Nebivolol in the treatment of chronic heart failure

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Abstract: Nebivolol is a highly selective beta1-adrenergic blocker that also enhances nitric oxide bioavailability via the L-arginine-nitric oxide pathway, leading to vasodilation and decreased peripheral vascular resistance. It is marketed in Europe for the treatment of hypertension and heart failure and is currently being reviewed for use in the US by the Food and Drug Administration. Nebivolol appears to be well tolerated with an adverse event profile that is at least similar, if not better, than that of other beta-adrenergic blockers. Studies suggest that long-term therapy with nebivolol improves left ventricular function, exercise capacity, and clinical endpoints of death and cardiovascular hospital admissions in patients with stable heart failure. To date, it is one of the only beta-adrenergic blockers that have been exclusively studied in elderly patients. Additionally, the unique mechanism of action of nebivolol makes it a promising agent for treatment of chronic heart failure in high-risk patient populations, such as African Americans. This article will review the pharmacologic and pharmacokinetic properties of nebivolol as well as clinical studies assessing its efficacy for the treatment of heart failure.

Keywords: nebivolol, beta-adrenergic blockers, heart failure

Introduction

The pathophysiology of chronic heart failure involves a process of left ventricular remodeling, whereby molecular changes occur within the myocardium in response to mechanical stresses induced by underlying diseases, such as hypertension, ischemic heart disease, cardiomyopathies, and valvular abnormalities. Structural changes that occur, including left ventricular hypertrophy and/or dilation, typically result in decreased left ventricular diastolic or systolic function (Jessup and Brozena 2003; Opie et al 2006). This remodeling process is accelerated by the activation of a number of endogenous neurohormonal systems including, but not limited to, the sympathetic nervous system, which releases high levels of the adrenergic substance norepinephrine and stimulates the release of renin in the kidney (Jessup and Brozena 2003; Hunt et al 2005). The resultant increase in heart rate, contractility, peripheral vasoconstriction, and blood volume, as well as the direct toxic effects of norepinephrine on myocytes, increases cardiac workload and further impairs cardiac performance (Hunt et al 2005). Beta-adrenergic blockers, by suppressing the deleterious effects of norepinephrine have become routine therapy for the treatment of chronic heart failure (Hunt et al 2005; McMurray et al 2005; Swedberg et al 2005).

The benefits of beta-adrenergic blockers in the treatment of chronic heart failure are exclusive to those agents that have demonstrated a survival benefit in clinical trials and should not, therefore, be considered a class effect. In the US, three beta-adrenergic blockers are currently available for use in chronic heart failure based on evidence demonstrating a survival benefit: carvedilol, which blocks alpha1-, beta1-, and beta2-receptors; and sustained-release metoprolol succinate and bisoprolol, which both selectively block beta1-receptors (Packer et al 1996; CIBIS II Investigators 1999; Hjalmarson et al 1999; Packer et al 2001; Hunt et al 2005). Nebivolol is a third-generation beta-adrenergic blocker that has been marketed and used in Europe...
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for the treatment of hypertension and heart failure (A. Menarini Pharmaceuticals 2005; McMurray et al. 2005). It is currently under FDA review in the US for hypertension and it is anticipated that an indication for heart failure will be pursued in the near future.

