Blocking the RAAS at different levels: an update on the use of the direct renin inhibitors alone and in combination

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Abstract: The renin–angiotensin–aldosterone system (RAAS), an important regulator of blood pressure and mediator of hypertension-related complications, is a prime target for cardiovascular drug therapy. Angiotensin-converting enzyme inhibitors (ACEIs) were the first drugs to be used to block the RAAS. Angiotensin II receptor blockers (ARBs) have also been shown to be equally effective for treatment. Although these drugs are highly effective and are widely used in the management of hypertension, current treatment regimens with ACEIs and ARBs are unable to completely suppress the RAAS. Combinations of ACEIs and ARBs have been shown to be superior than to either agent alone for some, but certainly not all, composite cardiovascular and kidney outcomes, but dual RAAS blockade with the combination of an ACEI and an ARB is sometimes associated with an increase in the risk for adverse events, primarily hyperkalemia and worsening renal function. The recent introduction of the direct renin inhibitor, aliskiren, has made available new combination strategies to obtain a more complete blockade of the RAAS with fewer adverse events. Renin system blockade with aliskiren and another RAAS agent has been, and still is, the subject of many large-scale clinical trials and furthermore, is already available in some countries as a fixed combination.

Keywords: angiotensin II receptor blockers, renin–angiotensin–aldosterone system, hypertension, angiotensin-converting enzyme inhibitors

Introduction: the blockade of the renin–angiotensin–aldosterone system

In the last decades, renin–angiotensin–aldosterone system (RAAS) has become, unquestionably, a key point in the pathophysiology of cardiovascular and kidney diseases.

Activation of the RAAS occurs following the release of renin by the kidney. Renin is a protease enzyme that catalyzes cleavage of angiotensinogen to form angiotensin I (Ang I). Ang I is itself inactive but is converted to the biologically active peptide angiotensin II (Ang II) by angiotensin-converting enzyme (ACE), which is produced in the lungs and cleaves the Ang I molecule.

Ang II binds with the type 1 Ang II receptors (AT1) in the smooth muscle cells of the peripheral blood vessels causing vasoconstriction, and hence increased peripheral vascular resistance and increased blood pressure (BP; Figure 1). Ang II is recognized to be a major contributor to the progression of cardiovascular and kidney diseases.1–4

Activation of AT1 receptors by Ang II also stimulates release of the mineralocorticoid, aldosterone, from the adrenal gland, situated superior to the kidney. Aldosterone promotes sodium and water retention along the nephron, further increasing the BP.
Plasma renin activity (PRA) is a measure of the level of activity in the RAAS, and consequently, it can be used as an indicator to detect dysregulation of the RAAS. As a matter of fact, PRA reflects the capacity of circulating renin to cleave angiotensinogen to form Ang I, and thus provides a measure of the level of activity in the RAAS.

The RAAS is targeted at different places by the existing antihypertensive therapies. Angiotensin-converting enzyme inhibitors (ACEIs) reduce the production of Ang II by inhibiting the conversion of Ang I to Ang II by these enzymes. Angiotensin receptor blockers (ARBs) antagonize AT1 receptors and prevent Ang II from binding. However, ACEIs and ARBs are associated with a feedback loop that increases PRA, as loss of stimulation of AT1 receptors on juxtaglomerular cells in the kidney (ie, removal of negative feedback) leads to a compensatory increase in renin release. The resulting increase in PRA may limit the organ protection offered by these drugs (Figure 1).

The whole RAAS is, therefore, upregulated although AT1 receptor-mediated effects of the effector molecule Ang II are blocked. Direct renin inhibitors (DRI) target the RAAS at its point of activation, resulting in the reduction of PRA (Figure 2).

As DRI reduces PRA, the production of Ang I decreases, resulting in less substrate availability for conversion to Ang II by ACE or other enzymes. In doing so, direct renin inhibition produces effective overall RAAS suppression.

The functional renin receptor plays a pivotal role in Ang II production and cellular responses to renin. The binding of renin to its receptor triggers intracellular signals by rapid activation of mitogen-activated protein kinases, ie, extracellular signal-regulated protein kinases 1 and 2, and may increase plasma activator inhibitor-1 – a prothrombotic agent – levels via this mechanism. Furthermore, the bound renin has increased catalytic efficiency for the cleavage of angiotensinogen to form Ang I, facilitating Ang II production (by ACE or vascular muscle smooth chymase) at the cell surface. This may have consequences in terms of organ damage and progression of cardiovascular disease.

