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REVIEW

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How Do I Manage Hypertension in Patients with Advanced Chronic Kidney Disease Not on Dialysis? Perspectives from Clinical Practice

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Erietta Polychronopoulou¹ Gregoire Wuerzner ^{1,2} Michel Burnier^{1,2}

¹Service of Nephrology and Hypertension, University Hospital, Lausanne, Switzerland; ²Hypertension Research Foundation, Saint-Légier, Switzerland

Correspondence: Michel Burnier Service of Nephrology and Hypertension, University Hospital, Rue du Bugnon 17, Lausanne 1011, Switzerland Email michel.burnier@chuv.ch



Abstract: In the general population, the prevalence of moderate and severe chronic kidney disease (CKD) is usually below 5% but this figure is often higher in specific groups of patients such as those with type 2 diabetes. Patients with advanced CKD (CKD stage 3b and 4) are at high or very high cardiovascular risk, and their risk of progressing towards end-stage kidney disease (CKD stage 5) and the need of renal replacement therapy are elevated. Hypertension is a major cause of poor cardiovascular and renal outcomes in severe CKD. Therefore, an adequate control of blood pressure (BP) is mandatory. However, normalizing BP is often challenging in these patients because the clinical management of hypertension in advanced CKD is not well defined and rarely supported by large randomized controlled trials. In the present review, we discuss the characteristics of hypertension in advanced CKD, excluding dialysis, and its management integrating data from recent clinical studies and a pragmatic approach enriched by a long-standing clinical experience.

Keywords: hypertension, chronic kidney disease, CKD stage 3b-4, blockers of the reninangiotensin, diuretics, calcium antagonists, SGLT2 inhibitors

Introduction

Chronic kidney disease (CKD) is defined as abnormalities of kidney structure or function present for > 3 months leading to a sustained reduction in glomerular filtration rate (GFR) and/or to the occurrence of kidney damage markers, such as albuminuria.^{1,2} Today, CKD is classified in five stages according to the level of GFR (G1 to G5) and three categories according to the absence or presence of albuminuria (A1, A2 and A3). Worldwide, the prevalence of CKD ranges between 7% and 15%.^{3,4} The high and rising prevalence of subjects with CKD stages 1 to 3a and a mild to moderate reduction in estimated GFR (eg above 45 mL/min/1.73 m²), is due in part to the ageing population but also to the growing prevalence of diabetes and hypertension.^{3,5} Fortunately, the prevalence of moderate to severe (stage 3b) and advanced (stage 4) CKD with estimated GFR below 30 mL/min/ 1.73m² is lower, usually below 5%, but with large variations depending on countries⁶ and populations studied. In the US population, the percentage of NHANES 2013–2016 participants with CKD stages 3b and 4 was 2.1%.⁷ In the Swiss population,⁸ 1.6% of the 1353 participants randomly enrolled in the Swiss Survey on Salt had CKD stages 3b and 4. However, in a similar survey among Swiss patients with type 2 diabetes followed by general practitioners, the

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Patients with advanced CKD not on dialysis are at high or very high cardiovascular risk according to hypertension guidelines.¹¹ Indeed, the risk of myocardial infarction, heart failure, stroke or peripheral artery disease is significantly higher in patients with CKD than in the non-CKD population and, as expected, the cardiovascular risk rises further as kidney function declines.^{1,5,11,12} Hence, patients with advanced CKD have a greater risk to die from any cardiovascular event than to progress towards end-stage kidney disease (ESKD).¹³ In this respect, analyses made by the Chronic Kidney Disease Prognosis Consortium have demonstrated that both estimated GFR and the urinary albumin/creatinine ratio (ACR) are independent prognostic factors for mortality in the general population:¹⁴ the lower the GFR and the higher the ACR, the greater the hazard ratio for all-cause and cardiovascular mortality.¹⁴

