Reduced kidney function is associated with BMD, bone loss and markers of mineral homeostasis in older women: a 10-year longitudinal study

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Abstract
Summary Kidney function decreases with age; however, the long-term influence on bone density (BMD) in older women already at risk of osteoporosis is unknown. We followed kidney function and bone loss for 10 years. Declining kidney function was adversely associated with bone loss and mineral homeostasis in old women, though it attenuated with advanced aging.

Introduction Existing studies do not fully address the relationship between kidney function and bone metabolism with advanced aging in Caucasian women. This study describes the association between kidney function, BMD, bone loss and bone metabolism in older women and provides a review of the available literature for context.

Methods We studied participants from the OPRA cohort with follow-up after 5 and 10 years. Using plasma cystatin C (cysC), estimated glomerular function rate (eGFR) was evaluated at age 75 (n = 981), 80 (n = 685) and 85 (n = 365). Women were stratified into “normal” function (CKD stages 1–2), “intermediate” (stage 3a) and “poor” (stages 3b–5), and outcome measures—BMD, bone loss and markers of mineral homeostasis—were compared.

Results Femoral neck (FN) BMD positively associated with kidney function at 75 years old (β = 0.001, p = 0.028) and 80 years old (β = 0.001, p = 0.001), although with small effect size. Prevalence of osteoporosis (FN T-score ≤ −2.5) did not differ with kidney function. Measured at age 75, women with poor kidney function had higher annual percentage bone loss over 5 years compared to those with normal function (2.3%, 95% CI 1.8–2.8 versus 1.3%, 95% CI 1.1–1.5, p = 0.007), although not when measured from age 80 or 85. Additionally, markers of mineral homeostasis (PTH, phosphate, vitamin D, calcium), CRP and osteocalcin differed by kidney function.

Conclusions In old women, kidney function is associated with BMD, bone loss and altered mineral homeostasis; probably, a relationship attenuated in the very elderly.

Keywords BMD · eGFR · Elderly · Kidney disease · Mineral homeostasis · Women

Introduction
In the coming years, we face a sizeable increase in the older population [1], and consequently, the prevalence of many age-related conditions will increase. This includes osteoporosis and, also of importance for bone health, reduced kidney function and chronic kidney disease (CKD) [2, 3]. Chronic kidney disease [4] can lead to disturbed mineral homeostasis, increasing or decreasing the circulating levels of calcium, phosphorus, FGF-23, vitamin D and parathyroid hormone. This effect on mineral homeostasis may be associated with increased bone fragility [5].
It is yet to be determined, however, how reduced kidney function affects bone health in a “normal” population, not diagnosed with CKD. We have previously shown that close to 95% of women after the age of 75 have a mild to moderate reduction of kidney function (CKD stages 2–3), with an accelerated loss between ages 80 and 85 [6]. This indicates a potentially large population at risk, in the event that even mild-to-moderate reduction of kidney function has a negative impact on bone health in older women. To correctly estimate kidney function in the elderly is challenging: glomerular filtration rate can be estimated (eGFR) using either plasma creatinine or cystatin C (cysC) [7]. However, both have drawbacks; diet and muscle mass affect creatinine values, while cysC levels can be affected by high doses of corticosteroids and possibly inflammation [8].

The most accurate tool for eGFR is proposed to be cysC [9], for risk stratification overall [7] and also in elderly individuals [9, 10], possibly because they typically have a low muscle mass. Although other studies have investigated the association between kidney function and bone density (BMD), they are mostly in defined kidney disease [5]. In reviewing the literature, few studies include older women; most are cross-sectional, ethnically diverse and based on creatinine. Subsequently, there is a gap in knowledge regarding kidney function in the old and bone health. Furthermore, no longitudinal studies have evaluated whether reduced kidney function is associated with low BMD, bone markers and deranged mineral homeostasis in a normal population of older women followed over many years. Although poor kidney function is associated with mortality [6], for older women who survive into advanced age despite having poor kidney function, little is known about their bone health.

