DIRECT RENIN INHIBITORS IN ARTERIAL HYPERTENSION TREATMENT

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Abstract. High blood pressure was the leading cause of death and disability-adjusted life years worldwide. Despite progress in treatment of hypertension, a number of people with uncontrolled or resistant hypertension increases. Hypertensive disorders are strongly linked with an overactive renin-angiotensin aldosterone system. The renin-angiotensin-aldosterone system has been a highly successful pharmacologic target, as the system is strongly implicated in the development of hypertension-related target organ damage. Renin-angiotensin aldosterone system renin inhibitors are the first of a new class of completely non-peptide, low-molecular-weight, orally active transition-state renin inhibitors and only approved for the treatment of hypertension.

Keywords: hypertension, renin-angiotensin aldosterone system, renin inhibition.

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Introduction

One of the key risk factors for cardiovascular disease (CVD) is hypertension (HT)—or raised blood pressure (BP), defined as SBP > 140 mm Hg or DBP > 90 mm Hg (> 130 mm Hg or > 80 mm Hg by High Blood Pressure Clinical Practice Guideline (ACC/AHA, 2017). High BP was the leading cause of death and disability-adjusted life years worldwide. Hypertension is a global public health issue, already affects one billion people worldwide, leading to heart attacks, strokes, kidney failure and premature mortality and disability. In the United States, hypertension (ACC/AHA, 2017) accounted for more CVD deaths than any other modifiable CVD risk factor and was second only to cigarette smoking as a preventable cause of death for any reason. Researchers have estimated that raised blood pressure currently kills nine million people every year (WHO, 2013).

The prevalence of hypertension appears to be around 30-45%in adults over 25 years old of the general population, with a steep increase with ageing. There also appear to be noticeable differences in the average BP levels across countries. Western European countries exhibit a downward trend, in contrast to eastern European countries, which show a clear-cut increase in death rates from stroke (one of the most important hypertension outcomes) (ESH/ESC Guidelines, 2013). The increasing prevalence of hypertension is attributed to population growth, ageing and behavioural risk factors, such as unhealthy diet, harmful use of
alcohol, lack of physical activity, excess weight and exposure to persistent stress. In addition, there are several metabolic factors including diabetes and high cholesterol, that increase the risk of complications of hypertension (WHO, 2013).

Evidence favouring the administration of BP-lowering drugs to reduce the risk of major clinical cardiovascular (CV) outcomes (fatal and non-fatal stroke, myocardial infarction, heart failure and other CV deaths) in hypertensive individuals results from a number of randomized controlled trials (RCTs) - mostly placebo-controlled (ESH/ESC Guidelines, 2013).

According to current ESH (European Society of Hypertension)/ESC (European Society of Cardiology) guidelines, physician have at their disposal five main groups of antihypertensives (as also referred to as basic antihypertensives) for use in either monotherapy or in combination: diuretics, beta-blockers (BBs), calcium ion channel blockers (CCBs), angiotensin-converting enzyme inhibitors (ACEIs) and angiotensin II type I (AT₁) receptor blockers (ARBs) – sartans (Vaclavik, Sliva, 2014).

Despite progress in treatment of HT, a number of people with uncontrolled or resistant HT increases. There is a problem of inefficiency of therapy or lack of patients' adherence to treatment. Possibly, poor compliance on the part of the patients is responsible. Another possibility is that awareness of prevention strategies and their implementation by primary physicians may not always be optimal. Finally, some lay the blame at the feet of the health care system or third-party payers. Undoubtedly, all of these factors play a role. However, an additional explanation might be that we need better drugs for patients with essential hypertension. Therefore, search for new approaches to treatment of HT continues (Popov, Bulanov, Ivanov, 2012). In any given patient, several concomitant pharmacologic targets must be addressed. Indeed, patients who require four or more classes of drugs to achieve the current treatment guidelines are no rarity (Müller, Luft, 2006).

**Renin-angiotensin aldosterone system**

Hypertensive disorders are strongly linked with an overactive renin-angiotensin aldosterone system (RAAS), and RAAS inhibitors (Ferrari, 2013). The renin-angiotensin aldosterone system plays an integral role in the homeostatic control of arterial pressure, tissue perfusion, and extracellular volume (Atlas, 2007; Katsi et al., 2015). The renin produced by the juxtaglomerular cells in the kidney cleaves angiotensinogen, which is produced by the liver, to release the inactive polypeptide angiotensin I (Ang I). Ang I is converted by angiotensin-converting enzyme into the polypeptide angiotensin II (Ang II). Ang II is the effector enzyme of the cascade and most biological actions of Ang II are mediated primarily through type I (AT₁), and type II (AT₂) receptors. In human activation of AT₁ receptors causes vasoconstriction via activation of phospholipase and inhibition of adenylate cyclase. Increased blood pressure, promotes adrenal aldosterone secretion, renal sodium reabsorption and release of catecholamine from the adrenal medulla and prejunctional nerve endings (Macit, Mercanoglu, 2015). The induced effects include vasoconstriction, sodium and water retention, decreased renal perfusion and cardiac and vascular hypertrophy (Morishita, Kusano, 2013).

