The value of maintaining normokalaemia and enabling RAASi therapy in chronic kidney disease

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Abstract

Background: People with chronic kidney disease (CKD) are at an increased risk of developing hyperkalaemia due to their declining kidney function. In addition, these patients are often required to reduce or discontinue guideline-recommended renin-angiotensin-aldosterone system inhibitor (RAASi) therapy due to increased risk of hyperkalaemia. This original research developed a model to quantify the health and economic benefits of maintaining normokalaemia and enabling optimal RAASi therapy in patients with CKD.

Methods: A patient-level simulation model was designed to fully characterise the natural history of CKD over a lifetime horizon, and predict the associations between serum potassium levels, RAASi use and long-term outcomes based on published literature. The clinical and economic benefits of maintaining sustained potassium levels and therefore avoiding RAASI discontinuation in CKD patients were demonstrated using illustrative, sensitivity and scenario analyses.

Results: Internal and external validation exercises confirmed the predictive capability of the model. Sustained potassium management and ongoing RAASi therapy were associated with longer life expectancy (+2.36 years), delayed onset of end stage renal disease (+5.4 years), quality-adjusted life-year gains (+1.02 QALYs), cost savings (£3135) and associated net monetary benefit (£23,446 at £20,000 per QALY gained) compared to an absence of RAASI to prevent hyperkalaemia.

Conclusion: This model represents a novel approach to predicting the long-term benefits of maintaining normokalaemia and enabling optimal RAASi therapy in patients with CKD, irrespective of the strategy used to achieve this target, which may support decision making in healthcare.

Keywords: Chronic kidney disease, Potassium, Hyperkalaemia, Renin-angiotensin-aldosterone system inhibitor, Economic modelling

Background

Chronic kidney disease (CKD) is defined as a condition that impairs kidney function, causing kidney damage to worsen over several months or years. It can progress to end stage renal disease (ESRD), which is fatal without dialysis or kidney transplantation. International guidelines recommend renin-angiotensin-aldosterone system inhibitors (RAASi), such as angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARBs), as first-line agents to prevent CKD progression [1, 2]. Treatment with RAASi has been shown to reduce blood pressure and proteinuria [3], delay estimated glomerular filtration rate (eGFR) decline [4], and lower the risks of kidney failure, cardiovascular (CV) morbidity and all-cause mortality in the CKD population [5]. Significant benefits have also been observed in patients requiring adjunctive treatment with mineralocorticoid receptor antagonists (MRAs) - an established treatment option for patients with persistent proteinuria, an important surrogate marker of ESRD [6]. However, despite their benefits, these drugs can compound the risk for hyperkalaemia among CKD patients [7–10]. Hyperkalaemia is often a major barrier for the optimal use of RAASI and/or MRAs in an already high-risk CKD population contributing to significant discrepancies between

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guideline recommendations and real-world practice in the use of RAASI and/or MRAs [8–11].

Hyperkalaemia is a clinically important electrolyte abnormality defined as a serum/plasma potassium level above the normal physiological range: 3.5–5.0 mmol/L (3.5–5.0 mEq/L) [12, 13]. Thresholds such as > 5.5, > 6.0 or > 7.0 mmol/L (mEq/L) are used to indicate severity [14]. As renal secretion is the main route of potassium elimination, patients with renal and metabolic comorbidities such as those with CKD or diabetes are at increased risk of hyperkalaemia. Whilst mild hyperkalaemia is usually asymptomatic, high levels of potassium are of concern to healthcare providers treating patients with CKD as it can induce electrophysiological disturbances that may cause life-threatening cardiac arrhythmias as well as muscle weakness or paralysis [15–18]. In the CKD population, hyperkalaemia is also associated with increased risk of hospitalisation, cardiac-related morbidity, and mortality [19–27].

There is currently a paucity of reliable treatment options for long-term serum potassium management and it has been suggested that best practice is to normalise serum potassium by down-titrating or discontinuing RAASI and/or MRA therapy in patients with cardiorenal disorders [28], despite evidence to suggest that submaximal dosing and discontinuation of RAASi are associated with increased risks of adverse outcomes in these patients [7, 29–31]. Furthermore, withholding RAASI treatment may lead to incremental healthcare costs associated with poor outcomes, for example, earlier onset of ESRD, hospitalizations due to cardiovascular causes, and cardiovascular mortality. Novel strategies that safely and effectively manage serum potassium levels, whilst allowing the optimal delivery of established renoprotective therapies, may therefore have significant potential to improve long-term outcomes in the CKD population.

With upcoming advances in pharmacological management of potassium levels for patients with cardiorenal syndromes, there is an increasing need to quantify the long-term health economic burden of hyperkalaemia in CKD, the consequences of suboptimal RAASI therapy, and the associated value of maintaining normokalaemia in this population. In the absence of long-term epidemiological evidence to describe hyperkalaemia in CKD, computer simulation modelling represents a valuable tool to simulate disease progression over a lifetime horizon, predict long-term outcomes, and inform clinical and reimbursement decision-making. Even though models characterising the natural history of CKD and predicting health economic outcomes associated with progression towards ESRD have been reported previously [32–37], none have modelled the inter-relationships between CKD progression, serum potassium levels, RAASI use, and the risk of adverse outcomes. The present study aimed to develop a novel model of CKD progression that incorporates serum potassium and RAASI use as variables to inform long-term health economic outcomes, and to quantify the value of maintaining normokalaemia and optimising RAASI therapy in advanced CKD patients at risk of hyperkalaemia.