In the context of kidney diseases, hypertension plays a special role, as it may be a cause as well as a consequence. Among the main clinical complications associated with the progression of kidney diseases such as anemia, metabolic acidosis, metabolic bonne disease, hypertension is the most prevalent. Indeed, in CKD stage 2, about one third of patients are hypertensive and when GFR is below 60 mL/min/1.73m², the prevalence of hypertension is up to >80% and reaches almost 100% in advanced CKD (stage 4 and 5 not on dialysis).¹⁵ Hypertension is an important determinant of CKD progression and a major risk factor for cardiovascular events. Therefore, lowering blood pressure (BP) to recommended targets represents a major task in the management of patients with CKD,¹⁶ the two primary objectives being: 1) to prevent the occurrence of cardiovascular events and the associated mortality and 2) to protect against the progression of CKD towards ESKD needing dialysis or transplantation.

Blood Pressure Profile and Targets in Advanced CKD

Two important patterns characterized the hypertensive phenotype in people with advanced CKD. The first is the high prevalence of nocturnal hypertension¹⁷ and uncontrolled masked hypertension due mainly to elevated BP values at night.^{18,19} The second is the high proportion of

patients with apparent or truly resistant hypertension.²⁰ The elevated nighttime BP in patients with advanced CKD can be explained in several ways including an increased prevalence of sleep apnea syndrome due to sodium and water retention.^{21,22} an increased sympathetic nerve activity and a reduced capacity to excrete sodium during daytime as GFR declines.²³ Indeed, a high nocturnal BP enables to maintain the 24 h sodium balance when kidney function is impaired. In this respect, Fukuda et al have nicely shown that as GFR declines sodium excretion, albuminuria and BP increase during the night.²⁴ Regarding resistant hypertension, several studies have demonstrated a high prevalence in patients with advanced CKD. In a recent review written by Georgianos and Agarwal, the prevalence of resistant hypertension in treated hypertensive patients with CKD ranged between 28.1% and 40.4%.²⁰ Both nocturnal and resistant hypertension are associated with an increased risk of cardiovascular events.18,19

The practical implication of these two observations is that out-of-office BP measurements providing both daytime and nighttime BP values should be performed more regularly in patients with hypertension and CKD. Ambulatory blood pressure monitoring should be used not only to confirm the eventual diagnosis of resistant hypertension but also to ascertain that, under treatment, BP control is achieved around the clock.²⁵ In this respect, as recommended by guidelines, only validated devices should be used.

Besides these two patterns of hypertension, one should mention that systolic hypertension is frequent in these patients, particularly in those with uncontrolled hypertension.²⁶ A high systolic BP seemed to account for most of the risk of progression to ESKD.²⁷ Another feature to consider is BP variability. Indeed, short-term systolic BP-variability is common in advanced CKD and analyses suggest that it may be involved in the sharp elevation of cardiovascular risk with worsening renal function.²⁸

According to the recent European Hypertension guidelines, the threshold values of BP for the treatment of hypertension in CKD are >140/90 mmHg and the recommended target is 130–140 mmHg systolic and 70–80 diastolic BP.¹¹ Guidelines coming from the United States are stricter with a proposed target BP at < 130/80 mmHg irrespective of the degree of albuminuria.²⁹ In 2013, the Kidney Disease Improving Global Outcomes (KDIGO) consensus conference recommended that BP be <140/90

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mmHg in all CKD patients without proteinuria and $\leq 130/80$ mmHg in patients with CKD and proteinuria.³⁰ However, KDIGO is now reconsidering these values, as new clinical trials have become available.³¹ So far, the KDIGO group did not publish any new official recommendations regarding target BPs.

Interestingly, there are no specific recommendations for BP targets in patients with advanced CKD and the most recent conclusions of KDIGO emphasized the need to obtain more data on the optimal management of hypertension in CKD stage 3a to stage 5 not on dialysis.³¹ In fact, whereas the benefits of a strict control of BP on cardiovascular outcomes is well recognized, the potential benefits of a well-controlled BP on CKD progression in advanced cases are still discussed. The main concerns are that an intensive lowering of BP in severe CKD might lead to more frequent hypotensive episodes and might eventually worsen kidney function. These concerns are particularly relevant in elderly and in CKD patients with comorbidities such as stroke or coronary heart disease. Table 1 summarizes the target BP recommended by various actual guidelines.

