2017 ACC/AHA/HFSA Focused Update of the 2013 ACCF/AHA **Guideline for the Management of Heart Failure**

A Report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Failure Society of America

Developed in Collaboration With the American Academy of Family Physicians, American College of Chest Physicians, and International Society for Heart and Lung Transplantation

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Preamble

Since 1980, the American College of Cardiology (ACC) and American Heart Association (AHA) have translated scientific evidence into clinical practice guidelines (guidelines) with recommendations to improve cardiovascular health. These guidelines, which are based on systematic methods to evaluate and classify evidence, provide a cornerstone for quality cardiovascular care. The ACC and AHA sponsor the development and publication of guidelines without commercial support, and members of each organization volunteer their time to the writing and review efforts. Guidelines are official policy of the ACC and AHA.

Intended Use

Practice guidelines provide recommendations applicable to patients with or at risk of developing cardiovascular disease. The focus is on medical practice in the United States, but guidelines developed in collaboration with other organizations may have a global impact. Although guidelines may be used to inform regulatory or payer decisions, their intent is to improve patients' quality of care and align with patients' interests. Guidelines are intended to define practices meeting the needs of patients in most, but not all, circumstances and should not replace clinical judgment.

Clinical Implementation

Guideline recommended management is effective only when followed by healthcare providers and patients. Adherence to recommendations can be enhanced by shared decision making between healthcare providers and patients, with patient engagement in selecting interventions based on individual values, preferences, and associated conditions and comorbidities.

Methodology and Modernization

The ACC/AHA Task Force on Clinical Practice Guidelines (Task Force) continuously reviews, updates, and modifies guideline methodology on the basis of published standards from organizations including the Institute of Medicine (1, 2) and on the basis of internal reevaluation. Similarly, the presentation and delivery of guidelines are reevaluated and modified on the basis of evolving technologies and other factors to facilitate optimal dissemination of information at the point of care to healthcare professionals. Given time constraints of busy healthcare providers and the need to limit text, the current guideline format delineates that each recommendation be supported by limited text (ideally, <250 words) and hyperlinks to supportive evidence summary tables. Ongoing efforts to further limit text are underway. Recognizing the importance of cost-value considerations in certain guidelines, when appropriate and feasible, an analysis of the value of a drug, device, or intervention may be performed in accordance with the ACC/AHA methodology (3).

To ensure that guideline recommendations remain current, new data are reviewed on an ongoing basis, with full guideline revisions commissioned in approximately 6-year cycles. Publication of new, potentially practice-changing study results that are relevant to an existing or new drug, device, or management strategy will prompt evaluation by the Task Force, in consultation with the relevant guideline writing committee, to determine whether a focused update should be commissioned. For additional information and policies regarding guideline development, we encourage readers to consult the ACC/AHA guideline methodology manual (4) and other methodology articles (5-8).

Selection of Writing Committee Members

The Task Force strives to avoid bias by selecting experts from a broad array of backgrounds. Writing committee members represent different geographic regions, sexes, ethnicities, races, intellectual perspectives/biases, and scopes of clinical practice. The Task Force may also invite organizations and professional societies with related interests and expertise to participate as partners, collaborators, or endorsers.

Relationships With Industry and Other Entities

http://www.acc.org/guidelines/about-guidelines-and-clinical-documents/guidelines-and-documents-task-forces.



Evidence Review and Evidence Review Committees

When developing recommendations, the writing committee uses evidence-based methodologies that are based on all available data (4-7). Literature searches focus on randomized controlled trials (RCTs) but also include registries, nonrandomized comparative and descriptive studies, case series, cohort studies, systematic reviews, and expert opinion. Only key references are cited.

An independent evidence review committee (ERC) is commissioned when there are 1 or more questions deemed of utmost clinical importance that merit formal systematic review. This systematic review will strive to determine which patients are most likely to benefit from a drug, device, or treatment strategy and to what degree. Criteria for commissioning an ERC and formal systematic review include: a) the absence of a current authoritative systematic review, b) the feasibility of defining the benefit and risk in a time frame consistent with the writing of a guideline, c) the relevance to a substantial number of patients, and d) the likelihood that the findings can be translated into actionable recommendations. ERC members may include methodologists, epidemiologists, healthcare providers, and biostatisticians. When a formal systematic review has been commissioned, the recommendations developed by the writing committee on the basis of the systematic review are marked with "^{SR}".

Guideline-Directed Management and Therapy

The term *guideline-directed management and therapy* (GDMT) encompasses clinical evaluation, diagnostic testing, and pharmacological and procedural treatments. For these and all recommended drug treatment regimens, the reader should confirm the dosage by reviewing product insert material and evaluate the treatment regimen for contraindications and interactions. The recommendations are limited to drugs, devices, and treatments approved for clinical use in the United States.

Class of Recommendation and Level of Evidence

The Class of Recommendation (COR) indicates the strength of the recommendation, encompassing the estimated magnitude and certainty of benefit in proportion to risk. The Level of Evidence (LOE) rates the

quality of scientific evidence that supports the intervention on the basis of the type, quantity, and consistency of data from clinical trials and other sources (Table 1) (4-6).

Glenn N. Levine, MD, FACC, FAHA Chair, ACC/AHA Task Force on Clinical Practice Guidelines

> American Heart Association

Circulation

1. Introduction

The purpose of this focused update is to update the "2013 ACCF/AHA Guideline for the Management of Heart Failure" (9) (2013 HF guideline) in areas in which new evidence has emerged since its publication. For this update and future heart failure (HF) guidelines, the Heart Failure Society of America (HFSA) has partnered with the ACC and AHA to provide coordinated guidance on the management of HF.

The scope of the focused update includes revision to the sections on biomarkers; new therapies indicated for stage C HF with reduced ejection fraction (HFrEF); updates on HF with preserved ejection fraction (HFpEF); new data on important comorbidities, including sleep apnea, anemia, and hypertension; and new insights into the prevention of HF.

This focused update represents the second part of a 2-stage publication; with the first part having been published as the "2016 ACC/AHA/HFSA Focused Update on New Pharmacological Therapy for Heart Failure" (10), which introduced guidance on new therapies, specifically for the use of an angiotensin receptor–neprilysin inhibitor (ARNI) (valsartan/sacubitril) and a sinoatrial node modulator (ivabradine). That focused update was published concurrently with the European Society of Cardiology's complete guideline, "2016 ESC Guidelines for the Diagnosis and Treatment of Acute and Chronic Heart Failure" (11).

1.1. Methodology and Evidence Review

To identify key data that influence guideline recommendations, the Task Force and members of the 2013 HF guideline writing committee reviewed clinical trials that were presented at the annual scientific meetings of the ACC, AHA, and European Society of Cardiology and other scientific meetings and that were published in peer-reviewed format from April 2013 through November 2016. The evidence is summarized in tables in the Online Data Supplement

(http://circ.ahajournals.org/lookup/suppl/doi:10.1161/CIR.0000000000000000/-/DC2). All

recommendations (new, modified, and unchanged) for each clinical section are included to provide a comprehensive assessment. The text explains new and modified recommendations, whereas recommendations from the previous guideline that have been deleted or superseded no longer appear. Please consult the full-text version of the 2013 HF guideline (9) for text and evidence tables supporting the unchanged recommendations and for clinical areas not addressed in this focused update. Individual recommendations in this focused update will be incorporated into the full-text guideline in the future. Recommendations from the prior guideline that remain current have been included for completeness, but the LOE reflects the COR/LOE system used when the recommendations were initially developed. New and modified recommendations in this focused update reflect the latest COR/LOE system, in which LOE

B and C are subcategorized for greater specificity (4-6). The section numbers correspond to the full-text guideline sections.

1.2. Organization of the Writing Group

For this focused update, representative members of the 2013 HF guideline writing committee were invited to participate. They were joined by additional invited members to form a new writing group, which is referred to as the 2017 HF focused update writing group. Members were required to disclose all RWI relevant to the data under consideration. The group was composed of experts representing general cardiologists, HF and transplantation specialists, electrophysiologists, pharmacists, and general internists. The 2017 HF focused update writing group included representatives from the ACC, AHA, and HFSA, as well as the American Academy of Family Physicians, American College of Chest Physicians, American College of Physicians, and International Society for Heart and Lung Transplantation.

1.3. Document Review and Approval



The focused update was reviewed by 2 official reviewers each nominated by the ACC, AHA, and HFSA; 1 reviewer each from the American Academy of Family Physicians, American College of Chest Physicians, and International Society for Heart and Lung Transplantation; and 19 individual content reviewers. Reviewers' RWI information is published in this document (Appendix 2).

This document was approved for publication by the governing bodies of the ACC, AHA, and

HFSA.

Table 1. Applying Class of Recommendation and Level of Evidence to Clinical Strategies, Interventions, Treatments, or Diagnostic Testing in Patient Care* (Updated August 2015)

CLASS (STRENGTH) OF RECOMMENDATION			
CLASS I (STRONG)	Benefit >>> Risk		
 Suggested phrases for writing recommendations Is recommended Is indicated/useful/effective/beneficial Should be performed/administered/other Comparative-Effectiveness Phrases†: Treatment/strategy A is recommended/incorpreference to treatment B Treatment A should be chosen over treatment 	s: dicated in lient B		
CLASS IIa (MODERATE)	Benefit >> Risk		
Suggested phrases for writing recommendations Is reasonable Can be useful/effective/beneficial Comparative-Effectiveness Phrases†: Treatment/strategy A is probably recommend preference to treatment B It is reasonable to choose treatment A over treatment B	:: nded/indicated in		
CLASS IIb (WEAK)	$\textbf{Benefit} \geq \textbf{Risk}$		
Suggested phrases for writing recommendations May/might be reasonable May/might be considered Usefulness/effectiveness is unknown/unclear or not well established	:: /uncertain		
CLASS III: No Benefit (MODERATE) (Generally, LOE A or B use only)	Benefit = Risk		
Suggested phrases for writing recommendations Is not recommended Is not indicated/useful/effective/beneficial Should not be performed/administered/other 	r		
CLASS III: Harm (STRONG)	Risk > Benefit		
Suggested phrases for writing recommendations Potentially harmful Causes harm 			

- Associated with excess morbidity/mortality
- Should not be performed/administered/other

LEVEL (QUALITY) OF EVIDENCE‡

LEVEL A

- High-quality evidence‡ from more than 1 RCT
- Meta-analyses of high-quality RCTs
- One or more RCTs corroborated by high-quality registry studies

LEVEL B-R

- Moderate-quality evidence‡ from 1 or more RCTs
- Meta-analyses of moderate-quality RCTs

LEVEL B-NR

(Nonrandomized)

(Randomized)

- Moderate-quality evidence‡ from 1 or more well-designed, well-executed nonrandomized studies, observational studies, or registry studies
- Meta-analyses of such studies

LEVEL C-LD

- Randomized or nonrandomized observational or registry studies with limitations of design or execution
- Meta-analyses of such studies
- Physiological or mechanistic studies in human subjects

LEVEL C-EO

Consensus of expert opinion based on clinical experience

COR and LOE are determined independently (any COR may be paired with any LOE).

A recommendation with LOE C does not imply that the recommendation is weak. Many important clinical questions addressed in guidelines do not lend themselves to clinical trials. Although RCTs are unavailable, there may be a very clear clinical consensus that a particular test or therapy is useful or effective.

- * The outcome or result of the intervention should be specified (an improved clinical outcome or increased diagnostic accuracy or incremental prognostic information).
- † For comparative-effectiveness recommendations (COR I and IIa; LOE A and B only), studies that support the use of comparator verbs should involve direct comparisons of the treatments or strategies being evaluated.
- ‡ The method of assessing quality is evolving, including the application of standardized, widely used, and preferably validated evidence grading tools; and for systematic reviews, the incorporation of an Evidence Review Committee.

COR indicates Class of Recommendation; EO, expert opinion; LD, limited data; LOE, Level of Evidence; NR, nonrandomized; R, randomized; and RCT, randomized controlled trial.

6. Initial and Serial Evaluation of the HF Patient

6.3. Biomarkers

Assays for BNP (B-type natriuretic peptide) and NT-proBNP (N-terminal pro-B-type natriuretic peptide), which are both natriuretic peptide biomarkers, have been used increasingly to establish the presence and severity of HF. In general, both natriuretic peptide biomarker values track similarly, and either can be used in patient care settings as long as their respective absolute values and cutpoints are not used interchangeably. Notably, BNP, but not NT-proBNP, is a substrate for neprilysin. Therefore, ARNI increases BNP levels (12) but not NT-proBNP levels (13). Note that the type of natriuretic peptide assay that has been performed must be considered during interpretation of natriuretic peptide biomarker levels in patients on ARNI. In 2 studies with ARNI, NT-proBNP levels were reduced (12, 14), with the reduction in 1 study being associated with improved clinical outcomes (12).

A substantial evidence base exists that supports the use of natriuretic peptide biomarkers to assist in the diagnosis or exclusion of HF as a cause of symptoms (e.g., dyspnea, weight gain) in the setting of chronic ambulatory HF (15-21) or in the setting of acute care with decompensated HF (22-30), especially when the cause of dyspnea is unclear. The role of natriuretic peptide biomarkers in population screening to detect incident HF is emerging (31-37). Elevated plasma levels of natriuretic peptide biomarkers are associated with a wide variety of cardiac and noncardiac causes (Table 2) (38-42). Obesity may be associated with lower natriuretic peptide concentrations, and this may modestly reduce diagnostic sensitivity in morbidly obese patients (42).

Because of the absence of clear and consistent evidence for improvement in mortality and cardiovascular outcomes (43-62), there are insufficient data to inform specific guideline recommendations related to natriuretic peptide–guided therapy or serial measurements of BNP or NT-proBNP levels for the purpose of reducing hospitalization or deaths in the present document.

Like natriuretic peptides, cardiac troponin levels may be elevated in the setting of chronic or acute decompensated HF, suggesting myocyte injury or necrosis (63). Troponins I and T respond similarly for acute coronary syndromes and acute decompensated HF. Elevations in either troponin I or T levels in the setting of acute HF are of prognostic significance and must be interpreted in the clinical context (64).

In addition to natriuretic peptides and troponins (65-67), multiple other biomarkers, including those of inflammation, oxidative stress, vascular dysfunction, and myocardial and matrix remodeling, have been implicated in HF (68-71). Biomarkers of myocardial fibrosis, soluble ST2 receptor, and galectin-3 are predictive of hospitalization and death and may provide incremental prognostic value over natriuretic peptide levels in patients with HF (72-74). Strategies that combine multiple biomarkers may

ultimately prove beneficial in guiding HF therapy in the future, but multicenter studies with larger derivation and validation cohorts are needed (75, 76). Several emerging biomarkers await validation with well-defined outcome measures and prognostic accuracy before they can reach the clinical arena (77-84).

This section categorizes the role of biomarkers into prevention, diagnosis, prognosis, and added risk stratification to clarify evidence-based objectives of their use in clinical practice.

Table 2. Selected Potential	Causes of Elevated Natriuretic Pe	ptide Levels (38-4	11)

Cardiac	
HF, including RV syndromes	
Acute coronary syndromes	
Heart muscle disease, including LVH	
Valvular heart disease	
Pericardial disease	
Atrial fibrillation	
Myocarditis	
Cardiac surgery	
Cardioversion	
Toxic-metabolic myocardial insults, including cancer chemotherapy	
Noncardiac	American Hoart
Advancing age	Association
Anemia	Ť
Renal failure	
Pulmonary: obstructive sleep apnea, severe pneumonia	
Pulmonary hypertension	
Critical illness	
Bacterial sepsis	
Severe burns	
HF indicates heart failure; LVH, left ventricular hypertrophy; and RV, right ventricular.	
Modified from Table 8 of the 2013 HF guideline (9).	

6.3.1. Biomarkers for Prevention: Recommendation

Biomarkers: Recommendation for Prevention of HF				
COR	LOE	Recommendation	Comment/Rationale	
IIa	B-R	For patients at risk of developing HF, natriuretic	NEW : New data suggest	
		peptide biomarker-based screening followed by	that natriuretic peptide	
		team-based care, including a cardiovascular	biomarker screening and	
See Onli	ne Data	specialist optimizing GDMT, can be useful to	early intervention may	
Supplement	and B.	prevent the development of left ventricular	prevent HF.	
		dysfunction (systolic or diastolic) or new-onset HF		
		(85, 86).		
In a large-scale unblinded single-center study (STOP-HF [The St Vincent's Screening to Prevent Heart Failure])				
(85), patients at risk of HF (identified by the presence of hypertension, diabetes mellitus, or known vascular				
disease [e.g., stage A HF]), but without established left ventricular systolic dysfunction or symptomatic HF at				
baseline, were randomly assigned to receive screening with BNP testing or usual primary care. Intervention-				
group participants with BNP levels of ≥50 pg/mL underwent echocardiography and were referred to a				
cardiovascular specialist who decided on further investigation and management. All patients received further				
coaching by a specialist nurse who emphasized individual risk and the importance of adherence to medication				

and healthy lifestyle behaviors. BNP-based screening reduced the composite endpoint of asymptomatic left ventricular dysfunction (systolic or diastolic) with or without newly diagnosed HF (85). Similarly, in another small, single-center RCT, accelerated up-titration of renin-angiotensin-aldosterone system antagonists and beta blockers reduced cardiac events in patients with diabetes mellitus and elevated NT-proBNP levels but without cardiac disease at baseline (86). Developing a standardized strategy to screen and intervene in patients at risk of HF can be difficult because of different definitions of HF risk, heterogeneity of prevalence in different populations, variable duration until clinical HF or left ventricular dysfunction develops, and variable interventions for risk factor modification or treatment. Further studies are needed to determine cost-effectiveness and risk of such screening, as well as its impact on quality of life (QoL) and mortality rate.

6.3.2. Biomarkers for Diagnosis: Recommendation

Biomarkers: Recommendation for Diagnosis				
COR	LOE	Recommendation	Comment/Rationale	
_		In patients presenting with dyspnea, measurement	MODIFIED: 2013 acute	
1	Α	of natriuretic peptide biomarkers is useful to	and chronic	
	D	support a diagnosis or exclusion of HF (15-24, 28-	recommendations have	
See Onli	ne Data	30).	been combined into a	
Supplement	ts A and B.		diagnosis section.	
Natriuretic peptide biomarker testing in the setting of chronic ambulatory HF provides incremental diagnostic				
value to clinical judgment, especially when the etiology of dyspnea is unclear (15-21). In emergency settings,				
natriuretic peptide biomarker levels usually have higher sensitivity than specificity and may be more useful for				
ruling out than ruling in HF (20). Although lower values of natriuretic peptide biomarkers exclude the presence				
of HF, and higher values have reasonably high positive predictive value to diagnose HF, clinicians should be				
aware that elevated plasma levels for both natriuretic peptides have been associated with a wide variety of				
cardiac and noncardiac causes (Table 2) (38-41).				

6.3.3. Biomarkers for Prognosis or Added Risk Stratification: Recommendations

Biomarkers: Recommendations for Prognosis			
COR	LOE	Recommendations	Comment/Rationale
I	Α	Measurement of BNP or NT-proBNP is useful for establishing prognosis or disease severity in chronic HF (16, 87-92).	2013 recommendation remains current.
Ι	A	Measurement of baseline levels of natriuretic peptide biomarkers and/or cardiac troponin on	MODIFIED: Current recommendation
See Online Data Supplements A and B.		admission to the hospital is useful to establish a prognosis in acutely decompensated HF (27, 93-100).	admission levels of natriuretic peptide biomarkers that are useful.
Higher levels of natriuretic peptide biomarkers on admission are usually associated with greater risk for clinical			

Higher levels of natriuretic peptide biomarkers on admission are usually associated with greater risk for clinical outcomes, including all-cause and cardiovascular mortality, morbidity, and composite outcomes, across different time intervals in patients with decompensated HF (20, 27, 29, 93-101). Similarly, abnormal levels of circulating cardiac troponin are commonly found in patients with acute decompensated HF, often without obvious

myocardial ischemia or underlying coronary artery disease (CAD), and this is associated with worse clinical outcomes and higher risk of death (95, 99, 102, 103).

Studies have demonstrated incremental prognostic value of these biomarkers to standard approaches of cardiovascular disease risk assessment (29, 95). However, there were differences in the risk prediction models, assay cutpoints, and lengths of follow-up (29). Furthermore, not all patients may need biomarker measurement for prognostication, especially if they already have advanced HF with established poor prognosis or persistently elevated levels of biomarkers in former settings. Therefore, assays of natriuretic peptide biomarkers for incremental prognostication should not preclude good clinical judgment; an individualized approach to each patient is paramount.

IIa	B-NR	During a HF hospitalization, a predischarge	NEW : Current
		natriuretic peptide level can be useful to establish a	recommendation reflects
See Onli	ne Data	postdischarge prognosis (93, 96, 104-113).	new observational studies.

Supplements A and B.

Predischarge natriuretic peptide biomarker levels and the relative change in levels during hospital treatment are strong predictors of the risk of death or hospital readmission for HF (93, 96, 104-113). Several studies have suggested that predischarge natriuretic peptide biomarker levels had higher reclassification and discrimination value than clinical variables in predicting outcomes (96, 106, 108-111). Patients with higher predischarge levels and patients who do not have a decrease in natriuretic peptide biomarker levels during hospitalization have worse outcomes (96, 106, 108-111). Although observational or retrospective studies have suggested that patients with natriuretic peptide biomarker reduction had better outcomes than those without any changes or with a biomarker rise (93, 107, 112, 113), targeting a certain threshold, value, or relative change in these biomarker levels during hospitalization may not be practical or safe for every patient and has not been tested in a prospective large-scale trial. Clinical assessment and adherence to GDMT should be the emphasis, and the prognostic value of a predischarge value or relative changes does not imply the necessity for serial and repeated biomarker measurements during hospitalization.

	÷ .		
		In patients with chronic HF, measurement of other	MODIFIED: 2013
llb	B-NR	clinically available tests, such as biomarkers of	recommendations have
		myocardial injury or fibrosis, may be considered for	been combined into
See Online Data		additive risk stratification (27, 95, 98, 99, 103, 114-	prognosis section,
Supplements A and B.		119).	resulting in LOE change
			from A to B-NR.

Biomarkers of myocardial fibrosis (e.g., soluble ST2 receptor, galectin-3, high-sensitivity cardiac troponin, and others) are predictive of hospitalization and death in patients with HF and also are additive to natriuretic peptide biomarker levels in their prognostic value (117, 119-126). A combination of biomarkers may ultimately prove to be more informative than single biomarkers (127).





Colors correspond to COR in Table 1.

*Other biomarkers of injury or fibrosis include soluble ST2 receptor, galectin-3, and high-sensitivity troponin. ACC indicates American College of Cardiology; AHA, American Heart Association; ADHF, acute decompensated heart failure; BNP, B-type natriuretic peptide; COR, Class of Recommendation; ED, emergency department; HF, heart failure; NT-proBNP, N-terminal pro-B-type natriuretic peptide; NYHA, New York Heart Association; and pts, patients.

7. Treatment of Stages A to D

7.3. Stage C

7.3.2. Pharmacological Treatment for Stage C HF With Reduced Ejection Fraction:

Recommendations

(See Figure 2 and Table 3).

7.3.2.10. Renin-Angiotensin System Inhibition With Angiotensin-Converting Enzyme Inhibitor
or Angiotensin Receptor Blocker or ARNI: Recommendations

Recommendations for Renin-Angiotensin System Inhibition With ACE Inhibitor or ARB or ARNI				
COR	LOE	Recommendations	Comment/Rationale	
		The clinical strategy of inhibition of the renin-	NEW : New clinical	
	ACE-I: A	angiotensin system with ACE inhibitors (Level of	trial data prompted	
		Evidence: A) (128-133), OR ARBs (Level of	clarification and	
		Evidence: A) (134-137), OR ARNI (Level of	important updates.	
Ι	ARB: A	<i>Evidence: B-R</i>) (138) in conjunction with evidence-		
		based beta blockers (9, 139, 140), and aldosterone		
		antagonists in selected patients (141, 142), is		
	ARNI: B-R	recommended for patients with chronic HFrEF to		
		reduce morbidity and mortality.		
		Angiotensin-converting enzyme (ACE) inhibitors reduce	morbidity and	
		mortality in heart failure with reduced election fraction (HFrEF) Randomized	
		controlled trials (RCTs) clearly establish the benefits of	ACE inhibition in	
		nations with mild moderate or severe symptoms of HE	and in nationts with or	
		without coronary artery disease (128, 133) ACE inhibitor	rs can produce merican	
		angioadama and should be given with coution to patients	with low systemication	
		blood progrupos renal insufficiency, or elevated sorum p	ataggium ACE	
		inhibitors also inhibit lininggo and increase levels of hrow	dultinin which can	
		inhibitors also inhibit kininase and increase levels of bradykinin, which can		
		induce cough but also may contribute to their beneficial effect through		
		vasodilation.		
		Angiotensin receptor blockers (ARBs) were developed with the rationale		
		that angiotensin II production continues in the presence of ACE inhibition,		
		driven through alternative enzyme pathways. ARBs do not inhibit kininase and		
		are associated with a much lower incidence of cough and angloedema than ACE		
See O	nline Data	inhibitors; but like ACE inhibitors, ARBs should be given	n with caution to	
Supple	ments 1, 2,	patients with low systemic blood pressure, renal insufficiency, or elevated		
1	8-20.	serum potassium. Long-term therapy with ARBs produces hemodynamic,		
		neurohormonal, and clinical effects consistent with those expected after		
		interference with the renin-angiotensin system and have been shown in RCTs		
		(134-137) to reduce morbidity and mortality, especially in ACE inhibitor–		
		intolerant patients.		
		In ARNI, an ARB is combined with an inhibitor of neprilysin, an enzyme		
		that degrades natriuretic peptides, bradykinin, adrenomedullin, and other		
		vasoactive peptides. In an RCT that compared the first approved ARNI,		
		valsartan/sacubitril, with enalapril in symptomatic patients with HFrEF		
		tolerating an adequate dose of either ACE inhibitor or ARB, the ARNI reduced		
		the composite endpoint of cardiovascular death or HF hospitalization		
		significantly, by 20% (138). The benefit was seen to a similar extent for both		
		death and HF hospitalization and was consistent across subgroups. The use of		
		ARNI is associated with the risk of hypotension and renal insufficiency and		
		may lead to angioedema, as well.		

		The use of ACE inhibitors is beneficial for	2013 recommendation	
т	ACE I. A	patients with prior or current symptoms of	repeated for clarity in	
L	ACE-I: A	chronic HF <i>r</i> EF to reduce morbidity and mortality	this section.	
		(128-133, 143).		
		ACE inhibitors have been shown in large RCTs to redu	ce morbidity and	
		mortality in patients with HFrEF with mild, moderate,	or severe symptoms of	
		HF, with or without coronary artery disease (128-133).	Data suggest that there	
		are no differences among available ACE inhibitors in the	neir effects on symptoms	
		or survival (143). ACE inhibitors should be started at lo	ow doses and titrated	
		upward to doses shown to reduce the risk of cardiovascular events in clinical		
		trials. ACE inhibitors can produce angioedema and sho	uld be given with	
		caution to patients with low systemic blood pressures, r	enal insufficiency, or	
		elevated serum potassium (>5.0 mEq/L). Angioedema	occurs in <1% of	
See On	line Data	patients who take an ACE inhibitor, but it occurs more	frequently in blacks and	
Supple	ment 18.	women (144). Patients should not be given ACE inhibit	tors if they are pregnant	
		or plan to become pregnant. ACE inhibitors also inhibit	t kininase and increase	
		levels of bradykinin, which can induce cough in up to 2	20% of patients but also	
		may contribute to beneficial vasodilation. If maximal doses are not tolerated,		
		intermediate doses should be tried; abrupt withdrawal of ACE inhibition can ion		
		lead to clinical deterioration and should be avoided.		
		Although the use of an ARNI in lieu of an ACE inhibitor for $HFrEF$ has		
		been found to be superior, for those patients for whom ARNI is not appropriate,		
		continued use of an ACE inhibitor for all classes of HF	rEF remains strongly	
_		aavisea.	2012	
		The use of ARBS to reduce morbially and mortality	2015 recommondation	
т		symptoms of chronic HErEE who are intolerent to	repeated for clarity	
I	AND, A	ACE inhibitors because of cough or angioedema	in this section	
		(134-137, 145, 146).	in this section.	
		ARBs have been shown to reduce mortality and HF hos	spitalizations in patients	
		with HFrEF in large RCTs (134-137). Long-term thera	py with ARBs in patients	
		with HFrEF produces hemodynamic, neurohormonal, and clinical effects		
		consistent with those expected after interference with the renin-angiotensin		
		system (145, 146). Unlike ACE inhibitors, ARBs do not inhibit kininase and are		
		associated with a much lower incidence of cough and angioedema, although		
See Online Data		kininase inhibition by ACE inhibitors may produce beneficial vasodilatory		
Supplem	ents 2 and	effects.		
19.		Patients intolerant to ACE inhibitors because of cou	ugh or angioedema	
		should be started on ARBs; patients already tolerating A	ARBs for other	
		indications may be continued on ARBs if they subseque	ently develop HF. ARBs	
		should be started at low doses and titrated upward, with	an attempt to use doses	
		shown to reduce the risk of cardiovascular events in clinical trials. ARBs should		
		be given with caution to patients with low systemic blo	od pressure, renal	
		insufficiency, or elevated serum potassium (>5.0 mEa/L). Although ARBs are		

		alternatives for patients with ACE inhibitor-induced angioedema, caution is		
		advised because some patients have also developed angioedema with ARBs.		
		Head-to-head comparisons of an ARB versus ARNI for HF do not exist.		
		For those patients for whom an ACE inhibitor or ARNI is	s inappropriate, use of	
		an ARB remains advised.		
		In patients with chronic symptomatic HFrEF	NEW : New clinical	
т	ARNI: R-R	NYHA class II or III who tolerate an ACE inhibitor	trial data necessitated	
-	ARIA. D-R	or ARB, replacement by an ARNI is recommended	this recommendation.	
		to further reduce morbidity and mortality (138).		
		Benefits of ACE inhibitors with regard to decreasing HF	progression,	
		hospitalizations, and mortality rate have been shown con-	sistently for patients	
		across the clinical spectrum, from asymptomatic to sever	ely symptomatic HF.	
		Similar benefits have been shown for ARBs in population	ns with mild-to-	
		moderate HF who are unable to tolerate ACE inhibitors.	In patients with mild-	
		to-moderate HF (characterized by either 1) mildly elevate	ed natriuretic peptide	
		levels, BNP [B-type natriuretic peptide] >150 pg/mL or N	NT-proBNP [N-	
		terminal pro-B-type natriuretic peptide] ≥600 pg/mL; or 2) BNP ≥100 pg/mL or		
		NT-proBNP \geq 400 pg/mL with a prior hospitalization in the preceding 12		
		months) who were able to tolerate both a target dose of enalapril (10 mg twice		
		daily) and then subsequently an ARNI (valsartan/sacubiti	ril; 200 mg twice daily,	
See Or	line Data	with the ARB component equivalent to valsartan 160 mg), hospitalizations and	
Suppler	ents 1 and	mortality were significantly decreased with the valsartan/	sacubitril compound	
Supple	18	compared with enalapril. The target dose of the ACE inhibitor was consistent		
	10.	with that known to improve outcomes in previous landmark clinical trials (129).		
		This ARNI has been approved for patients with symptom	atic HFrEF and is	
		intended to be substituted for ACE inhibitors or ARBs. HF effects and potential		
		off-target effects may be complex with inhibition of the neprilysin enzyme,		
		which has multiple biological targets. Use of an ARNI is	associated with	
		hypotension and a low-frequency incidence of angioeden	na. To facilitate	
		initiation and titration, the approved ARNI is available in	3 doses that include a	
		dose that was not tested in the HF trial; the target dose used in the trial was		
		97/103 mg twice daily (147). Clinical experience will provide further		
		information about the optimal titration and tolerability of ARNI, particularly		
		with regard to blood pressure, adjustment of concomitant	HF medications, and	
		the rare complication of angioedema (14).		
		ARNI should not be administered concomitantly	NEW : Available	
		with ACE inhibitors or within 36 hours of the last	evidence	
		dose of an ACE inhibitor (148, 149).	demonstrates a	
III:	B-R		potential signal of	
Harm			narm for a	
			concomitant use of	
			ACE inhibitors and	
			ARNI.	

