Clinical utility of fixed-dose combinations in hypertension: evidence for the potential of nebivolol/valsartan

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Abstract: Despite significant advances in pharmacologic approaches to treat hypertension during the last decades, hypertension- and hypertension-related organ damage are still a high health and economic burden because a large proportion of patients with hypertension do not achieve optimal blood pressure control. There is now general agreement that combination therapy with two or more antihypertensive drugs is required for targeted blood pressure accomplishment and reduction of global cardiovascular risk. The goals of combination therapies are to reduce long-term cardiovascular events by targeting different mechanism underlying hypertension and target organ disease, to block the counterregulatory pathways activated by monotherapies, to improve tolerability and decrease the adverse effects of up-titrated single agents, and to increase persistence and adherence with antihypertensive therapy. Multiple clinical trials provide evidence that fixed-dose combinations in a single pill offer several advantages when compared with loose-dose combinations. This review discusses the advances in hypertension control and associated cardiovascular disease as they relate to the prospect of combination therapy targeting a third-generation beta (β) 1-adrenergic receptor (nebivolol) and an angiotensin II receptor blocker (valsartan) in fixed-dose single-pill formulations.

Keywords: blood pressure control, hypertension, β1-adrenergic receptor, renin angiotensin system, fixed-dose combination therapy, nebivolol, valsartan

Introduction

Hypertension, having a high prevalence worldwide,1,2 is estimated to affect approximately 78 million persons in the United States.3,4 Although high blood pressure (BP) is recognized as an important cardiovascular risk factor for the development of coronary heart disease and stroke,5 effective hypertension control can reduce heart failure (HF) incidence by 50%.6

Despite increased awareness of both patients and physicians as to the consequences of uncontrolled BP and the availability of new pharmacologic approaches to treat hypertension during the last decades,7 hypertension-related and, consequently, hypertension-related organ damage still make up a high health and economic burden because a large proportion of patients with hypertension do not achieve optimal BP control. Thus, there is an imperative for efficient control of hypertension, as the economic burden to society associated with hypertension was estimated to a have cost of US$73.4 billion in 2009.8 Although numerous initiatives from the medical community across the world continue to adjust guidelines and policies to assist in hypertension prevention, detection, and treatment,6 the need for focus management of hypertension and its associated risk factors using appropriate medicines and effective comanagement of cardiometabolic...
risk factors remains unattained. This review discusses the advances in hypertension control and associated cardiovascular (CV) disease as they relate to the prospect of combination therapy targeting beta (β1)-adrenergic receptor (nebivolol) and the renin angiotensin system (RAS), angiotensin II (Ang II) type 1 receptor (valsartan) in fixed-dose single pill.

BP control/cardiovascular risks/target organ diseases: a need for combination therapy

An emerging view suggests that hypertension results from an imbalance between vasoconstrictor and vasodilatory mechanisms of diverse hormonal, neurogenic, and local vascular nature. The heterogeneous, often time-dependent, contribution of altered mechanisms influencing BP and, hence, tissue perfusion, accounts for the successful development and availability of classes of pharmacological options: thiazide and thiazide-like diuretics, selective (β1-) and alpha (α1-) adrenergic receptor blockers, calcium channel blockers (CCBs), various drugs inhibiting RAS components (i.e., direct renin inhibitors, angiotensin converting enzyme inhibitors [ACEIs], and Ang II receptor blockers [ARBs]), and vasodilators. To achieve optimal BP control, the Seventh Report of the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure (JNC 7) endorses the use of combination therapy rather than increasing the dose of an agent. Although approaches geared toward the prevention of hypertension and its clinical complications through lifestyle modifications, increased physical activity, and dietary changes are critically important, antihypertensive therapy will continue to be the mainstay of therapy.

