New therapeutic targets for ACE inhibitors and angiotensin receptor blockers

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Abstract: Angiotensin-converting enzyme inhibitors (ACE-I) and angiotensin receptor blockers (ARB) possess multiple beneficial effects such as cardioprotection, cerebroprotection, nephroprotection which provide opportunity to select the most suitable drug for the target vascular bed (e.g. coronary, or cerebral circulation). In some clinical settings, combined therapy ACE-I with ARB (double blockage of the renin–angiotensin–aldosteron system) may appear the most effective. These drugs (especially ARB) may successfully prevent atrial fibrillation and play a protective role in metabolic syndrome. Recently, it has been demonstrated that losartan is able to inhibit vasodilatation of the aorta in Marfan syndrome, which might prevent sudden death due to aorta rupture. An increasing role of ARB is most beneficial in hipotensive therapy (inhibition/regression of hypertension-related organ damage). With particular interest, results of the ONTARGET study are being awaited. This study is focused on the effect of dual blockade (ramipril and telmisartan) on reduction of the occurrence of myocardial infarction, stroke, and heart failure.

Key words: ACE inhibitors, angiotensin, angiotensin receptor blockers

INTRODUCTION

History of the pharmacotherapy of renin–angiotensin–aldosteron (RAA) axis dates back over a quarter of century. This history had some surprising, but also very beneficial reversals. Angiotensin-converting enzyme inhibitors (ACE-I), drugs originally intended to treat arterial hypertension, turned out to be a key to neurohormonal (readaptive) treatment of congestive heart failure. After it had occurred that the ACE-I are multi-potential and irreplaceable drugs (a milestone in cardio development), a potential rival appeared – angiotensin II receptor blockers (ARB), which in certain cases act better (e.g., cerebral protection in primary and secondary stroke prevention). Of note, instead of competition, the cooperation between those drugs was achieved – an interesting new trend (gaining an increasing acceptance) of the RAA axis dual blockade by the simultaneous use of the ACE-I and ARB. In some clinical cases this simultaneous (two-level) therapy – a remedy for the “escape from RAA system” – is very useful (nephroprotection). The next success of the RAA blocking drugs is their efficacy in the prevention of atrial fibrillation and treatment of metabolic syndrome. In both diseases of an increasing frequency (as a result of population ageing) especially the ARB are very efficient in prevention.

The role of ARB in stroke prevention

Among the trials concerning hypertensive patients in which the usefulness of the ARB in primary cerebral stroke was assessed, the LIFE [1] and SCOPE [2] are of special interest. In the Losartan Intervention For Endpoint trial (LIFE) reduction in hypertension was observed in the elderly patients suffering from isolated systolic hypertension. After losartan treatment stroke frequency was decreased by 25% in comparison to atenolol. The similar efficacy was shown by candersartan in the SCOPE trial. The MOSES trial showed that in secondary prevention of stroke eprosartan was successful [3]. Paying attention to the special efficiency of ARB, other hypotensive drugs were analyzed (some of them known as “therapeutic pillars” in treatment of cardiovascular diseases and their complications, e.g. cerebral stroke). A valuable summary of β-adrenolytics efficacy in stroke prevention, published in Lancet, was made on 6825 patients by Carlberg et al. [4]. Surprisingly, comparing to other hypotensive drugs, atenolol increased by 30% the risk of stroke. Drugs which decrease the risk of cerebral stroke were suggested to be called “cerebroprotective drugs”. Besides thiazide diuretics, the ARB were also included to this group [4]. The high efficacy of the SARB was mentioned above; another important observation is that if instead of the AT$_2$ receptor blockade, the ACE-I are used, the cerebroprotective effect is lost (a complex role of AT$_2$ receptor was described below in detail). For example, in the PROGRESS trial [5] perindopril in monotherapy decreased blood pressure by 5 mmHg and the risk of stroke by 5% – which was statistically insignificant. Confronted to the Post-stroke Antihypertensive Treatment Study (PATS) [6], the use of indapamide in monotherapy also
Angiotensin II (ANG II) is the most important element of the RAA axis, since it plays a pivotal role in the development of cardiovascular diseases, both through the hemodynamic and direct influence on vessels, as well as indirectly through aldosterone stimulation [11,12]. The ANG II derives from the ANG I in reaction catalyzed, among others, by the ACE. Through the activation of angiotensin receptors, biological action of the ANG II is transduced. The basal and the best known is the AT1 receptor, however, several others were also detected (AT2, AT3, AT4); currently, the clinical impact of the AT1 receptor is shown [13,14]. By the activation of the AT1 receptor, the ANG II mediates several processes: the release of aldosterone with the consequent retention of sodium and water, increased release of epinephrine and endothelin (neurohormonal effect), vasoconstriction, increased myocardial contractility, cell proliferation and hypertrophy, as well as oxidative and inflammatory action. Recently, the proatherogenic action of the ANG was shown, since the activation of the AT1 receptor causes unfavorable changes of lipid and coagulation parameters, which leads to atherosclerotic plaque development. The effect of the ANG II by the AT1 receptors is partially opposite to the effects mediated by the AT1 receptors – the activation of the AT1 receptors causes vasodilation and antimitogenic as well as apoptotic rate decreasing actions. However, the role of the AT1 receptors is more complex; in certain cases they send signals similar to the AT2 receptors, causing the expression of the nuclear factor κB (NF-κB), which is responsible for inflammatory and fibrotic reactions.