		Oral neprilysin inhibitors, used in combination with ACE inhibitors can lead to			
		angioedema and concomitant use is contraindicated and should be avoided. A			
		medication that represented both a neprilysin inhibitor an	d an ACE inhibitor,		
Saa Onli	na Data	omapatrilat, was studied in both hypertension and HF, bu	it its development was		
See Olli	mont 2	terminated because of an unacceptable incidence of angio	bedema (148, 149) and		
Supplet	nent 5.	associated significant morbidity. This adverse effect was	thought to occur		
		because both ACE and neprilysin break down bradykinin	, which directly or		
		indirectly can cause angioedema (149, 150). An ARNI sh	nould not be		
		administered within 36 hours of switching from or to an	ACE inhibitor.		
III:	C EO	ARNI should not be administered to patients with a	NEW : New clinical		
Harm	C-EO	history of angioedema.	trial data.		
		Omapatrilat, a neprilysin inhibitor (as well as an ACE inl	hibitor and		
		aminopeptidase P inhibitor), was associated with a higher frequency of			
		angioedema than that seen with enalapril in an RCT of patients with HFrEF			
		(148). In a very large RCT of hypertensive patients, omapatrilat was associated			
		with a 3-fold increased risk of angioedema as compared with enalapril (149).			
		Blacks and smokers were particularly at risk. The high in	cidence of angioedema		
N/.	A	ultimately led to cessation of the clinical development of omapatrilat (151,ican			
		152). In light of these observations, angioedema was an exclusion criterion in			
		the first large trial assessing ARNI therapy in patients with hypertension (153)			
		and then in the large trial that demonstrated clinical benefit of ARNI therapy in			
		HFrEF (138). ARNI therapy should not be administered in patients with a			
		history of angioedema because of the concern that it will increase the risk of a			
		recurrence of angioedema.			
7.3.2.11. Ivabradine: Recommendation					

7.3.2.11. Ivabradine: Recommendation

Recommendation for Ivabradine						
COR	LOE	Recommendation	Comment/Rationale			
		Ivabradine can be beneficial to reduce HF	NEW : New clinical trial			
		hospitalization for patients with symptomatic	data.			
		(NYHA class II-III) stable chronic HFrEF				
Ha	B-R	(LVEF ≤35%) who are receiving GDEM*,				
		including a beta blocker at maximum tolerated				
		dose, and who are in sinus rhythm with a heart				
		rate of 70 bpm or greater at rest (154-157).				
		Ivabradine is a new therapeutic agent that selectively inhibits the I_f current in				
		the sinoatrial node, providing heart rate reduction. One RCT demonstrated the				
		efficacy of ivabradine in reducing the composite endpoint of cardiovascular				
See Onli	ine Data	death or HF hospitalization (155). The benefit of ivabradine was driven by a				
Supplement 4.		reduction in HF hospitalization. The study included patients with HFrEF				
		(NYHA class II-IV, albeit with only a modest representation of NYHA class IV				
		HF) and left ventricular ejection fraction (LVEF) \leq 35%, in sinus rhythm with a				
		resting heart rate of \geq 70 beats per minute. Patients enrolled included a small				
		number with paroxysmal atrial fibrillation (<40% of	the time) but otherwise in			

sinus rhythm and a small number experiencing ventricular pacing but with a
predominant sinus rhythm. Those with a myocardial infarction within the
preceding 2 months were excluded. Patients enrolled had been hospitalized for
HF in the preceding 12 months and were on stable GDEM* for 4 weeks before
initiation of ivabradine therapy. The target of ivabradine is heart rate slowing
(the presumed benefit of action), but only 25% of patients studied were on
optimal doses of beta-blocker therapy (9, 139, 140, 155). Given the well-proven
mortality benefits of beta-blocker therapy, it is important to initiate and up
titrate these agents to target doses, as tolerated, before assessing the resting
heart rate for consideration of ivabradine initiation (155).

*In other parts of the document, the term "GDMT" has been used to denote guideline-directed management and therapy. In this recommendation, however, the term "GDEM" has been used to denote this same concept in order to reflect the original wording of the recommendation that initially appeared in the "2016 ACC/AHA/HFSA Focused Update on New Pharmacological Therapy for Heart Failure: An Update of the 2013 ACCF/AHA Guideline for the Management of Heart Failure" (10).

Circulation

Figure 2. Treatment of HFrEF Stage C and D



Colors correspond to COR in Table 1. For all medical therapies, dosing should be optimized and serial assessment exercised.

*See text for important treatment directions.

[†]Hydral-Nitrates green box: The combination of ISDN/HYD with ARNI has not been robustly tested. BP response should be carefully monitored.

\$See 2013 HF guideline (9).

§Participation in investigational studies is also appropriate for stage C, NYHA class II and III HF.

ACEI indicates angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor-blocker; ARNI, angiotensin receptor-neprilysin inhibitor; BP, blood pressure; bpm, beats per minute; C/I, contraindication; COR, Class of Recommendation; CrCl, creatinine clearance; CRT-D, cardiac resynchronization therapy–device; Dx, diagnosis; GDMT, guideline-directed management and therapy; HF, heart failure; HF*r*EF, heart failure with reduced ejection fraction; ICD, implantable cardioverter-defibrillator; ISDN/HYD, isosorbide dinitrate hydral-nitrates; K+, potassium; LBBB, left bundle-branch block; LVAD, left ventricular assist device; LVEF, left ventricular ejection fraction; MI, myocardial infarction; NSR, normal sinus rhythm; and NYHA, New York Heart Association.

Drug	Initial Daily Dose(s)	Maximum Doses(s)	Mean Doses Achieved in Clinical Trials	References
ACE inhibitors				L
Captopril	6.25 mg TID	50 mg TID	122.7 mg QD	(158)
Enalapril	2.5 mg BID	10–20 mg BID	16.6 mg QD	(129)
Fosinopril	5–10 mg QD	40 mg QD	N/A	
Lisinopril	2.5–5 mg QD	20–40 mg QD	32.5-35.0 mg QD	(130)
Perindopril	2 mg QD	8–16 mg QD	N/A	
Quinapril	5 mg BID	20 mg BID	N/A	
Ramipril	1.25–2.5 mg QD	10 mg QD	N/A	
Trandolapril	1 mg QD	4 mg QD	N/A	
ARBs	-			
Candesartan	4–8 mg QD	32 mg QD	24 mg QD	(137)
Losartan	25–50 mg QD	50–150 mg QD	129 mg QD	(136)
Valsartan	20–40 mg BID	160 mg BID	254 mg QD	(134)
ARNI				
Sacubitril/ valsartan	49/51 mg BID (sacubitril/valsartan) (therapy may be initiated at 24/26 mg BID)	97/103 mg BID (sacubitril/valsartan)	375 mg QD; target dose: 24/26 mg, 49/51 mg OR 97/103 mg BID	He (138) Association
If channel inhibit	or			
Ivabradine	5 mg BID	7.5 mg BID	6.4 mg BID (at 28 d) 6.5 mg BID (at 1 y)	(155-157)
Aldosterone antag	gonists			
Spironolactone	12.5–25 mg QD	25 mg QD or BID	26 mg QD	(142)
Eplerenone	25 mg QD	50 mg QD	42.6 mg QD	(159)
Beta blockers				
Bisoprolol	1.25 mg QD	10 mg QD	8.6 mg QD	(160)
Carvedilol	3.125 mg BID	50 mg BID	37 mg QD	(161)
Carvedilol CR	10 mg QD	80 mg QD	N/A	
Metoprolol succinate extended release (metoprolol CR/XL)	12.5–25 mg QD	QD 200 mg QD 159 mg QD		(139)
Isosorbide dinitra	te and hydralazine			
Fixed-dose combination	20 mg isosorbide dinitrate / 37.5 mg hydralazine TID	40 mg isosorbide dinitrate / 75 mg hydralazine TID	90 mg isosorbide dinitrate / ~175 mg hydralazine QD	(162)
Isosorbide dinitrate and hydralazine	20–30 mg isosorbide dinitrate / 25–50 mg hydralazine	40 mg isosorbide dinitrate TID with 100 mg hydralazine TID	N/A	(163)

Table 3. Drugs Commonly Used for HFrEF (Stage C HF)

 TID or QD

 Modified (Table 15) from the 2013 HF guideline (9).

L

ACE indicates angiotensin-converting enzyme; ARB, angiotensin receptor blocker; ARNI, angiotensin receptorneprilysin inhibitor; BID, twice daily; CR, controlled release; CR/XL, controlled release/extended release; HF, heart failure; HF*r*EF, heart failure with reduced ejection fraction; N/A, not applicable; QD, once daily; and TID, 3 times daily.

Recommendations for Stage C HFpEF					
COR	LOE	Recommendations	Comment/Rationale		
I	В	Systolic and diastolic blood pressure should be controlled in patients with HFpEF in accordance with published clinical practice guidelines to prevent morbidity (164, 165).	2013 recommendation remains current.		
I	С	Diuretics should be used for relief of symptoms due to volume overload in patients with HFpEF.	2013 recommendation remains current.		
IIa	С	Coronary revascularization is reasonable in patients with CAD in whom symptoms (angina) or demonstrable myocardial ischemia is judged to be having an adverse effect on symptomatic HF <i>p</i> EF despite GDMT.	2013 recommendation remains current.		
IIa	С	Management of AF according to published clinical practice guidelines in patients with HF <i>p</i> EF is reasonable to improve symptomatic HF.	2013 recommendation remains current (Section 9.1 in the 2013 HF guideline).		
IIa	С	The use of beta-blocking agents, ACE inhibitors, and ARBs in patients with hypertension is reasonable to control blood pressure in patients with HFpEF.	2013 recommendation remains current.		
IIb	B-R	In appropriately selected patients with HFpEF (with EF \geq 45%, elevated BNP levels or HF admission	NEW: Current recommendation reflects		
See Online Data Supplement C.		>30 mL/min, creatinine <2.5 mg/dL, potassium <5.0 mEq/L), aldosterone receptor antagonists might be considered to decrease hospitalizations (83, 166, 167).	new KC I uata.		

			-		
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1.3.3. FI	alliacological	<i>Healineni</i> IOI		рег. кес	commenuations

Mechanistic studies have suggested that mineralocorticoid receptor antagonists can improve measures of diastolic function in patients with HF*p*EF, possibly by a similar effect on remodeling (83, 168).

The TOPCAT (Treatment of Preserved Cardiac Function Heart Failure With an Aldosterone Antagonist) trial (166) investigated the effects of spironolactone on a combined endpoint of death, aborted cardiac death, and HF hospitalization in patients with HFpEF. A small reduction (HR=0.89) in this composite endpoint did not reach statistical significance, although HF hospitalization was reduced (HR=0.83); known side effects of hyperkalemia and rising creatinine were seen more commonly in the treatment group (166). An unusual amount of regional variation was seen in this trial, prompting a post-hoc analysis (167) that showed that rates of the primary endpoint were 4-fold lower in Russia/Georgia than in North America and South America (the Americas). Rates in the Americas were comparable to those in other HFpEF trials (169, 170). The post-hoc analysis showed efficacy in the Americas (HR=0.83) but not in Russia/Georgia (HR=1.10). Moreover, a sample of the Russia/Georgia population, despite having been in the active treatment arm, had nondetectable levels of

the metabolite of spironolactone. These post-hoc analyses have significant limitations, but they suggest that in appropriately selected patients with symptomatic HF*p*EF (with ejection fraction [EF] \geq 45%, elevated BNP level or HF admission within 1 year, estimated glomerular filtration rate >30 mL/min creatinine <2.5 mg/dL, and potassium <5.0 mEq/L), particularly in those with elevated BNP levels, use of spironolactone might be considered with close monitoring of potassium and renal function. Confirmatory studies are required.

With regard to the use of mineralocorticoid receptor antagonists, creatinine should be <2.5 mg/dL in men or <2.0 mg/dL in women (or estimated glomerular filtration rate >30 mL/min) and potassium should be <5.0 mEq/L. Careful monitoring of potassium, renal function, and diuretic dosing represents best practices at initiation and during follow-up thereafter to minimize risk of hyperkalemia and worsening renal function.

IIb	В	The use of ARBs might be considered to decrease hospitalizations for patients with HF <i>p</i> EF (169).	2013 recommendation remains current.
III: No Benefit	B-R	Routine use of nitrates or phosphodiesterase-5 inhibitors to increase activity or QoL in patients	NEW : Current recommendation reflects
See Online Data Supplement C.		with HFPEF is ineffective (171, 172).	new data from RC1s.

Nitrate therapy can reduce pulmonary congestion and improve exercise tolerance in patients with HF*r*EF. However, the NEAT-HF*p*EF (Nitrate's Effect on Activity Tolerance in Heart Failure With Preserved Ejection Fraction) trial (171) randomized 110 patients with EF \geq 50% on stable HF therapy, not including nitrates, and with activity limited by dyspnea, fatigue, or chest pain, to either isosorbide mononitrate or placebo and found no beneficial effects on activity levels, QoL, exercise tolerance, or NT-proBNP levels. On the basis of this trial, routine use of nitrates in patients with HF*p*EF is not recommended. This recommendation does not apply to patients with HF*p*EF and symptomatic CAD for whom nitrates may provide symptomatic relief. Phosphodiesterase-5 inhibition augments the nitric oxide system by upregulating cGMP activity. The RELAX (Phosphodiesterase-5 Inhibition to Improve Clinical Status and Exercise Capacity in Heart Failure with Preserved Ejection Fraction) trial (172) randomized 216 patients with EF \geq 50% on stable HF therapy and with reduced exercise tolerance (peak observed VO₂ <60% of predicted) to phosphodiesterase-5 inhibition with sildenafil or placebo. This study did not show improvement in oxygen consumption or exercise tolerance.

III: No	C	Routine use of nutritional supplements is not	2013 recommendation
Benefit	C	recommended for patients with HFpEF.	remains current.

9. Important Comorbidities in HF

9.2. Anemia: Recommendations

Recommendations for Anemia						
COR	COR LOE Recommendations		Comment/Rationale			
IIb	IIb B-R In patients with NYHA class II and III HF and iron		NEW : New evidence			
		deficiency (ferritin <100 ng/mL or 100 to 300 ng/mL	consistent with			
See Onli	ne Data	if transferrin saturation is <20%), intravenous iron	therapeutic benefit.			
Suppler	nent D.	replacement might be reasonable to improve	-			
		functional status and QoL(173, 174).				
Routine baseline assessment of all patients with HF includes an evaluation for anemia in addition to other						
baseline labo	ratory measu	rements. Anemia is independently associated with HF disea	se severity, and iron			

deficiency appears to be uniquely associated with reduced exercise capacity. When iron deficiency is diagnosed and after full evaluation for cause, intravenous repletion of iron, especially in the setting of concomitant hepcidin deficiency in HF, may improve exercise capacity and QoL. Studies examining correction of iron deficiency in HF have demonstrated improvement in surrogate endpoints, such as QoL, NT-proBNP, and LVEF; however, controlled trials have been underpowered to detect reductions in hard clinical endpoints. The FAIR-HF (Ferric Carboxymaltose Assessment in Patients With Iron Deficiency and Chronic Heart Failure) trial (173) demonstrated improvements in NYHA class and functional capacity over a short-term exposure. The CONFIRM-HF (Ferric Carboxymaltose Evaluation on Performance in Patients With Iron Deficiency in Combination with Chronic Heart Failure) trial (174) included a larger cohort of patients (n=304) and demonstrated improvements in 6-minute walk test. A meta-analysis of 5 prospective controlled studies (631 patients) evaluated the effect of intravenous iron on deaths, hospitalizations, and other events in patients with HF and iron deficiency (175). Patients receiving intravenous iron experienced limited but statistically significant improvements in functional capacity and LVEF but no reduction in mortality rate. The FAIR-HF 2 trial is underway to further address the potential benefit of intravenous iron in HF associated with iron deficiency. Therefore, a strong recommendation for intravenous iron repletion must await the results of an appropriately powered trial on morbidity and mortality. There is an uncertain evidence base for oral iron repletion in the setting of anemia associated with HF.

III: No	B-R	In patients with HF and anemia, erythropoietin-	NEW: Current
Benefit		stimulating agents should not be used to improve	recommendation reflects
See Onli Suppler	ne Data nent D.	morbidity and mortality (176).	demonstrating absence of therapeutic benefit.

Small studies evaluating the treatment of anemia in patients with HF have suggested a trend toward improvement in functional capacity and reduction in hospitalization with the use of erythropoietin-stimulating agents (177-182), but results have varied (183) and have been limited because of sample size. Although a meta-analysis of 11 RCTs (n=794) comparing erythropoietin-stimulating agents to control in patients with HF demonstrated significant improvements in 6-minute walk, exercise duration, peak VO₂, NYHA functional status, EF, BNP, HFrelated hospitalizations, and QoL (184), in the STAMINA-HeFT (Study of Anemia in Heart Failure) trial (183), darbepoetin alfa was not associated with significant clinical benefits. In the largest RCT to date (n=2,278), correction of anemia with darbopoetin alfa did not result in benefit and resulted in a significant increase in the risk of thromboembolic events and a nonsignificant increase in fatal and nonfatal strokes, supporting findings from other trials (176, 185-188). In summary, the strongest evidence on erythropoietin-stimulating agent therapy in HF suggests lack of benefit and increased adverse events. Therefore, erythropoietin-stimulating agent therapy cannot be recommended in patients with HF and anemia.

9.5. Hypertension (New Section)

9.5.1. Treating Hypertension to Reduce the Incidence of HF: Recommendation

Recommendation for Prevention					
COR	LOE	Recommendations	Comment/Rationale		
Ι	B-R	In patients at increased risk, stage A HF, the optimal	NEW : Recommendation		
		blood pressure in those with hypertension should be	reflects new RCT data.		

See Online Data	less than 130/80 mm Hg (189-193).				
Supplements E and F.					
A large RCT demonstrated that in those with increased cardiovascular risk (defined as age >75 years, established					
vascular disease, chronic renal disease, or a Framingham Risk Score >15%), control of blood pressure to a goal					
systolic pressure of <120 mm Hg, as determined by blood pressure assessment as per research protocol, was					
associated with a significant reduction in the incidence of HF (191) and an overall decrease in cardiovascular					
death. Blood pressure measurements as generally taken in the office setting are typically 5 to 10 mm Hg higher					
than research measurements; thus, the goal of <130/80 mm Hg is an approximation of the target blood pressure					
in conventional practice. Targeting a significant reduction in systolic blood pressure in those at increased risk					
for cardiovascular disease is a novel strategy to prevent HF.					

9.5.2. Treating Hypertension in Stage C HFrEF: Recommendation

Recommendation for Hypertension in Stage C HFrEF						
COR	LOE	Recommendation	Comment/Rationale			
_		Patients with HFrEF and hypertension should be	NEW : Recommendation			
1	С-ЕО	prescribed GDMT titrated to attain systolic blood	has been adapted from			
		pressure less than 130 mm Hg (191).	recent clinical trial data			
See Online Data			but not specifically tested			
Supplements E and F.			per se in a randomized			
			trial of patients with HF.			
Clinical trials evaluating goal blood pressure reduction and optimal blood pressure-lowering agents in the setting						
of HFrEF and concomitant hypertension have not been done. However, it is apparent that in those patients at						

higher risk, blood pressure lowering is associated with fewer adverse cardiovascular events. GDMT for HFrEF with agents known to lower blood pressure should consider a goal blood pressure reduction consistent with a threshold now associated with improved clinical outcomes but not yet proven by RCTs in a population with HF.

9.5.3. Treating Hypertension in Stage C HFpEF: Recommendation

Recommendation for Hypertension in Stage C HFpEF						
COR	LOE	Recommendation	Comment/Rationale			
I	C-LD	Patients with HFpEF and persistent hypertension after management of volume overload should be	NEW : New target goal blood pressure based on			
See Online Data Supplements E and F.		prescribed GDM1 titrated to attain systolic blood pressure less than 130 mm Hg (167, 169, 170, 194- 199).	recent clinical trial data.			

The use of nitrates in the setting of HFpEF is associated with a signal of harm and, in most situations, should be avoided. For many common antihypertensive agents, including alpha blockers, beta blockers, and calcium channel blockers, there are limited data to guide the choice of antihypertensive therapy in the setting of HFpEF (172). Nevertheless, RAAS inhibition with ACE inhibitor, ARB (especially mineralocorticoid receptor antagonists), and possibly ARNI would represent the preferred choice. A shared decision-making discussion with the patient influenced by physician judgment should drive the ultimate choice of antihypertensive agents.

9.6. Sleep Disordered Breathing: Recommendations

(Moved from Section 7.3.1.4, Treatment of Sleep Disorders in the 2013 HF guideline.)

Recommendations for Treatment of Sleep Disorders					
COR	LOE	Recommendations	Comment/Rationale		
	CID	In patients with NYHA class II–IV HF and suspicion	NEW : Recommendation		
11a	C-LD	of sleep disordered breathing or excessive daytime	reflects clinical necessity		
See Onli	ine Data	sleepiness, a formal sleep assessment is reasonable	to distinguish obstructive		
Suppler	nent G.	(200, 201).	versus central sleep apnea.		
Sleep disorde	ers are commo	on in patients with HF. A study of adults with chronic HF tr	eated with evidence-based		
therapies fou	nd that 61% h	ad either central or obstructive sleep apnea (202). It is clini	cally important to		
distinguish o	bstructive slee	ep apnea from central sleep apnea, given the different respo	nses to treatment. Adaptive		
servo-ventila	tion for centra	al sleep apnea is associated with harm (203). Continuous po	sitive airway pressure		
(CPAP) for c	bstructive sle	ep apnea improves sleep quality, reduces the apnea-hypopr	lea index, and improves		
nocturnal oxy	ygenation (20	0, 201).			
IIb	рр	In patients with cardiovascular disease and	NEW : New data		
110	D-K	obstructive sleep apnea, CPAP may be reasonable to	demonstrate the limited		
See Onli	ine Data	improve sleep quality and daytime sleepiness (204).	scope of benefit expected		
Suppler	ment G.		obstructive sleep appea		
In patients w	ith sleep apne	a, a trial evaluated the impact of CPAP with usual therapy	versus usual therapy alone		
on subsequer	nt cardiovascu	lar events, including HF (204). In this RCT of >2,700 patie	ents, there was no evidence		
of benefit on	cardiovascula	r events at a mean follow-up of 3.7 years for CPAP plus us	sual care compared with		
usual care alo	one. Improven	nents in sleep quality were noteworthy and represented the	primary indication for		
initiating CP.	AP treatment	(204). However, in patients with atrial fibrillation (AF) (a f	requent comorbidity noted		
with HF), the use of CPAP for obstructive sleep apnea was helpful. In a trial of 10,132 patients with AF and					
obstructive sleep apnea, patients on CPAP treatment were less likely to progress to more permanent forms of AF					
than were par	tients without	CPAP (205).			
III. Harm	рр	In patients with NYHA class II–IV HFrEF and	NEW : New data		
III: Harm	D-N	central sleep apnea, adaptive servo-ventilation	demonstrate a signal of		
See Online Data		causes harm (203).	narm when adaptive		
Supplement G.			for central sleep appea.		
Mortality rate (all cause and cardiovascular) was higher with adaptive servo-ventilation plus GDMT than with					
GDMT alone in a single RCT to test the addition of adaptive servo-ventilation (\geq 5 hours/night, 7 days/week) to					
GDMT in patients with HFrEF and central sleep apnea (203). A similar risk has been seen in another trial, and a					
third trial of	third trial of adaptive servo-ventilation in central sleep apnea and HF was aborted because of ethical concerns.				
The weight of evidence does not support the use of adaptive servo-ventilation for central sleep apnea in HFrEF.					

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 natriuretic peptide biomarker

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Appendix 1. Author Relationships With Industry and Other Entities (Relevant)—2017 ACC/AHA/HFSA Focused Undate of the 2013 ACCF/AHA Guideline for the Management of Heart Failure (December 2015)

I ocuseu e punte		ne for the Manage	mene of fieu	i e i anui e (Dee	ember 2013)			
Committee Member	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness	Voting Recusals By Section*
Clyde W. Yancy (<i>Chair</i>)	Northwestern University Feinberg School of Medicine, Division of Cardiology—Professor of Medicine and Chief; Diversity and Inclusion—Vice Dean	None	None	None	None	None An	None	None
Mariell Jessup (Vice Chair)	Fondation Leducq—Chief Scientific Officer	None	None	None	None	None As	None	None
Biykem Bozkurt	Baylor College of Medicine, Department of Medicine — Professor of Medicine; Cardiology Section, DeBakey VA Medical Center — Chief; The Mary and Gordon Cain Chair & W.A. "Tex" and Deborah Moncrief, Jr. — Chair; Winters Center for Heart Failure Research — Director; Cardiovascular Research Institute — Associate Director	None	None	None	• Novartis	None	None	7.3.2.10, 7.3.2.11, 7.3.3, and 9.5.
Javed Butler	Stony Brook University— Division Chief of Cardiology	 Bayer† Boehringer Ingelheim CardioCell† Luitpold Medtronic Merck† Novartis† Relypsa† Takeda Trevena† Z Pharma 	• Novartis†	None	• Amgen (DSMB)†	None	None	7.3.2.10, 7.3.2.11, 7.3.3, and 9.5.

		• Zensun						
Donald E. Casey, Jr	Thomas Jefferson College of Population Health— Faculty; Alvarez & Marsal IPO4Health— Principal and Founder	None	None	None	None	None	None	None
Monica M. Colvin	University of Michigan— Associate Professor of Medicine, Cardiology	None	None	None	None	None	None	None
Mark H. Drazner	University of Texas Southwestern Medical Center—Professor, Internal Medicine	None	None	None	None	None	None	None
Gerasimos S. Filippatos	National and Kapodistrian University of Athens; Attikon University Hospital, Department of Cardiology, Heart Failure Unit—Professor of Cardiology	None	None	None	 Bayer† Bayer (DSMB) Novartis† Servier Pharmaceuticals† Vifor 	None Am	None erican art sociation。	7.3.2.10, 7.3.2.11, 7.3.3, 9.2, and 9.5.
Gregg C. Fonarow	Ahmanson-UCLA Cardiomyopathy Center— Director; UCLA Division of Cardiology—Co-Chief	 Amgen Janssen Pharmaceuticals Novartis† 	None	None	Novartis†	None	None	7.3.2.10, 7.3.2.11, 7.3.3, and 9.5.
Michael M. Givertz	Brigham and Women's Hospital—Professor of Medicine	Merck Novartis	None	None	None	None	None	7.3.2.10, 7.3.2.11, 7.3.3, and 9.5.
Steven M. Hollenberg	Cooper University Hospital— Director, Coronary Care Unit, Professor of Medicine	None	None	None	None	None	None	None
JoAnn Lindenfeld	Vanderbilt Heart and Vascular Institute—Director, Advanced Heart Failure and Transplant Section—Professor of Medicine	 Abbott Janssen Pharmaceuticals Novartis Relypsa† ResMed† 	None	None	 AstraZeneca Novartis† 	None	None	6.3, 7.3.2.10, 7.3.2.11, 7.3.3, 9.5, and 9.6.
Frederick A. Masoudi	University of Colorado, Anschutz Medical Campus—Professor of Medicine, Division of Cardiology	None	None	None	None	None	None	None
Patrick E. McBride	University of Wisconsin School of Medicine and Public Health— Professor of Medicine and Family	None	None	None	None	None	None	None

	Medicine; Associate Director, Preventive Cardiology							
Pamela N. Peterson	University of Colorado, Denver Health Medical Center— Associate Professor of Medicine, Division of Cardiology	None	None	None	None	None	None	None
Lynne Warner Stevenson	Brigham and Women's Hospital Cardiovascular Division— Director, Cardiomyopathy and Heart Failure Program	None	None	None	 Novartis— PARENT trial (PI) NHLBI— INTERMACS (Co–PI) 	None	None	7.3.2.10, 7.3.2.11, 7.3.3, and 9.5.
Cheryl Westlake	Azusa Pacific University, School of Nursing, Doctoral Programs— Professor	None	None	None	None	None Ha	None art sociation.	None

This table represents the relationships of committee members with industry and other entities that were determined to be relevant to this document. These relationships were reviewed and updated in conjunction with all meetings and/or conference calls of the writing committee during the document development process. The table does not necessarily reflect relationships with industry at the time of publication. A person is deemed to have a significant interest in a business if the interest represents ownership of \geq 5% of the voting stock or share of the business entity, or ownership of \geq \$5,000 of the fair market value of the business entity, or if funds received by the person from the business entity exceed 5% of the person's gross income for the previous year. Relationships that exist with no financial benefit are also included for the purpose of transparency. Relationships in this table are modest unless otherwise noted.

According to the ACC/AHA, a person has a *relevant* relationship IF: a) The *relationship or interest* relates to the same or similar subject matter, intellectual property or asset, topic, or issue addressed in the *document*; or b) the *company/entity* (with whom the relationship exists) makes a drug, drug class, or device addressed in the *document*, or makes a competing drug or device addressed in the *document*; or c) the *person or a member of the person's household* has a reasonable potential for financial, professional, or other personal gain or loss as a result of the issues/content addressed in the *document*.

*Writing committee members are required to recuse themselves from voting on sections to which their specific relationships with industry and other entities may apply. †Significant relationship.

ACC indicates American College of Cardiology; ACCF, American College of Cardiology Foundation; AHA, American Heart Association; DCRI, Duke Clinical Research Institute; DSMB, data safety monitoring board; HFSA, Heart Failure Society of America; NHLBI, National Heart, Lung, and Blood Institute; INTERMACS, Interagency Registry for Mechanically Assisted Circulatory Support; PARENT, Pulmonary artery pressure reduction with entresto; UCLA, University of California, Los Angeles; and VA, Veterans Affairs.