**Pharmacology**

Nebivolol is a racemic mixture containing equal amounts of 2 isomers, d-nebivolol and l-nebivolol. D-nebivolol provides selective beta₁-adrenergic receptor blockade while both d- and l-nebivolol cause nitric oxide-induced vasodilation (Cockcroft et al. 1995; Van Neuten 1998). Nebivolol, which has no intrinsic sympathomimetic activity, is considered a highly selective beta₁-adrenergic blocker due to its 321-fold higher affinity for human cardiac beta₁-receptors versus beta₂-receptors; it is also more selective for beta₁-receptors than any other agent in its class (Brixius et al. 2001; Bristow et al. 2005). Unlike other beta-adrenergic blockers with vasodilatory properties, nebivolol has no alpha-blocking effects (Bowman et al. 1994; Van Bortel et al. 1997). The vasodilatory action of nebivolol is mediated via the L-arginine-nitric oxide pathway, whereby nitric oxide production by endothelial nitric oxide synthases is enhanced (Bowman et al. 1994; Cockcroft et al. 1995; Ignarro 2004). There is evidence to suggest that this mechanism is in part due to agonist activity of nebivolol at endothelial beta₁-adrenergic receptors (Figure 1) (Gauthier et al. 1998; Gosgnach et al. 2001; Dessy et al. 2005). This was recently tested and confirmed by Dessy and colleagues who established that nebivolol relaxation of human coronary microarteries that were precontracted with endothelin-1 was significantly inhibited by bupranolol, a beta₁,₂,₃-receptor blocker, but not...

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**Figure 1** The effect of nebivolol on the L-arginine-nitric-oxide pathway. Reprinted with permission Veverka A, Nuzum DS, Jolly JL. 2006. Nebivolol: a third-generation beta-adrenergic blocker. *Ann Pharmacother, 40*:1353–60. Copyright © 2006, Harvey Whitney Books.

**Abbreviations:** NOS, nitric oxide synthase; GTP, guanosine triphosphate; cGMP, cyclic guanosine monophosphate.
Nebivolol significantly inhibited by nadolol, a beta₁,₂-receptor blocker (Dessy et al. 2005). Nitric oxide bioavailability may also be augmented with nebivolol treatment due to decreased inactivation by reactive oxygen entities (Janssen et al. 2001; Cominacini et al. 2003; Pasini et al. 2005).

This unique mechanism of nebivolol is particularly important due to the critical role of nitric oxide in the pathophysiology of cardiovascular diseases, including heart failure. In addition to causing vasodilation, nitric oxide also inhibits platelet aggregation, atherosclerosis and proliferation of vascular smooth muscle cells (Mason 2006). When endothelial dysfunction occurs, nitric oxide production and function is impaired, leading to increased peripheral resistance and a pro-thrombotic and pro-atherogenic environment (Panza et al. 1990; Moncada and Higgs 1993; Kinlay et al. 2001; Mason 2006). To distinguish the vasodilatory action of nebivolol from other beta-adrenergic blockers that selectively inhibit beta₁-receptors, Lekakis and colleagues tested the effect of nebivolol and atenolol on flow-mediated dilation of the brachial artery (Lekakis et al. 2005). Following 4 weeks of drug therapy, patients treated with nebivolol had significantly increased flow-mediated dilation, while those treated with atenolol had no change compared to baseline.

The increased bioavailability of nitric oxide with nebivolol treatment may prove to be particularly useful for treating African American patients with cardiovascular disease. It has been proposed that decreased endothelial nitric oxide bioavailability may be more prevalent as an underlying cause of cardiovascular disease in this patient subgroup, and previous studies of African American patients with heart failure have shown a favorable response to therapies that increase nitric oxide availability (Taylor et al. 2004). The mechanism of decreased nitric oxide bioavailability in African American patients may be due to oxidative stress caused by upregulation of NAD(P)H-dependent oxidases and subsequent increases in production of superoxide (O₂⁻). O₂⁻ can react with nitric oxide, decreasing its bioavailability and increasing production of the oxidant peroxynitrite (ONOO⁻) (Kalinowski et al. 2004). Mason and colleagues compared the activity of nebivolol and atenolol on nitric oxide release from endothelial cells of age-matched African American and Caucasian donors with comparable cardiovascular risk histories (Mason et al. 2005). Levels of nitric oxide, as well as ONOO⁻ and O₂⁻, the primary components of nitrooxidative and oxidative stress in the vascular system, were measured to assess endothelial function. At baseline, release of nitric oxide was 5 times slower, and release of both ONOO⁻ and O₂⁻ was 2–4 times faster in African Americans compared to Caucasians. While atenolol had no effect on nitric oxide, ONOO⁻, and O₂⁻ levels in either white or black patients, nebivolol treatment increased nitric oxide and reduced ONOO⁻ and O₂⁻ levels in African Americans to similar levels documented in Caucasian patients.

