ABSTRACT

Objective To evaluate the effects of drug interventions that may modify the progression of chronic kidney disease (CKD) in adults with CKD stages 3 and 4.

Methods Systematic review and meta-analysis.

Data sources Searching MEDLINE, EMBASE, Database of Abstracts of Reviews of Effects, Cochrane Central Register of Controlled Trials, Cochrane Database of Systematic Reviews, International Clinical Trials Registry Platform, Health Technology Assessment, Science Citation Index, Social Sciences Citation Index, Conference Proceedings Citation Index and Clinical Trials Register, from March 1999 to July 2018, we identified randomised controlled trials (RCTs) of drugs for hypertension, lipid modification, glycaemic control and sodium bicarbonate, compared with placebo, no drug or a drug from another class, in ≥40 adults with CKD stages 3 and/or 4, with at least 2 years of follow-up and reporting renal function (primary outcome), proteinuria, adverse events, maintenance dialysis, transplantation, cardiovascular events, cardiovascular mortality or all-cause mortality. Two reviewers independently screened citations and extracted data. For continuous outcomes, we used the ratio of means (ROM) at the end of the trial in random-effects meta-analyses. We assessed methodological quality with the Cochrane Risk of Bias Tool and confidence in the evidence using the Grading of Recommendations Assessment, Development and Evaluation (GRADE) framework.

Results We included 35 RCTs and over 51 000 patients. Data were limited, and heterogeneity varied. Final renal function (estimated glomerular filtration rate) was 6% higher in those taking glycaemic control drugs (ROM 1.06, 95% CI 1.02 to 1.10, I²=0%, low GRADE confidence) and 4% higher in those taking lipid-modifying drugs (ROM 1.04, 95% CI 1.00 to 1.08, I²=88%, very low GRADE confidence). For RCTs of antihypertensive drugs, there were no significant differences in renal function. Treatment with lipid-modifying drugs led to a 36% reduction in cardiovascular disease and 26% reduction in all-cause mortality.

Strengths and limitations of this study

► By focusing on patients with chronic kidney disease (CKD) stages 3 and 4, we provide a primary care perspective on the management of CKD.

► We completed an extensive and comprehensive search including common drug mis-spellings, with no restriction on language or publication type.

► Generalisability of our findings is limited by the paucity of trials, high statistical heterogeneity, high clinical heterogeneity, in terms of comorbidities and by the effect of drugs on CKD being secondary in all the studies.

► Lack of data restricted our ability to explore the baseline characteristics of studies, the effects of drugs, and the link between treatment effect and drug dose, and assess the potential for publication bias.

Conclusions Glycaemic control and lipid-modifying drugs may slow the progression of CKD, but we found no pooled evidence of benefit nor harm from antihypertensive drugs. However, given the data limitations, further research is needed to confirm these findings.

PROSPERO registration number CRD42015017501.

INTRODUCTION

Chronic kidney disease (CKD) is a long-term loss of renal function and is a global health problem, with an estimated prevalence of 11%–13%. 1 CKD is an independent risk factor for cardiovascular disease, 2 and people with mild-to-moderate CKD are at increased risk of mortality and progression to renal failure. 3 Historically, CKD was categorised by its cause, which was predominantly diabetes,
hypertension, glomerulonephritis or polycystic kidney disease. Treatment could be recommended, but often opportunities were missed for early detection of the disease or prevention of clinical complications and further deterioration. The National Kidney Foundation addressed this challenge by developing a five-stage classification system based on the glomerular filtration rate (GFR) (online supplementary material), which was subsequently modified to take into account degrees of proteinuria (online supplementary material). A UK audit in 2012 estimated that 90% of cases of CKD are at stage 3, and of these 84% are at stage 5a (estimated GFR (eGFR) 45–59 mL/min/1.73²). Patients with CKD who are managed in primary care include those in stages 1 and 2, with an emphasis on lifestyle and diet changes. Patients in stages 3 and 4 may be managed either in primary care or shared care with specialist nephrology services, with a greater use of medication, primarily for associated long-term conditions, such as diabetes and hypertension. Both published data for Oxfordshire and unpublished national data from our group show that some patients with CKD stage 4 have their renal function monitored in primary care.

Four different classes of drugs may modify CKD progression: antihypertensives, lipid-modifying drugs, glycaemic control medications in patients with diabetes, and sodium bicarbonate (M). Sodium bicarbonate addresses metabolic acidosis, a prevalent complication in moderate and late-stage CKD, and may offer a simple and low-cost treatment to slow CKD progression. The effects of drugs on CKD are secondary, as these drugs are primarily used to treat hypertension, hypercholesterolaemia or diabetes.

Systematic reviews have been compiled to summarise the effects of drug interventions on the rate of CKD progression. However, while two have reported results based on eGFR, none has reported results with reference to CKD staging. Furthermore, no systematic review has compared the effects of different drug classes.

Our objective was to carry out a systematic review and meta-analysis of randomised controlled trials (RCTs) to examine and compare the effects on the progression of CKD of the four classes of drugs, focusing on patients who can be managed in primary care or shared care with specialist nephrology services.

**METHODS**

Our systematic review protocol was registered with PROSPERO (International Prospective Register of Systematic Reviews) and reported in line with the recommendations from the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement.

**Search strategy**

The search strategy was designed by an information specialist (NR) with advice from clinicians including nephrologists. Medical Subject Headings (MeSH) terms included terms for population, study design and intervention drugs (randomised control trials, chronic kidney, renal, drug classes and specific drug names, including common misspellings). Our included drug list is shown in the online supplementary material.

Searches were conducted in Ovid MEDLINE (online supplementary material), Cochrane Central Register of Controlled Trials, Cochrane Database of Systematic Reviews, Database of Abstracts of Reviews of Effects, EMBASE, Health Technology Assessment, Science Citation Index, Social Sciences Citation Index, Conference Proceedings Citation Index, Clinical Trials Register, and the WHO International Clinical Trials Registry Platform to 6 July 2018 to identify relevant studies, regardless of their publication type. No language restriction was applied. All publications from March 1999 onwards were considered, to align with the publication of the first equation to calculate eGFR, which had led to the introduction of CKD staging. Reference lists were also examined.

**Study selection**

Two reviewers (KST, JM, NP and SF; in pairs) reviewed the title and abstract of each reference and identified potentially relevant references. Considering the full texts of these studies, two reviewers independently selected studies to be included in the review using predetermined inclusion criteria. Disagreements about study inclusion were resolved by a third reviewer.

