Understanding the early mortality benefit observed in the PARADIGM-HF trial: considerations for the management of heart failure with sacubitril/valsartan

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Abstract: This review aims to elucidate the optimal dosing of angiotensin receptor-neprilysin inhibitor (ARNI) therapy in the heart failure (HF) treatment paradigm through examination of the trial population characteristics and the mortality benefit observed in the Prospective Comparison of ARNI with angiotensin-converting enzyme inhibitor (ACEI) to Determine Impact on Global Mortality and Morbidity in Heart Failure (PARADIGM-HF; NCT01035255) trial. Considerations regarding the initiation and titration of sacubitril/valsartan, a first-in-class ARNI, will also be addressed. The approval of sacubitril/valsartan heralded the first novel pharmacological class in over a decade for the treatment of heart failure with reduced ejection fraction (HFrEF). The PARADIGM-HF trial showed that treatment with valsartan/valsartan reduced the risk of first occurrence of either cardiovascular death or HF-related hospitalization (composite primary endpoint) by 20% compared with enalapril in patients with HFrEF. The incremental benefits of treatment with valsartan/valsartan over enalapril demonstrated in the PARADIGM-HF trial led to strong recommendations for its use over ACEIs or angiotensin receptor blockers to further reduce morbidity and mortality in the 2016 and 2017 American College of Cardiology/American Heart Association/Heart Failure Society of America updates to the guidelines for the management of HF. Although the optimal timing for the initiation of valsartan/valsartan has yet to be determined, its early use is likely to have a positive impact on patient outcomes.

Keywords: angiotensin receptor-neprilysin inhibitor, cardiovascular death, patient outcomes, treatment recommendations

Introduction

Heart failure (HF) represents a substantial health care burden in the United States, related to 1 in 9 deaths.1,2 From 2012 to 2030, its prevalence has been projected to increase by 46%, affecting more than 8 million Americans.1,2 The pathophysiologic mechanisms underlying HF development and progression are complex, predominantly involving increased activation of both the renin–angiotensin–aldosterone system (RAAS) and the sympathetic nervous system (SNS). The activation of these systems is counterbalanced by endogenous peptides (eg, natriuretic peptides), which are released in response to myocardial stretch that results from excessive neurohormonal activity.3–5

Lifestyle interventions aimed at risk reduction comprise an important strategy for preventing HF and delaying or reversing disease progression following its onset.6 However, when symptomatic HF develops, pharmacotherapy is typically warranted.6–8
Response to pharmacologic therapies for HF differs depending on cardiac function, which is determined by measurement of ejection fraction (EF). Accordingly, EF has been used to classify patients as having either HF with reduced EF (EF<40%; HFrEF) or HF with preserved EF (EF>50%; HFpEF), although many HF clinical trials have used a lower threshold of EF≤35% to define HFrEF. In symptomatic patients with HFrEF, pharmacologic therapies targeting the overactive RAAS and SNS become necessary. Agents targeting these pathways, including angiotensin-converting enzyme inhibitors (ACEIs), angiotensin receptor blockers (ARBs), β-blockers, and mineralocorticoid receptor antagonists (MRAs), have been the mainstays of HFrEF treatment since the 1990s.

Although mortality rates associated with HF have improved with the development of these therapeutic options and the use of guideline-directed evaluation and management (GDEM), approximately 50% of the patients with HF still do not survive 5 years past diagnosis. Thus, new therapies that can further improve survival rates and patient outcomes in HFrEF are required. The aim of this review is to discuss the implications of the introduction of one such new therapy—the angiotensin receptor-neprilysin inhibitor (ARNI) pharmacological class—and to explore whether use of this therapy earlier in the HFrEF treatment paradigm may ultimately lead to an overall improvement in patient outcomes, particularly mortality.

Established conventional therapies for HF
The traditional pharmacological treatments of GDEM for HFrEF—β-blockers, ACEIs/ARBs, and MRAs—inhibit the activation of the SNS and RAAS, effectively resisting the development and progression of HF in two main ways: 1) control of blood pressure and left ventricular (LV) afterload and 2) reversing impairments caused by the effects of the over-activated RAAS and SNS on pathologic ventricular remodeling. In addition, to decrease the risk of hospitalization for HF, digoxin and/or the If current inhibitor ivabradine can be added to standard GDEM in patients who have persistent symptoms of HFrEF.

