity (Supplemental Figure 3), whereas for the other cohorts, heterogeneity was not explained by metaregression (Supplemental Figure 4).

For the CKD cohorts, absolute risk of ACM was higher with greater antecedent decline in eGFR, but current eGFR was relatively more important in determining the absolute mortality risk. Absolute risk of ACM in the other cohorts was low (Supplemental Table 7).

**Risk of ACM Associated with an Increase in eGFR**

ACM risk associations of antecedent eGFR increase were at least as strong as those for eGFR decline and mortality (Figure 5) or when the model was stratified by RMSE (Supplemental Figure 6). Although weight loss of $>2.0$ kg was associated with increased odds of eGFR rise, excluding subjects who lost $>2.0$ kg during the antecedent 3 years did not alter the U-shaped relationship between antecedent eGFR slope and ACM (Supplemental Figure 7). Excluding patients with diabetes and either adjusting for or stratifying by use of renin-angiotensin system–inhibiting medications in the antecedent period made no meaningful difference in the risk associations (Supplemental Figures 8–10).

Analyses using percentage change of eGFR rather than slope are shown in Supplemental Figure 11. Because a given absolute change in eGFR represents a higher percentage change for persons with lower current eGFR values and because the CKD cohorts had, in general, lower current eGFR, the distribution of percentage decline is shifted to the left for the CKD relative to the other cohorts, such that a greater number of persons in the CKD cohorts experienced a $\geq 30\%$ reduction in eGFR over 3 years. Nonetheless, risk associations were similar to slightly stronger when prior eGFR trajectory was assessed as a percentage change rather than slope (Supplemental Figure 11). Compared with subjects with no change in eGFR over the antecedent 3-year period, a slope of $+6$ ml/min per 1.73 m$^2$ per year was associated with HRs for ACM of 1.58 (95% CI, 1.29 to 1.95) for the CKD cohorts and 1.43 (95% CI, 1.11 to 1.84) among members of the other cohorts (Figure 1, Supplemental Table 6). The risk associated with an eGFR slope of $+6$ ml/min per 1.73 m$^2$ per year over the 3-year antecedent period showed heterogeneity across both CKD and other cohorts (Figure 4). The absolute risk of ACM was higher among members of the CKD versus the other cohorts, with current eGFR being a more important risk factor than antecedent slope (Supplemental Table 7).

The association of eGFR increase and mortality remained significant in all sensitivity analyses. Participants with positive eGFR slopes in the other cohorts had a trend toward higher risk of both cardiovascular and noncardiovascular mortality, although risk associations were attenuated (Table 2). Similarly, the increased risk of ACM associated with a positive eGFR slope in the antecedent period persisted when we included a measure, the root mean squared error (RMSE), of each individual’s variation around the eGFR slope line as a covariate in the Cox model (Supplemental Table 7).

**Figure 1.** HRs of ACM and change in eGFR. Analyses are shown for (A) CKD cohorts and (B) other (general population and high cardiovascular risk) cohorts. C depicts the adjusted HRs for the open circles in A and B. The upper panels of A and B depict metaanalyzed HRs for ACM associated with various annualized rates of eGFR. The reference group for calculation of HRs was patients with stable eGFR values (i.e., slope = 0 ml/min per 1.73 m$^2$ per year). Black circles indicate statistical significance compared with the reference (diamonds). The HR for eGFR slope was adjusted for age, sex, race (black versus nonblack), systolic BP, total cholesterol, diabetes, history of CVD, and current (last) eGFR. The lower panels of A and B illustrate histograms of the distribution of eGFR slopes among members of the CKD and other cohorts.
estimates for the resulting change of c statistics were 0.003 (95% CI, −0.000 to 0.007) and 0.002 (95% CI, 0.001 to 0.004) for the CKD and other cohorts, respectively (Supplemental Table 8).