It is now clear that the blockade of the RAAS not only reduces the BP but also decreases the risk of cardiovascular events and slows the progression of renal disease in different types of patients. Nevertheless, some patients do not respond adequately to the conventional renin–angiotensin system blockade, raising a clinical problem that qualifies the patient still at high risk for cardiovascular or renal diseases; in a significant number of patients, in fact, even if the target of lowering BP is obtained, the RAAS blockade with an ACE inhibitor or an ARB does not reduce the risk
of experiencing cardiovascular events and chronic heart and kidney disease progress despite the therapy.\textsuperscript{9,12–16} Incomplete blockade of the RAAS at recommended doses may be one explanation for this observation.\textsuperscript{17,18} Not withstanding the evolution from ACE inhibitors to ARBs, the RAAS will not be fully blocked using just these molecules alone.\textsuperscript{17,19,20} Accordingly, one strategy to improve the efficacy of RAAS blockade is to combine ACEIs and ARBs. ACEIs and ARBs combination has been widely studied in patients with hypertension, nephropathy, diabetes, myocardial infarction (MI), and heart failure.\textsuperscript{21}

This review intends to examine the rationale for RAAS blockade at different levels using combinations of different molecules acting on this system; we will consider at first the characteristics of ACEI/ARBs combinations, following a pathway that will bring us to the actual use of DRIs alone or in fixed combination with ARBs.

**ACEI/ARB combination therapy**

Although the theoretical benefits of combined ACEI-based and ARB-based therapy may enhance BP control and improve clinical outcomes in some subpopulations of patients, in 2006, Weber and Giles\textsuperscript{22} already observed that, for the most patients, these RAAS-based therapies do not reduce the risk for major cardiovascular events to the expected extent based on the vital role of the RAAS in the pathophysiology of hypertension and cardiorenal disease. They speculated that this may indicate that the RAAS is not as widely dysregulated in hypertension as previously believed, or that the current strategies and/or agents available for RAAS suppression are not as effective as they could or should be. With respect to the latter argument, dual RAAS blockade at different sites in the RAAS pathway theoretically would have provided additive protective effects by further reducing systemic and local levels of some or all angiotensin peptides, which could further inhibit formation of the Ang II.\textsuperscript{23–26} On the same topic, combinations of ACEIs and ARBs have generally been shown to produce small but significant additional reductions in BP compared with either monotherapy;\textsuperscript{27,28} although several studies have shown that ACEI/ARB combinations result in modest improvement in BP and proteinuria compared with treatment with only one of these drug classes,\textsuperscript{2,13,29–31} others do not support this combination because of concerns about efficacy (modest improvement in BP with limited or no additional benefit on outcomes) and safety.\textsuperscript{32} For instance, in a crossover study of 64 hypertensive patients not achieving BP control with full-dose valsartan, the addition of amlopidine reduced ambulatory 24-hour BP to a greater extent than the addition of benazepril (−15.2/−9.9 mmHg vs −8.6/−6.3 mmHg, respectively).\textsuperscript{33} Similarly, ACEI/ARB combination therapy was less effective for lowering

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**Figure 2** Direct renin inhibition acts at the point of activation of the renin system and neutralizes the PRA increase.

**Abbreviations:** ACE, angiotensin-converting enzyme; PRA, plasma renin activity; Ang I, angiotensin I; Ang II, angiotensin II; AT\textsubscript{1}, type I Ang II receptor.
BP than ARB/thiazide diuretic combination therapy in a study of 327 hypertensive patients uncontrolled by ARB monotherapy and in a study of 88 African Americans with hypertension.

The Valsartan in Acute Myocardial Infarction (VALIANT) study of 14,703 elderly patients with left ventricular systolic dysfunction, congestive heart failure (CHF), or both after MI reported similar rates of all-cause mortality, death from cardiovascular events, recurrent MI, and hospitalization for heart failure for all three of its treatment groups (ACEI, ARB, and ACEI/ARB), accompanied by significantly ($P < 0.05$) more adverse events in the combination therapy group.

