Integrated control of hypertension by olmesartan medoxomil and hydrochlorothiazide and rationale for combination

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Abstract: Hypertension affects nearly one-third of all individuals in the US, yet one-half of all treated patients achieve blood pressure (BP) controlled to recommended goals. The percentage of patients with uncontrolled BP is likely to be much higher when considering the number of patients who are not even aware of their hypertensive state. Elevated BP is associated with increased risks of cardiovascular events and end-organ damage. Antihypertensive monotherapy is not always sufficient to achieve BP goals, and thus more aggressive treatment regimens need to be considered. Antihypertensive combination therapy, which may improve tolerability, offers the benefit of targeting different mechanisms of action. Numerous outcomes studies support the use of a renin–angiotensin system inhibitor as a first-line choice in antihypertensive therapy. This review discusses the benefits of combination therapy with the angiotensin type II receptor blocker olmesartan medoxomil (OM) paired with the thiazide diuretic hydrochlorothiazide (HCTZ). The pharmacokinetic properties of OM will be reviewed in addition to efficacy studies that support OM + HCTZ combination therapy over other possible antihypertensive combinations. Finally, a rationale for choosing HCTZ over another diuretic, chlorthalidone, will also be discussed based on pharmacokinetic differences, clinical concerns, and trends in use.

Keywords: antihypertensives, blood pressure, combination therapy, HCTZ

Introduction
Hypertension, a highly prevalent condition that affects 29% of the population in the US, is a modifiable risk factor for cardiovascular (CV) morbidity and mortality, stroke, and renal failure. Indeed, there is a linear correlation between blood pressure (BP) and the risk of death from ischemic heart disease and stroke, regardless of age; this risk is doubled for each 20- or 10-mmHg increase in systolic BP (SBP) and diastolic BP (DBP), respectively. An increasing number of patients with hypertension in the US are receiving treatment. However, approximately 50% of patients receiving treatment fail to attain recommended BP goals of <140/90 mmHg or <130/80 mmHg for patients with diabetes mellitus or chronic renal disease. Current practice guidelines are based on clinical trial evidence, demonstrating that treating patients with hypertension to defined BP thresholds, or goals, improves long-term outcomes. Achieving BP control in a greater proportion of patients with hypertension will require treating more patients, treating them earlier, and intensifying their therapy when treatment goals are unmet.

An important component of ensuring successful treatment is the use of more aggressive treatment strategies. BP goals are reached in only one-third of patients receiving monotherapy. As a result, combination therapy is required to achieve recommended
BP goals in the majority of patients with hypertension, particularly those with stage 2 hypertension, and treatment guidelines emphasize the importance of starting antihypertensive combination therapy in patients with a BP level that exceeds the goal by >20/10 mmHg.3

Combination therapy should comprise different classes of agents with complementary mechanisms of action, which may provide an antihypertensive effect greater than either component alone, and with a favorable tolerability profile.3 Blockade of the renin–angiotensin system (RAS) pathway by angiotensin type II receptor blockers (ARBs) provides an antihypertensive effect that can be enhanced by the addition of hydrochlorothiazide (HCTZ).7 HCTZ acts in the kidney by blocking the reabsorption of sodium and chloride in the distal portion of the kidney tubule.8 In addition, HCTZ is believed to have direct (vasodilation) or indirect effects on the blood vessel itself, although the exact mechanism explaining this is unknown.9 Use of HCTZ alone causes volume contraction that has been shown to cause an increase in RAS pathway activity to compensate.7 The synergistic addition of a RAS inhibitor to HCTZ blunts this physiological response to diuresis, thereby achieving volume contraction with decreased RAS activity.7

ARBs are a well-tolerated drug class. Olmesartan medoxomil (OM) is a widely prescribed ARB that has been shown in some head-to-head studies to have greater BP-lowering efficacy than older ARBs such as losartan potassium (LOS),10,11 valsartan, and irbesartan.10 In a 12-week randomized, double-blind, forced-titration study, patients received LOS, OM, or valsartan.12 At week 8, reductions in seated cuff DBP (SeDBP) were significantly greater in the OM 40-mg group compared with the LOS group (100 mg once daily). By week 12, however, there were no significant differences in BP-lowering efficacy between OM (40 mg), valsartan (320 mg), and LOS (50 mg twice daily). In a subgroup analysis of this study in Black patients, OM demonstrated greater efficacy by week 8 compared with LOS; however, all drugs had similar antihypertensive effects by week 12.13 Recently, the newest member of the ARB family, azilsartan medoxomil, which has been approved for use in the US, has superior efficacy at its highest dose compared with OM and valsartan.14