There is now general agreement that monotherapy is less likely to be sufficient to achieve optimal BP control in a majority of patients, and combination therapy with two or more antihypertensive drugs is usually required for targeted BP accomplishment and reduction of CV risk. Moreover, according to JNC 7 guidelines and the guidelines put forth jointly by the International Society of Hypertension and the European Society of Hypertension, BP-lowering treatment should be initiated with a combination of two drugs in patients with a systolic pressure higher than 20 mmHg and/or a diastolic pressure higher than 10 mmHg of the targeted goal, as well as in the patients with multiple CV risk factors such as metabolic syndrome, diabetes, and heart and renal disease. The recommendation to target multiple pathophysiological mechanisms underlying the development and progression of hypertension using combination therapy of two or more complementary, yet distinct, drugs is supported by the evidence that in comparison with monotherapy, combination therapy raises the probability of achieving a BP target in a shorter amount of time, and combination therapy is several times more potent than increasing the dose of the single drug.

Indeed, the use of combination therapy at diagnosis contributed to a greater CV risk reduction when compared with monotherapy. According to a global survey conducted by the International Society of Hypertension, the preferred combination across different countries is RAS inhibition combined with CCB (particularly for the patients with type 2 diabetes) or diuretics. The successful combination of drugs also reflects the ability of the components to block the counterregulatory pathways activated by each other. Thus, when CCBs and diuretics activate RAS, the addition of RAS inhibitors will offer more favorable outcomes for the combination therapy than for CCB or diuretics alone. Undeniably, convincing evidence from large clinical studies such as the Avoiding Cardiovascular Events Through Combination Therapy in Patients Living with Systolic Hypertension (ACCOMPLISH), the Valsartan Antihypertensive Long-term Use Evaluation (VALUE), the Action in Diabetes and Vascular Disease: Preterax and Diamicron MR Controlled Evaluation (ADVANCE), and the Perindopril Protection Against Recurrent Stroke Study (PROGRESS) trials clearly support the use of combination therapies to initiate antihypertensive treatment. An additional rationale for combination regimes is based on improved tolerability and decreased adverse effects of up-titrated single agents. In addition to combination therapy given separately, fixed-dose combinations (FDCs) in a single pill offer several advantages reflected in greater adherence and improved treatment outcomes when compared with loose-dose combinations.

β-blockers in hypertension management

Increased sympathetic nervous system activity is a major contributor to the development and progression of hypertension. Agents that block cardiac β-adrenergic receptors have been used as an alternative approach to diuretics for the initial treatment of hypertension in many countries for years. However, evidence from a recent meta-analysis suggests that β-blockers, apart from coronary heart disease, are less effective in preventing CV disease, particularly in the elderly. Thus, they are no longer regarded as first-choice drugs in the treatment of elderly patients or patients with metabolic syndrome, diabetes, or renal disease. Although the panelists appointed to the Eighth Joint National Committee (JNC 8)
did not recommend the use of β-blockers, specifically atenolol or metoprolol, for the initial treatment of hypertension on the basis of the results obtained in the Losartan Intervention for Endpoint Reduction in Hypertension (LIFE) Study,17 we directly quote the JNC statement acknowledging that “[…] compared a β-blocker to the 4 recommended drug classes, the β-blocker performed similarly to the other drugs (question 3, evidence statement 8) or the evidence was insufficient to make a determination (question 3, evidence statements 7, 12, 21, 23, and 24).”56 The exclusion of primarily atenolol for initial hypertension treatment was based on the results obtained in the Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack (ALLHAT) Trial,38 but even in this situation, the conclusion is muddied by the fact that most patients assigned to the chlorthalidone group of the study received a β-blocker as the second drug for appropriate BP control. These limitations do not apply to the more advanced cases of hypertension with concomitant comorbidities, particularly HF. Thirty-nine years after Waagstein’s original report of the beneficial effects of β-blockers in HF,29 these agents have become a well-established treatment component of HF management. This is an important consideration, as HF admissions to community hospitals pose a significant burden to the health care system, given the fact that readmission rates average 90 days postdischarge and 30-day mortality rates remain quite high.40