DISCUSSION

In this analysis of >1.2 million subjects and >100,000 deaths, we found that antecedent eGFR slope over a 3-year period, whether positive or negative, exhibited a statistically significant association with ACM, cardiovascular mortality, and noncardiovascular mortality. These associations were observed even after adjustment for current eGFR (last eGFR in the antecedent period), suggesting that there is modest incremental information in the prior eGFR trajectory beyond eGFR measured at a single time point. In general, large changes in eGFR were unusual (11% for <−5 ml/min per 1.73 m² per year and 5% for >5 ml/min per 1.73 m² per year), but associated with the highest risk of mortality, whereas lesser changes were more common, but associated with smaller risks. Antecedent improvement in eGFR was associated with a mortality risk similar in magnitude to antecedent decline. This association persisted in numerous sensitivity analyses, suggesting that rapid change in creatinine-based eGFR—whether for the worse or the better—may be a poor prognostic sign. The relationship between antecedent eGFR slope and ACM was apparent across the entire spectrum of current eGFR, but at least within CKD cohorts, current eGFR had a much greater effect on absolute mortality risk than did prior trajectory.

Previous studies have shown that low eGFR measured at a single time point is an important risk factor for ACM.3,5,17–19 We sought to evaluate whether prior change in eGFR contributes independently to ACM prognosis in the clinical setting, where last eGFR value is known. Previous studies have investigated this association from a clinical trial perspective, adjusting for the first eGFR. The latter is relevant for the situation where two subjects begin a clinical trial at the same eGFR value, but one maintains a stable eGFR, whereas the other subject’s eGFR either falls or rises.20 In contrast, adjustment for last eGFR during the antecedent period, as per this analysis, replicates the clinical scenario, whereby ACM risk is compared between two patients who present with the same eGFR value, but one has had a stable eGFR, and the other has either fallen or risen to that value. Similar to previous work, in which adjustments were made for either the first or last eGFR in the antecedent period, we found a U-shaped relationship between eGFR slope and subsequent ACM risk.11,13–16 Direct, quantitative comparison between the results of these investigations and our own investigations are hampered by different indices of renal function change, different antecedent periods, and the use of rates, in some studies, rather than HRs to quantify mortality risk. However, Turin et al.12 found adjusted HRs for ACM of 1.14 and 1.68 for 4-ml/min per 1.73 m² per year declining and increasing slopes, respectively, compared with subjects with a stable eGFR value in a Canadian population-based study. These values are qualitatively similar to those for the other (general population and high risk) cohorts in this analysis. The small quantitative difference may be caused by differences in the set of adjustment factors used in the two studies. Note that data from the latter cohort were included in this analysis.

Several mechanisms may underlie the association of antecedent change in eGFR and mortality. In principle, change in creatinine-based eGFR may reflect either change in true GFR—caused by progression or remission of CKD or onset or recovery from acute kidney disease—or change in nonfiltration determinants of serum creatinine, such as muscle wasting or malnutrition. A steeper antecedent eGFR decline has been traditionally held to signify past decline in true GFR. Thus,
in this study, the true GFR for individuals with a steeper eGFR decline in the antecedent period may have continued to decline in the follow-up period below the last or current true GFR, and lower true GFR per se is expected to be associated with mortality. Alternatively, an antecedent decline in true GFR may simply reflect a more severe comorbidity profile. For example, although we adjusted for diabetes and our findings were qualitatively similar across different models of care, we did not adjust for severity of diabetes, a key determinant of both true GFR decline and mortality risk. Similarly, episodes of acute coronary syndrome or congestive heart failure may increase the risk of death and also, cause true GFR decline. However, we observed similar associations of antecedent eGFR decline with non-CVD mortality as we did with CVD mortality (albeit in the limited cohorts with these data). Previous investigations have suggested that variability in the eGFR itself may be associated with higher ACM risk. However, in this study, with individual residual eGFR variation expressed as the RMSE, we found little attenuation of the effect of decreasing (or increasing) eGFR slope on ACM.

The association between increasing eGFR and mortality is less intuitive. Rather than indicating improving true GFR, a rising eGFR may be an indicator of declining muscle mass or malnutrition, with the latter being responsible for the increase in ACM risk. However, exclusion of subjects who lost weight attenuated the risk of ACM on both ends of the eGFR slope spectrum but did not eliminate the U shape. Furthermore, a previous study reported an association between higher ACM risk and positive eGFR slope using cystatin C as a filtration marker, although cystatin C levels are less affected by muscle mass than creatinine, suggesting that a rising eGFR may reflect a rising true GFR. A rising prior true GFR may be caused by recovery from acute kidney disease associated with an acute illness, and it was the latter that was responsible for the observed increase in ACM risk rather than the rising true GFR per se. Finally, a rising true GFR could be seen with hyperfiltration in remnant nephrons, which could be associated with subsequent kidney disease progression, but it is not generally hypothesized to be associated with mortality. Because single-nephron GFR cannot be measured in humans, this mechanism remains speculative.