Pharmacokinetic differences such as higher angiotensin II receptor type 1 (AT₁) receptor affinity, longer terminal elimination half-life, and slower AT₁ receptor disassociation help contribute to the efficacy of OM.15,16 This article briefly reviews the efficacy and safety of combining the ARB OM with HCTZ in the management of hypertension and provides an update on current findings from recent clinical studies.

**OM/HCTZ: pharmacokinetics and pharmacodynamics**

OM is a prodrug that is hydrolyzed in the gastrointestinal tract to form its active metabolite, olmesartan.17 Once absorbed, the metabolite does not undergo further changes, and 35%–50% of the absorbed dose is excreted in the urine.17 Peak plasma concentrations are achieved in 1–2 hours, followed by an elimination half-life of 13 hours.17 Steady-state plasma concentrations are achieved in 3–5 days with once-daily dosing.17 HCTZ is not metabolized and is rapidly eliminated by the kidney, with a plasma half-life between 5.6 and 14.8 hours.17 No significant pharmacokinetic drug–drug interactions occur when OM and HCTZ are coadministered.18 The pharmacokinetics of OM 20 mg + HCTZ 25 mg in healthy subjects were similar to OM and HCTZ monotherapy at the same doses with regards to area underneath the concentration–time curve and maximum plasma concentration values at steady state and time to maximum plasma concentration values.18

OM selectively binds the AT₁ receptor with high affinity, slow disassociation, and a high degree of insurmountable antagonism.15 OM is a more potent inhibitor of angiotensin II receptor binding than LOS and its active metabolite19 and dissociates from the receptor more slowly than telmisartan.15 OM inhibits the pressor effects of angiotensin I at doses of 2.5–40 mg; this inhibitory effect is dose-dependent.17 HCTZ combined with OM causes diuresis to begin within 2 hours of administration and peaks at approximately 4 hours, with a duration of 6–12 hours.17

**Efficacy of OM and HCTZ combination therapy**

Several studies have demonstrated the efficacy of OM + HCTZ for lowering BP and enabling the achievement of BP goals. A summary of OM + HCTZ efficacy studies are presented in Table 1.

In a multicenter, randomized, double-blind factorial design study, 502 patients were assigned to placebo, OM monotherapy (10, 20, or 40 mg/day), HCTZ monotherapy (12.5 or 25 mg), and OM/HCTZ combination therapy (10/12.5, 10/25, 20/12.5, 20/25, 40/12.5, and 40/25 mg). All six combinations of OM + HCTZ produced statistically significant reductions in BP from baseline relative to placebo, and all OM + HCTZ combinations had greater BP reductions than their individual components.20 The BP reduction achieved at the maximum dose of OM/HCTZ...
Clinical trials assessing the antihypertensive efficacy of OM/HCTZ combination therapy

<table>
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<th>Study</th>
<th>Patients (N)</th>
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<th>SeSBP/SeDBP reduction (mmHg)</th>
<th>% achieved SeSBP target</th>
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<td>PLA</td>
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<td>3.3/8.2</td>
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Notes: *DBP reductions not reported; bbaseline at end of OM 40 mg run-in period; *all patients with T2DM.

Abbreviations: AML, amloidipine; BEN, benazepril; BP, blood pressure; DBP, diastolic blood pressure; HCTZ, hydrochlorothiazide; OM, olmesartan medoxomil; PLA, placebo; SBP, systolic blood pressure; SeSBP, seated cuff blood pressure; SeDBP, seated cuff diastolic blood pressure; SeSBP, seated cuff systolic blood pressure; T2DM, type 2 diabetes mellitus.

40/25 mg was 26.8/21.9 mmHg from a baseline BP of 153.6/103.4 mmHg. This study reported individual SBP and DBP goals, and at the maximum OM/HCTZ 40/25 mg dose, 87.2% achieved SBP <140 mmHg, while 79.5% achieved DBP <90 mmHg.