The wiser approach appears to be the individualization of BP targets (but below 140/90 mmHg) taking into account age, gender, ethnicity, comorbidities, the patient's acceptation and tolerance of treatments and other ongoing pharmacological therapies. Taken together, the diagnosis and management of hypertension in advanced CKD should take into account the following points:

- 1. Elevated nighttime BP is common and represents an important BP component of the cardiovascular and renal risk.
- 2. Ambulatory BP monitoring should be used more frequently to ascertain BP control around the clock. Use only validated devices.
- 3. Blood pressure should be lowered at least <140/90 mmHg in all patients and eventually lower (<130/80 mmHg) in young and/or proteinuric patients.
- 4. BP targets should however be individualized according to the patient's characteristics, comorbidities and concomitant treatments.

Non-Pharmacological Treatment of Hypertension in Severe CKD

All hypertension guidelines recommend lifestyle changes as the initial step of the management of hypertension. Of

Table I	Target Blood	Pressures	in the	General	Population	and
CKD and	d Diabetes Ac	cording to	Various	s Guideli	nes	

Society	General Population	СКД	Diabetes
ESC/ESH 2018 ¹¹	< 65y: <130 - 120 systolic <80 - 70 diastolic > 65y: <140 - 130 systolic <80 - 70 diastolic	<140 - 130 systolic <80 - 70 diastolic	< 65y: <130 - 120 systolic <80 - 70 diastolic > 65y <140 - 130 systolic <80 - 70 diastolic
ACC/AHA 2017 ²⁹	<130 systolic <80 diastolic	<130 systolic <80 diastolic	<130 systolic <80 diastolic
KDIGO 2012 ⁸⁵		Without albuminuria <140 systolic <90 diastolic With albuminuria <130 systolic <80 diastolic	
NICE 2019 ⁸⁶	< 80y: <140 systolic <90 diastolic > 80y: <150 systolic <90 diastolic	<130 systolic <80 diastolic	Without albuminuria <140 systolic <90 diastolic With albuminuria <130 systolic <80 diastolic
Hypertension Canada 2020 ⁸⁷	Low CV risk to to high risk <140 systolic <90 diastolic	High risk (severe CKD) < 120 systolic	<130 systolic <80 diastolic

Note: All values are in mmHg.

Abbreviations: Y, years; CV, cardiovascular; CKD, chronic kidney disease.

course, these lifestyle adaptations should be maintained when drug treatments are introduced. In patients with CKD, limitation of sodium intake to 5–6 g NaCl/day is an effective intervention to lower BP and albuminuria through every stage of CKD as confirmed in a Cochrane meta-analysis and a recent review of randomized clinical studies.^{32,33} For example, in the LowSALT CKD study, a double blind, placebo-controlled randomized crossover trial conducted in 20 adult patients with hypertension and CKD stages 3 and 4, BP was significantly lower on the low than on the high sodium diet, with a mean difference of -9.7 mmHg for systolic and -3.9 mmHg for diastolic BP.³⁴ In addition to lowering BP, salt restriction potentiates the antihypertensive effect and reinforce the ability of drugs such as blockers of the renin-angiotensin system or diuretics to lower albuminuria.³⁵ How much salt patients with advanced CKD should ingest daily is not well defined and whether a reduction of sodium intake enables to retard the progression of kidney diseases remains controversial, despite several observational studies supporting this hypothesis.³⁶ Nevertheless, an accepted recommendation is to obtain a sustained reduction of salt intake to about 6 g NaCl per day, which is not an easy objective to achieve.