The following were our study inclusion criteria:

1. RCTs of drugs that may modify the progression of CKD: antihypertensives, lipid-modifying drugs, glycaemic control drugs and sodium bicarbonate.
2. Control arms were either given a placebo drug, no drug intervention or a comparator drug from one of the three other classes.
3. Adult participants.
4. Patients with stage 3 and/or stage 4 CKD. Studies additionally including patients with stage 1 and/or stage 2 CKD were also included to increase the number of potential studies, but we would expect the treatment effect to be overestimated in these studies, as patients with stage 1 and 2 CKD would respond better to treatment.
5. Populations with at least 40 participants.
6. Follow-up of at least 2 years.
7. Publications reporting a quantitative summary of effect as a change from baseline, or an endpoint measure of at least one of seven prespecified outcomes.
   i. Renal function, which could be measured by GFR, eGFR, creatinine clearance (CrCl) or estimated CrCl. This was our primary outcome of interest.
   ii. Proteinuria, which could be measured by protein excretion rate (PER), protein creatinine ratio (PCR), albumin excretion rate (AER) or albumin creatinine ratio (ACR).
   iii. Adverse events.
   iv. Commencement of maintenance dialysis or kidney transplantation.
   v. Cardiovascular events.
Cardiovascular mortality.

All-cause mortality.

We excluded trials that reported only data about populations that included people with CKD stage 5 to avoid underestimating the treatment effect, as we would expect stage 5 patients to have more comorbidities, which might lead to worse outcomes.

Data extraction and quality assessment

Data were extracted independently by two reviewers (KST, JM, NP and JYV; in pairs) using a data extraction form, which was piloted on a sample of five studies.

Extracted data included study details (setting, intervention dose and frequency, follow-up period), information on study participants (age, gender, ethnicity, smoking status, blood pressure, CKD stage, existing comorbidities) and outcome data. For one study, we extracted endpoint eGFR data from an earlier point than the longest follow-up as the patient numbers at the longest follow-up were so small (<7% of the population). We contacted study authors to seek clarification about data and to request data that could be included in our study.

The methodological quality of studies was assessed independently by two reviewers (KST, JM, NP and JYV; in pairs) using the Cochrane Risk of Bias Tool. This tool considers the method of random sequence generation and allocation concealment (selection bias), blinding of participants, personnel and outcomes (performance and detection biases), the levels and balance of missing outcome data (attrition bias), and completeness of reporting (selective reporting bias).

Disagreements about extracted data or methodological assessments were resolved by a third reviewer.

Data synthesis and analysis

We expected high statistical heterogeneity, but pooled data to give an indication of the average effect. We also pooled data when there were only two or three studies, although this could be potentially misleading, in order to give an indication of potential trends had there been more study data. We carried out random-effects analyses based on the DerSimonian and Laird method, assessed statistical heterogeneity using the I² statistic and calculated approximate 95% prediction intervals to estimate the likely effects in a clinical setting using the methods of Higgins et al.

For the continuous outcomes (renal function and proteinuria), we used a ratio of means (ROM) effect measure, which allowed us to combine data from trials that reported different measures of renal function or proteinuria in a single analysis. Using the ROM effect measure overcomes limitations of the well-established standardised mean difference (SMD), by providing units that are more easily interpreted than SDs, and avoiding the problems associated with presenting results with SDs, which can produce misleading results by deflating treatment effects for studies of heterogeneous populations and inflating treatment effects in homogeneous populations. We calculated the ROM as the ratio of the final eGFR in the intervention group to the final eGFR in the comparator group, so ROM >1 favoured the intervention group for eGFR, where higher values are better than lower values. If a study of renal function reported reductions from baseline or rates of decline rather than endpoint data, we estimated the ROM of endpoints by the ratio of the reduction from baseline or rate of decline in the comparator group to that in the intervention group (the reciprocal) to ensure the direction of effect was consistent across all studies, as higher values of declining renal function are worse. ROM <1 favoured the intervention group for the proteinuria outcomes (ACR, AER, PER, PCR), where higher values are worse.

For a few studies it was necessary to make further estimates by approximating a mean using the median; estimating SD from the IQR; or estimating SE of ROM by imputation using the mean value of the known SEs of the other studies.

Meta-analysis was carried out on the log-transformed ROM and its SE. For the primary analysis, eGFR measured by the Modification of Diet in Renal Disease (MDRD) equation was selected over Chronic Kidney Disease Epidemiology Collaboration (CKD-EPI) data, as MDRD is the main equation advocated in clinical guidelines and CKD-EPI was not introduced until 2009. eGFR was also selected over CrCl. When analysing the effects of treatments on proteinuria, we pooled data for PER, PCR, ACR and AER using a ROM effect measure. For the dichotomous outcomes, we specified the relative risk (RR) as the effect measure.

We performed separate sensitivity analyses to ensure that our estimates of the effect of treatment in studies of patients with CKD stages 3 and 4 only (our main focus) were not sensitive to our choice of model, assumptions, quality of the studies, estimations made or our choices of data.

The following were the prespecified sensitivity analyses:

- Fixed-effect analyses based on the inverse-variance method were carried out for the continuous outcomes and based on the Mantel-Haenszel method for the dichotomous outcomes.
- Excluding studies rated as low quality, which we defined as having a high risk of attrition bias or allocation concealment bias; studies rated as ‘unclear’ were not excluded.
- Excluding studies without intention-to-treat analysis. This was prespecified on the assumption that we would be dealing with primary studies of the progression of CKD, but as the studies were secondary or post-hoc this analysis was no longer relevant.

The following were further, standard set of sensitivity analyses:

- Excluding studies with estimates.
- Using more conservative estimates, considering those of the SE of ROM by imputing with the 75% percentile value of the known SEs rather than the mean value, reflecting greater uncertainty.
► Using alternative data, considering CKD-EPI data instead of MDRD data, for studies that measured eGFR using both equations.
► Excluding studies that measured PCR and PER in the analysis of proteinuria, as urine albumin is a more accurate measure of glomerular damage.
► Excluding each individual study, one by one, to see if the level of heterogeneity depended on a particular study.

The following were post-hoc sensitivity analyses:
► Using the Hartung-Knapp-Sidik-Jonkman variance correction to calculate 95% CIs reflecting the uncertainty in heterogeneity estimates.
► Excluding studies with non-placebo controls.

Our prespecified subgroup analyses were by age (<65 or ≥65 years), ethnicity, smoking status, intensity of intervention, high blood pressure (≥140/90 mm Hg), low blood pressure (<90/60 mm Hg), method of eGFR measurement, degree of proteinuria in the renal function analysis, drug type within each drug class and by CKD stages 3 and 4 separately. Heterogeneity was assessed using the I² statistic. Publication bias was explored using funnel plot analysis with contour plots.

We used the Grading of Recommendations Assessment, Development and Evaluation (GRADE) framework to report the overall quality of evidence for the primary outcome (final eGFR). The certainty in the evidence for each outcome was graded as high, moderate, low or very low.