Methods

Model development

A model was designed to characterise the natural progression of CKD patients (stage 3a onwards) and predict long-term health economic outcomes, which were associated with serum potassium levels and/or RAASI use. The model structure and inputs were informed by a targeted review of contemporary literature, UK clinical practice and National Institute for Health and Care Excellence (NICE) guidance where available. An advisory panel of four clinical experts, who were selected for their experiences in nephrology, cardiology, diabetes and general medicine, guided model conceptualisation and established the face validity of the model outputs.

Model structure

A patient-level simulation was developed in Microsoft Excel to capture the complex and chronic nature of CKD over a lifetime horizon (using a monthly cycle). Natural progression of CKD was modelled via a gradual decline in eGFR, where RAASI use impacted the rate of decline [4]. Similar to health states in previous CKD models [32–34], simulated patients progressed sequentially through CKD stages until ESRD onset (CKD stage 5; eGFR < 15 mL/min/1.73 m²), the initiation of renal replacement therapy (RRT) (RRT; eGFR < 8.6 mL/min/1.73 m² [38]) and death from disease-specific or general causes (Fig. 1).

Whilst not influential to our illustrative analysis, the model can capture fluctuations in serum potassium commonly observed in CKD patients (further details in Additional file 1), acute hyperkalaemia events when predicted serum potassium levels exceed a specified threshold (e.g. > 6.5 mEq/L) and changes in RAASI use (discontinuation, down-titration or up-titration) according to serum potassium levels [11, 29].

Clinical events, and the effect of serum potassium levels and/or RAASI use on event risks, were modelled prior to RRT according to published relationships (Table 1). Baseline probabilities of hospitalisation, CV events and mortality were sourced by CKD stage [39] and the likelihood of arrhythmia among CKD patients was assumed equivalent to the heart failure population [40]. Associations between serum potassium and arrhythmia, CV events, hospitalisation and mortality were modelled using incidence rate ratios (IRRs) [11],
Fig. 1 Flow diagram summarising CKD health states (shaded) and events (unshaded) applied in the model. Disease progression and the incidence of modelled events may be modified according to serum potassium level (grey circles), RAASi usage (white triangles) and/or CKD stage (black squares). CKD: chronic kidney disease; RAASi: renin-angiotensin-aldosterone system inhibitor

Table 1 Model inputs to inform cardiovascular, hospitalisation and mortality event rate predictions in CKD patients

| CKD Stage | Baseline Annual Event Rate | Odds Ratio by RAASi Use | Incidence Rate Ratio by Serum Potassium Subgroup | Source |
|-----------|----------------------------|-------------------------|-------------------------------------------------|--------|
| CV event  | Hospitalisation            | Mortality               |                                                  |        |
| CKD 1–2  | 0.0211 (0.0012)            | 0.1354 (0.0045)         | 0.0076 (0.0002)                                 | Go et al. [39] |
| CKD 3a    | 0.0365 (0.0012)            | 0.1722 (0.0045)         | 0.0108 (0.0004)                                 |        |
| CKD 3b    | 0.1129 (0.0012)            | 0.4526 (0.0067)         | 0.0476 (0.0007)                                 |        |
| CKD 4     | 0.218 (0.0024)             | 0.8675 (0.0090)         | 0.1136 (0.0018)                                 |        |
| CKD 5 (ESRD) | 0.366 (0.0048)         | 1.4641 (0.0090)         | 0.1414 (0.0031)                                 |        |

Odds ratio by RAASi use; mean (SE)²

Any versus none 0.820 (0.054) 1 (0)* 0.870 (0.069) Xie et al.[5]³

Incidence rate ratio by serum potassium subgroup; mean (SE)²

| eGFR (mL/min/1.73m²) | Any | < 30 | 30–40 | 40–50 | 50–60 | Any |
|----------------------|-----|------|-------|-------|-------|-----|
| K⁺ < 3.5 mEq/L       | 1.89 | 1.93 (0.45) | 1.77 (0.34) | 2.24 (0.38) | 2.06 (0.27) | 3.05 (0.29) |
| K⁺ 3.5–3.9 mEq/L     | 1.27 | 1.65 (0.22) | 1.35 (0.16) | 1.23 (0.13) | 1.13 (0.08) | 1.49 (0.08) |
| K⁺ 4.0–4.4 mEq/L     | 1.04 | 0.93 (0.09) | 0.99 (0.08) | 1.08 (0.07) | 1.01 (0.05) | 1.06 (0.04) |
| K⁺ 4.5–4.9 mEq/L     | 1 (0) | 1 (0) | 1 (0) | 1 (0) | 1 (0) | 1 (0) |
| K⁺ 5.0–5.4 mEq/L     | 1.01 | 1 (0.10) | 0.96 (0.09) | 1.07 (0.09) | 1 (0.07) | 1.14 (0.06) |
| K⁺ 5.5–5.9 mEq/L     | 1.12 | 1.34 (0.18) | 1.07 (0.14) | 1.23 (0.18) | 0.81 (0.11) | 1.6 (0.13) |
| K⁺ ≥ 6.0 mEq/L       | 1.88 | 3.65 (0.58) | 1.82 (0.44) | 1.91 (0.56) | 1.07 (0.30) | 3.31 (0.46) |

CKD: chronic kidney disease, CV: cardiovascular, eGFR: estimated glomerular filtration rate, ESRD: end stage renal disease, RAASi: renin-angiotensin-aldosterone system inhibitor, SE: standard error.