The goal of optimal management of comorbid hypertension is the reduction of afterload on the heart, thereby preventing the cardiac remodeling and fibrosis that results from impaired ventricular function. Studies have demonstrated that effective hypertensive control reduces the incidence of HF by approximately 50%, with the most effective therapeutic classes in this regard being β-blockers, ACEIs, and diuretics. Even once HF is established, hypertension management remains a critical treatment goal. The use of β-blockers and ACEIs has also been shown to reverse impairments in LV function caused by pathologic remodeling, which has been shown to contribute to HF progression and risk of cardiovascular (CV) events. This reversal has been demonstrated in symptomatic and asymptomatic patients with LV dysfunction but no overt HFrEF. Morbidity and mortality benefits from guideline-directed pharmacological therapies for HFrEF have been demonstrated in clinical trials (summarized in Table 1).

### β-blockers
β-blockers—specifically bisoprolol, carvedilol, and long-acting metoprolol—have consistently been shown to significantly and incrementally lower mortality risk when added to optimized GDEM for patients with HFrEF.

### ACEIs and ARBs
Although enalapril was the first ACEI to demonstrate mortality benefits for patients with HF, a meta-analysis conducted in 1995 found similar benefits with other ACEIs, including captopril, ramipril, quinapril, and lisinopril. The use of ACEIs in patients with normal or controlled blood pressure, but who are at high risk for developing HF (stage A HF), has also been shown to reduce the risk for developing HF, as was observed in the Studies of Left Ventricular Dysfunction (SOLVD) prevention trial (NCT00000516). In this trial, patients with asymptomatic LV dysfunction had an annualized 10% risk for developing HF and an 8% risk for death or HF-related hospitalization despite the absence of significant symptoms. Intervention with the ACEI enalapril versus placebo at this early stage, prior to development of overt HF, prevented the development of HF and improved clinical outcomes in patients with LV dysfunction.

In 2000, losartan became the first ARB to demonstrate similar mortality benefits and improved tolerability compared with ACEIs in patients with New York Heart Association (NYHA) class II–IV HFrEF. Soon after, the addition of the ARB valsartan to background therapy (which included an ACEI in approximately 83% of the patients) was also shown to reduce the risk for the combined endpoint of morbidity and mortality compared with placebo, although it did not reduce the risk for all-cause mortality. However, the high number of discontinuations with ARB therapy added to ACEI therapy led to recommendations against their combined use. Accordingly, treatment with ARBs is typically recommended for...
Table 1 Morbidity and mortality benefits of pharmacologic therapies for heart failure

| Pharmacological therapy | Mortality endpoint | Other key endpoints |
|-------------------------|--------------------|---------------------|
| β-blockers              |                    |                     |
| Carvedilol vs placebo   | RR, 0.35; 95% CI, 0.20–0.61; P<0.001 | Mortality or hospitalization for CV disease: RR, 0.62; 95% CI, 0.47–0.82; P<0.001 |
| Metoprolol CR/XL vs placebo | RR, 0.66; 95% CI, 0.53–0.81; P=0.00009 | CV-related mortality: RR, 0.62; 95% CI, 0.50–0.78; P=0.00003 |
| Bisoprolol vs placebo   | HR, 0.66; 95% CI, 0.54–0.81; P<0.0001 | CV death or CV-related hospitalization: HR, 0.79; 95% CI, 0.69–0.90; P=0.0004 |
| ACEIs                   |                    |                     |
| Enalapril vs placebo    | RR, 0.73; P=0.003 | Mortality due to HF: RR, 0.50; P<0.001 |
| Enalapril vs placebo    | RR, 0.84; 95% CI, 0.74–0.95; P<0.0072 | Mortality or HFH: RR, 0.74; 95% CI, 0.66–0.82; P<0.001 |
| ARBs                    |                    |                     |
| Losartan vs captopril   | HR; 1.13; 95.7% CI, 0.95–1.35; P=0.16 | Mortality or hospitalization: HR, 1.07; 95% CI, 0.97–1.19; P=0.18 |
| valsartan vs placebo    | RR, 1.02; 98% CI, 0.88–1.18; P=0.80 | Morbidity or mortality: RR, 0.87; 97.3% CI, 0.77–0.97; P=0.009 |
| MRAs                    |                    |                     |
| Spironolactone vs placebo | RR, 0.70; 95% CI, 0.60–0.82; P<0.001 | Cardiac-related mortality or hospitalization: RR, 0.68; 95% CI, 0.59–0.78; P<0.001 |
| Eplerenone vs placebo   | Adjusted HR, 0.76; 95% CI, 0.62–0.93; P=0.008 | CV-related mortality or HFH: adjusted HR, 0.63; 95% CI, 0.54–0.74; P<0.001 |
| Cardiac glycoside       |                    |                     |
| Digoxin vs placebo      | RR, 0.99; 95% CI, 0.91–1.07; P=0.80 | Hospitalization for worsening HF: RR, 0.72; 95% CI, 0.66–0.79; P<0.001 |
| HCN channel blocker     |                    |                     |
| Ivabradine vs placebo   | HR, 0.90; 95% CI, 0.80–1.02; P=0.092 | CV death or hospitalization for worsening HF: HR, 0.82; 95% CI, 0.75–0.90; P<0.0001 |
| ARNI                   |                    |                     |
| valsartan/valsartan vs enalapril | HR, 0.84; 95% CI, 0.76–0.93; P<0.001 | CV-related mortality or first HFH: HR, 0.80; 95% CI, 0.73–0.87; P<0.001 |
| Sacubitril/valsartan vs enalapril | HR, 0.66; 95% CI, 0.30–1.48; P not reported | Composite of serious clinical events:① HR, 0.54; 95% CI, 0.37–0.79; P not reported |