### Pharmacokinetics and drug interactions

Table 1 summarizes the general pharmacokinetic properties of nebivolol and other beta-adrenergic blockers typically used in the management of heart failure (Frishman and Alwarshetty 2002; Eon Labs, Inc. 2004; AstraZeneca.

| Characteristic       | Bisoprolol (Zebeta) | Carvedilol (Coreg) | Metoprolol succinate (Toprol XL) | Nebivolol |
|----------------------|---------------------|--------------------|----------------------------------|-----------|
| Absorption           | Bioavailability     | 80%                | 25%–35%                          | 50%       |
|                      | First-pass elimination | Small             | Significant                       | Moderate |
|                      | Effect of food      | None               | Decreases rate but not extent     | None      |
|                      | Protein binding     | 30%                | 95%–98%                          | 12%–96%   |
|                      | Half-life (hours)   | 9–12               | 6–10                             | 98%       |
| Hepatic metabolism   | 50% to inactive metabolites via N-dealkylation and O-dealkylation | Extensive primarily by CYP450 2D6 and 2C9 to active and inactive metabolites<sup>a</sup> | Extensive via CYP450 2D6 to active and inactive metabolites<sup>a</sup> | Extensive via CYP450 2D6 to active and inactive metabolites<sup>a</sup> |
| Renal excretion      | 50% as unchanged drug, 50% as metabolites | <2% as unchanged drug | 95%, <5% as unchanged drug | <1% unchanged in urine |
| Other excretion      | <2% in feces        | Primarily in bile and feces | Minimal                           |           |

<sup>a</sup>CYP450 = cytochrome P450.
<sup>b</sup>Carvedilol is metabolized to a lesser extent by CYP 450 3A4, 2C19, 1A2, and 2E1.
<sup>c</sup>Bioavailability and first-pass elimination are dependant on cytochrome P450 2D6 genetic polymorphism.
L.P. 2006; GlaxoSmithKline 2007). Nebivolol is rapidly absorbed following oral administration, reaching peak plasma concentrations within 0.5–4 hours after a dose (Sule and Frishman 2006). Food has a minimal impact on absorption and therefore nebivolol may be taken without regard to meals (Shaw et al 2003a). Nebivolol is extensively metabolized via hydroxylation in the hepatic system to active and inactive metabolites. The oral bioavailability of nebivolol is dependent on cytochrome P450 2D6 genetic polymorphism and so ranges from 12% in extensive metabolizers to 96% in poor metabolizers. Similarly, the half-life of nebivolol is approximately 10 hours in extensive metabolizers but can be prolonged up to 30–50 hours in poor metabolizers (Van Peer et al 1991; A. Menarini Pharmaceuticals 2005). Despite genetic differences in metabolism of nebivolol, the clinical response to the drug appears to be similar (Lefebvre et al 2006). Nebivolol displays linear kinetics across a dose range of 2.5–20 mg, demonstrated by dose-proportional changes in maximum concentrations (Cmax) and area under the drug concentration curve (AUC) (Shaw 2003b). The average volume of distribution of nebivolol is 10 L/kg and this does not appear to be affected by patient weight (Cheymol et al 1997). Less than 1% of the drug is excreted unchanged in the urine and so adjustments of doses in patients with chronic renal failure are unnecessary (A. Menarini Pharmaceuticals 2005).