A multicenter, double-blind study by Kereiakes et al randomized 191 patients with stage 2 hypertension to an OM + HCTZ or benazepril (BEN) + amloidipine (AML) combination treatment regimen for 12 weeks. Doses were up-titrated in a stepwise fashion from OM or BEN monotherapy if BP was ≥120/80 mmHg. Titration steps in the OM treatment group were OM 20 mg, OM 40 mg, OM/HCTZ 40/12.5 mg, OM/HCTZ 40/25 mg, and OM/HCTZ 40/25 mg, while titration steps in the AML treatment group were BEN 10 mg, BEN 20 mg, BEN/AML 20/5 mg, and BEN/AML 20/10 mg. The OM + HCTZ treatment arm was associated with a greater reduction in SBP from baseline than BEN + AML at 32.5 mmHg vs 26.5 mmHg ($P < 0.024$). A cumulative BP goal of <140/90 mmHg was achieved by 66.3% of patients treated with OM + HCTZ compared with 44.7% of patients in the BEN + AML treatment arm ($P < 0.006$).

The Benicar Efficacy: New Investigative Findings showed Olmesartan Medoxomil Safely and Effectively Reduced Blood Pressure Compared With Placebo in a Clinical Evaluation of Patients With Stage 1 and Stage 2 Hypertension (BENFORCE) study was a 12-week, randomized, double-blind, placebo-controlled, titration study in 276 patients with stage 1 or stage 2 hypertension.22 Patients were randomized to placebo or an OM treatment regimen for a period of 12 weeks. If BP was ≥120/80 mmHg, patients were up-titrated in a stepwise fashion from monotherapy to a maximum of OM/HCTZ 40/25 mg. The titration steps were OM 20 mg, OM 40 mg, OM/HCTZ 40/12.5 mg, and OM/HCTZ 40/25 mg. The OM-based treatment regimen provided significantly greater least-squares mean reductions in seated BP (SeSBP) from baseline compared with placebo (22.3/12.1 vs 0.1/−0.8 mmHg; $P < 0.0001$).22 The achievement rate of a cumulative BP goal of <140/90 mmHg was significantly higher in OM-based treatment vs placebo recipients (74.1% vs 30.7%; $P < 0.0001$).22 Furthermore, BP normalization (<120/80 mmHg) was also achieved by more patients treated with the OM-based regimen vs placebo (27.3% vs 1.5%; $P < 0.0001$) (Figure 1).22 Recently, a subgroup analysis of BENFORCE indicated that the significant improvements in BP lowering were achieved with OM-based therapy vs placebo, regardless of race, age, or sex (Figure 2).

A recent European study investigated the safety and tolerability of OM/HCTZ in 1226 patients with stage 2 hypertension.24 Patients entered an 8-week open-label period and were treated with OM 40 mg per day. Patients who failed to achieve BP control (trough seated cuff SBP [SeSBP] of 140–180 mmHg and SeDBP of 90–115 mmHg,
and mean 24-hour DBP $\geq 80$ mmHg and $\geq 30\%$ of daytime DBP $>85$ mmHg) entered a randomized double-blind treatment phase of 8 weeks. Patients were randomized in a 2:2:2:1 scheme to OM 40 mg, OM/HCTZ 20/12.5 mg, OM/HCTZ 40/12.5 mg, and OM/HCTZ 40/25 mg. The primary endpoint was the change from baseline in SeDBP from week 8 to the end of week 16. For the primary endpoint for the highest dosage of OM/HCTZ 40/25 mg, the change in SeDBP was $-11.2$ mmHg compared with $-5.7$ mmHg for patients who remained on OM 40 mg ($P < 0.0001$). The change in SeSBP for the same time period was $-16.2$ mmHg for OM/HCTZ 40/25 mg compared with $-8.9$ mmHg for OM

Figure 1 Efficacy results from the BENIFORCE trial. (A) Mean change from baseline to week 12 or last observation carried forward in seated cuff BP by titration step in the total efficacy cohort. (B) Proportion of patients who achieved BP $< 140/90$ mmHg in the total efficacy cohort. $^{11}$

Notes: $^* P < 0.05$, $^† P < 0.01$, $^‡ P < 0.001$, $^§ P < 0.0001$ for within-group comparisons between study baseline and end of treatment using paired t-test.