Notes: ①This endpoint included death, rehospitalization for heart failure, implantation of a left ventricular device, and inclusion on the list of patients eligible for heart transplantation.
Abbreviations: ACEI, angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor blocker; ARNI, angiotensin receptor-neprilysin inhibitor; CI, confidence interval; CR, controlled-release; CV, cardiovascular; HCN, hyperpolarization-activated cyclic nucleotide; HF, heart failure; HFH, heart failure hospitalization; HR, hazard ratio; MRA, mineralocorticoid receptor antagonist; RR, relative risk; XL, extended-release.

patients unable to receive ACEIs, due to poor tolerance or contraindications to therapy.6–8

MRAs
MRAs are recommended in patients with NYHA class II–IV HFrEF to reduce morbidity and mortality unless contraindications are present.6 Mortality benefits were first observed in 1999 with spironolactone, and subsequently, in 2011 with eplerenone.13,33 The overwhelming benefits observed with eplerenone for reducing mortality and hospitalization risks support introducing an MRA earlier in the continuum of HF progression, consistent with guideline recommendations.6,33

Digoxin
Digoxin is recommended to reduce risk of HF hospitalization for patients with HFrEF who continue to experience persistent symptoms during GDEM or for patients with severe symptoms at initial presentation who have not yet responded symptomatically to GDEM.6 In clinical trials,
long-term treatment with digoxin has not been shown to decrease risk of mortality, but modest reductions have been observed for risks of composite endpoints, including hospitalization for HF and CV or HF mortality.34,35

**Ivabradine**

Ivabradine is a specific inhibitor of the If current in the sinusoidal node that is recommended to reduce risk of hospitalization for HF in patients with NYHA class II–III HFrEF who are receiving GDEM that includes a β-blocker at the maximally tolerated dose and who are in sinus rhythm with a resting heart rate ≥70 bpm.7,8 Ivabradine has been shown to reduce the risk of the composite endpoint of CV death or HF hospitalization in patients with HFrEF, but statistical significance was not reached compared with placebo for reduction in all-cause mortality.36

**Sacubitril/valsartan and the PARADIGM-HF trial**

Sacubitril/valsartan: the first ARNI

Even among patients receiving standard-of-care management, HFrEF continues to be associated with high rates of mortality, as the disease typically progresses despite treatment.2,37,38 The number of deaths attributable to HF was approximately as high in 2013 as it was in 1995,2 likely due to the lack of therapeutic advancements. A number of novel treatments have been investigated to improve clinical outcomes for patients with HF. Sacubitril/valsartan is the first-in-class ARNI for the treatment of HFrEF.7,39 The inhibition of neprilysin prevents the breakdown of endogenous compensatory peptides, including the natriuretic peptides (ie, ANP, BNP, CNP) and angiotensin II, allowing them to exert their mitigating effects.40 Neprilysin inhibition from the sacubitril component complements the blockade of angiotensin receptors with valsartan to synergistically improve clinical outcomes in patients with HFrEF compared with standalone RAAS inhibition.40–42

The ARNI sacubitril/valsartan was approved in 2015 by the US Food and Drug Administration (FDA) and the European Medicines Agency for the treatment of chronic HF and HFrEF.43,44 The granted approval was supported by the positive results of the phase 3 Prospective Comparison of ARNI with ACEI to Determine Impact on Global Mortality and Morbidity in Heart Failure (PARADIGM-HF; NCT01035255) trial (Table 1).45