Appendix 2. Reviewer Relationships With Industry and Other Entities (Comprehensive)—2017 ACC/AHA/HFSA

Focused Update of the 2013 ACCF/AHA Guideline for the Management of Heart Failure (October 2016)

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership / Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness
Kim K. Birtcher	Official Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines	University of Houston College of Pharmacy—Clinical Professor	• Jones & Bartlett Learning	None	None	None	None	None
Akshay S. Desai	Official Reviewer—HFSA	Brigham and Women's Hospital—Director, Heart Failure Disease Management, Advanced Heart Disease Section, Cardiovascular Division; Associate Professor of Medicine, Harvard Medical School	 Medscape Cardiology* Merck Novartis* Relypsa* St. Jude Medical* 	None	None	None	 Novartis* Thoratections 	None
Anita Deswal	Official Reviewer—AHA	Michael E. DeBakey VA Medical Center—Chief of Cardiology; Director, Heart Failure Program; Baylor College of Medicine— Professor of Medicine	None	None	None	• NIH*	 AHA AHA (GWTG Steering Committee)† HFSA† 	None
Dipti Itchhaporia	Official Reviewer—ACC Board of Trustees	Newport Coast Cardiology— Robert and Georgia Roth Endowed Chair for Excellence in Cardiac Care; Director of Disease Management	None	None	None	None	• St. Jude Medical	None
Ileana L. Piña	Official Reviewer—AHA	Montefiore Medical Center— Associate Chief for Academic Affairs, Cardiology; Professor of Medicine & Epidemiology and Population Health— Albert Einstein College of Medicine	• Relypsa	None	None	None	None	None
Geetha	Official	University of Missouri-Kansas	None	None	None	None	None	None

Raghuveer	Reviewer—ACC Board of Governors	City School of Medicine— Professor of Pediatrics; Children's Mercy Hospital—						
James E. Udelson	Official Reviewer—HFSA	Pediatric Cardiology Tufts Medical Center—Chief, Division of Cardiology	• Lantheus Medical Imaging	None	None	 Gilead (DSMB) GlaxoSmithKline (DSMB) NHLBI Otsuka 	 Abbott Laboratories AHA* Circulation / Circulation: Heart Failure† HFSA (Executive Council)† Pfizer/ GlaxoSmithKline Sunshine Heart 	None
Mary Norine Walsh	Official Reviewer—ACC Board of Trustees	St Vincent Heart Center of Indiana—Medical Director, Heart Failure and Cardiac Transplantation	None	None	None	None	 Corvia Medical Otsuka PCORI Thoratec 	None
David A. Baran	Organizational Reviewer—ISHLT	Newark Beth Israel Medical Center—Director of Heart Failure and Transplant Research	 Maquet Otsuka* 	• Novartis	None	• XDx* • NIH*	None	None
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Appendix 3. Abbreviations

ACE = angiotensin-converting enzyme ARB = angiotensin-receptor blocker ARNI = angiotensin receptor-neprilysin inhibitor BNP = B-type natriuretic peptide BP = blood pressureCOR = Class of Recommendation CPAP = continuous positive airway pressure EF = ejection fractionGDMT = guideline-directed management and therapy $HF_{p}EF =$ heart failure with preserved ejection fraction HFrEF = heart failure with reduced ejection fraction LOE = Level of Evidence LVEF = left ventricular ejection fraction NT-proBNP = N-terminal pro-B-type natriuretic peptide tio QoL = quality of life RCT = randomized controlled trial





2017 ACC/AHA/HFSA Focused Update of the 2013 ACCF/AHA Guideline for the Management of Heart Failure: A Report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Failure Society of America Clyde W. Yancy, Mariell Jessup, Biykem Bozkurt, Javed Butler, Donald E. Casey, Jr, Monica M. Colvin, Mark H. Drazner, Gerasimos S. Filippatos, Gregg C. Fonarow, Michael M. Givertz, Steven M. Hollenberg, JoAnn Lindenfeld, Frederick A. Masoudi, Patrick E. McBride, Pamela N. Peterson, Lynne Warner Stevenson and Cheryl Westlake

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*Significant relationship.

†No financial benefit.

ACC indicates American College of Cardiology; AHA, American Heart Association; ABIM, American Board of Internal Medicine; AHRQ, Agency for Healthcare Research and Quality; DCRI, Duke Clinical Research Institute; DSMB, data safety monitoring board; GWTG, Get With The Guidelines; HF, heart failure; HFSA, Heart Failure Society of America; HRSA, Heath Resources and Services Administration; HSAG, Health Services Advisory Group; IMPROVE-HF, Registry to Improve the use of Evidence-Based Heart Failure Therapies in the Outpatient Setting; INTERMACS, Interagency Registry for Mechanically Assisted Circulatory Support; JAHA, Journal of the American Heart Association; PCORI, Patient Centered Outcomes Research Institute; PI, principal investigator; PRT, pharmaceutical round table; NIDDK, National Institute of Diabetes and Digestive and Kidney Diseases; NIH, National Institutes of Health; NHLBI, National Heart, Lung, and Blood Institute; NIAID, National Institute of Allergy and Infectious Diseases; UCLA, University of California, Los Angeles; and VA, Veterans Affairs.

Appendix 1. Author Relationships With Industry and Other Entities (Relevant)-2017 ACC/AHA/HFSA Focused

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Update of the 2013 ACCF/AHA Guideline for the Management of Heart Failure (December 2015)

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ACC indicates American College of Cardiology; AHA, American Heart Association; DCRI, Duke Clinical Research Institute; DSMB, data safety monitoring board; HFSA, Heart Failure Society of America; INTERMACS, Interagency Registry for Mechanically Assisted Circulatory Support; PARENT, Pulmonary Artery Pressure Reduction With Entresto; UCLA, University of California, Los Angeles; and VA, Veterans Affairs.

(Section numbers correspond to the 2013 full-text guideline.)

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Key Search Terms: Heart Failure, Angiotensin Receptor-Neprilysin Inhibitor, Ivabradine, Angiotensin Receptor Blockers, Angiotensin-Converting Enzyme Inhibitors, Beta Blockers, Angioedema, Natriuretic Peptides, Ferric Carboxymaltose, Iron deficiency, hypertension, sleep apnea, natriuretic peptide biomarker.

Master Abbreviation List:

1° indicates primary; 2°, secondary; ~, approximately; 6MWT, 6 min walk test; ACE, angiotensin-converting enzyme; ACEI indicates angiotensin-converting-enzyme inhibitor; ACS, acute coronary syndrome; ADHF, acute decompensated heart failure; ADHERE, Acute Decompensated Heart Failure National Registry; AF, atrial fibrillation; AHI, apneahypopnea index; AHRQ, Agency for Healthcare Research and Quality; AIRE, Acute Infarction Ramipril Efficacy; ALLHAT, Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial; ALT, alanine aminotransaminase; AMI; acute myocardial infarction; APE, acute pulmonary embolism; ARB, angiotensin-receptor blocker; AKI/ARF, acute kidney injury/acute renal failure; ARNI, angiotensin receptor-neprilysin inhibitor; ASA, aspirin; AST, aspartate transaminase; ATLAS, Assessment of Treatment with Lisinopril and Survival; AUC, area under the curve; AV, atrioventricular; ; BEAUTIFUL, Morbidity-Mortality Evaluation of the If Inhibitor Ivabradine in Patients With Coronary Disease and

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Left-Ventricular Dysfunction; BID, twice a day; BL, baseline; BNP, plasma B-type natriuretic peptide; BP, blood pressure; bpm, beats per minute; BSA, body surface area; CABG, coronary artery bypass graft; CAD, coronary artery disease; CANPAP, Canadian Continuous Positive Airway Pressure for Patients with Central Sleep Apnea and Heart Failure Trial; CCB, calcium channel blockers; CKD, chronic kidney disease; cGMP, cyclic guanosine monophosphate; CHARM, Candesartan in Heart Failure Assessment of Reduction in Mortality and Morbidity; CHD, chronic heart disease; CI, confidence interval; CM, contrast media; CONFIRM-HF, Ferric carboxymaltose evaluation on performance in patients with iron deficiency in combination with chronic heart failure: CONSENSUS Cooperative North Scandinavian Enalapril Survival Study; COPD, chronic obstructive pulmonary disease; CPAP, Continuous positive airway pressure; Cr, creatinine; CRT, cardiac resynchronization therapy; CSA, central sleep apnea; cTnl, cardiac troponin I; CTR, cardiothoracic ratio; CV, cardiovascular; CVA, cerebrovascular accident; CVD, cardiovascular disease; C/W, compared with; DBP, diastolic blood pressure; DM, diabetes mellitus; DOSE-AHF, Diuretic Optimization Strategy Evaluation in Acute HF; DPB, diastolic blood pressure; ECG, electrocardiography; ED, emergency department; EF, ejection fraction; eGFR, estimated glomerular filtration rate: ELAN-HF, European Collaboration on Acute Decompensated Heart Failure: ESRD, end-stage renal disease: EMPHASIS, Eplerenone in Mild Patients Hospitalization and Survival Study in Heart Failure; EQ-5D, EuroQoL five dimensions guestionnaire; ET, ; FAIR-HF, Ferinject Assessment in Patients with Iron Deficiency and Chronic Heart Failure; FCM, ferric carboxymaltose; FU, follow-up; GDEM, guideline-directed evaluation and management; GDMT, guideline-directed management and therapy; GP, ; HCM, ; HDL, high density lipoprotein; HEAAL study, effects of high-dose versus low-dose losartan on clinical outcomes in patients with heart failure; HF, heart failure; HF, DEF, Heart failure with preserved ejection fraction; h/o, history of; HF, EF, Heart failure with reduced ejection fraction; HR, hazard ratio; hs-CRP, high sensitivity Creactive protein; HTN, hypertension; HYVET, Hypertension in the Very Elderly Trial; Hx, history; ICD, implantable cardioverter defibrillator; ID, iron deficiency; IDI, integrated discrimination improvement; IHD, ischemic heart disease; IMPRESS, Comparison of Vasopeptidase Inhibitor, Omapatrilat, and Lisinopril on Exercise Tolerance and Morbidity; I-PRESERVE, Irbesartan in Heart Failure with Preserved Ejection Fraction Study; IQR, interguartile range; ITT, intent to treat; IV, intravenous; KCCQ, Kansas City Cardiomyopathy Questionnaire; LCZ, ; LV, left ventricular; LVD, Left ventricular dysfunction; LVEF, left ventricular ejection fraction; LVEDD; left ventricular end-diastolic dimension; LVH, left ventricular hypertrophy; MACE, major adverse cardiac event; MI, myocardial infarction; MR-proANP, ; MR-proADM, ; MRA, mineralocorticoid receptor antagonists; MTD, maximal tolerated dose: MV, mitral valve; MWT, minute walk test: N/A, not available; NEAT-HFpEF, Nitrate's Effect on Activity Tolerance in Heart Failure With Preserved Ejection Fraction; NEP, neutral endopeptidase; NNH, number needed to harm; NNT, number needed to treat; NP, natriuretic peptide; NRI, net reclassification improvement; NS, nonsignificant; NSTEMI, non-ST-elevation myocardial infarction; NT-proBNP, N-terminal pro-B-type natriuretic peptide; NYHA, New York Heart Association; OCTAVE, The Omapatrilat Cardiovascular Treatment vs. Enalapril; ONTARGET, The Ongoing Telmisartan Alone and in Combination With Ramipril Global Endpoint Trial; OPTIMIZE-HF, Organized Program to Initiate Lifesaving Treatment in Hospitalized Patients with Heart Failure; OR, odds ratio; OSA, obstructive sleep apnea; OVERTURE, Omapatrilat Versus Enalapril Randomized Trial of Utility in Reducing Events; PAD, peripheral artery disease; PARADIGM-HF, Prospective Comparison of ARNI With ACE to Determine Impact on Global Mortality and Morbidity in Heart Failure: PARAMOUNT, Prospective Comparison of ARNI With ARB on Management of Heart Failure With Preserved Ejection Fraction; PAP, positive airway pressure; PCI, percutaneous coronary intervention; PCP, Primary Care Physician; PDE, phosphodiesterase; PEP-CHF, Perindopril in Elderly People With Chronic Heart Failure; PGA, patient global assessment; PPM, permanent pacemaker; PSG, polysomnography; PTCA, percutaneous transluminal coronary angioplasty; PONTIAC, NT-proBNP Guided Primary Prevention of CV Events in Diabetic Patients; PRIMA, Can Pro-Brain-Natriuretic Peptide Guided Therapy of Chronic Heart Failure Improve Heart Failure Morbidity and Mortality?: PROTECT, ProBNP Outpatient Tailored Chronic Heart Failure Therapy; pts. patients: PVD, peripheral vascular disease; QoL, guality of life: RAAS, renin-angiotensinaldosterone system; RAS, renin-angiotensin system; RCT, randomized controlled trial; RED-HF, Reduction of events by darbepoetin alfa in heart failure; RELAX, Phosphodiesterase-5 Inhibition to Improve Clinical Status and Exercise Capacity in Heart Failure With Preserved Ejection Fraction; ROC, receiver-operating characteristic; RR, relative risk; SBP, systolic blood pressure; SCr, serum creatinine; SERVE-HF, Adaptive Servo-Ventilation for Central Sleep Apnea in Systolic Heart Failure; SHEP, Systolic Hypertension in the Elderly Program; SHIFT, Systolic Heart Failure Treatment with the If Inhibitor Ivabradine Trial; SIGNIFY, Study Assessing the Morbidity–Mortality Benefits of the If Inhibitor Ivabradine in Patients with Coronary Artery Disease; SOB, shortness of breath; SPRINT, Systolic Blood Pressure Intervention Trial; SR, systematic review; SSS, sick sinus syndrome; STARBRITE, the Strategies for Tailoring Advanced Heart Failure Regimens in the Outpatient Setting; STARS-BNP, Systolic Heart Failure Treatment Supported by BNP; STEMI, ST-elevation myocardial infarction; STOP-HF, St. Vincent's Screening to Prevent Heart Failure; SUPPORT, Supplemental Benefit of ARB in Hypertensive Patients With Stable Heart Failure Using Olmesartan; SURVIVE, Survival of Patients with Acute Heart Failure in Need of Intravenous Inotropic Support; TIA, transient ischemic attack; TIME-CHF, ; TOPCAT, Treatment of Preserved Cardiac Function Heart Failure With an Aldosterone Antagonist; TRANSCEND, the Telmisartan Randomised Assessment Study in ACE Intolerant Subjects With Cardiovascular Disease; TSAT, transferrin saturation; UA, unstable angina; UL, ; UPSTEP, Use of Peptides in Tailoring Heart Failure Project; VF, ventricular fibrillation; VHD, valvular heart disease VT, ventricular tachycardia; and w/o, without.

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Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% Cl)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Biomarker Studies Per	tinent to Stage A / B HI	Patients			
PONTIAC Huelsmann et al. 2013 (1) 23810874 • Medical University of Vienna • Roche Pharma AG	Aim: To assess the effectiveness of neurohumoral therapy for the prevention of cardiac events in pts with type 2 DM with increased biomarker NT- proBNP Study type: RCT Size: 300	Inclusion criteria: Pts with type 2 DM, age ≥18 y, elevated NT-proBNP (≥125 pg/mL) Exclusion criteria: Free of heart disease, chronic infections or malignancies, systemic cortisone treatment, renal replacement therapy, nondiabetic conditions that lowered life expectancy to <1 y and absence of reliable contraception in women of	Intervention: Individualized up-titration of RAS antagonists and beta blockers in addition to diabetes treatment (150), treated at cardiology clinic Comparator: "Control" group treated for diabetes, (150), treated at diabetes care units	 <u>1° endpoint</u>: Hospitalization or death due to cardiac disease following 24 mo Results: Significant reduction of 1° endpoint in intervention group (HR: 0.351; 95% CI: 0.127–0.975; p=0.044) <u>1° Safety endpoint</u>: BP was significantly reduced in both intervention and control (p<0.05); heart rate was only reduced in the intensified group (p=0.004) 	 All-cause hospitalizations, HF hospitalizations and unplanned CV hospitalizations or death (p<0.05 reduction) Study limitations: Absence of pt randomization for treatment, pt population mainly Caucasian, statistical analysis done without adjustment of co-variates Pts treated with a RAS antagonist/beta-blocker and the dosage reached higher in intensified group (p<0.0001) No difference in NT-proBNP levels
STOP-HF Ledwidge et al. 2013 (2) <u>23821090</u>	Aim: To establish efficacy of BNP screening <u>and</u> collaborative care in at-risk	Inclusion criteria: Pts ≥40 y, and history of HTN (on meds ≥1 mo), hypercholesterolemi	Intervention: BNP screening at BL and annually and protocol referral for BNP ≥50 pg/mL for echocardiography and	 <u>1° endpoint</u>: LV dysfunction (systolic: LVEF <50% or diastolic: E/E' ratio >15) with or without newly diagnosed 	 Emergency hospitalizations for major MACE [40 vs. 22 (0.60 OR; 95% CI: .45-0.81; p=.002)]] CV investigations more likely to be done in the intervention group with
 Heartbeat Trust, Health Research 	population in reducing newly	a, obesity, vascular disease including	collaborative care. (697)	HF(with symptoms of HF requiring admission to	BNP levels ≥50 pg/mL Increase in RAAS agents in the

Data Supplement A. RCTs and Meta-analyses With Biomarkers (Section 6.3)

Board of the Irish Government; and European Commission Framework Programme. The Heartbeat Trust received unrestricted grants from Pfizer, A. Menarini, Alere, Roche, Takeda, Abbott, Covidien, and Servier.	diagnosed HF and prevalence of significant LV systolic and /or diastolic dysfunction. <u>Study type</u> : RCT (unblinded) <u>Size</u> : 1,374	CAD,, cerebrovascular disease or peripheral vascular disease, DM, arrhythmia therapy, or moderate to severe valvular disease Exclusion criteria: Established LV systolic dysfunction, symptomatic HF, diagnosis compromising survival	Comparator: Usual 1° care (677)	hospital, confirmed by d/c summary) • 59 (8.7 %) vs. 37 (5.3%) (0.55 OR; 95% CI: 0.37– 0.82; p=0.003)	 intervention group In the subgroup with BNP levels ≥50 pg/mL, increase in BNP levels in the intervention group was ~1/2 of that in the control group The results might not be applicable to general population (single center), non-blinding introduces bias. Event rate was lower than expected. Cost-effectiveness unclear. Incremental value of and cut-off of BNP may change in population studied.
Meta-Analyses or SRs	of RUIS of NP Guided	Therapy in Stage C HF			
Brunner-La Rocca et al. 2015 (3) <u>26419999</u>	Aim: To assess which HF pts benefit from NT-pro BNP therapy Study type: Meta-Analysis Size: 2,137 pts from 8 NT- proBNP trials	Inclusion criteria: Studies that included individual pt data HF <i>p</i> EF and HFrEF. EF ≤45% Exclusion criteria: Pts with unknown LVEF, STARBRITE study, 1° meta- analyses that aggregated data	Intervention: (NT-pro)BNP-guided therapy and HF/EF (1,731) <u>Comparator</u> : (NT-pro)BNP- guided therapy and HF <i>p</i> EF (301)	 <u>1° endpoint:</u> All-cause mortality and admission for HF Results: Lower mortality in HF/EF with guided treatment (HR: 0.78; 95% CI: 0.62–0.97; p=0.03). Lesser HF admissions in HF/EF (HR: 0.80; 95% CI: 0.67–0.97; p=0.02) 	 NT pro BNP-guided treatment harmful in HF<i>p</i>EF without HTN and in pts with renal failure Limitations: Bias due to exclusion of aggregate data, Lack of specific testing for diagnosis of comorbidities, absence of comorbidity index, insufficient sample size for pts with HF<i>p</i>EF, treatment management aspects unaddressed and statistical tests are not powerful
Don-Wauchope et al. 2015 (4) <u>25448029</u>	Aim: Review evidence of SRs regarding utility of NPs in clinical practice. Study type: Review of SRs	Inclusion criteria: SRs that authors were aware of through their participation in an AHRQ comparative effectiveness review.	Intervention: NP-guided therapy <u>Comparator:</u> Clinically-guided care	 <u>1° endpoint</u> 8 SRs assessed all-cause mortality and "generally found there was a benefit." 4 SRs examined all cause-hospitalization and did not find decrease with NP- 	 Underlying SRs largely comprised analysis of the same RCTs. Results were qualitative.

	<u>Size</u> : 9 reviews	<u>Exclusion criteria</u> : N/A		guided therapy • 4 SRs assessed HF hospitalization and "consistently" found a significant reduction with NP-quided therapy	
Xin W. et al. 2015 (5) <u>24888383</u>	Aim: To assess the effects of NP- guided treatment of chronic HF on outcomes Study type: Meta-analysis Size: 14 studies, 3,004 pts	Inclusion criteria: Prospective RCTs with adult HF pts comparing the effects of BNP or NT- proBNP-guided therapy with clinically guided therapy	Intervention: BNP or NT-proBNP-guided therapy (1,503) Comparator: Clinically guided therapy (1,501)	 <u>1° endpoints</u>: All-cause mortality, HF hospitalization, all-cause hospitalization, safety (adverse events) <u>Results</u>: Compared with clinical group, BNP-guided treatment significantly decreased the risk of HF-related hospitalization (RR: 0.79; 95% CI: 0.63–0.98; p=0.03), although did not significantly affect the risk of all-cause mortality (RR: 0.94, 95% CI: 0.81–1.08, p=0.39) or all-cause hospitalization (RR: 0.97; 95% CI: 0.89–1.07; p=0.56). <u>1° Safety endpoint</u>: NP-guided therapy was not associated with increased risk for serious adverse events. 	 BNP-guided therapy improved LV systolic function in HF pts (LVEF: weighted mean difference=2.80%, 95% CI: 0.90–4.69%; p=0.01), But did not significantly affect NYHA class or QoLs (p=ns)
Troughton RW et al. 2014 (6) <u>24603309</u>	Aim: To assess the effects of NP- guided treatment of chronic HF on outcomes Study type: Meta-analysis	Inclusion criteria: RCTs reporting all- cause mortality and comparing BNP- guided treatment of HF with clinically guided treatment and 1 study (PROTECT trial) that did not	Intervention: BNP-guided therapy (1,006) Comparator: Clinically guided therapy (994)	1° endpoint: • All-cause mortality Results: • All-cause mortality was significantly reduced by NP- guided treatment [HR: 0.62 (0.45–0.86); p=0.004]	 HF hospitalizations were reduced in the NP-guided group, compared with clinically guided pts [HR: 0.80 (0.67–0.94); p=0.009] as were CV admissions [HR: 0.82 (0.67–0.99); p=0.048] Each of the included RCTs was relatively small and 2 trials did not

	<u>Size:</u> 11 studies, 2,000 pts	report mortality (11 studies, 9 with individual pt data) Exclusion criteria: For 2 studies, data from the 3rd ('usual care') groups were not included.		 Significant interaction between age and treatment efficacy (p=0.028), with a survival benefit for BNP- guided vs. clinical treatment in pts <75 y [HR: 0.62 (0.45– 0.85); p=0.004] but not in pts ≥75 y [HR: 0.98 (0.75–1.3); p=ns] 	provide individual pt data.
De Vecchis et al. 2014 (7) <u>24522083</u>	Aim: To assess the effects of NP- guided treatment of chronic HF on outcomes Study type: Meta-analysis Size: 6 studies, 1,775 pts	Inclusion criteria • RCT to a strategy of titrating drug therapy based on the level of a circulating NP (BNP or NT-proBNP) compared to clinical conventional criteria, and they reported all-cause mortality. Should have included >60 pts and its follow-up should have been longer than 90 d.	Intervention: BNP or NT-proBNP-guided therapy Comparator: Clinically guided therapy	 <u>1° endpoint</u>: Combined endpoint of all-cause mortality and HF hospitalization <u>Results:</u> NP-guided therapy for outpatients with HF was shown to be associated with a decreased risk of death and HF hospitalizations (OR: 0.64; 95% CI: 0.43–0.95; p=0.026) 	 Limitations: Each of the included RCTs was relatively small Benefit was not seen in some of the studies
Balion et al. 2014 (8) <u>25074674</u>	Aim: To assess the effects of NP- guided treatment of chronic HF on outcomes Study type: SR Size: 9 RCTs; 2,104 pts	Meta-analysis was not done due to study heterogeneity.	Intervention: BNP or NT-proBNP-guided therapy (1,503) Comparator: Clinically guided therapy (1,501)	 <u>1° Outcome:</u> Review: Overall, there was a wide variation in study design and how parameters were reported including pt selection, BL characteristics, therapy goals, BNP/NT-proBNP cutpoint, and outcome types. The strength of evidence for the outcome of mortality, reported in 7 studies, was found to be low due to inconsistency and 	N/A

				imprecision.	
Savarese et al. 2013 (9) <u>23472172</u>	Aim: To determine whether NP-guided (BNP or NT- proBNP) therapy, compared to clinically guided therapy, improves outcomes Study type: Meta-analysis Size: 12 trials enrolling 2,686 participants (730 in BNP, 1,956 in NT-proBNP related trials)	Inclusion criteria: All randomized trials reporting clinical endpoints (all- cause mortality and/or HF related hospitalization and/or all-cause hospitalization) with comparison of BNP or NT-proBNP guided therapy vs. a control group in chronic HF pts	Intervention: • BNP-guided therapy: BNP-guided: 373 • NT-proBNP guided: 872 Comparator: Clinically guided therapy • BNP group control 357 • NT-proBNP group control 1,084 • Separate analyses on pts ≤ or >75 y using data reported in 3 trials.	 <u>1° endpoints</u> All-cause mortality, all-cause hospitalization, HF hospitalization Results: NP-guided therapy (either BNP or NT-proBNP) significantly reduced all-cause mortality (OR: 0.738; 95% CI: 0.596–0.913; p=0.005) and HF related hospitalization (OR: 0.554; 95% CI: 0.399–0.769; p=0.000), but not all-cause hospitalization (OR: 0.803; 95% CI: 0.629–1.024; p=0.077) 	• When separately assessed, NT- proBNP-guided therapy reduced all-cause mortality (OR: 0.717; 95% CI:0.563–0.914; p=0.007) and HF hospitalization (OR: 0.531; 95% CI: 0.347–0.811; p=0.003), but not all-cause hospitalization (OR: 0.779; CI:0.414–1.465; p=0.438), whereas BNP-guided therapy did not significantly reduce all-cause mortality (OR: 0.814; CI:0.518–1.279; p=0.371), HF related hospitalization (OR: 0.599; 95% CI: 0.303–1.187; p=0.14) or all-cause hospitalization (OR: 0.726; 95% CI:0. 0.509 – 1.035; p=0.077) • Analysis from 3 trials showed the composite outcome of all-cause mortality and HF hospitalization was significantly reduced by NP- guided therapy in younger pts (\leq 75 y) (OR: 0.449; 95% CI: 0.207– 0.973; p=0.043), but not in older pts ($>$ 75 y) (OR: 0.800; 95% CI: 0.423–1.513; p=0.5).
(10) <u>23602555</u>	To assess the effects of NP- guided treatment of chronic HF on all- cause mortality and HF hospitalization	Studies with >40 pts and involved comparison of BNP- guided vs. guideline-guided drug therapy of the pts with chronic HF in the outpatient	BNP-guided therapy <u>Comparator:</u> Clinically guided therapy	 Combined end point of all- cause mortality and HF hospitalization Results: Significantly decreased risk of all-cause mortality (RR: 0.83; 95% CI: 0.69–0.99; p=0.035; and HF 	rehospitalization was significantly decreased in the pts <70 y (RR: 0.45; 95% CI: 0.33–0.61; p=0.000; or with BL higher BNP (≥2114 pg/mL) (RR: 0.53; 95% CI: 0.39–0.72; p=0.000)

	Study type: Meta-analysis Size: 11 studies, 2,414 pts	setting		rehospitalization (RR: 0.75; 95% CI: 0.62–0.91; p=0.004; in the BNP-guided therapy group.	
Felker et al. 2009 (11) 19699866	Aim: To determine whether titration of therapy based on NP measurements improves mortality in chronic HF Study type: Meta-analysis Size: 6 studies; 1,627 pts	Inclusion criteria Prospective RCTs of pts with chronic HF randomized pts to a strategy of titrating medical therapy based on the level of a circulating biomarker compared to a parallel control group, reporting all- cause mortality	Intervention: BNP-guided therapy <u>Comparator:</u> Clinically guided therapy	 <u>1° endpoint</u>: All-cause mortality Results: Significant mortality advantage for biomarker- guided therapy (HR: 0.69, 95% CI: 0.55–0.86) compared to control 	N/A
Porapakkham et al. 2010 (12) <u>20308637</u> RCTs of NP Guided Th	Aim: To determine whether BNP guided therapy improves CV outcomes in chronic HF <u>Study type:</u> Meta-analysis <u>Size:</u> 8 studies; 1,726 pts	Inclusion criteria Eligible RCTs were those that enrolled >20 pts and involved comparison of BNP- guided drug therapy vs. usual clinical care of the pt with chronic HF in an outpatient setting	Intervention: BNP-guided therapy <u>Comparator:</u> Clinically guided therapy	<u>1° endpoint:</u> • All-cause mortality Results: Significantly lower risk of all- cause mortality (RR: 0.76; 95% CI: 0.63–0.91; p=0.003) in the BNP-guided therapy group compared with the control group	 In pts <75 y, all-cause mortality was significantly lower in the BNP-guided group (RR: 0.52; 95% CI: 0.33–0.82; p=0.005). No reduction in mortality with BNP-guided therapy in pts ≥75 y (RR: 0.94; 95% CI: 0.71–1.25; p=0.70). All-cause hospitalization and survival free of any hospitalization was not significantly different between groups (RR: 0.82; 95% CI: 0.64–1.05; p=0.12 and RR: 1.07; 95% CI: 0.85–1.34; p=0.58, respectively). Additional % pts achieving target doses of ACE-inhibitors and beta blockers 21% and 22% in the BNP group and 11.7% and 12.5% in the control group, respectively.