¹SE estimated from digitised plots showing 95% confidence intervals.
²SE estimated from 95% confidence intervals.
³Cardiovascular event defined in Go et al. [39] as: hospitalisation for coronary heart disease, heart failure, ischaemic stroke, and peripheral arterial disease.
⁴Cardiovascular event defined in Xie et al. [5] as: composite of fatal or nonfatal myocardial infarction, stroke, heart failure, cardiovascular death; or comparable definitions used by individual authors in studies included in the network meta-analysis.
⁵Luo et al. [11] reported incidence rate ratios (IRRs) for a major adverse cardiovascular event (MACE); these values were applied to the risk of both arrhythmia and cardiovascular events.

*Null value; no evidence identified.
while the impact of RAASi use on mortality and CV event incidence was modelled using odds ratios [5]. Background all-cause mortality was applied if it exceeded CKD-specific probabilities (adjusted for serum potassium level and/or RAASi use) according to gender-specific life tables [41].

This study aimed to estimate the value of maintaining normokalaemia irrespective of the strategy used to achieve this target, therefore costs and utilities related to pharmacological serum potassium management were not considered. For all other costs and benefits applied in the illustrative analyses, a UK healthcare payer perspective was adopted. Healthcare resource costs were obtained from published sources [1, 40, 42–46] and inflated to 2014–15 GBP [47]. Health-related quality of life was estimated via the multiplicative application of published health state and event utilities [48–55] to an age-dependent baseline value [56].

A summary of the methods used to model CKD progression and events is provided in Additional file 2: Table S1, an illustration of modelled cumulative event incidence for different patient characteristics in Additional file 3: Figure S1, and the inputs applied to modelled health states and events in Additional file 2: Table S2.

Model validation

To assess the validity of the model’s predictions, the modelled incidence of death and major adverse cardiovascular events (MACE) were used to derive modelled IRRs as a function of serum potassium level, which were compared to IRRs published by Luo et al. [11] (internal validation) and unadjusted IRRs derived from a retrospective, observational cohort study of CKD patients listed on the UK Clinical Practice Research Datalink (CPRD) [57, 58] (external validation).

Model application

The model was used to estimate the consequences of discontinuing RAASi therapy to maintain normal potassium levels in advanced CKD patients in terms of lifetime healthcare costs, life-expectancy and quality-adjusted life years (QALYs). Analysis was conducted for a cohort of CKD stage 3a patients (eGFR 52.5 mL/min/1.73m²), who were aged 60 years at baseline. Serum potassium was maintained at 4.5 mEq/L for all patients. Though the treatment arm represented a cohort of patients who received optimal serum potassium management to enable the continuation of RAASi therapy, the cost of such strategies (pharmacological and/or monitoring) was not included. All other costs and benefits were discounted at 3.5% per annum [59].

The health economic value of maintaining normokalaemia and optimising RAASi therapy was summarised in terms of incremental net monetary benefit (NMB) which was derived using willingness-to-pay (WTP) thresholds of £20,000–30,000 per QALY gained, in line with UK assessments of cost-effectiveness. In this analysis, incremental NMB represents the amount of money that could be spent on strategies to maintain normokalaemia that would be deemed good value for money.

Sensitivity and scenario analyses

The sensitivity of model predictions to changes in the following variables were assessed: time horizon (5, 10, 15 or 20 years); age (60 ± 20 years); CKD stage 3a (eGFR 52.5 mL/min/1.73m²), 3b (eGFR 37.5 mL/min/1.73m²) or 4 (eGFR 22.5 mL/min/1.73m²); male or female; RAASi efficacy for CKD progression (0–100% of base case), events (0–100% of base case), or both (base case ±20%); transplant probability, RRT mortality rate, event costs, event disutility, CKD 3–5 costs, CKD 3–5 utility, RRT costs and RRT utility (all base case ±20%).

Additional scenario analyses were conducted to investigate the impact of RAASi dosing on predicted outcomes. In the base case, patients continuing RAASi therapy were assumed to receive optimal dosing and consequently achieve the full benefit of RAASi observed in intention-to-treat trial populations [4, 5]. Though suboptimal dosing is common in clinical practice, there is a paucity of published data to describe the efficacy associated with such dosing; therefore, model parameters describing the cost and efficacy of RAASi treatment (related to CKD progression, CV and mortality risk) were scaled from 0 to 100% of base case values.

Results

Model validation

The model reproduced the associations between serum potassium and incident clinical outcomes from observational studies (Fig. 2). Internal validation exercises found that predicted IRRs for death and MACE were consistent with adjusted IRRs reported by Luo et al. [11] (R²: 0.995 and 0.995, respectively). External validation exercises against CPRD data showed that the model had a high predictive capability when estimating associations between serum potassium and mortality (R²: 0.961), while differences between predicted and observed IRRs for MACE may be due to the potential under-reporting of fatal MACE in the CPRD (R²: 0.743).