Additional findings from the PARADIGM-HF trial

In an extrapolation of survival data from the PARADIGM-HF trial, it was projected that the long-term treatment effect of sacubitril/valsartan could be a mean increase in survival time of 1.4 years for a 55-year-old patient and 1.3 years for a 65-year-old patient—higher than any increase imparted by enalapril treatment.46 Furthermore, a pharmaeconomic analysis suggested that, compared with the ACEI lisinopril, sacubitril/valsartan treatment provided 0.69 additional life-years and 0.62 additional quality-adjusted life-years.47 This evidence highlights the value of adding ARNIs in place of ACEIs—when not contraindicated—in the treatment of HFrEF. A detailed analysis of the mode of death in the PARADIGM-HF trial showed that, compared with enalapril, sacubitril/valsartan also reduced the risk for sudden cardiac death and death due to worsening HF (Figure 1), whereas the risk for death attributed to stroke or myocardial infarction was not significantly affected.48 The observation of reduced risk of sudden cardiac death with sacubitril/valsartan is noteworthy given the relatively low rate (~15%) of implantable cardioverter-defibrillator (ICD) use in the trial, which was in line with clinical practice.45,49
The mechanism by which sacubitril/valsartan reduces CV mortality is not fully understood. However, it has been hypothesized that neprilysin inhibition may reduce the risk for fatal ventricular arrhythmias (which are a common cause of sudden cardiac death) by reducing myocardial fibrosis and ventricular hypertrophy or attenuating progressive ventricular remodeling. Hyperkalemia is also a known risk factor for CV mortality and sudden cardiac death. Thus, it is plausible that some of the reduced mortality benefits associated with sacubitril/valsartan could be associated with reduced risk of hyperkalemia compared with enalapril (discussed in detail below), possibly as a result of membrane stabilization.

Improvements in safety outcomes were also observed in the PARADIGM-HF trial. Hyperkalemia (serum potassium ≥6.0 mmol/L) was observed less frequently in patients receiving sacubitril/valsartan (4.3%) than in patients taking enalapril (5.6%); however, the incidence of symptomatic hypotension was higher in the sacubitril/valsartan group (14.0%) than in the enalapril group (9.2%). Other adverse events that were observed less frequently with sacubitril/valsartan compared with enalapril included cough (11.3% vs 14.3%) and elevated serum creatinine (≥2.5 mg/dL; 3.3% vs 4.5%). Angioedema was reported in 19 patients receiving sacubitril/valsartan and 10 patients receiving enalapril. Additionally, fewer patients discontinued sacubitril/valsartan than discontinued enalapril because of an adverse event (10.7% vs 12.3%) or renal impairment (0.7% vs 1.4%).

Neprilysin inhibition and RAAS blockade have also been shown to be renal protective in a meta-analysis of 4 studies, including the PARADIGM-HF and Prospective Comparison of ARNI withARB on Management of Heart Failure with Preserved Ejection Fraction (PARAMOUNT; NCT00887588) trials (sacubitril/valsartan in HFpEF and HFrEF, respectively) and the International Multicentre Prevalence Study on Sepsis (IMPRESS) and Omapatrilat Versus Enalapril Randomized Trial of Utility in Reducing Events (OVERTURE) studies (omapatrilat in HFrEF). This analysis showed that the synergistic effect of inhibiting neprilysin and the RAAS provides a 32% reduction in the risk of decline in renal function, defined as: 1) elevated serum creatinine or 2) >50% reduction in glomerular filtration rate, over a mean follow-up time of 51 weeks.

Patients receiving sacubitril/valsartan were also less likely to experience worsening of NYHA functional class within 8 months of randomization versus those receiving enalapril (5.4% vs 7.0%; P=0.004). Likewise, fewer patients receiving treatment with valsartan/valsartan compared with those receiving enalapril experienced symptom worsening, as measured by a ≥5-point reduction in the Kansas City Cardiomyopathy Questionnaire (KCCQ) total symptom score within 4 months (25.1% vs 28.3%; P=0.002), 8 months (28.2% vs 31.8%; P=0.001), and 12 months (29.0% vs 31.5%; P=0.03) of randomization (Figure 2). Among patients who did experience clinical progression, significantly fewer patients receiving sacubitril/valsartan versus enalapril required intensification of therapy for HF (ie, addition of a new drug, intravenous therapy, or increase in diuretic dose; hazard ratio (HR), 0.84; 95% confidence interval (CI), 0.74–0.94; P=0.003).