When describing the included trials, we refer to ‘studies’ to encompass single trials and publications that report more than one trial. When reporting analyses of the effects of the drugs, we refer to populations as some studies reported data for CKD subgroups or dose-dependent subgroups. All results of meta-analysis are displayed in forest plots and/or tabulated. Results are divided into those for studies of patients with CKD stages 3 and 4 only and studies of patients with CKD stages 3 and 4 mixed with patients with stages 1 and 2.

All analyses were carried out using STATA v.14.2, except the risk of bias figures which were created in RevMan v.5.3.

**Patient involvement**

Members of a patient and public involvement group were part of the Stakeholder Group and Steering Committee of the National Institute for Health Research (NIHR) Programme Grant that inspired this study. No patients or patient representatives were involved in setting the outcome measures, nor were they directly involved in developing plans for the design or implementation of the study. Three members of this group commented on our manuscript and we thank them for their help in the Acknowledgement section. A 1-day dissemination event is planned to report the results of all the studies funded by the NIHR Programme Grant, including this study. Members of the patient and public involvement group will be invited to this event.

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**RESULTS**

**Search results**

Through database registry and hand searches, we identified 1333 records (figure 1), of which 35 studies (51 155 patients) satisfied our eligibility criteria.

**Study characteristics**

Most of the 35 studies reported data about a single trial, while 3 studies reported data about a pooled analysis of more than one trial. Of the 35 studies, 1 was from China, 1 from Tasmania, 3 from Japan, 6 from USA/Canada, 9 from Europe, either in single or multiple countries, and the other 15 studies were multinational (online supplementary material). Nineteen studies provided data for populations with CKD stages 3 and/or 4 only. These studies had between 108 and 3094 patients, and the mean or median follow-up was between 24 and 66 months.

One study had a 2×2 factorial design involving an antihypertensive drug and a lipid-modifying drug, and this study is included in our analysis of both antihypertensive drugs and lipid-modifying drugs. Another study had a 2×2 factorial design involving two antihypertensive drugs and a lipid-modifying drug, and this study is only included in our analysis of lipid-modifying drugs because all patients received antihypertensives. The other 33 studies were 12 studies of antihypertensives, 13 of lipid-modifying drugs, 6 of glycaemic control agents (and 2 of sodium bicarbonate online supplementary material). Four studies had a single comorbidity, and 4 studies had more than one
comorbidity. Comorbidities were cardiovascular in 19 studies9-19 and type II diabetes in 9 studies19-22, including all 6 studies of glycaemic control drugs; and a renal disorder or damage (idiopathic chronic glomerulonephritis, advanced chronic renal insufficiency without diabetes, advanced autosomal dominant polycystic kidney disease, renal impairment and metabolic acidosis) in 7 studies.13,44,49-51,58-70. CKD, chronic kidney disease; MDRD, Modification of Diet in Renal Disease; ROM, ratio of means.

### Risk of bias

We considered the methodological quality of all 35 eligible studies (online supplementary material), based on the design of the trial(s) from which data were obtained. Eight studies19-22 and for the other 16 studies, baseline data were only provided for either a different subset of the trial population or the whole trial population (online supplementary material).

### Renal function

Most of the available data were on eGFR. The pooled ROM of eGFR for antihypertensive drugs versus comparators at the end of the studies is shown in Figure 2. There was no significant effect in studies of populations with CKD stages 3 and 4, and heterogeneity between studies was substantial at 94% (yielding a prediction interval of 0.79–1.34), and this was not attributable to a single study. Adding data from the studies of patients that included those with CKD stages 1 and/or 2 increased the benefit to the intervention group but not significantly. All except one study (Bianchi et al26) had placebo controls. All the sensitivity analyses produced similar results of no significant treatment effect and heterogeneity remained substantial (online supplementary material).

The only subgroup analysis that was possible, of those prespecified, was by drug type within the antihypertensive drug class. Among studies of CKD stages 3 and 4 only, there were no significant treatment effects within the different drug types (online supplementary material), except in the case of calcium channel blockers, where the endpoint eGFR was 14% higher in the intervention group (ROM 1.14, 95% CI 1.01 to 1.27), but this was based on a single study.53,54

In studies of patients with CKD stages 3 and 4, there was a marginally higher benefit to those taking the lipid-modifying drugs (Figure 3), as the endpoint eGFR was 4% higher (ROM 1.04, 1.00 to 1.08, statistically significant, I²=88%; STATA’s metaninf command produced output that indicated that the lower confidence bound was 1.00 to 4 decimal places) with a prediction interval of 0.91–1.18. This result was not robust as the results of sensitivity analysis were variable (online supplementary material). The high degree of heterogeneity was attributed to the Study of Heart and Renal Protection102-104 of the CKD stage 4 subgroup, as removing this study reduced I² from 88.3% to 45.3%. The intervention for this study was a higher dose of statin than in the other studies, and another lipid-modifying drug, ezetimibe, was also used. Adding data from the studies of patients including those with CKD stages 1 and/or 2 increased the benefit to the intervention group but not significantly, and there was substantial heterogeneity between studies (I²=98%). Three trials had non-placebo controls (ASUCA,48 MEGA49-51 and GREACE76,77).
There was a significant benefit to those taking glycaemic controlling drugs (figure 4) as the mean eGFR at the end of the study was 6% higher in the intervention group (ROM 1.06, 1.02 to 1.10). Heterogeneity between studies was low (I²=0%), yielding a prediction interval of 1.01–1.11. Sensitivity analyses showed that this result was robust (online supplementary material). Adding the CKD stage 1–4 studies, there was no significant difference between the intervention and comparator groups, and the heterogeneity between studies was high (I²=85%). In all studies the comparator groups received placebo drugs. We assessed the outlier41–43 as having high attrition bias, as of the 1887 who were randomised only 487 completed the 104-day study period. After excluding this study, there was a significant benefit to the intervention group, which was similar to that for the studies of patients with CKD stages 3 and 4.

A study44 of patients with CKD stage 3 reported a 22% higher endpoint eGFR from taking sodium bicarbonate (ROM 1.22, 1.10 to 1.36). Another study45 of patients with CKD stage 4 reported that taking sodium bicarbonate produced no significant effect on renal function in terms of CrCl (ROM 0.94, 0.85 to 1.05). Pooling these data, there was no significant treatment effect based on renal function (online supplementary material). In both studies the comparator groups received routine care.

Figure 4 summarises the effects of treatment on renal function by drug class for the four drug classes.

The GRADE confidence in these estimates was very low for our analysis of antihypertensive drugs, lipid-regulating drugs and sodium bicarbonate, and low for our analysis of glycaemic control drugs.