Troughton et al. 2000 (13)	<u>Aim</u> : To assess the	Inclusion criteria: Ambulatory pts with	Intervention: (NT-pro)BNP-guided therapy	 <u>1° endpoints</u>: Death, CV hospitalization 	Changes in LVEF, QoL, renal function, and adverse events were
<u>10791374</u>	effects of NT- proBNP-guided	LVEF <40% and symptomatic HF	with a target of NT-proBNP level	and outpatient HF event	similar in both groups.
	treatment of chronic HF on outcomes	(NYHA II-IV)	<200 pmol	Results: • Fewer CV events (death, begyitate or HE	reduced total CV events, and delayed time to first event
	Study type: RCT	Pts with unknown LVEF	Comparator: Standardized clinical assessment	decompensation) in the NT- proBNP group than in the	NP was reduced significantly and
	<u>Size</u> : 69 pts	Follow up : Minimum 6 mo	(clinical group)	clinical group (19 vs. 54; p=0.02) • At 6 mo. 27% of pts in the	NP guidance changed therapy
		(median 9.5 mo)		BNP group and 53% in the clinical group had experienced a first CV event (n=0.034)	
STARS-BNP Jourdain et al. 2007 (14) <u>17448376</u>	Aim: To evaluate the prognostic impact of a therapeutic strategy using plasma BNP Study type: RCT Size: 220 pts	Inclusion criteria: Ambulatory NYHA class II to III pts considered optimally treated Exclusion criteria: N/A Follow up : median 15 mo	Intervention: BNP-guided therapy Target : BNP <100 pg/mL <u>Comparator:</u> Medical treatment according to either current guidelines (clinical group)	 <u>1° endpoint</u> HF-related death or hospital stay for HF Results: Mean dosages of ACE inhibitors and beta blockers significantly higher in the BNP group (p<0.05), BNP-guided strategy reduced the risk of HF related death or hospital stay for HF (24% vs. 52%, p<0.001), mainly obtained through an increase in ACE inhibitor and beta blocker 	 NP guidance changed therapy Unknown whether BNP-guided therapy resulted in reduction in BNP levels
TIME-CHF Pfisterer et al. 2009 (15) <u>19176440</u>	Aim: To compare 18-mo outcomes of N- terminal BNP- guided vs. symptom guided HF therapy	Inclusion criteria: Ambulatory HF pts 60 y with systolic HF (LVEF ≤45%), NYHA class of II or greater, prior HF hospitalization within	Intervention: Uptitration of guideline- based treatments to BNP level of ≤2 times of UL (BNP-guided therapy) Targets:	1° endpoints: 18 mo survival free of all- cause hospitalizations Results: N-terminal BNP and	 Survival free of hospitalization for HF was higher among those in the N-terminal BNP-guided group (72% vs. 62%, respectively; HR: 0.68 [95% CI: 0.50–0.92]; p=0.01). N-terminal BNP-guided therapy

	Study type: RCT Size: 499 pts	1 y, and N-terminal BNP level of ≥2 times the upper limit of normal.	NT-proBNP <400 pg/mL if age <75 y, NT-proBNP <800 pg/mL if 75 y Comparator: Uptitration of guideline- based treatments to reduce symptoms to NYHA class of II or less (symptom guided therapy)	 symptom-guided therapy resulted in similar rates of survival free of all-cause hospitalizations (41% vs. 40%, respectively; HR: 0.91 [95% CI: 0.72–1.14]; p=0.39) BNP guidance changed therapy (higher doses of ACE inhibitors, ARB, Beta blockers and higher use of spironolactone) NT-ProBNP levels were not different between groups 	 improved outcomes in pts 60 to 75 y of age but not in those ≥75 y of age (p<0.02 for interaction). QoL improvements were similar in both the N-terminal BNP-guided and symptom guided strategies
BATTLESCARRED Lainchbury et al. 2009 (16) 20117364	Aim: to compare the effects of NT- proBNP)-guided therapy with those of intensive clinical management and with usual care Study Type: RCT (Australia hospitals) Size: 364 pts	Inclusion criteria: Pts admitted to a single hospital with HF, NT-proBNP >50 pmoL/l or 400 pg/mL.(included HF <i>p</i> EF)	Intervention: Outpatient post d/c therapy guided by NT-proBNP levels Target: NT-proBNP <150 pmoL/l (1,270 pg/mL) Comparators: Therapy guided by intensive clinical management, <u>or</u> according to usual care	<u>1° endpoints</u> : Mortality Results: 1-y mortality was less in both the hormone (9.1%) and clinically-guided (9.1%) groups compared with usual care (18.9%; p=0.03)	 3 y mortality was selectively reduced in pts ≤75 y receiving hormone guided treatment (15.5%) compared with either clinically managed treatment (30.9%; p=0.048) or usual care (31.3%; p=0.021). NP guidance changed therapy NT-ProBNP levels were not different between groups
Berger et al. 2010 (17) <u>20170790</u>	Aim: To investigate whether the addition of NT- proBNP-guided, intensive pt management to multidisciplinary care improves outcome in pts following hospitalization due to HF	Inclusion criteria: Pts admitted to a hospital with HF, NYHA III or IV on admission, Cardiothoracic Index>0.5 or LVEF <40%	Intervention: Outpatient post discharge discontinue • BM: NT-proBNP-guided, intensive up-titration of medication by HF specialists in high-risk pts. • Target: NT-proBNP (<2,200 pg/mL)	 <u>1° endpoints:</u> Hospitalization Results: Pt management reduced HF hospitalization (488 D) compared with the multidisciplinary care (1254 D) and usual care (1,588 d) groups (p<0.0001) Combined end point of death or HF rehospitalization was lower 	 NT-ProBNP levels were not different between groups: Pt management group had the highest proportion of RAAS inhibition triple-therapy Death rate was similar between the pt management (22%) and multidisciplinary care groups (22%), but was lower compared with the usual care group (39%; vs. pt management: p<0.02; vs. multidisciplinary care: p<0.02)

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	Study Type: RCT (8 Viennese hospitals) Size: 278 pts		specialist-therapeuticrecommendations andhome care by a HF nurseUsual care	 in the BM (37%) than in the multidisciplinary care group (50%; p<0.05) and in the multidisciplinary care than in the usual care group (65%; p=0.04) NT-ProBNP levels were lowered in guided pt management arm 	
PRIMA Eurlings et al. 2010 (18) <u>21144969</u>	Aim: To assess whether management by an individualized NT- proBNP target would lead to improved outcome compared with HF management guided by clinical assessment alone Study Type: RCT Size: 345 pts	Inclusion criteria: Hospitalized HF pts with for decompensated, symptomatic HF with NT-proBNP levels >1,700 pg/mL at admission (included HF <i>p</i> EF)	Intervention: After discharge discontinue out pt management guided by an individually set NT- proBNP (n=174) defined by the lowest level at discharge or 2 wk thereafter. Comparators: Clinically-guided outpatient management (n=171)	<u>1° endpoints:</u> Number of d alive outside the hospital after index <u>Results:</u> _Management guided by NT- proBNP target did not significantly improve the 1° endpoint p=0.49)	 In the NT-proBNP-guided group mortality was lower, as 46 pts died (26.5%) vs. 57 (33.3%) in the clinically guided group, but this was not statistically significant (p=0.21) Individualized NT-proBNP target increased the use of HF medication (p=0.006)
SIGNAL HF Trial Persson et al. 2010 (19) 20876734	Aim: To investigate if NT- proBNP-guided therapy in HF pts in 1° care would improve clinical outcomes over and above treatment according to guidelines Study Type: RCT (Sweden 1° care centers)	Inclusion criteria: Ambulatory HF pts NYHA class II-IV, LVEF <50% and NT-proBNP levels males >800, females >1,000 ng/	Intervention: Structured treatment of HF according to guidelines with or without NT-proBNP monitoring • Target: At least a 50% reduction from BL NT- proBNP	1° endpoints: Composite endpoint of d alive, d out of hospital and symptom score Results: There were no differences between the groups concerning either the 1° endpoint (p=0.28) or its components (CV) death, p=0.93; CV hospitalization, p=0.28	Treatment doses of beta blockers and RAS blockers were markedly increased towards target doses a similar degree in both groups

	<u>Size:</u> 252 pts				
STARBRITE Trial Shah et al. 2011 (20) <u>21807321</u>	Aim: Whether outpatient diuretic management guided by BNP and clinical assessment better compared with clinical assessment alone Study Type: Multicenter (3) RCT Size: 130	Inclusion criteria Hospitalized HF pts with LEVF ≤35% Exclusion criteria Serum creatinine >3.5 mg/dL and ACS	Intervention: Outpatient post discharge BNP and clinical assessment guided therapy Comparator: Clinical assessment alone.	<u>1° endpoints</u> : Composite endpoint of d alive and d out of hospital, <u>Results:</u> No significant difference HR: 0.72; 95% CI: 0.41–1.27; p=0.25	 Change in serum creatinine, or change in SBP not different BNP strategy pts received significantly more ACE inhibitors, beta blockers
PROTECT Study Gaggin et al. 2012 (21) 22858078	Aim: Whether elders benefit from NP- guided HF care Study Type: Single center RCT Size: 151	Inclusion criteria Chronic HF pts with LV systolic dysfunction	Intervention: Management guided by NT- proBNP with a goal to lower NT-proBNP ≤1000 pg/mL over 10 mo Comparator: Standard of care	1° endpoints: Total CV events in 2 age categories 75 and ≥75 y Results: Pts ≥75 y with NT-proBNP management had lowest rate of CV events (1.76 events per pt with standard of care vs. 0.71 events per pt with NT-proBNP guide, p=0.03)	 Improvement in QoL, LVEF, and indices of LV volume with guided approach NP guidance changed therapy: greater use of aldosterone antagonists and lesser use of loop diuretics in the guided therapy group (no difference in ACE inhibitors or beta blockers)
UPSTEP-study group Karlstrom et al. 2011 (22) <u>21715446</u>	Aim: To determine whether BNP- guided HF treatment improves morbidity and/or mortality	Inclusion criteria Ambulatory HF NYHA II-IV, LVEF <40% and elevated BNP levels	Intervention: BNP-guided (BNP) with a goal <150 or 300 ng/L for elderly Comparator: Conventional (CTR) HF treatment	1° endpoints:Combined death and worsening/hosp for HFResults:No significant differences 1° outcome (p=0.18)	 No differences for d out of hospital, and younger vs. elderly. Subgroup analysis: improved survival (p<0.0001 for the 1° outcome) among responders with >30% decrease in BL BNP value vs. nonresponders.

	Study Type: Multicenter RCT- probe design Size: 279				
Maisel et al. 2002 (23) <u>12124404</u>	Aim: To validate and characterize the use of BNP in the diagnosis of HF in pts with dyspnea Study type: Prospective, blinded, diagnostic accuracy study Size: 1,856	Inclusion criteria: Pts who came to the emergency department with acute dyspnea Exclusion criteria: Age <18 y and those whose dyspnea was clearly not secondary to HF (i.e., those with trauma or cardiac tamponade), acute myocardial infarction, unstable angina, or renal failure	Intervention: Comparisons of BNP values among diagnostic groups including HF and non HF pts Comparator: Non-HF pts such as pulmonary disease, cor pulmonale	<u>1° endpoint:</u> Diagnostic accuracy of BNP at a cutoff of 100 pg/mL was 83.4%. The negative predictive value of BNP <50 pg/mL was 96%. <u>Secondary endpoint :</u> In multiple logistic-regression analysis, measurements of BNP added significant independent predictive power to other clinical variables in models predicting which pts had HF	•Used in conjunction with other clinical information, rapid measurement of BNP is useful in establishing or excluding the diagnosis of acute HF failure in pts with acute dyspnea
van Kimmenade et al. 2006 (24) <u>16860029</u>	Aim: To analyze the role of NT-pro-BNP in diagnosis of HF in pts presenting with dyspnea, the so- called natriuretic peptide gray zone. NT-pro-BNP concentrations, clinical characteristics, and 60-d mortality were studied in acutely dyspneic pts from an international	Inclusion criteria: Acutely dyspneic pts Exclusion criteria: With trauma or cardiac tamponade), acute myocardial infarction, unstable angina, or renal failure	Intervention: Comparisons of NT-pro- BNP among diagnostic groups including HF and non-HF pts Comparator: Non-HF pts such as pulmonary disease, cor pulmonale	<u>1° endpoint:</u> Subjects with HF and diagnostically elevated NT- pro-BNP concentrations had the highest mortality rates, subjects without HF and NT- pro-BNP concentrations < 300 ng/L had the lowest mortality rates, and subjects with gray- zone NT-pro-BNP had intermediate outcomes, irrespective of their final diagnoses.	•Adding specific clinical information to NT-pro-BNP improves diagnostic accuracy in subjects with intermediate NT-pro-BNP concentrations. Mortality rates in subjects with intermediate NT-pro- BNP concentrations are lower than in those with NT-pro-BNP concentrations diagnostic for HF but are higher than in subjects with NT-pro-BNP concentrations less than the gray zone

	multicenter study				
	Study type: Prospective, blinded, diagnostic accuracy study Size: 1.256				
Maisel et al. 2004 (25) <u>15364340</u>	Aim: To examine the relationships among BNP levels and HF severity, clinical decision making, and outcomes Study type: Multicenter, prospective, blinded, diagnostic accuracy study Size: 464	Inclusion criteria:Pts over the age of 18y presenting to theED with HF andwho receivedtreatment in the EDor hospitaladmission for HFwere included.Exclusion criteria:Current MI or ACSwith ST-segmentdeviation of ≥1 mm,renal failurerequiring dialysis, orpts with a baselineBNP concentrationof ≤100 pg/mL wereexcluded	Intervention: Physicians were blinded to the actual BNP level and subsequent BNP measurements. Comparator: Comparison between severity of HF determined by physicians or BNP and outcomes	<u>1° endpoint:</u> ED doctor's intention to admit or discharge a pt had no influence on 90-d outcomes, while the BNP level was a strong predictor of 90-d outcome. The 90-d combined event rate (HF visits or admissions and mortality) in the group of pts admitted with BNP <200 pg/mL and >200 pg/mL was 9% and 29%, respectively (p=0.006).	 In pts presenting to the ED with HF, there is a disconnect between the perceived severity of HF by ED physicians and severity as determined by BNP levels. The BNP levels can predict future outcomes and thus may aid physicians in making triage decisions about whether to admit or discharge pts. Emerging clinical data will help further refine biomarker-guided outpatient therapeutic and monitoring strategies involving BNP.
O'Connor et al. 2010 (26) <u>20185037</u>	Aim: To identify high-risk HF pts at hospital discharge Study type: Predictive modeling using variables obtained during hospitalization in the ESCAPE trial	Inclusion criteria: hospitalized with severe HF, LVEF ≤30%, SBP ≤125 mmHg, Exclusion criteria: creatinine >3.5 mg/dL, prior inotrope use	Derivation cohort: ESCPAPE trial, n=423 Validation cohort: FIRST trial, n=471	 <u>1° endpoint</u>: 6-mo mortality and death or rehospitalization rates (64%) Multivariate discharge predictors of death included: BNP, per doubling (HR: 1.42), cardiac arrest or mechanical ventilation, yes/no (HR: 2.54), BUN, per 20 mg/dL increase (HR: 1.22) and sodium, per unit mEq/L increase (HR: 0.93) 	 A simplified discharge score discriminated mortality risk from 5% (score=0) to 94% (score=8). Bootstrap validation demonstrated good internal validation for the model (c-index 0.78) Limitations: ESCAPE represented pts with severe LV dysfunction and advanced symptoms (not the general population of acute HF) managed at experienced centers; exclusion of pts with characteristics

	<u>Size</u> : 423				known to be associated with worse outcomes (e.g., creatinine >3.5 mg/dL, requiring inotropes)
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Search Terms and Date: natriuretic peptides, heart failure, human, last 5 years. Last search done on April 18, 2016.

Data Supplement B. Nonrandomized Trials/ Observational Studies/ Registries for Changes in or Discharge NP Levels in ADHF – Biomarkers (Section 6.3)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Primary Endpoint and Results (P values, OR or RR & 95 % CI)	Summary / Conclusion / Comments
Bayés-Genís et al. 2005 (27) <u>15948093</u>	Aim: Percentage of NT- proBNP reduction during admission and its prognostic significance Study type: NR Prospective cohort Size: 74 pts	Inclusion criteria: Pts diagnosed with acute HF in emergency department and who had follow-up evaluation for 6 & 12 mo after admission Follow up :12 mo	 <u>1° endpoints:</u> Percent reduction in NT-proBNP and its association with CV mortality <u>Results:</u> The area under the ROC curve for % NT-proBNP reduction to predict CV death was 0.78 (95% CI: 0.66–0.90; p=0.002) 	 30% NT-proBNP reduction percentage cutoff value had 75% accuracy for the identification of high-risk pts and was the only variable that was associated with CV death in multivariate analysis (OR: 4.4; 95% CI: 1.12–17.4; p=0.03). Study relatively old and small
Verdiani et al. 2008 (28) <u>18545069</u>	Aim: To evaluate the prognostic significance of NT-proBNP % reduction during ADHF Study type: Prospective cohort Size: 120 pts	Inclusion criteria: Pts consecutively admitted with ADHF Follow up: 6 mo	 <u>1° endpoint</u> Percent reduction in NT-proBNP and its association with CV mortality <u>Results:</u> In ROC, the mean AUC for NT-ProBNP % reduction was 0.63 (95% CI: 0.51–0.75; p=0.04) for the composite endpoint (death or readmission), and 0.81 (95% CI: 0.65–0.97, p=0.01) for CV mortality at risk of events. 	 NT-ProBNP reduction percentage <30% was the best cut off for the identification of pts Study relatively old and small

Bettencourt et al. 2004 (29) <u>15451800</u>	Aim: To compare 18 mo outcomes of NT-BNP- guided vs. symptom guided HF therapy Study type: Prospective cohort single center study Size: 182 pts	Inclusion criteria: Consecutive ADHF pts defined by ESC or Framingham criteria Follow up: 6 mo	 <u>1° endpoints</u>: Death or readmission <u>Results:</u> Pts were classified into 3 groups: (1) decreasing NT-proBNP levels by at least 30% (n=82), (2) no significant modifications on NT-proBNP levels (n=49), and (3) increasing NT-proBNP levels by at least 30% (n=25). Among the 64 pts discharged without volume overload, a positive association between change in NT-proBNP and outcome was observed (HR: 2.66; 95% CI: 0.77–9.18 for change <30%; HR: 16.04; 95% CI: 9.49 – 52 02 for increase ≥30% compared with 	 Pts demonstrating a ≥30% increase in NT-proBNP levels during the course of their admission had the most adverse prognosis Study relatively old and small
Kociol et al. 2013 (30) 23250981	Aim: Examine_relationship between markers of decongestion and symptom relief and clinical outcomes Study type: retrospective analysis of the RCT, DOSE- AHF Size: 308 pts Aim:	Inclusion criteria: Pts enrolled in DOSE-AHF Follow up: 60 d	 b2:02 for indicade 200% compared with those with decreasing NT-proBNP by at least 30% <u>1° endpoints</u>: Time to death, first rehospitalization or emergency department visit <u>Results:</u> Of the weight loss, fluid loss, and NT-proBNP reduction, only % reduction in NT-proBNP was significantly associated with symptom relief (r=0.13; p=0.04). Reduction in NT-proBNP Associated with better outcome (NT-proBNP HR: 0.95; 95% CI: 0.91–0.99 per 10% reduction). The discharge BNP had the best 	Favorable changes in each of the 3 markers of decongestion were associated with improvement in time to death, rehospitalization, or emergency department visit at 60 d
Kociol et al. 2011 (31) <u>21743005</u>	Aim: To examine if admission, discharge, or change from admission to discharge BNP measure is the most important predictor of long-term outcomes	Inclusion criteria: Linked pts ≥65 y of age from hospitals in OPTIMIZE-HF to Medicare claims Follow up: 1 y	• The discharge BNP had the best performance and was the most important characteristic for predicting 1 y mortality (HR for log transformation: 1.34; 95% CI: 1.28–1.40) and 1 y death or rehospitalization (HR: 1.15; 95% CI: 1.12– 1.18).	• Compared with a clinical variables, discharge BNP model improved risk reclassification and discrimination in predicting each outcome (1 y mortality: NRI: 5.5%, p<0.0001; IDI: 0.023, p<0.0001; 1-y mortality or rehospitalization: NRI: 4.2%, p<0.0001; IDI: 0.010, p<0.0001)

Flint KM et al. 2014 (32) <u>24922626</u>	Study type: Retrospective analysis -from OPTIMIZE HF Trial Size: 7,039 pts Aim: To examine if admission, discharge, or change from admission to discharge BNP measure is the most important predictor of outcomes Study type: Retrospective analysis from VA database Size: 109,875 pts	Inclusion criteria: All hospital discharges with a 1° diagnosis of HF in the Veterans Affairs Health Care System from 2006 to 2009. Follow up: 30 d	1° endpoints: • 30 d readmission rate for HF Results: 30 d HF readmission was associated with elevated admission BNP, elevated discharge BNP, and smaller percent change in BNP from admission to discharge. • Pts with a discharge BNP ≥1,000 ng/L had an unadjusted 30 d HF readmission rate over 3 times as high as pts whose discharge BNP was ≤200 ng/L (15% vs. 4 1%)	 Discharge BNP had the greatest effect (C-statistic, 0.639–0.664 [p<0.0001]; NRI, 9% [p<0.0001]). Large sample size
ELAN-HF Score Salah et al. 2014 (33) 24179162	Aim: To examine if admission, discharge, or change from admission to discharge BNP measure is the most important predictor of outcomes Study type: Individual pt data meta- analyses of prospective cohort studies Size: 1,301 pts	Inclusion criteria: Pts from 7 prospective cohorts with pts admitted because of clinically validated ADHF, discharged alive, and NT-proBNP measurements available at admission and at discharge Follow up: 180 d	 <u>1° endpoints:</u> All-cause mortality and a composite of all- cause mortality and/or first readmission for CV reason within 180 d after discharge <u>Results:</u> NT-proBNP levels at discharge and the changes in NT-proBNP during hospitalization yielded the best C-statistic (AUC: 0.78; 95% CI: 0.74–0.82). 	 In pts hospitalized for ADHF, the addition of the discharge NT- proBNP values as well as the change in NT-proBNP to known risk markers, generates a relatively simple yet robust discharge risk score that importantly improves the prediction of adverse events

Cohen-Solal et al. 2009 (34) <u>19539144</u>	Aim: Examine whether decreases in BNP levels during the first few d of hospitalization were associated with greater survival in pts with ADHF Study type: Detrecoective analysis	Inclusion criteria: Of 1,327 SURVIVE pts, this analysis included 1,038 who had BNP samples at both BL and d 5 Follow up: 180 d	 <u>1° endpoints:</u> All-cause mortality and a composite of all- cause mortality and/or first readmission for CV reason within 180 d after discharge <u>Results:</u> A pt was classified as a "responder" if the follow-up BNP level was ≥30% lower than BL BNP Short-term 30 d mortality risk reduction was 	• Pts with lowered BNP on treatment for ADHF had reduced mortality risks (31- and 180-d) compared to those with little or no BNP decrease
	of SURVIVE <u>Size:</u> 1,327 pts		67% in d 5 BNP responders compared with nonresponders, whereas long-term (180-d) all-cause mortality risk reduction was 47%	
Logeart et al. 2004 (35) <u>14975475</u>	Aim: To determine the value of BNP predicting post-discharge outcome of pts admitted for ADHF Study type: Prospective cohort Size: 105 pts	Inclusion criteria: Serial BNP measurements were performed from admission to discharge in 2 samples of consecutive pts	1° endpoints: Combined death or first re-admission for HF Results: The predischarge BNP assay had the best discriminative power (AUC for ROC=0.80) and remained the lone significant variable in multivariate analysis (HR: 1.14; 95% CI: 1.02–1.28; p=0.027	 High predischarge BNP assay is a strong, independent marker of death or readmission after decompensated HF, more relevant than common clinical or echocardiographic parameters and more relevant than changes in BNP levels during acute cares Study relatively old and small
O'Brien et al. 2003 (36) <u>12921811</u>	Aim: To determine the value of BNP predicting post-discharge outcome of pts admitted for ADHF Study type: Prospective cohort Size: 96 pts	Inclusion criteria: NT-proBNP was measured at admission in 96 pts hospitalized with acute LVF Follow up: 180 d	<u>1° endpoints:</u> Combined death or HF <u>Results:</u> Only pre-discharge plasma NT-proBNP (OR: 15.30; 95% CI: 1.4–168.9], p=0.026) was independently predictive of the composite endpoint. The AUC ROC curve for pre- discharge NT-proBNP was superior to that for admission NT-proBNP for prediction of death or HF (AUC ROC 0.87 cf 0.70), for death (0.79 cf 0.66), LVF hospitalization (0.78 cf 0.70) or HF as an outpatient (0.71	 Plasma NT-proBNP measured pre- discharge provides useful prognostic information following hospitalization with acute LVF. Study relatively old and small
			of 0.61	
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Richards et al. 2001 (37) <u>11401111</u>	Study type: Observational study within a randomized trial	Inclusion criteria: Ischemic CM, EF<45%, chronic stable CHF, NYHA II-III or prior IIIV	<u>1° endpoint:</u> Association of plasma N-BNP and adrenomdeullin with mortality and HF events at 18 mo	NT-proBNP and adrenomedullin levels are independently associated with outcome in pts with heart failure from an ischemic cardiomyopathy
	<u>Size</u> : 297	Exclusion criteria: Current NYHA IV, HR<50 bpm, BP<90 or >160/100, coronary event/procedure last 4 weeks, IDDM, CKD, hepatic/renal disease, sick sinus syndrome, 2 nd or 3 rd degree heart block, treatment with beta-blocker, beta-agonist or verapamil	 <u>Results:</u> Above median proBNP increased risk of mortality (HR: 4.7; Cl 2–10.9) and HF admission (HR: 4.7; Cl: 2–10) Above median adrenomedullin increased risk of mortality (HR 3.9,Cl 1.8-8.7) and HF admission (HR 2.4, Cl 1.3-4.5) Associations persist in multivariable modeling 	
Tang et al. 2003 (38) <u>14662703</u>	Study type: Retrospective, observational Size: 558	Inclusion criteria: Chronic systolic HF >3 mo duration, stable medical therapy, LVEF<50%, NYHA class I-III, followed in outpatient HF clinic at a single center who had BNP obtained at clinic visit Exclusion criteria: Congenital heart disease, cardiac transplant, primary valvular disease, active ischemia requiring urgent revascularization	 <u>1° endpoint:</u> Prevalence, clinical characteristics, and characteristics of a BNP<100 pg/mL in a HF clinic population <u>Results:</u> 21% of symptomatic HF pts had BNP <100 pg/mL Characteristics associated with this phenotype include younger age, female gender, nonischemic etiology, better preserved cardiac and renal function, less have atrial fibrillation 	 A sizeable minority (21%) of ambulatory pts with chronic HF have a BNP <100 pg/mL This phenotype (HF with non- diagnostic BNP) is associated with identifiable clinical characteristics
Januzzi et al. 2008 (39) <u>18243855</u>	Study type: Review paper regarding utility of NT-proBNP testing for diagnosis or exclusion of HF in pts with acute HF Size: N/A	Inclusion criteria: Studies using NT-proBNP assays used commercially Exclusion criteria: N/A	 <u>1° endpoint</u>: N/A <u>Results:</u> NT-proBNP had comparable sensitivity/specificity to BNP for diagnosis of acute HF in dyspneic pts NT-proBNP testing may be superior to 	 NT-proBNP testing can help with the diagnosis and triage of the patients with acute dyspnea."

			clinical assessment in diagnosing HF	
Santaguida et al. 2014 (40) <u>25052418</u>	Study type: Systematic review Size: 7 publications included	Inclusion criteria: Study assessing incremental value of BNP or NT-proBNP for predicting morbidity and mortality in acute decompensated HF Exclusion criteria: Studies of stable HF; natriuretic peptide could not be included in base model to allow assessment of incremental value	 <u>1° endpoint</u>: BNP or NT-proBNP improved prognostic model performance for mortality as assessed by discrimination and or likelihood statistics <u>Results:</u> 5 BNP publications consistently predicted all-cause mortality in short (3–6 mo) and long (9,12 mo) beyond base model but not all statistically significant Two NT-proBNP publications both showed incremental value at 22 mo and 6.8 y with 1 being statistically significant 	Clinical heterogeneity precluded formal meta-analysis
Hill et al. 2014 (41) <u>24957908</u>	Study type: Systematic review Size: 76 publications included (37 BNP alone, 25 NT- proBNP alone, 14 both)	 Inclusion criteria: Age >18 y presenting to ED or urgent care center with signs/symptoms suggestive acute HF English language articles from 1989-2012 FDA-approved assays Exclusion criteria: Studies with pts who had conditions that may impact NP levels (transplant, HCM, valvular) 	1° endpoint: Test performance characteristics Results: •BNP pooled sensitivity=95%, 95% CI: 93– 97%), specificity 67% (58–75%) • NT-proBNP pooled sensitivity 91% (95% CI: 88–93), specificity 67% (50–80%)	 Both BNP and NT-proBNP had high sensitivity but low specificity Overall strength of evidence for sensitivity and all decision cutpoints for both peptides was high; strength of evidence for specificity rated as moderate. Both BNP and NT-proBNP performed well to rule out, but less well to rule in, for the diagnosis of heart failure among patients presenting to the ED or urgent care centers.
Zaphiriou et al. 2005 (42) <u>15921792</u>	Study type: Diagnostic accuracy study (observational) Size: 306 pts	Inclusion criteria: Pts with new symptoms suggestive of HF referred by GP to rapid access HF clinics at 5 centers in UK between 201 and 2003	 <u>1° endpoint:</u> Sensitivity, specificity, PPV, NPV, LR, AUC for diagnosis of HF <u>Results:</u> 104 (34%) of pts had HF 	 2 of 5 sites withdrew after recruiting 18 and 14 pts Both BNP and NT-proBNP are useful for ruling out HF in pts presenting to PCP with possible HF symptoms

	Study types	Exclusion criteria: None listed	 AUC BNP 0.84 (95% CI: 0.79–0.89), Nt-proBNP 0.85 (0.81–0.9) BNP: NPV: 0.87, PPV: 0.59 NT-proBNP NPV: 0.97, PPV: 0.44 	
Son et al. 2012 (43) <u>22564550</u>	Study type: Observational, decision making model using rough set and decision tree approaches Size: 159 subjects (71 HF, 88 control)	 Inclusion criteria: ED presentation for dyspnea (HF vs. Noncardiac control) Complete medical records Exclusion criteria: HF excluded if other diagnosis made 	 <u>Production</u> <u>Results:</u> NT-proBNP was one of 6 variables identified in decision-tree rough set and one of 4 variables in logistic regression model 	N I -proBNP identified as a critical variable for decision making of HF in pts with dyspnea presenting to ED
Kelder et al. 2011 (44) <u>22104551</u>	Study type: Cross-sectional, diagnostic accuracy (observational) Size: 721 subjects	Inclusion criteria: Pts presenting with signs/symptoms of HF who were referred to 1 of 8 rapid access clinics in the Netherlands Exclusion criteria: Known, established HF Acute HF requiring immediate therapeutic intervention	<u>1° endpoint:</u> Diagnosis of HF <u>Results:</u> • 207/721 (29%) had HF • C-statistic without proBNP =0.83 • C-statistic with proBNP =0.86 NRI 69%	• NT-proBNP had utility beyond the history and physical for diagnosing HF among primary care outpatients presenting with signs/symptoms of HF
Booth et al. 2014 (45) <u>24969534</u>	Study type: Systematic review Size: 12 BNP publications; 20 NT-proBNP publications	 Inclusion criteria: Pts presenting with signs or symptoms of HF or were at risk of HF a time of presentation Primary care setting Exclusion criteria: Studies with subjects with: Age <18 y Acute HF Known exacerbation of chronic stable HF 	 <u>1° endpoint:</u> Diagnostic accuracy of BNP or NT-proBNP <u>Results:</u> BNP pooled sensitivity (lowest cutpoint 0.85, optimal 0.8, manufacturer 0.74) and specificity (0.54, 0.5, 0.58, respectively) NT-proBNP pooled sensitivity (lowest cutpoint 0.90, optimal 0.86, manufacturer 0.82) and specificity (0.5, 0.58, 0.58, respectively) 	 Both BNP and NT-proBNP have good diagnostic utility for diagnosing HF in the primary care setting in those with signs/symptoms of HF or at risk of developing HF Tests have better sensitivity than specificity Authors felt that it was unlikely that further studies will change these conclusions