In patients with hypertension and a preserved GFR (GFR>60 mL/min/1.73m²), there is now high-quality evidence demonstrating that increasing the potassium content of the diet lowers BP and reduces the incidence of stroke.³⁷ Experimentally, potassium has been shown to counterbalance the hypertensive effects of a high sodium diet.³⁸ However, the situation is very different in CKD patients with stages 3b–4 who may already have slightly elevated serum potassium levels at baseline. In these patients, the risk of hyperkalaemia is very high, specifically in patients with type 2 diabetes. As hyperkalaemia significantly increases the risk of death, physicians should avoid prescribing potassium supplements or drugs increasing serum potassium (such as non-steroidal anti-inflammatory drugs) in patients with an estimated GFR below 45 mL/min/1.73m².

Hypertension Drug Therapy in Severe CKD

In advanced CKD, recommended BP targets are rarely reached prescribing a single antihypertensive class. This is the reason why European guidelines propose to start immediately with a single pill combination associating a blocker of the renin-angiotensin system (RAS) and a calcium channel blocker (CCB) or a diuretic in CKD patients.¹¹ Then, if BP remains uncontrolled, the therapy should be intensified by combining the three drug classes (RAS blocker + CCB + diuretic), possibly in a single pill. However, the use of antihypertensive drug classes in patients with a low GFR (<30 mL/min/1.73m²) deserves some comments:

Blockers of the Renin-Angiotensin System

Blockers of the renin-angiotensin system are the mainstays of the management of hypertension in CKD because they enable to lower simultaneously systemic BP and the

albumin/creatinine ratio. Hence, RAS blockers may contribute to reduce both the cardiovascular and the renal risk of CKD patients.³⁹⁻⁴¹ RAS blockers are recommended as first-line therapy by most international hypertension, heart failure, diabetes and kidney diseases guidelines.^{11,42,43} ACC/AHA guidelines 2017 recommend that hypertensive adults with CKD (stages 3 or higher) and albuminuria receive an angiotensin converting enzyme inhibitor (ACEi), or an angiotensin receptor blocker (ARB) in case of ACE intolerance, to slow kidney disease progression.²⁹ Yet, there are two issues with the use of RAS blockers in advanced CKD. The first is the risk of hyperkalaemia, which is known to increase the occurrence of cardiac events and sudden deaths. In CKD stages 4 and 5, the incidence of hyperkalaemia defined as a plasma K >5.5 mmol/L can reach 30%. The second is the acute, reversible, decline in GFR at initiation of therapy, generally in the order of 10-20% depending on the baseline level of GFR. This decline in GFR can be a major concern in patients with advanced CKD and it is today a frequent cause of withdrawal of RAS blockers in order to gain some kidney function and delay the need of renal replacement therapy. Recent analyses have suggested that any increase in serum creatinine induced by ACEIs or ARBs is associated with a higher risk of CV events even when the increase is modest (<30%).⁴⁴ Moreover, the evidence that ACEI or ARBs still provide an effective cardiovascular and renal protection in advanced CKD is still weak. Thus, the issue arose of knowing whether RAS blockers should be maintained or stopped in advanced CKD.^{45,46} Today, there are no clear answers to this question and studies are ongoing assessing the benefits and risks of maintaining or stopping RAS blockers in advanced CKD. In one survey, the use of suboptimal doses of RAS inhibitors or withdrawing them in CKD stages 3 and 4, was associated with a two-fold increase in mortality when compared to patients maintained on maximum doses irrespective of comorbidity status.47 This would rather support the safety of RAS blockers in advanced CKD and favour the maintenance of RAS blockers when GFR is low. In fact, ACEIs and ARBs should be discontinued only when serious clinical problems arise, such as a substantial GFR decline, severe hyperkalaemia that cannot be readily controlled otherwise, persistent symptomatic hypotension, severe acidosis in the setting of very low GFR, and, potentially, in the very elderly. Today, the availability of potassium binders that can be prescribed chronically, such as patiromer, can help solving the issue