Two meta-analyses of patients with CHF or left ventricular dysfunction (LVD; including CHARM-Added, Val-HeFT, and VALIANT) yet showed that ACEI/ARB combination therapy significantly increases the risk for adverse events (eg, hypertension, worsening renal function, and hyperkalemia), inducing treatment discontinuation. On the other hand, in the Randomized Evaluation of Strategies for Left Ventricular Dysfunction pilot study, ACEI/ARB combination therapy, compared with monotherapy, significantly limited the increases in end-diastolic and end-systolic volumes ($P < 0.01$) and reduced brain natriuretic peptide, a biomarker of heart failure.

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However, some recent large trials have failed to find better cardiovascular outcomes with the ACEI/ARB combination despite better BP reductions.

The Valsartan Heart Failure Trial determined whether valsartan could further reduce morbidity and mortality in patients with heart failure, who already receiving optimal therapy (including ACEIs in 93% of patients and β-blockers in 35% of patients). The primary end point of mortality was similar for the valsartan and placebo groups, whereas the combined primary end point of morbidity and mortality was significantly reduced ($P = 0.009$) in patients receiving valsartan plus optimal therapy compared with the placebo group. This benefit was primarily due to a 24% reduction in hospitalizations for heart failure in valsartan-treated patients. A subgroup analysis of patients on different background therapies revealed that valsartan had a favorable effect on the combined primary end point in those receiving an ACEI ($P = 0.002$), a β-blocker ($P = 0.037$), or no background therapy ($P = 0.003$). In contrast, in patients receiving both an ACEI and a β-blocker, valsartan had an adverse effect on mortality ($P = 0.009$), suggesting that this particular approach to comprehensive blockade of neurohormone systems in heart failure may be detrimental.

In the Ongoing Telmisartan Alone and in Combination With Ramipril Global Endpoint Trial, combination therapy with telmisartan plus ramipril produced no greater reduction in the primary end point of death from cardiovascular events, MI, stroke, or hospitalization for heart failure than either component monotherapy in high-risk patients with cardiovascular disease or diabetes but without heart failure. Combination therapy was associated with an increased risk of hypotension ($P < 0.001$), syncope ($P = 0.03$), hyperkalemia ($P < 0.001$), and acute renal impairment ($P < 0.001$). The reasons for the lack of additional benefits with combination therapy, despite an additional reduction in systolic BP of 3.4 mmHg, compared with ACEI monotherapy are unknown. As the investigators pointed out, the majority of patients were also receiving statins, β-blockers, and antiplatelet medications so that additional RAAS blockade with the ACEI/ARB combination therapy resulted in little additional clinical benefit compared with the ACEI therapy alone.

Although it is clear that monotherapy with ACE inhibitors or ARBs is effective in reducing cardiovascular mortality and morbidity in patients with heart failure, the reasons for the different cardiovascular outcomes in trials examining ACEI/ARB combinations may relate to different patient populations, previous or concurrent successful treatment with other drugs, or study design. As noted by Arici and Erdem, many clinical studies have been small and of short duration, and most used submaximal doses of ACEIs and ARBs both alone and in combination. Most combination studies were not designed to maximize BP control and in fact, achieved only modest improvement in BP (∼3–4 mmHg) over monotherapy with an ACEI or ARB. In addition, many early studies used once-daily dosing with short-acting ACEIs. Therefore, it is possible that low ACEI concentrations at trough in combination studies using short-acting ACEIs could have increased the likelihood of both acute (method related) and chronic (mechanistic mediated) ACE escape. Administration of diuretics also has resulted in increases in PRA, and the use of diuretics as concurrent...
medications usually is permitted in ACEI and ARB studies. Increases in both PRA and ACE escape have been associated with adverse clinical outcomes in patients on ACEI or ARB therapy. For example, in a study of 70 patients with CHF, elevated PRA despite 6 months of treatment with an ACEI was an independent predictor of elevated Ang II levels \( (P = 0.0004) \), and elevated plasma Ang II levels were an independent predictor of death or worsening CHF \( (P = 0.002) \). In another study, 699 patients with CHF underwent a complete clinical and biochemical workup at baseline and were monitored for a median of 23 months; 81% of them were receiving an ACEI or ARB. Elevated baseline PRA was an independent predictor of death or the need for cardioversion in patients with implantable cardioverter devices \( (P < 0.001) \), and PRA was higher in patients on RAAS inhibitors relative to those not receiving RAAS inhibitors \( (P = 0.017) \). In Val-HeFT, analysis of 4,300 patients with CHF, who had neurohormonal measurements, showed that increased baseline PRA was an independent predictor of all-cause mortality \( (P = 0.011) \) and of combined mortality and morbidity \( (P = 0.0025) \). This was observed despite the fact that the majority of patients were receiving treatment with an ACEI and approximately 50% were also receiving treatment with an ARB. A recent post hoc analysis of Val-HeFT data showed that PRA was 3.7 times higher in patients receiving ACEIs (vs those not receiving them), and that higher PRA at baseline was associated with greater mortality among patients receiving ACEIs \( (P = 0.0005) \).