There are well-established drugs that interfere with RAAS at several sites, including (1) ACEIs, (2) ARBs, (3) direct renin inhibitors (DRIs), (4) mineralocorticoid receptor antagonists (MRAs), and even (5) beta blockers, the last of which may be considered partial inhibitors (Ghazi, Drawz, 2017; Ferrari, 2013). ACEIs and ARBs have been the cornerstone
of RAAS inhibition for years and are key therapeutic options in patients with hypertension, reducing cardiovascular morbidity and mortality and improving renal outcomes (Ghazi, Drawz, 2017). However, inhibition of RAAS with ACEIs or ARBs has proven effective for controlling hypertension most important handicap of these agents is incomplete blockage of RAAS. A fundamental reason is reduced feedback inhibition of renin release, triggering a reactive rise in plasma renin activity. With an ACEIs, the reactive rise in plasms renin activity causes compensatory increase in Ang I, which partially restores Ang II production by both ACE-dependent and independent pathways (ACE-escape phenomenon) (Macit, Mercanoglu, 2015).

Fig. 1. Schematic representation of the renin-angiotensin-aldosterone system and its pharmacological blockade (Pantzaris, Karanikolas, Tsiotsios, Velissaris, 2017)

Renin secretion, a rate-limiting step, is the first step of the RAAS cascade, and thus represents the most logical target for inhibition of the renin system. Whereas angiotensinogen is present at plasma concentrations of approximately 500 to 600 pmol, Ang I is present in the
50- to 100-fmol range and Ang II at approximately half that. From a pharmacologist’s perspective, the renin step would be the one earning the target focus, as it is the step with the largest step down in concentration (Müller, Luft, 2006). Renin has a unique specificity for its substrate angiotensin. Inhibition of renin provides an attractive option to inhibit the RAAS from its origin (Ghazi, Drawz, 2017). By complete blockage of RAAS at its origin, decrease in both Ang I and Ang II levels can be achieved by DRIs. Although blocking feedback inhibition causes reactive rise in renin secretion, plasma renin activity, the enzymatic activity of renin is markedly reduced by the DRI (DRI binds directly to the catalytic site of renin). Therefore, through this more complete RAAS inhibition, DRIs can offer greater protection from hypertensive complications (Müller, Luft, 2006; Musini et al., 2017).

The development of DRI started more than 30 years ago (Ghazi, Drawz, 2017). The first synthetic renin inhibitor was pepstatin, which was followed by first-generation agents that were active but required parenteral administration. In 1980s, first orally active direct renin inhibitor compounds were developed, such as enalkiren, remikiren, and zankiren. Because of their poor bioavailability (less than 2%), short half-lives and weak antihypertensive activity they had limited clinical use (Macit, Mercanoglu, 2015).

Blood Pressure Lowering with Aliskiren

Aliskiren, an octanamide, is the first (approved in 2007) of a new class of completely non-peptide, low-molecular-weight (609.8 Da), orally active transition-state renin inhibitors and only approved for the treatment of hypertension (Pantzaris, Karanikolas, Tsiotsios, Velissaris, 2017; Nissenson, Fine, 2017), and a significant BP reduction has been demonstrated in patients with essential HT with no rebound effects on BP after the withdrawal of the treatment (Ghazi, Drawz, 2017; Wal, Rai, Dixit, 2011).

By binding to renin with high affinity, aliskiren blocks the conversion of angiotensinogen to angiotensin I, which subsequently results in a reduction in angiotensin II concentrations. Unlike the angiotensin-converting enzyme inhibitors and the angiotensin II receptor blockers, which reactively stimulate an increase in plasma renin activity, aliskiren suppresses the effects of renin and leads to a reduction in plasma renin activity (Sanoski, 2009).