Model application

In modelled CKD patients who received optimal serum potassium management to enable ongoing RAASi therapy, predicted mean life expectancy was extended by 2.36 years when compared with patients who discontinued RAASi to maintain normokalaemia (Table 2). Ongoing RAASi treatment was associated with delayed CKD progression; average time spent in CKD stage 3a, 3b and 4 was longer in the treatment arm than in the
control arm (by 1.05, 1.69 and 0.86 years, respectively), while fewer patients receiving RAASi were predicted to reach ESRD within their lifetime (40% versus 49% in the control arm). Of those who progressed to ESRD, the estimated time to ESRD onset was delayed by 5.4 years in patients who received optimal serum potassium management and continued RAASi therapy (16.2 years versus 10.8 years without RAASi treatment). Similarly, the estimated time to RRT initiation was delayed by 6.2 years in the treatment arm, and the percentage of patients predicted to reach RRT over a lifetime was reduced by 9%.

Over a lifetime horizon, optimisation of both serum potassium levels and RAASi therapy was associated with per-patient cost savings of £3135 (£35 undiscounted), and QALY gains of 1.02 (1.43 undiscounted) (Table 2). As shown in Fig. 3, predicted cost savings in this cohort were largely driven by the avoidance of RRT (£14,143 per patient), while additional per-patient costs related to CKD management (+£8091), arrhythmia (+£327) and hospitalisation (+£2129) were attributed to improved survival and extended exposure to such risks. Overall, predicted cost savings and QALY gains corresponded to incremental NMB of £23,446 and £33,601, which represent the amount that could be spent on cost-effective strategies to maintain normokalaemia, at WTP thresholds of £20,000 and £30,000 per QALY gained, respectively (Table 2).

### Table 2 Predicted value of optimal serum potassium management to enable ongoing RAASi therapy in CKD patients

| Discounting | Outcome | Treatment arm (ongoing RAASi) | Control arm (no RAASi) | Incremental benefit | Incremental NMB |
|-------------|---------|-------------------------------|------------------------|---------------------|-----------------|
|             | Cost    | £108,685                      | £108,720               | -£35                | £28,570         | £42,837         |
|             | QALY    | 8.60                          | 7.17                   | 1.43                |                 |                 |
|             | LY      | 15.12                         | 12.75                  | 2.36                |                 |                 |
| Discounted  | Cost    | £72,067                       | £75,202                | -£3135              | £23,446         | £33,601         |
|             | QALY    | 6.68                          | 5.67                   | 1.02                |                 |                 |
|             | LY      | 11.32                         | 9.77                   | 1.56                |                 |                 |

CKD chronic kidney disease, ESRD end stage renal disease, LY life-year, NMB net monetary benefit, QALY quality-adjusted life-year, RAASi renin-angiotensin-aldosterone system inhibitor, WTP willingness-to-pay threshold

Simulated patients were aged 60 years at baseline, with CKD stage 3a (eGFR 52.5 mL/min/1.73m²) and serum potassium of 4.5 mEq/L; analyses were conducted independent of costs and utilities related to pharmacological potassium management.
**Sensitivity and scenario analysis**

Figure 4 summarises the results of deterministic sensitivity analyses. The predicted value of maintaining normokalaemia and RAASi therapy was greatest for CKD patients aged 40 years at baseline (£24,979 incremental NMB at £20,000 per QALY), and progressively decreased with increasing cohort age (£19,037 at 80 years of age); this was largely attributed to lower QALY gains predicted for older patients. Similarly, lower predicted QALY gains led to lower incremental NMB in patients with more advanced CKD at baseline (£23,446, £16,532 and £10,139 in CKD stage 3a, 3b and 4, respectively). As the modelled time horizon increased from 5 to 20 years, progressively greater QALY gains were captured; however, the most favourable cost savings (−£12,942) and incremental NMB (£31,831) were estimated at 15 years, driven by the timing of ESRD and initiation of RRT. Predicted incremental costs, QALYs and NMB were also sensitive to the modelled efficacy of RAASi therapy, particularly in relation to CKD progression. When the cost and efficacy of RAASi were scaled from 0 to 100% of base case values, discounted QALYs, life-years and NMB improved with increasing RAASi efficacy, while the incidence of ESRD and RRT, and associated costs, decreased with improved RAASi efficacy (Additional file 3: Figure S2).

Sensitivity analyses suggested that the timing and cost of ESRD, and subsequent RRT, were significant drivers of health economic outcomes. Figure 5 illustrates the evolution of mean health state utility (age-adjusted), undiscounted health state costs, and associated absolute NMB, which represents the undiscounted monetary value of one additional year in each health state. Among surviving CKD patients in the treatment arm, estimated NMB was £3057–£10,585 in the 18.8 years prior to RRT initiation, resulting from comparatively high mean health state utility (0.42–0.70) and low health state costs (£3404–£5311) associated with CKD stages 3a–5 (pre-dialysis). In contrast, the initiation of RRT was associated with lower health state utility and significantly increased health state costs, resulting in negative NMB (−£66,175 NMB at £20,000 per QALY). Estimated NMB among surviving RRT patients was observed to increase over time, as fewer patients received dialysis due to increased transplantation and death. Nevertheless, negative NMB estimated following the initiation of RRT served to significantly offset the value associated with increased survival prior to ESRD.