All together, these findings strongly indicate that PRA is related to adverse clinical outcomes even for patients receiving treatment with ACEIs and/or ARBs and further raises the possibility that DRIs may be useful alone or in combination with ACEIs or ARBs for reducing the risk for these outcomes.

**Direct renin inhibitor**

**Pharmacokinetic and pharmacodynamic**

Aliskiren, an octanamide, is the first representative of a new class of nonpeptide, low molecular weight, orally effective transition-state renin inhibitors. It is a DRI that has a high binding affinity for renin, and this may be explained by a number of interactions with the enzyme’s active site. In healthy humans, aliskiren of doses between 40 and 640 mg exerts a dose-dependent reduction in PRA, Ang I, and Ang II levels. The molecule is superior in reducing PRA compared with ARBs. Again, aliskiren at a dose of 300 mg decreases PRA in hypertensive patients by approximately 50%–80% and reduces PRA and plasma levels of Ang I and Ang II for 48 hours. Furthermore, urinary aldosterone was reduced at a dose of 80 mg or more, and sodium extraction was increased to 91% at a dose of 640 mg. Compared with valsartan, aliskiren more strongly decreases the activity of renin in the circulation and reduces the excretion of urinary aldosterone for a longer period. Following oral administration, peak plasma concentrations of aliskiren are reached within 1–3 hours, and the plasma half-life of aliskiren in humans shows a slow terminal elimination at 23–70 hours and approximately 47%–51% of aliskiren is bound by plasma proteins in humans, independent of the concentration. Based on *in vitro* studies, the major enzyme responsible for its metabolism appears to be Cytochrome P450 (CYP3A4). Aliskiren does not inhibit the CYP450 isoenzymes (CYP1A2, CYP2C8, CYP2C19, CYP2D6, CYP2E1, and CYP3A), and the main elimination route of aliskiren is via feces in its unmetabolized form. Approximately one-fourth of the absorbed dose also appears in the urine as unchanged compound; the pharmacokinetic and pharmacodynamic differences of aliskiren between Caucasians and Japanese are minimal and no clinically important pharmacokinetic differences were observed between patients with type 2 diabetes and normal population: the half-life of this drug was 40 hours and 44 hours in healthy subjects and patients with diabetes, respectively.

**Clinical features**

Aliskiren is well tolerated by all age groups, including the very elderly, and there are no indications to change the recommended dose of aliskiren in patients with hepatic and renal insufficiency because the peak concentration, area under the curve (AUC), and half-life were only slightly greater in patients with hepatic dysfunction. Aliskiren exposure was also increased slightly in patients with renal function impairment, but these changes did not correlate with creatinine clearance. All agents that inhibit the RAAS activate the negative feedback loop that leads to a compensatory increase in plasma renin concentration. When this increase occurs during treatment with ACEIs and ARBs, the result is increased levels of PRA but during treatment with aliskiren, the effect of increased renin levels is blocked, so the levels of Ang I and Ang II, as well as PRA, are all reduced. The clinical trials do not report any major adverse effects of aliskiren compared with ARBs or...
ACEs being generally well tolerated by all patients. The most common adverse events of aliskiren are diarrhea, headache, nasopharyngitis, dizziness, fatigue, back pain, gastrointestinal disorders, rash, and renal stones.\textsuperscript{52,53,62,65} Because aliskiren directly inhibits the RAAS, theoretically adverse events such as coughing and angioedema may also occur; as a matter of fact, few cases of edema involving the face, lips, tongue, hands, and whole body have been reported to involve more than 1% of patients treated with aliskiren, which also occur at a similar or greater extent in patients receiving placebo.\textsuperscript{52,62,65} Aliskiren has the same contraindications as ACEIs and ARBs, including hypersensitivity reactions to the molecule, pregnancy, and bilateral renal-artery stenosis.