Nebivolol is highly protein bound intravascularly, predominantly to albumin. Studies assessing drug interactions with nebivolol in healthy volunteers have found no significant interactions with spironolactone, hydrochlorothiazide, digoxin, warfarin, losartan, and ramipril (Lawrence et al 2003; Morton et al 2003, 2005; Lawrence et al 2005a, b, c). Co-administration with cimetidine, a potent inhibitor of cytochrome P450 3A4, increased the bioavailability of nebivolol, however this interaction did not influence the extent to which nebivolol reduced heart rate and blood pressure (Kamali et al 1997). Similarly, fluoxetine, a cytochrome P450 2D6 inhibitor, resulted in peak plasma concentrations of nebivolol that were three times higher than normal (Shaw 2005). Although the clinical impact of cytochrome P450 drug interactions with nebivolol is unclear, caution should be exercised when inhibitors or inducers of 2D6 and 3A4 are used in conjunction with this agent. At this time, it is also unknown whether nebivolol is a substrate of p-glycoprotein and if there is a risk of drug interactions at this protein.

Clinical studies
Earlier studies assessing the utility of nebivolol in chronic heart failure were limited by small patient populations. These studies did suggest, however, that nebivolol would improve left ventricular function and mechanics; improve patient functional capacity assessed by New York heart association (NYHA) classification; and would at least have a stabilizing effect on exercise capacity (Uhlir et al 1997; Brehm et al 2002). More recently, the ENECA (efficacy of nebivolol in the treatment of elderly patients with chronic heart failure as add-on therapy to ACE inhibitors or angiotensin II receptor blockers, diuretics, and/or digitalis) study performed by Edes and colleagues evaluated whether nebivolol therapy improves left ventricular ejection fraction (LVEF) compared with placebo in 260 patients with chronic heart failure (Edes et al 2005). The study design also included a secondary endpoint to assess the safety and tolerability of nebivolol in elderly patients, defined in the study as age greater than 65. In addition to the age requirement, patients qualified for enrollment in the study if they met the following criteria: NYHA class II, III, or IV; LVEF less than or equal to 35%; stable clinical status; and stable therapy for at least 2 weeks prior to randomization with angiotensin-converting-enzyme inhibitors (ACEIs) and/or angiotensin receptor blockers (ARBs), diuretics, and/or digitalis. Patients were randomized to therapy with nebivolol 1.25 mg daily, titrated to a target dose of 10 mg, or placebo and followed for a period of 8 months. Intention to treat analysis showed that nebivolol therapy significantly increased LVEF compared with placebo (improvement of 6.51 ± 9.15% vs 3.97 ± 9.2% from baseline, respectively; p = 0.027), and this was consistent across all subgroups examined. Quality of life and changes in NYHA functional class were not significantly improved with nebivolol therapy in this trial. While nebivolol was generally well tolerated in this elderly population and did not result in an increased number of patients experiencing adverse events compared with placebo (81 vs 78, respectively; p = 0899), drug-related adverse events were more commonly reported with nebolol vs placebo (40 vs 14; p < 0.0001). The most frequent of these were hypotension, bradycardia, and dizziness. Despite proving in the ENECA study that nebivolol is superior to placebo in improving surrogate endpoints of chronic heart failure, this study was underpowered to assess the effect of nebolol on clinical endpoints such as overall survival, cardiovascular death, or need for cardiovascular hospital admission.