Abbreviations: BENIFORCE, Benicar Efficacy: New Investigative Findings Showed Olmesartan Medoxomil Safely and Effectively Reduced Blood Pressure Compared With Placebo in a Clinical Evaluation of Patients With Stage 1 and Stage 2 Hypertension; BP, blood pressure; HCTZ, hydrochlorothiazide; OM, olmesartan medoxomil; SeDBP, seated cuff diastolic blood pressure; SeSBP, seated cuff systolic blood pressure.
40 mg ($P < 0.0001$). The SeBP target of $<140/90$ mmHg ($<130/80$ mmHg for patients with diabetes) was achieved by 42.1% of patients who received OM/HCTZ 40/25 mg compared with 24.8% of those treated with OM 40 mg.

Assessment of 24-hour ambulatory BP efficacy with OM/HCTZ combination in patients with difficult-to-treat hypertension

Two classes of patients with hypertension that is often difficult to treat are patients with type 2 diabetes mellitus (T2DM) and the elderly. Diabetes mellitus affects an estimated 25.8 million US residents of all ages. T2DM is associated with higher risks of CV disease, nephropathy, and retinopathy – hard endpoints that are associated closely with BP control. It is recommended that patients with T2DM and hypertension be treated to a more aggressive BP goal of $<130/80$ mmHg, which will often require two or more antihypertensive agents. In the Action to Control Cardiovascular Risk in Diabetes (ACCORD) trial, patients treated to an SBP of 119.3 mmHg required an average of 3.4 medications, while patients treated to an SBP $<133.3$ mmHg required an average of 2.1 medications. The elderly are more likely to have treatment-resistant hypertension due to physiological changes in the arterial vasculature that occur naturally with aging. Hypertension is more prominent in the elderly than in any other age group, with an estimated prevalence of 65% in men and 75% in women. Treating BP in the elderly has been associated with decreased incidence of various CV endpoints. BP naturally fluctuates and exhibits a diurnal variation over a 24-hour period. In the morning hours, CV events are more common due to a morning surge in BP. This increase in BP towards the end of the sleep period may be related to circadian upregulation of the RAS during the nighttime. Ambulatory BP monitoring (ABPM) can provide clinicians with additional information to diagnose hypertension, make more informed treatment decisions, and to gauge the effectiveness of antihypertensive therapy over 24 hours. BP control over a 24-hour dosing period has been demonstrated in several efficacy studies with OM/HCTZ.

Patients with T2DM: the BENIFICIARY trial

The BENIficar safety and effIcacy evaluation: an open-label, single-ARm, titration study in patients with hypertension and type 2 diabetes (BENIFICIARY) study assessed 24-hour BP control in patients initiated on OM 20 mg up-titrated to OM 40 mg, OM/HCTZ 40/12.5 mg, and OM/HCTZ 40/25 mg if BP was $\geq 120/70$ mmHg. ABPM was performed at baseline and at the end of week 12. The primary endpoint was the change from baseline in mean 24-hour ambulatory SBP at week 12. At 12 weeks, 24-hour ambulatory BP was reduced.
by 20.4/11.1 mmHg ($P < 0.0001$ vs baseline), and ambulatory BP targets of $<130/80$, $<125/75$, and $<120/80$ mmHg were achieved by 61.6%, 47.1%, and 39.0% of patients, respectively (Figure 3). Of special interest is that BP control was maintained during the last 6, 4, and 2 hours of the dosing interval when the normal morning rise in BP occurred. The SeBP reduction from baseline was 21.8/9.9 mmHg in patients titrated to OM/HCTZ 40/25 mg intensified, and 41.1% of patients achieved the cumulative guideline-recommended BP goal of $<130/80$ mmHg.