Treatment with sacubitril/valsartan also resulted in reduced hospitalization rates. In the PARADIGM-HF trial, hospitalization due to worsening HF was 23% less likely for patients treated with sacubitril/valsartan compared with enalapril (HR, 0.77; 95% CI, 0.67–0.89; P<0.001); the difference between treatment groups was evident as early as 30 days after randomization. This finding is particularly important because hospitalization due to acute decompensation is associated with disease progression. Data from studies of patients hospitalized for acute HF suggest a subsequent increased post-discharge mortality risk compared with those who were hospitalized for other reasons or not at all. Of note, the episodes of acute decompensation observed in PARADIGM-HF were less severe for patients treated...
with sacubitril/valsartan compared with enalapril. Among surviving patients from the PARADIGM-HF trial who were hospitalized, those receiving sacubitril/valsartan versus enalapril were less likely to need intensive care (HR, 0.87; 95% CI, 0.78–0.98; \( P=0.019 \)) or intravenous positive inotrope therapy (HR, 0.69; 95% CI, 0.57–0.85; \( P<0.001 \)), and also had a decreased risk for visiting the emergency department due to HF (HR, 0.66; 95% CI, 0.52–0.85; \( P=0.001 \)).

Similarly, repeated hospitalization for HF has been associated with worse post-discharge outcomes. Compared with enalapril, sacubitril/valsartan therapy reduced the rates of 30-day rehospitalization for any cause (21.0% vs 17.8%, respectively; \( P=0.031 \)) and of 30-day rehospitalization for HF (13.4% vs 9.7%; \( P=0.006 \)). Overall, patients treated with sacubitril/valsartan were 29% less likely to be repeatedly hospitalized for HF compared with those treated with enalapril (\( P=0.001 \)).

The early mortality benefit observed with sacubitril/valsartan in the PARADIGM-HF trial
An important finding from the PARADIGM-HF trial was that the mortality benefit with sacubitril/valsartan was conferred early during treatment, as reflected by the early and persistent divergence of sacubitril/valsartan from enalapril on Kaplan–Meier curves for both the combined primary outcome (ie, CV death or hospitalization for HF) and for CV death alone (Figure 3).

This early mortality benefit was sustained throughout treatment, as the difference in favor of sacubitril/valsartan for the primary outcome was observed at each of the interim analysis time points. The trial was stopped early after a median follow-up of only 27 months for reaching the statistical threshold of an overwhelming benefit prior to the anticipated end of the trial.

The precise mechanisms by which sacubitril/valsartan prevents or delays CV death are uncertain; however, emerging evidence suggests that the beneficial effects of this therapy are related to specific amelioration of HF disease pathways. In a pharmacodynamics study of patients with HFrEF who received sacubitril/valsartan, plasma levels of aldosterone, endothelin-1, and N-terminal-proBNP (NT-proBNP) were significantly decreased after 21 days of treatment (all \( P<0.05 \)), suggesting an early hemodynamic benefit from ARNI therapy.

The 2016 and 2017 American College of Cardiology (ACC)/American Heart Association (AHA)/Heart Failure Society of America (HFSA) guideline updates
The early clinical benefits associated with sacubitril/valsartan underscore the importance of its initiation early in the HF treatment paradigm. Accordingly, the 2016 and 2017 ACC/AHA/HFSA guideline updates added ARNi to the list of recommendations to reduce HF morbidity and mortality risks for patients with chronic HFrEF, irrespective of prior therapy, and did not specify different recommendations for those who are naïve to therapy with ACEI or ARBs. Furthermore, the guidelines recommend the replacement of ACEI or ARB therapy that is tolerated by patients with chronic symptomatic NYHA class II or III HFrEF with ARNI therapy to further reduce morbidity and mortality. The recommendation to switch from the standard of care to this novel pharmacologic class in patients with NYHA class II or III HFrEF, in particular, is in recognition of the value of the early use of ARNI therapy. These guidelines also offer important safety considerations for the use of ARNI therapy; specifically, that it should not be used concomitantly (or within 36 hrs) of a dose of an ACEI and that it should not be prescribed to patients with a history of angioedema.