**Discussion**

Hyperkalaemia has been associated with increased risks of mortality and morbidity and with the dose down-titration or discontinuation of RAASi among CKD patients, which in turn is associated with worsening clinical outcomes. To our knowledge, this is the first study to describe the development of a natural history
model for CKD that utilised serum potassium and RAASi use as variables to inform long-term health economic outcomes. The model represents a valuable resource to assess the health economic burden of hyperkalaemia in CKD, quantify the value of serum potassium management, and inform future decision making. Findings from applications of the model are consistent with those recently reported by Bakhai et al., which demonstrated the value of potassium management to avoid hyperkalaemia, enable RAASi therapy and improve long-term health economic outcomes in patients with heart failure [60].

The accuracy and generalisability of the model predictions are limited by the inputs used to inform its development and several assumptions were necessary due to a paucity of published literature. For this reason, the model does not capture associations between RAASi use and hospitalisation or arrhythmia; and our base case analyses reflect a scenario in which patients received RAASi doses recommended in treatment guidelines for heart failure [42]. This study focussed on RAASi treatment due to its use as first line therapy, and the use of MRAs was not included in analyses. Consequently, the increased risk of hyperkalaemia associated with adjunctive MRA treatment and the value of maintaining normokalaemia to enable appropriate prescription of MRAs has not been captured.

The true value of optimal serum potassium management in CKD patients may be underestimated by current model predictions, as other clinical events and downstream benefits (such as the avoidance of implantable cardiac devices through reduced arrhythmia incidence) were not considered due to lack of published evidence but may be modelled if such data become available. Studies used to inform the incidence of hospitalisation did not stratify hospitalisations by cause [11, 39]; consequently, total admissions may be overestimated due to the inclusion of hospitalisations and events that may require hospitalisation (e.g. cardiovascular events) without available evidence to adjust for any potential double counting. Furthermore, an average linear decline was applied to model progression of eGFR. Factors such as age, sex, hypertension, acute kidney injury, diabetes
or proteinuria may influence rates of renal function decline and associated non-linear trajectories of eGFR are an area for further research and expansion of the model scope. Presence or absence of proteinuria was not modelled, however is also relevant considering guideline recommendations for prescription of RAASi therapy in CKD with urine albumin excretion > 30 mg/24 h (diabetics) and > 300 mg/24 h (non-diabetics) [2].

The presented validation exercises were limited to a comparison of modelled IRRs against observed values, due to a current paucity of literature available to validate model predictions of absolute outcomes. However, as new data describing the relationships between serum potassium, clinical events and long-term outcomes in CKD become available, ongoing model refinement and validation will ensure the continuing accuracy of its

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Fig. 5 Evolution of mean health state utility, undiscounted health state costs and associated absolute NMB (WTP at £20,000 per QALY gained) over 20 years in surviving CKD patients. CKD chronic kidney disease, ESRD end stage renal disease, NMB net monetary benefit, QALY quality-adjusted life-year, RRT renal replacement therapy, WTP willingness-to-pay threshold.
predictions. Finally, our analyses were conducted independent of pharmacological potassium management and/or monitoring costs, making the analysis relevant to any intervention, noting that interventions aimed to maintain normokalaemia in CKD are likely to incur additional expenditure. Extending the model to incorporate treatment costs and effects will allow the value of specific serum potassium management strategies to be quantified in future health economic analyses.

To illustrate the functionality of the model, we showed that patients who maintained normokalaemia and received ongoing RAASi therapy were predicted to benefit from delayed CKD progression, ESRD onset and RRT initiation, improved life expectancy, overall cost savings, and QALY gains, compared to those who discontinued RAASi to achieve equivalent serum potassium levels. However, as RAASi treatment is a known cause of hyperkalaemia careful monitoring for hyperkalaemia may be required to ensure a timely intervention should potassium levels rise. This requirement to monitor potassium may add to the burden of healthcare providers and patients and could hinder RAASi use in clinical practice.

Currently available approaches to restore potassium homeostasis in patients treated with RAASi are limited to dietary potassium restriction, correction of metabolic acidosis with use of bicarbonate solutions and increasing doses of loop diuretics to enhance renal potassium secretion, whilst recent advances include potassium-exchanging resins to bind potassium in the gut [61–63]. Potassium-binding agents can improve hyperkalaemia management whilst providing the opportunity for patients with CKD to benefit from the full cardio- and renoprotective effects of RAASi [64]. The model developed in this study has the potential to inform broad decision making on interventions for hyperkalaemia, as cost is an increasingly important consideration. To allow generalisability across a range of clinical scenarios, the presented model application did not focus on a single intervention and estimates of NMB from this study may be compared against the cost of any intervention to maintain normokalaemia and enable optimal RAASi therapy.