Given the hypothesized association between kidney function and skeletal health, this investigation in older women aims to be of relevance for clinicians with the following objectives: (1) to evaluate the association between kidney function and BMD; (2) to identify differences in BMD, bone loss and bone markers in women stratified by kidney function; and (3) to describe bone-related characteristics of elderly women surviving up to age 85 and whose kidney function remained stable during the 10-year follow-up. To understand the current knowledge base, we started with a non-systematic literature review. We then estimated kidney function using cysC with the CKD-EPI formula in 1044 Caucasian women from the population-based Osteoporosis Prospective Risk Assessment (OPRA) cohort. All women were 75 years at baseline with follow-up visits at ages 80 and 85.

Material and methods

Subjects

The OPRA cohort is a single-center, longitudinal population-based cohort of Caucasian women, randomly selected from city files of Malmö from 1995 to 1999 [11]. Aiming to capture the average community-dwelling older woman, no exclusion criteria were applied. A total of 1604 women received an invitation on their 75th birthday, and 1044 (65%) attended at baseline with 715 and 382 returning for the 5- and 10-year follow-up, respectively. At age 80, reasons for non-attendance included the following: illness (31%); sheltered living (5%); no reason (53%); other reasons (social reasons, walking problems or migration (11%)). At age 85: illness (56%); sheltered living (7%); no reason (23%); other reasons (14%).

The analyses in this study are based only on data from women with eGFRcysC-values, corresponding to 981 at baseline, 685 at age 80 and 365 at age 85. Reasons for missing analytical values include the following: lack of serum, could not provide a blood sample, hemolysis and failed analysis; reflecting a random loss across the cohort and not a systematic error or selection.

The study was approved by the Regional Ethical Review Board in Lund and performed in accordance with the Helsinki declaration. Participants provided written informed consent.

Kidney function

Plasma-cysC from all visits was analyzed in batch in 2015 at the Department of Clinical Chemistry, Malmö, Skåne University Hospital using Cobas auto-analyzer (CV 2.2–1.1%). Estimated glomerular function rate (mL/min/1.73 m²) was calculated using the CKD-EPI\textsubscript{cysC}-formula [7]. Comparative analyses were also performed with creatinine-based eGFR calculations using the CG- and MDRD-formulas, as previously described [6].

BMD and bone loss

Using dual-energy X-ray absorptiometry (DXA), measured with a Lunar DPX-L (GE Lunar, Madison, WI), areal BMD (g/cm²) was determined at femoral neck (FN) and total body (TB) at Skåne University Hospital Malmö. Precision of DXA was assessed by duplicate measurements on healthy individuals. Precision error was 0.009–0.010 g/cm² at FN and 0.011–0.030 g/cm² at TB. No drifts in phantom-measured results were observed [12]. Osteoporosis was defined as a T-score ≤ −2.5 at the FN and osteopenia as a T-score between −1 and −2.5. Rate of bone loss was calculated individually for every woman as annual percentage change in BMD or as absolute bone loss over 5 years (between 75 and 80, \(n = 685\); between 80 and 85, \(n = 365\)) and 10 years (between 75 and 85, \(n = 365\)).

Blood biochemistry

Non-fasting blood samples were collected between 8 a.m. and 1 p.m., centrifuged and stored at −80 °C. Plasma calcium and
alkaline phosphate (ALP) were analyzed using a Beckman
synchron LX20–4 auto-analyzer or Cobas auto-analyzer
(Roche). At age 75, serum levels of PTH were analyzed using
Elecsys PTH immunoassay (Roche Diagnostics, Mannheim,
Germany). The normal range spanned from 1.6 to 6.9 pmol/L
with intra- and inter-assay CVs of 1.6 and 5.7%, respectively.
At the 5- and 10-year follow-up, PTH was analyzed using
Cobas auto-analyzer by Roche (intra-assay CV, 7–5%). To
ensure inter-assay comparability between methods, consisten-
cy of PTH values was assessed by duplicate measurements
[13]. Plasma phosphate from all visits was analyzed in batch
according to routine methods. Serum 25(OH)D (vitamin D)
was analyzed using liquid chromatography mass spectropho-
tometry (LC-MS) (inter-assay CV 3–6%). CRP was analyzed
using Roche Diagnostics (Cobas) with a CV of 3.6–4.1%. All
assays were performed according to routine methods. Bone
turnover markers S-TRACP5b, serum bone-specific alkaline
phosphatase (s-BALP) and total osteocalcin were analyzed as
previously described [14].