of hyperkalaemia and should prevent from withdrawing RAS blockers.⁴⁸

Diuretics

Regarding the use of diuretics in patients with advanced CKD (GFR<30 mL/min/1.73m²), the general concept is that thiazide and thiazide-like diuretics are quasi ineffective and most guidelines recommend the use of short (furosemide) or long-acting (torasemide) loop diuretics. However, this dogma has been challenged as an increasing number of small studies have demonstrated that thiazide diuretics can lower BP even in advanced CKD.⁴⁹ In one of them, long-acting furosemide (60 mg) and hydrochlorothiazide (25 mg) were compared. Initially, the two diuretics were given alone for 3 months and then both diuretics were given simultaneously for another 3 months period in 23 patients with hypertension and stages 4 or 5 CKD.⁵⁰ Hydrochlorothiazide was as effective as furosemide in reducing BP and combining the thiazide with the loop diuretic had a synergistic effect. In another randomized trial, amiloride/hydrochlorothiazide (5 mg/50 mg daily) was compared with dietary sodium restriction (60 mmol per day) in patients with hypertension and CKD stages 3 and 4. The diuretic combination produced a greater decrease in BP than sodium restriction.⁵¹ Thus, although loop diuretics remain the preferred diuretics in advanced CKD to control volemia and reduce BP as recommended in guidelines, one may now consider combining them with thiazides. This might be particularly effective in patients with CKD and difficult to control hypertension. Of note, when using thiazides as well as loop diuretics in advanced CKD, higher doses must be prescribed to obtain the desired therapeutic efficacy. Today, studies evaluating the efficacy of chlorthalidone in CKD stage 4 are ongoing. The use of aldosterone antagonists will be discussed below.

Calcium Channel Blockers

Dihydropyridine (DHP) calcium channel blockers (CCB) are potent vasodilators, which efficaciously lower BP in hypertensive patients with or without CKD. However, their ability to protect kidney function has long be questioned.⁵² Indeed, the classical first- and second-generation DHP calcium antagonists, such as nifedipine or amlodipine, increased rather than decreased intraglomerular pressure and hence possibly increased GFR but also proteinuria unless they induced a marked reduction in BP. In addition, in large CKD clinical trials, where

they usually served as control group, their impact on kidney progression was repeatedly inferior to that of RAS blockers, despite a comparable reduction in BP.^{53,54} In contrast, DHP calcium antagonists of the third generation such as lercanidipine, have a different effect on kidney function as they lower proteinuria.⁵³ At last, the most prevalent side effect of calcium antagonists is the development of peripheral edema. The incidence of edema with CCB is relatively high in patients with advanced CKD as these patients are already prone to retain sodium and water because of the reduced kidney function.55 Noteworthy, diltiazem, a non-DHP calcium antagonist, lowers proteinuria and BP in patients with CKD.⁵² However, diltiazem is a weaker vasodilator than DHP calcium antagonists are and in advanced CKD, physicians may consider the numerous drug interactions (more than 600 interactions described) reported with diltiazem.

Today, there is still a lack of specific trials demonstrating the benefits of calcium antagonist in patients with CKD beyond their impact on BP. Nonetheless, according to hypertension guidelines, CCB have an important place in the management of hypertension in CKD. This is due mainly to the results of the ACCOMPLISH (Avoiding Cardiovascular Events Through Combination Therapy in Patients Living with Systolic Hypertension) trial conducted in patients with hypertension and a high cardiovascular risk. In this trial, despite a comparable effect on BP, the benazepril-amlodipine combination was surprisingly superior to the benazepril-hydrochlorothiazide combination not only in reducing cardiovascular events⁵⁶ but also in preventing the progression of kidney diseases.55 Clinically, one of the limitation in CKD patients was the tolerability profile. Indeed the most frequent adverse event in patients with CKD was peripheral oedema (33.7% in the benazepril plus amlodipine group and 16.0% in the benazepril plus hydrochlorothiazide group).⁵⁵