The strengths of this analysis include its large sample size with geographically diverse general population, high CVD risk, and CKD cohorts with current eGFR values that spanned a wide spectrum. We used an index of eGFR change that is commonly used in the clinical setting, the annualized eGFR slope, and in sensitivity analyses, the percentage change in eGFR variation expressed as the RMSE, we found little attenuation of the effect of decreasing (or increasing) eGFR slope on ACM.

Figure 3. Forest plot of HRs associated with a 6 ml/min per 1.73 m² per year decline in eGFR (an eGFR slope of −6 ml/min per 1.73 m² per year) over a 3-year antecedent period. Analyses are shown for (A) CKD cohorts and (B) other (general population and high cardiovascular risk) cohorts. Adjusted HRs within each cohort for ACM associated with an annualized decline of the eGFR of 6 ml/min per 1.73 m² per year are depicted. The reference group for calculation of HRs was patients with stable eGFR values (i.e., a slope = 0 ml/min per 1.73 m² per year). The HR for eGFR slope was adjusted for age, sex, race (black versus nonblack), systolic BP, total cholesterol, diabetes, history of CVD, and current (last) eGFR. AASK, African American Study of Kidney Disease and Hypertension; ADVANCE, The Action in Diabetes and Vascular Disease: Preterax and Aldomet Randomized Controlled Trial; AASK, African American Study of Kidney Disease and Hypertension; ADVANCE, The Action in Diabetes and Vascular Disease: Preterax and Aldomet Randomized Controlled Trial; AICH, Aich Werker’s Cohort; AKDN_Dipstick, Alberta Kidney Disease Network; ARIC, Atherosclerosis Risk in Communities Study; BC CKD, British Columbia CKD Study; CARE, The Cholesterol and Recurrent Events Trial; CCF, Cleveland Clinic CKD Registry Study; CHS, Cardiovascular Health Study; CIRCs, Circulatory Risk in Communities Study; Framingham, Framingham Heart Study; Geisinger, Geisinger CKD Study; GLOMMS 1, Grampian Laboratory Outcomes, Morbidity and Mortality Studies 1; IPHS, Ibaraki Prefectural Health Study; KP Hawaii, Kaiser Permanente Hawaii Cohort; KPNW, Kaiser Permanente Northwest; KSHS, Kangbuk Samsung Health Study; MASTERPLAN, Multifactorial Approach and Superior Treatment Efficacy in Renal Patients with the Aid of a Nurse Practitioner; MDRD, Modification of Diet in Renal Disease Study; MESA, Multi-Ethnic Study of Atherosclerosis; MRFIT, Multiple Risk Factor Intervention Trial; NephroTest, NephroTest Study; NZDCS, New Zealand Diabetes Cohort Study; Ohasama, Ohasama Study; Pima, Pima Indian Study; Rancho Bernardo, Rancho Bernardo Study; RENALD, Reduction of Endpoints in Non-Insulin Dependent Diabetes Mellitus with the Angiotensin II Antagonist Losartan; Severance, Severance Cohort Study; Sunnybrook, Sunnybrook Cohort; Taiwan MJ, Taiwan MJ Cohort Study; VA CKD, Veterans Administration CKD Study; ZODIAC, Zwolle Outpatient Diabetes Project Integrating Available Care.
cohort. Our study also has limitations. The general/high-risk cohorts enrolled generally younger persons and were less representative with respect to elderly individuals than the CKD cohorts. As in all observational studies, residual confounding is possible, and we captured only certain comorbidities. Laboratory assays were not uniform, but where possible, serum creatinine measures were calibrated to isotope dilution mass spectrometry standards. Variation in cohort study design as well as study population might introduce heterogeneity, but the relative consistency across cohorts, despite these variations, points toward the robustness of our findings. Finally, \( P \) values close to the nominal level of significance may be prone to type 1 error given the number of statistical tests involved in our analyses.

In conclusion, compared with patients with a stable eGFR, those with either an antecedent rise or fall in values were at increased risk of subsequent mortality. Prior change of eGFR over 3 years contributed additional information regarding mortality risk beyond the current eGFR itself. However, these incremental risks were clinically meaningful only for large eGFR changes, which were uncommon. Future research could focus on new filtration markers or direct GFR measurement to
help to elucidate the nature of the relationship between rising eGFR and mortality risk.