Other variables

Using standardized methods, weight (kg) and height (cm)
were measured and BMI (kg/m²) calculated. Lifestyle-
related information was available from questionnaires.

Literature search

For the purpose of comparison to other published studies, we
made a non-systematic literature search in PubMed using the
following: (((“Renal Insufficiency”[All Fields] OR “Renal
Insufficiency, Chronic”[Mesh] OR “estimated glomerular fil-
tration rate”[All Fields]) AND “Bone Density”[All Fields])
AND (“female”[MeSH Terms] AND “aged”[MeSH Terms])
AND “longitudinal studies”[MeSH Terms]). This only resulted
in 10 articles; therefore, the search term “longitudinal” was
removed, resulting in 309 articles, of which 20 were deemed
relevant.

Statistics

Analyses were performed using just the available data
(Online resource 1a). Descriptive data is presented as
mean with standard deviation or median with interquartile
range as appropriate. To identify association between kid-
ey function (eGFR) and BMD, multivariate linear regres-
sion analyses with eGFR (independent) and BMD
(dependent) as continuous variables was performed, ad-
justed for weight (model 1) and weight, smoking and vi-
tamin D (model 2). In each model, we present partial
explained variation (unadjusted $R^2$) for eGFR. The anal-
yses were also performed after excluding steroid
($n = 29$ at age 75; $n = 41$ at age 80; $n = 22$ at age 85)
and bisphosphonate ($n = 33$ at age 75; $n = 58$ at age 80;
$n = 55$ at age 85) users. A previous report from the OPRA
cohort indicated that having ever used a potent estrogen
($n = 49, 5\%$) has no effect on BMD at the age of 75 [15],
and therefore, current users were not excluded from the
analyses.

Participants were staged into categories of kidney func-
tion based on their eGFR alone as follows: (1) normal-
mild reduction of kidney function, eGFR $\geq 60$ mL/min/
1.73 m² (CKD stages 1–2, “normal” kidney function), (2)
mild-moderate reduction, eGFR from 45 to 59 mL/min/
1.73 m² (CKD stage 3A, “intermediate” function) and (3)
moderate-severe reduction, eGFR < 45 mL/min/1.73 m²
(CKD stages 3B–5, “poor” function) [4].

Power analyses for bone density were performed prior
to baseline inclusion; based on the supposition of 0.13 g/
cm² SD in BMD, this study has over 80% power to detect
a 0.056 g/cm² difference between equal groups at a 5%
significance level. Differences in bone loss and character-
istics according to kidney function were calculated using
ANOVA (for normally distributed data) or Kruskal-Wallis
(for non-normally distributed data). Differences between
the women who maintained a normal versus poor kidney
function throughout follow-up were compared using inde-
pendent sample t test or Mann-Whitney U-test.

All data was analyzed with SPSS (IBM SPSS v22.0;
Armonk, NY: IBM Corp.). A $p$ value $< 0.05$ was consid-
ered nominally significant. When difference between
three groups is compared, significance is presented as $p$-
values for trend.

Results

Characteristics of the OPRA population cross-sectionally
and longitudinally for those continuing follow-up are shown
in Table 1. Cross-sectionally, eGFR declined, as
did FN and TB T-score at each assessment point and mir-
rored in those with who attended all visits. The number of
women with osteoporosis increased from 28 to 49% dur-
ing the follow-up. As described previously [6], women
who attended follow-up tended to have higher baseline
eGFR compared to non-attendees. Calcium and/or vitamin
D supplement intake did not differ between categories of
kidney function (Online resource 1b).

Association between kidney function and bone mineral
density

Kidney function (eGFR by CKD-EPIcysC) was positively as-
associated with FN BMD at both 75 and 80 years old in the fully
adjusted models ($\beta = 0.001, p = 0.028$ and $\beta = 0.001,$
But not in advanced old age (85 years). Overall, the contribution was small; however, the partial explained variation, 0.4% at 75 years old and 1.6% at age 80 years.