Proteinuria

Data pooling for proteinuria was possible only for antihypertensive drugs, lipid-regulating drugs and sodium bicarbonate, and low for our analysis of glycaemic control drugs.

There was a significant benefit to those taking lipid-modifying drugs versus comparator (boxes) and pooled estimates across studies (diamonds) calculated by the random-effects DerSimonian and Laird method, split by CKD stage. ASCOT-LLAa, atorvastatin and amlodipine versus placebo and amlodipine; ASCOT-LLb, atorvastatin and atenolol versus placebo and atenolol; GISSI-HFa, CKD stage 3; GISSI-HFb, CKD stage 4; GREACEa, atorvastatin versus usual care for the group with metabolic syndrome; GREACEb, atorvastatin versus usual care for the group without metabolic syndrome; SHARPa, CKD stages 1–3; SHARpb, CKD stage 4. CKD, chronic kidney disease; ROM, ratio of means; SHARP, Study of Heart and Renal Protection.

Proteinuria

Data pooling for proteinuria was possible only for antihypertensive drugs, lipid-regulating drugs and sodium bicarbonate, and low for our analysis of glycaemic control drugs.

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5–3545 mg/24 hours for the comparator group). The PREVEND IT lipid study of patients with CKD stages 1–3 reported that treatment slightly increased urinary albumin excretion by quoting medians and IQRs (21.8 and 11.6–41.9 mg/24 hours for the intervention group, and 20.3 and 12.5–40.5 mg/24 hours for the control group).

One glycaemic control study presented proteinuria as an outcome. The CANVAS programme reported that treatment significantly reduced albuminuria compared with the placebo group. ACR was expressed on the geometric mean scale, as 13% lower in the intervention group compared with the controls (95% CI 1% to 24%).

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Adverse events
There were no significant differences in the risk of adverse events in the studies of antihypertensives, lipid-modifying drugs and glycaemic control agents (online supplementary material). Neither of the bicarbonate studies reported adverse events.

Maintenance dialysis and kidney transplantation
Two studies of antihypertensive drugs reported the number of patients who deteriorated and required maintenance dialysis or kidney transplantation. The risk in the intervention group was 31% lower than in the control group in the HALT PKD study of patients with stages 3 and 4 CKD (RR 0.69, 95% CI 0.50 to 0.96). In the RENAAL study, which was based on patients with CKD stages 2–4, treatment with antihypertensives led to a 23% reduction in risk of deterioration to kidney failure (RR 0.77, 95% CI 0.64 to 0.93). There were no significant differences in this outcome in the studies of lipid-modifying drugs or glycaemic control drugs (online supplementary material).

Cardiovascular events
There were no significant differences between the intervention and control groups in the risk of cardiovascular events in the studies of antihypertensives, whereas treatment with lipid-modifying drugs led to a reduction in risk of 36% (RR 0.64, 95% CI 0.52 to 0.80, I²=30%) (figure 6).

All the sensitivity analyses yielded similar results and heterogeneity generally remained low (online supplementary material). The reduction in risk was slightly higher when data were added from the studies of patients that included those with CKD stages 1 and/or 2, but this reduction in risk was not significant (online supplementary material).

The TECOS glycaemic control study reported the number of patients who experienced cardiovascular events in patients with CKD stage 3. They reported no significant differences between the intervention and control groups (online supplementary material).

Mortality
In both our analyses of the studies of antihypertensive drugs and glycaemic control drugs, we found no significant differences, between the intervention and comparator groups, in the risk of cardiovascular mortality and all-cause mortality (online supplementary material).

Two studies reported the effect of lipid-modifying drugs on the risk of cardiovascular mortality in patients with CKD stages 3 and 4. In the FIELD study, the risk was 47% lower in the intervention group (RR 0.53, 95% CI 0.44 to 0.65).
D (figure all-cause mortality by 26% (RR 0.74, 95% CI 0.56 to 0.98) treatment with lipid-modifying drugs reduced the risk of there was no significant difference.

the findings of Sandhu, et al, who also consistent with those of Lewis et al, who reported the preliminary results of a review that considered the effect on kidney function of lowering low-density lipoprotein cholesterol with statins. Their preliminary findings suggested that for patients taking statins there was a highly significant reduction in the annual rate of CKD progression compared with control groups (based on the annual rate of change in eGFR). Both of these studies considered general populations with CKD. Our finding that lipid-modifying drugs reduce the risk of cardiovascular events and all-cause mortality also concurs with other reviews of patients with CKD.106–110

Figure 7 Relative risk (RR) of all-cause mortality during trials of lipid-modifying drugs versus comparator (boxes) and pooled estimates across trials (diamonds) calculated by the random-effects DerSimonian and Laird method, split by CKD stage. GREACEa, atorvastatin versus usual care for the group with metabolic syndrome; GREACEb, atorvastatin versus usual care for the group without metabolic syndrome. CKD, chronic kidney disease.

0.30 to 0.92), and in the AFCAPS/TEXCAPS study there was no significant difference.

In the studies of patients with CKD stages 3 and 4 only, treatment with lipid-modifying drugs reduced the risk of all-cause mortality by 26% (RR 0.74, 95% CI 0.56 to 0.98) (figure 7 and online supplementary material).

DISCUSSION
Main findings
We found limited data on the effects of antihypertensive, lipid-modifying and glycaemic control drugs and sodium bicarbonate on the progression of renal function in patients with CKD stages 3 and 4. There was heterogeneity of patient populations in the included studies. Pooling data provided no pooled evidence of benefit or impairment of renal function in these patients attributable to antihypertensive drugs. There was some suggestion of benefit to renal function from taking glycaemic control drugs and lipid-modifying drugs, and a clear reduction in cardiovascular disease and all-cause mortality by taking lipid-modifying drugs. There was insufficient evidence to draw any conclusions about sodium bicarbonate. There were only sufficient data to study the effects of antihypertensive drugs on proteinuria and we found no significant treatment effect.