As discussed above, the combination of a DHP calcium antagonist and a RAS blocker is now recommended as first-line therapy for the management of hypertension including in CKD but in advanced CKD most patients will need the triple association of a RAS blocker, a CCB and a diuretic. Calcium antagonists are particularly useful to lower BP in CKD patients of African ancestry alone or in association with diuretics.⁵⁷ Indeed, in a study published in 2019, amlodipine plus hydrochlorothiazide was as effective as amlodipine plus perindopril and more effective than perindopril plus hydrochlorothiazide at lowering BP in black patients in sub-Saharan Africa.⁵⁸

Beta-Blockers

In many recent hypertension guidelines, beta-blockers have been relegated to the 4th or even 5th line of therapy, unless there is a cardiac indication such as coronary heart disease or heart failure. Yet, the prescription of betablockers remains high in the general population as well as in patients with CKD even in the absence of cardiac comorbidity. This may in part be justified by the fact that the activity of the sympathetic nervous system is increased in all CKD stages and particularly when GFR is < 30 mL/ min/1.73m^{2.59,60} Therefore, although they are not proposed as first-line therapy, beta-blockers are still useful when BP is difficult to control and other recommended drug classes such as mineralocorticoid antagonists, for example, cannot be prescribed.

Mineralocorticoid Antagonists

In addition to increasing sodium reabsorption and potassium excretion in the late distal tubule and collecting duct, aldosterone is implicated in renal inflammatory and fibrotic processes.⁶¹ Thus, in the cardiovascular system, aldosterone has specific hypertrophic and fibrotic effects and can alter endothelial function. In the kidney, aldosterone may induce podocyte injury and mesangial cell proliferation. In line with experimental findings, several clinical studies have demonstrated that aldosterone antagonists such as aldactone or eplerenone, not only lower BP in hypertension but also lower proteinuria. In addition, they may retard CKD progression in diabetic or non-diabetic nephropathies on top of a RAS blocker.^{62,63} Moreover, in resistant hypertension, Williams et al have demonstrated the efficacy of aldactone as 4th line therapy after a RAS blocker, a calcium antagonist and a diuretic.⁶⁴

With these data in hand, one would be tempted to prescribe mineralocorticoid antagonists in advanced CKD, often characterized by a difficult to control BP. In this context, however, the main concern is the occurrence of severe hyperkalaemia taking into account that most of these patients are also receiving a RAS blocker. Wisely, the high risk of hyperkalaemia has been the main limiting factor for a widespread use of aldosterone antagonists in advanced CKD. Aldosterone antagonists might be used in patients with diuretic-induced hypokalaemia or CKD and heart failure but a careful monitoring of serum potassium is mandatory as hyperkalaemia can occur any time during the follow-up of these patients.⁴⁷

Very recently, finerenone, a new nonsteroidal, selective mineralocorticoid receptor antagonist has been reported to lower significantly the risk of CKD progression and cardiovascular events when compared to placebo in patients with CKD stages 3 and 4 and type 2 diabetes.⁶⁵ These cardiovascular and renal benefits were obtained with a very modest effect on systemic BP. Indeed, finerenone decreased mean systolic BP from baseline to month 1 and to month 12 by only -3.0 and -2.1 mmHg, respectively. As expected, the incidence of hyperkalaemia-related discontinuation of the trial regimen was slightly higher with finerenone than with placebo (2.3% and 0.9%, respectively) but it remained relatively low. Thus, finerenone will probably be useful to reduce the cardiovascular and renal complications of type 2 diabetes, but not necessarily to ameliorate the BP control of patients with diabetic nephropathy and an uncontrolled BP.