Excluding women on bisphosphonates did not substantially alter the results, but with exclusion of steroid users, the association at age 75 was lost. Notably, kidney function and TB BMD were inversely associated at all three ages, although abolished with weight adjustment (Table 2).

Comparative analyses using the creatinine-based formulas were essentially the same (Online resource 2).

### Table 1 Clinical data at ages 75, 80 and 85 presented as cross-sectional and as longitudinal for those who successfully underwent long-term follow-up.

| Cross-sectional | Longitudinal at age 80 (n = 685) | Longitudinal at age 85 (n = 365) |
|-----------------|---------------------------------|---------------------------------|
|                 | Age 75 (n = 981) | Age 80 (n = 685) | Age 85 (n = 365) | Age 75 | Age 80 | Age 85 | MIC (SD) | Age 75 | Age 80 | Age 85 | MIC (SD) | Age 75 | Age 80 | Age 85 | MIC (SD) |
| Age             | Mean (SD) | 75.2 (0.1) | 80.2 (0.2) | 85.2 (0.14) | 75.2 | 80.2 | 85.2 | + 5.0 (0.1) | 75.2 | 80.2 | 85.2 | + 5.0 (0.1) |
| Body mass index | Mean (SD) | 26.3 (4.2) | 26.0 (4.2) | 25.4 (4.0) | 26.3 | 26.0 | - 0.3 (2.1) | 26.1 | 25.9 | 25.4 | - 0.5 (1.8) |
| Weight          | Mean (SD) | 68 (12) | 66 (11) | 64 (11) | 68 | 66 | - 2 (5) | 67 | 66 | 64 | - 2 (5) |
| Height          | Mean (SD) | 161 (6) | 159 (6) | 158 (6) | 161 | 159 | - 1 (1) | 161 | 160 | 158 | - 1 (1) |
| Femoral neck T-score | Mean (SD) | - 1.8 (1.1) | - 2.2 (1.1) | - 2.4 (1.1) | - 1.7 | - 2.2 | - 0.5 (0.7) | - 1.7 | - 2.2 | - 2.4 | - 0.2 (0.8) |
| Total body T-score | Mean (SD) | - 1.6 (1.2) | - 1.7 (1.2) | - 1.7 (1.3) | - 1.5 | - 1.7 | - 0.2 (0.4) | - 1.5 | - 1.7 | - 1.7 | - 0.1 (0.4) |
| Osteopenia (n)  | Mean (SD) | 464 (49%) | 299 (44%) | 144 (39%) | 343 | 299 | - 44 | 174 | 160 | 144 | - 16 |
| Osteoporosis (n) | Mean (SD) | 269 (28%) | 300 (44%) | 181 (49%) | 180 | 300 | + 120 | 94 | 150 | 181 | + 31 |

**Bone loss and markers of bone metabolism stratified by kidney function**

To compare differences in bone loss and markers of bone metabolism, women were stratified into three categories (normal, intermediate, poor) based on their kidney function. To evaluate age-related associations, bone loss was compared during two different 5-year periods (between ages 75–80 and 80–85). Individually calculated annual percentage bone loss differed between the three categories (FN p = 0.001, Fig. 1); TB p = 0.014); a higher rate was
Table 2 Association between kidney function (eGFR) and BMD at femoral neck and total body at age 75, 80 and 85