Nussberger et al\textsuperscript{56} first reported the clinical effects of aliskiren compared with enalapril and placebo. Subjects were randomized into three 8-day treatments with enalapril 20 mg/d or placebo, aliskiren 40 or 160 mg/d, and aliskiren 80 or 640 mg/d.

Aliskiren produced dose-dependent reductions in PRA, Ang I, and Ang II vs placebo. Inhibition of Ang II production on day 8 was superior to that of placebo with aliskiren 160 mg, aliskiren 640 mg, and enalapril. Reductions in plasma and urine aldosterone concentrations were similar between enalapril and aliskiren doses greater than 40 mg/d. All subjects were normotensive upon enrollment, and no changes in BP or heart rate were observed. Adverse events were similar among all treatment groups, including enalapril and placebo. Orthostasis and dizziness following blood draws were reported in all treatment groups, with a tendency toward increased dizziness with aliskiren 640 mg (actual incidence not reported).

Stanton and colleagues\textsuperscript{66} performed a randomized, double-blind, parallel-group study in 226 patients aged 21–70 years with mild to moderate essential hypertension comparing the 4-week BP-lowering effects and safety of aliskiren 37.5, 75, 150, and 300 mg/d with those of losartan 100 mg/d and showed dose-dependent decreases in daytime ambulatory systolic BP and PRA in patients treated with aliskiren. There was no difference in BP lowering of daytime ambulatory systolic BP between aliskiren 75, 150, and 300 mg/d and losartan 100 mg/d. The BP-lowering activity of the 300 mg dose persisted for 24 hours. With aliskiren, PRA fell dose-dependently by 50%–80%, whereas it doubled with losartan. The authors concluded that aliskiren, through inhibition of renin, was an effective and safe orally active BP-lowering drug.\textsuperscript{66}

In a double-blind, randomized parallel-group trial, Gradman and colleagues\textsuperscript{67} assigned 652 hypertensive patients once-daily doses of 150 mg, 300 mg, or 600 mg aliskiren, 150 mg irbesartan, or placebo. The reductions in BP (systolic/diastolic) measured at trough after 8 weeks of follow-up and adjusted for baseline values and the difference from placebo averaged 6.1/9, 1.5/4, 1.4/2, and 7.2/2.5 mmHg on the increasing doses of aliskiren and irbesartan, respectively. The 2 highest aliskiren doses lowered diastolic BP significantly more than irbesartan ($P = 0.01$). The frequency of adverse effects was similar for placebo and aliskiren with the exception of diarrhea associated with the dose 600 mg/d (1.5% vs 6.9%).

**Dual blockade DRI/ARB**

Because aliskiren is more likely, at least initially, to be added into existing therapy than to be used for newly diagnosed patients, much of the phase 3 clinical studies have investigated its use as add-on therapy to the major classes. Indeed, aliskiren has been studied both in comparison to and in combination with each of the RAAS-blocking drugs, ACEIs, and ARBs.

An early study in healthy human volunteers explored the effect of aliskiren in combination with an ARB on components of the RAAS, comparing the effects of aliskiren, valsartan, and their combination on PRA, Ang II, and aldosterone.\textsuperscript{53} A single dose of 300 mg aliskiren inhibited PRA by 95.6% compared with baseline at 4 hours, and it remained below placebo for 48 hours ($P < 0.05$). Similarly, Ang II decreased and remained below placebo for 48 hours ($P < 0.05$). In contrast, 160 mg valsartan increased PRA and Ang II levels within 4 and 6 hours of dosing, respectively, and they were still higher than placebo at 24 and 48 hours ($P < 0.05$). The combination of 150 mg aliskiren and 80 mg valsartan decreased PRA below placebo values at 4 hours ($P < 0.05$), with a subsequent return to baseline by 12 hours. All the three treatments significantly ($P < 0.05$) decreased plasma levels and urinary excretion of aldosterone for at least 4 hours after dosing compared with placebo. Aliskiren alone or in combination with valsartan reduced 24-hour cumulative urinary aldosterone excretion to a greater extent ($P < 0.05$) than did valsartan alone. Overall, in this study, the low-dose combination of aliskiren and valsartan produced enhanced blockade of the components of the RAAS that were equal to those produced by the higher dose of aliskiren and higher to those produced by the standard dose of valsartan.