β-blockers are a heterogeneous class of drugs consisting of agents with different adrenergic receptor selectivity, lipophilicity, and intrinsic sympathimimetic activity, as well as vasodilatory and antioxidant properties (Table 1). Thus, the BP-lowering effect of traditional β-blockers with no vasodilatory ability (ie, atenolol, metoprolol, propranolol) is achieved through a decrease in central nervous system sympathetic outflow,31 reduced cardiac output consequent to decreased heart rate and contractility, and reduced renin release.41 The magnitude of the antihypertensive response may be offset by compensatory vasoconstriction in the peripheral vasculature31,42 and negative effects on metabolic parameters and lipid profile,43 as well as increased venous return mediated by β2-adrenergic receptor blockade enlarging the left ventricle.42

Recent advances in the development of β-blockers led to the introduction of agents with additional vasodilatory abilities resulting from their concomitant blockade of α1-adrenergic receptors (labetalol, carvedilol) or the intriguing action of nebivolol in stimulating vascular endothelium-mediated release of nitric oxide (NO). Nebivolol is a racemic mixture of D- and L-enantiomers. D-nebivolol is responsible for β1-antagonistic properties, whereas both D- and L-nebivolol activate NO synthase (NOS) predominantly through β2 agonist activation.44 In addition, nebivolol can increase NO bioavailability by decreasing levels of asymmetric dimethylarginine, an endogenous inhibitor of NOS, which elevated levels in plasma are shown to correlate with endothelial dysfunction in different CV diseases45 and/or by its antioxidant properties targeting nicotinamide adenine dinucleotide phosphate-oxidase46 or more directly scavenging reactive oxygen species.47,48 Moreover, nebivolol offers additional protective effects on endothelial dysfunction and vascular remodeling with supplementary antiinflammatory properties49 associated with neutral effects on metabolic parameters in patients with hypertension.50 In contrast to traditional β-blockers, nebivolol preserves cardiac output51 and reduces central aortic BP.52 Maintenance of cardiac output is related to the mild effects of the drug on heart rate coupled with the unique ability of this drug to induce peripheral venodilation, thus maintaining venous return.53 It has been also shown that nebivolol, but not metoprolol, improved left ventricular dysfunction in an experimental model of myocardial infarction54 and that these cardioprotective effects may be mediated by β3-receptors and NOS55 not only on the coronary vessels but also on the cardiac myocytes as well. Nebivolol also improves diastolic dysfunction and myocardial remodeling in Ang II-dependent hypertension,46 as well as in an obese insulin-resistant rodent model.56 These later findings are of particular relevance, as left ventricular diastolic dysfunction, prevalent in hypertensive women, has proven highly resistant to treatment with ACEIs or even ARBs.40

### Table 1 Primary characteristics of β-adrenergic receptors

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>β1-adrenergic receptor</th>
<th>β2-adrenergic receptor</th>
<th>β3-adrenergic receptor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of amino acids</td>
<td>477</td>
<td>413</td>
<td>408</td>
</tr>
<tr>
<td>Introns</td>
<td>–</td>
<td>–</td>
<td>2</td>
</tr>
<tr>
<td>Phosphorylation by pKα and βARK</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Most potent catecholamine</td>
<td>Norepinephrine</td>
<td>Epinephrine</td>
<td>Norepinephrine</td>
</tr>
<tr>
<td>Selective agonists</td>
<td>Xamoterol, dobutamine</td>
<td>Zinterol, procatel</td>
<td>BLR 37344, CGP 12177</td>
</tr>
<tr>
<td>Selective antagonists</td>
<td>CGP 20712A, Metoprolol</td>
<td>ICI 118551</td>
<td>SR 59230A</td>
</tr>
<tr>
<td>G protein</td>
<td>Gs</td>
<td>Gs/Gi</td>
<td>Gs/Gi</td>
</tr>
<tr>
<td>Effector</td>
<td>Adenylate cyclase</td>
<td>Adenylate cyclase</td>
<td>Adenylate cyclase, NO synthase</td>
</tr>
</tbody>
</table>