| BMD-FN       | CKD-EPI cysC |
|--------------|-------------|
| ■β (95% CI)  | p-value     | Partial explained variation |
| Age 75 (n = 981) |             |                          |
| Unadjusted   | -0.2 × 10^{-3} (-0.7 × 10^{-3} to 0.3 × 10^{-3}) | 0.478 | 0.1% |
| Model 1      | 0.4 × 10^{-3} (-0.01 × 10^{-3} to 0.9 × 10^{-3}) | 0.056 | 0.3% |
| Model 2      | 0.5 × 10^{-3} (0.06 × 10^{-3} to 1.0 × 10^{-3})  | 0.028 | 0.4% |
| Age 80 (n = 685) |             |                          |
| Unadjusted   | 0.3 × 10^{-3} (-0.3 × 10^{-3} to 1.0 × 10^{-3})  | 0.323 | 0.1% |
| Model 1      | 1.1 × 10^{-3} (0.5 × 10^{-3} to 1.7 × 10^{-3})   | <0.001 | 1.7% |
| Model 2      | 1.1 × 10^{-3} (0.5 × 10^{-3} to 1.8 × 10^{-3})   | 0.001 | 1.6% |
| Age 85 (n = 366) |             |                          |
| Unadjusted   | -0.2 × 10^{-3} (-1.2 × 10^{-3} to 0.9 × 10^{-3}) | 0.778 | 0.0% |
| Model 1      | 0.6 × 10^{-3} (-0.4 × 10^{-3} to 1.6 × 10^{-3})  | 0.219 | 0.4% |
| Model 2      | 0.6 × 10^{-3} (-0.4 × 10^{-3} to 1.7 × 10^{-3})  | 0.238 | 0.3% |
| BMD-TB       | CKD-EPI cysC |
| ■β (95% CI)  | p-value     | Partial explained variation |
| Age 75 (n = 981) |             |                          |
| Unadjusted   | -0.4 × 10^{-3} (-0.8 × 10^{-3} to -0.3 × 10^{-3}) | 0.034 | 0.5% |
| Model 1      | 0.03 × 10^{-3} (-0.3 × 10^{-3} to 0.3 × 10^{-3}) | 0.830 | 0.0% |
| Model 2      | 0.04 × 10^{-4} (-0.3 × 10^{-3} to 0.3 × 10^{-3}) | 0.796 | 0.0% |
| Age 80 (n = 685) |             |                          |
| Unadjusted   | -0.6 × 10^{-3} (-1.1 × 10^{-3} to -0.1 × 10^{-3}) | 0.019 | 0.8% |
| Model 1      | 0.1 × 10^{-3} (-0.3 × 10^{-3} to 0.6 × 10^{-3})  | 0.559 | 0.1% |
| Model 2      | 0.1 × 10^{-3} (-0.3 × 10^{-3} to 0.6 × 10^{-3})  | 0.610 | 0.0% |
| Age 85 (n = 366) |             |                          |
| Unadjusted   | -1.0 × 10^{-3} (-1.8 × 10^{-3} to -0.2 × 10^{-3}) | 0.012 | 1.7% |
| Model 1      | -0.2 × 10^{-3} (-0.8 × 10^{-3} to 0.5 × 10^{-3}) | 0.609 | 0.1% |
| Model 2      | -0.3 × 10^{-3} (-1.0 × 10^{-3} to 0.4 × 10^{-3}) | 0.410 | 0.2% |

Model 1 adjusted for: weight
Model 2 adjusted for: weight, smoking, Vitamin-D
P-values calculated using linear regression analyses
§Significant when bisphosphonate users excluded (n = 32 at age 75, n = 47 at age 80, n = 44 at age 85)
§§Non-significant when bisphosphonate users excluded
¤ Significant when steroid users excluded (n = 29 at age 75, n = 33 at age 80, n = 17 at age 85)
¤¤ Non-significant when steroid users excluded

Apparent in women with intermediate (stage 3A) and poor (stages 3B–5) kidney function in the first time period, at both measurement sites. In the second period, bone loss did not differ at the FN (p = 0.264). However, a difference between the three categories was observed at TB (p = 0.044), and the lowest bone loss was seen in women with intermediate kidney function. During the first 5-year time period, analyzing absolute bone loss (g/cm²/5 years), the results were similar (data not shown).

Excluding bisphosphonate or steroid users at baseline did not significantly change the bone loss results between ages 75 and 80. However, at age 80, with exclusion of current bisphosphonate users, FN bone loss (between 80 and 85), differed between the groups (p for trend 0.048).

Table 3 shows characteristics of women stratified by kidney function at each age (75, 80 and 85). Although women with reduced kidney function (i.e. intermediate and poor) tended to have higher TB BMD (p < 0.05), there was no
Markers of mineral homeostasis also differed by kidney function; at all ages, PTH was higher as kidney function decreased ($p < 0.001$), although mean values were well within normal range. However, phosphate and ALP only differed at age 75 ($p < 0.001$), although women with poor function developed higher calcium levels. Furthermore, at all ages, women with reduced kidney function had higher values for weight, the inflammation marker CRP and the bone formation marker osteocalcin ($p < 0.001$ for all). Other bone markers, BALP and TRACP5, did not differ with kidney function.