**Elderly patients: the BeniSILVER trial**

The Benicar Efficacy: New Investigation Shows Olmesartan Medoxomil Treatment Increasingly Leads Various Elderly Populations to Safe BP Reductions (BeniSILVER) study was a 12-week, open-label, multicenter trial. This study was conducted in 178 patients aged ≥65 years, and similar to BENIFICIARY, patients were initiated on OM 20 mg and up-titrated to OM/HCTZ 40/25 mg in a stepwise fashion if SeBP was ≥120/70 mmHg. The primary endpoint was the change in mean 24-hour ambulatory SBP from baseline to week 12. At study end, mean 24-hour ambulatory BP decreased by 25.7/12.3 mmHg ($P < 0.0001$ vs baseline) (Figure 4) from a mean baseline BP of 148.8/80.9 mmHg. After 12 weeks, the achievement of 24-hour ambulatory BP targets was also assessed in this study. Twenty-four hour ambulatory BP targets of $<130/80$, $<125/75$, and $<120/80$ mmHg were achieved by 73.3%, 56.7%, and 44.0% of patients, respectively. BP control was maintained throughout the 24-hour dosing interval with significant BP reductions from baseline observed during the last 6, 4, and 2 hours before re-dosing ($P < 0.0001$). A subgroup analysis in patients aged >75 years showed that 24-hour ambulatory BP targets of $<130/80$, $<125/75$, and $<120/80$ mmHg were achieved by 67.5%, 52.5%, and 40.0% of patients, respectively. Based on these results, a treatment algorithm using OM ± HCTZ appears to be effective in providing 24-hour BP control in a range of patients with hypertension, including those with T2DM and the elderly. There is currently no consensus on ambulatory BP goal values; however, the American Heart Association recommends normal 24-hour, daytime, and nighttime ambulatory BP values in adults of $<130/80$, $<135/85$, and $<120/70$ mmHg, respectively.

**Safety and tolerability of OM and HCTZ**

A fixed-dose combination of OM + HCTZ is associated with an overall adverse event (AE) rate that is similar to placebo, including when race or sex are considered. AEs
that occurred at a higher frequency than placebo in ≥2% of patients in pivotal trials include dizziness, upper respiratory tract infection, hyperuricemia, and nausea.43 Lending further support to placebo-like tolerability of OM + HCTZ are safety data reported in the BENIFORCE and BENIFICIARY trials. In BENIFORCE, the incidence of at least one AE across titration steps in the OM + HCTZ treatment arm ranged from 15.9% to 28.4% compared with 15.9%–26.2% during the placebo run-in period.22 Drug-related AEs ranged from 2.2% to 7.6% across titration steps in the OM + HCTZ treatment arm compared with 2.1%–9.5% in the placebo arm. Most adverse effects were mild to moderate in intensity, with dizziness being the most commonly reported AE at 3.4%.22 In the randomized double-blind period of the European study, treatment-emergent AEs (TEAEs) occurred in 11.8%–15.3% of patients across the treatment groups.24

In the BENIFICIARY study, where all patients had T2DM, the incidence of one or more TEAE was 13.5%–25.7% across all titrations steps, slightly lower than in BENIFORCE.29 Drug-related TEAEs ranged from 0.5% to 7.6% across the titration steps. The most commonly reported TEAE in BENIFICIARY was arthralgia and extremity pain at 2.1%.30 The occurrence of dizziness reported in BENIFICIARY was lower than in BENIFORCE at 0.7%.29

The treatment of hypertension in the elderly may result in relatively large BP reductions, especially in SBP. These large SBP reductions may be associated with dizziness and hypotension. In the BeniSILVER study, conducted in patients aged >65 years, 32.6% of patients reported an AE during the entire 12-week active treatment period, of which 11.8% were drug related.38 Incidences of drug-related dizziness and hypotension were 3.4% and 2.2%, respectively.

The use of HCTZ as monotherapy has been associated with hypokalemia, hyponatremia, hyperuricemia, and elevated blood glucose.44 In a study by Izzo et al, the maximum dose of OM/HCTZ 40/25 mg was not found to be associated with clinically significant decreases in sodium or potassium.45 Glucose and uric acid levels were found to be increased, with a mean uric acid level of 7.38 mg/dL (baseline value = 6.03 mg/dL) and mean glucose value of 109.0 mg/dL (baseline value = 103.9 mg/dL).45 These were within normal limits and were not clinically significant events associated with these laboratory elevations.