**Note:** Copyright ©2004. Medicina (Kaunas). Reproduced from Skeberdis VA. Structure and function of β1-adrenergic receptors. Medicina. 2004;40(5)407-413.52

**Abbreviations:** pKα, protein kinase A; βARK, beta-adrenergic receptor kinase; NO, nitric oxide.
Studies in animal models show that nebivolol augments vascular NO release, and data obtained from individuals with essential hypertension indicate that nebivolol treatment dilates forearm vasculature and reverses endothelial dysfunction. Nebivolol has also been shown to reduce QT dispersion in hypertensive patients with left ventricular hypertrophy after 4 weeks of treatment, an effect that did not correlate with BP reduction.

The beneficial action of nebivolol in terms of facilitating production/release of NO may account for the observation that sexual dysfunction is not observed in male patients medicated with this drug. In one study, atenolol resulted in sexual dysfunction in 12.8% of patients compared with 3% of placebo-treated patients. A second study found that atenolol was associated with sexual dysfunction in 17.3% of patients compared with 3% of lisinopril-treated patients. The effects of 12 weeks of therapy with nebivolol 5 mg once daily or metoprolol succinate 95 mg on sexual function in stage 1 hypertensive men were studied in a double-blind, randomized crossover study. The primary endpoint at the end of each 12 week treatment period was the assessment of changes in the International Index of Erectile Function. The index is a reliable, validated, and widely used assessment scale in most trials evaluating erectile function. At the end of the 12 week period and before crossover to the other drug, the erectile function subscore showed no significant changes from baseline in male patients with stage 1 hypertension who were treated with nebivolol, whereas the results for the mean changes from baseline of all secondary efficacy criteria derived from the International Index of Erectile Function (orgasmic function, sexual desire, intercourse satisfaction, and overall satisfaction subscores) favored nebivolol. An additional study in which patients were assigned to the combination of chlorthalidone/atenolol, atenolol, or nebivolol only confirmed the essential lack of sexual dysfunction in only those patients who were randomized to nebivolol 5 mg daily. Moreover, when nebivolol was compared with losartan, there was no difference in their effects on sexual function.

In the United States, nebivolol has been approved for the treatment of stage 1–2 hypertension, and in Europe and many countries around the world, it has been approved for the treatment of HF as well. The efficacy and tolerability of nebivolol monotherapy in patients with hypertension have been demonstrated in three pivotal phase 3 trials; results of another phase 3, randomized, double-blind, placebo-controlled trial in patients with stage 1–2 hypertension suggest that nebivolol, when added to an ongoing treatment with an ACEI, ARB, or diuretic, further lowers BP and improves BP response rate. A large German observational postmarketing study, carried out in 6,376 individuals with hypertension (mean systolic BP/diastolic BP, 173/101 mmHg; CV disease, 26.3%; diabetes, 20.3%; left ventricular hypertrophy, 20.4%; HF, 9.1%), demonstrated that a 6-week treatment with nebivolol resulted in DBP normalization (<90 mmHg) in 62.2% of the patients. The beneficial action of nebivolol in reducing central BP suggests improvement in vascular compliance that may be related to the observations that the drug can reverse cardiac and vascular fibrosis.