		Conditions that may interfere with NP levels (heart transplant, obesity, HCM, valvular lesion)		
Dao et al. 2001 (46) <u>11216950</u>	Study type: Observational, convenience sample at 1 VA urgent care center Size: 250	 Inclusion criteria: SOB as prominent complaint Exclusion criteria: Dyspnea clearly not from HF ACS (unless predominant presentation was HF 	 <u>1° endpoint:</u> Diagnostic utility of point-of-care BNP for diagnosis of HF <u>Results:</u> BNP C-statistic =0.98 Treating physician C statistic =0.88 BNP remained independently associated with HF diagnosis in multivariable model beyond H+P, xray, ECG 	BNP had diagnostic utility for HF diagnosis in the urgent care setting
Davis et al. 1994 (47) <u>7905953</u>	Aim: Assessed value of ANP and BNP in pts presenting with dyspnea Study type: Observational Size: 52	Inclusion criteria: Suspected HF among elderly pts presenting with acute dyspnea requiring admission Exclusion criteria: Pneumonia, pulmonary thromboembolism, or pneumothorax	1° endpoint: Strong negative correlations between LVEF and log BNP (r=-0.7; p<0.001) and log ANP (r=-0.59; p<0.001).	 One of the original studies that showed that plasma BNP was raised in dyspneic pts with HF But not in acutely breathless pts with lung disease Rapid BNP assays may assist in the diagnosis of pts with acute dyspnea
Cheng et al. 2001 (48) <u>11216951</u>	Aim: To determine if BNP levels predict outcomes of pts admitted with decompensated HF Study type: Observational Size: 72	Inclusion criteria: Pts admitted with decompensated NYHA class III to IV HF, measuring daily BNP levels Exclusion criteria: Lack of levels	 <u>1° endpoint:</u> Association between initial BNP and the predischarge or premoribund BNP measurement and subsequent death and 30-d readmission <u>Results:</u> In pts surviving hospitalization, BNP discharge concentrations were strong predictors of subsequent readmission (area under the receiver operator curve of 0.73). 	 In pts admitted with decompensated HF, changes in BNP levels during treatment are strong predictors for mortality and early readmission. BNP levels might be used successfully to guide treatment of pts admitted for decompensated HF
Fonarow et al. 2008 (49) <u>18178412</u>	Aim: To determine additive prognostic value of	Inclusion criteria: Hospitalizations for HF from April 2003 to December	1° endpoint: BNP above the median and increased Tn were associated with significantly increased	 Admission BNP and cardiac Tn levels are significant, independent predictors of in-hospital mortality in

	admission BNP and Tn levels in acutely decompensated HF <u>Study type</u> : Registry analysis	2004 entered into ADHERE were analyzed. BNP assessment on admission was performed in 48,629 (63%) of 77,467 hospitalization episodes	risk of in-hospital mortality (OR: 2.09 and 2.41 respectively, each p<0.0001).	acutely decompensated HF.
	<u>Size</u> : 48,629	Exclusion criteria: Absence of BNP levels		
Zairis et al. 2010 (50) <u>19157603</u>	Aim: To investigate the combined prognostic value of admission serum levels of BNP, cTnI and hs-CRP, in pts hospitalized because of acutely decompensated severe (NYHA class III/IV) low-output chronic HF. Study type: Multicenter Prospective cohort Size:	Inclusion criteria: Consecutive hospitalized acute decompensated HF pts with NYHA class III/IV recruited in the 5 participating centers Exclusion criteria: Competing diagnoses of renal failure, MI	<u>1° endpoint</u> : Cardiac mortality by 31 d <u>Results:</u> There was a significant gradual increased risk of 31-d cardiac death with increasing in the number of elevated biomarkers (p<0.001). By multivariate Cox regression analysis, elevated serum levels of BNP (p=0.002), cTnl (p<0.001) and hs-CRP (p=0.02) were independent predictors of the study end point.	 In pts hospitalized for acute decompensation of severe (NYHA III/IV) low-output HF, BNP, cTnI and hs-CRP upon admission offers enhanced early risk stratification.
Peacock et al. 2008	577 Aim:	Inclusion criteria:	1º endpoint:	 In pts with acute decompensated
(51) <u>18480204</u>	Describe the association between elevated cardiac troponin levels and adverse events in hospitalized pts with acute decompensated HF	Hospitalizations for acute decompensated HF between 2001 and 2004 in ADHERE. Entry criteria included a troponin level that was obtained at the time of hospitalization	Overall, 4,240 pts (6.2%) were positive for troponin. Results: Pts who were positive for troponin had lower SBP on admission, a lower EF, and higher in-hospital mortality (8.0% vs. 2.7%, p<0.001) than those who were negative for	HF, a positive cardiac troponin test is associated with higher in-hospital mortality, independently of other predictive variables.
	Study type: Registry analysis	Exclusion criteria: Pts with a serum creatinine level ≥ 2.0 mg per deciliter	troponin. The adjusted odds ratio for death in the group of pts with a positive troponin test was 2.55 (95% CI: 2.24–2.89; p<0.001)	

	Sine: 67.004	(177 micromol por liter)		
	<u>512e</u> : 07,924			
Lee et al. 2012 (52) <u>22665814</u>	Aim: To derive and validate a model for acute HF mortality applicable in the ED. Study type: Multicenter Registry analysis Size: 12,591	Inclusion criteria: Population-based random sample of 12,591 pts presenting to the ED from 2004 to 2007 Exclusion criteria: No lab availability	1° endpoint: Death within 7 d of presentation Results: Mortality risk increased with higher triage heart rate (OR: 1.15; [95% CI: 1.03–1.30] per 10 beats/min) and creatinine concentration (OR: 1.35; [CI: 1.14–1.60] per 1 mg/dL [88.4 micro mol/L]), and lower triage SBP (OR: 1.52 [CI: 1.31–1.77] per 20 mm Hg) and initial oxygen saturation (OR, 1 16 [CI: 1.01–1.33] per 5%)	• A multivariate index comprising routinely collected variables stratified mortality risk with high discrimination in a broad group of pts with acute HF presenting to the ED.
Dhaliwal et al. 2009 (53) <u>19398076</u>	Aim: Compare the relationship between absolute and relative changes in BNP with future clinical events, and whether serial BNP measurements add prognostic information in pts treated for decompensated HF Study type: Retrospective registry analysis	Inclusion criteria: Pts hospitalized for acute decompesated HF by Framingham criteria Exclusion criteria: Renal failure, severe lung disease, acute coronary syndrome	 1.10 [ci. 1.01–1.35] per 5 %). <u>1° endpoint:</u> For the combined end point of total mortality or readmission for HF <u>Results:</u> Increasing tertiles of BNP levels after treatment had a hazard ratio of 1.4 (1.1–1.7, p<0.01) and increasing tertiles of percent reduction in BNP, had a HR:0.7 (0.6–0.9; p=0.005), respectively, for the combined end point of total mortality or readmission for HF Follow-up BNP performed better than did baseline BNP or percent reduction in BNP. More BNP measurements other than the follow-up BNP did not improve the fit of the model further. 	 Both lower absolute BNP levels and greater percentage reduction in BNP with treatment of decompensated HF are associated with better event-free survival. Advocating a threshold BNP to which pts should be treated may not be possible given that high BNP levels tend not to decrease to levels associated with better outcomes during the short period of treatment. More BNP measurements do not add prognostic information beyond that provided by a single BNP level after treatment.
Alonso-Martinez et al. 2002 (54) <u>12034159</u>	Aim: To determine usefulness of CRP in predicting need for readmission in HF	Inclusion criteria: Intervention group: admission with HF; control group: admission with syncope	 <u>1° endpoint:</u> 18-mo HF readmission CRP levels were higher in pts with HF compared to syncope (3.94 vs. 0.84, p<0.0007) 	 Multivariate predictors of readmission were CRP levels, NYHA class and plasma K on discharge Limitation: small, single-center

	Study type: Observational Size: 76	Exclusion criteria: Clear cause for elevated CRP (e.g., inflammation, infection)	Higher CRP levels were associated with higher NYHA class, increased risk of HF readmission, shorter time to readmission, and increased mortality <u>Safety endpoint:</u> NYHA class on discharge and death	observational study
Dieplinger et al. 2010 (55) <u>20153308</u>	Aim: To evaluate the prognostic value of established and novel biomarkers in pts with acute dyspnea Study type: Observational Size: 251	Inclusion criteria: Pts presenting to ED with acute dyspnea Exclusion criteria: STEMI, NSTEMI or ACS troponin pos. Biomarkers: BNP, MR-proANP, MR- proADM, copeptin, C- terminal pro-ET-1, soluble ST2, chromogranin A (CgA), adiponectin, proguanylin, prouroguanylin	 <u>1° endpoint:</u> All-cause mortality at 1 y 25% died within 1 y At baseline, decedents (n=62) had higher median plasma concentrations of all 10 biomarkers than survivors (n=189) In multivariate model, only MR-proANP (RR: 1.6), ST2 (RR: 1.7) and CgA (RR: 1.5) were independent predictors of death 	 Low systolic BP and advanced age were also independent predictors of 1-y mortality Limitations: post-hoc analysis; sub- group (87 of 251) had dyspnea due to acute HF alone; single-center, majority men (94%)
Ilva et al. 2008 (56) <u>18599345</u>	Aim: To evaluate prevalence and prognostic significance of elevated cTnl and cTnT in acute HF <u>Study type:</u> Observational substudy <u>Size</u> : 364	Inclusion criteria: Hospitalized with acute HF Exclusion criteria: ACS pts; missing sample for cardiac TnI/TnT Biomarkers on admission and 48 hours: cTnT, cTnI, cystatin C, NT- proBNP	 <u>1° endpoint</u>: 6 -mo mortality 51% of pts had +cTnI and 30% had +cTnT 6-mo all-cause mortality was 18.7% Both cTnI (OR: 2.0; 95% CI: 1.2–3.5) and cTnT (OR: 2.6; 95% CI: 1.5–4.4) were associated with adverse outcome in pts with previous, but no de novo HF 	 On multivariable analysis, cystatin C (OR: 6.3; 95% CI: 3.2–13), logNT-proBNP (OR: 1.4; 95% CI: 1.0–1.8) and SBP on admission (/10 mm Hg increase; OR: 0.9; 95% CI: 0.8–0.9) were independent risk predictors, whereas troponins were not Mortality was proportional to troponin release Limitations: exclusion of pts with ACS was based on clinician judgment; cut-off values for troponins was based on 2000 ESC/ACC guidelines
Januzzi et al. 2007 (57)	Aim: To examine the value of	Inclusion criteria: Pts presenting to ED with	<u>1° endpoint:</u> •death at 1 y	ST2 levels were higher in pts with HF/EF (0.67 ng/ml; IQR 0.31–1.50)
<u>17692745</u>	measuring ST2 in pts	acute dyspnea	• ST2 levels were significantly higher in pts	vs. HFpEF (0.42 ng/ml; IQR 0.22-

	with acute dyspnea Study type: Observational Size: 593 (pts with acute HF 209, other causes of acute dyspnea 384)	Exclusion criteria: Not reported	 with acute HF (0.50 ng/ml; IQR 0.27–1.22) vs. those without (0.15 ng/ml; IQR 0.06– 0.42) 1-y mortality was 15.7% ST2 levels were significantly higher in decedents than survivors (1.03 vs. 0.18 ng/ml; p<0.001) In multivariable analysis, ST2 ≥0.20 ng/ml strongly predicted death at 1 y 	 0.90) A multi-marker approach with both ST2 and NT-proBNP levels identified subjects with the highest risk for death Limitations: single-center study; biologic role of ST2 in acute HF poorly understood
Manzano-Fernandez et al. 2011 (58) <u>21211603</u>	Aim: To determine whether risk of mortality associated with ST2 differs in pts with acute HF,pEF vs. HF/EF Study type: Observational study combining 3 databases (Boston, MA; Linz, Austria; Murcia, Spain) Size: 447	Inclusion criteria: Acute HF Exclusion criteria: N/A Biomarkers: ST2, troponin T, NT-proBNP, CRP	 <u>1° endpoint</u>: 1 y vital status During 1-y follow-up, 117 pts (26%) died ST2 levels were higher among deceased than survivors (median 0.80 ng/ml vs.0.38 ng/ml; p<0.001); and this pattern was true for HF<i>r</i>EF and HF<i>p</i>EF On multivariate analysis, elevated ST2 levels were associated with greater risk of 1-y mortality for HF<i>p</i>EF (HR: 1.41; 95% CI: 1.14–1.76) than HF<i>r</i>EF (HR: 1.20; 95% CI: 1.10–1.32) 	 Pts with HF/EF had higher ST2 levels than HFpEF (median 0.55 ng/ml vs. 0.38 ng/ml; p<0.001) Addition of ST2 to NT-proBNP improved C statistic and both net reclassification improvement and integrated discrimination improvement, regardless of LVEF Limitations: pooled multinational analysis that lacked predefined endpoints and complete echocardiographic measures; no pre-discharge ST2 levels
Rehman et al. 2008 (59) <u>19017513</u>	Aim: To examine patient- specific characteristic of ST2 in pts with acute HF Study type: Observational study combining 2 databases (Boston, MA; Linz, Austria) Size: 346	Inclusion criteria: Acute HF Exclusion criteria: N/A Biomarkers: ST2, BNP, NT-proBNP, CRP	 <u>1° endpoint</u>: •ROC curves and multivariable Cox proportional hazards analyses • ST2 levels correlated with severity of HF (p<0.001), LVEF and creatinine clearance • ST2 levels correlated with BNP, NT-proBNP and CRP • In a multivariable model, ST2 remained a predictor of mortality (HR: 2.04; 95% CI: 1.30–3.24) 	 Pts with HFpEF had lower ST2 levels compared to HF/EF 1-y mortality was 42% among 116 pts with elevation in both ST2 and BNP/NT-proBNP In the presence of a low ST2 level, BNP/NT-proBNP did not predict mortality Limitations: lack of serial measures of ST2; biologic role of ST2 in acute HF poorly understood

Shah et al. 2010 (60) <u>20525986</u>	Aim: To determine the relationship between galectin-3 and cardiac structure and function in pts with acute dyspnea Study type: Observational Size: 115	Inclusion criteria: PT presenting to ED with acute dyspnea, detailed echo exams during admission Exclusion criteria: N/A Biomarkers: galectin-3, NT-proBNP	 <u>1° endpoint:</u> Association between galectin-3 and echo and clinical indices Higher levels of galectin-3 associated with older age, poorer renal function, and higher NT-proBNP Significant relationship between galectin-3 and poorer RV function, higher RV systolic pressure and more severe MR and TR 	 Galectin-3 levels higher in pts who died at 1 and 4 y In multivariate analysis, galectin-3 remained a significant predictor of 4-y mortality independent to echocardiographic markers of risk Limitations: delay between collection of biomarkers and echocardiograms; small, single- center cohort
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Search Terms and Date: natriuretic peptides, heart failure, human, last 5 years. Last search done on April 18, 2016.

Data Supplement 1. RCTs Comparing ARNI (Section 7.3.2.10)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% Cl)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
PARAMOUNT Solomon et al. 2012 (61) 22932717	Aim: To address safety and efficacy of LCZ696 (ARNI) in pts with HF <i>p</i> EF Study type: RCT Size: 308	Inclusion criteria: Pts ≥40 y of age, LVEF ≥45%, NYHA class II-III HF, NT-pro BNP >400 pg/mL. Exclusion criteria: Right HF due to pulmonary disease, dyspnea due to noncardiac causes, valvular/myocardial disease, CAD or CVD needing revascularization within 3 mo of screening.	Intervention: LCZ696 (149) target dose 200 mg BID achieved in 81% Comparator: Valsartan (152) target dose 160 mg BID achieved in 78%	 <u>1° endpoint</u>: Change from BL at 12 wk for NT-proBNP Results: Reduction in LCZ696 group vs. valsartan (ratio of change from BL: 0.77, 95% CI: 0.64–0.92; p=0.005) <u>1° Safety endpoint</u>: LCZ-696 well tolerated. Serious adverse events: 15% in LCZ696 vs. 20% in valsartan group 	 No difference in change in NT-proBNP from BL at 36 wk BP reduced in the LCZ696 group vs. valsartan at 12 wk (p=0.001 for SBP and p=0.09 for DBP) Change in BP correlated poorly with the change in pro-BNP No difference in improvement in NYHA class at 12 wk (p=0.11) and 36 wk (p=0.05). No difference in KCCQ scores Trial not powered to ascertain clinical outcomes. Further studies needed to assess safety and efficacy in HF<i>p</i>EF pts.
PARADIGM-HF McMurray et al. 2014	Aim: To compare survival rates with the use of	Inclusion criteria: ≥18 y of age, NYHA class II, III, IV; EF ≤35%, BNP of at least 150	Intervention: LCZ696 (4,187) target dose 200 mg BID (mean	 <u>1° endpoint:</u> Composite of death (CV causes) or a first 	 Less CV death in LCZ696 arm (558 vs. 693) HR: 0.8 (95% CI: 0.71–0.89; p<0.001)

(62)	LCZ696 with	pg/mL, hospitalized for HF <12 mo	375 <u>+</u> 71 mg daily)	hospitalization for HF	• Less HF hospitalizations in LCZ696 arm
<u>25176015</u>	enalapril in HF	(≥BNP100 pg/mL), on ACE			(537 vs. 658) HR: 0.79 (95% CI: 0.71–
	O f i i h h h h h h h h h h	inhibitors or ARBs ≥4 wk before	Comparator:	Results: Composite less in	0.89; p<0.001)
	Study type:	screening, required to take stable	Enalapril (4,212) target 10	LCZ696 group vs.	Less death from any cause in LCZ696
	KUI	inhibitor (or ARB) equal to 10mg of	mg daily)	enalapril, 914 (21.8%) vs. 1,117, (26.5%) HR: 0.80	arm (711 vs. 835), HR: 0.84 (95% CI: 0.76–0.93; p<0.001)
	<u>Size</u> : 8,442	enalapril. Prior to randomization pts were required to complete 2 wk each of enalapril 10 mg BID and LCZ 100 BID. Exclusion criteria: Symptomatic hypotension, SBP <95 mm Hg, eGFR <30 mL/min/min/1.73m ² of body surface area, serum K level >5.2 mmol/L, angioedema history, unacceptable		(95% Cl: 0.73–0.87; p<0.001)	 The change from baseline to 8 mo in the score on the KCCQ in LCZ696 arm (2.99 points reduction vs. 4.63 points), HR: 1.64 (95% CI: 0.63–2.65; p=0.001) No difference in new onset of AF (84 vs. 83; p=0.84) No difference in protocol defined decline in renal function, HR: 0.86 (95% CI: 0.65–1.13; p=0.28). More symptomatic hypotension (14% vs. 9.2%; p<0.001)
		side effects of ACE inhibitors or ARBs			 No difference in angioedema, 19 vs.10 (p=0.13)

Search Terms and Date: 3 trials identified by chairs in December 2015.

			1		
Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% Cl)	Relevant 2º Endpoint (if any); Study Limitations; Adverse Events
ONTARGET ONTARGET Investigators et al. 2008 (63) <u>18378520</u>	Aim: Compare ACE (ramipril), ARB (telmisartan), and combination ACE/ARB in pts with CVD or high- risk DM Study Type: RCT Size: 25,620	Inclusion Criteria: Pts >55 y of age, CAD, PVD, previous stroke, or high-risk DM with end-organ damage Exclusion Criteria: HF at trial entry, ACE or ARB intolerance, revascularization planned or <3 mo	Intervention: Run in, then randomization to ramipril (8,576) target dose 10 mg daily, telmisartan (8,542) target dose 80 mg daily or combination (8,502), titrated to BP	 <u>1° endpoint:</u> Composite of CV death, MI, stroke, or HF hospitalization at 5 y Results: No difference in outcome (16.5% ACE, 16.7% ARB, 16.3% combination; CI: ARB RR: 1.01 (95% CI: 0.94–1.09) 	 Compared to the ramipril arm: Telmisartan had more hypotensive symptoms (p<0.001); less cough (p<0.001) and angioedema (p=0.01); same syncope. Combination arm had more hypotensive symptoms (p<0.001); syncope (p=0.03); and renal dysfunction (p<0.001) BP fell by 6.4/7.4/9.8 mm Hg Less angioedema with telmisartan
TRANSCEND Yusuf et al. 2008 (64) <u>18757085</u>	Aim: To assess the effectiveness of ARB in ACE- intolerant pts with CVD or high-risk DM Study Type: RCT Size: 5,926	Inclusion Criteria: ACE-intolerant pts with CAD, PVD, previous stroke, or high-risk DM with end-organ damage Exclusion Criteria: HF at trial entry, revascularization planned or <3 mo	Intervention: Run in, then randomization to telmisartan titrated to 80 mg as tolerated (2,954) Comparator: Titration of other mediations as needed to control BP (2,944)	 <u>1° endpoint</u>: Composite of CV death, MI, stroke, or HF hospitalization at 5 y <u>Results</u>: No significant difference RR: 0.92 (95% CI: 0.81–1.05); p=0.216 	 No difference in 2° outcomes; ARB was safe in this pt population - no angioedema
SUPPORT Sakata et al. 2015 (65) 25637937	<u>Aim:</u> Discover whether addition of ARB to ACE and beta blockers in pts with chronic HF will	Inclusion Criteria: Pts 20–79 y of age with hypertension, NYHA class II-IV, stable on ACE ± beta blockers	Intervention: Randomization to olmesartan (578) titrated up to 40 mg as tolerated (578) (mean dose achieved at 5 y, 17.9	 <u>1° endpoint</u>: Composite of all-cause death, MI, stroke, or HF hospitalization at 4.4 y <u>Results</u>: No significant difference RR: 1.18 (95% CI: 0.96–1.46); p=0.11 	Pts on triple therapy with ACE/ARB/Beta blocker had more of 1° composite outcome, 38.1 vs. 28.2%, HR: 1.47 (95% CI: 1.11– 1.95; p=0.006); all-cause death, 19.4 vs. 13.5%, HR: 1.50 (95% CI:

Data Supplement 2. RCTs Comparing RAAS Inhibition (Section 7.3.2.3)

	improve clinical outcomes Study Type: Open label blinded endpoint Size: 1,147	Exclusion Criteria: Creatinine >3.0, MI or, revascularization within 6 mo	mg/d) <u>Comparator:</u> Titration to control BP without use of an ARB (568)		1.01–2.23; p=0.046); and renal dysfunction (21.1 vs. 12.5%, HR: 1.85 (95% CI: 1.24–2.76; p=0.003).
Mineralocorticoids An	tagonist Trials				
EMPHASIS subgroup analysis Eschalier et al. 2013 (66) 23810881	Aim: Investigate the safety and efficacy of eplerenone in pts at high risk for hyperkalemia Study Type: Prespecified subgroup analysis of RCT Size: 2,737	Inclusion Criteria: Pts enrolled in EMPHASIS at high risk for hyperkalemia of worsening renal function (>75 y, DM, eGFR <60, or SBP <123) Exclusion Criteria: eGFR<30	Intervention: Randomization to eplerenone Comparator: Placebo	 <u>1° endpoint</u>: Efficacy: Hospitalization for HF or worsening renal failure. Safety: K >5.5, >6.0, <3.5, hospitalization for significant hyperkalemia, hospitalization for worsening renal function <u>Results:</u> Efficacy: reduced composite endpoint. Safety: increased risk of K+ >5.5 mmol/L, hospitalization for hyperkalemia or discontinuation of study medication due to adverse events. No differences from the main trial results in the high-risk subgroups. K >5.5 was increased in the whole cohort and the subgroups, but K >6.0, clinically significant hyperkalemia, and change in eGFR were not substantially higher. 	The beneficial effects of eplerenone were maintained in the high-risk subgroups.
RALES Pitt et al. 1999 (67) <u>10471456</u>	Aim: To investigate the effect of spironolactone on mortality and morbidity in pts with severe HF. Study Type:	Inclusion Criteria: NYHA class III, IV; HF≤6 mo, Left EF≤35%, On ACE inhibitors, loop diuretic. Digitalis and vasodilators allowed. Exclusion Criteria: 1° operable VHD (other than	Intervention: Spironolactone 25 mg daily (822) Comparator: Placebo (841)	 <u>1° endpoint:</u> Death from all causes <u>Results:</u> Placebo vs. Spironolactone group (46% vs. 35%; RR: 0.70; 95% Cl: 0.60–0.82; p<0.001) Trial stopped early due to favorable results at 24 mo. 	 Reduction in death from cardiac causes and Hospitalization for cardiac causes (p<0.001) Improvement in NYHA class (p<0.001) No clinically important safety concerns for electrolytes. Gynecomastia/breast pain more frequent in the spironolactone

RCT	mitral or tricuspid), ACHD,		group (p<0.001)
	unstable angina, 1° heaptic		
<u>Size:</u>	failure, active cancer, life		
1,663	threatening disease, heart		
	transplant, serum Cr ≥2.5		
	mg/dĹ, serum K ≥5.0 mmoL/L		

The ARB evidence table from the 2013 Heart Failure Guideline is included at the end of this document.

The ACE inhibitor evidence table from the 2013 Heart Failure Guideline is also included at the end of this document.

The Beta Blocker evidence table from the 2013 Heart Failure Guideline is included at the end of this document.

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% CI)	Relevant 2° Endpoint; Study Limitations; Adverse Events
IMPRESS Rouleau et al. 2000 (68) <u>10968433</u>	Aim: Determine if inhibition of neutral endopeptidase and ACE with the vasopeptidase inhibitor omapatrilat is better than ACE inhibition alone with lisinopril Study type: Double blind RCT Size: 573 pts	Inclusion criteria: • Informed consent • Age ≥18 • Stable (>3 mo) symptomatic HF (NYHA class II–IV HF) • Decreased LVEF ≤40 • ≥4 wk dose of ACE inhibitors • Seated SBP ≥90 mm Hg Exclusion criteria: • Uncontrolled hypertension • Acute coronary events within 3 mo • Revascularization within 3 mo • Serum potassium <3.5 or >5.3 mmol/L • Creatinine >221 mcmol/L • Transaminases >2 upper limit of normal • Leucocytes <3.0x10 ⁹ /L, neutrophils <1. • Sx10 ⁹ /L, or platelets <120x10 ⁹ /L	Intervention: Omapatrilat (289) target dose 40 mg daily Comparator: Lisinopril (284) target dose 20 mg daily	1° endpoint: Change in exercise duration from baseline to wk 12 Results: Similar exercise duration at 12 wk (p=0.45)	 <u>2° endpoint</u>: No difference in combined endpoint of death and admission for worsening HF (p=0.52) Combined endpoint of death and comorbidity for worsening HF was better for omapatrilat HR: 0.52 (95% CI: 0.28–0.96; p=0.035) Angioedema occurred in no pts taking omapatrilat vs. 1 taking enalapril <u>Comments</u>: Vasopeptidase inhibitor omapatrilat did not improve exercise tolerance compared with ACE inhibitor lisinopril

Data Supplement 3. RCTs Comparing Pharmacological Treatment for of ARNI With ACE (Section 7.3.2.10)

		● Lise of beta blockers <6 mo			
		Calcium channel blockers for use other than			
		• Pts included in previous RCTs of omanatrilat			
OVERTURE	Aim:	Inclusion criteria:	Intervention:	1° endpoint: Combined	 Omapatrilat reduced risk of death
Packer et al. 2002	Determine dual ACE	NYHA class II–IV HF due to non/ischemic	Omapatrilat (2,886),	risk of death or	and hospitalization for chronic HF
(69)	and NEP inhibitors	cardiomyopathy for ≥2 mo, or	target dose 40 mg daily	hospitalization for HF	HR: 0.89 (95% CI: 0.82–0.98;
<u>12186794</u>	provides greater	 LVEF ≤30% and hospitalized for HF within 12 	achieved 82.5%	requiring IV treatment	p=0.012). For this analysis, pts were
	benefit in pts with HF	mo			treated with intensification of oral
	than ACE inhibitors		Comparator: Enalapril	Results: No significant	medications.
	alone	Exclusion criteria:	(2,884) target dose 10	difference HR: 0.94 (95%	
		Surgically correctable or reversible cause of	mg BID achieved 86.4%	CI: 0.86–1.03; p=0.187)	 More frequent angloedema with
	Study type:	HF			omapatrilat (0.8% vs. 0.5%)
	Double blind RCT	 Likely to receive cardiac transplant or left 			
		ventricular assist device			
	<u>Size</u> :	• Severe 1° pulmonary, renal, or hepatic disease			
	5,770 pts	 Hx of intolerance to ACE inhibitors 			
		ACS within 1 mo			
		 Coronary revascularization or an acute 			
		cerebral ischemic event within 3 mo			
		 Hx of ventricular tachycardia, ventricular 			
		fibrillation, or sudden death who did not have an			
		ICD placed and had not fired within 2 mo			
		• Hx or hospitalization or intravenous therapy			
		for HF within 48 h			
		 IV positive inotropic agent within 2 wk 			
		• SBP >180 or <90 mm Hg			
		Heart rate >130 bpm			
		 Serum creatinine >2.5 mg/dL 			
		• Serum potassium <3.5 or >5.2 mmol/L			
OCTAVE	Aim:	Inclusion criteria:	Intervention:	1° endpoints:	2° endpoints:
Kostis et al. 2004	Compare safety and	• Age ≥18	Omapatrilat target dose	Reduction in SBP at wk	 Reduction in DBP at wk 8
(70)	efficacy of dual ACE	• 3 separate BP criteria for 3 groups: Group 1	80 mg daily	8	Reduction in SBP and DBP at wk
<u>14751650</u>	and NEP inhibitors to	untreated hypertension (SBP ≥140 mm Hq or		 Need for new 	24
	ACE inhibitors alone	DBP ≥90 mm Hg); Group 2 hypertension and	Comparator: Enalapril	adjunctive	• BP control (SBP <140 mm Hg and
		persistent mild hypertension (trough SBP 140-	target dose 40 mg daily	antihypertensive therapy	DBP <90 mm Hg) at wk 8 and 24
	Study type:	159 mm Hg and DBP <100 mm Hg, or trough		by wk 24	3,
	Double blind RCT	DBP 90–99 mm Hg and SBP <160 mm Hg);			Comments:

<u>Size</u> : 25,302 pts	Group 3 hypertension with persistent moderate to severe hypertension (trough SBP 160–179 mm Hg and DBP <110 mm Hg, or trough DBP 100–109 mm Hg and SBP <180 mm Hg) Exclusion criteria:	 Greater reductions in BP in omapatrilat within each study (p<0.001) Overall mean reduction in SBP ≥3.6 mm Hg Larger reductions in BP in black
	Contraindication to therapy with ACE inhibitors or angiotensin II receptor antagonists Hy of angioedema, anaphylaxis, drug, induced	pts with omapatrilat than with enalapril. But overall reduction
	or chronic urticarial, or multiple drug sensitivities	subgroups.
	• Recent hospitalization for MI, unstable angina, stroke, TIA or COPD	Adverse events, serious adverse events, and deaths were the same
	 Recent treatment for malignancy, chronic renal disease 2° to autoimmune disease, or end-stage renal disease of any etiology 	 More angioedema with omapatrilat (2.17% vs. 0.68%)
	• Hypertensive pts treated with ACE inhibitors whose BP placed them in study group 3	• More angioedema in blacks with omapatrilat (5.54% vs. 1.62%) and current smokers (3.93% vs. 0.81%)

Search Terms and Date: March 2016, angioedema, neprilysin inhibitors, omapatrilat.

Data Supplement 4. RCTs Comparing Pharmacological Treatment for Stage C HF/EF (Section 7.3.2.11)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% Cl)	Relevant 2º Endpoint (if any); Study Limitations; Adverse Events
SHIFT HF Böhm et al. 2015 (71) <u>26508709</u>	Aim: To assess influence of comorbidities on outcomes and ivabradine treatment effect of heart rate reduction in stable HF. Study type: Post hoc analysis of RCT	Inclusion criteria: Pts ≥18 y of age in sinus rhythm, heart rate at rest ≥70 bpm, MTD for HF meds Exclusion criteria: N/A	Intervention: Ivabradine Comparator: Placebo	 <u>1° endpoint</u>: CV death or HF hospitalization rate increased with the comorbidity load (p<0.0001) with most events in pts with >3 comorbidities for both drug and placebo. Hospitalization rate lower for comorbidity loads of ivabradine 	 Number of comorbidities was related to outcomes Heart rate reduction with Ivabradine is conserved at all comorbidity loads

	<u>Size</u> : 6,505				
SHIFT Swedberg K et al. 2010 (72) 20801500 Ivabradine and outcomes in chronic HF (SHIFT)	Aim: To assess the effect of heart rate reduction by the selective sinus- node inhibitor ivabradine on outcomes in HF <u>Study type:</u> randomized, double-blind placebo-controlled trial. 677 centers 37 countries <u>Size:</u> 6,558 6,505 analyzed 3,241 ivabradine 3,264 placebo	Inclusion criteria: O ver 18 y of age, in sinus rhythm, resting heart rate of ≥70 bpm, stable symptomatic chronic HF (NYHA class II-IV) for ≥4 wk, previous admission to the hospital for HF within 12 mo, LVEF ≤35% Exclusion criteria: HF due to congenital heart disease or 1° severe valvular disease. MI within 2 mo, ventricular or AV pacing for ≥40% of the d, AF or flutter, symptomatic hypotension The following treatments not allowed during study: • diltiazem and verapamil (nondihydropyridine CCB) • class I antiarrhythmics • strong inhibitors of CYP450 3A4	Intervention: Ivabradine Comparator: Placebo	 <u>1° endpoint</u>: Composite of CV death or hospital admission for worsening HF Primary endpoint: ivabradine better. Event rate 24% vs. 29%. HR 0.82 (0.75–0.90); p<0.0001 Hospitalization for worsening HF: ivabradine better. 16% vs 21%, HR: 0.74 (95% CI: 0.66–0.83; p<0.001) Death from HF: ivabradine better. 3% vs. 5%; HF: 0.74 (0.58–0.94); p=0.014 	 Composite of CV death or hospital admission for worsening HF among those receiving at least 50% of target beta blocker dose at time of randomization. All cause death; any CV death; HF hospitalization; death from HF; composite of CV death HF hospitalization, death from HF; composite of CV death HF hospitalization, nonfatal MI. No difference in all-cause mortality or CV mortality Ivabradine better for all-cause hospitalization, HF hospitalization, CV hospitalization, and composite 2° endpoint Analyzed as time to first event. Median follow-up of 22.9 mo In subgroup analysis, effect limited to those with higher baseline heart rate (≥77 bpm) Use of devices was low (CRT in 1% and ICD in 4%) Mean age 61 y When added to GDEM, including beta blocker at optimal dose, ivabradine reduced adverse events, driven largely by HF mortality or HF hospitalization Adverse Effects: 1% withdrew due to bradycardia (p<0.001) Phosphenes 3% (p<0.001)
Fox et al. 2014 (73)	Assess the mortality-morbidity	Stable CAD without clinical HF and heart rate of ≥70	Ivabradine (n=9,550)	Composite of CV death and nonfatal MI	 Adverse Events: Increased bradycardia, AF, phosphenes and cardiac disorders.