The safety and tolerability of dual RAAS blockade with aliskiren for treatment of heart failure was examined in the Aliskiren Observation of Heart Failure Treatment study
(ALOFT). In this study, patients (n = 302) were randomized to receive aliskiren (150 mg daily) or placebo for 3 months in addition to standard therapy. A 98% of the placebo group and 99% of the aliskiren group were also receiving an ACEI or an ARB. In patients receiving aliskiren, the incidences of renal dysfunction, hypotension, and hyperkalemia were similar to placebo. In this study, N-terminal prohormone brain natriuretic peptide, an important biomarker of heart failure prognosis and the primary efficacy outcome measure, increased with placebo but decreased with aliskiren (P = 0.01). Serum levels of the related biomarker brain natriuretic peptide decreased with both placebo and aliskiren, ie, to a significantly greater extent with aliskiren than with placebo (P = 0.02). Aliskiren also caused a greater reduction in PRA (P < 0.0001) and 24-hour urinary aldosterone excretion (P = 0.02) compared with placebo (Table 1).

In the Aliskiren Left Ventricular Assessment of Hypertrophy (ALLAY) study, in which patients with hypertension and left-ventricular hypertrophy (LVH) were randomized to aliskiren (300 mg), losartan (100 mg), or their combination, aliskiren was noninferior to losartan in reducing LV mass index, even if the combination was not superior to losartan alone. These findings suggest that aliskiren is an effective and well-tolerated treatment option for patients with LVH. The lack of additional benefit with combination therapy may be related to the design of the study in which there was no significant difference in BP reduction between groups. LV mass reduction seems to be directly related to BP reduction. Tolerability and safety were similar in all treatment groups.

In some recent important trials, aliskiren has also shown renoprotective potential in patients with type 2 diabetes and albuminuria, the best available surrogate parameter in the treatment of diabetic nephropathy, associated with risk of renal and cardiovascular events.

In the Aliskiren in the Evaluation of Proteinuria in Diabetes (AVOID) trial, Parving et al evaluated the effects of dual blockade of the RAAS with aliskiren and losartan in patients (n = 599) with hypertension and type 2 diabetes with nephropathy. Patients were maintained on losartan (100 mg daily) for the duration of the study and were randomized to receive a 6-month treatment with aliskiren (150 mg/d for 3 months, then 300 mg/d for 3 months) or placebo. After 3 months of treatment with aliskiren (150 mg/d), the urinary albumin/creatinine ratio (UACR) had decreased by 11% compared with placebo (P = 0.02). Increasing the dose of aliskiren to 300 mg/d caused a further decrease in the UACR to 20% (P < 0.001). A nonsignificant difference in BP was seen between the treatment groups, suggesting that the renoprotective effect of aliskiren was independent of

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**Table 1 Aliskiren and ACEIs or ARBs combination therapy**

<table>
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<tr>
<th>Patients</th>
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<th>Duration</th>
<th>Treatment (mg/d)</th>
<th>Results</th>
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<td>Heart failure</td>
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<td>3 mo</td>
<td>Aliskiren (150) or placebo</td>
<td>Aliskiren decreased plasma NT-proBNP and BNP</td>
<td>McMurray et al⁶⁸ The ALOFT Study</td>
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<td>Left-ventricular mass reduced to a similar extent in all treatment groups</td>
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<td>Losartan/Aliskiren (100/300)</td>
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¹Change in mean sitting systolic/mean sitting diastolic blood pressure (mmHg).

**Abbreviations:** ACEIs, angiotensin-converting enzyme inhibitors; ARBs, angiotensin receptor blockers; UACR, urinary albumin/creatinine ratio; NYHA, New York Heart Association; NT, N-terminal; BNP, brain natriuretic peptide.
BP. There was no difference in the rates of adverse events, such as hyperkalemia, or discontinuation rates between the 2 groups.

In a subsequent double-blind, randomized, crossover study, 26 patients with type 2 diabetes, hypertension, and albuminuria (>100 mg/d) were randomly assigned to four 2-month treatment periods in random order with placebo, 300 mg aliskiren once daily, 300 mg irbesartan once daily, or the combination of two drugs with identical doses. Persson and colleagues yet demonstrated that with the combination of 300 mg aliskiren and 300 mg irbesartan, the reduction in albuminuria was enhanced compared with the reduction obtained with the 2 monotherapies. The added antiproteinuric effect with combination treatment compared with aliskiren alone was 31%.