Comparison with other reviews
Our review is comparable with two reviews of drug interventions in patients with CKD, as they report changes in eGFR as outcome measures. Our findings on the protective effect of lipid-modifying drugs are consistent with the findings of Sandhu et al, who concluded that statins produce a small reduction in the progression of CKD in patients with cardiovascular disease. Our results are also consistent with those of Lewis et al, who reported the preliminary results of a review that considered the effect on kidney function of lowering low-density lipoprotein cholesterol with statins. Their preliminary findings suggested that for patients taking statins there was a highly significant reduction in the annual rate of CKD progression compared with control groups (based on the annual rate of change in eGFR). Both of these studies considered general populations with CKD. Our finding that lipid-modifying drugs reduce the risk of cardiovascular events and all-cause mortality also concurs with other reviews of patients with CKD.106–110

Strengths and limitations
Our study is a first attempt to summarise the literature on drug treatment for patients with CKD types 3 and 4. We have identified, evaluated and compared the effects of different drug interventions on reducing the rate of CKD progression in patients with CKD stages 3 and 4. By focusing on this patient group, we took a primary care perspective on the management of CKD. Our search was extensive and comprehensive, with no restrictions on language or publication type. By using the ROM as the effect measure for continuous outcomes, we were able to pool data on different measures in a single analysis and overcome the limitations of the more established SMD. This provided an overall indication of the treatment effect, as did pooling data in cases where there were few studies or a high degree of heterogeneity. The generalisability of our findings is limited by the paucity of studies, the high degree of statistical heterogeneity, and clinical heterogeneity in terms of comorbidities and treatment interventions. We concluded that clinical heterogeneity was inevitable in studies in these patients, as most were undergoing treatment with background medications. Paucity of data restricted our ability to explore the baseline characteristics of studies and the effects of lipid-modifying drugs, glycaemic control drugs and bicarbonates, particularly on proteinuria and our other secondary outcomes. We were also restricted in our ability to investigate the link between treatment effect and drug dose and assess publication bias fully. There was a wide variation of study characteristics (in terms of population size and follow-up period) and baseline characteristics (in terms of male/female split, smoking status, ethnicity and hypertensive status). The generalisability of the findings is also limited by the effect of drugs on CKD being secondary in all the studies. For example, antihypertensives target hypertension and glycaemic control drugs target diabetes. Our ability to assess the quality of the included studies was restricted, because for nearly half the studies we could not assess allocation concealment bias, and for around a quarter of the studies we could not ascertain the risk of attrition bias or whether the participants, personnel and outcomes assessors had been blinded. Further bias may have arisen as most of the reports in our included studies

Table 8

| Study | N | Follow-up (months) | RR (95% CI) |
|-------|---|--------------------|-------------|
| CKD stages 3/4 only | | | |
| ASCUSA | 334 | 24 | 0.99 (0.96, 1.02) |
| FIELD | 510 | 60 | 0.80 (0.54, 1.19) |
| VA-HIT | 369 | 62 | 0.91 (0.52, 1.62) |
| MEGA | 2976 | 64 | 0.48 (0.27, 0.87) |
| Subtotal (inception + 3.7%, p = 3.43%) | | | |
| with estimated predictive interval | | | |
| Mixed with CKD stages 3/4 | | | |
| ASCOT-LA 10350 | 0 | | 0.87 (0.71, 1.08) |
| LODE | 123 | 29 | 1.12 (0.29, 4.29) |
| GREACEa | 885 | 36 | 0.44 (0.21, 0.92) |
| GREACEb | 712 | 36 | 0.41 (0.20, 0.86) |
| PREVEND 1054a | 44 | | 1.49 (0.42, 5.29) |
| Subtotal (inception + 47.1%, p = 0.109) | | | |
| with estimated predictive interval | | | |
| Overall (inception + 24.6%, p = 0.225) | | | |
| with estimated predictive interval | | | |

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were subgroup or post-hoc analyses of trials. The balance achieved by randomising the whole trial population may not apply to subgroups of patients. However, we were unable to check this as few of these studies reported the necessary baseline characteristics, split by intervention group. Few studies reported the use of maintenance dialysis and kidney transplantation, which may be considered to be more appropriate survival outcomes for patients with CKD stages 3 and 4, compared with mortality. Limitations may also arise from the possible impact on patient management decisions following the introduction of CKD staging in 2002. Before that, patients were treated on the basis of the perceived cause of CKD rather than the CKD itself. A number of the included studies had recruitment periods that began before 2002, so their populations would be a mixture of patients managed under the old and new guidance. However, in spite of all these limitations, the results of our analyses are consistent with those of other studies.

**Implications for clinical practice and future research**

In populations with CKD stages 3 and 4, lipid-modifying drugs and glycaemic control drugs may improve renal function, and lipid-modifying drugs are associated with a reduction in cardiovascular events. Although antihypertensive drugs and sodium bicarbonate are presumed to reduce CKD progression, we did not find a significant treatment effect. However, the paucity of data and high heterogeneity do not allow strong recommendations for clinical practice. Therefore, future research is needed into the effects of drugs on the progression in CKD, in particular in studies that focus on CKD progression as a primary outcome.

Our finding of limited data is important given that CKD is a common progressive disease. The drugs we considered are commonly used, but studies have concentrated on cardiovascular outcomes and not CKD outcomes.

**CONCLUSIONS**

This review suggests that glycaemic control and lipid regulation may improve renal function in patients with moderate CKD. There was no pooled evidence of either benefit or harm from antihypertensive drugs. However, the data were limited and heterogeneity was high for many of the analyses. Therefore, our findings are tentative.

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**REFERENCES**

1. Hill NR, Fatoba ST, Oke JL, et al. Global Prevalence of Chronic Kidney Disease - A Systematic Review and Meta-Analysis. *PLoS One* 2016;11:e0158765.
2. Go AS, Chertow GM, Fan D, et al. Chronic kidney disease and the risks of death, cardiovascular events, and hospitalization. *N Engl J Med* 2004;351:1296–305.
3. National Institute for Health and Care Excellence. *Chronic kidney disease: early identification and management of chronic kidney disease in adults in primary and secondary care (CG182)*. London: National Clinical Guideline Centre, 2014.
4. Levey AS, Eckardt K-U, Tsai KT, et al. Definition and classification of chronic kidney disease: a position statement from kidney disease: improving global outcomes (KDIGO). *Kidney Int* 2005;67:2089–100.
5. Levin A, Stevens PE. Summary of KDIGO 2012 CKD guideline: behind the scenes, need for guidance, and a framework for moving forward. *Kidney Int* 2014;85:49–61.
de Luisignan S, Tomson C, Harris K, et al. UK Prevalence of Chronic Kidney Disease for the Adult Population Is 6.76% Based on Two Creatinine Readings. *Nephron Clin Pract* 2012;120:107–c07.

Oke J, Shire B, McFadden E, et al. Trends in serum creatinine testing in Oxford, UK, 1993–2015: a population-based cohort study. *BMJ Open* 2015;5:e009459.

Peterson JC, Adler S, Burkart JM, et al. Blood pressure control, proteinuria, and the progression of renal disease, the modification of diet in renal disease study. *Ann Intern Med* 1995;123:754–62.

Sandelu S, Wiebe N, Fried LF, et al. Statins for improving renal outcomes: a meta-analysis. *J Am Soc Nephrol* 2006;17:2006–16.

Wang PH, Lau J, Chalmers TC. Meta-Analysis of effects of intensive blood-glucose control on late complications of type I diabetes. *Lancet* 1993;341:1306–9.