Nebivolol is a safe and effective treatment for elderly patients with HF improving diastolic function and reducing mortality. In addition, at doses of 5–20 mg/day, nebivolol is very well tolerated. Higher doses of 40 mg/day were associated with more adverse effects, probably because of loss of selectivity for β receptors. Thus, in addition to better tolerability and greater persistence, this novel β-blocker may be particularly useful in difficult-to-treat patients such as African Americans (5–40 mg/day dose range), elderly patients (5 mg/d), and Hispanics with hypertension. Their favorable hemodynamic effect in reducing peripheral vascular resistance, rather than cardiac output, may account for the absence of fatigue as a significant adverse reaction to the drug. An obese patient with HF is expected to show some level of fatigue, and fatigue has frequently been considered to be associated with β-blocker treatment. However, a 2002 meta-analysis of randomized trials of β-blockers used in the treatment of myocardial infarction, hypertension, and HF demonstrated a statistically significant, yet small, annual risk for fatigue that was equivalent to one additional report per 57 patients treated with a β-blocker, with the effect being more pronounced with older-generation β-blockers. A similar meta-analysis that looked at HF trials only failed to demonstrate an absolute risk for fatigue associated with β-blocker treatment. At the time of these meta-analyses, no nebivolol trial satisfying the inclusion criteria (randomized, placebo-controlled studies with at least 100 enrolled patients and a noncrossover design) was completed, but no participants in the 8 month study involving elderly subjects with HF (N=260) reported fatigue, and in the 2-year trial involving the elderly with HF (N=2,128), fatigue was reported by 6.7% nebovicol-treated and 5.8% placebo-treated participants. In the three pivotal US trials of nebivolol (12 weeks, total enrollment of 2,016 patients), no nebovicol-treated individual reported fatigue as an adverse event. There is currently a study underway (ClinicalTrials.gov NCT00999102) designed to explicitly examine the effects of nebivolol and metoprolol.
on fatigue and quality of life, which should provide helpful clinical evidence regarding the risk for fatigue with these two β-blockers.

Rationale for nebivolol/valsartan fixed-dose combination

It is well known that Ang II exerts its vasoconstrictor/hypertrophic/profibrotic effects by activating AT₁ receptors on vasculature, heart, and kidneys. In this way, Ang II acts as a major contributor to the development of hypertension and target organ damage. Thus, the combination of drugs that target Ang II synthesis (via inhibition of renin or ACE) or action (by blockade of AT₁ receptors) with hydrochlorothiazide or CCB has a long-standing history of additive BP lowering effects, as well as reduction in CV risk factors. In contrast, effectiveness of the combination of β-blockers and RAS system inhibitors have been questioned for a long time because of the potential overlap in their mechanisms of action. Indeed, evidence showed that blockade of β receptor halts RAS activity by reducing renin release, whereas inhibition of RAS decreased sympathetic nervous activity. However, an increasing body of evidence testifies that in hypertensive patients, nebivolol, in doses ranging from 5 to 40 mg/day, offers an additional BP lowering effect when it was administered in a combination therapy or as an add-on drug to the ongoing therapy with ACEI or ARB. Part of the favorable outcomes for the combination therapy could be attributable to the possibility that nebivolol may ameliorate the increase in plasma renin activity induced by elimination of the negative feedback on renin release mediated by blockade of AT₁ receptors. Increased Ang II observed with ARB treatment may also lead to “aldosterone escape”, and a reduction in aldosterone with the nebivolol therapy has been noted. Furthermore, in contrast to the added hydrochlorothiazide, little or no effect was shown on glucose metabolism and lipid profile in hypertensive patients with prediabetes when nebivolol was added on the background of ACEI or ARB initiating treatment. Consistently, several randomized studies demonstrated a favorable or neutral effect of nebivolol in patients with dysglycemia.

We here review data from the first, and to the best of our knowledge only, study on the efficacy and safety of nebivolol and valsartan in a FDC in hypertensive patients. The results from this phase 3, randomized, double-blind, placebo-controlled, multicentre study were recently published in the Lancet and were commented on in editorial coverage focusing on hypertension as the biggest contributor to global morbidity and mortality.