All these studies suggest that aliskiren plus an ARB may provide cardiorenal benefits that go beyond those solely attributable to lowering of BP. The DRI approach to RAAS suppression is still very recent and the ultimate role of aliskiren in combination therapies with ACEIs, ARBs, and other antihypertensives will be better defined through future studies, which are being conducted as part of the ASPIRE HIGHER clinical program.

On the other hand, the dual blockade in post-MI seems not be as useful as in the previous reported conditions. As a matter of fact, the results of the recently presented study ASPIRE77 (designed to test the hypothesis that aliskiren on top of standard therapy would improve remodeling in post-MI patients with LV dysfunction) have shown the following:

- The primary endpoint (LVESV) was nonsignificant; however, consistent trends toward smaller ventricles were seen in the aliskiren group and early post hoc analyses suggest a potential benefit in post-MI patients with diabetes.
- Secondary composite endpoints were not different, with no statistical difference in any of the components.
- There was a nonsignificant trend toward higher mortality in the aliskiren group, although total mortality (3%) is low compared with the other post-MI trials.
- Hyperkalemia, hypotension, and renal dysfunction were increased in the aliskiren group overall.

Summarizing, based on echocardiographic markers, the ASPIRE study does not support the addition of aliskiren to an ACEI or an ARB in post-MI patients with impaired left ventricular function. Although early post hoc analyses may point toward a beneficial effect in favor of aliskiren in some subgroups (eg, patients with diabetes), ASPIRE as a hypothesis-generating trial does not support a morbidity and mortality study in post-MI patients.

Conclusion

Even though awareness and treatment of hypertension have increased over the years, substantial improvements in BP control rates are still lacking, with approximately two-thirds of hypertensive adults aged 35–64 years failing to reach a BP target of <140/90 mmHg. Nowadays, it is well known that the majority of hypertensive patients require 2 or more agents to reach BP goal. Not only epidemiological or observational researches but also important and wide clinical trials, including Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial (ALLHAT), Avoiding Cardiovascular Events through Combination Therapy in Patients Living with Systolic Hypertension (ACCOMPLISH), The International Verapamil SR/Trandolapril Study (INVEST), and The Losartan Intervention For Endpoint reduction (LIFE), have reported that 23%–54% patients require 3 or more antihypertensive agents for BP control and target-level maintenance (<140/90 or <130/80 mmHg depending on cardiovascular risk). In patients with BP ≥20/10 mmHg above goal, on average, 3.2 drugs might be needed to achieve BP control.

The problem of the comorbidity and its consequence – the coadministration of multiple drugs – may be substantial for elderly patients. In a recent trial conducted by Yarrows and colleagues, the aliskiren/valsartan combination therapy had been studied in elderly patients too compared with those younger than 65 years of age. The combination provided significantly greater reductions in SBP systolic BP and BP control rates than did valsartan monotherapy in the elderly patients, and the effect of aliskiren/valsartan and high-dose valsartan on pulse pressure was greater in elderly patients than in younger patients. Aliskiren monotherapy and aliskiren/valsartan combinations were equally well tolerated in the elderly as in younger patients.

Following the above-reported data and several clinical observations, in 2009, the US Food and Drug Administration approved the fixed combination of aliskiren/valsartan at the dosages of 150/160 mg and 300/320 mg for the treatment of hypertension in patients not adequately controlled on aliskiren or ARB monotherapy and as initial therapy in patients likely to need multiple drugs to achieve their BP goals.

Disclosure

Dr Destro has received consulting and lecture fees from Novartis Pharma, Dr Destro has received Boehringer
Ingelheim, AstraZeneca, Servier, Menarini IFR, Schering-Plough, Guidotti, Pfizer, Knoll, Bayer, Chiesi, Sankyo, Merck Sharp and Dohme, Malesci and Errekapppa; he has received research support as a study investigator from Novartis Pharma, Boehringer Ingelheim, Menarini IFR, Guidotti, Bayer and Chiesi. Dr Preti received consulting and lecture fees and research support as study investigator from Novartis Pharma. Dr Cagnoni and Dr Rossi Ricci received research support as study coinvestigators from Novartis Pharma. No other potential conflict of interest relevant to this article was reported.

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