Sympatholytic Agents

The place of sympatholytic agents in the management of patients with hypertension and advanced CKD is poorly defined as no large randomized controlled trial has demonstrated that this class of antihypertensive agents is able to reduce cardiovascular events or to slow the progression of kidney diseases. Yet there are some indications suggesting that some agents of this drug class may be helpful to complete the treatment of CKD patients with uncontrolled BP. Thus, moxonidine 0.3 mg/d was found to be as effective as 20 mg nitrendipine in lowering BP on top of a RAS blocker in 177 patients with advanced CKD.⁶⁶ In this study, the increase in serum creatinine was significantly greater with nitrendipine than with moxonidine. Some authors have proposed to revise the place of low doses reserpine in the management of uncontrolled hypertension.⁶⁷ In a small group of patients with refractory hypertension, defined as uncontrolled BP despite > 5 antihypertensive drugs including a mineralocorticoid antagonist, reserpine reduced mean systolic and diastolic automated office BP by 29.3 \pm 22.2 and 22.0 \pm 15.8 mm Hg, respectively.⁶⁸ These results need to be confirmed in a larger number of patients but they suggest that reserpine at low dose, thereby reducing side effects, might be effective in refractory hypertension. Centrally acting drugs such as clonidine or methyldopa are relatively safe in patients with CKD. They may be used in some patients with

resistant hypertension although there are no specific data in advanced CKD. In patients with resistant hypertension and normal renal function, clonidine was found to be as effective as spironolactone to lower BP.⁶⁹

New Therapies to Lower BP in Hypertension

In the last 5 years, several new therapeutic options have become available to improve the management of CKD patients reducing their tendency to develop cardiovascular events and to progress towards ESKD. These new therapies concerned essentially patients with type 2 diabetes and a diabetic nephropathy with the development of inhibitors of sodium-glucose cotransporter 2 (SGLT2) or glucagon-like peptide-1 receptor agonists (GLP-1-RA). In patients with type 2 diabetes and kidney disease, both SGLT2-inhibitors and GLP-1-RA demonstrated significant beneficial effects on cardiovascular, mortality, and kidney outcomes.^{70,71} In patients with type 2 diabetes and stages 3b-4 CKD, dapagliflozin did not decrease HbA1c but decreased albuminuria, BP and body weight to a clinically meaningful extent.⁷² In large randomized clinical trials, the marked cardiovascular and renal benefits of SGLT2 inhibitors and GLP-1-RA were obtained with relatively modest decreases in BP suggesting other mechanisms of cardiac and renal protection. Nonetheless, SGLT2 inhibitors have been reported to lower BP effectively even on top of RAS blockers or diuretics.⁷³

Whether SGLT2 inhibitors provide the same clinical benefits in patients with diabetic and non-diabetic nephropathies was the main purpose of the DAPA-CKD trial.⁷⁴ DAPA-CKD enrolled 4304 CKD patients with an estimated GFR of 25-75 mL/min/1.73 m² and albuminuria of 200-5000 mg/g.⁷⁵ Two-thirds of patients had type 2 diabetes and the median follow-up was 2.4 years. The mean baseline eGFR was 43 mL/min/1.73 m² and 14.5% of patients had a GFR $<30 \text{ mL/min}/1.73 \text{ m}^2$. The trial was stopped early for overwhelming efficacy. Dapagliflozin reduced significantly the risk of the primary combined endpoint of >50% eGFR decline, onset of ESKD or kidney or CV death with a hazard ratio of 0.61 (95% CI 0.51–0.72).⁷⁶ The renal benefits were comparable in patients with diabetic and non-diabetic kidney diseases. The impact on cardiac events and mortality were also significant. In heart failure, patients on empagliflozin were also found to be at lower risk of cardiovascular death and hospitalization for heart failure or kidney

outcomes than those on placebo, regardless of the presence or absence of diabetes.⁷⁷