<u>25176136</u>	benefits of Ivabradine in pts with stable CAD without clinical HF <u>Study type</u> : RCT <u>Size</u> : 19,102	bpm and in sinus rhythm, persistence and confirmation of ≥1 CV risk factors <u>Exclusion criteria</u> : Serum creatinine >200 mcmol /L, significant anemia, ALT or AST >3 times upper normal value, unstable CV condition, LVEF ≤40%; MI, coronary revascularization, stroke ≤3 mo.	Comparator: Placebo (n=9,552)	 Results: No significant difference in incidence of 1° endpoint (HR: 1.08; 95% CI: 0.96–1.20; p=0.20), death from CV causes (HR: 1.10; 95% CI: 0.94–1.28; p=0.25), nonfatal MI (HR: 1.04; 95% CI: 0.90–1.21; p=0.60) and rate of death (HR: 1.06; 95% CI: 0.94–1.21; p=0.35) <u>1° Safety endpoint:</u> Incidence of bradycardia higher in Ivabradine group (p=0.001) 	 Significant interaction between ivabradine and presence of angina in a subgroup analysis (p=0.02).
BEAUTIFUL Fox et al. 2008 (74) <u>18757088</u>	Aim: Assess the mortality-morbidity benefits of Ivabradine in pts with CAD and LV systolic dysfunction Study type: Randomized, double-blind, placebo-controlled Size: 10,917 5,479 ivabradine 5438 placebo	 Inclusion criteria: Pts ≥55 y of age with stable CAD defined as: previous MI, previous revascularization (PCI or surgery), or angiographic evidence of ≥1 stenosis of ≤50%) AND LVEF <40% and end diastolic internal dimension of >56 mm. Sinus rhythm with resting heart rate of ≥60 bpm. Angina and HF symptoms stable for 3 mo Appropriate conventional CV medication for 1 mo. Exclusion criteria: MI or coronary revascularization within the previous 6 mo; stroke or TIA within 3 mo, PPM or ICD, valvular disease likely to 	Intervention: Ivabradine n=5,479 Comparator: • Placebo in addition to appropriate CV medication n=5,438	 <u>1° endpoint</u>: Composite of CV death, admission for MI and admission for HF No difference in composite 1° endpoint (22.5% vs. 22.8%; HR: 1.00; 0.91–1.1; p=0.94) No differences in any prespecified subgroup. 	 <u>2° endpoints</u>: All-cause mortality Cardiac death (death from MI or HF or related to a cardiac procedure) CV death (death from a vascular procedure, presumed arrhythmic death, stroke death, other vascular death or sudden death of unknown cause) or admission for HF, Composite of admission for fatal and nonfatal MI or UA Coronary revascularization CV death CV death No differences in 2° endpoints in overall population. In <u>subgroup with heart rate of ≥70</u>, ivabradine reduced admission for AMI (fatal and nonfatal) (HR 0.64; 0.49–0.84; p=0.001) composite of admission for AMI or UA (HR 0.78; 0.62–0.97; p=0.023)

need surgery within 3 y,	 coronary revascularization (HR 0.7; 0.52–0.93;
SSS, sinoatrial block,	p=0.16)
congenital long QT,	
complete AV block, severe	 28% in Ivabradine group discontinued medication
or uncontrolled	(vs. 16%), largely due to bradycardia (13% vs. 2%)
hypertension, NYHA class	(,
IV HF	No significant difference in adverse effects (23% vs.
	220/(0.70)
	[23%; p=0.70]

Search Terms and Date: studies identified by chairs in December 2015, one study added by Jan 2016.

Data Supplement C. RCTs Comparing Pharmacologic Treatment for HF*p*EF: Recommendations (Section 7.3.3)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% Cl)	Relevant 2º Endpoint (if any); Study Limitations; Adverse Events
HYVET Beckett et al. 2008 (75) <u>18378519</u>	Aim: To determine whether treatment of HTN is beneficial in the elderly. Study type: RCT Size: 3,845	Inclusion criteria: Age >80, persistent HTN (SBP >160) Exclusion criteria: Known HF, creatinine >150 µmol/L (1.7 mg/dL), CVA <6 mo	Intervention: Indapamide + perindopril if needed for BP control. Target 150/80 mm Hg (1,933) Comparator: Placebo (1,912)	 <u>1° endpoint</u>: Fatal or nonfatal stroke. Trend for improved outcome with active treatment 51 strokes (12.4/1,000 pt-y) vs. placebo 69 (17.7/1,000 pt-y), HR: 0.70; 95% Cl: 0.49–1.01; p=0.06) and significantly reduced fatal stroke 27 (6.5/1000 pt-y) vs. placebo 42 (10.7/1000 pt-y), HR: 0.61; 95% Cl: 0.38–0.99; p=0.046) 	 Significantly reduced all-cause death HR: 0.79 (95% CI: 0.65–0.95; p=0.02) and HF incidence HR: 0.36 (95% CI: 0.22–0.58, p<0.001) with active treatment Trend for decreased CV and HF death (p=0.06 for both)
ALLHAT Long-term Follow-up Piller et al. 2011 (76) 21969009	Aim: To compare diuretic- based to ACE- inhibitor or CCB- based treatment of HTN Study type: RCT	Inclusion criteria: Age >55, HTN (SBP ≥140, DBP≥90), at least 1 CV risk factor (MI, stroke, LVH, diabetes, low HDL, PVD) Exclusion criteria:	Intervention: Amlodipine (8,898) 572 with in-trial HF, Lisinopril (8,904); 469 with in-trial HF Comparator: Chlorthalidone (15,002); 720 with in-trial HF	 <u>1° endpoint</u>: Adjusted mortality risk Increased mortality with intrial incident HF, both HF<i>p</i>EF: HR: 2.42 (95% CI: 2.08–2.81, p<0.001) and HFrEF: HR: 3.06; 95% CI: 2.67–3.51; p<0.001) 	 Increased HF mortality with incident HF, both HF<i>p</i>EF: HR: 3.81 (95% CI: 2.18–6.67, p<0.001) and HF<i>r</i>EF: HR: 6.80; 95% CI: 4.36–10.62; p<0.001) No difference in mortality in pts with incident HF by drug treatment

	<u>Size</u> : 32,804	Symptomatic HF, EF <35% at trial entry			
SHEP HF Results Kostis et al. 1997 (77) <u>9218667</u>	Aim: To assess the effect of antihypertensive treatment in isolated systolic HTN Study type: RCT Size: 4,736	Inclusion criteria: Age > 60, SBP 160– 219, DBP<90 Exclusion criteria: Recent MI or CABG, pts with DM, stroke, AF	Intervention: Antihypertensive therapy: step 1, chlorthalidone, step 2, atenolol (2,365) Comparator: Placebo (2,371)	 <u>1° endpoint:</u> Incident HF Active treatment decreased BP from mean of 170/77 to mean of and decreased HF events from 105 (4.4%) with placebo to 55 (2.3%) RR: 0.51 (95% CI: 0.37–0.71, p<0.001) at 4.5 y 	 1° results of SHEP showed decreased stroke risk with active treatment 149 (8.2%) with placebo to 96 (5.4%) RR: 0.64 (95% CI: 0.49–0.82, p=0.003) at 4.5 y LV function was not measured
CHARM-Preserved Yusuf et al. 2003 (78) <u>13678871</u>	Aim: To ascertain efficacy of candesartan in pts with HF <i>p</i> EF. Study type: RCT Size: 3,023	Inclusion criteria: HF pts in NYHA class II-IV with EF >40% Exclusion criteria: Creatinine >265 µmol/L (3.0 mg/dL), potassium >5.5 mmol/L, MI, stroke, or open-heart surgery in the previous 4 wk	Intervention: Candesartan (1,514) <u>Comparator</u> : Placebo (1,509)	 <u>1° endpoint</u>: CV death or admission for HF. No difference for candesartan 333 (22%) vs. placebo 366 (24%) at 3.5 y, HR: 0.89; 95% CI: 0.77–1.03; p=0.12) covariate adjusted HR: 0.86 (95% CI: 0.74–1.00); p=0.051) 	 No differences for 2° endpoints except for covariate adjusted risk of HF admission HR: 0.84 (95% CI: 0.70–1.00; p=0.047). CV death 11.2 vs. 11.3% HR: 0.99 (95% CI: 0.80–1.22; p=0.918). Adverse effects requiring discontinuation: hypotension (2.4 vs. 1.1%; p=0.009; increased creatinine, 4.8 vs. 2.4%; p=0.005; hyperkalemia 1.5 vs. 0.6%; p=0.029) Limitations: Some pts may have had previous EF <40%.
PEP-CHF Cleland et al. 2003 (79) <u>16963472</u>	Aim: To ascertain efficacy of perindopril in pts with HF <i>p</i> EF. Study type: RCT Size:	Inclusion criteria: Age ≥70, Rx with diuretics for clinical diagnosis of HF, echo criteria for diastolic dysfunction Exclusion criteria:	Intervention: Perindopril (424) Comparator: Placebo (426)	 <u>1° endpoint:</u> All-cause mortality or admission for HF. No difference for perinopril 107 (25.1%) vs. placebo 131 (23.6%) at 3 y, HR: 0.92; 95% CI: 0.70– 1.21; p=0.5. 	 HF hospitalization lower at 1 y with perindopril: 34 events (8.0%) vs. placebo 53 (12.4%), HR: 0.63; 95% CI: 0.41–0.97; p=0.033). Limitations: Many pts withdrew (40% by 18 mo), often to take open-label ACE inhibitors (36% by study end).

	850	Creatinine >200 µmol/L (2.3 mg/dL), potassium > 5.4 mmol/L			
I-PRESERVE Massie et al. 2008 (80) <u>19001508</u>	Aim: To ascertain efficacy of irbesartan on in pts with HF <i>p</i> EF. Study type: RCT Size: 4,128	Inclusion criteria: Age > 60, HF pts in NYHA class II-IV with EF >45% Exclusion criteria: Previous EF <40%, creatinine >222 μmol/L (2.5 mg/dL) ACS, stroke, or revascularization in the previous 3 mo	Intervention: Irbesartan (2,067) Comparator: Placebo (2,061)	 <u>1° endpoint</u>: CV death or hospitalization for CV cause. No difference for irbesartan vs. placebo (742 (36%) vs. 763 (37%), HR: 0.95; 95% CI: 0.86 – 1.05; p=0.35) 	 No differences for mortality or any other 2° endpoints Minnesota living with HF scale improved in both, groups to the same No difference in BNP levels No difference in adverse effects requiring discontinuation: doubling of creatinine, 6% vs. 4%; p<0.001; K >6.0 3% vs. 2%; p=0.01) Limitations: Study drug discontinuation in 34% of pts by end of study. High rate of concomitant ACE-I (40%)
NEAT-HF <i>p</i> EF Redfield et al. 2015 (81) <u>26549714</u>	Aim: To ascertain efficacy of isosorbide mononitrate on daily activity in pts with HF <i>p</i> EF. Study type: Double-blind crossover Size: 110	Inclusion criteria: Age ≥50 y on stable HF therapy, EF ≥50%, activity limited by dyspnea, fatigue, or chest pain Exclusion criteria: SBP <110mm Hg and >180 mm Hg, current nitrates or PDE-5 inhibitors	Intervention: Isosorbide mononitrate (110) Comparator: Placebo (110)	 <u>1° endpoint</u>: Average daily activity assessed by accelerometer units during 120 mg phase. Nonsignificant trend for lower daily activity in the treatment group. (-381 accelerometer units; 95% Cl: -780–17; p=0.06) and significant decrease in h of activity/d (-0.30 h; 95% Cl: -0.55– -0.05; p=0.02) 	 No differences for any of the 3 doses on QoL scores, 6MWT and levels of NT-proBNP (trend unfavorable for nitrates) Limitations: Rapid dose escalation of study drug.
Redfield et al. 2013 (82) <u>23478662</u>	Aim: To ascertain effects of sildenafil on exercise capacity in pts with HF <i>p</i> EF. Study type:	Inclusion criteria: Age ≥18 on stable HF therapy, EF ≥50%, peak VO ₂ <60% normal and either nt-proBNP >400 or elevated	Intervention: Sildenafil (113) Comparator: Placebo (103)	 <u>1° endpoint:</u> Change in peak VO₂ from BL at 24 wk No difference between sildenafil (-0.20, IQR -1.7–1.11) and placebo (-0.20, 	 No differences in clinical rank score or 6-min walk Limitations: Urinary cGMP levels were not increased in sildenafil group, raising questions about dosing. High prevalence of

	Double-blind <u>Size</u> : 216	PCWP <u>Exclusion criteria</u> : Systolic BP <110mm Hg and >180 mm Hg, MMI or revascularization within 60 d, eGFR <20 mL/min		IQR -0.70–1.0) • More worsening of renal function in sildenafil group (p=0.047)	chronotropic incompetence in study population.
TOPCAT Pitt et al. 2014 (83) <u>24716680</u> • New England Research Institutes Post-hoc analysis that captures differences in outcomes by geography - for reference list only	Aim: To assess the effects of spironolactone in pts with HF <i>p</i> EF. Study type: RCT Size: 3,445	Inclusion criteria: Symptomatic HF, Age ≥50y, LVEF ≥45% stratified according to - HF Hospitalization within past y - Elevated NPs Exclusion criteria: Renal disease (eGFR <30 or creatinine >22 µmol/L (2.5 mg/dL), systemic illness with life expectancy <3 y. Specific co- existing conditions, meds, and acute events	Intervention: Spironolactone (1,722) Comparator: Placebo (1,723)	 <u>1° endpoint and results:</u> Composite of CV mortality, HF hospitalization, or aborted cardiac arrest. No difference with spironolactone vs. placebo 320 (18.6%) vs. 351 (20.4%), HR: 0.89; 95% Cl: 0.77–1.04; p=0.138) 	 HF hospitalization was reduced with spironolactone 206 (12.0%) vs. 245 (14.2%), HR: 0.83; 95% CI: 0.69–0.99; p=0.04) Increased hyperkalemia (18.7% vs. 9.1%), decreased hypokalemia (16.2% vs. 22.9%) and more doubling of creatinine (10.2% vs. 7.0%) with spironolactone

TOPCAT Regional Analysis Pfeffer et al. 2015 (84) 25406305 Post-hoc analysis that captures differences in outcomes by geography	Aim: To assess regional differences in the effects of spironolactone in pts with HF <i>p</i> EF. Study type: RCT Size: 3,445	Inclusion criteria: Symptomatic HF, Age ≥50y, LVEF ≥45% stratified according to • HF Hospitalization within past y • Elevated NPs Exclusion criteria: Renal disease (eGFR <30 or creatinine >22 µmol/L (2.5 mg/dL), systemic illness with life expectancy <3 y. Specific co- existing conditions, meds, and acute events	Intervention: Spironolactone (1,722) Comparator: Placebo (1,723)	 <u>1° endpoint and results</u>: Composite of CV mortality, HF hospitalization, or aborted cardiac arrest across regions. 1° outcome events in 522 (29.5%) pts in the Americas and 149 (8.9%) in Russia/Georgia. 1° outcome event rates with spironolactone and placebo 10.4/100 pt y and 12.6/100 pt y in the Americas and 2.5/100 pt y and 2.3/100 pt y in Russia/Georgia. HR spironolactone vs. placebo 0.82; 95% Cl: 0.69–0.98; p=0.026) in the Americas and 1.10 95% Cl: 0.79– 1.51; p=0.12) in Russia/Georgia. 	 Spironolactone had markedly greater effects on BP (4.2 mm Hg drop vs. 0.6 mm Hg; p<0.001, potassium change relative to placebo (0.26 mmol/L vs. 0.08 mmol/L), and increase in creatinine (0.10 vs. 0.02 mg/dL; p<0.001) Limitations: post-hoc analysis
Chen et al. 2015 (85) 25598008	Aim: To assess effects of MRAs in pts with HF <i>p</i> EF. Study type: Meta-analysis Size: 14 RCTs with 6,428 pts	Inclusion criteria: Prospective, RCTs that enrolled adult pts with LVEF ≥40% (including post-MI and those with symptomatic or asymptomatic HF) with a study duration of ≥4 mo that assessed at least 1 clinical outcome of interest.	Intervention: MRAs (3,249) Comparator: Placebo (2,861) Or standard therapy (301) Or active comparator (31)	 <u>1° endpoint and results:</u> All-cause mortality and HF hospitalization No difference in all-cause mortality (RR: MRAs vs. placebo 0.90; 95% CI: 0.78–1.04; p=0.17) Reduced risk of HF hospitalization (RR: MRA vs. placebo 0.83; 95% CI: 0.70–0.98; p=0.03) <u>1° Safety endpoint :</u> More hyperkalemia with MRAs (12.2% vs. 6.2%, p<0.001) 	 MRAs improved QOL (weighted mean difference -5.2; 95% CI: -8.02.3). MRA's improved echo indices of LV function: E/e', E/A ratio, deceleration time, interventricular relaxation time Renal failure in 1.19% of pts with MRAs vs. 0.39% Gynecomastia in 2.81%R vs. 0.3% Limitations: discrepancies in definitions of HF<i>p</i>EF in different trials; heterogeneity of trial outcomes and their assessment, including follow-up duration; 1° outcome results driven by

					TOPCAT
	Aim	Inclusion criteria:	Intervention:	1º endnoint:	Effect percisted after adjustment
Solomon et al. 2012 (61) <u>22932717</u>	To address safety and efficacy of LCZ696 in pts with HF <i>p</i> EF. Study type: RCT Size: 308	Pts ≥40 y of age, LVEF ≥45%, NYHA class II-III HF, NT-pro BNP >400 pg/mL Exclusion criteria: Previous EF <45%, isolated right HF, noncardiac dyspnea, CAD or CVD needed revascularization <3 mo Right HF due to pulmonary disease, dyspnea due to noncardiac causes, valvular/myocardial disease, CAD or CVD needing revascularization within 3 mo of screening.	LCZ696 (149) Comparator: Valsartan (152)	 Change in BNP at 12 wk Greater reduction with LCZ696 (ratio of change compared to valsartan 0.77; 95% CI: 0.64–0.92; p=0.001) <u>1° Safety endpoint</u>: Serious adverse events 15% in LCZ676 group and 20% in valsartan group (p=NS) 	 for more lowering of BP in LCZ676 group Improvement in NYHA class at 36 wk in LCZ676 group compared to valsartan. Reduction of LA size at 36 wk in LCZ676 group compared to valsartan. BNP levels higher than in other HF<i>p</i>EF trials, perhaps because this was an entry criterion.

Date: Some studies added by chairs in December 2015, others added by the writing committee.

Data Supplement D. RCTs Comparing Anemia (Section 9.2)

Study Acronym;	Aim of Study;	Patient Population	Study Intervention	Endpoint Results	Relevant 2° Endpoint (if any);
Author;	Study Type;		(# patients) /	(Absolute Event Rates,	Study Limitations;
Year Published	Study Size (N)		Study Comparator	P values; OR or RR; &	Adverse Events
			(# patients)	95% CI)	

CONFIRM-HF	Aim:	Inclusion criteria:	Intervention:	1° endpoint:	2°Endpoints:
Ponikowski et al. 2015	To assess benefits	Pts at least 18 y,	FCM (152)	Change in 6MWT distance	Changes in NYHA class
(86)	and safety of long	NYHA class II or III,		from BL to wk 24	• PGA
<u>25176939</u>	term FCM in iron-	LVEF≤45%,	Comparator:	Results: Change in 6MWT	 6MWT distance
	deficient pts with	elevated NPs, ID	Placebo (152)	distance FCM vs. placebo of	Fatigue score
	HF	defined as ferritin		33±11 m (p=0.002)	• KCCQ
		<100 ng/mL, or		, , , , , , , , , , , , , , , , , , ,	• FQ-5D
 Vifor Inc. 	Study type:	ferritin 100–			 Assessed at wk 6, 12, 24, 36, 52
 ICON Clinical 	RCT (1:1)	300 ng/mL if TSAT			• Rate of any hospitalization, rate of
Research		<20%, Hb <15			hospitalization for any CV reason
	<u>Size</u> :	mg/dL			and rate of hospitalization due to
	304				worsening HF ⁻
		Exclusion criteria:			 Time to first hospitalization for any
		Pts in need of			reason, time to first hospitalization
		transfusion, if not			for any CVCV reason and time to
		able to complete			first hospitalization due to worsening
		6MW I, uncontrolled			HF;
					 Time to death for any reason, time
		impaired liver or			to death for any CV reason, and time
		renal function			to death due to worsening HF.
					<u>Results:</u>
					 Significant improvements in NYHA
					class, PGA, QoL and Fatigue
					scores, 6 MWD up to 52 wk
					 Significant reduction in the risk of
					hospitalizations for deteriorating HF,
					HR: 0.39 (95% CI: 0.19–0.82)
					(p=0.009)
					Preserved treatment effect across
					subgroups
					No differences in adverse events
					when compared to placebo
					 Study was not designed to test
					morbidity and mortality outcomes of
					the ID therapy with FCM

FAIR-HF Anker et al. 2009 (87) <u>19920054</u>	Aim: To evaluate the effects of intravenous iron (FCM) on HF symptoms in pts with systolic HF and ID, with and without anemia. Study type: RCT (2:1) Size: 459	Inclusion criteria: • Chronic HF • NYHA class II or III, • LVEF ≤40% (for pts in NYHA class II) or ≤45% (for pts in NYHA class III), • Hemoglobin level 95–135 g/L • ID Exclusion criteria: • Uncontrolled HTN • Other clinically significant heart disease • Inflammation • Clinically significantly impaired liver or renal function.	Intervention: Ferric carboymaltose 200 mg weekly until hemoglobin was corrected (n=304) Comparator: Placebo (n=155)	 <u>1° endpoint</u>: PGA at 24 wk Results: improvement in the FCM group compared to placebo 50% much or moderately improved vs. 28% (OR for being in a better rank, 2.51; 95% Cl: 1.75–3.61; p<0.001) NYHA class at 24 wk Results: improvement in the FCM arm compared to placebo 47% with NYHA I or II vs. 30% in the placebo arm (OR for improvement by 1 class, 2.40; 95% Cl: 1.55–3.71; p<0.001) <u>1° Safety endpoint</u>: Trend towards fewer HF hospitalizations in the FCM group (p=0.08) 	 Improvement in the FCM group in PGA and NYHA at wk 4 and 12 (p<0.001) Mean improvement in 6MWT of 35±8m at 24 wk (p<0.001); also significant improvements at 4 and 12 wk Significant improvement in the EQ-5D and in KCCQ
RED-HF Swedberg et al. 2013 (88) <u>23473338</u> • Amgen	Aim: To assess effects of darbepoetin alfa on pts with systolic HF and anemia. Study type: RCT Size: 2,278	Inclusion criteria: NYHA class II, III, or IV HF; LVEF≤40%; Hgb: 9.0–12.0 g/dL; on guideline- recommended HF treatment. Exclusion criteria: Transferrin saturation <15%, bleeding or other causes of anemia, serum creatinine >3 mg/dL, BP	Intervention: Darbepoetin alfa (1,136) Comparator: Placebo (1,142)	 <u>1° endpoint</u>: Composite of death from any cause or hospitalization for worsening HF Results: 1° outcome occurred in 576 pts in the darbepoeitin alfa group vs. 562 in the placebo group (HR: 1.01; 95% CI: 0.90– 1.13; p=0.87) <u>1° Safety endpoint</u>: Increased thromboembolic adverse events in the treatment group (p=0.01); 	Limitation: pts with severe anemia were excluded

>160/100 mm Hg.	No significant increase in	
	fatal/nonfatal strokes in	
	treatment group and similar	
	cancer-related adverse	
	events between groups	

Date: Chairs selected trials in December 2015. One trial added by writing committee.

Data Supplement E. RCTs Comparing HTN (Section 9.5)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% Cl)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Xie et al. 2016 (89) <u>26559744</u>	Aim: To assess the efficacy and safety of intensive BP lowering strategies. SR and meta- analysis Size: 19 trials with 44,989 pts; 3.8 y of follow- up.	Inclusion Criteria: RCTs with different BP targets or different BP changes between more vs. less intense therapy with at least 6 mo follow-up. Exclusion Criteria: Trials that did not assess a different target or relevant outcome.	5 RCTs (6,960 pts) enrolled only pts with DM and 6 trials (2,809 pts) specifically recruited pts with CKD.	 <u>1° Outcomes</u>: Major CV events, defined as MI, stroke, HF or CV death, separately and combined; nonvascular and all-cause mortality; ESRD; and adverse events; new onset microalbuminuria/macroalbuminu ria or change from micro- to macroalbuminuria and retinopathy in pts with DM. <u>Results</u>: Pts in the more intensive BP- lowering treatment group had mean BP 133/76 mm Hg compared with 140/81 mm Hg in the less intensive group. Intensive BP-lowering treatment achieved RR reductions for major CV events: 14% (95% CI: 4, 22), MI: 13% (95% CI: 0, 24), stroke: 22% (95% CI: 10, 32), albuminuria: 10% (95% CI: 3, 16), and retinopathy progression: 19% (95% CI: 0–34). However, more 	 Study Limitations: Only 6,960 pts with DM were included in the total study size of 44,989 pts. Conclusions: The absolute CV benefits were greatest in trials in which all enrolled pts had vascular disease, renal disease or DM. However, only 6,960 of the 44,989 pts had DM and no subanalysis for DM was provided; however, the outcome benefits were qualitatively most striking for pts with DM, CKD and/or vascular disease.

				intensive treatment had no clear	
				effects on HF: RR: 15% (95% CI:	
				-11, 34), CV death: 9% (-11, 26),	
				total mortality: 9% (95% CI: -3,	
				19), or ESRD: 10% (95% CI: -6,	
				23). The reduction in major CV	
				events was consistent across pt	
				groups, and additional BP	
				lowering had a clear benefit even	
				in pts with SBP <140 mm Hg. The	
				absolute benefits were greatest in	
				trials in which all enrolled pts had	
				vascular disease, renal disease,	
				or DM. Serious adverse events	
				associated with BP lowering were	
				only reported by 6 trials and had	
				an event rate of 1.2% per v in	
				intensive BP lowering group pts.	
				compared with 0.9% in the less	
				intensive treatment group (RR	
				1 35 (95% CI: 0 93 1 97))	
				Severe hypotension was more	
				frequent in the more intensive	
				treatment regimen (RR: 2.68	
				$(95\% \text{ Cl} \cdot 1.21.5.89) \text{ n=}0.015)$	
				but the absolute excess was	
				small (0.3%) vs. 0.1% par at v for	
				the duration of follow up)	
SDDINT	Aim:	Inclusion criteria:	Intervention:	19 Endpoint	Summary:
Wright at al. 2015	To tost the	$\frac{111Clusion Chiena.}{SPD > 130 mm Ha}$	Intervention.	<u>1º Enapoint:</u>	<u>Summary.</u>
(00)	offectiveness of a	$SDF \ge 150$ mini Hy,	treatment to goal SPD	• Composite of IVI, non-IVI ACS,	• More intensive SBP lowering to
(90)			120 mm Hg (4.679)	Stroke, ADHF, CV death; HR:	a goal of < 120 mm Hg with
20001272	yuai SDF <120	of pro-trial DD	<120 mm Hg (4,070)	0.75 (95% CI: 0.64, 0.89)	achieved mean of ~121 mm Hg
	$\frac{11111}{2} = \frac{1}{2} = $	or pre-triar BP-	Compariaon		resulted in less CVD and lower
	SBP < 140 mm Hg	iowening meas	Comparison:	 Lower BP target reduced 	total mortality over 3.26 y in
	tor the prevention	Increased.	Standard BP lowering	composite outcome 243 pts	comparison with a goal SBP
		Age ≥50 y	treatment to goal SBP	(1.65%/y) vs. higher target 319	<140 mm Hg and achieved SBP
	588 ≤130 mm Hg	Presence of at least	<140 mm Hg (4,678)	(2.19%/y), HR: 0.75; 95% CI:	ot ~135 mm Hg.
	at BL.		 Net treatment difference 	0.64–0.89; p<.001) and death:	 There were small increases in
		Clinical or	~3 drugs (2.8) on average	lower target 155 vs. 201, HR:	some expected SAEs. Perhaps
	Study type:	subclinical CVD	vs. 2 drugs (1.8) on	0.73; 95% CI: 0.60–0.90;	unexpected, a sizable increase

RCT	 CKD stage 3 or 	average	p=0.003)	in reduced eGFR in the non-CKD
	greater	 During the trial, mean 		group and AKI/ARF overall was
Size:	• Age ≥75	SBP was 121.5 vs. 134.6.		observed in the intensive group.
9361 pts followed	Framingham		Other endpoints:	While of uncertain etiology and
median of 3.26 y.	General CVD risk		 Total deaths HR: 0.73 (95% CI: 	significance, there is speculation
	≥15% in 10 y		0.60–0.90)	this could be an acute
			 1° or death HR: 0.78 (95% CI: 	hemodynamic effect, especially
	Exclusion criteria:		0.67–0.90)	given the findings regarding
	DM, history of		 Components of 1° composite 	albuminuria.
	stroke, ESRD		mostly consistent in direction	 Low target significantly reduced
	(eGFR <20		other than ACS – no difference.	HF: HR: 0.62 (95% CI: 0.45–
	mL/min),			0.84; p=0.002)
	anticipated survival		CKD outcomes:	 No difference in composite or
	<3 y		 1° in CKD pts: reduction in GFR 	individual renal outcomes with
			of ≥50% or ESRD HR: 0.89 (95%	lowering of BP
			Cl: 0.42, 1.87)	
			 Incident albuminuria HR: 0.72 	1
			(95% 0.48, 1.07)	Limitations:
			In pts without CKD: reduction in	rew pis were unitedied at DL
			GFR \geq 30% and to <60	~9%, SUSERINT provides inflie if
			• HR: 3.49 (95% CI: 2.44–5.10)	BP lowering medication initiation
			Incident albuminuria HR: 0.81	for untreated people with SRP
			(95% CI: 0.63–1.04)	130–139
				100 100.
			Adverse events:	
			• SAEs: 1.04, p=0.25	
			 Significant absolute increases 	
			seen in intensive group for	
			hypotension (1%), syncope	
			(0.6%), electrolyte abnormality	
			(0.8%), AKI/ARF (1.6%) over the	
			study period.	
			• 1.7% fewer pts had orthostatic	
			hypotension in intensive group,	
			p=0.01.	