Aliskiren has good water solubility (this property is important prerequisite for improved oral bioavailability) and low level of lipophilicity and is resistant to biodegradation by peptidases found in the intestine, circulation and/or the liver (Stanton, Jensen, Nussberger, O’Brien, 2003). Although aliskiren exhibits oral bioavailability compared with other previously synthesized renin inhibitors, it is still poorly absorbed (oral bioavailability, ~2.5%) (Wal, Rai, Dixit, 2011). Aliskiren is 50% protein bound, and the apparent volume of distribution is 135 L (Waldmeier et al., 2007). The plasma concentrations of aliskiren peak between 2 and 4 h following its administration and steady state is reached after 5–8 days of once-daily administration (Staessen, Li, Richart, 2006; Pool, 2007). Aliskiren demonstrates dose-linear pharmacokinetics over the dose range 75–600 mg in healthy volunteers (Sica, 2009). It has a long half-life (approximately 40 h) and provides a 24-hour antihypertensive effect with once-daily dosing with less potential for loss of efficacy between doses than shorter acting agents (Saseen, 2013). Following oral administration, aliskiren undergoes minimal metabolism (based on in vitro studies, aliskiren is metabolized by CYP 3A4) and is mainly eliminated as unchanged (mostly unabsorbed) compound in the feces. The pharmacokinetics of aliskiren are not effected by moderate to
severe chronic kidney disease or hepatic impairment. Aliskiren has a low potential for drug interactions (including warfarin, digoxin, statins and other antihypertensive agents, celecoxib, cimetidine) relating to its modest protein binding, limited metabolism as well as its lack of effect on a wide range of CYP450 isozymes (Sica, 2009, Macit, Mercanoglu, 2015). Coadministration of aliskiren with furosemide reduced the AUC of furosemide by 28% and C\text{max} by 49%, but clinical significance of this remains uncertain.

**Effects of other drugs on aliskiren**

Irbesartan: Coadministration of irbesartan reduced aliskiren C\text{max} up to 50% after multiple dosing. P-glycoprotein Effects: Pgp (MDR1/Mdr1a/1b) was found to be the major efflux system involved in absorption and disposition of aliskiren in preclinical studies. The potential for drug interactions at the Pgp site will likely depend on the degree of inhibition of this transporter.

Atorvastatin: Coadministration of atorvastatin resulted in about a 50% increase in aliskiren C\text{max} and AUC after multiple dosing.

Ketoconazole: Coadministration of 200 mg twice-daily ketoconazole with aliskiren resulted in an approximate 80% increase in plasma levels of aliskiren. A 400-mg once-daily dose was not studied but would be expected to increase aliskiren blood levels further.

Itraconazole: Coadministration of 100 mg itraconazole with 150 mg aliskiren resulted in approximately 5.8-fold increase in C\text{max} and 6.5-fold increase in AUC of aliskiren. Concomitant use of aliskiren with itraconazole is not recommended.

Cyclosporine: Coadministration of 200 and 600 mg cyclosporine with 75 mg aliskiren resulted in an approximately 2.5-fold increase in C\text{max} and fivefold increase in AUC of aliskiren. Concomitant use of aliskiren with cyclosporine is not recommended.

Verapamil: Coadministration of 240 mg of verapamil with 300 mg aliskiren resulted in an approximately twofold increase in C\text{max} and AUC of aliskiren. However, no dosage adjustment is necessary (Wal, Rai, Dixit, 2011).

Renin inhibition with aliskiren reaches above 99% in the first hours following administration and remains above 95% 24 h later (Pantzaris, Karanikolas, Tsiotsios, Velissaris, 2017). Aliskiren was generally well tolerated, and produced sustained BP reductions in patients with hypertension during 6 months, greater than those with ramipril-based therapy.

The incidence of adverse events with aliskiren and the number of study discontinuations as a result of adverse events during aliskiren treatment have been relatively low and were similar to results obtained in patients treated with placebo (Allikmets, 2007). Most commonly reported adverse effects are headache, diarrhea, dizziness, fatigue, and back pain. Furthermore because of having no interaction with metabolism of substance P and bradykinin, cough and angioedema like adverse effects caused by the use of ACEIs, do not occur with aliskiren treatment (Macit, Mercanoglu, 2015). Aliskiren had no clinically important effects on total cholesterol, high-density lipoproteins, fasting triglycerides, or fasting glucose. Laboratory abnormalities that may occur in some patients include a minor increase in blood urea nitrogen and serum creatinine, small reductions in hemoglobin and hematocrit, an increase in serum potassium greater than 5.5 mEq/L, elevated uric acid levels, and renal stones (Wal, Rai, Dixit, 2011).

Aliskiren is available in 150- and 300-mg tablets. The usual recommended starting dose of aliskiren is 150 mg QD. Doses >300 mg did not provide an increased BP response but did increase the rate of diarrhea (Pantzaris, Karanikolas, Tsiotsios, Velissaris, 2017). The antihypertensive effect of a given dose of aliskiren is attained after 2 weeks of therapy. No
dosage adjustment is required when used in elderly patients (i.e., those aged >65 years) or those with mild to severe renal impairment (creatinine clearance, <80 mL/min) or hepatic impairment (Child-Pugh Clinical Assessment score, 5–15) (Wal, Rai, Dixit, 2011).