Sacubitril/valsartan post-approval experience and implementation in clinical practice
Target patient profile
To translate the benefits associated with sacubitril/valsartan therapy seen in the PARADIGM-HF trial into clinical practice, the target patient populations for this treatment should be identified. This is accomplished by: 1) comparing the patient’s clinical profile with the indications, contraindications, and safety warnings stated in the approved prescribing information for the prescriber’s jurisdiction; 2) by considering the expert guidance of the American and/or European cardiology guidelines on all appropriate treatment options for this patient; and 3) by evaluating the patient’s characteristics in comparison to those of patients enrolled in the PARADIGM-HF trial (Table 2).

Baseline and demographic characteristics of patients from the PARADIGM-HF trial are summarized in Table 2. Despite use of GDEM recommended at the time, patients exhibited persistent signs and symptoms of HF. As discussed previously, approximately 15% (n=1243) of enrolled
patients had an ICD. Although this number is relatively low, it is comparable to the percentage of patients with an ICD enrolled in previous HF trials. Moreover, only approximately 50% of the eligible patients receive ICD therapy in clinical practice.

Across the various analyses of the PARADIGM-HF trial that explored the potential effects of different patient characteristics on treatment benefits, very little heterogeneity in the sacubitril/valsartan treatment outcomes was observed. An examination of the PARADIGM-HF trial results by age showed similar benefits with sacubitril/valsartan over enalapril therapy across the broad range of ages (18–96 years) entered in the trial, even after adjustment for baseline differences. Based on the post hoc analysis of PARADIGM-HF trial results by baseline EF, which ranged from 5% to 42%, the degree of LV dysfunction also did not appear to impact outcomes. Therefore, patient age and degree of LV dysfunction should not be a priori factors considered for the selection of patients who may benefit from ARNI therapy.

Although most patients in the PARADIGM-HF trial had mild HF symptoms according to NYHA classification, numerous patients in the trial were at high risk for adverse outcomes when evaluated according to HF risk scores, including the Meta-Analysis Global Group in Chronic Heart Failure (MAGGIC) and the Eplerenone in Mild Patients Hospitalization and Survival Study in Heart Failure (EMPHASIS-HF) risk scores. The benefits of sacubitril/valsartan in the PARADIGM-HF trial were observed across all risk categories, with the greatest absolute benefit reported in patients deemed to be at highest risk.

Although this discourse has advocated the selection of patients having characteristics resembling the PARADIGM-HF trial population, it is important to note that the approved prescribing information may include patients who would have been excluded from the PARADIGM-HF trial. For example, the FDA approved sacubitril/valsartan for the treatment of patients naïve to ACEIs and ARBs, although such patients were excluded from the PARADIGM-HF trial. Similarly, the benefits of sacubitril/valsartan therapy observed in populations composed mostly of patients with NYHA class II HFrEF could be extrapolated to suggest that its use early in the disease process is likely to have a positive impact on clinical outcomes. Ideally, cardiologists would treat HFrEF immediately following the onset of symptoms; however, the optimal timing for the initiation of sacubitril/valsartan has yet to be determined.

Hospitalization presents an opportunity for the initiation of sacubitril/valsartan therapy once patients are stabilized. The safety and efficacy of this approach cannot be determined from the results of the PARADIGM-HF trial, as this trial did not include patients recently hospitalized for HF (due to the requirement for patients to be taking stable doses of a β-blocker and an ACEI or ARB for at least 4 weeks prior to randomization). To support recommendations regarding the use of sacubitril/valsartan therapy following hospitalization for HF, the Comparison of Sacubitril/valsartan Versus Enalapril on Effect

| No. at Risk | Days Since Randomization | Cumulative Probability |
|-------------|--------------------------|------------------------|
| A            | Primary Endpoint         |                        |
| LCZ696      | 4187 3922 3663 3018 2257 1544 896 249 | 0.01 0.30 0.55 0.70 0.80 0.87 0.93 0.97 |
| Enalapril   | 4212 3883 3579 2922 2123 1488 853 236 | 0.01 0.30 0.55 0.70 0.80 0.87 0.93 0.97 |
| B            | Death From Cardiovascular Causes |                    |
| LCZ696      | 4187 4056 3891 3282 2478 1716 1005 280 | 0.01 0.30 0.55 0.70 0.80 0.87 0.93 0.97 |
| Enalapril   | 4212 4051 3860 3231 2410 1726 994 279 | 0.01 0.30 0.55 0.70 0.80 0.87 0.93 0.97 |

Figure 3 PARADIGM-HF Kaplan-Meier curves for the primary endpoint (A) and death from cardiovascular causes (B), according to study group.