The activation of the AT1 receptors in the circulatory system leads to an increase of nitric oxide, prostaglandin I2 and bradykinin synthesis; it also leads to a decrease in the type T calcium channels activity and the opening of potassium channels [13,14]. The AT1 receptors stimulation causes small vessels vasodilatation and recruits collaterals in the ischemic region; it also protects against the consequences of anoxia (by increasing neuronal resistance to oxygen deficiency). In contrast to the AT2 receptors selective recruitment action in ischemic region, bradykinin (increased by the ACE-I) causes vasodilatation also in non-ischemic regions [15]. In consequence, vasodilative competition can occur between ischemic and non-ischemic regions, with the subsequent ischemic region steal syndrome.

The ARB inhibit the RAA axis activity, but the mechanism of action is different from that of the ACE-I. The ACE-I decrease the ANG II production, whereas the ARB act by competitive and selective blocking of the ANG II action. It

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**Fig.** Cerebroprotection mechanism of selected hypotensive drugs

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**Revision Articles**

**Dual action of the ARB – the role of angiotensin receptor types**

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The ARB inhibit the RAA axis activity, but the mechanism of action is different from that of the ACE-I. The ACE-I decrease the ANG II production, whereas the ARB act by competitive and selective blocking of the ANG II action. It
was suggested that the AT$_2$ receptor blockade acts in cerebroprotective mode, since it diminishes the vasoconstriction of large cerebral vessels (fig.) and restricts inflammation, connected with ischemic-reperfusion damage [16]. By this receptor blockade seemingly a better inhibition of the RAA axis, than by the ACE-I, was achieved. However, there is a drawback of the AT$_2$ receptor blockade: by the inhibition of the "feedback loop", the ARB cause the increase of the ANG II concentration.

The ACE-I act in another way, since they decrease the ANG II concentration. In a group of patients with a low concentration of the ANG II (e.g., hypertensive Afro-Americans), this difference in clinical effects between the two classes of drugs, as a result of an opposite action on the ANG II concentration, is of a special importance. For example, in the ALLHAT trial the risk of stroke in this population was by 40% higher in the lizinopril-treated comparing to the chlortalidone (diuretic)-treated group, despite a very similar degree of blood pressure reduction. A question arises why the increase of the ANG II concentration causes cerebroprotection. The increase of the ANG II concentration causes a non-inhibited stimulation of the AT$_2$ receptors – number of which is additionally increased. This phenomenon is called up-regulation.

The AT$_1$ receptor which plays a key role in cerebroprotection is also stimulated by diuretics and dihydropiridylne calcium antagonists (fig.) [9,10]. The direct blockade of the AT$_1$ receptor, caused exceptionally by sartans, influences also cerebral vessel vasodilatation. β-adrenolytics, dihydropiridylne calcium antagonists and the ACE-I lack this favorable effect.

In the experimental model of stroke – the closure of cervical artery (preceded by drug administration), a decrease of mortality after the ARB, in comparison to the ACE-I was obtained [17]. It was accompanied by a faster partial reflow on the stroke-induced side. The exceptional role of the ARB is confirmed by the observation that in the case of combined administration of the ARB and ACE-I, there was no decrease of mortality detected. Coming from bench to bedside, it was revealed that in patients after cerebral ischemia, the expression of the AT$_2$ receptors is increased (fig.). That is why sartans may occur especially beneficial in secondary stroke prevention.