**CONCISE METHODS**

**Cohort Selection Criteria**
The Chronic Kidney Disease Progression Consortium includes cohorts in which the presence of CKD was required for cohort entry and those in which entry was determined by factors other than CKD (general population and high-CVD risk cohorts; i.e., other cohorts).3–5,8,18 This study involved 35 cohorts (13 CKD and 22 other) and included subjects ≥18 years of age who had repeated serum creatinine measurements during antecedent intervals from 1 to 3 years in duration. For the main analysis, we included 34 cohorts (12 CKD and 22 other) that could provide data for a 3-year antecedent period. This study was approved by the Institutional Review Board at the Johns Hopkins Bloomberg School of Public Health.

**Antecedent Change in eGFR**
eGFR was calculated using the Chronic Kidney Disease Epidemiology Collaboration 2009 creatinine equation.26 In cohorts without standardization of creatinine measurement to isotope dilution mass spectrometry, reported creatinine levels were multiplied by 0.95.27 For standardization of creatinine measurement to isotope dilution mass spectrometry, reported creatinine levels were multiplied by 0.95.27 For each participant, annualized eGFR slope (milliliters per minute per 1.73 m² per year) was derived from ordinary least squares10 regression using all eGFR measurements available during the antecedent period. The last eGFR measurement was taken at 3 years after the first available eGFR. All eGFR slopes were defined as antecedent slopes of < −5, −5 to +5, and > +5 ml/min per 1.73 m² per year, respectively.28

**Assessment of Baseline Covariates**
Within the antecedent period, we considered the last eGFR as the current eGFR. The last eGFR measurement was taken at 3 ± 0.5 years (i.e., between 2.5 and 3.5 years after the first available eGFR). All covariates were assessed within 1 year before the last eGFR measurement during the antecedent period. Diabetes was defined as fasting glucose ≥ 7.0 mmol/L (126 mg/dl), nonfasting glucose ≥ 11.1 mmol/L (200 mg/dl), hemoglobin A1c ≥ 6.5%, use of antglycemic drugs, or self-reported diabetes. Prior myocardial infarction, coronary revascularization, heart failure, or stroke was considered as a history of CVD. Albuminuria was categorized as none, moderately increased, or severely increased.29

**Table 2. Adjusted HRs for cardiovascular mortality and noncardiovascular mortality subsequent to an eGFR slope during a 3-year antecedent period for the other (general/high risk) cohorts (among 14 cohorts with available data)**

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Slope Change in eGFR (ml/min per 1.73 m² per yr) during the 3-yr Antecedent Period</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>−9</td>
</tr>
<tr>
<td>CV mortality Other cohorts</td>
<td>1.33 (1.17 to 1.52)</td>
</tr>
<tr>
<td>Non-CV mortality Other cohorts</td>
<td>1.29 (1.17 to 1.43)</td>
</tr>
</tbody>
</table>

Data are presented as adjusted HR (95% CI). CV, cardiovascular. The HR for eGFR slope was adjusted for age, sex, race (black versus nonblack), systolic BP, total cholesterol, diabetes, history of CVD, and current (last) eGFR.

**Assessment of Outcomes**
The primary study outcome was ACM occurring subsequent to the antecedent time period, with time at risk starting at the last measurement of eGFR (current). In Supplemental Material, we analyzed cardiovascular and noncardiovascular mortality when data were available (i.e., for 14 of the other cohorts).

**Statistical Analyses**
We performed two-stage meta-analyses, whereby each cohort was first analyzed separately and then pooled using random effect models (Supplemental Appendix 1). We imputed missing values of covariates (except eGFR) using cohort–specific mean values. Covariates that were completely missing for a particular cohort were excluded from the regression model for that cohort. We assessed heterogeneity with the I² statistic8 and random effects meta–regression analyses. Because the distributions of antecedent eGFR slope may be different among other and CKD cohorts, we a priori designed the meta-analyses to be stratified by cohort type.