SPRINT Senior	Aim:	Inclusion:	Intervention:	1 endpoint:	Limitations:
Williamson et al.	Intensive SBP goal	Men and women age	Medications and dietary	Composite CVD outcome (AMI,	Does not apply to nursing home
2016	<120mmHq) vs	75+; mean age	advice to achieve SBP of	non-MI ACS, Stroke, HF, CVD	patients or those with dementia
(91)	standard (SBP	79.8 v; 38%	<120 mm Hg	death.	
27195814	goal <140)	women: 17%	ů – Č		Conclusions:
	5	black, 74%	Comparator:	Results: 102 events in the	Intensive SBP is safe and effective
	Study Type:	Caucasian:	Medications and dietary	intensive treatment group vs 148	for lowering CVD events and
	RCT	Exclusions:	advice to achieve SBP of	events in the standard treatment	total mortality in persons age 75
		Nursing home	<140 mm Ha	aroup: HR: 0.66:	and older
	Size:	residents:	5	95%CI: 0.51–0.85 and all-cause	
	2.636	diabetes, Stroke,	Achieved SBP:	mortality (73 deaths vs. 107	
	,	symptomatic HF in	Intensive= 123.4 mm Hg	deaths, respectively; HR; 0.67;	
	30% met criteria for	past 6 mo or EF	Standard= 134.8 mm Hg	95%CI: 0.49-0.91. No significant	
	being classified as	<35%, dx or	Ũ	difference in falls, orthostatic	
	ambulatory frail	treatment of		hypotension, or overall SAEs.	
	,	dementia,		NNT for primary outcome=27 and	
	Mean follow-up:	unintentional wt		NNT for all-cause mortality=41	
	3.1 y	loss >10% in past			
		5 mo. SBP<110			
		after standing 1			
		min, expected			
		survival <3y			
TOPCAT Regional	Aim:	Inclusion criteria:	Intervention:	1° endpoint and results:	Spironolactone had markedly
Analysis	To assess regional	Symptomatic HF,	Spironolactone (1,722)	Composite of CV mortality, HF	greater effects on BP (4.2 mm
Pfeffer et al. 2015	differences in the	Age ≥50y, LVEF		hospitalization, or aborted cardiac	Hg drop vs. 0.6 mm Hg; p<0.001,
(84)	effects of	≥45% stratified	Comparator:	arrest across regions.	potassium change relative to
25406305	spironolactone in	according to	Placebo (1,723)	• 1° outcome events in 522	placebo (0.26 mmol/L vs. 0.08
	pts with HF <i>p</i> EF.	 HF Hospitalization 		(29.5%) pts in the Americas and	mmol/L), and increase in
		within past y		149 (8.9%) in Russia/Georgia, 1°	creatinine (0.10 vs. 0.02 mg/dL;
Post-hoc analysis that	Study type:	 Elevated NPs 		outcome event rates with	p<0.001)
captures	RCT			spironolactone and placebo	Limitations: post-hoc analysis
differences in		Exclusion criteria:		10.4/100 pt v and 12.6/100 pt v in	
outcomes by	<u>Size</u> :	Renal disease		the Americas and 2.5/100 pt v	
geography	3,445	(eGFR <30 or		and 2.3/100 pt v in	
		creatinine >22		Russia/Georgia. HR	
		µmol/L (2.5		spironolactone vs. placebo 0.82:	
		mg/dL), systemic		95% CI: 0.69–0.98; p=0.026) in	
		illness with life		the Americas and 1.10 95% CI:	
		expectancy <3 y.		0.79–1.51; p=0.12) in	
		Specific co-existing		Russia/Georgia.	

		conditions, meds.			
		and acute events			
Law et al., 2009	Study type:	Inclusion criteria:	1° endpoint:	With the exception of the extra	
(92)	Meta-analysis of use	The database	CAD events; stroke	protective effect of beta blockers	
<u>19454737</u>	of BP lowering	search used		given shortly after a MI and the	
	drugs in	Medline (1966-	Results:	minor additional effect of CCBs in	
	prevention of CVD	Dec. 2007 in any	In 37 trials of pts with a	preventing stroke, all the classes	
	from 147	language) to	history of CAD, beta	of BP lowering drugs have a	
	randomized trials	identify	blockers reduced CAD	similar effect in reducing CAD	
		randomized trials	events 29% (95% CI:	events and stroke for a given	
	<u>Size</u> :	of BP lowering	22%–34%). In 27 trials in	reduction in BP.	
	Of 147 randomized	drugs in which	which beta blockers were		
	trials of 464,000	CAD events or	used after acute MI, beta		
	pts, 37 trials of	strokes were	blockers reduced CAD		
	beta blockers in	recorded. The	events 31% (95% CI:		
	CAD included	search also	24%-38%), and in 11		
	38,892 pts, and 37	included the	trials in which beta		
	trials of other	Cochrane	blockers were used after		
	antihypertensive	Collaboration and	long term CAD, beta		
	drugs in CAD	Web of Science	blockers insignificantly		
	included 85,395	databases and the	reduced CAD events		
	pts	citations in trials	13%. In 7 trials, beta		
		and previous meta-	blockers reduced stroke		
		analyses and	17% (95% CI: 1%–30%).		
		review articles.	CAD events were reduced		
			14% (95% CI: 2%–25%)		
		Exclusion criteria:	in 11 trials of thiazide		
		Trials were	diuretics, 17% (95% CI:		
		excluded if there	11%–22%) in 21 trials of		
		were <5 CAD	ACE inhibitors,		
		events and strokes	insignificantly 14% in 4		
		or if treatment	trials of angiotensin		
		duration was <6	receptor blockers, and		
		mo.	15% (95% CI: 8%–22%)		
			in 22 trials of CCBs.		
			Stroke was reduced 38%		
			(95% CI: 28%–47%) in 10		

			trials of thiazide diuretics, 22% (95% CI: 8%–34%) in 13 trials of angiotensin- converting enzyme inhibitors, and 34% (95% CI: 25%–42%) in 9 trials of CCBs.		
Aronow et al. 1997 (93) <u>9230162</u>	Aim: To determine effect of propranolol vs. no propranolol on mortality plus nonfatal MI in pts with prior MI and HF <i>p</i> EF	Inclusion criteria: Pts ≥62 y with MI and LVEF ≥40% and HF NYHA class II or III treated with diuretics and ACE inhibitors for 2 mo	Intervention: 79 pts were randomized to treatment with propranolol Comparator: 79 pts were randomized to no propranolol. All pts continued diuretic and ACE inhibitor therapy.	<u>1° endpoint</u> : At 32-mo mean follow-up, multivariate Cox regression analysis showed that compared with no propranolol, propranolol reduced mortality 35% (p=0.03) and mortality plus nonfatal MI 37% (p=0.018)	Relevant 2° Endpoint: At 1-y follow-up, LVEF was increased by propranolol from 57% to 63% (p<0.001) and LV mass was decreased by propranolol from 312 grams to 278 grams (p=0.001) Propranolol was stopped because of adverse effects in 11 of 79 pts (14%)
Van Veldhuisen et al. 2009 (94) <u>19497441</u>	<u>Aim</u>: To determine the effect of nebivolol vs. placebo in pts with HF <i>r</i> EF and HF <i>p</i> EF	Inclusion criteria: Pts ≥70 y history of HF and HF/EF or HF <i>p</i> EF	Intervention/Comparator: 1,359 pts with a history of HF/EF and 752 pts with a history of HF/EF were randomized to nebivolol or to placebo	<u>1° endpoint</u> : At 21-mo follow-up, the primary endpoint of all-cause mortality or CV hospitalization was reduced by nebivolol 14% (95% CI: 0.72– 1.04) in pts with HF/EF and 19% (95% CI: 0.63, 1.04) in pts with HF/DEF	Relevant 2° Endpoint: HR for reduction of all-cause mortality by nebivolol: 0.84 (95% Cl: 0.66–1.08) for HF <i>r</i> EF and 0.91 (95% Cl: 0.62–1.33) for HF <i>p</i> EF
Yusuf et al. 2003 (78) <u>13678871</u>	Aim: To determine the effects of candesartan vs. placebo in pts with HF <i>p</i> EF	Inclusion criteria: 3,023 pts, mean age 67 y, with HF <i>p</i> EF and NYHA class II-IV HF	Intervention/Comparator: 3,023 pts were randomized to candesartan or placebo	<u>1° endpoint:</u> At 36.6 m follow-up, the primary outcome of CV death or hospitalization for HF was reduced 11% (p=0.118) by candesartan	Relevant 2° Endpoint: Hospitalization was reduced 16% (p=0.047) by candesartan
Massie et al. 2008 (80) <u>19001508</u>	<u>Aim</u> : To determine the effect of irbesartan vs. placebo on all- cause mortality or hospitalization for a CV cause in pts with HF <i>p</i> EF	Inclusion criteria: Pts 60 y and older with HF <i>p</i> EF and NYHA class II, III, or IV HF	Intervention/Comparator 4,128 pts were randomized to irbesartan or placebo	<u>1° endpoint</u> : At 49.5-mo follow-up, the primary outcome of all-cause mortality or hospitalization for CV cause was reduced 5% by irbesartan (p=0.35)	Relevant 2° Endpoint: Irbesartan did not significantly reduce the secondary outcomes of death from HF or hospitalization for HF, death from any cause and from CV causes, and quality of life

Piller LB, et al., 2011 (76) <u>21969009</u>	Aim: To determine mortality rates in pts who developed HF in ALLHAT	Inclusion criteria: 1,761 pts, mean age 70 y, developed HF during ALLHAT	Intervention/Comparator At 8.9-y mean follow-up, 1,348 of 1,761 pts (77%) with HF died	<u>1° endpoint:</u> Post-HF all-cause mortality was similar for pts treated with chlorthalidone, amlodipine, and lisiopril. 10-y adjusted rates for mortality were 86% for amlodipine, 87% for lisinopril, and 83% for chlorthalidone	Relevant 2° Endpoint: All-cause mortality rates were similar for those with HF <i>r</i> EF (84%) and for those with HF <i>p</i> EF (81%) with no significant differences by randomized treatment arm
Lv et al. 2013 (95) <u>23798459</u>	MA of RTC that randomly assigned individuals to different target BP levels	15 trials including a total of 37,348 pts.	 7.5/4.5 mm Hg BP difference. Intensive BP lowering achieved. <u>RR for</u> Major CV events: 11%; 95% CI: 1%–21%) MI: 13%; 95% CI: 0%– 25% Stroke: 24%; 95% CI: 8%–37% ESRD: 11%; 95% CI: 3%–18% Albuminuria: 10%; 95% CI: 4%–16% Retinopathy 19%; 95% CI: 0%–34% p=0.051 	More intensive strategy for BP control reduced cardio-renal end point	

Date: Chairs selected trials in October 2016.

Data Supplement F. Nonrandomized Trials for Hypertension (Section 9.5)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Primary Endpoint and Results (P values, OR or RR & 95 % CI)	Summary / Conclusion / Comments
Thomopoulos et al. 2016 (96) <u>26848994</u>	Meta-analysis of RCT's of more versus less intense BP control	16 trials (52,235 pts) compared more vs. less intense treatment 34 (138,127 pts) active vs. placebo	 More intense BP Stroke RR: 0.71; 95% CI: 0.60–0.84) Coronary heart disease RR: 0.80; 95% CI: 0.68–0.95) Major CV events RR: 0.75; 95% CI: 0.68–0.85 CV mortality RR: 0.79; 95% CI: 0.63–0.97 Stratification of SBP cutoffs (150, 140 and 130 	 Intensive BP reduction improves CV outcomes compared to less intense Achieved BP of <130/80 mm Hg may be associated with CV benefit.

	mmHg) showed that a SBP/DBP difference of _10/_5mmHg across each cutoff reduced risk of all outcomes	

Date: Chairs selected trials in October 2016.

Data Supplement G. RCTs Comparing Treatment of Sleep Disorders (CPAP makers) (Section 9.6)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% Cl)	Relevant 2° Endpoint; Study Limitations; Adverse Events
SAVE McEvoy et al. 2016 (97) <u>27571048</u>	Aim: To whether treatment with CPAP prevents major CV events. Study type: RCT with 1 wk run-in on sham CPAP Size: n=2,717	 Inclusion criteria: Adults 45 - 75 y of age Moderate-to-severe OSA Coronary or cerebrovascular disease Exclusion criteria: 	Intervention: CPAP treatment plus usual care (CPAP group) Comparator: Usual care alone (usual-care group)	 <u>1° endpoint:</u> Composite of death from CVD, MI, stroke, or hospitalization for UA, HF, or TIA <u>Results:</u> Duration of CPAP=3.3 h/night; AHI events/h decreased from baseline to end of follow up at 3.7 y, 29.0–3.7 events/h Primary endpoint – no significant difference in CPAP vs usual-care group (n=229, 17.0% vs. n=207; 15.4%; HR: 1.10 with CPAP; 95% CI: 0.91–1.32; p=0.34). No significant difference in any individual or other composite CV end point. CPAP significantly reduced snoring and daytime sleepiness and improved health-related quality of life and mood. 	 Secondary end points: Other CV outcomes Health-related quality of life Snoring symptoms Daytime sleepiness Mood Study Limitations: Primarily men with moderate-to-severe OSA and minimal sleepiness Adverse Events:
ORBIT-AF Holmovist et al. 2015	<u>Aim</u> : 1) Define	Inclusion criteria: • >18 years of age	Intervention: N/A	<u>1° endpoint:</u> • All-cause mortality:	Secondary end points: N/A
(98) <u>25965712</u>	frequency of diagnosed	Electrocardiographic evidence of AF	Comparator: N/A	 First all-cause hospitalization; Composite of first event of CV 	Study Limitations:

004				
USA among		Multicenter,	death, stroke/non-central	 Voluntary, observational
nationwide	Exclusion criteria:	ambulatory-based	nervous system embolism, TIA,	study - selection &
AF	 Life expectancy of <6 months or 	registry	or MI;	reporting biases
population;	AF secondary to reversible		First major bleed within 2 years	 No randomization -
2) Determine	conditions		of baseline enrollment in registry	Voluntary, observational
whether OS	A			study - selection &
is associate	d		Results:	reporting biases
w/·	-		Frequency of diagnosed OSA	\circ OSA diagnosis made on
a) Worse			among nationwide AE population	basis of physician report
	.c.		• 18% (n =1 841)	& medical records
b) Arrhythn	ic		OSA associations w/ outcomes	No data on average
			USA associations w/ outcomes	duration of CDAD upo por
AF			• Figher fisk of.	uuralion or CFAF use per
progress			 Hospitalization (43 vs 35 	night Maturation sharped in
			events/100 patient-years	 Maturation – changes in
3) Determine			among patients without USA	subjects over 2 years not
whether			[adjusted hazard ratio (HR),	accounted for in data
СРАР			1.12; 95% confidence interval	
treatment is			(Cl), 1.03-1.22; p=.0078]	Adverse Events:
associated	v/		No higher risk of:	N/A
outcomes ir			 Death (HR, 0.94; 95% CI, 	
patients w/			0.77-1.15; p=.54);	
AF & OSA.			 Composite of CV death, 	
			stroke/non-central nervous	
Study type:			system embolism, TIA, or MI	
Prospective			(HR. 1.07: 95% Cl. 0.85-1.34:	
descriptive.			p=.57):	
correlationa	1		\circ First major bleeding (HR 1 18)	
comparative			95% CL 0 96-1 46: n= 11)	
time-series	,		OSA associations w/ AF	
design			progression	
Data			Not associated w/ higher risk of	
collection at			AE progression (HR 1.06: 95%	
enrollment			$CI = 0.89 \pm 28$; n = 51)	
6 month			CPAP treatment association w/	
o-monul intervale for			<u>CFAF ilediment association w/</u>	
	2			
	<u> </u>			
years			Less likely to progress to more	
			permanent forms of AF versus	
<u>Size</u> : National	y		patients w/out CPAP (HR, 0.66;	
representative			95% Cl, 0.46-0.94; p=.021).	

				-	
	sample enrolled consecutively • n=10,132 w/ AF o n=1,841 w/ AF & OSA o n=1,837 patients w/ OSA & complete CPAP data o n =1,763 patients w/ OSA & 2- year outcomes data o n=937 patients w/ AF, OSA, & CPAP treatment Sites: 176 national sites that w/ provider & geographic heterogeneity				
SERVE-HF Cowie et al. 2015	Aim: Effects of	Inclusion criteria: • Chronic HF (defined as ≥12 wk	Intervention: Adaptive servo	 <u>1° endpoint</u>: Death from any cause 	<u>2° Endpoint</u> • CV death
(99) 26323938	adaptive servo- ventilation in	since diagnosis) according to current ESC guidelines	ventilation use ≥5h/night, 7d/wk.	Lifesaving CV intervention (cardiac transplantation	 Unplanned hospitalization from any cause
	HF pts with	• LVEF ≤45%	(n=666)	implantation of a ventricular assist	 Time to death from CV
ResMed The Oliviael	reduced EF	 Hypopnea index of ≥10/h 	Comparator:	device, resuscitation after sudden	causes
 The Clinical Research Institute 		 Stable, GDMT NYHA class III or IV, or NYHA 	GDMT (n=659)	cardiac arrest, or appropriate	 Change in NYHA class Change in 6 MWT (both at
GmbH	Study type:	class II with ≥1 hospitalization for		Unplanned hospitalization for HF	 Gliow-up visits).
	RCT	HF in the last 24 mo		Significant Results	General QoL (EuroQOL)
	Size:	 No hospitalization for HF in 4 wk prior to eprolment 		 All-cause mortality was higher with the intervention (34.8%) than 	HF-specific QoL (MLWHF)
					 Daytime sleepiness

1,325	 Optimized GDMT 	control (29.3%; HR: 1.28; 95% CI:	(Epworth Sleepiness Scale)		
	No new class of disease-	1.06–1.55; p=0.01).			
	modifying drug for prior >4 wk	CV mortality was higher with the	Limitations:		
	$\Delta H > 15/b$ with $\geq 50\%$ control	intervention (29.9%) than control	 Unblinded study - more 		
	• AFI > 15/11 With > 50 % Certifal	(24.0%· HP· 1.34· 95% CI- 1.00	likely to favor treatment		
	events and a central API \geq 10/11	(24.0%, 110, 1.04, 90% 01, 1.09 - 1.65, p=0.006)	group, particularly for OOL		
		1.03, p=0.000).	group, particularly for QOL,		
	Exclusion criteria:	• 6IVIVV I decreased over time and			
	 Significant COPD with a forced 	were significantly lower with the	seen		
	expiratory volume in 1 s in 4 wk	intervention than with the control	 HF pts with reduced EF only 		
	before randomization	(p=0.02).	 HF pts with predominantly 		
	 O₂ saturation ≤90% at rest during 	 Daytime sleepiness decreased 	CSA not obstructive sleep		
	d	over time and was significantly	apnea.		
	 Currently receiving PAP therapy 	lower with the intervention than	 Sample had very limited # of 		
	Cardiac surgery, PCI, MI or UA	with the control (p<0.001).	women but reflects		
	within the previous 6 mo		epidemiology of CSA with		
	Cardiac resynchronization	Non-Significant Results	HF/EF		
	therapy implantation scheduled or	Unplanned hospitalization for HF			
	performed within 6 mo prior to	was not significantly higher with			
	randomization	the intervention (43.1%) than			
		control (41.3% HR 1.13 95% CI			
	TIA of stroke within the previous	0.95–1.33: n=0.16)			
	3 mo	• Of the lifesaving CV interventions			
	 1° hemodynamically-significant 	• Of the mesaving OV interventions,			
	uncorrected VHD (obstructive or	the intervention then control			
	regurgitant) or any valvular				
	disease expected to require	(p=0.08–0.61)			
	surgery during the trial;	Unplanned nospitalization for any			
	 Acute myocarditis/pericarditis 	cause was not significantly lower			
	within the previous 6 mo	with the intervention (67.9%) than			
	 Untreated or therapy-refractory 	control (68.0%; HR: 1.05; 95% CI:			
	restless leas syndrome	.92–1.20; p=0.47)			
	 Contraindication to the use of 	 The NYHA class change was not 			
	AutoSet CS2 because of	significantly different with the			
	symptomatic hypotonsion or	intervention than with the control			
	symptomatic mypotension of	(p=0.46)			
	significant intravascular volume	General QoL trends were not			
	depietion or prieumothorax or	significantly higher with the			
	prieumomediastinum	intervention than with the control			
	Pregnancy	(n=0.09)			
		HE specific Ool trands were not			
		• IIF-specific QUL fields were not			
		significantly higher with the			
				intervention than with the control $(p=0.92)$	
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CANPAP Arzt et al. 2007 (100) <u>17562959</u>	Aim: Investigate whether suppression of CSA below threshold by CPAP would LVEF & ht tx- free survival. Study type: Post hoc analysis of RCT Size:100	Inclusion criteria: • Age 18 to 79 y • NYHA II-IV • HF due to ischemic, hypertensive, or idiopathic DCM • Stabilized w/ optimal medical therapy for ≥1 mo • LVEF <40% • CSA Exclusion criteria: • Pregnancy • MI • Unstable angina • Cardiac surgery w/in 3 mo of enrollment • OSA	Intervention: • CPAP=CSA suppressed, n=57 • CPAP=CSA suppressed, n=43 <u>Comparator:</u> Control, n=110:	 Intervention than with the control (p=0.92). <u>1° endpoint</u>: Transplant free survival - Combined rate of all-cause mortality & ht tx Significant Results <u>1° endpoint</u>: Transplant free survival Significantly different between 3 groups (p=0.016) Significantly higher in CPAP-suppressed vs. control group (p<0.043) No difference between CPAP-unsuppressed vs. control group (p<0.26) <u>2° endpoint</u>: AHI AHI significantly > reduction in both CPAP-suppressed (p<0.001) and CPAP-unsuppressed (p<0.001) and CPAP-unsuppressed (p<0.001) and CPAP-unsuppressed (p<0.001) and CPAP-unsuppressed (p<0.002) than control groups AHI significantly > reduction in CPAP-suppressed (p<0.002) than control groups Mean nocturnal SaO2 Mean nocturnal SaO2 significantly > increased in CPAP-suppressed vs. control group (p<0.001) No significant difference between CPAP-suppressed vs. control group (p<0.001) 	 <u>2° endpoint:</u> AHI Mean nocturnal SaO2 LVEF <u>Eimitations:</u> Post hoc analysis Stratification of CPAP-treated pts based on polysomnogram performed 3 mo after randomization. Because suppressed and unsuppressed status could not be ascertained until completion of PSG, events that occurred during the first 3 mo could not be included The CPAP-CSA-suppressed group was younger, had a lower AHI, and had a slightly lower proportion of central events than the CPAP CSA-unsuppressed group
				LVEF	

				 LVEF significantly increased over time in CPAP-suppressed group (p<0.001) LVEF significantly increased in CPAP-suppressed vs. CPAP- unsuppressed (p=0.006) and vs. control (p<0.001) groups. No significant difference between CPAP-unsuppressed and control group (p=0.984) 	
CPAP for CSA & HF (CANPAP) Bradley et al. 2005 (101) 16282177	Aim: Test long-term treatment of CSA w/ CPAP in HF pts receiving optimal medical therapy on combined rates of death & ht tx. Study type: 11 center RCT Size: 258	Inclusion criteria: • 18-79 y • NYHA II-IV • HF due to ischemia • HTN, Idiopathic DCM • Stable condition • Optimal medical therapy for 1+ mon • LVEF <40% • CSA w/ ≥15 AHI >50% of AHI had to be central. Exclusion criteria: • Pregnancy • MI • UA • Cardiac surgery within prior 3 mon, OSA	Intervention: CPAP n=128 Comparator: No CPAP n=130	 <u>1° endpoint</u>: Transplant free survival No significant difference in transplant free survival between CPAP and control groups (p=0.54) <u>2° endpoints:</u> Hospitalizations: No significant difference between CPAP and control groups (p=0.45) EF: Significant increase in EF between CPAP vs. control groups (p=0.02) Frequency of apnea and hypopnea episodes Significant increase between CPAP vs. control groups (p=0.01) Mean Nocturnal SaO2 Significant increase between CPAP vs. control groups (p≤0.001) Mean Nocturnal SaO2 Significant increase in 6MWT between CPAP vs. control groups (p≤0.001) QoL: No significant difference between CPAP and control groups (p=0.74) 	 <u>2° endpoints:</u> Hospitalizations EF Frequency of apnea and hypopnea episodes Mean nocturnal SaO₂ 6MWT QoL Neurohormones – norepinephrine and atrial NP <u>Limitations:</u> Underpowered because trial stopped early for low enrollment

Ruttanaumpawan et al. 2009 (102) <u>19189783</u>	Aim: To determine whether attenuation of CSA by CPAP in pts w/ HF reduces the frequency of arousals from sleep or improves sleep structure. Study type: RCT Size: 205	Inclusion criteria: Age 18 - 79 y of age; NYHA II -IV HF due to ischemic, hypertensive, or idiopathic DCM, stabilized on optimal medical therapy ≥1 mo LVEF <40% by radionuclide angiography CSA defined as an AHI ≥15, w/ >50% central apneas & hypopneas Exclusion criteria: Pregnancy MI UA Cardiac surgery within 3 mo of enrollment OSA	Intervention: CPAP n=97 Comparator: Control n=108	 Neurohormones: Norepinephrine Significant reduction in CPAP vs. control groups (p=0.009) Atrial NP: No significant difference between CPAP and control groups <u>1° endpoint</u>: AHI (central and obstructive) Mean and lowest SaO₂ <u>Significant Results</u> In the CPAP group. Central and obstructive AHI decreased significantly <u>over BL</u> and vs. the control group (p<0.001) Mean and lowest SaO₂ improved in both the CPAP (p<0.001) and control (p<0.04) but the improvement was significantly better in the CPAP vs. the control group (p<0.001). <u>2° endpoints</u>: No significant improvement in arousals from sleep or sleep structure within or between groups (p=0.14–0.99) 	 <u>2° endpoints</u>: Arousals from sleep Sleep structure (time in bed, sleep period time, total sleep time, sleep efficiency, sleep onset latency, percentage in each sleep stage, periodic leg movement index) <u>Limitations:</u> 2° analysis of CANPAP data Did not classify arousals as being respiratory or non-respiratory related, and did not examine their timing.
Kaneko et al. 2003 (103) <u>12660387</u>	Aim: To determine the effect of CPAP on LVEF when awake and daytime BP in pts with HF and OSA <u>Study type</u> : RCT	 Inclusion criteria: HF due to ischemic or nonischemic dilated CM for >6 mo; LVEF <45% by radionuclide angiography NYHA class II–IV; Absence, in last 3 mo, of HF exacerbations while receiving optimal pharmacologic therapy at highest tolerated doses; 	Intervention: CPAP n=12 Comparator: Control n=12	1° endpoint: LVEF when awake LVEDD LVESD Heart rate Daytime BP Significant Results 1° endpoint: LVEE when awake	 <u>2° endpoint:</u> BMI Episodes of apnea and hypopnea Total Obstructive Central Desaturation index (# hr of sleep) Lowest oxyhemoglobin caturation (%)

	 OSA defined as ≥20 episodes of 	Significant increase in CPAP	Total sleep time
<u>Size</u> : 24	apnea and hypopnea /h of sleep of	(p<0.001) but not control group	Stage I and II sleep (% of
	which >50% were obstructive	and difference between groups	total sleep time)
	Evolution evitoria	was significant (p=0.009)	• Stage III and IV sleep (% of
	Exclusion criteria:		total sleep time)
	Presence of implanted cardiac	No significant difference for either	REM sleep (% of total sleep time)
	pacemaker:	group or between groups	Arousals/hr of sleep
	• UA;		
	• MI:	LVESD	Limitations:
	 Cardiac surgery within 3 mo of 	Significant reduction in CPAP	 No placebo
	enrollment	(p=0.009) but not control group and difference between groups	Small sample size
		was significant (p=0.02)	 Pts unblinded to group
		Heart Rate	
		Significant decrease in CPAP	
		(p=0.007) but not control group	
		was significant (p=0.02)	
		Daytime BP	
		Significant decrease in systolic BP	
		in CPAP (p=0.02) but not control	
		groups was significant (p=0.008)	
		No significant difference in	
		diastolic BP for either group or	
		between groups	
		<u>2° endpoint:</u>	
		BMI	
		No significant difference for either group or between groups	
		Episodes of apnea and hypopnea	
		Total	
		Significant reduction in CPAP	
		and difference between groups	

		was significant (p=0.002)	
		Obstructive	
		Significant reduction in CPAP	
		(p<0.001) but not control group	
		and difference between groups	
		was significant (p<0.001)	
		Central	
		No significant difference for CPAP	
		group or between groups	
		Desaturation index (# hr of sleep)	
		Significant reduction in CPAP	
		(p<0.001) but not control group	
		and difference between groups	
		was significant (p=0.008)	
		Lowest oxyhemoglobin	
		saturation (%)	
		 Significant increase in CPAP 	
		(p=0.004) but not control group	
		and difference between groups	
		was significant (p=0.01)	
		Total sleep time	
		No significant difference for CPAP	
		group or between groups	
		Stage I and II sleep (% of total	
		sleep time)	
		No significant difference for CPAP	
		group or between groups	
		Stage III and IV sleep sleep (% of	
		total sleep time)	
		No significant difference for CPAP	
		group or between groups	
		REM sleep (% of total sleep time)	

				No significant difference for CPAP	
				group or between groups	
				Arousals/h of sleep • Significant reduction in CPAP (p=0.003) but not control group and difference between groups was significant (p=0.03)	
Mansfield et al. 2004 (104) <u>14597482</u>	Aim: To assess long- term effect of OSA treatment with nocturnal CPAP on systolic heart function, sympathetic activity, BP, and QoL in pts with HF Study type: RCT Size: 44	 Inclusion criteria: HF due to ischemic or nonischemic dilated CM for >6 mo; LVEF <45% by radionuclide angiography NYHA class II–IV; Absence, in last 3 mo, of HF exacerbations while receiving optimal pharmacologic therapy at highest tolerated doses; OSA defined as ≥20 episodes of apnea and hypopnea /h of sleep of which >50% were obstructive Exclusion criteria: 1° valvular heart disease; Presence of implanted cardiac pacemaker; UA; MI: Cardiac surgery within 3 mo of enrollment 	Intervention: CPAP X 3 mo n=19 Comparator: Control n=21	 <u>1° endpoint</u>: LVEF Overnight urinary norepinephrine excretion BP QoL <u>Significant Results</u> <u>1° endpoint</u>: LVEF Significant improvement in CPAP group (p<0.001) and vs. control group (p=0.04) Overnight urinary norepinephrine excretion Significant reduction in CPAP group (p<0.05) and vs. control group (p=0.036) BP No significant difference in CPAP group or between groups QoL Significant improvements in most domains within CPAP group SF-36 Significant improvements between groups in 4/8 domains Physical (p=0.03) Vitality (p=0.02) Social (p=0.03) Mental health (p=0.01) 	 <u>2° endpoint:</u> Peak Vo₂ NYHA class Epworth sleepiness scale BMI AHI events per h Minimum SpO₂ saturation <u>Limitations:</u> No placebo Significant difference between groups in peak Vo₂ and mean BP at BL Dropout rate = 27% Higher than expected death rate Higher than expected rate of interventions initiated that may have effected end points Small sample size with only 3 females

	Chronic HF questionnaire	
	 Significant improvements between 	
	groups in 3/4 domains	
	\circ Fatigue (p=0.01)	
	\circ Emotional well-being (p=0.02)	
	$_{\odot}$ Disease mastery (p=0.02)	
	с, (р)	
	2° endpoint:	
	Peak Vo ₂	
	No significant difference in CPAP	
	aroun or between arouns	
	No significant difference CPAP	
	aroun or between arouns	
	Enworth cleanings scale	
	• Significant reduction in CPAP vs.	
	control group (p=0.01)	
	BMI	
	No significant difference CPAP	
	group or between groups	
	AHI events per n	
	Significant reduction in CPAP	
	group (p<0.001) and vs. control	
	group (p<0.001)	
	Minimum SpO ₂ saturation	
	 Significant improvement in CPAP 	
	group (p<0.001) and vs. control	
	aroup (p=0.001)	

Date: Study selected by the chairs in December 2015 and some trials added by the writing committee.