Dual ACE-I and ARB blockade

Another novelty in the RAA blockade, after the cerebroprotective effect, is the ACE-I and ARB combined treatment concept [18]. This idea was born on the basis of several observations. The main premise was that no drug used in monotherapy (despite constant improvement of drugs: increased biological availability, extended time of action, improvement of penetration to tissues) is able to block totally the RAA axis. Importantly, the effect of blockade is eminent, since in the peak first action of the ACE-I, the RAA activity is decreased by 60–80% [18]. However, this submaximal blockade represents only the initial effect of treatment and unfortunately, during the next days of therapy, the RAA activity is unfavo-
the dual blockade (losartan and trandolapril), in 23% treated only with losartan (p = 0.016) and in 23% administered trandolapril monotherapy (p = 0.018).

As regards heart failure, one must realize that in this disease the RAA system activity is augmented and the increased ANG concentration is an unfavorable prognosis factor. The dual blockade decreased the NYHA level and restricted the detrimental myocardial remodelling. In the Val-HeFT trial [29] sartan addition to the previously used ACE-I decreased the occurrence of the complicated end-point risk (death + morbidity due to heart failure). In the CHARM-Added trial [30] the dual blockade decreased a risk of cardiovascular complications. In patients treated with the dual blockade hospitalization due to heart failure was rare (p <0.01) and death from a cardiovascular cause was also uncommon.

**ARB in metabolic syndrome**

Nowadays metabolic syndrome becomes epidemic. It is frequently accompanied by hypertension (one of syndrome’s factors) which makes physicians maximize pharmacotherapy. The multidirectional effect of sartans makes them a valuable treatment tool. Discussing the role of sartans in hypertensive patients with metabolic syndrome, it was shown that decreased insulin resistance is a key factor of the special usefulness of these hypotensive drugs. The advantages of sartans, improving metabolic disturbances, are:

1) decrease of insulin resistance, shown in the ICARUS trial [36],
2) agonistic effect of some ARB on PPAR-γ receptors which improves insulin sensitivity,
3) neutral effect on carbohydrate and lipid metabolism, documented in several trials including the ALPINE trial [37],
4) demonstration in trials (LIFE with losartan, CHARM and VALUE) a reduction of new non-insulin-dependent diabetes cases,
5) nephroprotective action, e.g. the decrease of microalbuminuria in patients with non-insulin-dependent diabetes.

The ARB demonstrate also other – extrametabolic, equally desired effects: a beneficial effect on regression of left ventricle hypertrophy, a high hypotensive efficiency and an excellent tolerance by patients.

**Prevention of atrial fibrillation**

The better and better effectiveness of electrophysiological procedures allows the effective treatment of supraventricular and ventricular causes of cardiac arrhythmias. Unfortunately, the “Achilles’ heel” of this invasive method is atrial fibrillation treatment, since these operations are less effective and more expensive. Another disadvantage is the possibility of complications with the subsequent implantation of cardiodiostimulator. Therefore, pharmacotherapy is the main treatment of atrial fibrillation, however, commonly used classical antiarrhythmic drugs (mainly from class I, II and IV) have a limited efficacy and tolerance, especially during prolonged treatment (which is the usual therapeutic horizon perspective in atrial fibrillation).

Due to above restrictions, advisable is a search for new therapeutic modalities increasing efficacy of treatment to restore/maintain sinus rhythm. In currently commonly occurring diseases like hypertension and heart failure, atrial fibrillation could be a consequence of pathologically changed cell structure, increased neurohormonal activation and disturbed signal transduction among atrial (but also ventricular) cells [38]. Due to the beneficial effect of the drug blocking RAA system on diseases which are risk factors for atrial fibrillation (heart failure, hypertension, coronary artery disease) one may assume that the application of this type of drugs will decrease the risk of cardiac arrhythmias. In atrial fibrillation there are three types of remodeling: neurohormonal, electrical and structural. The potential mechanisms of the drug blocking RAA system protective effect include: the inhibition of structural remodeling by improving left ventricle and atrium hemodynamic parameters, decrease of atrial tension, inhibition of fibrosis, prevention of neurohormonal remodeling and a direct effect on ion channels (electrical remodeling) [39]. During fibrillation, atrial stretching with walls tension increases the local synthesis of the ANG II. The density of the AT receptors in atria is higher than in ventricles, therefore atria are more sensitive to the ANG II. Receptor stimulation causes the activation of mitogen activated protein kinase (MAP kinase) which in turn leads to the hypertrophy of myocytes, proliferation of fibroblasts and apoptosis [40]. Besides the structural remodeling, it was shown in experimental research [41] that the ANG II affects atrial “functional” remodeling, called electrical remodeling [41]. This pathophysiological process leads to significant disturbances. Thanks to protein kinase C activation and the phosphorylation of L channels, the calcium current through them is increased [42].