Psizabeth S, Khanna A, Simon J, et al. Amelioration of metabolic acidosis in patients with low GFR reduced kidney endothelin production and kidney injury, and better preserved GFR. *Kidney Int* 2010;77:617–23.

Mahajan A, Simon J, Sheather SJ, et al. Daily oral sodium bicarbonate preserves glomerular filtration rate by slowing its decline in early hypertensive nephropathy. *Kidney Int* 2010;78:303–9.

de Brito-Ashurst I, Varugamun M, Raferty MJ, et al. Bicarbonate supplementation slows progression of CKD and improves nutritional status. *J Am Soc Nephrol* 2009;20:2075–84.

Lv J, Ehteshami P, Schmid CH, et al. The PRISMA statement for systematic reviews and meta-analyses of controlled clinical trials with binary outcome. *BMJ* 2008;337:a2341.

Friedrich JO, Adhikari NKJ, Beyene J. Ratio of means for analyzing continuous outcomes: as an alternative to mean differences for analyzing continuous outcome variables in meta-analysis: a simulation study. *BMJ Med Res Methods* 2009;2:27.

Friedrich JO, Adhikari NKJ, Beyene J. Ratio of means as an alternative to mean differences for analyzing continuous outcomes in meta-analysis performed as well as mean difference methods. *J Clin Epidemiol* 2011;64:56–64.

Guyatt GH,Thorlund K, Oxman AD, et al. Grade guidelines: 13. preparing summary of findings tables and evidence profiles-continuous outcomes. *J Clin Epidemiol* 2013;66:73–93.

Levey AS, Stevens LA, Schmid CH, et al. The modification of diet in renal disease study. *J Am Soc Nephrol* 2001;20:3875–89.

Peters JL, Sutton AJ, Jones DR, et al. Contour-enhanced meta-analysis funnel plots help distinguish publication bias from other causes of asymmetry. *J Clin Epidemiol* 2008;61:991–6.

SzaboCorp. Stata statistical software: release 14 [program]. College Station, TX: StataCorp LP, 2015.

Cochrane Community. Review Manager (RevMan) [Computer program]. Version 5.3. Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration, 2014.

Hawkins NM, Jhund PS, Pozzi A, et al. Severity of renal impairment in patients with heart failure and atrial fibrillation: implications for non-vitamin K antagonist oral anticoagulant dose adjustment. *Eur J Heart Fail* 2016;18:1162–71.

Hillege HL, Nitash B, Pfeffer MA, et al. Renal function as a predictor of outcome in a broad spectrum of patients with heart failure. *Circulation* 2006;113:671–8.

Pfeffer MA, Swedberg K, Granger CB, et al. Effects of candesartan on mortality and morbidity in patients with chronic heart failure: the CHARM-Overall programme. *Lancet* 2003;362:759–66.

Leiter LA, Cefalu WT, de Bruin TW, et al. Long-term maintenance of efficacy of dapagliflozin in patients with type 2 diabetes mellitus and cardiovascular disease. *Diabetes Obes Metab* 2016;18:766–74.

Cefalu WT, Leiter LA, de Bruin TWA, et al. Dapagliflozin’s effects on glycemia and cardiovascular risk factors in high-risk patients with type 2 diabetes: a 24-week, multicenter, randomized, double-blind, placebo-controlled study with a 28-Week extension. *Diabetes Care* 2015;38:1218–27.

Leiter LA, Cefalu WT, de Bruin TW, et al. Dapagliflozin added to usual care in individuals with type 2 diabetes mellitus with persistently elevated blood glucose in existing cardiovascular disease: a 24-week, multicenter, randomized, double-blind, placebo-controlled study with a 28-week extension. *J Am Geriatr Soc* 2014;62:1252–62.

Hou FF, Zhang X, Zhang GH, et al. Efficacy and safety of benazepril for advanced chronic renal insufficiency. *N Engl J Med* 2006;354:191–40.

Fassett RG, Robertson IK, Ball MJ, et al. Effect of atorvastatin on kidney function in chronic kidney disease: a randomised double-blind placebo-controlled trial. *Atherosclerosis* 2010;213:218–24.

Sakata Y, Shibata N, Takahashi J, et al. Clinical impacts of additive use of olmesartan in hypertensive patients with chronic heart failure: the supplemental benefit of an angiotensin receptor blocker in hypertensive patients with stable heart failure using olmesartan (support) trial. *Eur J Heart Fail* 2015;17:915–23.

Sakata Y, Nochikoa K, Miura M, et al. Supplemental benefit of an angiotensin-receptor antagonist in hypertensive patients with stable heart failure using olmesartan (SUPPORT) trial: rationale and design. *J Cardiol* 2013;62:31–6.

Kimura G, Kasahara M, Ueshima K, et al. Effects of atorvastatin on renal function in patients with dyslipidemia and chronic kidney disease: assessment of clinical usefulness in CKD patients with atorvastatin (ASUCA) trial. *Clin Exp Nephrol* 2017;21:417–24.

Management of Elevated Cholesterol in the Primary Prevention Group of Adult Japanese (MENDA) Study Group. Design and baseline characteristics of a study of primary prevention of coronary events with pravastatin among Japanese with mildly elevated cholesterol levels. *Circ* 2004;68:860–7.

Nakamura H, Mizuno K, Ohashi Y, et al. Pravastatin and cardiovascular risk in moderate chronic kidney disease. *Circulation* 2009;2006:512–7.

Nakamura H, Arakawa K, Itakura H, et al. Primary prevention of cardiovascular disease with pravastatin in Japan (MENDA) study: a prospective randomised controlled trial. *Lancet* 2006;368:1155–63.

Torres VE, Abebe KZ, Chapman AB, et al. Angiotensin blockade in late autosomal dominant polycystic kidney disease. *N Engl J Med* 2014;371:2267–76.

Lewis EJ, Hunsicker LG, Clarke WR, et al. Renoprotective effect of the angiotensin-receptor antagonist irbesartan in patients with nephropathy due to type 2 diabetes. *N Engl J Med* 2001;345:851–60.

Evans M, Bain SC, Hogan S, et al. Irbesartan delays progression of nephropathy as measured by estimated glomerular filtration rate: post hoc analysis of the irbesartan diabetic nephropathy trial. *Nephrol Dial Transplant* 2012;27:2555–63.

Taylor KS, et al. *BMJ Open* 2019;9:e030596. doi:10.1136/bmjopen-2019-030596. Downloaded from http://bmjopen.bmj.com/ on September 17, 2023 by guest. Protected by copyright.
56. Lambers Heerspink HJ, Weldegigiorgis M, Inker LA, et al. Estimated GFR decline as a surrogate end point for kidney failure: a post hoc analysis from the reduction of end points in non-insulin-dependent diabetes with the angiotensin II antagonist losartan (RENAAL) study and irbesartan diabetic nephropathy trial (IDNT). Am J Kidney Dis 2014;63:244–50.