Subsequent to the ENECA study, Flather and colleagues published the results of the SENIORS (study of the effects of nebolol intervention on outcomes and rehospitalization in seniors with heart failure) trial (Flather et al 2005). This was the first and is the only randomized, double-blind,
placebo-controlled trial to date assessing the benefit of nebivolol therapy on morbidity and mortality. The trial enrolled 2135 elderly patients, defined as 70 years of age or older, with a clinical history of heart failure, defined as a documented LVEF less than or equal to 35% within the previous 6 months or hospitalization with a discharge diagnosis of chronic heart failure within the previous 12 months. Patients were enrolled provided they were not currently receiving therapy with a beta-adrenergic blocker or had had a contraindication to treatment. The primary outcome of the SENIORS trial was a composite of all-cause mortality or cardiovascular hospital admission. In the nebivolol arm, doses of 1.25 mg daily were initiated and titrated over a 16 week period to a target dose of 10 mg once daily. Over a mean treatment period of 21 months, 31.1% of patients in the nebivolol group reached the primary endpoint compared with 35.3% in the placebo group (hazard ration [HR] 0.86, 95% confidence interval [CI] 0.74–0.99, p = 0.039). These results imply that 24 patients with chronic heart failure would need to be treated with nebivolol for approximately 2 years to prevent one death or cardiovascular hospital admission. Of note, this benefit was observed as early as 6 months and was independent of baseline therapy with diuretics, ACEIs, digoxin, and/or spironolactone which were used by approximately 85%, 82%, 40%, and 38% of patients enrolled, respectively. Subgroup analysis determined that nebivolol was efficacious regardless of age, gender, ejection fraction, diabetes, or prior myocardial infarction. At the present time, the SENIORS trial is the only assessment of beta-adrenergic blocker therapy in an elderly population with chronic heart failure. This may be of particular importance in clinical practice since the prevalence of heart failure increases with age, from 2% to 3% at age 65 years to greater than 80% in patients aged 80 years and above (Hunt et al 2005). Additionally, treatment of elderly patients with beta-adrenergic blockers can be more challenging due to desensitization of beta-adrenergic receptors and variable pharmacokinetic responses that occur with age (potentially decreased absorption, metabolism and excretion) (Tregaskis and McDevitt 1990; Frishman and Alwarshetty 2002). While adverse outcomes have been documented when standard therapy for heart failure is insufficient, it is important to individualize therapy for each individual patient (Komajda et al 2005).

There are very few head-to-head comparisons of beta-adrenergic blockers for the treatment of chronic heart failure. Nebivolol has been compared with carvedilol in two small trials. Patrianakos and colleagues assessed the effects of carvedilol and nebivolol on left ventricular function and exercise capacity at 3 and 12 months. Seventy-two patients with NYHA class II or III heart failure, specifically non-ischemic dilated cardiomyopathy documented by a LVEF of less than 45% on echocardiogram within the previous 6 months (Patrianakos et al 2005) were included. Patients were randomized to double-blind therapy with either carvedilol 3.125 mg twice daily or nebivolol 1.25 mg once daily, with titration to carvedilol 25 mg twice daily or nebivolol 5 mg daily as tolerated. Additional requirements for enrollment included stable therapy with an ACEI or ARB for at least 4 weeks prior to randomization with no new drug therapies initiated within 6 weeks prior to randomization. No patients enrolled in the study had received prior treatment with a beta-adrenergic blocker. At 3 and 12 months, both nebivolol and carvedilol caused significant improvements in LVEF compared with baseline. Inter-group comparisons, however, revealed that carvedilol provided a greater change in LVEF than nebivolol at these time points (3 months: absolute improvement of 7.4% vs 4.8%; relative improvement of 32.1% ± 34.9% vs 15.3% ± 15.9%, mean difference –16.7% ± 16.5%, 95% CI –29.9 to –3.4, p = 0.004; 12 months: absolute improvement of 8.8% vs 6.1%; relative improvement of 35.5% ± 31.9% vs 20.7% ± 19.1%, mean difference –14.7% ± 6.4%, 95% CI –27.8 to –1.8, p = 0.02). Both agents significantly decreased left ventricular end-systolic volumes at 3 and 12 months and although only carvedilol improved left ventricular end-diastolic volumes compared to baseline, inter-group analysis showed no statistically significant differences in left ventricular volumes across the treatment period. Diastolic function, assessed by ventricular relaxation and filling patterns, was significantly improved at 12 months with both nebivolol and carvedilol therapy; however, only carvedilol demonstrated a benefit as early as 3 months.