**Characteristics of women whose CKD stage remained stable over 10 years**

To understand the skeletal implications of kidney function over time, we compared women who had, and maintained, poor kidney function (stages 3b–5, $n = 26$) throughout the 10-year observation period to those with normal function throughout (stages 1–2, $n = 52$). Women who survived despite poor function had higher annual percentage bone loss at the FN over the 10 years, although variability precluded statistical significance ($p = 0.097$) (Fig. 2). Using absolute bone loss values, the results were largely similar.

Continuous poor kidney function was associated with a higher TB BMD ($p < 0.05$ all time points). FN BMD was higher, and correspondingly, prevalence of osteoporosis lower, although only significant at baseline ($p = 0.039$ and $p = 0.017$, respectively; Table 4). In addition, the women with continuous poor function had higher PTH (increasing even above normal range at age 85) and were 10 kg heavier ($p < 0.05$ and $p < 0.001$, respectively, for all time points). Differences in calcium, phosphate, CRP, BALP and osteocalcin were also observed.

**Review of the literature**

To put our results into context and demonstrate the current knowledge base for kidney function and BMD or bone loss in elderly women, the findings from the literature review are summarized in Online resource 3 (in all 20 studies). A coherent picture of kidney function and BMD/mineral homeostasis is hampered by heterogeneity in study populations (wide age spans, ethnicity and number of participants). Only ten studies comprised more than 200 participants, and only four represent Caucasian women. Seven studies have been performed in patients with defined CKD, with a mean age considerably below 75 years. Furthermore, only three studies were longitudinal, and none of these had > 5 years of follow-up. Most studies used the creatinine-based MDRD or CG-formulas to estimate GFR, making comparisons difficult [16–35].

**Discussion**

Kidney function decreases with age [36], and although chronic kidney disease is associated with deranged mineral homeostasis [5] potentially affecting bone mass, this is to our knowledge the first study extending over a decade focusing on very elderly women, a group already at high risk of osteoporosis. In this study, we followed kidney function and bone loss for 10 years in the large population-based cohort and show that declining kidney function is adversely associated with bone
mass in older women, although the association appears to attenuate with advanced aging.

In trying to elucidate the implications of kidney function on BMD in elderly women, we began by performing a non-systematic literature search. Although lack of longitudinal studies and differences in study populations make comparison of results difficult, this search indicated an association between kidney function and BMD [17–21, 23, 28, 30], though whether mediated through differences in sex, age and weight is unclear [18, 19]. The homogeneity of the
OPRA cohort, followed for 10 years, allowed us to address this gap in knowledge and make reliable conclusions about the association between kidney function and BMD in this elderly age group.

In this cohort, kidney function explains only a small proportion of the variation in FN BMD. Since others report stronger associations [20, 21, 23], we initially assumed the discrepancy could be explained by the use of creatinine-based eGFR formulas; yet, our results using either creatinine or cysC were largely similar. It is also possible that the contribution from kidney function to BMD is manifested at a much earlier age, or that in women aged 75–85, BMD is already so low that a further reduction regardless of cause will be small. The lack of association between kidney function and BMD in advanced age (85 years) or bone loss suggests that the association between kidney function and bone diminishes with age, reflecting an influence of health bias or a reduction in power with follow-up.

In postmenopausal women, higher hip and spine BMD [37] and reduced risk of hip fractures are commonly associated with higher BMI [38]. Accordingly, and highlighting the importance of weight for older women, kidney function was inversely associated with TB BMD before adjustments for weight. Indeed, reduced kidney function was associated with higher weight, perhaps explaining why these women tended to have a lower prevalence of osteoporosis.