In this study, hypertensive adult patients with BP lower than 180/110 mmHg were randomly assigned to 4 weeks of double-blind treatment with nebivolol and valsartan FDC (5 and 80 mg/day, 5 and 160 mg/day, or 10 and 160 mg/day), nebivolol (5 mg/day or 20 mg/day), valsartan (80 mg/day or 160 mg/day), or placebo. After the first month, the doses were doubled for the next 4 weeks. After 8 weeks of receiving the respective treatment, it was observed that the reductions in diastolic and systolic BP from baseline, as well as the percentage of patients responding to treatment or achieving BP target goal (<140/90 mmHg in nondiabetics and <130/80 mmHg in diabetics), were greater in the highest FDC (20 and 320 mg/day) when compared with the highest doses of monotherapy, nebivolol 40 mg/day or valsartan 320 mg/day, respectively. A lower-dose combination also showed more favorable outcomes when compared with the respective monotherapy doses. Importantly, the rates of treatment-emerged adverse events were comparable across the treatment groups with the exception of the highest nebivolol dose (40 mg/day). Thus, the placebo-like adverse event profiles of both nebivolol and valsartan shown in previous studies have been maintained in the FDC as well. In addition, a high response to placebo in Hispanic and elderly patients, as well as the relatively small number of elderly participants, prevented a critical conclusion of the combination efficacy for these subgroups of patients. Nevertheless, the results of this study suggest that FDC of nebivolol and valsartan offer better BP efficacy and comparable tolerability to monotherapies. In accord with potential better adherence to the single-pill treatment, the study outcomes underline that these combinations should be consider as a safe and effective treatment for hypertensive patients.

Consideration for nebivolol/valsartan combination in the treatment of target organ disease

An increasing body of evidence suggests that target organ damage depends more on central than peripheral arterial pressure. In that regard, both nebivolol and valsartan have been shown to reduce central BP and central pulse pressure. In contrast to nebivolol, despite a similar reduction in brachial artery pressure in patients with uncomplicated hypertension, metoprolol and atenolol affected neither aortic pulse pressure nor augmentation index, which are important hemodynamic parameters particularly in the elderly population, in whom arterial compliance is already affected. Pulse wave velocity, as a measure of aortic stiffness, was similarly reduced by nebivolol and atenolol or
metoprolol, and the difference in central pressure may have mirrored a favorable effect on wave reflection as a result of less reduction in heart rate and peripheral vasodilatory effect by nebivolol. Yet in another study, the EFFORT study, with predominantly African-American patients with type 2 diabetes and hypertension who are treated with maximally tolerated doses of RAS blockade, nebivolol did not offer a significant reduction in aortic BP over metoprolol.62 Thus, further studies are necessary to clarify a difference between nebivolol and traditional β-blockers with respect to the beneficial effects on central arterial pressure.

A number of smaller studies suggest left ventricular ejection fraction (LVEF) benefits of nebivolol treatment in patients with HF,74,108,109 with one study describing an effect similar to that of carvedilol.109 Data from an 8 month randomized trial involving elderly patients with HF (mean LVEF, 26%; N=260) suggest a relative improvement of LVEF by approximately 36% among the nebivolol-treated patients compared with 19% in the placebo group.62 The Study of the Effects of Nebivolol Intervention on Outcomes and Rehospitalization in Seniors with Heart Failure (SENIORS trial)73 addressed for the first time the value of β-blockers in elderly subjects (≥70 years of age). Patients with HF (mean LVEF, 36%; N=2,128) and a mean follow-up duration of 21 months demonstrated that nebivolol treatment was associated with a significantly lower risk for all-cause mortality or CV-related hospital admission compared with placebo treatment (31.3% versus 35.3%; P=0.039). Of interest is the finding that the greatest benefits were found in women.73 The positive results achieved with nebivolol in the SENIORS trial led to the inclusion of nebivolol as one of four β-blockers recommended for HF in the 2008 guidelines of the European Society of Cardiology.110,111