Notes: From *N Engl J Med*, McMurray JJ, Packer M, Desai AS, et al, Angiotensin-neprilysin inhibition versus enalapril in heart failure, 371(11):993–1004. Copyright © (2014) Massachusetts Medical Society. Adapted with permission from Massachusetts Medical Society.
Table 2 Identification of candidates for sacubitril/valsartan treatment

| PARADIGM-HF patient characteristics, N=8442 |
|---------------------------------------------|
| • Mean age: 64 years                        |
| • Predominantly male (78%)                  |
| • Less severe disease (70% NYHA class II)   |
| • Majority with ischemic cardiomyopathy (60%)
| • Hospitalized for HF (63%)                 |
| • Comorbid hypertension (71%) and diabetes mellitus (35%) |
| • On GDEM (recommended at the time) prior to enrollment, treated with an ACEI (77%), ARB (22%), or both |
| • Background therapy included β-blockers (93%), diuretics (80%), and MRAs (60%) |
| • Mean LVEF <30%                            |
| • High NT-proBNP levels (mean concentration=1608 pg/mL) |
| • Peripheral edema (21%)                    |
| • Poor self-reported quality of life, KCCQ OSS=73 |

Evidence-based support for treatment

- Patients with HFrEF
- Patients with mild-to-moderate chronic HF, eg, NYHA class II-III
- Patients with stable HF already receiving guideline-directed therapy in other pharmacologic classes, as appropriate (β-blocker and MRA)
- Systolic blood pressure ≥100 mmHg
- eGFR ≥30 mL/min/1.73 m²
- Potassium ≤5.2 mmol/L

Evidence insufficient or not available to currently recommend treatment

- Patients with HFrEF
- Patients with severe HF, eg, NYHA class IV
- Hospitalized patients with acute decompensated HF
- Patients with specific disease etiology or comorbidity

Contraindicated for treatment

- History of angioedema related to previous ACEI or ARB therapy
- Concomitant use of ACEI or other ARB
- Concomitant use of aliskiren in patients with diabetes mellitus
- Severe hepatic impairment

Notes: "Data from these studies. 7-8,39-45,60

Abbreviations: ACEI, angiotensin-converting enzyme inhibitor; ARB, angiotensin II receptor blocker; ARNI, angiotensin receptor-neprilysin inhibitor; eGFR, estimated glomerular filtration rate; GDEM, guideline-directed evaluation and management; HF, heart failure; HFpEF, HF with preserved ejection fraction; HFrEF, HF with reduced ejection fraction; KCCQ OSS, Kansas City Cardiomyopathy Questionnaire overall summary score; LVEF, left ventricular ejection fraction; MRA, mineralocorticoid receptor antagonist; NT-proBNP, N-terminal pro-B-type natriuretic peptide; NYHA, New York Heart Association; PARADIGM-HF, Prospective Comparison of ARNI with ACEI to Determine Impact on Global Mortality and Morbidity in Heart Failure.

on NT-proBNP in Patients Stabilized From an Acute Heart Failure Episode (PIONEER-HF; NCT02554890) trial compared the efficacy and safety of sacubitril/valsartan and enalapril in patients who were hospitalized for acute decompensated HF and were hemodynamically stable prior to initiating treatment. 64 Sacubitril/valsartan therapy provided a significantly greater reduction in levels of NT-proBNP than enalapril, with a difference between treatment groups evident as early as week 1. In addition, the risk of the composite endpoint of serious clinical events (death, rehospitalization for HF, planned implantation of a left ventricular device, and listing for heart transplant) was lower in patients receiving sacubitril/valsartan compared with enalapril. The effect of inpatient sacubitril/valsartan therapy following an ischemic insult and revascularization (peri-myocardial infarction) or post-myocardial infarction evaluated by the Prospective ARNI vs ACE Inhibitor Trial to Determine Superiority in Reducing Heart Failure Events After MI (PARADISE-MI; NCT02924727) 65 trial.