Exercise duration, measured in seconds, significantly improved at 12 months with both nebivolol (894 ± 381 at baseline vs 994 ± 396 at 12 months; p = 0.01, 95% CI –181 to –18) and carvedilol (982 ± 475 at baseline vs 1124 ± 427 at 12 months; p = 0.01, 95% CI –248 to –36), with no statistically significant differences observed between the two groups. Of note, there was an initial decline in exercise capacity detected at 3 months with nebivolol (894 ± 381 at baseline vs 795 ± 392 at 3 months; p = 0.07, 95% CI –20 to –209). Although this was not statistically significant, this effect was not seen in the carvedilol group (982 ± 475 at baseline vs 1025 ± 419 at 3 months; p = 0.26, 95% CI –120 to –33) and compared with nebivolol, exercise capacity at 3 months was significantly better with carvedilol therapy (p = 0.002, 95% CI 0.03–0.48). One explanation for the
initial decline in exercise capacity with nebivolol could be too rapid titration of the drug to target doses, which was accomplished over 4 weeks, a much faster titration than used in the SENIORS trial. Although, this study appears more favorable for carvedilol, a subsequent trial published by Lombardo and colleagues found conflicting results (Lombardo et al 2006). A similar patient population, 70 patients with NYHA class II or III heart failure and LVEF less than or equal to 40%, were randomized to carvedilol and nebivolol at similar doses used in the aforementioned trial. Patients were evaluated at baseline, 3, and 6 months, but data for baseline and 6 months only were reported. In contrast to the study by Patrianakos and colleagues, increases in LVEF and decreases in left-ventricular end-systolic volumes observed at 6 months were not statistically different from baseline; nor was there a difference observed between groups. Both carvedilol and nebivolol showed a trend towards an increased exercise capacity at 6 months and there was no reported decline in exercise capacity with nebivolol at earlier assessments. Both of these trials enrolled a small number of patients and evaluated surrogate endpoints. The study by Patrianakos and colleagues was performed in patients with non-ischemic dilated cardiomyopathy so extrapolation to the general heart failure population is inappropriate given that ischemic heart disease is one of the most common causes of chronic heart failure. Additionally, target doses of nebivolol 5 mg used in these comparator trials is lower than the 10 mg target dose used in the ENECA study and SENIORS trial discussed above. Any differences between nebivolol and carvedilol on clinical endpoints of mortality and hospitalizations for heart failure cannot be inferred from these trials. Larger trials with head-to-head comparisons of nebivolol, carvedilol, metoprolol succinate, and bisoprolol are needed to further establish if one agent is any more beneficial than the others in increasing survival and decreasing hospitalizations for acute decompensated heart failure.

### Conclusion

Nebivolol is currently marketed in Europe for the treatment of hypertension and heart failure and is under FDA review for use in the US. Evidence shows that nebivolol, titrated to a maximum dose of 10 mg, is a potentially promising therapeutic option for the treatment of chronic heart failure when added to standard therapy. Its unique mechanism of selectively blocking beta1-receptors and decreasing peripheral vascular resistance by enhancing nitric oxide bioavailability distinguish it from other agents in its class; however the clinical significance of this still needs to be defined. Since not all beta-adrenergic blockers have proved to be effective for the treatment of heart failure, the addition of nebivolol to the current armamentarium of carvedilol, metoprolol succinate, and bisoprolol is encouraging and provides more options for individualizing patient therapy. Specifically, and in light of recent evidence for other agents known to work via the nitric oxide pathway, nebivolol may prove to be more useful than other beta-adrenergic blockers in African American patients and those suspected of having decreased nitric oxide bioavailability as an underlying pathophysiology of disease (Taylor et al 2004). Large-scale clinical trials are needed, however, to test this hypothesis. Additionally, without head-to-head trials assessing mortality or hospitalizations for decompensated heart failure with nebivolol, it is premature to comment on which beta-adrenergic blocker is preferred for heart failure management. Nebivolol is the only agent to date that has evidence supporting use of beta-adrenergic blockers in the treatment of elderly patients with chronic heart failure. At this time, numerous clinical studies with nebivolol are in progress and include: an assessment of the role of nebivolol in the treatment of diastolic heart failure.
failure; use of nebivolol in African American patients with hypertension; and a comparison of nebivolol and metoprolol in patients with subclinical left ventricular dysfunction (The Menarini Group 2007).

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