In patients with congestive HF, enhanced activity of both renin angiotensin and sympathetic system activities contribute to the evolution of the syndrome. Accordingly, a large body of clinical trials has shown the benefits of both β-blockers and RAS inhibitors in patients with cardiac remodeling or HF. The results of the Valsartan in Heart Failure Trial (Val-HeFT)112 and the Valsartan in Acute Myocardial Infarction (VALIANT)113 trial showed that valsartan is an effective treatment for patients with HF and in patients with postmyocardial infarction. On the basis of the results of the SENIOR study, nebivolol is safe and effective in elderly patients with HF, some of whom have mildly reduced or preserved ejection fraction.73,75,76 Taking into account the limited effectiveness of traditional RAS inhibitors in the treatment of HF with preserved ejection fraction, particularly in women,40 these results suggest a great potential of nebivolol in the treatment of these patients. Vinereanu and colleagues114 compared the effects of 6 months of therapy with either nebivolol or metoprolol on the left ventricular function of hypertensive patients. They found that mean longitudinal early diastolic velocity increased by 16% (P<0.05) with nebivolol compared with a nonsignificant 9% change with metoprolol.114 In addition, flow propagation velocity increased by 34% on nebivolol and did not change on metoprolol.114 Associated improvements in mean longitudinal displacement and ejection time were only observed in the hypertensive patients randomized to nebivolol treatment.114 Because the antihypertensive effect was comparable in both groups of treated subjects, the data suggest a unique action of nebivolol in reducing cardiac stiffness, and possibly cardiac fibrosis. This interpretation is in keeping with our demonstration of improved diastolic relaxation and myocardial fibrosis in a renin-dependent model of hypertension after treatment with nebivolol.56 Thus, combination therapy with nebivolol and valsartan may offer additional complementary mechanisms in decreasing afterload, improving coronary artery flow, and remodeling and preventing progression of HF.

Our recent studies115,116 on the BP-independent effects of nebivolol on cardiac structure and function of hypertensive rats fed a high-salt diet further promote the idea that nebivolol in combination with RAS blockade would have been more beneficial in preventing salt-related cardiac dysfunction. However, additional studies will be needed to validate these early clinical and experimental findings, as nebivolol did not improve impaired ventricular diastolic function, nor did it reduce left ventricular mass in an experimental model of hypertension induced by high salt feeding. In this experimental hypertension model, neither nebivolol nor Ang II blockade lowers BP, even though it has a significant effect in reversing cardiac collagen deposition.116

Consistent with small-scale clinical studies showing that nebivolol prevented contrast-induced nephropathy,117,118 and improved renal function in the postvascularization phase in patients with renal artery stenosis,119 we demonstrated that nebivolol improved renal blood flow and prevented excessive urinary protein loss, collagen deposition, and associated nitro-oxidative (nitrosative) stress while suppression plasma renin in salt-loaded spontaneously hypertensive rats.115 These studies agree with the demonstration that nebivolol improved proteinuria through reductions in renal nicotinamide adenine dinucleotide phosphate oxidase/reactive oxygen species and increases in bioavailable NO in a renin-dependent hypertension model.120 Valsartan is a powerful antihypertensive and renoprotective drug in an
experimental model of Ang II-dependent hypertension and renal injury. It seems that the combination of these two drugs may be advantageous in terms of renoprotection, considering that nebivolol may reduce the reactive increase in renin after Ang II blockade.

In summary, nebivolol, a third-generation β-blocker, is a potent antihypertensive agent that is essentially devoid of clinically important adverse effects. Distinctive mechanisms of action in terms of hemodynamics, metabolic response, and NO-dependent activation of cardiac and vascular vasodilator and antifibrotic mechanisms make this member of the β-blocker class unique. Emerging studies in which nebivolol is combined with an ARB in a FDC should be of significant value in terms of increased patient compliance, enhanced BP-lowering actions, and reversal of the adverse cardiorenal and vascular remodeling adaptation to hypertension.

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