Sensitivity analyses demonstrated that greater value was expected among simulated patients of younger age and/or earlier CKD stage. Scenario analyses highlighted the detrimental impact of suboptimal RAASi dosing on clinical and economic outcomes. Though maintenance of lower doses of RAASi therapy are expected to provide some benefit, reduced efficacy relative to optimal dosing was associated with modelled increases in incidence of ESRD and RRT, and associated costs, and decreases in discounted QALYs, life-years and NMB. The model has flexibility for future applications investigating other scenario analyses.

Prolonged survival among the CKD population may increase the number of patients reaching ESRD and requiring RRT; a modality of care that may be considered not cost-effective due to its high costs and poor quality of life. Data arising from this study, whereby negative NMB associated with RRT partially offset the benefit of treatment prior to ESRD, subsequently illustrate the potential for RRT to confound the cost-effectiveness of health technologies for CKD. This observation is consistent with the notion that the inclusion of RRT costs may distort the results of economic analyses, and thus deny the availability of efficacious treatments that extend the life of CKD patients but do not directly impact the requirement for dialysis [65]. While others present a case for excluding dialysis costs in such evaluations [65], we alternatively propose that health economic outcomes in CKD may be more accurately described when partitioned according to ESRD status.

Conclusions
The model developed in this study is the first to our knowledge to characterise the natural history of CKD as a function of serum potassium levels and adherence to RAASi therapy, and highlights the current health economic consequences of underutilising RAASi to avoid hyperkalaemia. The optimisation of both serum potassium and RAASi therapy was associated with delayed CKD progression and initiation of RRT, improved quality of life and survival, and overall cost savings. Analyses were not intended to quantify the cost-effectiveness nor support the adoption of a specific treatment approach, but rather support the notion that maintaining normokalaemia (using guideline-recommended treatments, based on best available evidence) is a valuable strategy to enable optimal RAASi therapy and improve long-term health economic outcomes in CKD patients.

Additional files

Additional file 1: Supplementary Methods Mixed-effects model for serum potassium profiles. (DOCX 15 kb)

Additional file 2: Table S1. Summary of methods employed to model disease progression and events in CKD patients. Table S2. Inputs applied to modelled health states and events. (DOCX 37 kb)

Additional file 3: Figure S1. Model-estimated cumulative events over five years, according to CKD stage (a–c), RAASi use (d–f) and incidence of hyperkalaemia (g–i). Figure S2. Impact of RAASI cost and efficacy (0–100% of base case values) on discounted lifetime per-patient costs, QALYs, life years and NMB at conventional WTP thresholds. (DOCX 453 kb)

Abbreviations
ACE: Angiotensin converting enzyme; ARB: Angiotensin receptor blockers; CKD: Chronic kidney disease; CPRD: Clinical Practice Research Datalink; CV: Cardiovascular; eGFR: Estimated glomerular filtration rate; ESC: European Society of Cardiology; ESRD: End stage renal disease; IRR: Incidence rate ratios; MACE: Major adverse cardiovascular events; MIMS: Monthly Index of Medical Specialities; MRA: Mineralocorticoid receptor antagonist; NICE: National Institute for Health and Care Excellence; NMB: Net monetary benefit; QALY: Quality-adjusted life year; RAASi: Renin-angiotensin-aldosterone system inhibitor; RRT: Renal replacement therapy; WPT: Willingness-to-pay
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Availability of data and material
The datasets analysed during the current study were sourced from and are available in the original publications referenced.

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Authors' contributions
ME, EP, HF, GL, LQ and AB were involved in the design, interpretation and writing of the manuscript. HB and PM were involved in the design, analysis, interpretation and writing of the manuscript. All named authors meet the International Committee of Medical Journal Editors (ICMJE) criteria for authorship for this manuscript, take responsibility for the integrity of the work as a whole, and have given final approval to the version to be published. As guarantor and corresponding author, H. Bennett takes full responsibility for the work as a whole, including the study design, access to data, and the decision to submit and publish the manuscript.

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
M. Evans declares no conflict of interest. E. Palaka is a full-time employee of AstraZeneca (as principal investigator), and modest speaker honoraria from Biotronik, Medtronic, Abbot, research grant funding from AstraZeneca in relation to this study. C. Linde has AstraZeneca (as principal investigator), and modest expert witness funding from AstraZeneca. H. Furuland has received modest research grant funding from M. Evans declares no conflict of interest. E. Palaka is a full-time employee of AstraZeneca. The funding agreement ensured the authors' independence in designing the study, interpreting the data, and preparing the manuscript for publication.