The second new therapeutic approach that has been developed to improve the control of BP in hypertension, including in CKD, is renal denervation. Initially, the efficacy of renal denervation was explored essentially in patients with resistant hypertension. In this context, effective renal denervation was found to be safe and to produce meaningful reductions in BP as assess by office or ambulatory BP.⁷⁸ A more recent study has demonstrated that renal denervation is also effective in untreated patients with essential hypertension.⁷⁹ As activation of the sympathetic nervous system is an important determinant of the cardiovascular and renal risk of patients with CKD, it was reasonable to investigate the potential benefits and risks of renal denervation in CKD. Experimental data in CKD models suggested that renal denervation was indeed effective to control hypertension.⁸⁰ Consequently, the procedure was also tested in patients with moderate to severe CKD and difficult to control BP. Hering et al reported a marked and significant decrease in office and ambulatory BP in 15 patients with resistant hypertension and stages 3-4 CKD (mean GFR, 31 mL/min/1.73 m²) with no deleterious impact on renal function.81 Post-hoc analyses of the SIMPLICITY program have also confirm the long-term renal safety of renal denervation in resistant hypertension. Thus, renal denervation represents an additional therapeutic option to control hypertension in advanced CKD but more large randomized controlled trials in hypertensive patients with CKD are needed before recommending a wider use of renal denervation in this population.⁸² In severe CKD, the indication for renal denervation should be evaluated on an individual basis balancing the potential benefits and risks of aggravating renal function. The discussion should preferably involve cardiologists (or radiologists) and nephrologists and treating physicians in an interdisciplinary approach.

Conclusions

The control of blood pressure in patients with CKD and hypertension is a major task for physicians as a normalization of BP contributes to reduce their cardiovascular and renal risk. However, in advanced CKD, the management of hypertension becomes more delicate for several reasons. Firstly, there are no specific recommendations on how hypertension should be treated in that specific context, because prospective randomized controlled trials are lacking. Secondly, the pharmacology of antihypertensive drugs and the tolerability profile change in advanced CKD and the

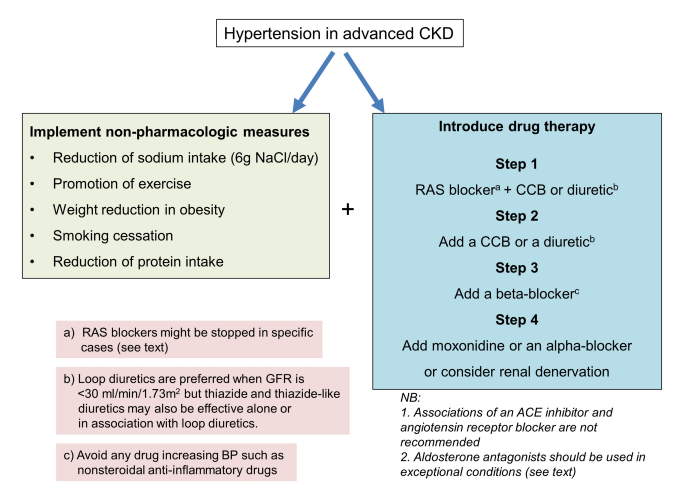


Figure I Schematic representation of a possible strategy to manage hypertension in severe CKD.

incidence of relevant drug-induced side effects (eg peripheral edema, hyperkalemia, worsening of GFR) increases. Thirdly, the cardiovascular and renal benefits of maintaining some drug classes such as RAS blockers are often called into question. Fourthly, patients with CKD belong to the group of subjects with one of the highest burdens of daily pill intake with up to >20 pills per day depending on the severity of their disease.⁸³ Therefore, poor medication adherence is a real issue in advanced CKD, which can lead to uncontrolled hypertension and a more rapid decline in kidney function.⁸⁴ Despite these limitations, physicians must control BP following a pragmatic approach (as illustrated in Figure 1) taking into account the complexity of the clinical situation, which is accentuated by comorbidities, other CKD complications (anemia, acidosis) and the high pill burden. While waiting for additional supporting evidence from trials, individualization of the management of hypertension in severe CKD appears to be the most reasonable and effective approach taking also into account the patient's preference of care and acceptance of treatments.

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The authors of this review have no conflict of interest to declare.

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