To further understand the clinical implications of kidney function on bone loss in older women, we used the CKD staging of kidney function. In agreement with the Rancho Bernardo Study [20], reduced kidney function (stages 3a and 3b–5) was associated with higher hip bone loss, though not in advanced age (80–85 years). The clinical relevance of this is uncertain and might be small in comparison with the contribution from other risk factors in this age group, since cross-sectionally, FN BMD did not differ between CKD stages. Why rate of bone loss was higher in the first (75–80 years) compared to the second (80–85 years) time period is unclear, but we assume that bone loss will not be linear over long time spans, with a steady state reached between periods of loss. The lack of association between kidney function and BMD at age 85 and subsequent bone loss indicates that at a certain age, identifying risk factors with pronounced contribution becomes more difficult.

In the OPRA cohort, PTH and osteocalcin were incrementally higher the lower the kidney function—of clinical relevance not just for bone health, since increased PTH has been associated with cardiovascular disease [39] and mortality independent of kidney function [40]. However, mean PTH was well within reference range at all time points, and although we hypothesized that even slightly higher levels of PTH would be associated with higher bone resorption markers, it is not certain that this is the case in early CKD, and it seems not to be the case in the OPRA. In addition, and demonstrating that even a moderate reduction in kidney function affects bone metabolism in elderly women, reduced kidney function was associated with increased markers of bone metabolism. Lower calcium levels are associated with reduced kidney function, although it is unlikely that levels will be affected until eGFR falls below 30 mL/min/1.73 m² (stages 4–5) [41] and very few women in our cohort reached that level of kidney disease. Opinions diverge regarding cysC and inflammation [8]; the women with reduced kidney function (stages 3a and 3b–5) had higher levels of CRP in this cohort.

We were interested to see what characterized women whose kidney function (either poor or normal) remained
stable as they aged. A poor compared to a normal kidney function over time (from ages 75–85) was associated with higher weight and higher levels of markers of bone metabolism. Furthermore, mean PTH increased over time in women who maintained CKD stages 3b–5 (n = 26) over 10 years. Although these results are based on a very small sample, this increase (despite a relatively stable eGFR) might represent a resistance to the activity of PTH, as previously described [5].

**Limitations**

Firstly, we acknowledge that participants might be healthier than women who declined to participate [42], although the random population-based sampling with a baseline attendance rate of 67%, which is comparatively very high, to an extent, reduces this. By the same token it might attract participants with an interest in osteoporosis and fracture. This limitation also applies to follow-up as the number is reduced from either mortality, indications of declining health or for various other reasons. In this study, this is mirrored in the eGFR values differing between attendees and non-attendees although, as previously reported not significantly between ages 75 and 80 (p = 0.123) but between ages 80 and 85 (p = 0.047) with eGFR as a predictor of mortality [6]. This is an inherent problem in all longitudinal studies of the elderly, which we have addressed by presenting both cross-sectional and longitudinal data. In addition, in an attempt to control for this, we have specifically evaluated those surviving, comparing those with good and poor function. Albeit, data needs to be interpreted with caution due to smaller numbers at the final follow-up, few other studies of same-aged women have been able to provide a similar analysis.

Secondly, the definition of normal kidney function in the “normal” older person is not without reservation since CKD staging may not fully apply. Reduced kidney function is a common state in the elderly, and many individuals in stage 3a without proteinuria might never reach stages 3b–5. We lack direct measurement, and data on urine protein would have been an advantage in this cohort. It is, however, plausible that cysC estimates of kidney function are more accurate compared to the
creatinine-based equations used in most previous studies of older people.

Thirdly, due to the observational study design, we cannot draw any conclusions about causality, but the 10-year follow-up and the absence of exclusion criteria enable reliable conclusions regarding skeletal changes over time that are generalizable to Caucasian women of old age. Importantly, the results are not substantially confounded by anti-osteoporotic medications. Hence, the study addresses the gap apparent from the literature review.

In summary, kidney function is positively associated with FN BMD in elderly women, although the association attenuates as aging progresses. A reduction of kidney function is furthermore associated with bone loss and affected mineral homeostasis. We therefore conclude that reduced kidney function is harmful to bone mass in older women.

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Compliance with ethical standards

Conflicts of interest None.

Ethical approval The study was approved by the Regional Ethical Review Board in Lund and performed in accordance with the Helsinki declaration.

Informed consent Participants provided written informed consent.

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