**Benefits beyond BP**

A number of ARBs have demonstrated the potential to provide benefits beyond their BP-lowering effects. In the Losartan Intervention for Endpoint Reduction (LIFE) trial, LOS monotherapy at 50 mg up-titrated to LOS 100 mg + HCTZ 25 mg over a period of 4 years resulted in a statistically significant decrease in the secondary endpoint of new-onset diabetes when compared with an atenolol + HCTZ regimen (13.0 vs 17.4 events/1000 patient-years; \( P = 0.001 \)).46 Beneficial effects of an ARB + HCTZ combination on the rate of new-onset
diabetes were also demonstrated in the Valsartan Antihypertensive Long-term Use Evaluation (VALUE) trial. Patients titrated to a maximum dose of valsartan 160 mg + HCTZ 25 mg had significantly fewer events of new-onset diabetes compared with an AML treatment regimen over a period of 4 years (32.1 vs 41.1 events/1000 patient-years; \( P < 0.0001 \)).57 New-onset diabetes was also a secondary endpoint in the VALUE trial. In the Randomized Olmesartan and Diabetes Microalbuminuria Prevention (ROADMAP) study, 4447 patients with T2DM were assigned to either OM 40 mg or placebo for a median of 3.2 years. Additional drugs (but not angiotensin-converting enzyme inhibitors or ARBs) were used as necessary to attain BP control \(<130/80 \text{ mmHg}\).48 The primary endpoint was the time to onset of microalbuminuria (MA). Overall, MA occurred in 8.2% of the OM group compared with 9.8% of the placebo group. The median time to onset of MA was 576 days for placebo compared with 722 days for OM (hazard ratio: 0.77; \( P = 0.01 \)), a risk reduction of 23%. Although no other study with ARBs has yielded similar results, ROADMAP provides evidence that pharmacological blockade with the ARB OM is highly effective in reducing the risk of developing MA and that the effect can be achieved through BP-dependent and BP-independent effects.49

**Rationale for combinations of HCTZ and OM**

Fixed-dose antihypertensive drug combination therapies that include a diuretic usually contain HCTZ rather than other agents such as chlorthalidone. Data reported by the Veterans Administration (VA) Cooperative study in 1967 is an early example whereby HCTZ demonstrated BP-reducing efficacy as well as reductions in CV events.50 In a cohort of high-risk male patients with DBP of 115–129 mmHg (\( N = 143 \)), HCTZ combined with reserpine and hydralazine reduced BP by an average of 43/30 mmHg after 24 months of treatment, and resulted in significantly reduced CV events compared with placebo (2 vs 27 total events; \( P < 0.001 \)). Three years later, the VA Cooperative Study reported data in a cohort of 380 male patients with lower risk diastolic hypertension (90–114 mmHg).51 An average reduction in BP of 27/17 mmHg was achieved after 4 months of combination therapy. The estimated 5-year risk of a morbid event was reduced with HCTZ-based treatment compared with placebo (18% vs 55%).

The preference for HCTZ over chlorthalidone may also be due to concerns about hypokalemia with chlorthalidone;52 however, hypokalemia is a class-wide effect for diuretics.53 The Multiple Risk Factor Intervention Trial (MRFIT) was a randomized primary prevention trial in 12,866 high-risk men with hypertension that compared a special intervention (SI) program (stepped-care combination therapy, smoking cessation counseling, and dietary advice) with usual care (UC) available within the community.54 Stepped-care therapy in the SI program began with a choice of HCTZ or chlorthalidone based on the preference at each treatment center. Reserpine, hydralazine, and guanethidine were able to be added on to the choice of diuretic, if required, to bring patients to the goal DBP of <90 mmHg. After 7 years of follow-up, a statistically nonsignificant difference of 7.1% in mortality from coronary heart disease (CHD) was observed in the SI care group compared with UC. Of interest, with regards to hypokalemia, was a finding in predefined subgroups that hypertensive men with baseline echocardiogram abnormalities had higher CHD mortality in the SI group compared with UC (36 vs 21 CHD deaths). This has led some to be concerned about the role that the choice of diuretics may have played in the increased mortality within this subgroup, particularly with regards to chlorthalidone.