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References
1. National Institute for Health and Care Excellence. Chronic kidney disease in adults: assessment and management. Clinical guideline (CG182). 2015. https://www.nice.org.uk/guidance/cg182. Accessed 28 July 2017.
2. Levin A, Stevens PE, Bilous RW, et al. Kidney disease: improving global outcomes (KDIGO) CKD work group. KDIGO 2012 clinical practice guideline for the evaluation and management of chronic kidney disease. Kidney Int Suppl. 2013;3(1):1-150.
3. Jafar TH, Schmid CH, Landa M, et al. Angiotensin-converting enzyme inhibitors and progression of nondiabetic renal disease: A meta-analysis of patient-level data. Ann Intern Med. 2001;135(2):73–87.
4. Evans M, Bain SC, Hogan S, Bilous RW. 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(6):2255–63.
5. Xie X, Liu Y, Perkovic V, et al. Renin-angiotensin system inhibitors and kidney and cardiovascular outcomes in patients with CKD: A Bayesian network meta-analysis of randomized clinical trials. Am J Kidney Dis. 2016;67(5):728–41.
6. Currie G, Taylor AH, Fujita T, et al. Effect of mineralocorticoid receptor antagonists on proteinuria and progression of chronic kidney disease: a systematic review and meta-analysis. BMC Nephrol. 2016;17(1):127.
7. Epstein M. Hyperkalemia constitutes a constraint for implementing renin-angiotensin-aldosterone inhibition: the widening gap between mandated treatment guidelines and the real-world clinical arena. Kidney Int Suppl. 2016(6):1–298.
8. Yildirim T, Arici M, Piskinapasa S, et al. Major barriers against renin–angiotensin–aldosterone system blocker use in chronic kidney disease stages 3–5 in clinical practice: a safety concern? Ren Fail. 2012;34(9):1095–9.
9. Preston RA, Afshartous D, Garg D, et al. Mechanisms of impaired potassium handling with dual renin-angiotensin-aldosterone blockade in chronic kidney disease. Hypertension. 2009;53(5):754–60.
10. Epstein M. Hyperkalemia as a constraint to therapy with combination renin–angiotensin system blockade: the elephant in the room. J Clin Hypertens. 2009;11(2):55–60.
11. Luo J, Brunelli SM, Jensen DE, Yang A. Association between serum potassium and outcomes in patients with reduced kidney function. Clin J Am Soc Nephrol. 2016;11(1):90–100.
12. Theisen-Toupal J. Hypokalemia and Hyperkalemia. Hosp Med Clin. 2015;4(1):34–50.
13. Viera AJ, Wouk N. Potassium Disorders: Hypokalemia and Hyperkalemia. Am Fam Physician. 2015;92(6):487–95.
14. Williams ME. Hyperkalemia. Crit Care Clin. 1991;7(1):155–74.
15. Esposito C, Bellotti N, Fasoli G, Plati A, Dall CA. Hyperkalemia-induced ECG abnormalities in patients with reduced renal function. Clin Nephrol. 2004;62(3):465–8.
16. Obalo CI, Ofili EQ, Mirza T. Hyperkalemia in CHF patients aged 63 to 85 years with subclinical renal disease. Am J Cardiol. 2003;90(6):663–5.
17. Mandal AK. Hypokalemia and hyperkalemia. Med Clin North Am. 1997;81(3):611–393.
18. Wiebe N, Klaarenbach SW, Allan GM, et al. Potentially preventable hospitalization as a complication of CKD: a cohort study. Am J Kidney Dis. 2014;64(2):230–8.
19. Collins AJ, Pitt B, Reaven N, et al. Association of serum potassium with all-cause mortality in patients with and without heart failure, chronic kidney disease, and/or diabetes. Am J Nephrol. 2017;46(3):213–21.
20. Nakhooul GN, Huang H, Amirjan S, et al. Serum potassium, end-stage renal disease and mortality in chronic kidney disease. Am J Nephrol. 2015;41(6):456–63.
21. Khanagavi J, Gupta T, Aronow WS, et al. Hyperkalemia among hospitalized patients and association between duration of hyperkalemia and outcomes. Arch Med Sci. 2014;10(2):251–7.
22. Jain N, Kotla S, Little BB, et al. Predictors of hyperkalemia and death in patients with cardiac and renal disease. Am J Cardiol. 2012;109(10):1510–3.
23. Hayes J, Kalantar-Zadeh K, Lu JL, et al. Association of hypo-and hyperkalemia with disease progression and mortality in males with chronic kidney disease: the role of race. Nephron Clin Pract. 2011;120(1):26–16.
24. Korgaonkar S, Tilea A, Gillespie BW, et al. Serum potassium and outcomes in CKD: insights from the RRI-CKD cohort study. Clin J Am Soc Nephrol. 2010;5(5):762–9.
25. Einhorn LM, Zhan M, Walker LD, et al. The frequency of hyperkalemia and its significance in chronic kidney disease. Arch Intern Med. 2009;169(12):1156–62.
26. Furuland H, McEwan P, Evans M, Linde C, Ayoubkhan D, Bahkai A, Palaka E, Bennett H, Qin L. Serum potassium as a predictor of adverse clinical outcomes in patients with chronic kidney disease: new risk equations using the UK clinical practice research datalink. BMC Nephrol. 2018;19(1):211.
27. Rosano G, Tamagno J, Kjeldsen KP, et al. Expert consensus document on the management of hyperkalemia in patients with cardiovascular disease.
treated with renin angiotensin aldosterone system inhibitors: coordinated by the working group on cardiovascular pharmacotherapy of the European Society of Cardiology. Eur Heart J Cardiovasc Pharmacother. 2018;4(3):180–8.

29. Epstein M, Reaven NL, Funk SE, et al. Evaluation of the treatment gap between clinical guidelines and the utilization of renin-angiotensin-aldosterone system inhibitors. Am J Manag Care. 2015;21(Suppl 11):s212–20.

30. Ouwerkerk W, Voors A, Anker S, et al. Determinants and clinical outcome of uptitration of ACE-inhibitors and beta-blockers in patients with heart failure: a prospective European study. Eur Heart J. 2017;38:1883–90.