Rebound hypertension has not emerged as a problem with aliskiren. Theoretically it is conceivable that long-term renin-inhibition therapy could induce pharmacologic tolerance with renin hypersecretion as well as the phenomenon of rebound hypertension after abrupt cessation of chronic therapy. However, clinical experience with aliskiren does not confirm this (Allikmets, 2007).

Several randomized control trials conducted showed significant dose-related BP lowering effects with aliskiren monotherapy, similar to those observed with losartan (LOS), valsartan (VAL), irbesartan (IRB), and lisinopril (LIS), and a placebo-like tolerability profile. Several meta-analyses published concluded that aliskiren is equally effective with ARBs with a similar adverse effects profile and is as effective as ARBs although it had higher control rates. Aliskiren was also proven superior to ACEIs in diastolic BP reductions, similar to hydrochlorothiazide (HCTZ), and inferior to CCBs in BP reduction and control rates (Pantzaris, Karanikolas, Tsiotios, Velissaris, 2017).

The combination therapy of aliskiren and CCBs, diuretics, ACEIs and ARBs may have synergistic anti-hypertensive effects (Morishita, Kusano, 2013). Aliskiren also neutralizes the reactive PRA increase caused by HCTZ. Studies of aliskiren/amlodipine combinations showed that doses of 300–150/10 mg are more effective than amlodipine 10 mg monotherapy and have a significantly lower incidence of peripheral edema. Liu et al., in 2014 showed that combination therapies aliskiren with amlodipine or HCTZ were more efficient than the respective monotherapies and that aliskiren/amlodipine produced significantly greater SBP/DBP reductions, and higher response and control rates. Triple combinations with aliskiren/amlodipine/HCTZ 300/10/25 mg have also shown similar tolerability and higher efficacy with significantly larger msSBP/msDBP reductions and higher control rates as compared to the components’ dual combinations in patients with moderate-to-severe hypertension (Pantzaris, Karanikolas, Tsiotios, Velissaris, 2017).

According to current data, the combination of aliskiren and ACEIs or ARBs is not recommended for hypertensive patients with diabetes, CVD or at least renal disease, especially when considering the risk of developing new adverse events such as hyperkalemia, renal dysfunction and hypotension (Sen, Ufuktepe, Ozunal, Uresin, 2014; Macit, Mercanoglu, 2015).

Hyperkalemia is the primary danger associated with RAAS blocking medications. The RAAS blockade leads to a decrease in aldosterone levels. Since aldosterone has a central role in the excretion of potassium, the RAAS blockers can cause retention of potassium. However, since the benefits of RAAS inhibitors outweigh the risks of hyperkalemia, there remains the need to overcome these challenges rather than withdraw treatment (ESC, 2016).

The addition of aliskiren to standard optimal therapy (ACEIs or ARBs and beta-blocker) in post-MI patients did not produce any change in left ventricular end-systolic volume compared to placebo. No benefit was shown by the use of aliskiren in prehypertensive individuals with coronary atherosclerosis (Pantzaris, Karanikolas, Tsiotios, Velissaris, 2017) or improvement in cardiovascular outcomes in patients hospitalized with heart failure (Ghazi, Drawz, 2017) was seen with aliskiren compared with placebo.
Conclusions and suggestions

Hypertension is the most common condition seen in primary care and leads to myocardial infarction, stroke, renal failure, and death if not detected early and treated appropriately. Evidence from randomized controlled trials has shown benefit of antihypertensive drug treatment in reducing important health outcomes in persons with high BP using diuretics, BBs, CCBs, ACEIs and ARBs. The renin-angiotensin aldosterone system plays an integral role in the homeostatic control of arterial pressure. However, adequate RAAS blockade cannot be achieved with ACE inhibitors or ARBs because of incomplete blockage of RAAS. The primary rate-limiting step in the RAAS now can be pharmacologically inhibited directly. Aliskiren, an octanamide, is the orally active transition-state renin inhibitors and only approved for the treatment of hypertension. Aliskiren is effective in reducing blood pressure and is well tolerated, with a side-effect profile similar to placebo or ACEIs and ARBs. The combination therapy of aliskiren and CCBs, diuretics, ACEIs and ARBs exhibits synergistic anti-hypertensive effects. Many patients often require multidrug antihypertensive therapy. Aliskiren can play an important role as a RAAS-blocking drug in combination therapy. There is a need for longitudinal studies assessing aliskiren alone and in combination in treatment of persons with diabetes mellitus, metabolic syndrome and resistant hypertension to identify the specific categories of patients that would benefit more from direct renin blockade.

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