57. Pfeffer MA, Braunwald E, Moyé LA, et al. Effect of captopril on mortality and morbidity in patients with left ventricular dysfunction after myocardial infarction. Results of the survival and ventricular enlargement trial. The SAVE Investigators. N Engl J Med 1992;327:697–7.

58. Downs JR, Clearfield M, Weis S, et al. Primary prevention of acute coronary events with lovastatin in men and women with average cholesterol levels: results of AFCAPS/TexCAPS. Air Force/Texas Coronary Atherosclerosis Prevention Study. JAMA 1998;279:1615–22.

59. Kendrick J, Shlipak MG, Talutzky et al. Effect of losartan on primary prevention of cardiovascular events in mild CKD and kidney function loss: a post hoc analysis of the air force/Texas coronary atherosclerosis prevention study. Am J Kidney Dis 2010;55:42–9.

60. Downs JR, Beere PA, Whittney E, et al. Design & rationale of the Air Force/Texas Coronary Atherosclerosis Prevention Study (AFCAPS/ TexCAPS). Am J Cardiol 1997;80:287–93.

61. Robins SJ, Collins D, Wittes JT, et al. Relation of gemfibrozil treatment and lipid levels with major coronary events: VA-HIT: a randomized controlled trial. JAMA 2001;285:1585–91.

62. Ronis RB, Ronis SJ, Ivane MK, et al. Rationale and design of the Department of Veterans Affairs high-density lipoprotein cholesterol intervention trial (HIT) for secondary prevention of coronary artery disease in men with low-high density lipoprotein cholesterol and desirable low-density lipoprotein cholesterol. Am J Cardiol 1993;71:45–52.

63. Goraya N, Munoz-Maldonado Y, Simon J. Treatment of metabolic acidosis in chronic kidney disease with fruits and vegetables but not sodium bicarbonate yields fewer adverse cardiovascular events after five-years follow up. Circulation Conference: Resuscitation Science Symposium RISS, 2017.

64. Goraya N, Simon J, C. O’H, et al. Treatment of metabolic acidosis in patients with stage 3 chronic kidney disease with fruits and vegetables or oral bicarbonate reduces urine angiotensinogen and preserves glomerular filtration rate. Kidney Int 2014;86:1031–8.

65. Bianchi S, Bigazzi R, Campese VM. Intensive versus conventional therapy to slow the progression of idiopathic glomerular diseases. Am J Kidney Dis 2010;55:671–81.

66. Alsebes BW, Diercks GFH, Hillegile HL, et al. Effects of fosinopril and pravastatin on cardiovascular events in subjects with microalbuminuria. Circulation 2004;110:1591–6.

67. Huskey J, Lindenfeld J, Cook T, et al. Effect of simvastatin on kidney function loss in patients with coronary heart disease: findings from the Scandinavian simvastatin survival study (4S). Atherosclerosis 2009;205:202–6.

68. Pedersen TR, Wilhelmsen L, Faergerman O, et al. Follow-up study of patients randomized in the Scandinavian simvastatin survival study (4S) of cholesterol lowering. Am J Cardiol 2000;86:257–62.

69. Design and baseline results of the Scandinavian simvastatin survival study of patients randomized for primary or secondary myocardial infarction. Am J Cardiol 1993;71:393–400.

70. Sever PS, Dahlof B, Poulter NR, et al. Prevention of coronary and stroke events with atorvastatin in hypertensive patients who have average or lower-than-average cholesterol concentrations, in the Anglo-Scandinavian Cardiac Outcomes Trial—Lipid Lowering Arm (ASCOT-LLA): a multicentre randomised controlled trial. Lancet 2003;361:1149–58.

71. Gupta A, Chang CL, Collier D. 758 the relationship between statin therapy and progression of renal damage among 10305 hypertensive patients randomised in the ASCOT-LIPID-Lowering arm (LLA). 79th European Atherosclerosis Society Congress, EAS 2011, Gothenburg Sweden, 2011:158–9.

72. Nanayakkara PWB, van Guldener C, ter Wee PM, et al. Effect of a treatment strategy consisting of pravastatin, vitamin E, and homocysteine lowering on carotid intima-media thickness, endothelial function, and renal function in patients with mild to moderate chronic kidney disease: results from the anti-oxidant therapy in chronic renal insufficiency (ATIC) study. Arch Intern Med 2007;167:1262–70.

73. Tavazzi L, Maggioni AP, Marchioli R, et al. Effect of rosvastatin in patients with chronic heart failure (the GISSI-HE-Fl trial): a randomised, double-blind, placebo-controlled trial. Lancet 2008;372:1231–9.

74. Tavazzi L, Tognoni G, Franzosi MG, et al. Rationale and design of the GISSI heart failure trial: a large trial to assess the effects of n-3 polysaturated fatty acids and rosuvastatin in symptomatic congestive heart failure. Eur J Heart Fail 2004;6:535–41.

75. Athyros VG, Mikhailidis DP, Liberooulos EN, et al. Effect of statin treatment on renal function and serum uric acid levels and their relation to vascular events in patients with coronary heart disease and metabolic syndrome: a subgroup analysis of the Greek atorvastatin and coronary heart disease evaluation (GREACE) study. Nephrol Dial Transplant 2007;22:11–27.

76. Athyros VG, Papageorgiou AA, Mercurios BR, et al. Treatment with atorvastatin to the National Cholesterol Educational Program goal versus ‘usual’ care in secondary coronary heart disease prevention. The GREeK Atorvastatin and Coronary-heart-disease Evaluation (GREACE) study. Curr Med Res Opin 2000;18:290–7.

77. Ford I, Bezyyk V, Stott DJ, et al. Reduced glomerular filtration rate and its association with clinical outcome in older patients at risk of vascular events: secondary analysis. PLoS Med 2009;6:e16.

78. Patel A, MacMahon S, Chalmers J, et al. Effects of a fixed combination of perindopril and indapamide on macrovascular and microvascular outcomes in patients with type 2 diabetes mellitus (the advance trial); a randomised controlled trial. Lancet 2007;370:829–40.

79. Heerspink HJJ, Nomiya M, Perkovic V, et al. Effects of a fixed combination of perindopril and indapamide in patients with type 2 diabetes and chronic kidney disease. Eur Heart J 2010;31:2888–96.

80. Peters R, Beckett N, Poulter R, et al. Kidney function in the very elderly with hypertension: data from the hypertension in the very elderly (HYVET) trial. Age Ageing 2013;42:253–8.