Dosing considerations

In the PARADIGM-HF trial, achievement and maintenance of the target doses of sacubitril/valsartan (97/103 mg twice daily) and enalapril (10 mg twice daily) were attempted, with mean total daily doses of sacubitril/valsartan (375 mg) and enalapril (18.9 mg) close to target at the last study assessment. 45 However, achieving and maintaining the target dose of an ACEI or sacubitril/valsartan can be challenging in some patient populations. 6 Among hospitalized patients with acute decompensated HF enrolled in the recent PIONEER-HF trial, 55.2% in the sacubitril/valsartan group and 60.8% in the enalapril group were receiving the target dose of the assigned trial drug by week 8. 66 Although it may take longer than 2 weeks for some patients to tolerate titration up to the target dose, ultimately, perseverance to achieve that target is important to obtain maximal therapeutic benefit. 6 In a post hoc analysis of the PARADIGM-HF trial, dose reductions by any amount in either treatment group were associated with a higher risk for the primary outcome of CV death or HF-related hospitalization (HR 2.5; 95% CI, 2.2–2.7). 66 Nonetheless, patients who received treatment with lower-than-target doses of sacubitril/valsartan still experienced a significant benefit compared with patients who received treatment with lower-than-target doses of enalapril (HR 0.80; 95% CI, 0.70–0.93). 66

The proportion of patients who required a dose reduction in the PARADIGM-HF trial was similar between the enalapril (43%) and sacubitril/valsartan (42%) groups, and the reasons for dose reduction were generally similar; however, significantly more patients taking enalapril required dose reduction due to cough (4.3% vs 1.8%; P<0.001) or hypotension (21.7% vs 16.3%; P<0.001). 66 Serum creatinine concentration was the best predictor of trial medication dose reduction (odds ratio 2.38; 95% CI, 2.01–2.82), 66 as explained by the relation of
renal function to common adverse events, including hyperkalemia, hypotension, and angioedema. Permanent discontinuation of treatment was rare, irrespective of therapy.\textsuperscript{45} In many cases, patients on either treatment were successfully re-uptitrated to a higher dose; however, more patients on sacubitril/valsartan were successfully re-uptitrated versus those on enalapril following hypotension (36% vs 27%; \textit{P}=0.026) or renal dysfunction (41% vs 28%; \textit{P}=0.018).\textsuperscript{66} The recent Multicenter, Randomized, Double-blind, Parallel Group Study to Assess the Safety and Tolerability of Initiating LCZ696 in Heart Failure Patients Comparing Two Titration Regimens (TITRATION; NCT01922089) compared the tolerability of initiating and titrating sacubitril/valsartan with a condensed (49/51 mg twice daily for 2 weeks, followed by 97/103 mg twice daily) versus conservative schedule (24/26 mg twice daily for 2 weeks, followed by 49/51 mg twice daily for 3 weeks, then 97/103 mg twice daily) in patients with varying levels of prior ACEI or ARB exposure.\textsuperscript{67} Notably, this trial differed from the PARADIGM-HF trial because it included ACEI-naïve, ARB-naïve patients.\textsuperscript{57} In the TITRATION trial, 78% and 84% of the patients on condensed versus conservative initiation schedules, respectively, successfully achieved and maintained the target dose of 97/103 mg sacubitril/valsartan twice daily for 12 weeks.\textsuperscript{67} Furthermore, 83% and 87% of the patients on condensed and conservative schedules, respectively, tolerated a dose of 97/103 mg twice daily for the last 2 weeks of the 12-week period, irrespective of previous dose reductions or interruptions.\textsuperscript{67} Patients initially on low-dose ACEI or ARB therapy had significantly higher success rates with the conservative titration approach compared with the condensed titration approach (85% vs 74%; \textit{P}=0.030).\textsuperscript{67} Although the titration period is often longer in clinical practice, this trial has demonstrated that a shorter titration period is generally well tolerated and can successfully be used to achieve target doses in some patients. Similarly, short titration periods (≤1 week) have been successful in previous ACEI and ARB clinical trials.\textsuperscript{14,26,68}

**Conclusion**

Sacubitril/valsartan represents the first of a breakthrough class of drugs for the treatment of HF. The early mortality benefit observed with sacubitril/valsartan compared with enalapril in the PARADIGM-HF trial and the recent guideline recommendations regarding the use of an ARNI in HFrEF management make a compelling case for the early use of sacubitril/valsartan in the HF treatment paradigm.

**Human and animal rights and informed consent**

This article does not contain any studies with human participants or animals performed by any of the authors.

**Data availability**

The data used to compile this review are available in the relevant source publications.

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**Disclosure**

Dr Sokos has served on the speakers’ bureaus of Bayer Pharmaceuticals, Actelion Pharmaceuticals, and Novartis Pharmaceuticals. He has served on the advisory boards of Gilead Sciences, Bayer Pharmaceuticals, and Novartis Pharmaceuticals. Dr Raina has received fees from Actelion, St. Jude, Bayer, and United Therapeutics, and has received nonfinancial support from Bellerophon and Oxford Pharmagenesis. The authors report no other conflicts of interest in this work.

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