Within each cohort, we estimated the adjusted HRs of ACM according to GFR slope with piecewise linear splines (knots at −10, −5, −3, −1, +1, and +3 ml/min per 1.73 m² per year). Cox models were adjusted for age, sex, race (black versus nonblack), systolic BP, total cholesterol, diabetes, history of CVD, and current eGFR. Adjustment for albuminuria was done only in secondary analyses, because albuminuria was not measured in conjunction with the last available eGFR in several cohorts. Forest plots of HR estimates at eGFR slopes of −6 and +6 ml/min per 1.73 m² per year were constructed (chosen as representative values within the rapid declining and rising eGFR slope categories, respectively). Differential effects of current eGFR and albuminuria on the relationship between change in eGFR and ACM were evaluated with interaction terms. We computed the base-case cumulative hazard of ACM at 1, 3, 5, and 10 years after baseline (Supplemental Appendix 2). Absolute risk was calculated by multiplying the meta–analyzed adjusted HRs for eGFR slopes of −6, −4, −2, 0, +2, +4, and +6 ml/min per 1.73 m² per year by the pooled base-case cumulative hazard. The improvement in discrimination with respect to ACM was assessed with the difference in c statistics for an adjusted model with and without eGFR slope as a covariate.

Because of an observed risk increase with antecedent increase in eGFR, we conducted several sensitivity analyses. First, we evaluated
the associations of antecedent eGFR slope with cardiovascular (death caused by myocardial infarction, heart failure, stroke, or sudden cardiac death) and noncardiovascular (all other etiologies) mortality. Second, we assessed the effect of individual residual eGFR variability. We used the RMSE as an indicator of the variation of an individual's eGFR values around his/her or her ordinary least squares regression line. The RMSE was included as a covariate and then, a stratifying variable (categorized as <3, 3–5, 5–10, and >10). Third, to explore whether increasing eGFR reflected weight loss, we excluded subjects with antecedent weight loss ≥2 kg over the 3-year period. Fourth, to evaluate whether the U–shaped risk relationship might represent diabetes–associated glomerular hyperfiltration, we repeated analyses excluding persons with diabetes mellitus. Fifth, analyses were repeated according to whether individuals had ever been exposed to renin–angiotensin–system blocking medications in the antecedent interval as a covariate in the Cox model and then, a stratifying variable. Analyses were performed using Stata/SE 13 software (StataCorp., College Station, TX; www.stata.com). P values <0.05 were considered statistically significant.

ACKNOWLEDGMENTS

The Chronic Kidney Disease Prognosis Consortium (CKD–PC) Data Coordinating Center is funded, in part, by a grant from the US National Kidney Foundation (funding sources include AbbVie and Amgen, Inc.) and National Institute of Diabetes and Digestive and Kidney Diseases Grant R01DK100446–01. A variety of sources have supported enrollment and data collection, including laboratory measurements, as well as follow-up in the collaborating cohorts of the CKD–PC. These funding sources include government agencies, such as the National Institutes of Health, and medical research councils as well as foundations and industry sponsors, and they are listed in Supplemental Appendix 3. Individual cohort and collaborator support is listed in Supplemental Appendix 3.

The funders had no role in the design, analysis, or interpretation of this study and did not contribute to the writing of this report or the decision to submit the article for publication.

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ULSAM: Johan Arnlöv, Uppsala University; Larsson, Uppsala University; and Anders Larsson, Uppsala University. Veterans Administration CKD Study: Csaba P. Kovessy, Memphis Veterans Affairs Medical Center and University of Tennessee Health Science Center; and Kamyar Kalantar-Zadeh, University of California Irvine Medicine Center. Zwolle Outpatient Diabetes Project Integrating Available Care: Henk J. Bilo, Isala Clinics; Nanno Kleefstra, Isala Clinics; Klaas H. Groenier, Isala Clinics; Hanneke Joosten, Isala Clinics; and I.D., Isala Clinics. CKD-PC Steering Committee: J.C., Johns Hopkins Bloomber School of Public Health; R.T.G., University Medical Center Groningen; P.E.d.J., University Medical Center Groningen; Kunioishi Iseki, University Hospital of the Ryukyus; Andrew S. Levey, Tufts Medical Center; K.M., Johns Hopkins Bloomber School of Public Health; Mark J. 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DISCLOSURES
None.

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This article contains supplemental material online at http://jasn.asnjournals.org/lookup/suppl/doi:10.1681/ASN.2015060688/-/DCSupplemental.

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