Study Name, Author, Year	Aim of Study	Study Type	Background Therapy	Study Size	Etiology	Patient	Population	I	Endpoints	Mortality	Trial Duration (Years)	Absolute Benefit	P Values & 95% CI:
			Pretrial standard treatment	N (Total) n (Experimental) n (Control)	Ischemic/ Nonlschemic	Inclusion Criteria	Exclusion Criteria	Primary Endpoint	Secondary Endpoint	1st Year Mortality			
CONSENSUS 1987 <u>2883575</u> (105)	To Evaluate influence of enalapril on prognosis of NYHA class IV HF	RCT	Diuretics (spironolactone 53%, mean dose 80mg), digitalis (93%), other vasodilators, except ACEI (ie, nitrates 46%)	253; 127;126	CAD 73%	Severe HF/symptoms at rest/NYHA class IV; Increased heart size >600 mL; BP: 120/75; HR: 80; AF 50%	APE; hemodynamically import aortic/MV stenosis; MI w/in prior 2 mo Unstable angina; planned cardiac surgery; right HF b/c of pulm disease; Cr >300 mmol/L	Mortality	Change in NYHA-FC, LV size, Cr level	52% placebo group and 36% enalapril group (6 mo mortality: 26% in enalpril group and 44% in placebo group)	0.51 y	N/A	Crude mortality at end of 6 mo (primary endpoint), 26% in enalapril group and 44% in placebo group—40% reduction (p =0.002). Mortality was reduced by 31% at 1 y (p=0.001)
10 y FU of CONSENSUS 1999 <u>10099910</u> (106)	Report on the survival at the 10-y follow up of the pts randomized in CONSENSUS. (1st study to show prognostic improvement by an ACEI. Pts in NYHA class IV HF treated with enalapril or placebo. After study completion all pts were offered open- label enalapril therapy).	10-y open- label follow- up study (via completion of a questionnaire) on the survival status of pts in CONSENSUS -a RCT.	All pts were offered open-label enalapril therapy	315; 77; 58		253 randomized pts included in analysis of time from randomization to death; Survivors (135) of the double-blind period included in analysis of the time from end of double-blind period to death; Severe, NYHA IV		Mortality			10 y		5 pts, all in the enalapril group, were long-term survivors (p=0.004). Averaged over the trial (double-blind plus open-label extension) risk reduction was 30% (p=0.008), 95% CI: 11% - 46%. At end of double-blind study period, mortality considerably higher among pts not receiving open ACEI therapy
SOLVD 1991 2057034 (107)	Study the effect of enalapril on mortality and hospitalization in pts with chronic HF and EF ≤35%	RCT	Diuretics + Digoxin	2569; 1285; 1284	Ischemic heart disease 72%	LVEF <35%; Mild to severe (11% class I/<2% class IV); LVEF 25%; BP: 125/77; HR: 80; AF: 8-12%	Age >80 y; Unstable angina; MI w/in past mo; Cr>2.0 mg/dL	Mortality	Hospitalizations; Incidence of MI; Mortality by specific causes; Combined mortality and morbidity from both SOLVD+/SOLVD-	15.70%	3.45 у	Treating 1000 SOLVD+ pts with enalapril for ~3 y would save ~50 premature deaths and 350 hospitalizations.	Reduced mortality by 16%; (95% Cl, 5-26%; p=0.0036)

2013 HF Guideline Data Supplement 18. ACE Inhibitors (Section 7.3.2.2)

SOLVD 1992 <u>1463530</u> (108)	Study effect of ACEIs on total mortality and mortality from CV causes, the development of HF, and hospitalization for HF in pts with EF <35%	RCT	No drug treatment for HF	4228; 2111; 2117	History of ischemic heart disease 85%	EF <35%; Asymptomatic; NYHA class I (67%) + II; EF: 28%; BP: 126/78; HR: 75; AF: 4%	As per SOLVD+	Mortality; Combined mortality and the incidence of HF and rate of hospitalization for HF	Incidence of HF and rate of hospitalization for HF	
SOLVD F/U 2003 <u>12788569</u> (109)	12-y FU of SOLVD to establish if the mortality reduction with enalapril among pts with HF was sustained, and whether a subsequent reduction in mortality would emerge among those with asymptomatic ventricular dysfunction.	12 y f/u of RCTs [SOLVD+ and SOLVD-]	N/A	6784; 3391; 3393	N/A	Participation in SOLVD+ and SOLVD- Asymptomatic to severe; NYHA I-IV	N/A	Mortality	N/A	N/A
ATLAS 1999 <u>10587334</u> (110)	To compare the efficacy and safety of low and high doses of ACEI on the risk of death and hospitalization in chronic HF. than the large doses that have been shown to reduce morbidity and mortality in pts with HF. AIM: Investigate if low doses and high doses of ACEIs have similar benefits.	RCT	N/A	3164; 1596 to the low- dose strategy and 1568 to the high- dose strategy.	CAD 65%	LVEF <=30%; NYHA class II, III, or IV, despite treatment with diuretics for ≥2 mo (Treatment for HF in ED or hospital within 6 mo required for pts in class II); Prior use of digitalis, ACEIs, or vasodilators allowed but not mandated; NYHA II-IV (mainly class II); LVEF 23%; SBP 126 mmHg; HR 80; NYHA class: III (few II and IV)	Acute coronary ischemic event or revascularization procedure within 2 mo; History of sustained or symptomatic ventricular tachycardia; Intolerant of ACEIs; SCr >2.5 mg/dL	Mortality from all causes	Combined risk of all- cause mortality and hospitalization for any reason; CV mortality, CV hospitalizations; All-cause mortality combined with CV hospitalizations; CV mortality combined with CV hospitalizations; Combined risk of fatal and nonfatal MI plus hospitalization for unstable angina	

3.12 у		Reduced mortality: p=0.30; 95% CI: -8-21%
N/A	Enalapril extended median survival by 9.4 mo in the combined trials (95% CI: 2.8–16.5, p=0.004).	In the prevention trial, 50.9% of the enalapril group had died c/w 56.4% of the placebo group (p=0.001). In the treatment trial, 79.8% of the enalapril group had died c/w 80.8% of the placebo group (p=0.01). Combined prevention and treatment trials: HR for death was 0.90 for the enalapril group c/w placebo group (95% CI: 0.84–0.95, p=0.0003).
5 y		High-dose group had 8% lower risk of all-cause mortality (p=0.128) and 10% lower risk of CV mortality (p=0.073) than low-dose group. Death or hospitalization for any reason, high-dose group had 12% lower risk than low-dose group, p=0.002. Total number of hospitalizations: high-dose group 13% fewer hospitalizations for any reason (p=0.021), 16% fewer hospitalizations for CV reason (p=0.05), and 24% fewer hospitalizations for HF (p=0.002).

SAVE, 1992 <u>1386652</u> (111)	To test the hypothesis that the long-term administration of captopril to survivors of acute MI who had baseline LV dysfunction but did not have overt HF requiring vasodilator therapy would reduce mortality, lessen deterioration in cardiac performance, and improve clinical outcome.	RCT	Beta-blockers 36%; Digitalis 26%; Nitrates 51%	2231; 1115; 1116	Ischemic 100%	Alive 3 d after MI; LVEF <40%; >21 y of age, but <80; Killip class I — 60% (60% of the ps did not have even transient pulmonary congestion at baseline/the time of their acute MI; EF 31%; BP 113/70; HR 78;	Failure to undergo randomization within 16 d after the MI; Relative contraindication to the use of an ACEIs or the need for such an agent; SCr > 2.5 mg/dI	Mortality from all causes	Mortality from CV causes; Mortality combined with a decrease in the EF of at least 9 units in surviving pts; CV morbidity (development of severe CHF or the recurrence of MI); Combination of CV mortality and morbidity; 2 endpoints of severe HF (treatment failure): 1st, development of overt HF necessitating treatment with ACEI and 2nd, hospitalization to treat CHD.	
AIRE 1993 <u>8104270</u> (112)	Investigated the effect of therapy with ACEI ramipril, on survival in pts who had shown clinical evidence of HF at any time after an acute MI. Also, to compare the incidences of progression to severe or resistant HF, nonfatal reinfarction and stroke between the 2 groups.	RCT		2006; 1014; 992		Aged ≥18 y, with a definite acute MI 3- 10 d before randomization; Clinical evidence of HF at any time since acute MI	Use of an ACEI considered to be mandatory	Mortality from all causes		

3.5 y	Mortality from all causes was significantly reduced in the captopril group (228 deaths, or 20%) as c/w the placebo group (275 deaths, or 25%); the RR: 19% (95% CI, 3-32%; p=0.019). RR:21% (95% CI, 5 -35%; p=0.014) for death from CV causes, 37% (95% CI, 20-50%; p<0.001) for the development of severe HF, 22% (95% CI, 4-37%; p=0.019) for CHF requiring hospitalization, and 25% (95% CI, 5-40%; p=0.015) for recurrent MI.
1.3 y	Mortality from all causes was significantly lower for pts on ramipril compared to pts on placebo. RR: 27%; 95% CI: 11- 40%; p=0.002. Prespecified secondary outcomes: risk reduction of 19% for the 1st validated outcome—namely, death, severe/resistant HF, MI, or stroke (95% CI: 5% - 31%; p=0.008).

TRACE 1995	To determine	RCT	Beta blocker 16%:	1749; 876; 873	Ischemic	Consecutive pts	Contraindication to	Death from	Death from a CV	The mortality from all	24 lives were saved	During the study period, 304 pts in
<u>7477219</u> (113)	whether pts who LV dysfunction soon after MI benefit from long-term oral ACE inhibition.	KU I	Calcium antagonist 28%; Diuretic 66%; Nitrates 53%; Digoxin 28%.	1749, 676, 873	100%	 >18 y hospitalized >18 y hospitalized with MI; Criteria for MI: chest pain or electrocardiographi c changes, accompanied by >2X increase in ≥1 cardiac enzymes; LV dysfunction (EF <35%); NYHA class 1 - 41%; BP 121/76; HR 81 	ACEI or a definite need for them; Severe, uncontrolled DM; Hyponatremia (<125 mmol/L); Elevated SCr level (2.3 mg/dL)	any cause	cause, sudden death; Progression to severe HF (hospital admission for HF, death due to progressive HF, or HF necessitating open- label ACEI); Recurrent infarction (fatal or nonfatal); Change in the wall- motion index (EF)	causes at 1 y was 24%.	after 1 mo of treating 1,000 pts	burning the study period, 304 pts in the trandolapril group died (34.7%), as did 369 in the placebo group (42.3%). RR: 0.78 (95% CI, 0.67 - 0.91; p=0.001). In every subgroup, treatment with trandolapril was associated with a reduction in risk.

ACEI indicates angiotensin-converting-enzyme inhibitor; AF, atrial fibrillation; AIRE, Acute Infarction Ramipril Efficacy; APE, acute pulmonary embolism; ATLAS, Assessment of Treatment with Lisinopril and Survival; BP, blood pressure; CAD, coronary artery disease; CHD, chronic heart disease; CHD, chronic heart disease; CHD, chronic heart disease; CHD, chronic heart disease; CHF, congestive heart failure; CONSENSUS Cooperative North Scandinavian Enalapril Survival Study; Cr, creatinine; CV, cardiovascular; C/W, compared with; DM, diabetes mellitus; ED, emergency department; FU, follow-up; HF, heart failure.

Study Name, Author, Year	Aim of Study	Study Type	Background Therapy	Study Size	Etiology	Patient Population		Severity	E	Endpoints	Mortality	Trial Duration (Y)	Statistical Results
CHARM	Discover	RCT	Pre-trial standard treatment. Diuretics,	N (Total) n (Experimental) n (Control) 2028; 1013;	Ischemic/ Non-Ischemic Ischemic 67-	Inclusion Criteria Symptomatic HF, EF	Exclusion Criteria	NYHA II-IV; mild to	Primary Endpoint Composite of CV	Secondary Endpoint CV death, hospital	1st Y Mortality	2.8 у	Absolute reduction of 7 major events per 100
Alternativ e; Granger et al; (2003) <u>13678870</u> (114)	whether ARB could improve outcome in pts not taking an ACEI (intolerant)		Beta-blockers (55%), spironolacton e 24%, Digoxin 45- 46%	1015	70%	<40%, no ACEI (b/c of intolerance)		severe (<4% class IV); EF: 30%; BP: 130/70; HR: 74-75; AF: 25-26%	death or hospital admission for CHF	admission for CHF or nonfatal MI; CV death, CHF admission, nonfatal MI, nonfatal stroke; CV death, CHF admission, nonfatal MI, nonfatal stroke, coronary revascularization; Death (any cause); New DM			pts threated - NNT 14 pts to prevent 1 CV death or hospitalization. HR: 0.77 (95% CI: 0.67-0.89); p=0.0004
CHARM- ADDED; McMurray et al; (2003) <u>13678869</u> (115)	To investigate if ARB + ACEI in pts with chronic HF improve clincal outcomes	RCT	Beta blocker- 55%; spironolacton e 17%; Digoxin 58- 59%	2548; 1276; 1272	Ischemic 62- 63%	Symptomatic HF; EF <40%; Treatment with ACEI; Age >18 y		NYHA class II-IV; mild to severe (<3% class IV); EF 28%; BP 125/75; HR 74; AF 27%	Composite of CV death or hospital admission for CHF	CV death, hospital admission for CHF or nonfatal MI; CV death, CHF admission, nonfatal MI, nonfatal stroke; CV death, CHF admission, nonfatal MI, nonfatal stroke, coronary revascularization; Death (any cause); New DM		3.4 y	Absolute reduction of 4.4 pts with events per 100 pts treated- NNT of 23 to prevent 1 first event of CV death or CHF hospitalization. RR: 0.85 (95% CI: 0.75-0.96); p=0.011

2013 HF Guideline Data Supplement 19. ARBs (Section 7.3.2.3)

VALIANT; Pfeffer et al; (2003) <u>14610160</u> (116)	Compare the effect of an ARB, ACEI and the combination of the 2on mortality	Randomize d double blind multicenter trial	Beta- blockers; ASA	14,703 Valsartan:490 9 Captopril-: 4909 VAL + CAP: 4885	Ischemic 100% (MI inclusion criteria)	Age >18 y; Acute MI complicated by HF; LV systolic dysfunct (EF <35%), (<40% on radionuclide ventriculography); SBP >100 mmHg; Cr <2.5 mg/dL	Prior intolerance or contra- indication to ACEI/ ARB	NYHA I-IV; asymptomatic- severe, EF 35%; BP: 123/72; HR: 76	Death from any cause		12.5% VAL 12.3% VALCAP 13.2% CAP	2.1 y	VAL and CAP: 1.0 (97.5% CI 0.90-1.11); p=0.98 ; VAL+CAP and CAP: 0.98 (97.5% CI 0.89- 1.09); p=0.73
Val-HeFT; Cohn et al; (2001) <u>11759645</u> (117)	Evaluate long term effects of adding ARB to standard therapy for HF	RCT	Diuretics; Digoxin 67%; Beta blocker 35%; ACEI 93%	5010; 2511; 2499	Ischemic 57%	Age >18 y; NYHA II, II, IV; At least 2 wk of background meds including ACEIs; EF <40% and LVID >2.9 cm/BSA		NYHA II-III, IV (only ~2% class IV); Mild to severe; EF 27%; BP 123/76; AF 12%	Mortality; Combined endpoint of mortality and morbidity	Change in EF; • NYHA class, QoL scores; Signs and symptoms of HF		1.92 y	Mortality similar for the 2 treatment groups. For the combined endpoint: RR: 0.87; 97.5% CI, 0.77-0.97; p=0.009
HEAAL study; Lancet 2009; 374: 1840-48. <u>19922995</u> (118)	Compared the effects of high-dose vs low-dose losartan on clinical outcomes in pts with HF.	RCT	Diuretic drugs (77%), beta blockers (72%), and ARBs (38%).	3846 losartan 150 mg (n=1927) or 50 mg daily (n=1919).	IHD 64%	>18 y; NYHA class II–IV; LVEF <40%, with stable CV medical therapy for at least 2 wk; Intolerance to ACEI; Investigators encouraged to start beta blocker and titrate to a maximum, whenever possible	Pregnancy or lactation; known intolerance to ARBs; Systolic arterial blood pressure <90 mm Hg; Significant stenotic valvular heart disease; Active myocarditis; active pericarditis; Planned heart transplantation w/in 6 mo; coronary angioplasty, CABG, acute MI, UA pectoris, cerebrovascular accident, or TIA within the previous 12 wk; Suspected significant renal artery stenosis	NYHA II-IV (70% II); EF: 33%; BP: 124/77; HR: 71; AF; 28%	Death or admission for HF	Composite endpoint of death or CV admission. Additional prespecified outcomes included: death, death or all-cause admission, CV death, all- cause admission, CV admission, admission for HF, and changes in the severity of heart disease		4.7 y median f/u	Treating pts with 150 mg dose instead of 50 mg dose would result in 1 additional pt w/out the primary event at 4 y for every 31 pts treated. Composite: 828 (43%) pts in 150 mg group vs. 889 (46%) in 50 mg group died or admitted for HF (HR: 0.90; 95% CI: 0.82-0.99 p=0.027) • Components: 635 pts in 150 mg group vs. 665 in 50 mg group died (HR: 0.94, 95% CI: 0.84-1.04; p=0.24), and 450 vs. 503 pts admitted for HF (0.87, 0.76–0.98; p=0.025)
CHARM- Overall <u>13678868</u> (116)	Aimed to find out whether the use of an ARB could reduce mortality and morbidity.	RCT- parallel, randomized , double- blind,	Diuretics 83% Beta blockers 55% ACEI 43% Spironolacton e 17% Digoxin 43%	7601 pts (7599 with data) 3803 3796		>18 y; NYHA class II–IV for at least 4 wk; 3 distinct populations: pts with LVEF <40% who were not receiving ACEIs (previous intolerance) or who were currently receiving ACE, and pts with LVEF >40%	SCr > 265 mcmol /L, serum potassium >5.5 mmol/L Bilateral renal artery stenosis; symptomatic hypotension Women of childbearing potential not using adequate contraception; Critical aortic or mitral stenosis; MI, stroke, or open-heart surgery in the previous 4 wk; Use of an ARB in the previous 2 wk	NYHA II-IV NYHA II-IV Only 3% class IV	The primary outcome of the overall program: all-cause mortality; For all the component trials: CV death or hospital admission for CHF.		The annual CV death rate among the placebo group who had reduced LVEF was around 9% and was only 4% in the placebo group of CHARM- Preserved.	3.1 y	886 (23%) pts in candesartan and 945 (25%) in placebo group died (unadjusted HR: 0.91; 95% Cl: 0.83–1.00; p=0.055; covariate aHR: 0.90 95% CU: 0.82–0.99; p=0.032) • Fewer CV deaths (691 [18%] vs 769 [20%], unadjusted HR: 0.88; 95% Cl: 0.79–0.97; p=0.012; covariate aHR: 0.87; 95% Cl: 0.78– 0.96; p=0.006) • Hospital admissions for CHF (757 [20%] vs 918 [24%], p<0.0001)

ACEI indicates angiotensin-converting-enzyme inhibitor; AF, atrial fibrillation; ARB, angiotensin receptor blockers; ASA, aspirin; BP, blood pressure; BSA, body surface area; CABG, coronary artery bypass graft; CHARM, Candesartan in Heart Failure: Assessment of Reduction in Mortality and Morbidity; CHD, chronic heart disease; CHF, congestive heart failure; Cr, creatinine; CV, cardiovascular; DM, diabetes mellitus; EF, ejection fraction; FU, follow-up; HEAAL study, effects of high-dose versus low-dose losartan on clinical outcomes in patients with heart failure; HF, heart failure; HR, heart rate; IHD, ischemic heart disease; LV, left ventricular; LVD, left ventricular dilatation; MI, myocardial infarction; MV, mitral valve; N/A, not applicable; NNT, number needed to treat; NYHA, New York Heart Association; QoL, quality of life; pts, patients; SBP, systolic blood pressure; RCT, randomized control trial; SCr, serum creatinine; TIA, transient ischemic attack; UA, unstable angina; Val-HeFT, Valsartan Heart Failure Trial; and VALIANT, Valsartan in Acute Myocardial Infarction.

2013 HF Guideline Data Supplement 20. Beta Blockers (Section 7.3.2.4)

Study Name, Author, Year	Aim of Study	Study Type	Background Therapy	Study Size	Etiology	Patie	ent Population	Severity		Endpoints		
				N (Total)								
				n (Experimental)		Inclusion			Primary			
				n (Control)		Criteria	Exclusion Criteria		Endpoint	Secondary Endpoin		

	Mortali	ity	Trial Duration	Statistical Results
nt	Annualized Mortality	1st Y Mortality		

CIBIS II CIBIS Il investigators and committee members (1999) <u>10023943</u> (119)	Investigate the efficacy of bisoprolol in decreasing all- cause mortality in chronic HF	RCT- multicenter double-blind randiomised placebo controlled trial (Europe)	Diuretics + ACEI; [amiodarone allowed14- I6%]	2647; 1327; 1320	Documented Ischemic 50%	NYHA class III or IV EF: <35% 18-80 y old	Uncontrolled HTN; MI/UA w/in previous 3 mo; PTCA/CABG w/in previous 6 mo; AV-block >1st degree w/o PPM; Heart rate < 60bpm; resting SBP <100mmHg; renal failure; Reversible obstruct lung disease; Use of beta blocker	Moderate to severe. Mean BP: 130/80; Mean HR: 80; Mean EF: 28%; Mean LVEDD: 6.7 cm; AF: 20%	All-cause mortality	All-cause hospital admissions All CV deaths Combined endpoints Permanent treatment withdrawal	13.2% Placebo group 8.8% Treatm't group	N/A	1.3 у	HR: 0.66 (95% CI: 0.54-0.81); p<0.0001
MERIT-HF; MERIT study Group; (1999) <u>10376614</u> (120)	Investigate whether Metoprolol CR/XL lowered mortality in pts with decreased EF and symptoms of HF	RCT multicenter double-blind randiomised placebo controlled trial (Europe + USA)	Diuretics + ACEI [Amiodarone NOT allowed]	3991; 1991; 2001	Ischemic 65%	NYHA II-IV; 40-80 y old; LVEF <40% (36- 40 if 6-min walk <450m); heart rate >68 bpm	MI/UA w/in 28 d; Contra-indication or current use of beta blocker; PTCA/CABG w/in 4 mo Planned transplant or ICD; Heart block >1 st degree w/o PPM; SBP <100mmHg	Mild to severe. Mean BP: 130/78; Mean HR: 78; Mean EF 28%; AF 16-17%	All-cause mortality All-cause mortality in combination with all-cause admission to hospital	N/A	11.0% Placebo group 7.2% Treatm't group	N/A	1 y	Treatment of 27 pt for 1 y can prevent 1 death. 0.66 (95% CI: 0.53- 0.81); p=0.00009
COPERNICUS ; Packer et al; (2002) <u>12390947</u> (121)	Investigate whether Carvadiolo is beneficial in severe HF	RCTdouble blind	Diuretics (PO or IV) + ACEI (or ARB); [Amiodarone allowed 17- 18%]	2289; 1156; 1133	Ischemic 67%	Euvolumic NYHA class IV; LVEF <25%; No positive inotropes or vasodilators w/in 4 d	Pt requiring hospitalized intensive care; Use of positive inotropes or IV; vasodilators w/in 4- d; Coronary revascularization/MI/CVA/ sign VT or VF w/in 2 mo; SBP < 85 mmHg, Heart rate <68, Cr >2.8 mg/dL	Severe Mean BP: 123/76; Mean HR: 83; Mean EF 20%;	All-cause mortality	Combined risk of death or hospitalization-any reason; Combined risk of death or hospitalizationCV reason; Combined risk of death or hospitalizationHF reason; Pt global assessment	19.7% placebo [24.0% in pts with recent or recurrent cardiac decompensations]	18.5% in placebo group 11.4% in Carvedilol group	10.4 mo	Treating 1000 pt for 1 y led to savings of 70 premature deaths p=0.0014
SENIORS; Flather et al; (2005) <u>15642700</u> (122)	Assess effects of the beta blocker Nebivolol in pts_70 y regardless of EF.	RCT	Diuretics + ACEI (+aldosterone antagonist in 29%)	2128; 1067; 1061	Prior h/o CAD in 69%	Age >70 CHF with 1 of the following: hospitalization with CHF w/in a year or EF <35% w/in the past 6 mo	New HF therapy w/in 6 wk or change in drug therapy w/in 2 wk Contraindication to beta blockers, current use of beta blockers Significant renal dysfunction CVA w/in 3 mo.	Mild to severe Mean BP: 139/81; Mean HR: 79; Mean EF 36% (1/3 with EF >35%);	Composite of all-cause mortality or CV hospital admission	All-cause mortality Composite of all-cause mortality or all-cause hospital admissions All cause hospital admissions CV hospital admissions CV mortality Composite of CV mortality or CV hospital admissions NYHA class assessment; 6 MWT	N/A	N/A	1.75 y	Absolute risk reductio 4.2%; 24 pts would need to be treated for 21 mo to avoid one event RR: 0.86; 95% CI: 0.74-0.99; p=0.039
A Trial of the Beta-Blocker Bucindolol in Pt with Advanced Chronic HF The Beta- Blocker Evaluation of Survival Trial Investigators <u>11386264</u> (123)	Designed to determine whether bucindolol hydrochloride, a nonselective beta- adrenergic blocker and mild vasodilator, would reduce the rate of death from any cause among pt with advanced HF	RCT	ACEIs (if tolerated) [91% ACE; 7% ARB], for at least 1 mo. Before the publication of the results of the DIG trial, 12 digoxin therapies were	2708; 1354; 1354	Ischemic 59%	NYHA class III or IV HF LVEF <35% >18 y	Reversible cause of HF present Candidates for heart transplantation Cardiac revascularization procedure within the previous 60 d UA Heart rate <50 bpm, SBP <80mmHg Decompensated HF.	NYHA III or IV (92% class III) EF 23%; HR 82; BP 117/71; AF 12%	Death from any cause	Death from CV causes (death due to pump failure or an ischemic event or sudden death) Hospitalization for any reason Hospitalization because of HF Composite of death or heart transplantation LVEF at 3 and 12 mo MI; QoL; and any change in	For pt in NYHA functional class III, the annual mortality rate was 16% in the placebo group; For pt with NYHA class IV, the annual mortality rate in the placebo group was 28% Overall : annual mortality of 17% in placebo group c/w	N/A	~2 y	449 pt in placebo group (33%) died, 411 in the bucindolol grou (30%; HR: 0.90; 95% CI, 0.78-1.02; unadjusted p=0.10; adjusted p=0.13)

	and to assess its effect in various subgroups defined by ethnic background and demographic criteria — specifically women and members of minority groups.		required, but thereafter its use became discretionary [DIG 94%].							the need for concomitant therapy	15% in the bucindolol group.			
COMET; Poole-Wilson et al; (2003) <u>12853193</u> (124)	To compare the effects of carvedilol and metoprolol on clinical outcome in pts with HF	RCT	Diuretics, ACEIs	3029; 1511 carvedilol; 1518 metoprolol tartrate	N/A	NYHA class II-IV EF <35% Previous CV admission	N/A	Mild to severe	All-cause mortality Composite endpoint of all- cause mortality, or all-cause admission	N/A	N/A	N/A	4.8 y	All-cause mortality 34% carvedilol and 40% metoprolol (HR: 0.83; 95% Cl 0.74- 0.93; p=0.0017)
(CIBIS) III; 2005 <u>16143696</u> (125)	Sufficient data do not currently exist to establish the optimum order of initiating chronic HF therapy (ACEI vs. beta blocker). This was the objective of the CIBIS III trial it compared the effect on mortality and hospitalization of initial monotherapy with either bisoprolol or enalapril for 6 mo, followed by their combination for 6 to 24 mo.	Multicenter, prospective, randomized, open-label, blinded endpoint evaluation (PROBE) trial,24 with 2 parallel groups.	Diuretics 84%; Digoxin 32%	1010 Bisoprolol 505; Enalapril 505	CAD 62%	>65 y, NYHA class II or III, and LVEF <35% (By echo within the 3 mo) Clinically stable HF (without clinically relevant fluid retention or diuretic adjustment within 7 d)	Treatment with an ACEI, an ARB, or a beta blocker for >7 d during the 3 mo before randomization Heart rate at rest <60 bpm without a functioning pacemaker Supine SBP <100 mm Hg at rest SCr≥220 mmol/L AV block>1° without a functioning pacemaker Obstructive lung disease contraindicating bisoprolol treatment	NYHA II or III; mild to moderate CHF LVEF 29%; Heart rate 79; SBP 134	The primary endpoint was time-to-the-first- event of combined all- cause mortality or all-cause hospitalization	Combined endpoint at the end of the monotherapy phase and the individual components of the primary endpoint, at study end and at the end of the monotherapy phase. CV death CV hospitalization	N/A	N/A	Mean of 1.22±0.42 y (maximum of 2.10 y).	In the ITT sample, 178 pt (35.2%) with a primary endpoint in th bisoprolol-1 st group, and 186 (36.8%) in the enalapril-1 st group (absolute difference - 1.6%; 95% CI: -7.6 to 4.4%; HR: 0.94; 95% CI: 0.77–1.16; noninferiority for bisoprolol-first versus enalapril-1 st treatment p=0.019)

ACEI indicates angiotensin-converting-enzyme inhibitor; AF, atrial fibrillation; ARB, angiotensin receptor blocker; AV, atrioventricular; BP, blood pressure; CABG, coronary artery bypass graft; CHF, congestive heart failure; CIBIS II, Cardiac Insufficiency Bisoprolol Study II; COMET, Carvedilol Or Metoprolol European Trial; COPERNICUS, carvedilol prospective randomized cumulative survival; Cr, creatinine; CR/XL, controlled release; CV, cardiovascular; CVA, cerebrovascular accident; c/w, compared with; DIG, Digitalis Investigation Group; EF, ejection fraction; HF, heart failure; h/o, history of; HR, hazard ratio; ICD, ICD, implantable cardioverter defibrillator; ITT, intent to treat; MERIT-HF, Metoprolol CR/XL Randomised Intervention Trial in Congestive Heart Failure; MI, myocardial infarction; MWT, minute walk test; NYHA, New York Heart Association; PPM, permanent pacemaker; PTCA, percutaneous transluminal coronary angioplasty; Pts, patients; QoL, quality of life; RCT, randomized control trial; RR, relative risk; SBP, systolic blood pressure; SCr, serum creatinine; UA, unstable angina; USA, United States of America; VF, ventricular fibrillation; VT, ventricular tachycardia; and w/o, without.

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