31. Komajda M, Cowie MR, Tavazzi L, et al. Physicians’ guideline adherence is associated with better prognosis in outpatients with heart failure with reduced ejection fraction: the QUALIFY international registry. Eur J Heart Fail. 2017;19(11):1414–23.

32. Hoerger TJ, Wittenborn JS, Segel JE, et al. A health policy model of CKD: 1. Model construction, assumptions, and validation of health consequences. Am J Kidney Dis. 2010;55(3):452–62.

33. Orlando LA, Belasco E, Patel UD, Matchar DB. The chronic kidney disease model: a general purpose model of disease progression and treatment. BMC Med Inform Decis Mak. 2011;11(1):41.

34. Gandjour A, Tschulena U, Steppan S, Gatti E. A simulation model to estimate cost-offsets for a disease-management program for chronic kidney disease. Expert Rev Pharmacoecon Outcomes Res. 2015;15(2):231–47.

35. Tangri N, Stevens LA, Griffith J, et al. A predictive model for progression of chronic kidney disease to kidney failure. JAMA. 2011;305(15):1553–9.

36. Kibard BÂ. Estimating the long term impact of kidney donation on life expectancy and end stage renal disease. Transplant Res. 2013;2(1):22.

37. Lee CP, Chertow GM, Zenios SA, et al. A health policy model to estimate the cost and effectiveness of alternative dialysis initiation strategies. Med Decis Mak. 2006;26(5):535–49.

38. UK Renal Association. UK Renal Registry 18th Annual Report 2016. https://www.renalreg.org/wp-content/uploads/2015/01/web_book_07-04-16.pdf. Accessed 8 Dec 2016.

39. Go AS, Chertow GM, Fan D, McCulloch CE, Hsu CY. Chronic kidney disease and end-stage renal disease. Transplant Res. 2013;2(1):22.

40. Lee AJ, Morgan CL, Conway P, Currie CJ. Characterisation and comparison of health-related quality of life among patients treated with renin-angiotensin-aldosterone system inhibitors in UK patients with CKD. Nephrol Dial Transplant. 2017;32(Suppl 3):ii73–4.

41. Office for National Statistics. National Life Tables, United Kingdom; based on data for the years 2013–2015. 2016. www.ons.gov.uk/peoplepopulationandcommunity/birthsdeathsandmarriages/lifeexpectancies/datasets/nationallifetables/unitedkingdomreferencec tables. Accessed 8 Dec 2016.

42. Ponikowski P, Voors AA, Anker SD, et al. 2016 ESC guidelines for the diagnosis and treatment of acute and chronic heart failure: the task force for the diagnosis and treatment of acute and chronic heart failure of the European Society of Cardiology (ESC). Eur Heart J. 2016;37(27):2129–200.

43. Haymarket Media Group Ltd. Monthly index of medical specialties. 2017. http://www.mims.co.uk/. Accessed 16 Nov 2017.

44. National Institute for Health and Care Excellence. Chronic kidney disease (stage 5); peritoneal dialysis. Clinical guideline [CG125]. 2011. https://www.nice.org.uk/guidance/cg125. Accessed 8 Dec 2016.

45. Department of Health. NHS reference costs 2014 to 2015. 2015. https://www.gov.uk/government/publications/nhs-reference-costs-2014-to-2015.

46. Curtis L, Burns A, Personal social services research unit (PSSRU) unit costs of health and social care 2015. 2015. https://www.pssru.ac.uk/public/uc/2015/p full.pdf. Accessed 8 Dec 2016.

47. Góhler A, Geisler BP, Manne JM, et al. Utility estimates for decision-analytic modeling in chronic heart failure – health states based on New York heart association classes and number of Rehospitalizations. Value Health. 2009; 12(1):185–7.

48. Sennfeld K, Magnusson M, Carlsson P. Comparison of hemodialysis and peritoneal dialysis—a cost-utility analysis. Perit Dial Int. 2002;22(1):39–47.

49. Novakowski T, McEwan P, Evans M, et al. The relationship between serum potassium and incidence rates of major adverse cardiovascular events and mortality in UK patients with CKD. Nephrol Dial Transplant. 2017;32(Suppl 3):iii654–5.

50. Office for National Statistics. National Life Tables, United Kingdom; based on data for the years 2013–2015. 2016. www.ons.gov.uk/peoplepopulationandcommunity/birthsdeathsandmarriages/lifeexpectancies/datasets/nationallifetables/unitedkingdomreferencec tables. Accessed 8 Dec 2016.

51. Gencer M, Guler MF, Kurtil S, et al. A proposed predictor of hospitalizations, quality of life and death. J Card Fail. 2010;16(2):150–6.

52. Lacey EA, Musgrave RJ, Freeman JV, Todd AM, Scott P. Psychological morbidity after myocardial infarction in an area of deprivation in the UK evaluation of a self-help package. Eur J Cardiovasc Nurs. 2004;3(3):219–24.

53. Lacey EA, Musgrave RJ, Freeman JV, Todd AM, Scott P. Psychological morbidity after myocardial infarction in an area of deprivation in the UK evaluation of a self-help package. Eur J Cardiovasc Nurs. 2004;3(3):219–24.