The beneficial effects of the ACE-I derive mainly from the inhibition of the ANG II synthesis, which directly causes the decrease of blood pressure, increase of large vessels compliance, decrease of hypertrophy and prevention of left ventricle remodeling. It is contributed by afterload and systolic tension reduction as well as the decrease of atrial pressure and tension which consequently lead the mitral insufficiency degree decrease. As regards neurohormonal re-adaptation, the concentration of catecholamines in blood and myocardium is diminished. The ACE-I application in patients with fixed atrial fibrillation decreases the expression and concentration of protein kinases, responsible for fibrosis activation [40]. It was shown in the experimental model of heart failure that enalapril inhibits both atrial fibrosis and remodeling. Consequently, the risk of atrial fibrillation as a complication of heart failure is also diminished. These data prove an important role of the RAA system in atrial remodeling [43].

Coming from bench to bedside, the prevention of the atrial fibrillation efficiency after the ACE-I was estimated retrospectively and prospectively. The retrospective analysis, concerning a group of hypertensive patients treated with the ACE-I or calcium antagonists showed significantly less atrial fibril-
lation attacks in the ACE-I group [44]. In the retrospective analysis of the SOLVD trial, in which significant benefits from enalapril treatment were proven in patients with heart failure, the reduction of frequency of atrial fibrillation occurrence by 18.6% was stated in the group treated with the ACE-I, comparing to placebo [45]. Further reports from the trial pointed to a much smaller number of hospitalizations due to atrial tachyarrhythmias during enalapril treatment [46].

Prospective trails are not finished yet; they concern also the evaluation of drugs from the ARB group (GISSI-AF, ACTIVE-AF, ANTIAPAF).

Another important issue is the pharmacological maintenance of sinus rhythm after a successful electrical cardioversion. Concerning also this secondary prevention, the retrospective analysis of results showed the decrease in the number of new atrial fibrillation attacks, in the group of patients treated with the ACE-I due to heart failure [47]. In the prospective trial the effects of associated amiodarone and enalapril treatment were compared to amiodarone alone, 28 days before planned cardioversion and after sinus rhythm restoration. Combined treatment (by classical antiarrhythmic drug and the ACE-I) is burdened with smaller risk of cardiac rhythm disturbances both in the early and late periods of observation [48]. In another prospective trial [49] patients with paroxysmal atrial fibrillation were randomly assigned to 3 groups with following schemes of treatment: monotherapy (amiodarone), two-drug therapy (amiodarone and losartan), and three-drug therapy (amiodarone, perindopril and losartan). In groups subjected to combined treatment, a significantly smaller number of atrial fibrillation episodes was shown in short- and long-term observation (p <0.006 and p <0.04). Moreover, in these groups a decrease of left atrium dimensions (p <0.001), making up the determinant of re-adaptation counterbalancing structural remodeling, was found. Importantly, there were no significant differences between groups treated with amiodarone and losartan and treated with amiodarone, losartan and perindopril [49].

Taking into consideration the long-term prophylaxis of atrial fibrillation, and simultaneously knowing the limited efficacy and potential adverse effects of antiarrhythmic drugs from groups I, III and IV, one can regard the ACE-I or the ARB as some of elements or even basis of pharmacotherapy. Therapy with these drugs is directed for the elimination of hypertensive organ complications, a fast regression of organ changes and leveling the effect on risk factors in patients from a cardiovascular high risk group. With special interest, the results from the ONTARGET study, designed on a large population of patients, are expected. In this trial the effect of the dual blockade (ramipril and telmisartan) on the reduction of the occurrence of myocardial infarction, cerebral stroke and heart failure episodes, will be assessed.

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