81. Beckett N, Peters R, Fletcher AE, et al. Treatment of hypertension in patients 80 years of age or older. N Engl J Med 2008;358:1887–98.

82. Buiplett C, Fletcher A, Beckett N, et al. Hypertension in the very elderly trial (HYVET): protocol for the main trial. Drugs Aging 2001;18:151–64.

83. Ting R-D, Keech AC, Drury PL, et al. Benefits and safety of long- term fenofibrate therapy in people with type 2 diabetes and renal impairment: the field study. Diabetes Care 2012;35:218–25.

84. FIELD Study Investigators. The need for a large-scale trial of fibrate therapy in diabetes: the rationale and design of the Fenofibrate Intervention and Event Lowering in Diabetes (FIELD) study. [ISRCTN64783481]. Cardiovasc Diabetol 2004;3:9.

85. Keech A, Simes RJ, Barter P, et al. Effects of long-term fenofibrate therapy on cardiovascular events in 9795 people with type 2 diabetes mellitus (the field study): randomised controlled trial. Lancet 2005;366:1849–61.

86. Zinman B, Wanner C, Lachin JM, et al. Empagliflozin, cardiovascular outcomes, and mortality in type 2 diabetes. N Engl J Med 2015;373:2117–28.

87. Wanner C, Inzucchi SE, Lachin JM, et al. Empagliflozin and progression of kidney disease in type 2 diabetes. N Engl J Med 2016;375:323–34.

88. Cornhol J, Bakris GL, Stevens SR, et al. Effect of sitagliptin on kidney function and respective cardiovascular outcomes in type 2 diabetes: outcomes from TECOS. Diabetes Care 2016;39:2304–10.

89. Green JB, Bethel MA, Armstrong PW, et al. Effect of sitagliptin on cardiovascular outcomes in type 2 diabetes. N Engl J Med 2015;373:232–42.

90. Zinman B, McMurray JJV, Krum H, et al. Eplenerone in patients with systolic heart failure and mild symptoms. N Engl J Med 2011;364:11–21.

91. Rossignol P, Dobre D, McMurray JJV, et al. Incidence, determinants, and prognostic significance of hyperkalemia and worsening renal function in patients with heart failure receiving the mineralocorticoid receptor antagonist eplerenone or placebo in addition to optimal medical therapy; results from the eplerenone in mild patients hospitalization and survival study in heart failure (EMPHASIS-HF). Circ Heart Fail 2014;7:51–8.

92. Brenner BM, Cooper ME, de Zeeuw D, et al. Effects of losartan on renal and cardiovascular outcomes in patients with type 2 diabetes and nephropathy. N Engl J Med 2001;345:861–9.

93. de Zeeuw D, Remuzzi G, Parving H-H, et al. Renoprotection in patients with type 2 diabetes and nephropathy: lessons from RENAL. Kidney Int 2004;65:2309–20.

94. Kowey PR, Dickson TZ, Zhang D, et al. Losartan and end- organ protection—lessons from the RENAL study. Clin Cardiol 2005;28:136–42.

95. Winkelmaier WC, Zhang Z, Shahinfar S, et al. Efficacy and safety of angiotensin II receptor blockade in elderly patients with diabetes. Diabetes Care 2006;29:2210–7.
97. Brenner BM, Cooper ME, de Zeeuw D, et al. The losartan renal protection study—rationale, study design and baseline characteristics of RENAAL (Reduction of Endpoints in NIDDM with the Angiotensin II Antagonist Losartan). *J Renin Angiotensin Aldosterone Syst* 2000;1:328–35.

98. Pitt B, Pfeffer MA, Assmann SF, et al. Spironolactone for heart failure with preserved ejection fraction. *N Engl J Med* 2014;370:1383–92.

99. Brenner BM, Cooper ME, de Zeeuw D, et al. The losartan renal protection study—rationale, study design and baseline characteristics of RENAAL (Reduction of Endpoints in NIDDM with the Angiotensin II Antagonist Losartan). *J Renin Angiotensin Aldosterone Syst* 2000;1:328–35.

100. Shepherd J, Blauw GJ, Murphy MB, et al. The design of a prospective study of pravastatin in the elderly at risk (prosper). PROSPER Study Group, prospective study of pravastatin in the elderly at risk. *Am J Cardiol* 1999;84:1192–7.

101. Shepherd J, Blauw GJ, Murphy MB, et al. Pravastatin in elderly individuals at risk of vascular disease (prosper): a randomised controlled trial. *Lancet* 2002;360:1623–30.

102. Baigent C, Landray MJ, Reith C, et al. The effects of lowering LDL cholesterol with simvastatin plus ezetimibe in patients with chronic kidney disease (study of heart and renal protection): a randomised placebo-controlled trial. *Lancet* 2011;377:2181–92.

103. Haynes R, Lewis D, Emberson J, et al. Effects of lowering LDL cholesterol on progression of kidney disease. *J Am Soc Nephrol* 2014;25:1825–33.

104. Sharp Collaborative Group. Study of heart and renal protection (sharp): randomized trial to assess the effects of lowering low-density lipoprotein cholesterol among 9,438 patients with chronic kidney disease. *Am Heart J* 2010;160:e10:785–94.

105. Bode B, Stenlöf K, Harris S, et al. Long-term efficacy and safety of canagliflozin over 104 weeks in patients aged 55–80 years with type 2 diabetes. *Diabetes Obes Metab* 2015;17:294–303.

106. Bode B, Stenlöf K, Sullivan D, et al. Efficacy and safety of canagliflozin treatment in older subjects with type 2 diabetes mellitus: a randomized trial. *Hosp Pract (1995)* 2013;41:72–84.

107. Kohan DE, Fioretto P, Tang W, et al. Long-term study of patients with type 2 diabetes and moderate renal impairment shows that dapagliflozin reduces weight and blood pressure but does not improve glycemic control. *Kidney Int* 2014;85:962–71.

108. Palmer SC, Navaneethan SD, Craig JC, et al. Hmg CoA reductase inhibitors (statins) for people with chronic kidney disease not requiring dialysis. *Cochrane Database Syst Rev* 2014;(5):CD007784.

109. Zhang X, Xiang C, Zhou Y-H, et al. Effect of statins on cardiovascular events in patients with mild to moderate chronic kidney disease: a systematic review and meta-analysis of randomized clinical trials. *BMC Cardiovasc Disord* 2014;14:19.

110. Herrington W, Emberson J, Mihaylova B, et al. Impact of renal function on the effects of LDL cholesterol lowering with statin-based regimens: a meta-analysis of individual participant data from 28 randomised trials. *Lancet Diabetes Endocrinol* 2016;4:329–39.