In a study of 233 hypertensive men, Siegel et al sought to determine the potassium-wasting effects of HCTZ at 50 mg/day, with and without potassium supplementation, or triamterene against chlorthalidone 50 mg and placebo.55 After 2 months of treatment, serum potassium levels were decreased to <3.5 mmol/L (threshold of hypokalemia) in 15% of patients treated with HCTZ 50 mg without potassium supplementation vs 33% of patients treated with chlorthalidone 50 mg \( (P < 0.01) \).55 Severe hypokalemia, defined as serum potassium levels <3.0 mmol/L, occurred in 10% of patients taking HCTZ without supplementation and 20% of patients taking chlorthalidone, which was not a statistically significant difference.55 However, the dosage of HCTZ that was used in the study was greater than the maximum dosage (25 mg) used in single-pill combination formulations.

Pharmacokinetic considerations also inform the rationale for using HCTZ over chlorthalidone. Chlorthalidone has an estimated half-life ranging from 40 to 72 hours,52 while HCTZ has a half-life ranging from 6 to 15 hours.44 In select patient populations with renal impairment, avoiding medications with long half-lives may help to reduce the likelihood of AEs. Drug labels for both HCTZ and chlorthalidone advise against administering to patients with renal impairment, and neither diuretic appears in the Beers criteria for inappropriate medication use in the elderly.56 The shorter half-life of HCTZ could potentially be a concern with regards to 24-hour BP control. However, the antihypertensive efficacy of OM/HCTZ combination therapy has been shown to be maintained throughout the 24-hour dosing interval in a variety of patient subgroups.57
Data from the ALLHAT study suggest that chlorthalidone may increase the incidence of new-onset diabetes. When compared with AML and lisinopril, chlorthalidone-treated patients had 43% and 65% higher incidences of new-onset diabetes, respectively. While there were no differences between chlorthalidone and AML or lisinopril in CV outcomes, the trial duration would not have been long enough to account for CV outcomes in patients with new-onset diabetes, as CV effects would not become manifest in the short timeframe of the study. Head-to-head outcomes studies between chlorthalidone and HCTZ have not been conducted, and thus it remains to be seen whether HCTZ would have had similar increases in new-onset diabetes.

An ABPM substudy of the Avoiding Cardiovascular Events through Combination Therapy in Patients Living with Systolic Hypertension (ACCOMPLISH) trial was recently conducted to identify any differences in 24-hour BP control between BEN plus AML and BEN plus HCTZ after 2 years of treatment. Mean 24-hour, daytime, and nighttime BP values were not significantly different between the two treatment groups (Figure 5); BP control rates were >80% in both groups. This indicates that 24-hour BP control is similar between the two treatment groups and supports the original conclusions of the ACCOMPLISH investigators that the improvement in CV outcomes seen in the AML-based regimen is most likely due to other putative cardioprotective properties of combining a RAS blocker with AML. AML has a terminal elimination half-life ranging from 30 to 50 hours, very similar to chlorthalidone; however, the longer half-life conferred no additional benefit over HCTZ with regards to 24-hour BP control.

The overall preference of HCTZ over chlorthalidone may simply be due to the availability of HCTZ as a component of fixed-dose, single-pill combinations. There are currently no fixed-dose ARB + chlorthalidone single-pill combination products available.

In view of the recent clinical evidence that demonstrates the efficacy and safety of treatment regimens based on OM + HCTZ, there is no reason that HCTZ should not remain as a preferred treatment option for use in combination with ARBs such as OM. However, there remains an unmet need for head-to-head outcomes studies that compare the relative efficacy and tolerability of HCTZ and chlorthalidone in order to provide evidence for informing clinical guidelines.

**Conclusion**

ARBs provide excellent efficacy and tolerability and are frequently used as first-line therapy, alone or in combination with diuretics. The combination of OM/HCTZ has been shown to be an effective and well tolerated treatment option.
that provides BP-lowering efficacy and improvements in BP control in patients with hypertension. BP reduction achieved through combination therapy has been associated with improvements in CV morbidity and mortality. The clinical evidence discussed in this review provides a rationale for the use of OMTZ combination therapy as an antihypertensive treatment strategy, regardless of patient age, sex, or race, or patients with common comorbidities such as diabetes.

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